Differential Gene Exprpression and Pathway Analysis

2024-03-25

BACKGROUND

ACESO Genomics and TIDREC collaboration on the SCOPE project. The example code below is meant to illustrate the process of standard gene expression and pathway analysis to the enable RNA-Seq analytical capabilities going forward.

CONTENT:

- PART1 Loading data, preparation and running of DESeq2 analysis
- PART2 Plotting data and Gene Set Enrichment Analysis
- PART3 Exploring results plots
- PART4 Statistics and Table ONE

PART1

 $\label{eq:Goal: Goal: Goal:$

1. Setup your environment

```
## Clean
rm(list = ls())
gc()
##
            used (Mb) gc trigger (Mb) limit (Mb) max used (Mb)
## Ncells 478681 25.6
                         1032958 55.2
                                              NA
                                                    669402 35.8
## Vcells 905192 7.0
                         8388608 64.0
                                          256000 1851706 14.2
##
## PACKAGES
##
## load basic packages
suppressPackageStartupMessages(suppressWarnings({
  library(data.table); library(parallel); library(tidyr); library(tidyverse)}))
## For plotting
suppressPackageStartupMessages(suppressWarnings({
  library(ggpubr);library(ggbeeswarm);library(RColorBrewer);library(ggdendro);
  library(ggridges);library(ggrepel)}))
```

```
## For clustering
suppressPackageStartupMessages(suppressWarnings({library(pheatmap)}))
## For DESeg analysis
suppressPackageStartupMessages(suppressWarnings({library(DESeq2)}))
#library(sva) <- could be used for batch normalization
## For GSEA
suppressPackageStartupMessages(suppressWarnings({
 library(clusterProfiler);library(msigdb);library(msigdbr);
  library(enrichplot); library(ggupset)}))
##
## DIRECTORIES
##
TAB.DIR <- "/Users/a_PGenzor/Documents/GITHUB/ahjf_scope/results/tables/"
FIG.DIR <- "/Users/a_PGenzor/Documents/GITHUB/ahjf_scope/results/figures/"
#SES.DIR <- "/Users/a_PGenzor/Documents/GITHUB/ahjf_scope/sessions/"
##
## VERSION AND CONTROLS
##
aSeed="1003"
set.seed(aSeed)
version.date = "10MAR24"
```

2. Load the data

##

1:

- Loading count data should be relatively simple since it should all be contained in the single matrix.
- Make sure that the column names in count matrix match the names in your metadata tables or that there is a way to calcualte them

```
##
## gene count table
count_table_path <- "/Users/a_PGenzor/Documents/GITHUB/ahjf_scope/data/from_kimkee/10MAR24/RNASeq_COVID</pre>
cnt.dt <- fread(count_table_path)</pre>
cnt.dt[1:5,1:3]
##
                                  gene_id
            ENSG00000223764.2|LINC02593
## 1:
## 2: ENSG00000272438.1|ENSG00000272438
## 3: ENSG00000230699.2|ENSG00000230699
## 4: ENSG00000241180.1|ENSG00000241180
## 5: ENSG00000288531.1|ENSG00000288531
##
      02_D0_DKDL230010052-1A_HNG3NDSX7_L1.nonovel.gtf
## 1:
                                                      10
                                                       0
## 2:
## 3:
                                                       0
## 4:
                                                       0
## 5:
                                                       9
```

06_D0_DKDL230011035-1A_HC2HKDSX7_L3.nonovel.gtf

```
## 2:
                                                        0
## 3:
                                                        9
## 4:
                                                        0
## 5:
                                                        0
## splitting complex names into pieces
colnames(cnt.dt)[1:5]
## [1] "gene id"
  [2] "02_D0_DKDL230010052-1A_HNG3NDSX7_L1.nonovel.gtf"
  [3] "06_D0_DKDL230011035-1A_HC2HKDSX7_L3.nonovel.gtf"
  [4] "100_D0_DKDL230011026-1A_HC2HKDSX7_L3.nonovel.gtf"
   [5] "11_D0_DKDL230011071-1A_HC2HKDSX7_L4.nonovel.gtf"
unlist(tstrsplit(colnames(cnt.dt),split="_",keep = 1))
##
     [1] "gene" "02"
                         "06"
                                "100"
                                        "11"
                                                "14"
                                                       "21"
                                                               "26"
                                                                       "31"
                                                                              "32"
    [11] "34"
                                        "39"
##
                 "35"
                         "36"
                                "37"
                                                "03"
                                                       "43"
                                                               "45"
                                                                       "46"
                                                                              "47"
    [21] "49"
                 "51"
                         "53"
                                "54"
                                        "56"
                                                "57"
                                                       "58"
                                                               "60"
                                                                              "62"
##
                                                                       "61"
##
    [31] "67"
                 "68"
                         "69"
                                "70"
                                        "72"
                                                "74"
                                                       "75"
                                                               "77"
                                                                       "78"
                                                                              "79"
                                                "87"
##
    [41] "80"
                 "81"
                         "82"
                                "85"
                                        "86"
                                                       "88"
                                                               "89"
                                                                       "90"
                                                                              "91"
    [51] "92"
                 "93"
                         "94"
                                "95"
                                        "98"
                                                "02"
                                                       "03"
                                                               "06"
                                                                       "11"
                                                                              "14"
##
    [61]
         "21"
                 "32"
                         "36"
                                "37"
                                        "39"
                                                "43"
                                                       "45"
                                                               "47"
                                                                       "49"
                                                                              "51"
##
##
                                                "62"
                                                                       "77"
                                                                              "78"
    [71] "54"
                 "56"
                         "57"
                                "58"
                                        "60"
                                                       "67"
                                                               "75"
    [81] "79"
                 "80"
                         "81"
                                "82"
                                        "85"
                                                "86"
                                                       "87"
                                                               "88"
                                                                       "89"
                                                                              "90"
    [91] "91"
                 "92"
                         "93"
                                "94"
                                        "98"
                                                "100"
                                                       "32"
                                                               "49"
                                                                              "92"
                                                                       "62"
##
   [101] "93"
                 "100"
                         "32"
                                "49"
                                        "62"
                                                "92"
                                                       "93"
unlist(tstrsplit(colnames(cnt.dt),split="_",keep = 2))
                                   "DO"
                                          "DO"
                                                 "DO"
                                                       "D0"
                                                                    "DO"
                                                                           "DO"
                                                                                  "DO"
##
     [1] "id"
                "DO"
                       "DO"
                             "DO"
                                                              "DO"
    [13] "D0"
                "DO"
                       "DO"
                             "DO"
                                    "DO"
                                          "DO"
                                                 "DO"
                                                       "DO"
                                                              "DO"
                                                                     "DO"
                                                                           "DO"
                                                                                  "DO"
##
                                          "DO"
##
    [25] "D0"
                "DO"
                       "DO"
                             "DO"
                                    "DO"
                                                 "DO"
                                                       "DO"
                                                              "DO"
                                                                    "DO"
                                                                           "DO"
                                                                                  "DO"
         "D0"
                "DO"
                       "DO"
                             "D0"
                                    "DO"
                                          "D0"
                                                 "DO"
                                                       "D0"
                                                              "DO"
                                                                    "DO"
##
    [37]
                                                                           "DO"
                                                                                  "DO"
##
    [49] "D0"
                "D0"
                       "DO"
                             "DO"
                                    "DO"
                                          "DO"
                                                 "DO"
                                                       "D28" "D28" "D28" "D28" "D28"
    [61] "D28" "D28" "D28" "D28" "D28" "D28" "D28" "D28"
                                                              "D28"
                                                                    "D28" "D28" "D28"
##
         "D28" "D28" "D28" "D28" "D28" "D28" "D28"
                                                       "D28" "D28" "D28" "D28" "D28"
##
    [73]
    [85] "D28" "D3"
##
    [97] "D3"
                "D3"
                       "D3"
                             "D3"
                                   "D3"
                                          "D7"
                                                "D7"
                                                       "D7"
                                                              "D7"
                                                                    "D7"
                                                                           "D7"
## make new colnames
cnt_new_col_names <- paste(unlist(tstrsplit(colnames(cnt.dt),split="_",keep = 2)),</pre>
      unlist(tstrsplit(colnames(cnt.dt),split="_",keep = 1)),
      sep = "_")
cnt_new_col_names <-gsub("id_gene", "gene_id", cnt_new_col_names)</pre>
colnames(cnt.dt) <- cnt_new_col_names</pre>
cnt.dt[1:5,1:5] ## <- ready to use</pre>
##
                                  gene_id D0_02 D0_06 D0_100 D0_11
## 1:
             ENSG00000223764.2|LINC02593
                                               10
                                                      7
                                                              7
## 2: ENSG00000272438.1|ENSG00000272438
                                                      0
                                                              0
                                                                    0
## 3: ENSG00000230699.2|ENSG00000230699
                                                0
                                                      9
                                                             17
                                                                    7
## 4: ENSG00000241180.1|ENSG00000241180
                                                      0
                                                              0
                                                                    0
## 5: ENSG00000288531.1|ENSG00000288531
                                                             21
                                                                   30
##
## metadata
metadata_path <- "/Users/a_PGenzor/Documents/GITHUB/ahjf_scope/data/from_kimkee/10MAR24/RNASeq_COVID/Me
```

```
meta.dt <- fread(metadata_path)</pre>
meta_cols_to_use <- c("IGU_Code", "sex", "pathogen", "disease", "time")</pre>
meta.clean.dt <- meta.dt[,.SD,.SDcols = meta_cols_to_use]</pre>
meta.clean.dt[,"subject":=tstrsplit(IGU_Code,split="_",keep = 1)]
meta.clean.dt[,"seq_id":= paste(time,subject,sep = "_")]
meta.clean.dt
##
        IGU Code
                            pathogen disease time subject seq_id
                     sex
##
     1:
           02 DO Female SARS-CoV-2 COVID19
                                                DO
                                                             D0 02
##
                                                         02 D28 02
     2:
          02 D28 Female SARS-CoV-2 COVID19
                                               D28
##
     3:
           03 DO Female SARS-CoV-2 COVID19
                                                D0
                                                         03
                                                             D0 03
##
     4:
           03 D28 Female SARS-CoV-2 COVID19
                                               D28
                                                         03 D28 03
##
     5:
           06_D0 Female SARS-CoV-2 COVID19
                                                DO
                                                         06
                                                             D0_06
##
## 102:
           98_D0
                    Male SARS-CoV-2 COVID19
                                                DO
                                                         98
                                                             D0_98
           98 D28
                                                         98 D28 98
## 103:
                    Male SARS-CoV-2 COVID19
                                               D28
  104:
           100_D0
                    Male SARS-CoV-2 COVID19
                                                D0
                                                        100 DO_100
##
##
   105:
           100_D3
                    Male SARS-CoV-2 COVID19
                                                D3
                                                        100 D3_100
##
   106:
           100_D7
                    Male SARS-CoV-2 COVID19
                                                D7
                                                        100 D7_100
```

3. Format the data

61902: ENSG00000275249.1|ENSG00000275249

61903: ENSG00000274792.1|ENSG00000274792

Once the data is loaded in a clean way, make sure that you format the data types to ones that can be used by DESeq - eg. matrix instead of table and so on - This is a good place to filter your data to remove uninformative genes - Here you will also be combining the sample information with the metadata so that they correspond to each other during analysis - NOTE: metadata and data alignment is a key for analysis.

```
## Filter raw counts
##
cnt.dt[1:5,1:5]
##
                                  gene_id D0_02 D0_06 D0_100 D0_11
## 1:
            ENSG00000223764.2|LINC02593
                                              10
                                                     7
                                                             7
                                                                  14
## 2: ENSG00000272438.1|ENSG00000272438
                                                     0
                                                             0
                                                                   0
                                               0
## 3: ENSG00000230699.2|ENSG00000230699
                                               0
                                                     9
                                                            17
                                                                   7
## 4: ENSG00000241180.1|ENSG00000241180
                                               0
                                                     0
                                                             0
                                                                   0
## 5: ENSG00000288531.1|ENSG00000288531
                                                     0
                                                            21
                                                                  30
## summarize raw counts
cnt.dt.sumarized <- cnt.dt[,list(max=max(.SD),</pre>
                                   min=min(.SD),
                                   mean=mean(unlist(.SD))), by=gene_id]
cnt.dt.sumarized
##
                                      gene_id max min
                                                               mean
##
                 ENSG00000223764.2|LINC02593
                                                38
       1:
                                                     0
                                                         6.40566038
##
       2: ENSG00000272438.1 | ENSG00000272438
                                                 2
                                                     0
                                                         0.01886792
##
       3: ENSG00000230699.2|ENSG00000230699
                                                49
                                                         7.40566038
                                                     0
##
       4: ENSG00000241180.1 | ENSG00000241180
                                                 0
                                                     0
                                                         0.00000000
##
       5: ENSG00000288531.1|ENSG00000288531
                                                95
                                                     0 12.57547170
##
```

