PAPER TITLE TO BE DEFINED (in common.yaml)

0-Microarray data analysis

$2021 - 11 - 15 \ 10 : 23 : 33 \ + 0100$

Abstract

Lung interstitium macrophages (IMs) are non-alveolar resident tissue macrophages which contribute to the lung homeostasis. These cells were reported to be heterogeneous by our group and other teams, which contains two main distinct subpopulations: CD206+ IMs and CD206- IMs. However, the exact origin of IMs and the transcriptional programs that control IM differentiation remains unclear. In recent report, we analyzed the refilled IMs in the course of time after induced IM depletion with single-cell RNA sequencing (10X Genomics Chromium) and bulk RNA sequencing.

Contents

1	Description	2
2	Load packages and data	2
3	Data preparration3.1 Make a list with genes to show in heatmap3.2 Make metadata table	3 3
4	Make heatmaps	6
5	Session information	12
6	References	12

1 Description

2 Load packages and data

```
# packages
library(ComplexHeatmap)
                                                                                 3
library(RColorBrewer)
library(circlize)
                                                                                 4
                                                                                 5
                                                                                 6
# data
dir.files <- "./downloaded_data"</pre>
files.list <- list.files(dir.files, pattern = "Gene_Expression_Activity.
   csv", full.names = T)
names.list <- sub(basename(files.list), pattern = "_in_Gene_Expression_</pre>
                                                                                 9
   Activity.csv",
    replacement = "")
                                                                                 10
```

read csv files and bind tables to one:

```
expr.table <- data.frame(read.csv(files.list[1]), row.names = 1)</pre>
expr.table <- expr.table[, -2]
                                                                                     2
                                                                                     3
n.rep <- length(2:ncol(expr.table))</pre>
                                                                                     4
names.rep <- paste(rep(names.list[1]), 1:n.rep, sep = "_")</pre>
                                                                                     5
                                                                                     6
colnames(expr.table)[2:ncol(expr.table)] <- names.rep</pre>
                                                                                     7
                                                                                     8
for (i in 2:length(files.list)) {
                                                                                     9
    tb <- read.csv(files.list[i])</pre>
    tb <- tb[, 4:ncol(tb)]
                                                                                     10
                                                                                     11
    n <- ncol(tb)
    repname <- paste(rep(names.list[i]), 1:n, sep = "_")
                                                                                     12
                                                                                     13
    colnames(tb) <- repname</pre>
                                                                                     14
    expr.table <- cbind(expr.table, tb)</pre>
                                                                                     15
    n.rep <- append(n.rep, n)
                                                                                     16
    names.rep <- append(names.rep, repname)</pre>
                                                                                     17
                                                                                     18
meta.sample <- data.frame(sampleName = names.list, n.rep = n.rep)</pre>
                                                                                     19
head(expr.table)
                                                                                     20
```

```
## # A tibble: 6 x 70
     Gene.Symbol GEXC_AMs_1 GEXC_AMs_2 GEXC_AMs_3 `GEXC_DC_Lu_CD1~ `
   GEXC_DC_Lu_CD1~
##
     <chr>
                        <dbl>
                                    <dbl>
                                                <dbl>
                                                                   <dbl>
                                                                                   3
               <dbl>
## 1 ---
                         12.2
                                     25.8
                                                -0.23
                                                                    64.1
                                                                                   4
                75.2
## 2 ---
                         17.3
                                     33.8
                                               -15.0
                                                                    40.0
                                                                                   5
                60.2
                         -2.6
                                     30.2
                                                -9.45
                                                                    45.3
                                                                                   6
## 3 ---
                59.4
                         23.3
                                     44.9
                                                16.6
                                                                    63.8
## 4 ---
                65.3
```

```
## 5 ---
                                             -11.1
                        16.4
                                    61.3
                                                                 22.4
                43.5
                        19.8
                                    44.8
                                              14.6
                                                                 75.4
                                                                               9
                82.6
     ... with 64 more variables: GEXC_DC_Lu_CD103+_3 <dbl>,
                                                                               10
       GEXC_DC_Lu_CD103+_4 <dbl>, GEXC_DC_Lu_CD103+_5 <dbl>,
                                                                               11
       GEXC_DC_Lu_CD24+_1 <dbl>, GEXC_DC_Lu_CD24+_2 <dbl>,
                                                                               12
       GEXC_DCLuLN_CD103+_1 <dbl>, GEXC_DCLuLN_CD103+_2 <dbl>,
                                                                               13
## #
       GEXC_DCLuLN_CD103+_3 <dbl>, GEXC_DCLuLN_CD11b+_1 <dbl>,
                                                                                14
##
       GEXC_DCLuLN_CD11b+_2 <dbl>, GEXC_DCLuLN_CD11b+_3 <dbl>, GEXC_L+WT_1
                                                                               15
    <dbl>,
       GEXC_L+WT_2 <dbl>, GEXC_L+WT_3 <dbl>, GEXC_LMIsW_1 <dbl>, ...
                                                                               16
## #
```

