2. Heatmaps of lab and field, facs data

Fay

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Import data

```
MICE <- read.csv("output_data/1.MICE_cleaned_data.csv")</pre>
```

Vectors for selecting genes

Heatmap on lab FACS data

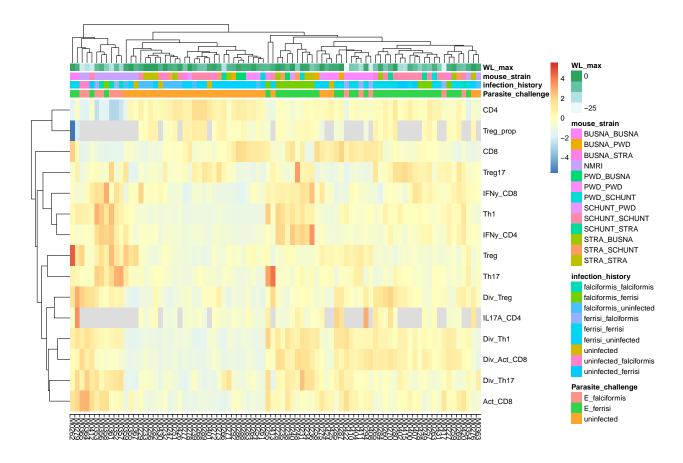
data cleaning

```
### Select the measurements from the mesenterial lymphnodes
### Prepare the annotation data frame for the heatma
annotation_df <- MICE %>%
 filter(origin == "Lab") %>% # filter the lab data
 filter(Position == "mLN") %>%
 filter(infection == "challenge") %>%
 drop na("CD4") %>%
 dplyr::select(c("Mouse_ID", "Parasite_challenge", "infection_history",
                    "mouse_strain", "WL_max"))
  ### Drop the columns that contain nas in the column CD4 of
#the facs columns
### Data tidying for the heatmap function
FACS <- MICE %>%
  dplyr::filter(origin == "Lab") %>% # filter the lab data
  filter(Position == "mLN") %>%
  filter(infection == "challenge", dpi == dpi_max) %>%
  drop_na("CD4") %>%
  dplyr::select(c(Mouse_ID, all_of(facs_lab)))
```

```
# turn the data frame into a matrix and transpose it. We want to have each cell
# type as a row name
FACS <- t(as.matrix(FACS))</pre>
#switch the matrix back to a data frame format
FACS <- as.data.frame(FACS)
# turn the first row into column names
FACS %>%
 row_to_names(row_number = 1) -> FACS
# Now further prepare the data frame for plotting by removing the first row
## and convert the column to row names with the cells
heatmap_data <- FACS
# turn the columns to numeric other wise the heatmap function will not work
heatmap_data[] <- lapply(heatmap_data, function(x) as.numeric(as.character(x)))</pre>
annotation_df <- unique(annotation_df) %>%
    dplyr::filter(Mouse_ID %in% colnames(heatmap_data))
### Prepare the annotation columns for the heatmap
rownames(annotation_df) <- annotation_df$EH_ID</pre>
# Match the row names to the heatmap data frame
rownames(annotation_df) <- colnames(FACS)</pre>
#remove the unecessary column
annotation_df <- annotation_df %>% dplyr::select(-Mouse_ID, )
heatmap_facs_LAB <- heatmap_data
```

Heatmap lab facs: Plot

```
heatmap_data %>%
  pheatmap(annotation_col = annotation_df, scale = "row")
```