16

0

2.42452830

1.30188679

```
## 61904: ENSG00000278510.1|ENSG00000278510 6 0 0.35849057
## 61905: ENSG00000277196.4|ENSG00000277196 19
                                                 0 1.67924528
## 61906:
                      ENSG00000277374.1|U1
                                                 0 0.54716981
## get gene names that have sufficient expression
## NOTE: this parameter is subjective and you can/should play with your cutoff value
## NOTE: sometimes, it makes more sense to not use min if you think some genes are on/off in subjects
## NOTE: counts are not like TPM, 10 counts per gene may still mean gene is off
cnt.dt.sumarized[mean > 0]
##
                                   gene id max min
##
               ENSG00000223764.2|LINC02593 38
                                                 0 6.40566038
      2: ENSG00000272438.1|ENSG00000272438
                                                 0 0.01886792
##
                                             2
      3: ENSG00000230699.2|ENSG00000230699 49
                                               0 7.40566038
##
      4: ENSG00000288531.1|ENSG00000288531 95 0 12.57547170
                  ENSG00000230368.2|FAM41C 73
                                                0 14.16981132
##
      5:
##
## 61696: ENSG00000275249.1|ENSG00000275249 16 0 2.42452830
## 61697: ENSG00000274792.1|ENSG00000274792 14
                                                0 1.30188679
## 61698: ENSG00000278510.1|ENSG00000278510 6
                                                0 0.35849057
## 61699: ENSG00000277196.4|ENSG00000277196 19
                                               0 1.67924528
                      ENSG00000277374.1|U1
                                                 0 0.54716981
cnt.dt.sumarized[mean > 10]
##
                                   gene_id max min
                                                          mean
      1: ENSG00000288531.1|ENSG00000288531
##
                                             95
                                                  0
                                                      12.57547
                  ENSG00000230368.2|FAM41C
##
                                             73
                                                  0
                                                      14.16981
##
                 ENSG00000187961.15 | KLHL17 547 28
                                                     200.42453
##
      4:
                ENSG00000187583.11 | PLEKHN1
                                             80
                                                 0
                                                      33.10377
##
                  ENSG00000188976.11|NOC2L 836 102
                                                     411.50000
##
## 24737: ENSG00000267793.1|ENSG00000267793
                                             75
                                                      14.43396
## 24738: ENSG00000260197.1|ENSG00000260197 412
                                                      93.06604
                  ENSG00000012817.16 | KDM5D 7361
                                                 0 1614.75472
## 24740: ENSG00000288049.1|ENSG00000288049 170
                                                      35.42453
## 24741:
                 ENSG00000198692.10|EIF1AY 1259
                                                  0 298.06604
cnt.dt.sumarized[mean > 50]
##
                                   gene_id
                                             max min
                 ENSG00000187961.15|KLHL17
##
      1:
                                             547
                                                  28
                                                      200.42453
##
                  ENSG00000188976.11|NOC2L
                                             836 102
                                                      411.50000
      3: ENSG00000272512.1|ENSG00000272512 1622
                                                   0
                                                       73.54717
##
                  ENSG00000188290.11 | HES4 2402
                                                   0 190.87736
##
      5:
                  ENSG00000187608.10 | ISG15 34685 51 1960.95283
##
## 16065:
                 ENSG00000215580.12|BCORP1
                                                   0 182.10377
                                             913
                 ENSG00000131002.14|TXLNGY 6766
                                                   0 1647.10377
## 16067: ENSG00000260197.1|ENSG00000260197
                                             412
                                                       93.06604
                 ENSG00000012817.16 | KDM5D 7361
## 16068:
                                                   0 1614.75472
                 ENSG00000198692.10 | EIF1AY 1259
## 16069:
                                                   0 298.06604
gene_ids_to_include <- cnt.dt.sumarized[mean > 50][["gene_id"]]
## filter data
```

```
cnt.filtered.dt <- cnt.dt[gene_id %in% gene_ids_to_include]</pre>
cnt.filtered.dt[1:5,1:5]
##
                                 gene_id D0_02 D0_06 D0_100 D0_11
## 1:
              ENSG00000187961.15 | KLHL17
                                           152
                                                 130
                                                         49
                                                                39
## 2:
               ENSG00000188976.11|NOC2L
                                                 380
                                           427
                                                         111
                                                               116
## 3: ENSG00000272512.1|ENSG00000272512
                                           362
                                                  46
                                                          6
                                                                12
## 4:
                ENSG00000188290.11 | HES4 1337
                                                 404
                                                          3
                                                                20
## 5:
               ENSG00000187608.10 | ISG15 11707 4699
                                                               150
                                                         105
##
## Make sample information table
##
## create a sample information table from cnt table
## NOTE: this will make sure you will always have the right samples present
si.dt <- data.table("seq_id"=colnames(cnt.dt)[-1])</pre>
si.dt[,"subject":=tstrsplit(seq_id,split="_",keep = 2)]
si.dt[,"time":=tstrsplit(seq_id,split="_",keep = 1)]
si.dt
##
        seq_id subject time
##
     1: D0 02
                    02
##
     2: D0_06
                    06
                         DO
##
     3: DO 100
                   100
                         D0
##
     4: DO_11
                         D0
                    11
        DO_14
                         D0
##
     5:
                    14
##
## 102: D7_32
                    32
## 103:
        D7_49
                         D7
                    49
        D7_62
                    62
                         D7
## 104:
                    92
                         D7
## 105: D7_92
## 106: D7_93
                    93
                         D7
## Load metadata and add to the sample information
##
# peak at ready metadata
meta.clean.dt
##
        IGU Code
                    sex pathogen disease time subject seq_id
##
     1:
           02 DO Female SARS-CoV-2 COVID19
                                              D0
                                                      02 D0 02
##
     2:
          02_D28 Female SARS-CoV-2 COVID19
                                             D28
                                                      02 D28_02
##
           03_D0 Female SARS-CoV-2 COVID19
                                                      03 D0 03
     3:
##
     4:
          03 D28 Female SARS-CoV-2 COVID19
                                             D28
                                                      03 D28 03
##
          06_D0 Female SARS-CoV-2 COVID19
                                                      06 D0_06
     5:
                                              D0
##
   ---
## 102:
          98_D0
                   Male SARS-CoV-2 COVID19
                                              D0
                                                      98 D0_98
          98_D28
## 103:
                   Male SARS-CoV-2 COVID19
                                             D28
                                                      98 D28_98
          100_D0
## 104:
                   Male SARS-CoV-2 COVID19
                                              D0
                                                     100 DO_100
## 105:
          100 D3
                                                      100 D3_100
                   Male SARS-CoV-2 COVID19
                                              D3
## 106:
          100_D7
                   Male SARS-CoV-2 COVID19
                                              D7
                                                     100 D7_100
# add to si.dt to make a master table (mt)
si.dt
```

```
##
        seq_id subject time
##
                          DO
     1: D0_02
                     02
     2: D0 06
##
                     06
                          D0
                   100
##
     3: D0_100
                          DO
##
     4: DO 11
                     11
                          D0
##
     5: DO 14
                     14
                          D0
##
## 102: D7_32
                     32
                          D7
        D7_49
## 103:
                     49
                          D7
                     62
                          D7
## 104: D7_62
## 105: D7_92
                     92
                          D7
## 106: D7_93
                     93
                          D7
si.mt.dt <- meta.clean.dt[si.dt,on=.(seq_id=seq_id,time=time,subject=subject)]</pre>
si.mt.dt[1:5,]
##
      IGU Code
                         pathogen disease time subject seq_id
## 1:
         02_D0 Female SARS-CoV-2 COVID19
                                             D0
                                                     02 D0_02
## 2:
         O6_DO Female SARS-CoV-2 COVID19
                                             DO
                                                     06 D0_06
## 3:
        100 D0
                 Male SARS-CoV-2 COVID19
                                             DO
                                                    100 DO_100
## 4:
         11_DO Female SARS-CoV-2 COVID19
                                             DO
                                                     11 DO_11
## 5:
         14 DO
                 Male SARS-CoV-2 COVID19
                                                     14 DO 14
                                             DO
## filter table to keep only comparison samples
si.mt.comp.dt <- si.mt.dt[time %in% c("D0","D28")]</pre>
si.mt.comp.dt[,.N,by=list(disease, time)]
##
          disease time N
## 1:
          COVID19
                    DO 30
## 2: Non-COVID19
                    DO 24
## 3:
          COVID19 D28 28
## 4: Non-COVID19 D28 12
si.mt.comp.dt[,.N,by=time]
##
      time N
## 1:
        D<sub>0</sub> 54
## 2: D28 40
##
## Filter count table to keep comparison columns
##
# present columns and their format
colnames(cnt.filtered.dt)
                              "D0 06"
##
     [1] "gene_id" "D0_02"
                                         "D0_100"
                                                   "D0_11"
                                                              "D0_14"
                                                                        "D0_21"
                    "D0_31"
                              "D0 32"
                                         "DO 34"
                                                   "D0 35"
                                                              "D0 36"
                                                                        "D0_37"
##
     [8] "D0 26"
##
    [15] "D0_39"
                    "D0 03"
                              "D0 43"
                                         "DO 45"
                                                   "D0 46"
                                                              "DO 47"
                                                                        "D0 49"
                                                              "D0_58"
    [22] "D0 51"
                    "D0 53"
                              "D0 54"
                                         "D0 56"
                                                   "D0 57"
                                                                        "D0 60"
##
   [29] "D0_61"
                    "D0_62"
                              "D0 67"
                                         "D0 68"
                                                   "D0_69"
                                                              "D0_70"
                                                                        "D0 72"
    [36] "D0_74"
                    "D0 75"
                              "DO 77"
                                         "D0 78"
                                                   "D0 79"
                                                              "D0 80"
                                                                        "D0 81"
##
                              "D0_86"
                                                              "D0 89"
##
   [43] "D0_82"
                    "D0 85"
                                         "D0 87"
                                                   "D0 88"
                                                                        "DO 90"
   [50] "D0 91"
                    "D0 92"
                              "D0 93"
                                         "DO 94"
                                                   "D0 95"
                                                              "D0 98"
                                                                        "D28 02"
    [57] "D28_03"
                                                   "D28_21"
                                                              "D28_32"
##
                    "D28_06"
                              "D28_11"
                                         "D28_14"
                                                                        "D28_36"
##
    [64] "D28_37"
                    "D28_39"
                              "D28 43"
                                         "D28 45"
                                                   "D28 47"
                                                              "D28 49"
                                                                        "D28 51"
   [71] "D28_54"
                    "D28_56"
                              "D28_57"
                                         "D28_58"
                                                   "D28_60"
                                                              "D28_62"
                                                                        "D28_67"
##
```

```
[78] "D28 75"
                   "D28 77"
                             "D28 78"
                                       "D28 79"
                                                  "D28 80"
                                                            "D28 81"
                                                                      "D28 82"
##
   [85] "D28 85"
                             "D28 87"
                                       "D28 88"
                                                  "D28 89"
                                                            "D28 90"
                                                                      "D28 91"
##
                   "D28 86"
  [92] "D28 92"
                   "D28 93"
                             "D28 94"
                                       "D28 98"
                                                  "D3 100"
                                                            "D3 32"
                                                                      "D3 49"
## [99] "D3_62"
                   "D3 92"
                             "D3_93"
                                       "D7_100"
                                                  "D7_32"
                                                            "D7_49"
                                                                      "D7_62"
## [106] "D7 92"
                   "D7 93"
# wanted columns and their format matching
si.mt.comp.dt[1:5,]
##
      IGU_Code
                        pathogen disease time subject seq_id
                  sex
         02_D0 Female SARS-CoV-2 COVID19
                                                    02 D0_02
## 1:
                                           DO
        06 DO Female SARS-CoV-2 COVID19
                                           DO
                                                    06 D0 06
## 3:
       100_D0
                 Male SARS-CoV-2 COVID19
                                           DO
                                                   100 DO_100
        11_DO Female SARS-CoV-2 COVID19
                                           DO
                                                    11 DO_11
         14_D0
## 5:
                 Male SARS-CoV-2 COVID19
                                                    14 DO_14
                                           DO
wanted_comp_columns <- si.mt.comp.dt[["seq_id"]]</pre>
wanted_comp_columns
                                                               "D0 26"
   [1] "D0 02"
                 "D0 06"
                          "D0 100" "D0 11"
                                            "DO 14"
                                                      "D0 21"
                                                                        "D0 31"
   [9] "D0_32"
                 "D0_34"
                          "D0 35"
                                   "D0 36"
                                             "D0_37"
                                                      "D0_39"
                                                               "D0 03"
                                                                        "D0 43"
##
## [17] "D0 45"
                 "DO 46"
                          "DO 47"
                                   "DO 49"
                                             "D0 51"
                                                      "D0 53"
                                                               "D0 54"
                                                                        "D0 56"
## [25] "D0_57"
                 "D0_58"
                          "D0 60"
                                   "DO 61"
                                            "D0_62" "D0_67"
                                                               "D0 68"
                                                                        "D0 69"
## [33] "D0 70"
                 "D0 72"
                          "DO 74"
                                   "D0 75"
                                            "DO 77"
                                                      "D0 78"
                                                               "D0 79"
                                                                        "D0 80"
                 "D0_82"
                          "D0_85"
                                             "D0_87"
                                                      "D0_88"
                                                               "D0_89"
## [41] "D0_81"
                                   "D0_86"
                                                                        "D0_90"
## [49] "D0 91" "D0 92"
                          "D0 93" "D0 94"
                                             "DO 95"
                                                      "D0 98"
                                                               "D28 02" "D28 03"
## [57] "D28 06" "D28 11" "D28 14" "D28 21" "D28 32" "D28 36" "D28 37" "D28 39"
## [65] "D28 43" "D28 45" "D28 47" "D28 49" "D28 51" "D28 54" "D28 56" "D28 57"
## [73] "D28_58" "D28_60" "D28_62" "D28_67" "D28_75" "D28_77" "D28_78" "D28_78"
## [81] "D28_80" "D28_81" "D28_82" "D28_85" "D28_86" "D28_87" "D28_88" "D28_89"
## [89] "D28 90" "D28 91" "D28 92" "D28 93" "D28 94" "D28 98"
## filter raw counts to keep the same samples
cnt.filtered.comp.dt <- cnt.filtered.dt[,.SD,.SDcols = c("gene_id",wanted_comp_columns)]</pre>
cnt.filtered.comp.dt[1:5,1:5]
                                gene_id D0_02 D0_06 D0_100 D0_11
##
## 1:
              ENSG00000187961.15 | KLHL17
                                           152
                                                 130
                                                               39
              ENSG00000188976.11|NOC2L
                                                 380
                                           427
                                                        111
                                                              116
## 3: ENSG00000272512.1|ENSG00000272512
                                           362
                                                 46
                                                          6
                                                               12
## 4:
               ENSG00000188290.11 | HES4 1337
                                                 404
                                                          3
                                                               20
               ENSG00000187608.10 | ISG15 11707 4699
## 5:
                                                        105
                                                              150
## Format into right types
##date
## counts need to be a matrix where rownames are gene_id
cnt.comp.mat <- as.matrix(x = cnt.filtered.comp.dt, rownames = "gene_id")</pre>
## sample information can remain a data table
si.mt.comp.dt[1:5,]
##
      IGU_Code
                  sex
                        pathogen disease time subject seq_id
## 1:
         O2_DO Female SARS-CoV-2 COVID19
                                           DO
                                                   02 D0 02
## 2:
                                           DO
         O6_DO Female SARS-CoV-2 COVID19
                                                    06 D0_06
               Male SARS-CoV-2 COVID19
                                           DO
## 3:
        100 D0
                                                   100 DO 100
```

```
## 4: 11_D0 Female SARS-CoV-2 COVID19 D0 11 D0_11 ## 5: 14_D0 Male SARS-CoV-2 COVID19 D0 14 D0_14
```