3 Data preparration

3.1 Make a list with genes to show in heatmap

The gene list IMvs(AM&DC).csv is calculated in ImmGen Datasets. We compared IM microarray data to both MA and DC and get the DE genes.

```
probset.DE <- read.csv("./IMvs(AM&DC).csv")
probset.toShow <- unique(as.character(probset.DE$ProbeSet_ID))

# the table is ordered by ratio, so take the top 100:
probset.toShow <- probset.toShow[1:100]
```

Then base on the intensity in IM, we choose the only top 50 probsets.

Take the to 50 probsets with highest intensity:

```
probset.DE <- probset.DE[order(probset.DE$Mean_A, decreasing = TRUE), ]
probset.top50 <- as.character(probset.DE[1:50, "ProbeSet_ID"])</pre>
```

```
probset.toShow <- intersect(probset.top50, probset.toShow)
```

subset expr.table

```
expr.table.toShow <- expr.table[probset.toShow,]
genes.toShow <- unique(expr.table.toShow$Gene.Symbol)
length(genes.toShow)

1
2
```

```
## [1] 50
```

As genes are unique to each probset, we can use gene symbols as rownames.

```
rownames(expr.table.toShow) <- expr.table.toShow$Gene.Symbol
```

3.2 Make metadata table

```
data.frame(meta.sample$sampleName, order = 1:nrow(meta.sample))
```

```
## # A tibble: 22 x 2
                                                                                   2
##
      meta.sample.sampleName
                                 order
                                                                                   3
##
      <chr>
                                 <int>
##
   1 GEXC_AMs
                                                                                   4
                                     1
    2 GEXC_DC_Lu_CD103+
                                     2
                                                                                   5
##
    3 GEXC_DC_Lu_CD24+
                                                                                   6
##
                                     3
    4 GEXC_DCLuLN_CD103+
                                                                                   7
##
                                     4
                                                                                   8
##
   5 GEXC_DCLuLN_CD11b+
                                                                                   9
##
    6 GEXC L+WT
                                     6
    7 GEXC LMIsW
                                     7
                                                                                   10
##
##
   8 GEXC_MF_BM
                                     8
                                                                                   11
   9 GEXC MF CNS
                                     9
                                                                                   12
## 10 GEXC_MF_Lu_CD11b+_CD24-
                                    10
                                                                                   13
                                                                                   14
## # ... with 12 more rows
```

```
meta.sample$cellType3 <- c("Mac_Alv_Lu", #1
                                   "DC L CD103+ Lu", #2
                                                                                                  2
                                   "DC_{\sqcup}CD24+_{\sqcup}Lu", #3
                                                                                                  3
                                   "DC_CD103+_LuLN", #4
                                                                                                  4
                                   "DC_{\perp}CD11b+_{\perp}LuLN", #5
                                                                                                  5
                                                                                                  6
                                   "Mo_{\perp}Ly6C+_{\perp}Lu", #6
                                                                                                  7
                                   "Mac_Int_Lu", #7
                                   "Mac<sub>□</sub>BM", #8
                                                                                                  8
                                                                                                  9
                                   "Mac⊔CNS", #9
                                   "Mac_{\sqcup}Int_{\sqcup}Lu", #10
                                                                                                  10
                                   "Mac_F4/80hi_PC", #11
                                                                                                  11
                                   "Mac_F4/80lo_PC", #12
                                                                                                  12
                                                                                                  13
                                   "Mac<sub>11</sub>SI", #13
                                   "Mac⊔SLN", #14
                                                                                                  14
                                   "Mac_{\sqcup}SP", #15
                                                                                                  15
                                   "Mo_{\square}Ly6C-_{\square}MHCII-_{\square}BL", #16
                                                                                                  16
                                   "Mo_{\square}Ly6C-_{\square}MHCII+_{\square}BL", #17
                                                                                                  17
                                   "Mo_Ly6C-_MHCIIint_BL", #18
                                                                                                  18
                                                                                                  19
                                   "Mo_{\square}Ly6C+_{\square}MHCII-_{\square}BL", #19
                                   "Mo_Ly6C+_MHCII+_BL", #20
                                                                                                  20
                                   "Mo_Ly6C-_MHCII-_BM", #21
                                                                                                  21
                                   "Mo_Ly6C+_MHCII-_BM" #22
                                                                                                  22
                  )
                                                                                                  23
                                                                                                  24
                                                                                                  25
meta.sample$cellType <- c("aMac", #1</pre>
                                                                                                  26
                  rep("DC", 4), # 2-5
                                                                                                  27
                  "Mo", # 6