Field data

data cleaning

```
### Select the measurements from the mesenterial lymphnodes
### Prepare the annotation data frame for the heatmap
annotation_df <- MICE %>%
    filter(origin == "Field") %>% # filter the lab data
    dplyr::select(c("Mouse_ID", "Sex", "HI", "delta_ct_cewe_MminusE")) %>%
    drop_na()

### Prepare the annotation columns for the heatmap
rownames(annotation_df) <- annotation_df$Mouse_ID

### Data tidying for the heatmap function

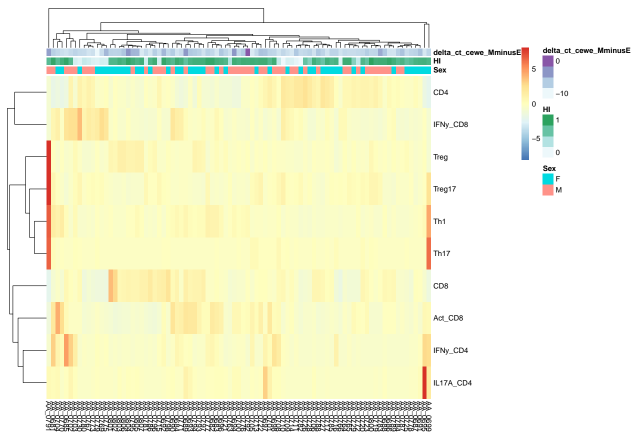
FACS <- MICE %>%
    dplyr::filter(origin == "Field") %>% # filter the lab data
    dplyr::select(c(Mouse_ID, all_of(facs_field)))

# turn the data frame into a matrix and transpose it. We want to have each cell
# type as a row name
FACS <- t(as.matrix(FACS))</pre>
```

```
#switch the matrix back to a data frame format
FACS <- as.data.frame(FACS)</pre>
# turn the first row into column names
FACS %>%
 row_to_names(row_number = 1) -> FACS
# Now further prepare the data frame for plotting by removing the first row
## and convert the column to row names with the cells
FACS -> heatmap_data
# turn the columns to numeric other wise the heatmap function will not work
heatmap_data[] <- lapply(heatmap_data, function(x) as.numeric(as.character(x)))
 # remove columns with only NAs
heatmap_data <- Filter(function(x)!all(is.na(x)), heatmap_data)</pre>
 #remove rows with only Nas
 heatmap_data <- heatmap_data[, colSums(is.na(heatmap_data)) !=</pre>
                                  nrow(heatmap_data)]
annotation_df <- unique(annotation_df) %>%
    dplyr::filter(Mouse_ID %in% colnames(heatmap_data))
#select the row names from the annotation df
heatmap_data <- heatmap_data %>%
  dplyr::select(row.names(annotation_df))
# Match the row names to the heatmap data frame
rownames(annotation_df) <- colnames(heatmap_data)</pre>
annotation_df <- annotation_df %>%
  dplyr::select(-Mouse_ID)
heatmap_facs_FIELD <- heatmap_data
```

Heatmap field facs: Plot

```
heatmap_data %>%
  pheatmap(annotation_col = annotation_df, scale = "row")
```



Lab + Field Heatmap combination

```
# select the same cells for both
names_rows_field <- row.names(heatmap_facs_FIELD)
heatmap_facs_LAB <- heatmap_facs_LAB[row.names(heatmap_facs_FIELD), ]
heatmap_lab_field <- cbind(heatmap_facs_LAB, heatmap_facs_FIELD)

#Prepare the annotation data frame
annotation_df <- MICE %%
    dplyr::select(origin, Mouse_ID)
annotation_df <- unique(annotation_df) %>%
    dplyr::filter(Mouse_ID %in% colnames(heatmap_lab_field))

### Prepare the annotation columns for the heatmap
rownames(annotation_df) <- annotation_df$Mouse_ID
annotation_df <- unique(annotation_df)
annotation_df <- as.data.frame(annotation_df)

# Match the row names to the heatmap data frame
rownames(annotation_df) <- colnames(heatmap_lab_field)</pre>
```

```
#remove the unecessary column
annotation_df <- annotation_df %>% dplyr::select(-Mouse_ID, )
```

Lab - Field data heatmap - plot

pheatmap(heatmap_lab_field, annotation_col = annotation_df, scale = "row")