4. Run DESeq2 analysis

- Once the data has been prepared, the DESeq package can be employed and comparative analysis performed. The analysis consists of three simple steps:
 - 1. Create a DESeq object using the raw counts and metadata from previous section. And specifying the comparison MODEL.
 - 2. Running the DESeq command.
 - 3. Retrieval of the result tables for plotting and analysis.

```
##
## Create a DESeq object
##
## Data
#cnt.comp.mat[1:5,1:5]
\#si.mt.comp.dt[1:5]
## load data into deseq object
dds <- DESeqDataSetFromMatrix(countData = cnt.comp.mat,</pre>
                               colData = si.mt.comp.dt,
                               design = ~time)
## Warning in DESeqDataSet(se, design = design, ignoreRank): some variables in
## design formula are characters, converting to factors
## add condition to the modeling
dds.sex <- DESeqDataSetFromMatrix(countData = cnt.comp.mat,</pre>
                               colData = si.mt.comp.dt,
                               design = ~time+sex)
## Warning in DESeqDataSet(se, design = design, ignoreRank): some variables in
## design formula are characters, converting to factors
## Run DESeq Analysis
##
## two modes - with and without sex consideration
dds <- DESeq(dds)</pre>
## estimating size factors
## estimating dispersions
## gene-wise dispersion estimates
## mean-dispersion relationship
## final dispersion estimates
## fitting model and testing
## -- replacing outliers and refitting for 403 genes
## -- DESeq argument 'minReplicatesForReplace' = 7
## -- original counts are preserved in counts(dds)
## estimating dispersions
```

```
## fitting model and testing
dds.sex <- DESeq(dds.sex)</pre>
## estimating size factors
## estimating dispersions
## gene-wise dispersion estimates
## mean-dispersion relationship
## final dispersion estimates
## fitting model and testing
## -- replacing outliers and refitting for 273 genes
## -- DESeq argument 'minReplicatesForReplace' = 7
## -- original counts are preserved in counts(dds)
## estimating dispersions
## fitting model and testing
##
## View and retrieve the results
##
## Look at results without sex consideration
resultsNames(dds)
## [1] "Intercept"
                        "time D28 vs D0"
res <- results(object = dds, name = "time_D28_vs_D0", alpha = 0.05)
summary(res)
##
## out of 16068 with nonzero total read count
## adjusted p-value < 0.05
## LFC > 0 (up)
                     : 931, 5.8%
## LFC < 0 (down)
                     : 2038, 13%
                     : 0, 0%
## outliers [1]
## low counts [2]
                     : 1, 0.0062%
## (mean count < 0)</pre>
## [1] see 'cooksCutoff' argument of ?results
## [2] see 'independentFiltering' argument of ?results
## Look at the results with sex consideration
resultsNames(dds.sex)
## [1] "Intercept"
                            "time_D28_vs_D0"
                                                  "sex_Male_vs_Female"
res.sex <- results(object = dds.sex, name = "time_D28_vs_D0", alpha = 0.05)
summary(res.sex)
##
## out of 16069 with nonzero total read count
## adjusted p-value < 0.05
## LFC > 0 (up)
                    : 934, 5.8%
## LFC < 0 (down)
                    : 1926, 12%
## outliers [1]
                    : 0, 0%
## low counts [2]
                     : 0, 0%
```

```
## (mean count < 2)
## [1] see 'cooksCutoff' argument of ?results
## [2] see 'independentFiltering' argument of ?results
## export a table of results for each
res.dt <- as.data.table(results(object = dds, name = "time_D28_vs_D0", alpha = 0.05), keep.rownames=TRU
## Warning in .local(x, row.names, optional, ...): Arguments in '...' ignored
colnames(res.dt) <- gsub("rn", "gene_id", colnames(res.dt))</pre>
res.sex.dt <- as.data.table(results(object = dds.sex, name = "time_D28_vs_D0", alpha = 0.05), keep.rown
## Warning in .local(x, row.names, optional, ...): Arguments in '...' ignored
colnames(res.sex.dt) <- gsub("rn", "gene_id", colnames(res.sex.dt))</pre>
## RESULT TABLES
res.dt <- res.dt[order(padj,log2FoldChange)]</pre>
##
                                     gene_id baseMean log2FoldChange
                                                                            lfcSE
##
                 ENSG00000108387.16|SEPTIN4 312.7207
                                                       -2.802464e+00 0.27604306
       1:
##
       2:
                   ENSG00000165949.13 | IFI27 1174.0043
                                                        -4.364281e+00 0.44337569
##
                   ENSG00000184979.11|USP18 830.0480
       3:
                                                        -3.040946e+00 0.32843781
##
                 ENSG00000196141.14|SPATS2L 995.0307
                                                        -2.607314e+00 0.28203196
       4:
##
                   ENSG00000187608.10 | ISG15 2014.3514
                                                        -3.131625e+00 0.35052613
       5:
##
## 16065:
                   ENSG00000155903.14|RASA2 3779.8496
                                                        -2.503228e-05 0.04456558
## 16066:
                  ENSG00000257246.2|PHB1P19 104.8990
                                                        -4.350574e-05 0.11009555
## 16067:
                ENSG00000286219.2|NOTCH2NLC 4280.7756
                                                        -9.717332e-06 0.08605497
                    ENSG00000165195.16 | PIGA 497.5808
                                                         3.578511e-06 0.05084592
## 16068:
                                                         0.000000e+00 0.00000000
## 16069: ENSG00000269693.1|ENSG00000269693
                                                0.0000
##
                              pvalue
                   stat
                                              padj
##
       1: -1.015227e+01 3.237387e-24 5.201833e-20
##
       2: -9.843303e+00 7.326482e-23 5.886095e-19
       3: -9.258817e+00 2.067101e-20 9.472012e-17
##
##
       4: -9.244747e+00 2.357982e-20 9.472012e-17
##
       5: -8.934070e+00 4.106169e-19 1.319558e-15
##
## 16065: -5.616955e-04 9.995518e-01 9.997385e-01
## 16066: -3.951635e-04 9.996847e-01 9.998092e-01
## 16067: -1.129201e-04 9.999099e-01 9.999438e-01
## 16068: 7.037950e-05 9.999438e-01 9.999438e-01
## 16069: 0.000000e+00 1.000000e+00
res.sex.dt <- res.sex.dt[order(padj,log2FoldChange)]
res.sex.dt
##
                                     gene_id
                                                baseMean log2FoldChange
                                                                              lfcSE
##
                   ENSG00000165949.13|IFI27 1174.00428 -4.321991e+00 0.44438975
##
       2:
                   ENSG00000184979.11 | USP18
                                               830.04803 -2.860923e+00 0.31575707
##
                   ENSG00000187608.10 | ISG15 2014.35142
                                                          -3.084138e+00 0.34314627
                  ENSG00000142089.17 | IFITM3 13186.71862 -2.023653e+00 0.23732131
##
       4:
                   ENSG00000161133.18|USP41
##
       5:
                                                72.44594 -2.609214e+00 0.30872184
##
## 16065:
                     ENSG00000172336.5|POP7
                                                85.59053 -4.234665e-05 0.09927925
```

```
## 16066:
                   ENSG00000063601.17 | MTMR1 1324.63352 -9.389151e-06 0.04372301
## 16067: ENSG00000276136.1|ENSG00000276136
                                               452.38103
                                                           2.133293e-05 0.10818643
## 16068:
                  ENSG00000196417.13 | ZNF765
                                               513.27454
                                                           2.447131e-05 0.08010933
## 16069:
                                                           1.316112e-04 0.19918038
                     ENSG00000134548.11|SPX
                                               60.77116
##
                   stat
                              pvalue
                                              padj
       1: -9.7256768058 2.343417e-22 3.765638e-18
##
       2: -9.0605202753 1.298298e-19 1.043118e-15
##
       3: -8.9878243218 2.521727e-19 1.350721e-15
##
##
       4: -8.5270582503 1.501163e-17 6.030547e-14
##
       5: -8.4516667939 2.871732e-17 8.620821e-14
##
## 16065: -0.0004265408 9.996597e-01 9.998427e-01
## 16066: -0.0002147417 9.998287e-01 9.998427e-01
## 16067: 0.0001971868 9.998427e-01 9.998427e-01
## 16068: 0.0003054739 9.997563e-01 9.998427e-01
## 16069: 0.0006607637 9.994728e-01 9.998427e-01
```

PART2

Goal: - Explore ways of plotting results from DESeq2 analysis - Use the results in Gene Set Enrichemnt Analysis

1. Plot PCA of the results

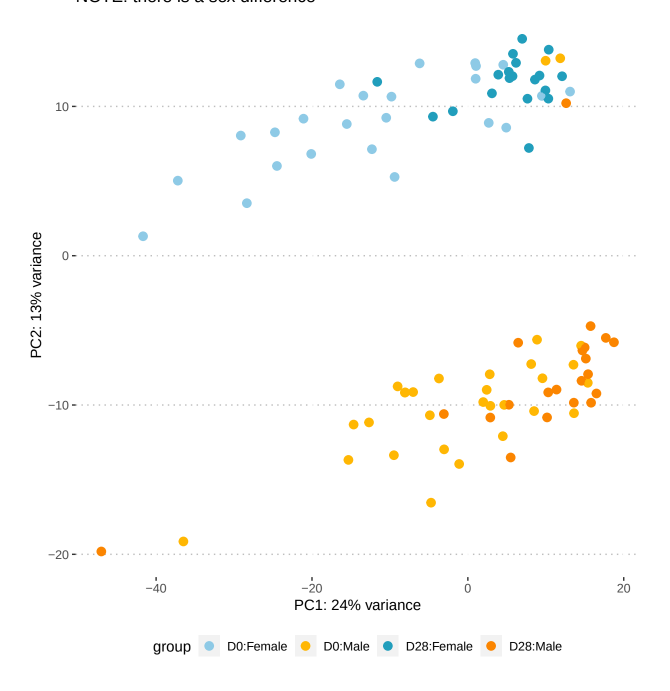
- PCA plot is one of the typical plots to evaluate whether there are any patterns in your data
- First, the data is normalized

```
# Input data
#dds.sex
#si.mt.comp.dt
## Stabilize the data using variance stabilizing transformation
vsd.sex <- vst(object = dds.sex)</pre>
vsd.sex
## class: DESegTransform
## dim: 16069 94
## metadata(1): version
## assays(1): ''
## rownames(16069): ENSG00000187961.15|KLHL17 ENSG00000188976.11|NOC2L ...
     ENSG00000012817.16 | KDM5D ENSG00000198692.10 | EIF1AY
##
## rowData names(27): baseMean baseVar ... replace dispFit
## colnames(94): D0_02 D0_06 ... D28_94 D28_98
## colData names(9): IGU_Code sex ... sizeFactor replaceable
## Use native DESeq PCA plotting capabilities
#?plotPCA
plotPCA(object = vsd.sex, intgroup = c("time", "sex"))
```



Modify the plot by saving into object and adjusting the ggplot parameters within it ## \rightarrow https://coolors.co/

PCA using sex-adjusted DESeq2 results NOTE: there is a sex difference



```
##
## PCA on dds (no sex adjustment)
## -> The PCA looks the same, however, the resulting genes are different due to different model [~time+
```

```
## Stabilize the data
#vsd <- vst(object = dds)

## Plot similar plot using custom ggplot
#pca.plot <- plotPCA(object = vsd, intgroup = c("time", "sex"))
#pca.plot + theme_pubclean() +
# geom_point(size = 2) +
# ggtitle(pasteO("PCA using NOT-adjusted for sex DESeq2 results\n",
# "NOTE: there is a sex difference")) +
# scale_colour_manual(values = four_colors) +
# theme(aspect.ratio = 1,
# legend.position = "right")</pre>
```

2. Plot Boxplot of the normalized counts

- It can be an important control or piece of data to look at the distribution of expression of a particular gene in your data
- To do so we extract the normalized counts (or use TPM data) and use boxplot that summarized number of imporatant statistics including median, quantiles and outliers

