

Day 2 – 02 Simple Heatmap (Exercises – Solutions)

Seminar practice worksheet

Below is one possible solution for the worksheet that builds the heatmap from `dataset2_subset.csv` / `dataset2_subset_long.csv` (three genomes). Feel free to compare this with your own approach.

1. Load packages and define paths

```
library(ComplexHeatmap)

## Loading required package: grid
## =====
## ComplexHeatmap version 2.24.1
## Bioconductor page: http://bioconductor.org/packages/ComplexHeatmap/
## Github page: https://github.com/jokergoo/ComplexHeatmap
## Documentation: http://jokergoo.github.io/ComplexHeatmap-reference
##
## If you use it in published research, please cite either one:
## - Gu, Z. Complex Heatmap Visualization. iMeta 2022.
## - Gu, Z. Complex heatmaps reveal patterns and correlations in multidimensional
##   genomic data. Bioinformatics 2016.
##
##
## The new InteractiveComplexHeatmap package can directly export static
## complex heatmaps into an interactive Shiny app with zero effort. Have a try!
##
## This message can be suppressed by:
##   suppressPackageStartupMessages(library(ComplexHeatmap))
## =====

library(circlize)

## =====
## circlize version 0.4.16
## CRAN page: https://cran.r-project.org/package=circlize
## Github page: https://github.com/jokergoo/circlize
## Documentation: https://jokergoo.github.io/circlize_book/book/
##
## If you use it in published research, please cite:
## Gu, Z. circlize implements and enhances circular visualization
##   in R. Bioinformatics 2014.
##
## This message can be suppressed by:
##   suppressPackageStartupMessages(library(circlize))
## =====
```

```

subset_path <- file.path('..', 'data', 'dataset2_subset.csv')
long_path <- file.path('..', 'data', 'dataset2_subset_long.csv')
pdf_path <- file.path('..', 'pdf', 'dataset2_heatmap.pdf')

```

2. Load/inspect the data

```

wide_df <- read.csv(subset_path, check.names = FALSE, stringsAsFactors = FALSE)
long_df <- read.csv(long_path, check.names = FALSE, stringsAsFactors = FALSE)

cat('Wide table dimensions:', nrow(wide_df), 'rows x', ncol(wide_df), 'columns\n')

## Wide table dimensions: 48 rows x 67 columns
cat('Long table dimensions:', nrow(long_df), 'rows x', ncol(long_df), 'columns\n')

## Long table dimensions: 3072 rows x 7 columns
head(wide_df[, c('Genome', 'snp_id', 'Position')])

##                                     Genome      snp_id Position
## 1 Akkermansia_muciniphila_YL44 239840-C-G  239840
## 2 Akkermansia_muciniphila_YL44 241793-A-G  241793
## 3 Akkermansia_muciniphila_YL44 355328-A-T  355328
## 4 Akkermansia_muciniphila_YL44 356291-C-A  356291
## 5 Akkermansia_muciniphila_YL44 2351445-C-T 2351445
## 6 Bacteroides_caecimuris_I48 1601848-T-C  1601848

head(long_df)

##           Genome      snp_id Position     value mouse_id day
## 1 Akkermansia_muciniphila_YL44 239840-C-G  239840 0.000000  1683  0
## 2 Akkermansia_muciniphila_YL44 241793-A-G  241793 0.049587  1683  0
## 3 Akkermansia_muciniphila_YL44 355328-A-T  355328 0.138182  1683  0
## 4 Akkermansia_muciniphila_YL44 356291-C-A  356291 0.000000  1683  0
## 5 Akkermansia_muciniphila_YL44 2351445-C-T 2351445 0.000000  1683  0
## 6 Bacteroides_caecimuris_I48 1601848-T-C  1601848 0.041609  1683  0
##   treatment_group
## 1          Control
## 2          Control
## 3          Control
## 4          Control
## 5          Control
## 6          Control

```

3. Choose a treatment group subset

```

target_group <- 'Control'
sample_meta <- unique(long_df[, c('mouse_id', 'day', 'treatment_group')])
sample_meta$sample_id <- paste(sample_meta$mouse_id, sample_meta$day, sep = '-')
keep_samples <- sample_meta$sample_id[sample_meta$treatment_group == target_group]
wide_df <- wide_df[, c('Genome', 'snp_id', 'Position', keep_samples)]