                                                                                                  28
                  "iMac", #7
                                                                                                  29
                  rep("Mac", 2), # 8-9
                                                                                                  30
                  "iMac", #10
                  rep("Mac", 5), # 11-15
                                                                                                  31
                                                                                                  32
                  rep("Mo", 7) # 16-22
                                                                                                  33
                                                                                                  34
meta.sample$organ <- c(rep("Lu", 3), #1-3
                                                                                                  35
              rep("LuLN", 2), #4-5
              rep("Lu", 2), #6-7
                                                                                                  36
              "BM", #8
                                                                                                  37
              "CNS", #9
                                                                                                  38
```

```
"Lu", #10
                                                                                   39
            rep("PC",2), #11-12
                                                                                   40
                                                                                   41
            "SI", #13
            "SLN", #14
                                                                                   42
            "SP", #15
                                                                                   43
            rep("BL",5),
                                                                                   44
            rep("BM", 2)
                                                                                   45
                                                                                   46
                                                                                   47
meta.sample$cellType2 <- c(</pre>
                                                                                   48
               "Mac", #1
                                                                                   49
               rep("DC", 4), # 2-5
                                                                                   50
               "Mo", # 6
                                                                                   51
               "Mac", #7
                                                                                   52
               rep("Mac", 2), # 8-9
                                                                                   53
                                                                                   54
               "Mac", #10
               rep("Mac", 5), # 11-15
                                                                                   55
                                                                                   56
               rep("Mo", 7) # 16-22
                                                                                   57
                                                                                   58
meta.sample$organ2 <- c(
             "Lu-Alv", #1
                                                                                   59
             rep("Lu", 2), #2-3
                                                                                   60
            rep("LuLN", 2), #4-5
                                                                                   61
            "Lu", #6
                                                                                   62
                                                                                   63
            "Lu-Int", #7
            "BM", #8
                                                                                   64
            "CNS", #9
                                                                                   65
            "Lu-Int", #10
                                                                                   66
                                                                                   67
            rep("PC",2), #11-12
            "SI", #13
                                                                                   68
            "SLN", #14
                                                                                   69
            "SP", #15
                                                                                   70
            rep("BL",5),
                                                                                   71
                                                                                   72
            rep("BM", 2)
                                                                                   73
                                                                                   74
meta.table <- data.frame(CellType = rep(meta.sample$cellType, meta.sample$</pre>
   n.rep),
    OrganType = rep(meta.sample$organ, meta.sample$n.rep), CellType2 = rep
```

```
meta.table <- data.frame(CellType = rep(meta.sample$cellType, meta.sample$ 1
    n.rep),
    OrganType = rep(meta.sample$organ, meta.sample$n.rep), CellType2 = rep (meta.sample$cellType2,
        meta.sample$cellType2,
        meta.sample$n.rep), OrganType2 = rep(meta.sample$organ2, meta.
        sample$n.rep),
    cellType3 = rep(meta.sample$cellType3, meta.sample$n.rep), row.names = 4
        names.rep)</pre>
```

```
## A HeatmapAnnotation object with 2 annotations

## name: heatmap_annotation_0

## position: column

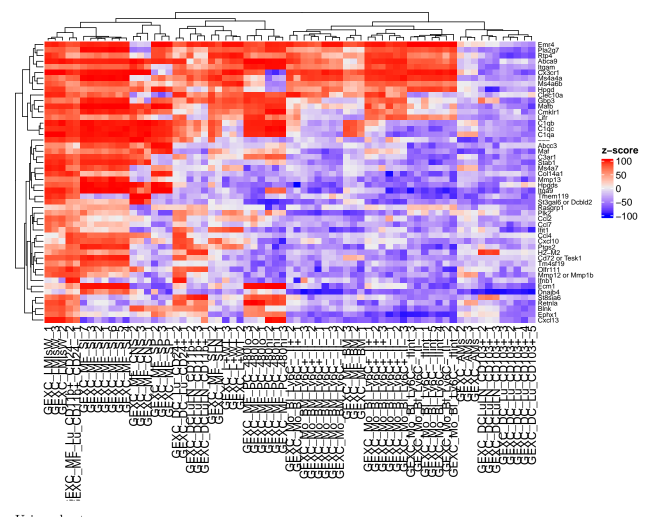
## items: 69
```

```
##
     width: 1npc
##
     height: 10.3514598035146mm
                                                                                6
                                                                                7
##
     this object is subsetable
##
     23.119166666667mm extension on the right
                                                                                8
                                                                                9
##
                                                                                10
##
          name annotation_type color_mapping height
                                                                                11
     Cell_type discrete vector
                                       random
                                                   5mm
    Organ_type discrete vector
                                                                                12
                                        random
                                                   5mm
```