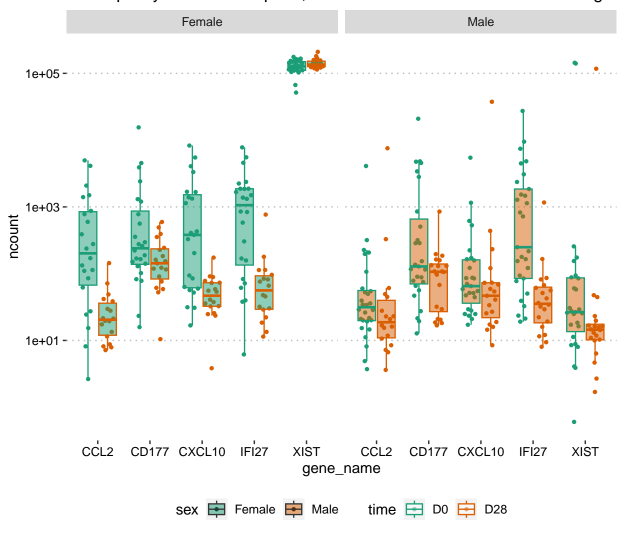
```
## Input
#dds.sex
#si.mt.comp.dt
## Extract normalized count data
# ?counts <- function that extracts normalized data from dds object
ncount.dt <- as.data.table(counts(dds.sex, normalized=TRUE), keep.rownames = TRUE)</pre>
colnames(ncount.dt) <- gsub("rn", "gene_id", colnames(ncount.dt))</pre>
ncount.dt[1:5,1:5]
##
                                              D0_02
                                                          DO 06
                                                                    D0_100
                                 gene_id
                                                                                DO_11
## 1:
              ENSG00000187961.15 | KLHL17
                                           148.5576 157.58938 122.268608 147.48129
## 2:
               ENSG00000188976.11|NOC2L
                                           417.3295
                                                      460.64587 276.975826 438.66230
## 3: ENSG00000272512.1 | ENSG00000272512
                                           353.8016
                                                       55.76239
                                                                 14.971666
                                                                            45.37886
## 4:
                ENSG00000188290.11|HES4 1306.7203 489.73929
                                                                  7.485833
                                                                            75.63143
## 5:
               ENSG00000187608.10|ISG15 11441.8660 5696.24984 262.004160 567.23573
## Re-arrange the table and get gene names
ncount.dtm <- melt.data.table(data = ncount.dt, id.vars = "gene_id",</pre>
                               variable.name = "subject",
                               value.name = "ncount")
ncount.dtm[,"gene_name" := tstrsplit(gene_id,split="\\|",keep = 2)]
ncount.dtm
##
                                       gene_id subject
                                                            ncount
                                                                          gene_name
##
         1:
                    ENSG00000187961.15 | KLHL17
                                                  D0 02
                                                          148.5576
                                                                             KLHL17
##
                     ENSG00000188976.11|NOC2L
                                                  D0_02
                                                          417.3295
                                                                              NOC2L
         2:
         3: ENSG00000272512.1|ENSG00000272512
                                                          353.8016 ENSG00000272512
##
                                                 D0 02
         4:
##
                      ENSG00000188290.11 | HES4
                                                  DO_02 1306.7203
                                                                               HES4
##
                     ENSG00000187608.10|ISG15
                                                  DO 02 11441.8660
                                                                              ISG15
##
## 1510482:
                    ENSG00000215580.12|BCORP1 D28 98
                                                          486.9312
                                                                             BCORP1
```

```
## 1510483:
                     ENSG00000131002.14|TXLNGY D28 98
                                                          3584.4561
                                                                              TXLNGY
## 1510484: ENSG00000260197.1|ENSG00000260197
                                                 D28 98
                                                           181.7779 ENSG00000260197
## 1510485:
                      ENSG00000012817.16 | KDM5D
                                                 D28 98
                                                          3306.3140
                                                                               KDM5D
## 1510486:
                     ENSG00000198692.10 | EIF1AY
                                                 D28 98
                                                           617.6069
                                                                              EIF1AY
## Combine the normalized counts with metadata
ncount.dtm <- ncount.dtm[si.mt.comp.dt,on=.(subject=seq_id)]</pre>
ncount.dtm[1:5]
##
                                 gene_id subject
                                                                    gene_name
                                                      ncount
## 1:
              ENSG00000187961.15 | KLHL17
                                            D0 02
                                                    148.5576
                                                                       KLHL17
               ENSG00000188976.11|NOC2L
                                            D0 02
                                                                         NOC2L
## 2:
                                                    417.3295
## 3: ENSG00000272512.1|ENSG00000272512
                                            D0 02
                                                    353.8016 ENSG00000272512
                                            D0_02 1306.7203
## 4:
                ENSG00000188290.11 | HES4
                                                                         HES4
                                            DO_02 11441.8660
## 5:
               ENSG00000187608.10 | ISG15
                                                                         ISG15
##
      IGU_Code
                         pathogen disease time i.subject
                   sex
## 1:
         02_D0 Female SARS-CoV-2 COVID19
                                             DO
                                                        02
                                                        02
## 2:
         02 DO Female SARS-CoV-2 COVID19
                                             DO
                                                        02
## 3:
         02_D0 Female SARS-CoV-2 COVID19
                                             D0
## 4:
         02_D0 Female SARS-CoV-2 COVID19
                                             DO
                                                        02
## 5:
         02_D0 Female SARS-CoV-2 COVID19
                                                        02
## select genes of interest
goi <- c("IFI27","CCL2","CD177","XIST","CXCL10")</pre>
## subset the count table
ncount.goi.dtm <- ncount.dtm[gene name %in% goi]</pre>
ncount.goi.dtm
##
                          gene_id subject
                                                 ncount gene_name IGU_Code
                                                                                sex
##
     1: ENSG00000169245.6 CXCL10
                                     D0_02
                                             4062.85445
                                                            CXCL10
                                                                       02_D0 Female
##
     2: ENSG00000165949.13|IFI27
                                     D0_02
                                                                       02_D0 Female
                                              581.52475
                                                             IFI27
##
     3: ENSG00000108691.10 | CCL2
                                     D0_02
                                             1400.54617
                                                              CCL2
                                                                       02_D0 Female
                                    D0_02
                                               78.18820
##
     4: ENSG00000204936.10 | CD177
                                                             CD177
                                                                      02_D0 Female
##
        ENSG00000229807.13 | XIST
                                    DO 02 142701.28745
                                                              XIST
                                                                      02_D0 Female
##
   466: ENSG00000169245.6 CXCL10
                                   D28 98
                                                                     98 D28
                                                                               Male
                                               73.00318
                                                            CXCL10
                                                                     98_D28
   467: ENSG00000165949.13|IFI27
                                   D28 98
                                                                               Male
                                                8.03035
                                                             IFI27
         ENSG00000108691.10 | CCL2
                                   D28 98
                                                                     98 D28
                                                                               Male
   468:
                                               16.79073
                                                              CCL2
                                   D28_98
   469: ENSG00000204936.10 CD177
                                               20.44089
                                                             CD177
                                                                     98 D28
                                                                               Male
                                                                     98 D28
        ENSG00000229807.13 | XIST
                                   D28 98
                                               44.53194
                                                              XIST
                                                                               Male
##
          pathogen disease time i.subject
##
     1: SARS-CoV-2 COVID19
                              D0
     2: SARS-CoV-2 COVID19
                              D0
                                         02
##
##
     3: SARS-CoV-2 COVID19
                              D0
                                         02
##
     4: SARS-CoV-2 COVID19
                              D0
                                         02
##
     5: SARS-CoV-2 COVID19
                                         02
##
## 466: SARS-CoV-2 COVID19
                             D28
                                         98
## 467: SARS-CoV-2 COVID19
                             D28
                                         98
## 468: SARS-CoV-2 COVID19
                             D28
                                         98
## 469: SARS-CoV-2 COVID19
                                         98
## 470: SARS-CoV-2 COVID19
                             D28
                                         98
## Boxplot with all the points
ggplot() + theme_pubclean() +
```

```
# plots all the points
  geom_quasirandom(data = ncount.goi.dtm,
                   aes(x = gene_name, y = ncount,
                      fill = sex, colour = time),
                   dodge.width = 0.8, size = 1) +
  geom_boxplot(data = ncount.goi.dtm,
               aes(x = gene_name, y = ncount,
                   fill = sex, colour = time),
               alpha = 0.5, outlier.shape = NA) +
  ggtitle(paste0("Boxplot with ggbeeswarm plot showing distribution of the counts\n",
                 "Data split by sex and timepoint; NOTE - there are few MALEs with high XIST expression
  scale_colour_brewer(palette = "Dark2") +
  scale_fill_brewer(palette = "Dark2") +
  # use wrap to conviniently re-arrange results
 facet_wrap(~sex) +
  scale_y_log10() +
  theme(aspect.ratio = 1.5,
       axis.text = element_text(colour = "black"),
       legend.position = "bottom")
## Warning: Transformation introduced infinite values in continuous y-axis
## Transformation introduced infinite values in continuous y-axis
## Warning: Removed 26 rows containing non-finite values (`stat_boxplot()`).
```

Warning: Removed 26 rows containing missing values (`geom_point()`).

Boxplot with ggbeeswarm plot showing distribution of the counts Data split by sex and timepoint; NOTE – there are few MALEs with high >



```
## NOTE: PRACTICE - Try identifying and plotting sex-specific
## genes that could help distinguish genetic sex
```

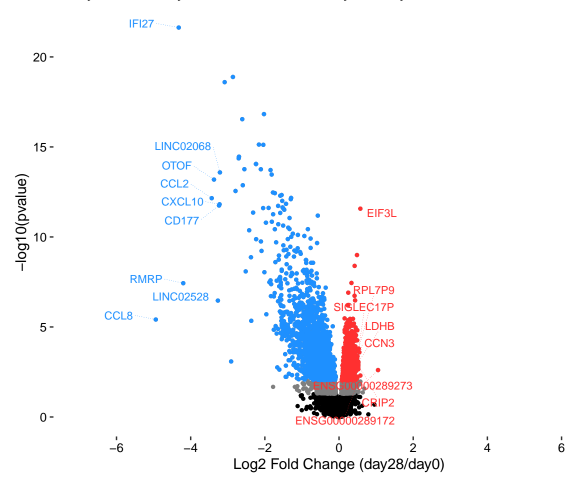
3. Volcano plot

- Volcanos are a common way to show overall change in gene expression in comparison of two conditions
- They combine statistical information with directional expression change information
- It is also nice to highlight few genes of interest on these plots

```
## Input data
#res.sex.dt
## Sig. up and down
## NOTE: ideally by padj value and can also be done by pvalue
res.up.dt <- res.sex.dt[padj <= 0.05][log2FoldChange > 0][order(-log2FoldChange)][1:10]
res.dn.dt <- res.sex.dt[padj <= 0.05][log2FoldChange < 0][order(log2FoldChange)][1:10]
## Volcano
ggplot() + theme_pubclean() +
  # plot non-significant points
  geom_point(data = res.sex.dt[pvalue > 0.05], ## non-significant genes - all
             aes(x = log2FoldChange, y = -log10(pvalue)),
             size=1, colour = "black") +
  # plot points by significant pvalue
  geom_point(data = res.sex.dt[pvalue <= 0.05],</pre>
             aes(x = log2FoldChange, y = -log10(pvalue)),
             size=1, colour = "grey50") +
  # plot only top significant - INCREASED
  geom_point(data = res.sex.dt[padj <= 0.05][log2FoldChange > 0],
             aes(x = log2FoldChange, y = -log10(pvalue)),
             size=1, colour = "firebrick1") +
  # add labels
  geom_text_repel(data = res.up.dt,
             aes(x = log2FoldChange, y = -log10(pvalue),
                 label = unlist(tstrsplit(gene_id,split="\\|",keep = 2))),
             size=3, colour = "firebrick1", segment.linetype = "dotted",
             nudge_x = 1,
             direction = "v",
             force = 2,
             force_pull = NA,
             vjust=1,
             hjust=1,
             segment.size = 0.2) +
  # plot only top signficant - DECREASED
  geom_point(data = res.sex.dt[padj <= 0.05][log2FoldChange < 0],</pre>
             aes(x = log2FoldChange, y = -log10(pvalue)),
             size=1, colour = "dodgerblue") +
  # add labels
  geom_text_repel(data = res.dn.dt,
             aes(x = log2FoldChange, y = -log10(pvalue),
```

```
label = unlist(tstrsplit(gene_id,split="\\|",keep = 2))),
             size=3, colour = "dodgerblue", segment.linetype = "dotted",
             nudge_x = -1,
             force = 3,
             force_pull = NA,
             vjust=0,
             direction = "y",
             segment.size = 0.2) +
  # add scales and extras
  scale_x_continuous(limits = c(-7,7), breaks = seq(-10,10,2)) +
  ggtitle(paste0("Volcano showing results of DGE analysis comparing d28 versus d0\n",
                 "red/blue indicate genes significantly changed in this comparison\n",
                 "many inflamatory markers decrease by 28 days")) +
  xlab("Log2 Fold Change (day28/day0)") +
  theme(aspect.ratio = 0.75,
       panel.grid.major.y = element_blank(),
        axis.text = element_text(colour = "black"))
## Warning: Removed 2 rows containing missing values (`geom_point()`).
## Warning: Removed 1 rows containing missing values (`geom_point()`).
## Warning: Removed 1 rows containing missing values (`geom_text_repel()`).
## Warning: Removed 1 rows containing missing values (`geom_point()`).
## Warning: Removed 1 rows containing missing values (`geom_text_repel()`).
## Warning: ggrepel: 1 unlabeled data points (too many overlaps). Consider
## increasing max.overlaps
```

Volcano showing results of DGE analysis comparing d28 versus d0 red/blue indicate genes significantly changed in this comparison many inflamatory markers decrease by 28 days



4. Gene Set Enrichemnt Analysis (GSEA)

- After identification of differentially changing genes, it is very informative to determine whether these changes amount to any systemic / pathway-specific changes
- It is much easier to interpret results of DGE in terms of pathway incread of single gene such analysis has more significance
- To do this we extract significantly changing genes from DESeq results
 - When there are too few *padj* significant genes it is possible to do GSEA with genes that pass *pvalue* significance
 - If there are not significant genes even by pvalue, analysis really looses significance
- Multiple curated sets can be found here: https://www.gsea-msigdb.org/gsea/msigdb/

```
## input data
#res.sex.dt
## First order the results by the log2FolChange from increasing to decreasing
## NOTE: write into new object to not overwrite the original results
res.sex.gsea.dt <- res.sex.dt[order(-log2FoldChange)]</pre>
res.sex.gsea.dt
##
                                     gene_id
                                                baseMean log2FoldChange
                                                                             lfcSE
       1: ENSG00000269693.1|ENSG00000269693
##
                                                16.47864
                                                              24.8272814 2.9422018
       2: ENSG00000289273.1|ENSG00000289273
##
                                                51.07667
                                                               1.0517758 0.3475690
##
       3:
                    ENSG00000253755.1 | IGHGP
                                                52.29388
                                                               0.9532807 0.7521117
##
       4:
               ENSG00000261796.1|ISY1-RAB43
                                                65.39494
                                                               0.7907526 2.1478173
##
       5:
                 ENSG00000184702.20|SEPTIN5
                                                96.50035
                                                               0.6822781 0.3072850
##
                    ENSG00000108691.10 | CCL2
                                                              -3.4328214 0.4781748
## 16065:
                                               266.02706
                                                              -4.2001709 0.7630199
##
  16066:
                     ENSG00000269900.3 RMRP 19150.27418
## 16067:
                   ENSG00000165949.13 | IFI27
                                              1174.00428
                                                              -4.3219911 0.4443898
## 16068:
                     ENSG00000108700.5 | CCL8
                                                69.89154
                                                              -4.9407004 1.0697586
## 16069: ENSG00000215472.10|RPL17-C18orf32
                                                15.85758
                                                             -10.6915338 2.6299579
                            pvalue
##
                stat
                                           padj
##
           8.4383340 3.218926e-17 8.620821e-14
       1:
##
           3.0260916 2.477372e-03 2.112998e-02
           1.2674723 2.049865e-01 3.949952e-01
##
##
           0.3681657 7.127497e-01 8.347187e-01
##
           2.2203429 2.639550e-02 1.031742e-01
##
## 16065: -7.1790102 7.021789e-13 4.339736e-10
  16066: -5.5046674 3.698659e-08 4.571827e-06
## 16067: -9.7256768 2.343417e-22 3.765638e-18
## 16068: -4.6185189 3.864887e-06 1.996941e-04
## 16069: -4.0652870 4.797338e-05 1.265820e-03
## Make sure gene_name is available, if not extract it to new column
res.sex.gsea.dt[,"gene_name" := tstrsplit(gene_id,split="\\|",keep = 2)]
res.sex.gsea.dt
##
                                     gene_id
                                                baseMean log2FoldChange
                                                                             1fcSE
##
       1: ENSG00000269693.1|ENSG00000269693
                                                16.47864
                                                              24.8272814 2.9422018
##
       2: ENSG00000289273.1|ENSG00000289273
                                                51.07667
                                                               1.0517758 0.3475690
```

