# 8. PCA FACS heatmap wild

Fay

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### Load libraries

```
library(tidyverse)
## -- Attaching packages -
                                                          ----- tidyverse 1.3.2 --
## v ggplot2 3.3.6
                                  0.3.5
                       v purrr
## v tibble 3.1.8
                        v dplyr
                                  1.0.10
## v tidyr
           1.2.1
                        v stringr 1.4.1
## v readr
           2.1.3
                        v forcats 0.5.2
## -- Conflicts ----
                                                   ----- tidyverse_conflicts() --
## x dplyr::filter() masks stats::filter()
## x dplyr::lag()
                     masks stats::lag()
library(dplyr)
library(stringr)
library(FactoMineR)
library(reshape2)
##
## Attaching package: 'reshape2'
## The following object is masked from 'package:tidyr':
##
##
       smiths
library(corrplot)
## corrplot 0.92 loaded
library(factoextra)
## Welcome! Want to learn more? See two factoextra-related books at https://goo.gl/ve3WBa
library(lmtest)
## Loading required package: zoo
##
## Attaching package: 'zoo'
## The following objects are masked from 'package:base':
##
##
       as.Date, as.Date.numeric
library(ggpubr)
library(janitor)
```

```
##
## Attaching package: 'janitor'
##
## The following objects are masked from 'package:stats':
##
## chisq.test, fisher.test
library(pheatmap)
```

### Load data

```
hm <- read.csv("output_data/imputed_mice.csv")</pre>
```

### vectors for selecting

```
<- c("IFNy", "CXCR3", "IL.6", "IL.13", "IL.10",
Gene_field
                "IL1RN", "CASP1", "CXCL9", "ID01", "IRGM1", "MP0",
                "MUC2", "MUC5AC", "MYD88", "NCR1", "PRF1", "RETNLB", "SOCS1",
                "TICAM1", "TNF") #"IL.12", "IRG6")
#add a suffix to represent changes in data file
Gene_field_imp <- paste(Gene_field, "imp", sep = "_")</pre>
Genes_wild <- c("IFNy", "CXCR3", "IL.6", "IL.13", "IL.10",
                  "IL1RN", "CASP1", "CXCL9", "ID01", "IRGM1", "MP0",
                  "MUC2", "MUC5AC", "MYD88", "NCR1", "PRF1", "RETNLB", "SOCS1",
                  "TICAM1", "TNF") #, "IL.12", "IRG6")
Genes_wild_imp <- paste(Genes_wild, "imp", sep = "_")</pre>
Facs_lab <- c("Position", "CD4", "Treg", "Div_Treg", "Treg17", "Th1",</pre>
                    "Div_Th1", "Th17", "Div_Th17", "CD8", "Act_CD8",
                    "Div_Act_CD8", "IFNy_CD4", "IFNy_CD8", "Treg_prop",
                    "IL17A_CD4")
Facs_wild <- c( "Treg", "CD4", "Treg17", "Th1", "Th17", "CD8",
                     "Act_CD8", "IFNy_CD4", "IL17A_CD4", "IFNy_CD8")
```

## PCA on the field facs -imputed

```
field <- hm %>%
    dplyr::filter(origin == "Field")

field <- unique(field)

#make a factor out of the melting curves (important for later visualization)
field <- field %>%
    dplyr::mutate(MC.Eimeria = as.factor(MC.Eimeria))

facs_mouse <- field %>%
```