4 Make heatmaps

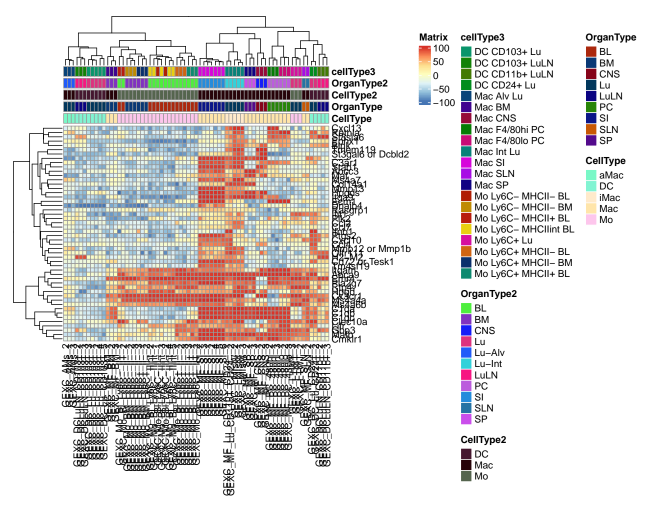
Use Heatmap:

```
Heatmap (
  as.matrix(expr.table.toShow[2:ncol(expr.table.toShow)]),
  # use_raster = FALSE, # use FALSE to export to vector image.
 name
                                = "z-score",
                                   = colorRamp2(seq(from=-2, to=2, length=11),
  # col
     rev(brewer.pal(11, "Spectral"))),
                                                                                6
  # show_row_names
                                   = TRUE,
  # show_column_names
                                   = FALSE,
                                                                                7
 row_names_gp
                                = gpar(fontsize = 7),
                                                                                8
                                                                                9
                                                                                10
  # row_title_rot
                                   = 0,
                                   = TRUE,
                                                                                11
  # cluster rows
                                                                                12
  # cluster_row_slices
                                  = FALSE,
  \#cluster\_columns
                                  = FALSE
                                                                                13
                                                                                14
```