```

4. Quick summaries

```
print(table(wide_df$Genome))

##
## Akkermansia_muciniphila_YL44    Bacteroides_caecimuris_I48
##                      5                  19
## Turicimonas_muris_YL45
##                      24

mouse_day_table <- with(long_df, table(mouse_id, day))
print(mouse_day_table)

##          day
## mouse_id  0  4  9 14 18 23 30 37 44 49 53 58 63 67 72 79
##      1683 48 48 48 48 48 48 48 48 48 48 48 48 48 48 48 48
##      1688 48 48 48 48 48 48 48 48 48 48 48 48 48 48 48 48
##      1692 48 48 48 48 48 48 48 48 48 48 48 48 48 48 48 48
##      1699 48 48 48 48 48 48 48 48 48 48 48 48 48 48 48 48

value_summary <- tapply(long_df$value, long_df$Genome, function(x) {
  c(min = min(x, na.rm = TRUE),
    median = median(x, na.rm = TRUE),
    max = max(x, na.rm = TRUE)))
})

value_summary <- do.call(rbind, value_summary)
print(value_summary)

##                   min     median       max
## Akkermansia_muciniphila_YL44   0 0.0523090 0.871245
## Bacteroides_caecimuris_I48     0 0.0338305 0.991218
## Turicimonas_muris_YL45        0 0.2649125 1.000000
```

5. Build the heatmap matrix

```
sample_cols <- setdiff(names(wide_df), c('Genome', 'snp_id', 'Position'))
heatmap_matrix <- as.matrix(wide_df[, sample_cols])
mode(heatmap_matrix) <- 'numeric'
rownames(heatmap_matrix) <- paste(wide_df$Genome, wide_df$snp_id, sep = ' | ')

sample_meta <- data.frame(sample_id = sample_cols, stringsAsFactors = FALSE)
split_ids <- strsplit(sample_meta$sample_id, '|', fixed = TRUE)
sample_meta$mouse_id <- vapply(split_ids, function(x) x[[1]], character(1))
sample_meta$day <- as.integer(vapply(split_ids, function(x) if (length(x) >= 2) x[[2]] else NA_character_))

order_idx <- order(sample_meta$mouse_id, sample_meta$day, sample_meta$sample_id)
sample_meta <- sample_meta[order_idx, ]
heatmap_matrix <- heatmap_matrix[, sample_meta$sample_id, drop = FALSE]
```

6. Colors and annotations

```
min_val <- min(heatmap_matrix, na.rm = TRUE)
max_val <- max(heatmap_matrix, na.rm = TRUE)
if (!is.finite(min_val)) min_val <- 0
if (!is.finite(max_val)) max_val <- 1
```

```

if (abs(max_val - min_val) < .Machine$double.eps) {
  max_val <- min_val + 1
}
mid_val <- (min_val + max_val) / 2
color_fun <- circlize::colorRamp2(c(min_val, mid_val, max_val),
                                    c('#0c2c84', '#f7fbff', '#b30000'))

mouse_levels <- unique(sample_meta$mouse_id)
mouse_colors <- setNames(grDevices::rainbow(length(mouse_levels)), mouse_levels)

min_day <- min(sample_meta$day, na.rm = TRUE)
max_day <- max(sample_meta$day, na.rm = TRUE)
if (min_day == max_day) {
  day_colors <- circlize::colorRamp2(c(min_day, min_day + 1), c('#fee8c8', '#e34a33'))
} else {
  day_colors <- circlize::colorRamp2(seq(min_day, max_day, length.out = 3),
                                       c('#fee8c8', '#fdbb84', '#e34a33'))
}
col_annotation <- HeatmapAnnotation(
  mouse = factor(sample_meta$mouse_id, levels = mouse_levels),
  day = sample_meta$day,
  col = list(mouse = mouse_colors, day = day_colors),
  annotation_name_side = 'left'
)