52.29388

0.9532807 0.7521117

ENSG00000253755.1 | IGHGP

##

3:

```
65.39494
##
               ENSG00000261796.1|ISY1-RAB43
                                                              0.7907526 2.1478173
##
                 ENSG00000184702.20|SEPTIN5
                                                96.50035
                                                              0.6822781 0.3072850
##
                    ENSG00000108691.10 | CCL2
                                               266.02706
                                                             -3.4328214 0.4781748
## 16065:
## 16066:
                     ENSG00000269900.3 RMRP 19150.27418
                                                             -4.2001709 0.7630199
## 16067:
                   ENSG00000165949.13 | IFI27
                                             1174.00428
                                                             -4.3219911 0.4443898
## 16068:
                     ENSG00000108700.5 | CCL8
                                                69.89154
                                                             -4.9407004 1.0697586
                                                15.85758
## 16069: ENSG00000215472.10|RPL17-C18orf32
                                                            -10.6915338 2.6299579
##
                stat
                           pvalue
                                           padj
                                                      gene_name
           8.4383340 3.218926e-17 8.620821e-14 ENSG00000269693
##
##
           3.0260916 2.477372e-03 2.112998e-02 ENSG00000289273
       3: 1.2674723 2.049865e-01 3.949952e-01
##
                                                          IGHGP
##
       4: 0.3681657 7.127497e-01 8.347187e-01
                                                     ISY1-RAB43
##
       5: 2.2203429 2.639550e-02 1.031742e-01
                                                        SEPTIN5
##
## 16065: -7.1790102 7.021789e-13 4.339736e-10
                                                           CCL2
## 16066: -5.5046674 3.698659e-08 4.571827e-06
                                                           RMRP
## 16067: -9.7256768 2.343417e-22 3.765638e-18
                                                          IFI27
## 16068: -4.6185189 3.864887e-06 1.996941e-04
                                                           CCL8
## 16069: -4.0652870 4.797338e-05 1.265820e-03 RPL17-C18orf32
## check if gene name is unique - duplicates cannot move further in analysis
summary(duplicated(res.sex.gsea.dt[["gene_name"]])) # <- there are 49 duplicates here</pre>
##
      Mode
             FALSE
                      TRUE
             16020
                        49
## logical
## what do duplicates look like?
dup_gnames <- unique(res.sex.gsea.dt[duplicated(res.sex.gsea.dt[["gene_name"]])][["gene_name"]])</pre>
length(dup_gnames)
## [1] 28
res.sex.gsea.dt[gene_name %in% dup_gnames][order(gene_name)][1:6]
##
                                gene_id baseMean log2FoldChange
                                                                       lfcSE
## 1:
           ENSG00000277739.1|5 8S rRNA 104.18887
                                                     -0.73001360 0.18702955
           ENSG00000275757.1|5_8S_rRNA 65.69818
## 2:
                                                     -0.74176688 0.18618235
           ENSG00000273730.1|5_8S_rRNA 63.50881
                                                     -0.77845299 0.19944816
## 4: ENSG00000197976.12_PAR_Y|AKAP17A 652.72880
                                                      0.05233269 0.05196628
## 5:
            ENSG00000197976.12|AKAP17A 638.01924
                                                      0.04448442 0.05398411
            ENSG00000231259.5 | ANAPC1P2 98.47333
                                                      0.18787592 0.45798714
## 6:
##
                       pvalue
                                      padj gene_name
            stat
## 1: -3.9031992 9.492952e-05 0.002128958 5 8S rRNA
## 2: -3.9840881 6.773975e-05 0.001644275 5_8S_rRNA
## 3: -3.9030342 9.499426e-05 0.002128958 5 8S rRNA
## 4: 1.0070509 3.139103e-01 0.515453149
                                             AKAP17A
## 5: 0.8240279 4.099237e-01 0.608111520
                                             AKAP17A
## 6: 0.4102210 6.816439e-01 0.814919674 ANAPC1P2
## Select the genes to use for GSEA
## Sets cutoff of significance
res_cutoff <- 0.05
```

```
## Check:Since there are more than few hundred sig. genes by padj use those
nrow(res.sex.gsea.dt[padj<res_cutoff])</pre>
## [1] 2860
## Check: Are any of 2860 duplicated? <- YES some are
res.sex.gsea.dt[padj<res_cutoff][duplicated(res.sex.gsea.dt[padj<res_cutoff][["gene_name"]])]
##
                                          baseMean log2FoldChange
                                gene_id
                                                                        1fcSE
##
   1:
               ENSG00000182162.11|P2RY8 1365.00269
                                                        0.2167833 0.04963399
##
   2: ENSG00000169100.14_PAR_Y|SLC25A6 1191.44637
                                                         0.1981591 0.06998477
##
                   ENSG00000276596.1|U2
                                        417.54855
                                                        -0.6324921 0.17393389
                   ENSG00000273709.1|U2 1095.03288
##
  4:
                                                        -0.6435656 0.21330581
                                         413.31202
##
   5:
                   ENSG00000277903.1|U2
                                                        -0.6443804 0.20056111
                   ENSG00000278774.1|U2
##
   6:
                                         480.69558
                                                        -0.6516084 0.17378676
                   ENSG00000274062.1|U2
                                                        -0.6577563 0.17479295
##
   7:
                                         417.40218
                                                        -0.6651420 0.17594991
##
   8:
                   ENSG00000275219.1|U2
                                         419.36283
   9:
                   ENSG00000274862.1|U2
                                                        -0.6855793 0.17461317
##
                                         442.21074
## 10:
                                                        -0.6930374 0.17706424
                   ENSG00000278591.1|U2 439.43502
                                                        -0.7116059 0.18416220
## 11:
                   ENSG00000274452.1|U2
                                         404.24922
## 12:
            ENSG00000275757.1|5 8S rRNA
                                           65.69818
                                                        -0.7417669 0.18618235
## 13:
            ENSG00000273730.1|5_8S_rRNA
                                           63.50881
                                                        -0.7784530 0.19944816
                                      padj gene_name
##
                       pvalue
##
       4.367637 1.255978e-05 0.0004693561
                                               P2RY8
   1:
                                              SLC25A6
##
       2.831460 4.633597e-03 0.0322325816
   3: -3.636393 2.764820e-04 0.0046039261
                                                   112
                                                   U2
  4: -3.017103 2.552029e-03 0.0215834504
## 5: -3.212888 1.314074e-03 0.0137742082
                                                   U2
   6: -3.749471 1.772080e-04 0.0033755343
                                                   U2
   7: -3.763060 1.678470e-04 0.0032487899
                                                   U2
  8: -3.780292 1.566447e-04 0.0031114009
                                                   U2
## 9: -3.926275 8.627145e-05 0.0019946705
                                                   U2
## 10: -3.914045 9.076270e-05 0.0020687459
                                                   U2
## 11: -3.864017 1.115374e-04 0.0023984195
                                                   U2
## 12: -3.984088 6.773975e-05 0.0016442749 5 8S rRNA
## 13: -3.903034 9.499426e-05 0.0021289578 5_8S_rRNA
res.sex.gsea.dt[padj<res_cutoff][gene_name %in% "P2RY8"]
##
                             gene_id baseMean log2FoldChange
                                                                   lfcSE
                                                                              stat
## 1: ENSG00000182162.11 PAR Y|P2RY8 1413.283
                                                   0.2170988 0.04945298 4.390005
            ENSG00000182162.11|P2RY8 1365.003
                                                    0.2167833 0.04963399 4.367637
## 2:
            pvalue
                           padj gene name
## 1: 1.133479e-05 0.0004275556
                                    P2RY8
## 2: 1.255978e-05 0.0004693561
                                    P2RY8
## Remove genes with _PAR_Y
res.sex.gsea.dt <- res.sex.gsea.dt[grep("_PAR_Y",gene_id, invert = TRUE)]
## Check: How do our duplicates look now?.... there are fewer
res.sex.gsea.dt[padj<res_cutoff][duplicated(res.sex.gsea.dt[padj<res_cutoff][["gene_name"]])]
##
                                     baseMean log2FoldChange
                           gene_id
                                                                  lfcSE
                                                                              stat
##
   1:
              ENSG00000276596.1|U2
                                   417.54855
                                                   -0.6324921 0.1739339 -3.636393
##
   2:
              ENSG00000273709.1|U2 1095.03288
                                                   -0.6435656 0.2133058 -3.017103
              ENSG00000277903.1|U2 413.31202
                                                   -0.6443804 0.2005611 -3.212888
```

```
##
              ENSG00000278774.1|U2 480.69558
                                                   -0.6516084 0.1737868 -3.749471
##
              ENSG00000274062.1|U2 417.40218
                                                   -0.6577563 0.1747929 -3.763060
  5:
              ENSG00000275219.1|U2 419.36283
                                                   -0.6651420 0.1759499 -3.780292
##
  6:
  7:
              ENSG00000274862.1|U2 442.21074
                                                   -0.6855793 0.1746132 -3.926275
##
##
   8:
              ENSG00000278591.1|U2
                                    439.43502
                                                   -0.6930374 0.1770642 -3.914045
  9:
              ENSG00000274452.1|U2 404.24922
                                                   -0.7116059 0.1841622 -3.864017
##
## 10: ENSG00000275757.1|5 8S rRNA
                                                   -0.7417669 0.1861824 -3.984088
                                     65.69818
                                                   -0.7784530 0.1994482 -3.903034
## 11: ENSG00000273730.1|5_8S_rRNA
                                     63.50881
##
             pvalue
                           padj gene_name
  1: 2.764820e-04 0.004603926
##
                                       U2
## 2: 2.552029e-03 0.021583450
                                       U2
                                       U2
## 3: 1.314074e-03 0.013774208
## 4: 1.772080e-04 0.003375534
                                       U2
## 5: 1.678470e-04 0.003248790
                                       U2
## 6: 1.566447e-04 0.003111401
                                       U2
## 7: 8.627145e-05 0.001994670
                                       U2
## 8: 9.076270e-05 0.002068746
                                       U2
## 9: 1.115374e-04 0.002398419
                                       U2
## 10: 6.773975e-05 0.001644275 5_8S_rRNA
## 11: 9.499426e-05 0.002128958 5_8S_rRNA
## Keep only unique gene names
res.sex.gsea.dt <- res.sex.gsea.dt[!duplicated(gene_name)]</pre>
## Extract log2FoldChange for these genes into a vector
geneVec <- res.sex.gsea.dt[padj <= res_cutoff][["log2FoldChange"]]</pre>
## Add names for each gene to the vector
names(geneVec) <- res.sex.gsea.dt[padj <= res_cutoff][["gene_name"]]</pre>
## Check if vector names are duplicated
summary(duplicated(names(geneVec))) ## NO MORE duplications
##
      Mode
             FALSE
## logical
              2844
##
## DO THIS IF YOU DID NOT FILTER BEFORE
## If above is yes: Find the duplicates and their names
#geneVec[duplicated(names(geneVec))]
#dup_names <- names(geneVec[duplicated(names(geneVec))])</pre>
#geneVec[names(geneVec) %in% dup names]
## For now remove duplicates BUT - work to avoid having duplicate gene names
#geneVec <- geneVec[!duplicated(names(geneVec))]</pre>
##
##
## gene vector to proceed with GSEA analysis
geneVec[1:10]
## ENSG00000269693 ENSG00000289273 ENSG00000289172
                                                              ETF31.
                                                                              KLRB1
                                                                          0.5488337
```