Using pheatmap.

```
pheatmap(
 as.matrix(expr.table.toShow[2:ncol(expr.table.toShow)]), annotation_col
     = meta.table
  # use_raster = FALSE, # use FALSE to export to vector image.
                                 = "z-score",
                                  = colorRamp2(seq(from=-2, to=2, length=11),
  # col
     rev(brewer.pal(11, "Spectral"))),
  # show_row_names
                                  = TRUE,
                                                                               6
                                 = FALSE,
  # show_column_names
                                                                               8
                                 = gpar(fontsize = 7),
  #row_names_gp
                                                                               9
                                                                               10
  # row_title_rot
                                  = 0,
  # cluster_rows
                                  = TRUE,
                                                                               11
  # cluster_row_slices
                                  = FALSE,
                                                                               12
                                 = FALSE
                                                                               13
  \#cluster\_columns
                                                                               14
```



Using Heatmap with annotations.

```
hp <- Heatmap(
                                                                                2
  as.matrix(expr.table.toShow[2:ncol(expr.table.toShow)]),
  # use_raster = FALSE, # use FALSE to export to vector image.
                                                                                3
  name
                                 = "z-score",
                                   = colorRamp2(seq(from=-2, to=2, length=11),
  # col
     rev(brewer.pal(11, "Spectral"))),
  # show_row_names
                                   = TRUE
                                                                                6
                                   = FALSE,
                                                                                7
  # show_column_names
                                                                                8
                                = gpar(fontsize = 7),
  row_names_gp
  column_names_gp
                                = gpar(fontsize = 7),
                                                                                9
                                                                                10
  #column_split = meta.table$CellType2,
                                                                                11
  column_split = factor(meta.table$CellType2, levels = c("Mac", "Mo", "DC"
                                                                                12
  top_annotation = HeatmapAnnotation(Organtype=meta.table $OrganType2,
                                                                                13
                                       col = list(Organtype = c(`Lu-Alv`="
                                                                                14
                                          #32a852",
                                                `Lu-Int`="#87c22f",
                                                                                15
                                                Lu = "#205c30",
                                                                                16
                                                LuLN="#265d69",
                                                                                17
                                                BM="#82622f",
                                                                                18
                                                                                19
                                                CNS="#4674e8",
```

```
PC="#a14bab", 20

SI="#dbed4e", 21

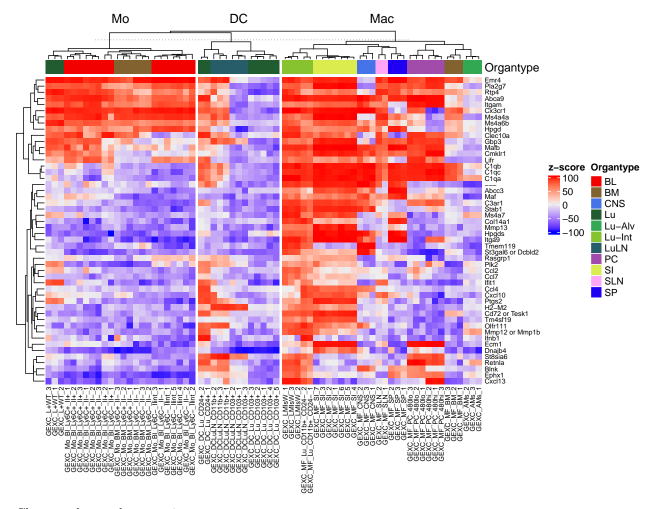
SLN="#ffa6f9", 22

SP="#2000f2", 23

BL="#f20000"))) 24

) 25

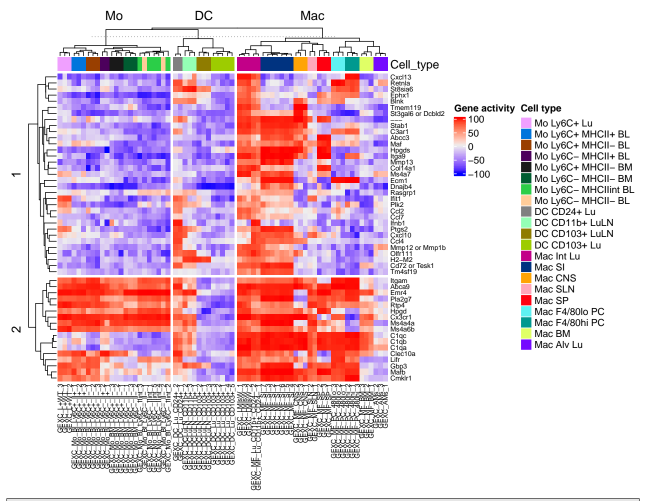
p <- draw(hp) 28
```