```

7. Draw and export the heatmap

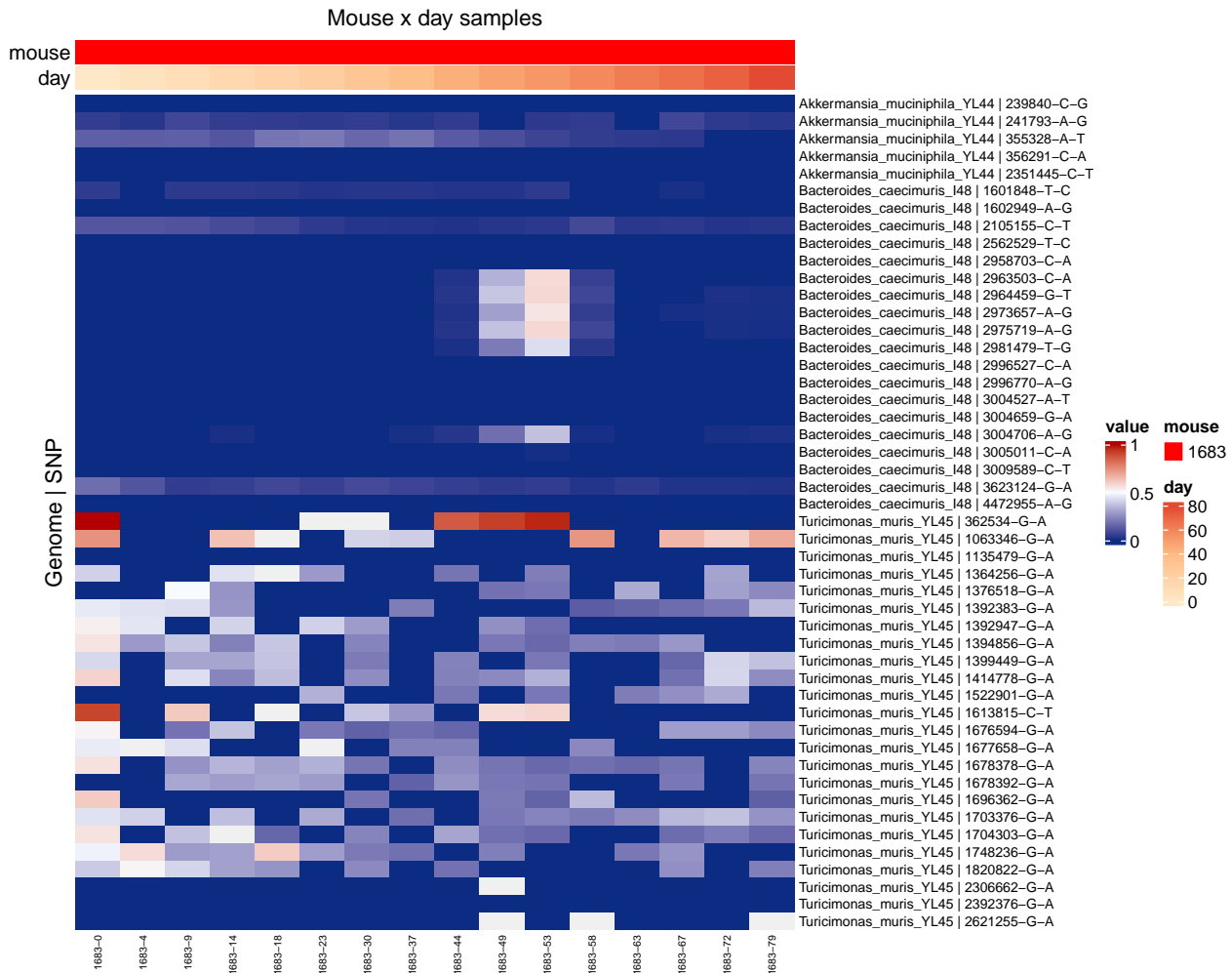
```

row_name_size <- max(5, min(8, 40 / log10(max(10, nrow(heatmap_matrix)))))
col_name_size <- max(6, min(10, 80 / ncol(heatmap_matrix)))

ht <- Heatmap(
  heatmap_matrix,
  name = 'value',
  col = color_fun,
  na_col = '#f0f0f0',
  top_annotation = col_annotation,
  column_split = factor(sample_meta$mouse_id, levels = mouse_levels),
  cluster_rows = FALSE,
  cluster_columns = FALSE,
  column_title = 'Mouse x day samples',
  row_title = 'Genome | SNP',
  show_row_names = TRUE,
  show_column_names = TRUE,
  row_names_gp = grid::gpar(fontsize = row_name_size),
  column_names_gp = grid::gpar(fontsize = col_name_size)
)

draw(ht, heatmap_legend_side = 'right', annotation_legend_side = 'right')

```



```
pdf_height <- max(6, min(18, 0.2 * nrow(heatmap_matrix) + 4))
pdf_width <- max(8, min(16, 0.2 * ncol(heatmap_matrix) + 6))

dir.create(dirname(pdf_path), recursive = TRUE, showWarnings = FALSE)
pdf(pdf_path, width = pdf_width, height = pdf_height)
draw(ht, heatmap_legend_side = 'right', annotation_legend_side = 'right')
dev.off()
```

```
## pdf  
## 2  
cat('Saved heatmap to', pdf_path, '\n')
```

```
## Saved heatmap to ../pdf/dataset2 heatmap.pdf
```

For extra practice, try adding `row_split` by Genome or experiment with a different color palette.