0.5757175

0.5752445

##

24.8272814

1.0517758

```
##
         SIGLEC17P
                            RPL7P9
                                               LDHB
                                                               CRIP2
                                                                                CCN3
##
         0.5299390
                         0.5253123
                                          0.5226034
                                                          0.5215145
                                                                           0.5213166
## Use the msigdbr package to load gene lists to compare with
##
#?msigdbr
#msigdbr(species = "Homo sapiens", category = "C7") # <- other gene sets
msig.H.dt <- as.data.table(msigdbr(species = "Homo sapiens", category = "H"))</pre>
msig.H.dt[1:5,]
##
      gs_cat gs_subcat
                                      gs_name gene_symbol entrez_gene
## 1:
           Η
                       HALLMARK_ADIPOGENESIS
                                                    ABCA1
## 2:
                                                    ABCB8
                                                                 11194
           Η
                       HALLMARK_ADIPOGENESIS
## 3:
                                                                 10449
           Η
                       HALLMARK_ADIPOGENESIS
                                                    ACAA2
## 4:
           Η
                                                                    33
                       HALLMARK_ADIPOGENESIS
                                                    ACADL
## 5:
                       HALLMARK_ADIPOGENESIS
                                                    ACADM
                                                                    34
         ensembl_gene human_gene_symbol human_entrez_gene human_ensembl_gene gs_id
## 1: ENSG00000165029
                                   ABCA1
                                                        19
                                                              ENSG00000165029 M5905
                                                     11194
## 2: ENSG00000197150
                                   ABCB8
                                                              ENSG00000197150 M5905
## 3: ENSG00000167315
                                   ACAA2
                                                     10449
                                                              ENSG00000167315 M5905
## 4: ENSG00000115361
                                                              ENSG00000115361 M5905
                                   ACADL
                                                        33
## 5: ENSG00000117054
                                   ACADM
                                                        34
                                                              ENSG00000117054 M5905
       gs_pmid gs_geoid gs_exact_source gs_url
## 1: 26771021
## 2: 26771021
## 3: 26771021
## 4: 26771021
## 5: 26771021
##
                                                            gs_description
## 1: Genes up-regulated during adipocyte differentiation (adipogenesis).
## 2: Genes up-regulated during adipocyte differentiation (adipogenesis).
## 3: Genes up-regulated during adipocyte differentiation (adipogenesis).
## 4: Genes up-regulated during adipocyte differentiation (adipogenesis).
## 5: Genes up-regulated during adipocyte differentiation (adipogenesis).
## View how many genes are involved in each category
msig.H.dt[,.N,by = gs name]
##
                                           gs_name
##
  1:
                            HALLMARK_ADIPOGENESIS 210
##
  2:
                     HALLMARK_ALLOGRAFT_REJECTION 335
                       HALLMARK_ANDROGEN_RESPONSE 102
## 4:
                            HALLMARK_ANGIOGENESIS
## 5:
                         HALLMARK_APICAL_JUNCTION 231
## 6:
                          HALLMARK_APICAL_SURFACE 46
## 7:
                               HALLMARK_APOPTOSIS 183
## 8:
                    HALLMARK_BILE_ACID_METABOLISM 114
## 9:
                 HALLMARK_CHOLESTEROL_HOMEOSTASIS 77
## 10:
                             HALLMARK COAGULATION 162
## 11:
                              HALLMARK_COMPLEMENT 237
## 12:
                              HALLMARK DNA REPAIR 170
## 13:
                             HALLMARK_E2F_TARGETS 218
## 14: HALLMARK_EPITHELIAL_MESENCHYMAL_TRANSITION 204
```

```
## 15:
                 HALLMARK ESTROGEN RESPONSE EARLY 216
## 16:
                  HALLMARK ESTROGEN RESPONSE LATE 218
## 17:
                  HALLMARK FATTY ACID METABOLISM 165
## 18:
                           HALLMARK_G2M_CHECKPOINT 204
## 19:
                               HALLMARK GLYCOLYSIS 215
## 20:
                      HALLMARK HEDGEHOG SIGNALING 36
## 21:
                          HALLMARK HEME METABOLISM 214
## 22:
                                  HALLMARK HYPOXIA 215
## 23:
                     HALLMARK_IL2_STAT5_SIGNALING 216
## 24:
                 HALLMARK_IL6_JAK_STAT3_SIGNALING 103
## 25:
                   HALLMARK_INFLAMMATORY_RESPONSE 222
## 26:
               HALLMARK_INTERFERON_ALPHA_RESPONSE 140
## 27:
               HALLMARK_INTERFERON_GAMMA_RESPONSE 286
## 28:
                       HALLMARK_KRAS_SIGNALING_DN 220
## 29:
                        HALLMARK_KRAS_SIGNALING_UP 220
## 30:
                         HALLMARK_MITOTIC_SPINDLE 215
## 31:
                        HALLMARK_MTORC1_SIGNALING 211
## 32:
                           HALLMARK MYC TARGETS V1 236
## 33:
                           HALLMARK_MYC_TARGETS_V2 60
## 34:
                               HALLMARK MYOGENESIS 212
## 35:
                         HALLMARK_NOTCH_SIGNALING
## 36:
               HALLMARK OXIDATIVE PHOSPHORYLATION 220
## 37:
                              HALLMARK_P53_PATHWAY 215
## 38:
                     HALLMARK PANCREAS BETA CELLS
## 39:
                               HALLMARK PEROXISOME 110
## 40:
                 HALLMARK PI3K AKT MTOR SIGNALING 118
## 41:
                        HALLMARK_PROTEIN_SECRETION
## 42:
         HALLMARK_REACTIVE_OXYGEN_SPECIES_PATHWAY
                                                    58
## 43:
                          HALLMARK_SPERMATOGENESIS 144
## 44:
                      HALLMARK_TGF_BETA_SIGNALING
                 HALLMARK_TNFA_SIGNALING_VIA_NFKB 228
## 45:
## 46:
               HALLMARK_UNFOLDED_PROTEIN_RESPONSE 115
## 47:
                           HALLMARK_UV_RESPONSE_DN 152
## 48:
                           HALLMARK_UV_RESPONSE_UP 191
## 49:
              HALLMARK WNT BETA CATENIN SIGNALING
## 50:
                   HALLMARK_XENOBIOTIC_METABOLISM 224
                                           gs name
## Extract only gene_name and pathway name columns
msig.H.t2g <- msig.H.dt[,.SD,.SDcols = c("gs_name", "gene_symbol")]</pre>
msig.H.t2g
##
                                 gs_name gene_symbol
##
                  HALLMARK_ADIPOGENESIS
                                               ABCA1
##
      2:
                  HALLMARK_ADIPOGENESIS
                                                ABCB8
##
                  HALLMARK ADIPOGENESIS
                                                ACAA2
##
      4:
                  HALLMARK_ADIPOGENESIS
                                               ACADL
##
                  HALLMARK_ADIPOGENESIS
                                               ACADM
##
## 8205: HALLMARK_XENOBIOTIC_METABOLISM
                                                UPB1
## 8206: HALLMARK_XENOBIOTIC_METABOLISM
                                                UPP1
## 8207: HALLMARK XENOBIOTIC METABOLISM
                                                VNN1
## 8208: HALLMARK XENOBIOTIC METABOLISM
                                                 VTN
## 8209: HALLMARK XENOBIOTIC METABOLISM
                                                 XDH
```

```
## Run GSEA
## => Much of the options can found online in the ClusterProfiler manual
?GSEA
agsea <- clusterProfiler::GSEA(geneList = geneVec,</pre>
                                TERM2GENE = msig.H.t2g,
                                minGSSize = 5, # minimum number of genes to match pathway
                                pvalueCutoff = 1, # this way all pathways are returned
                                pAdjustMethod = "BH", # many other methods are out there
                                by = 'fgsea',
                                seed = TRUE)
## preparing geneSet collections...
## GSEA analysis...
## leading edge analysis...
## done...
agsea.dt <- as.data.table(x = agsea)</pre>
## Clean up and organize the results table
agsea.dt <- agsea.dt[order(p.adjust)]</pre>
agsea.dt[,"sig" := p.adjust<=0.05]
agsea.dt[,"updown" := ifelse(NES<0,"down","up")]</pre>
agsea.dt[["ID"]] <- gsub("HALLMARK_","",agsea.dt[["ID"]])</pre>
agsea.dt[["ID"]] <- tolower(gsub("_"," ",agsea.dt[["ID"]]))</pre>
agsea.dt[["ID"]] <- factor(agsea.dt[["ID"]], levels = agsea.dt[["ID"]])</pre>
agsea.dt[["Description"]] <- NULL</pre>
agsea.dt[1:4]
##
                              ID setSize enrichmentScore
                                                                NES
                                                                           pvalue
## 1: interferon alpha response
                                      54
                                              -0.6742649 -2.032744 4.775842e-09
## 2: interferon gamma response
                                      88
                                              -0.6115491 -1.917812 4.273297e-09
## 3:
          inflammatory response
                                      64
                                              -0.4748911 -1.451781 1.144414e-02
## 4:
                                       7
                                              -0.7125717 -1.550740 1.829138e-02
            pancreas beta cells
##
          p.adjust
                         qvalue rank
                                                         leading_edge
## 1: 1.146202e-07 1.080849e-07 619 tags=70%, list=22%, signal=56%
## 2: 1.146202e-07 1.080849e-07 619 tags=57%, list=22%, signal=46%
## 3: 1.831063e-01 1.726660e-01 850 tags=53%, list=30%, signal=38%
## 4: 2.194965e-01 2.069814e-01 376 tags=57%, list=13%, signal=50%
##
## 1:
                                                                              SAMD9L/UBE2L6/NUB1/HELZ2/IFI
## 2: SAMD9L/C1R/UBE2L6/VAMP5/MX2/HELZ2/IFI30/TOR1B/LAP3/CDKN1A/TRAFD1/SOCS3/EIF2AK2/MX1/IFIT3/PLSCR1/F
## 3:
## 4:
##
        sig updown
## 1: TRUE
              down
## 2: TRUE
              down
## 3: FALSE
              down
## 4: FALSE
              down
```

```
## FINAL RESULT
agsea.dt[1:4]
##
                             ID setSize enrichmentScore
                                                               NES
                                                                          pvalue
## 1: interferon alpha response
                                     54
                                              -0.6742649 -2.032744 4.775842e-09
## 2: interferon gamma response
                                      88
                                              -0.6115491 -1.917812 4.273297e-09
## 3:
          inflammatory response
                                      64
                                              -0.4748911 -1.451781 1.144414e-02
## 4:
                                       7
                                              -0.7125717 -1.550740 1.829138e-02
            pancreas beta cells
##
          p.adjust
                         qvalue rank
                                                        leading_edge
## 1: 1.146202e-07 1.080849e-07
                                 619 tags=70%, list=22%, signal=56%
## 2: 1.146202e-07 1.080849e-07
                                 619 tags=57%, list=22%, signal=46%
## 3: 1.831063e-01 1.726660e-01
                                 850 tags=53%, list=30%, signal=38%
## 4: 2.194965e-01 2.069814e-01
                                 376 tags=57%, list=13%, signal=50%
##
## 1:
                                                                             SAMD9L/UBE2L6/NUB1/HELZ2/IFI
## 2: SAMD9L/C1R/UBE2L6/VAMP5/MX2/HELZ2/IFI30/TOR1B/LAP3/CDKN1A/TRAFD1/SOCS3/EIF2AK2/MX1/IFIT3/PLSCR1/F
## 3:
## 4:
##
        sig updown
## 1:
      TRUE
              down
## 2: TRUE
              down
## 3: FALSE
              down
## 4: FALSE
              down
```

PART 3

Input data

##

Goal: - Explore ways of plotting results from GSEA and DESeq - Other ways of visualizing data

1. Barplot of up and down-regulated pathways

- This plot highlights the pathways that have increased or decreased based on the significance cut-offs
- It is sometimes helpful to plot the pathways that are not part of the significance

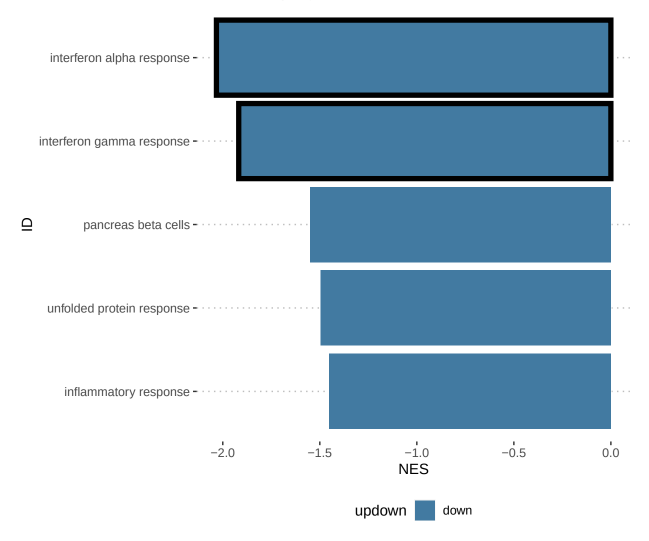
```
agsea.dt[1:5,]
##
                             ID setSize enrichmentScore
                                                               NES
                                                                         pvalue
## 1: interferon alpha response
                                     54
                                             -0.6742649 -2.032744 4.775842e-09
                                     88
                                             -0.6115491 -1.917812 4.273297e-09
## 2: interferon gamma response
## 3:
          inflammatory response
                                     64
                                              -0.4748911 -1.451781 1.144414e-02
## 4:
                                      7
                                              -0.7125717 -1.550740 1.829138e-02
            pancreas beta cells
## 5: unfolded protein response
                                     14
                                             -0.5841944 -1.495397 4.638619e-02
##
          p.adjust
                         qvalue rank
                                                        leading_edge
## 1: 1.146202e-07 1.080849e-07
                                 619 tags=70%, list=22%, signal=56%
## 2: 1.146202e-07 1.080849e-07
                                 619 tags=57%, list=22%, signal=46%
                                 850 tags=53%, list=30%, signal=38%
## 3: 1.831063e-01 1.726660e-01
                                 376 tags=57%, list=13%, signal=50%
## 4: 2.194965e-01 2.069814e-01
## 5: 4.453074e-01 4.199171e-01
```

SAMD9L/UBE2L6/NUB1/HELZ2/IFI ## 2: SAMD9L/C1R/UBE2L6/VAMP5/MX2/HELZ2/IFI30/TOR1B/LAP3/CDKN1A/TRAFD1/SOCS3/EIF2AK2/MX1/IFIT3/PLSCR1/F

19 tags=14%, list=1%, signal=14%

```
## 3:
## 4:
## 5:
##
        sig updown
## 1: TRUE
## 2: TRUE
              down
## 3: FALSE
              down
## 4: FALSE
              down
## 5: FALSE
              down
## Reorder to have NES reveresed
agsea.dt <- agsea.dt[order(-NES)]</pre>
## Change order of labels by adjusting the factor levels
class(agsea.dt[["ID"]])
## [1] "factor"
agsea.dt[["ID"]] <- factor(agsea.dt[["ID"]], levels = agsea.dt[["ID"]])</pre>
## Plot - subset
ggp.d28.gset.bar <- ggplot() + theme_pubclean() +</pre>
  geom_bar(data = agsea.dt[pvalue <= 0.05],</pre>
           aes(x = NES, y = ID, fill = updown),
           stat = "identity") +
   geom_bar(data = agsea.dt[p.adjust <= 0.05], ## SHOW SIGNIFICANT</pre>
            aes(x = NES, y = ID),
            colour = "black", lwd = 2,
            fill=NA,
           stat = "identity") +
  scale fill manual(values = c("#427AA1", "#CE8D99")) +
  ggtitle("Black box: padj <= 0.05") +</pre>
  theme(aspect.ratio =1,
        legend.position = "bottom",
        axis.text.y = element_text()); ggp.d28.gset.bar
```

Black box: padj <= 0.05

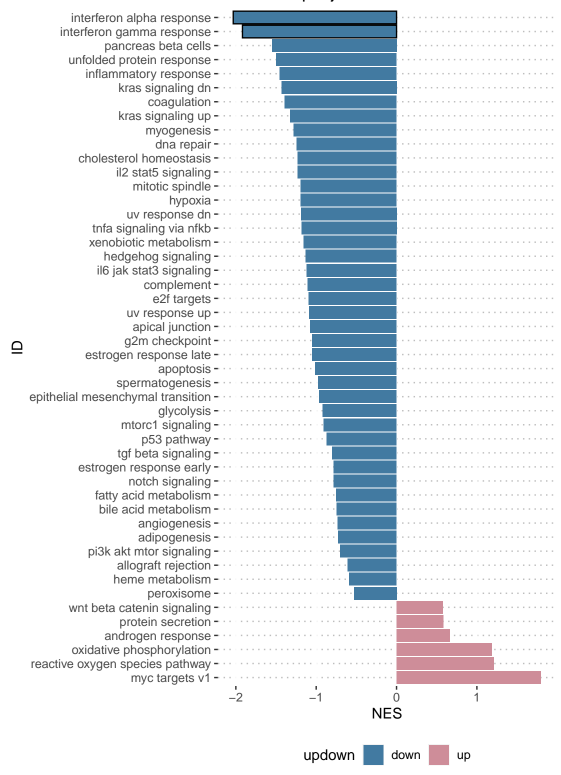


```
## Plot - all
ggp.d28.gset.bar <- ggplot() + theme_pubclean() +
geom_bar(data = agsea.dt, ## ALL PATHWAYS</pre>
```

```
aes(x = NES, y = ID, fill = updown),
    stat = "identity") +
geom_bar(data = agsea.dt[p.adjust <= 0.05], ## SHOW SIGNIFICANT
    aes(x = NES, y = ID),
    colour = "black", lwd = 0.51,
    fill=NA,
    stat = "identity") +
scale_fill_manual(values = c("#427AA1","#CE8D99")) +
ggtitle("Black box: padj <= 0.05") +

theme(aspect.ratio = 2,
    legend.position = "bottom",
    axis.text.y = element_text()); ggp.d28.gset.bar</pre>
```

Black box: padj <= 0.05



2. Leading edge plots - ClusterProfiler

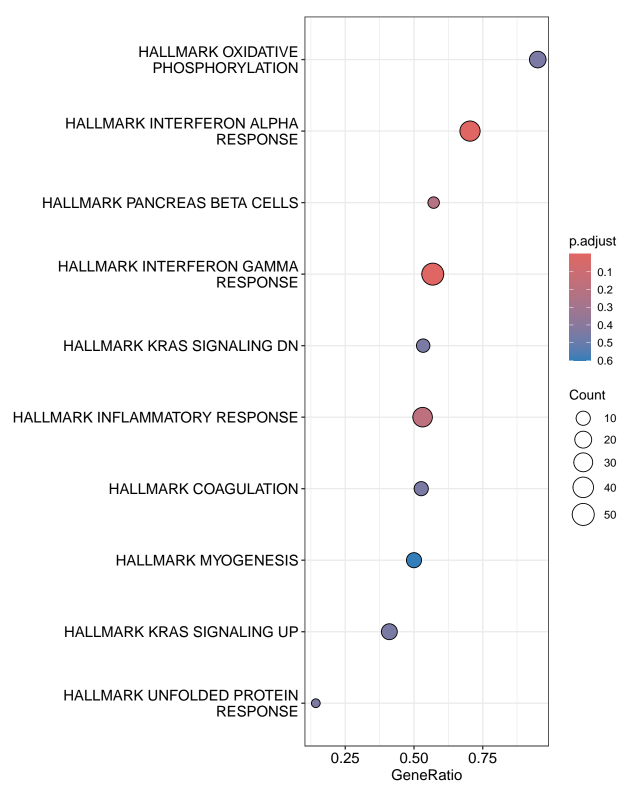
• These plots are helpful to demonstrate the distribution of the genes within the differential spectrum

- Secondly, it indirectly shows where are the genes responsible for the pathway direction located and how they contribute to enrichment
- Take your time to explore different plots in the book online and playig with them to make them look clear in R (most of the are ggplot objects)
 - See the book: https://yulab-smu.top/biomedical-knowledge-mining-book/enrichplot.html

```
## Input data
#agsea

##
## Below plots are all parth of the ClusterProfiler package
##

## Dotplot
dotplot(object = agsea)
```