Change colors and annotations

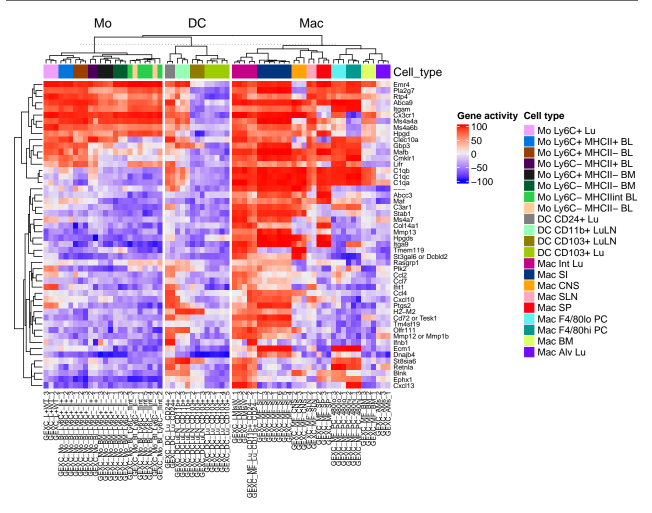
```
col.cellType3 <- read.csv("../0-Microarrays/colors_celltype3.csv", sep = "</pre>
   \t", header = FALSE, row.names = 1)
colors.cellType3 <- as.character(col.cellType3$V2)</pre>
names(colors.cellType3) <- rownames(col.cellType3)</pre>
                                                                                  3
genes.toSplit <- rownames(expr.table.toShow)</pre>
                                                                                  4
genes.toSplit <- genes.toSplit %in% c("Tmem119", "Cx3cr1")</pre>
                                                                                  5
                                                                                  6
# the one with row split:
hp2 <- Heatmap(
                                                                                  8
  as.matrix(expr.table.toShow[2:ncol(expr.table.toShow)]),
                                                                                  9
  # use_raster = FALSE, # use FALSE to export to vector image.
                                                                                  10
```

```
= "Gene \activity",
                                                                                 11
  name
                                                                                 12
                                   gpar(fontsize = 7),
  row_names_gp
                                 = gpar(fontsize = 7),
                                                                                 13
  column_names_gp
  column_split = factor(meta.table$CellType2, levels = c("Mac", "Mo", "DC"
                                                                                 14
     )),
                                                                                 15
  row_split = 2,
  top_annotation = HeatmapAnnotation(Cell_type=meta.table$cellType3,
                                                                                 16
                                                                                 17
                     col = list(
                                  Cell_type = colors.cellType3 ),
                                                                                 18
                     annotation_legend_param = list(
                                                                                 19
                                  Cell_type = list(title = "Cell_type",
                                                                                 20
                                                                                 21
                                                    at = names(colors.
                                                        cellType3),
                                                                                 22
                                                    labels = names(colors.
                                                        cellType3))) )
                                                                                 23
                                                                                 24
                                                                                 25
                                                                                 26
p2 <- draw(hp2)
```



```
# the one WITHOUT row split:
hp3 <- Heatmap(
   as.matrix(expr.table.toShow[2:ncol(expr.table.toShow)]),</pre>
1
2
3
```

```
# use_raster = FALSE, # use FALSE to export to vector image.
                                 = "Gene activity",
                                                                                5
  name
                                                                                6
                                 = gpar(fontsize = 7),
  row_names_gp
                                = gpar(fontsize = 7),
                                                                                7
  column_names_gp
  column_split = factor(meta.table$CellType2, levels = c("Mac", "Mo", "DC"
                                                                                8
  \#row\_split = 2,
                                                                                9
  top_annotation = HeatmapAnnotation(Cell_type=meta.table$cellType3,
                                                                                10
                     col = list(
                                                                                11
                                  Cell_type = colors.cellType3 ),
                                                                                12
                     annotation_legend_param = list(
                                                                                13
                                  Cell_type = list(title = "Cell_type",
                                                                                14
                                                    at = names(colors.
                                                                                15
                                                       cellType3),
                                                    labels = names(colors.
                                                                                16
                                                       cellType3))) )
                                                                                17
                                                                                18
                                                                                19
                                                                                20
p3 <- draw(hp3)
```