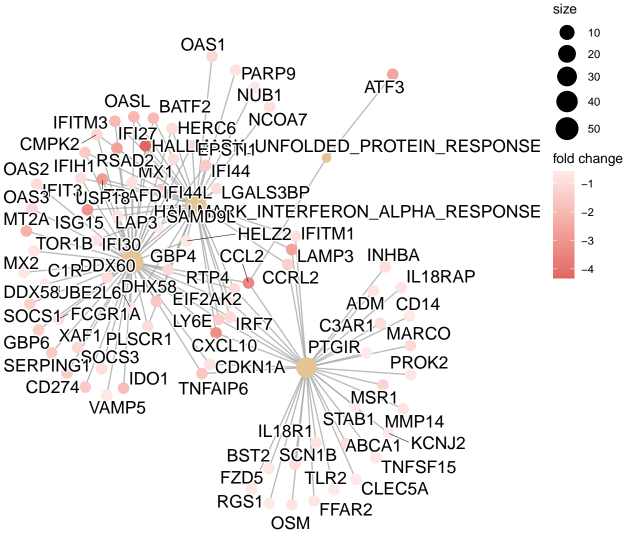


```
## Network plot
cnetplot(x = agsea, foldChange=geneVec, base.size =3 )
```

```
## Warning in cnetplot.enrichResult(x, ...): Use 'color.params = list(foldChange = your_value)' instead ## The foldChange parameter will be removed in the next version.
```

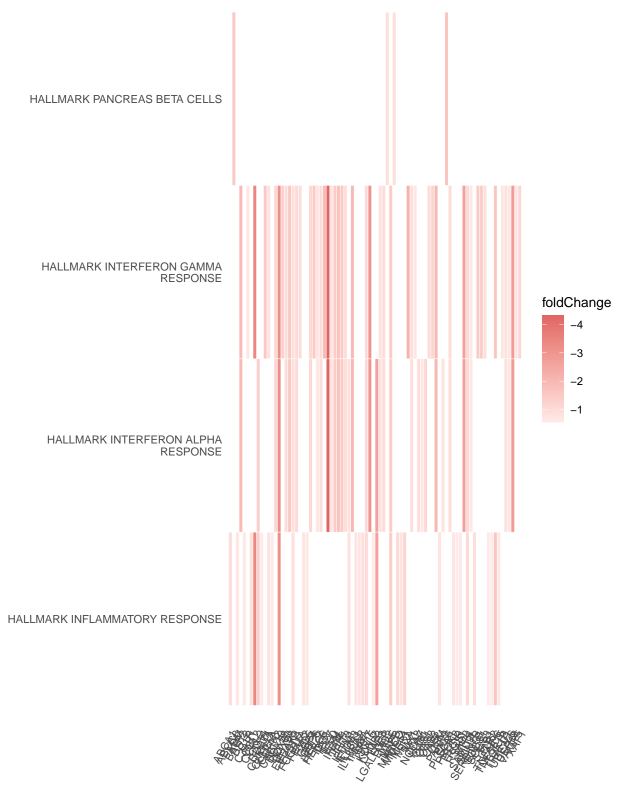
- ## Scale for size is already present.
- ## Adding another scale for size, which will replace the existing scale.
- ## Warning: ggrepel: 2 unlabeled data points (too many overlaps). Consider
- ## increasing max.overlaps

PAX6 LMO2 HALLMARK_PANCREAS_BETA_CELLS ABCC8



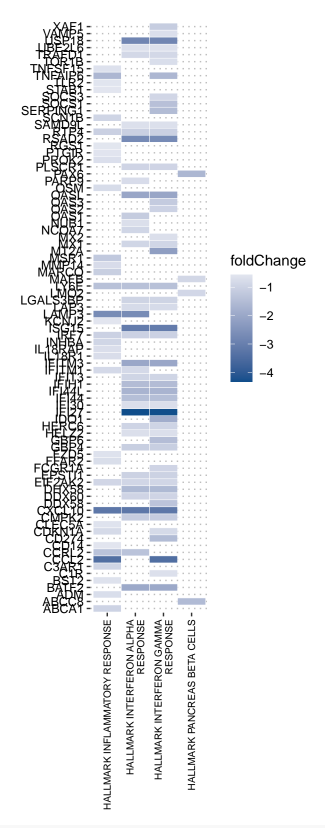
heatplot ## -> lets play with this one a little

```
##
## basic
heatplot(x = agsea, showCategory =4, foldChange = geneVec)
```



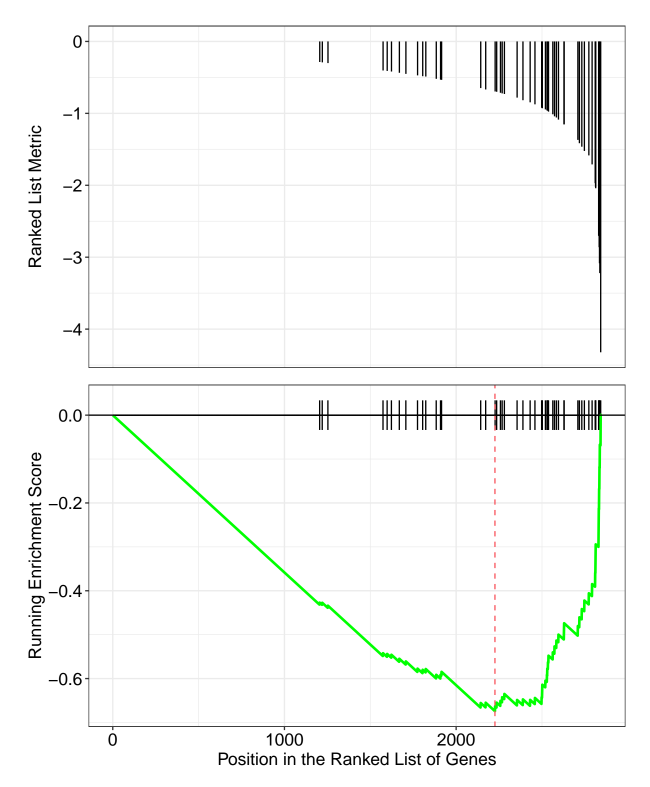
```
## Modified - This is a ggplot object
heatplot(x = agsea, showCategory =4, foldChange = geneVec) +
    theme_pubclean() +
```

Scale for fill is already present.
Adding another scale for fill, which will replace the existing scale.

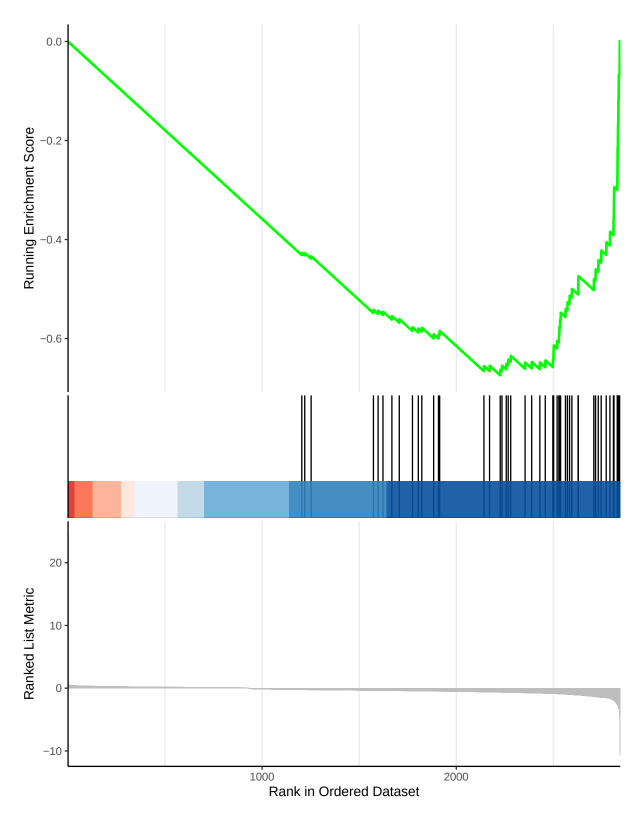


Plot using pre-built function
?gseaplot2

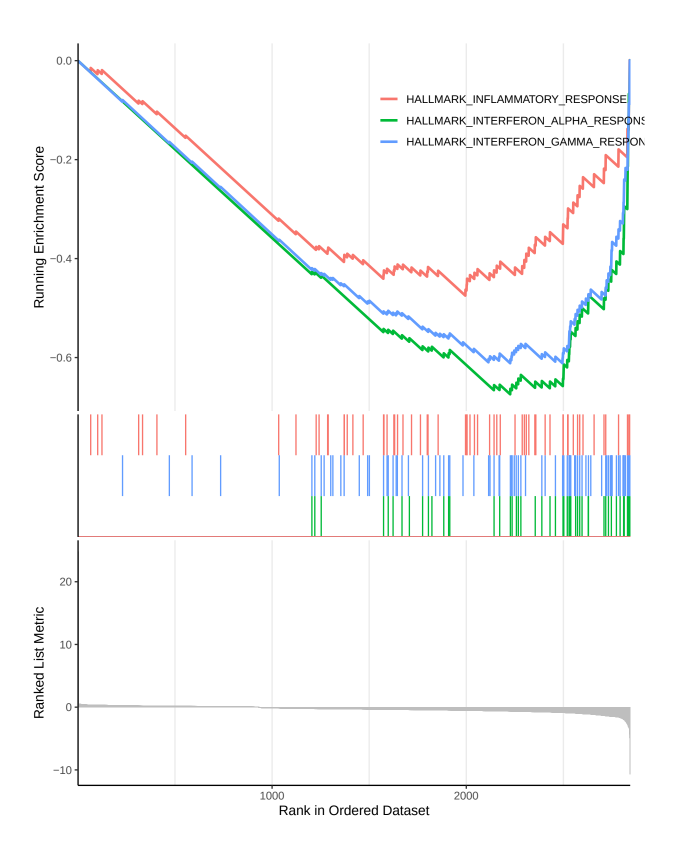
```
## Leading edge - simple
gseaplot(x = agsea, geneSetID = "HALLMARK_INTERFERON_ALPHA_RESPONSE")
```



Leading edge - more comlicated
gseaplot2(x = agsea, geneSetID = "HALLMARK_INTERFERON_ALPHA_RESPONSE")



```
## Leading edge - multiple gene sets
gseaplot2(x = agsea, geneSetID = 1:3)
```



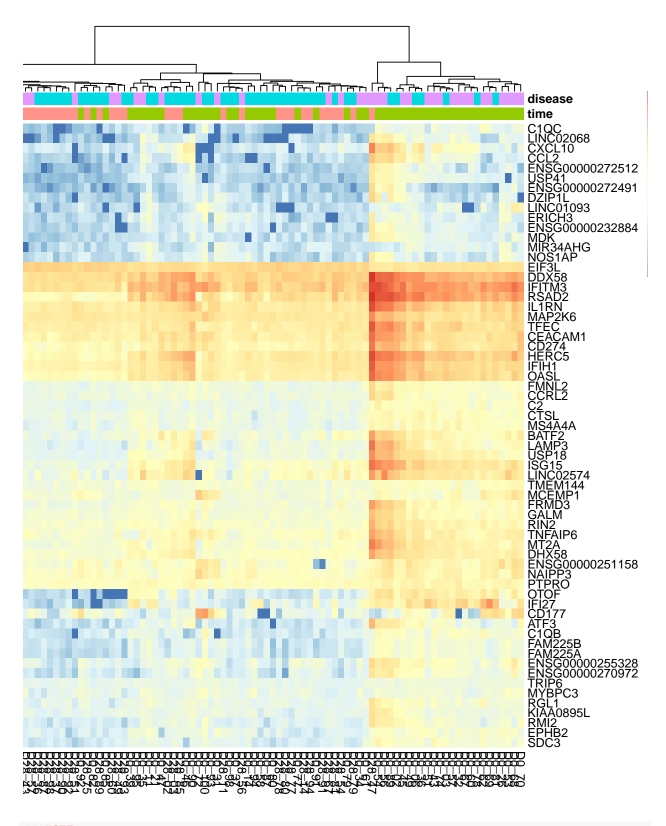
3. Clustering and heatmaps - Pheatmap

- The goal is to explore possible relationships between subjects using unsupervised clustering
- Pheatmap is a vewry useful package to visualize these relationships
- More information about using it can be found at: https://r-charts.com/correlation/pheatmap/

```
## library
library(pheatmap)
## Prepare data to compare
##
## data to plot
meta.clean.dt[1:3]
      IGU Code
                        pathogen disease time subject seq_id
                  sex
## 1:
         02 DO Female SARS-CoV-2 COVID19
                                                    02 D0 02
                                            D0
## 2:
        02 D28 Female SARS-CoV-2 COVID19
                                          D28
                                                    02 D28 02
## 3:
         03_D0 Female SARS-CoV-2 COVID19
                                            DO
                                                    03 D0_03
ncount.dt[1:3,1:3]
##
                                gene_id
                                            D0_02
                                                      D0_06
## 1:
              ENSG00000187961.15|KLHL17 148.5576 157.58938
## 2:
               ENSG00000188976.11|NOC2L 417.3295 460.64587
## 3: ENSG00000272512.1|ENSG00000272512 353.8016 55.76239
## to subset genes you can use results of DESeq - only significant ones
## a lot of genes can make the heatmap hard to read - select small set of relevant genes
res.sex.dt[1:3,]
##
                       gene id baseMean log2FoldChange
                                                            lfcSE
## 1: ENSG00000165949.13|IFI27 1174.004
                                           -4.321991 0.4443898 -9.725677
## 2: ENSG00000184979.11|USP18 830.048
                                             -2.860923 0.3157571 -9.060520
## 3: ENSG00000187608.10|ISG15 2014.351
                                             -3.084138 0.3431463 -8.987824
            pvalue
                           padj
## 1: 2.343417e-22 3.765638e-18
## 2: 1.298298e-19 1.043118e-15
## 3: 2.521727e-19 1.350721e-15
res.sex.topsig.dt <- res.sex.dt[padj < 0.0000001][baseMean > 50]
res.sex.topsig.names <- res.sex.topsig.dt[["gene_id"]]</pre>
## subset the normalized count table to your genes of interest
ncount.subset.dt <- ncount.dt[gene_id %in% res.sex.topsig.names]</pre>
ncount.subset.dt[1:5,1:5]
##
                                               D0 02
                                                          DO 06
                                                                   DO 100
                                gene_id
                                                                              DO 11
## 1: ENSG00000272512.1 | ENSG00000272512
                                           353.80161
                                                       55.76239
                                                                14.97167 45.37886
               ENSG00000187608.10|ISG15 11441.86601 5696.24984 262.00416 567.23573
## 2:
## 3:
             ENSG00000228526.8|MIR34AHG
                                            39.09410 187.89503 194.63166 336.55987
## 4:
               ENSG00000159189.13 | C1QC
                                            98.71261 163.65051 319.39555 102.10243
## 5:
                ENSG00000173369.18 | C1QB
                                           586.41151 269.11417 776.03137 113.44715
```

```
## cleanup the gene names
ncount.subset.dt[,"gene_id":=tstrsplit(gene_id,split="\\|",keep = 2)]
ncount.subset.dt[1:5,1:5]
              gene_id
                            D0 02
                                        DO 06
##
                                                 DO 100
                                                             DO 11
## 1: ENSG00000272512
                        353.80161
                                     55.76239 14.97167 45.37886
## 2:
                ISG15 11441.86601 5696.24984 262.00416 567.23573
             MIR34AHG
                         39.09410 187.89503 194.63166 336.55987
## 4:
                 C1QC
                         98.71261 163.65051 319.39555 102.10243
## 5:
                 C1QB
                        586.41151 269.11417 776.03137 113.44715
##
## Plot with pheatmap
#library(pheatmap)
#?pheatmap
## coerce your data to matrix
class(ncount.subset.dt)
## [1] "data.table" "data.frame"
ncount.subset.mat <- as.matrix(x = ncount.subset.dt, rownames="gene_id")</pre>
## prepare column information to use for clustering
col info dt <- data.table("samples" = colnames(ncount.subset.mat))</pre>
col_info_dt[1:5]
##
      samples
## 1:
        D0_02
## 2:
        D0 06
## 3: DO 100
## 4:
        D0 11
## 5:
        DO 14
col.info.dt <- col_info_dt[meta.clean.dt,on=.(samples=seq_id)][,.SD,.SDcols=c("samples","time","disease</pre>
rownames(col.info.dt)
                     "3"
                            "4"
                                  "5"
                                        "6"
                                              "7"
                                                    "8"
                                                           "9"
                                                                 "10"
##
     [1] "1"
               "2"
                                                                       "11"
                                                                             "12"
                                        "18"
               "14"
                     "15"
                           "16"
                                  "17"
                                              "19"
                                                    "20"
                                                          "21"
                                                                "22"
                                                                       "23"
                                                                             "24"
##
   [13] "13"
##
  [25] "25"
               "26"
                     "27"
                           "28"
                                  "29"
                                        "30"
                                              "31"
                                                    "32"
                                                          "33"
                                                                "34"
                                                                       "35"
                                                                             "36"
                     "39"
                                        "42"
                                              "43"
                                                    "44"
                                                                 "46"
                                                                       "47"
                                                                             "48"
## [37] "37"
               "38"
                           "40"
                                  "41"
                                                          "45"
                                  "53"
                                              "55"
  [49] "49"
               "50"
                     "51"
                           "52"
                                        "54"
                                                    "56"
                                                           "57"
                                                                 "58"
                                                                       "59"
                                                                             "60"
##
                                        "66"
                                              "67"
                                                    "68"
                                                           "69" "70"
                                                                       "71"
                                                                             "72"
##
    [61] "61"
               "62"
                     "63"
                           "64"
                                  "65"
##
   [73] "73"
               "74"
                     "75"
                           "76"
                                  "77"
                                        "78"
                                              "79"
                                                    "80"
                                                           "81"
                                                                "82"
                                                                       "83"
                                                                             "84"
                                              "91"
                                                                             "96"
   [85] "85"
               "86"
                     "87"
                           "88"
                                 "89"
                                        "90"
                                                    "92"
                                                          "93"
                                                                 "94"
                                                                       "95"
   [97] "97"
               "98"
                     "99"
                           "100" "101" "102" "103" "104" "105" "106"
rownames(col.info.dt) <- col.info.dt[["samples"]]</pre>
col.info.dt[["samples"]] <- NULL</pre>
## log transform your data
## Plot
pheatmap(mat = log10(ncount.subset.mat+1),
         annotation col = col.info.dt,
```

```
cluster_rows = TRUE,
clustering_method = "ward.D2",
cluster_cols = TRUE,
clustering_distance_rows = "canberra",
cellwidth = 5, cellheight = 8,
border_color = NA)
```



```
\ensuremath{\mbox{\#\#}} NOTE \ensuremath{\mbox{\#\#}} to export the plot into file use following structure
```

```
#pdf(file = filename.pdf)
#pheatmap(...)
#dev.off()
```

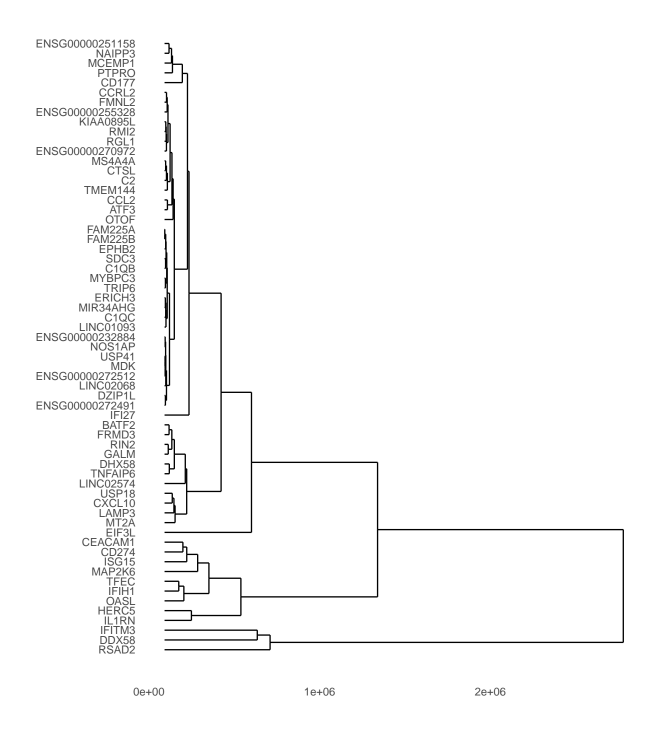
4. Clustering and heatmaps - Custom

- It is often usefull to have a full control of the clustering and plotting
- This scripts show how one can reproduce the pieces of the pheatmap plot using other R tools

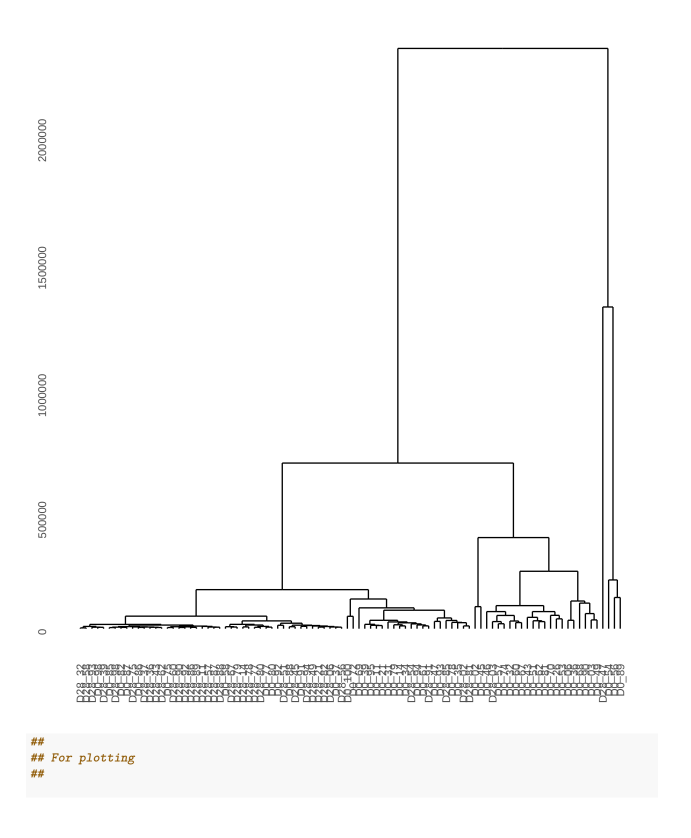
```
## Start with same pieces as with Pheatmap
ncount.subset.dt[1:5,1:5] # <- used to make ggplot</pre>
                             D0_02
                                        D0_06
                                                  D0_100
                                                             DO_11
              gene_id
## 1: ENSG00000272512
                         353.80161
                                     55.76239
                                               14.97167 45.37886
                ISG15 11441.86601 5696.24984 262.00416 567.23573
## 3:
                          39.09410 187.89503 194.63166 336.55987
             MIR34AHG
                          98.71261 163.65051 319.39555 102.10243
## 4:
                 C1QC
## 5:
                 C1QB
                         586.41151 269.11417 776.03137 113.44715
ncount.subset.mat[1:5,1:5] # <- matrix is used to cluster</pre>
                          D0 02
                                     DO 06
                                               DO 100
                                                          DO 11
                                                                     DO 14
## ENSG00000272512
                      353.80161
                                  55.76239 14.97167 45.37886 38.68685
## ISG15
                    11441.86601 5696.24984 262.00416 567.23573 567.40710
## MIR34AHG
                      39.09410 187.89503 194.63166 336.55987 54.16159
## C1QC
                      98.71261 163.65051 319.39555 102.10243 41.26597
## C1QB
                      586.41151 269.11417 776.03137 113.44715 121.21879
##
## Clustering
##
dist.meth <- "manhattan"</pre>
#dist.meth <- "eucledian"
clus.meth <- "ward.D2"</pre>
#clus.meth <- "ward.D"</pre>
## create a distance matrix
## - there are multiple ways of calculating this - see the help menu
?dist
row_dist <- dist(x = ncount.subset.mat, method = dist.meth)</pre>
col_dist <- dist(x = t(ncount.subset.mat), method = dist.meth)</pre>
## cluster the data
## - there are multiple ways of calculating this - see the help menu
?hclust
row_clust <- hclust(d = row_dist, method = clus.meth)</pre>
col_clust <- hclust(d = col_dist, method = clus.meth)</pre>
## extract the order for the data to be plotted
```

```
row_order_names <- row_clust$labels[row_clust$order]
col_order_names <- col_clust$labels[col_clust$order]

## plot dendogram - this is the way of the unsupervised clustering
row.dendo.ggp <- ggdendrogram(data = row_clust, rotate = 90)
row.dendo.ggp</pre>
```

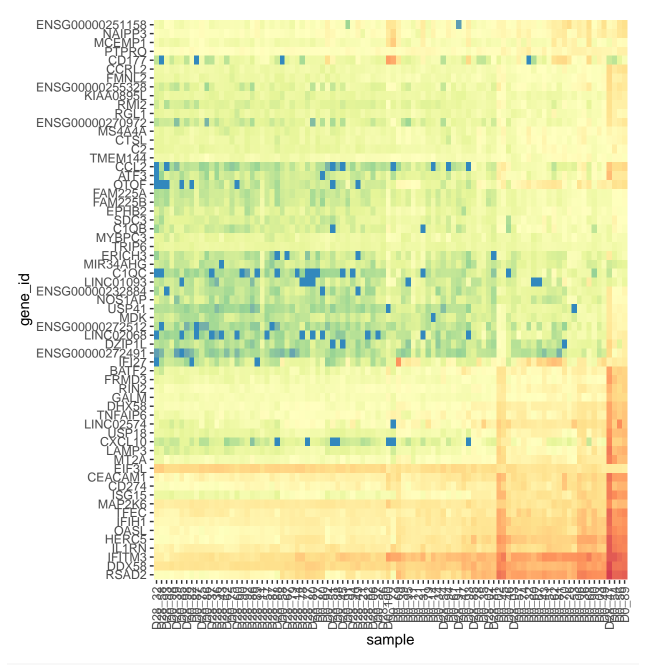


```
col.dendo.ggp <- ggdendrogram(data = col_clust)
col.dendo.ggp</pre>
```

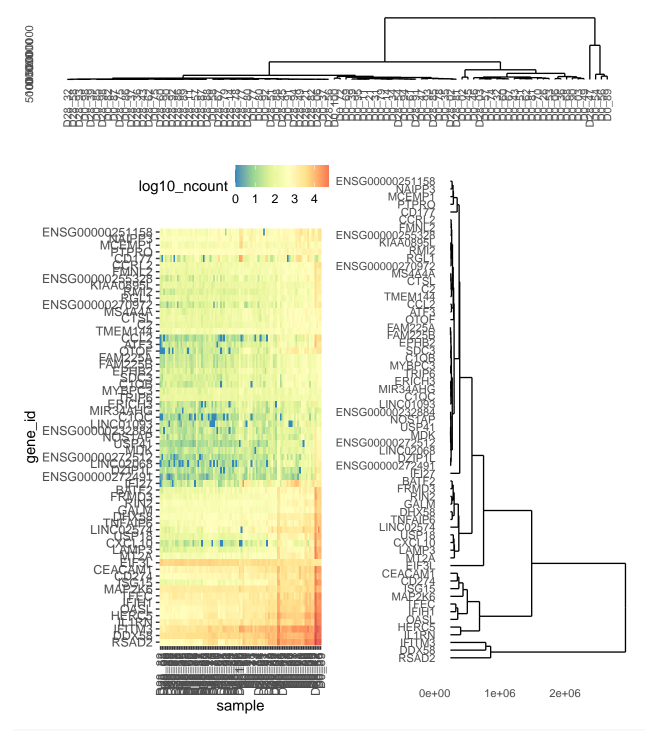


```
## melt into long format
ncount.subset.dtm <- melt.data.table(data = ncount.subset.dt,</pre>
                                      id.vars = "gene_id",
                                      variable.name = "sample",
                                      value.name = "ncount")
ncount.subset.dtm
##
                 gene_id sample
                                      ncount
                                  353.801614
##
      1: ENSG00000272512 D0_02
##
                   ISG15 D0_02 11441.866008
##
      3:
                MIR34AHG DO_02
                                   39.094101
                    C1QC D0_02
##
      4:
                                   98.712605
                    C1QB D0_02
##
      5:
                                 586.411515
##
     ___
## 5918:
                    RIN2 D28_98
                                  240.910490
## 5919: ENSG00000232884 D28_98
                                  18.250795
## 5920:
                   USP18 D28_98
                                   89.793910
## 5921:
                   USP41 D28 98
                                   6.570286
## 5922:
                   EIF3L D28_98 5088.321556
## log transform data
ncount.subset.dtm[,"log10 ncount":=log10(ncount+1)]
ncount.subset.dtm[,"sqrt_ncount":=sqrt(ncount+1)]
## give plot order
## - this is where the order we calculated above gets applied to data we want to plot
ncount.subset.dtm[["gene_id"]] <- factor(ncount.subset.dtm[["gene_id"]], levels = row_order_names)</pre>
ncount.subset.dtm[["sample"]] <- factor(ncount.subset.dtm[["sample"]], levels = col_order_names)</pre>
ggp.hm <- ggplot() + theme_pubclean() +</pre>
  geom_tile(data = ncount.subset.dtm,
            aes(x = gene_id, y = sample, fill = log10_ncount)) +
  scale_fill_distiller(palette = "Spectral") +
  coord_flip() +
  theme(axis.text.x = element_text(angle = 90, vjust = 0.5, hjust = 1)); ggp.hm
```





```
## arrange different plots together
arr1 <- ggarrange(plotlist = list(ggp.hm, row.dendo.ggp))
arr2 <- ggarrange(plotlist = list(col.dendo.ggp, arr1), nrow = 2, heights = c(1,4))
arr2</pre>
```



the plot can be saved with ggsave #ggsave(...)

NEXT

PART 4

1. Calculating statistical differences

• Comparison of gene expressions

```
## Goal compare

#ncount.dt
#meta.dt
```

2. Preparing a "Table 1 summary"

#meta.dt