5 Session information

R session:

```
sessionInfo()
```

```
## R version 4.0.3 (2020-10-10)
## Platform: x86_64-pc-linux-gnu (64-bit)
                                                                                 2
## Running under: Ubuntu 20.04.3 LTS
                                                                                 5
## Matrix products: default
           /usr/lib/x86_64-linux-gnu/openblas-pthread/libblas.so.3
                                                                                 6
## LAPACK: /usr/lib/x86_64-linux-gnu/openblas-pthread/liblapack.so.3
##
                                                                                 8
##
  locale:
                                                                                 9
                                                                                 10
##
    [1] LC_CTYPE=en_US.UTF-8
                                     LC_NUMERIC=C
##
    [3] LC_TIME=en_GB.UTF-8
                                     LC_COLLATE = en_US.UTF-8
                                                                                 11
    [5] LC_MONETARY=en_GB.UTF-8
                                                                                 12
##
                                     LC_MESSAGES=en_US.UTF-8
##
    [7] LC_PAPER=en_GB.UTF-8
                                     LC_NAME = C
                                                                                 13
    [9] LC_ADDRESS=C
                                     LC_TELEPHONE = C
                                                                                 14
  [11] LC_MEASUREMENT=en_GB.UTF-8 LC_IDENTIFICATION=C
                                                                                 15
                                                                                 16
## attached base packages:
                                                                                 17
## [1] grid
                             graphics grDevices utils
                                                                                 18
                  stats
                                                             datasets
                                                                        methods
## [8] base
                                                                                 19
                                                                                 20
                                                                                 21
## other attached packages:
## [1] circlize_0.4.13
                                                                                 22
                              RColorBrewer_1.1-2
                                                    ComplexHeatmap_2.6.2
                                                                                 23
##
                                                                                 24
## loaded via a namespace (and not attached):
                                                                                 25
##
    [1] Rcpp_1.0.7
                              highr_0.9
                                                   pillar_1.6.2
                                                                                 26
##
    [4] compiler_4.0.3
                              formatR_1.11
                                                   tools_4.0.3
                                                                                 27
##
    [7] digest_0.6.27
                                                   lifecycle_1.0.0
                              evaluate_0.14
## [10] tibble_3.1.3
                              clue_0.3-59
                                                   pkgconfig_2.0.3
                                                                                 28
                                                                                 29
## [13] png_0.1-7
                              rlang_0.4.11
                                                   rstudioapi_0.13
                                                                                 30
## [16] cli_3.0.1
                              magick_2.7.2
                                                   yaml_2.2.1
                                                                                 31
  [19] parallel_4.0.3
                              xfun_0.24
                                                   stringr_1.4.0
                                                                                 32
## [22] knitr 1.33
                              cluster_2.1.0
                                                   GlobalOptions_0.1.2
                                                                                 33
## [25] vctrs 0.3.8
                              S4Vectors 0.28.1
                                                   IRanges 2.24.1
## [28] stats4_4.0.3
                                                   fansi_0.5.0
                                                                                 34
                              GetoptLong_1.0.5
## [31] rmarkdown 2.9
                              magrittr_2.0.1
                                                   matrixStats 0.60.0
                                                                                 35
## [34] ellipsis_0.3.2
                              htmltools_0.5.1.1
                                                   BiocGenerics_0.36.1
                                                                                 36
                                                                                 37
                                                   utf8_1.2.2
## [37] shape_1.4.6
                              colorspace_2.0-2
                                                                                 38
                              crayon_1.4.1
                                                   rjson_0.2.20
## [40] stringi_1.7.3
                                                                                 39
## [43] Cairo_1.5-12.2
```

6 References