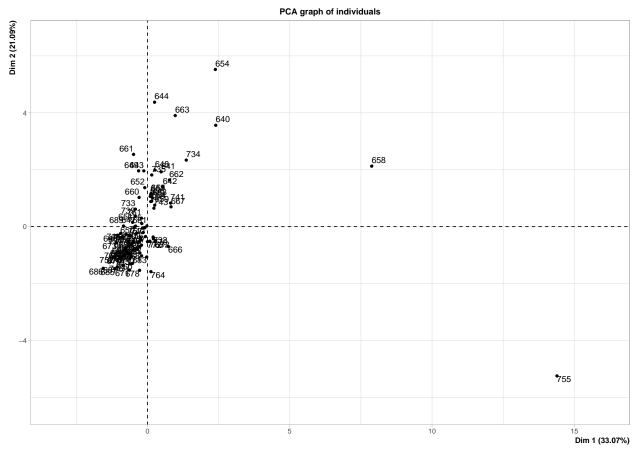
```
dplyr::select(c(Mouse_ID, all_of(Facs_wild)))
facs <- facs_mouse %>%
    dplyr::select(-Mouse_ID)

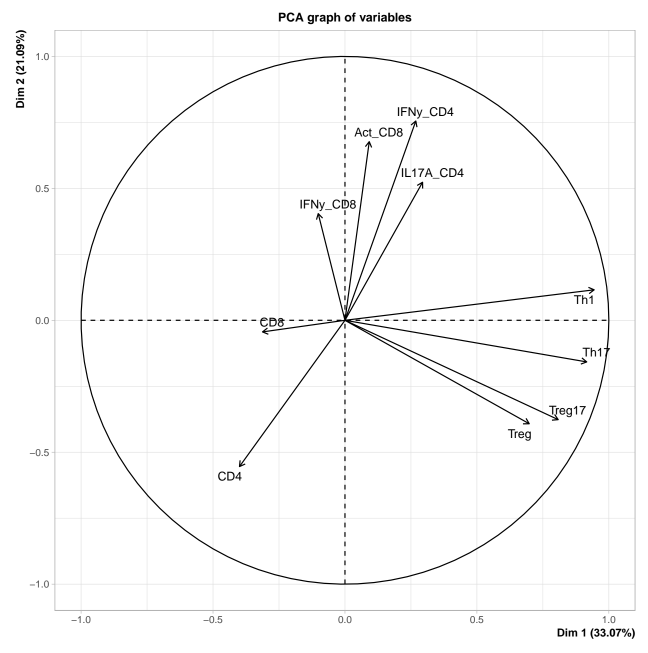
#remove rows with only nas
facs <- facs[,colSums(is.na(facs))<nrow(facs)]

#remove colums with only nas
facs <- facs[rowSums(is.na(facs)) != ncol(facs), ]

# select same rows in facs_mouse
facs_mouse <- facs_mouse[row.names(facs),]

# we can now run a normal pca on the complete data set
res.pca <- PCA(facs)</pre>
```





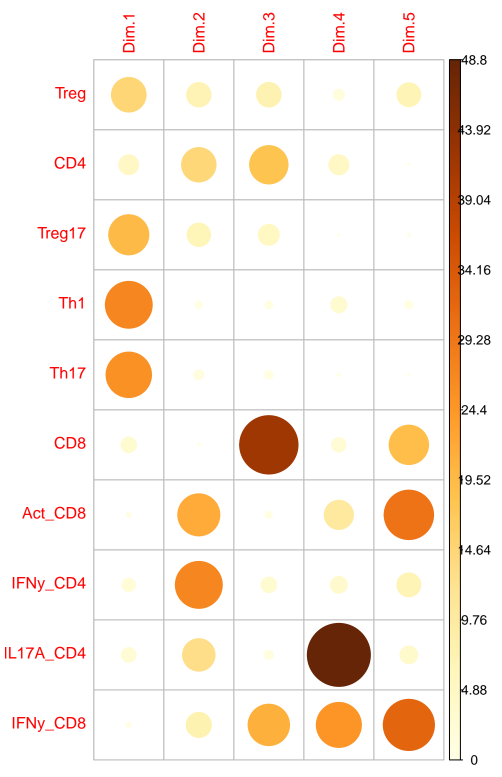
Caution: When imputing data, the percentages of inertia associated with the first dimensions will be overestimated.

Another problem: the imputed data are, when the pca is performed considered like real observations. But they are estimations!!

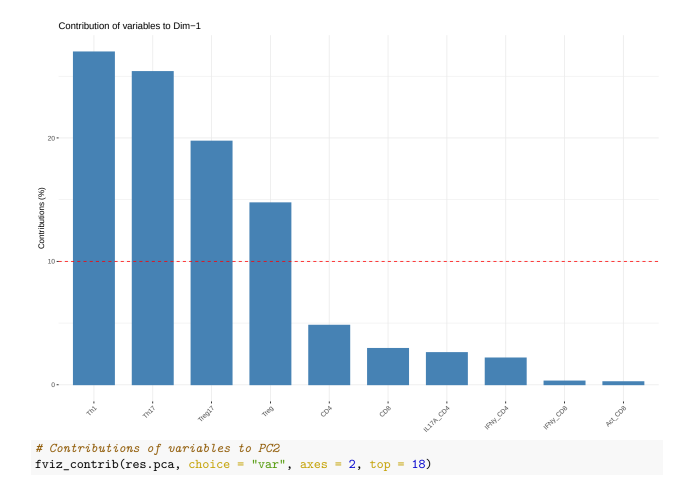
Visualizing uncertainty due to issing data:

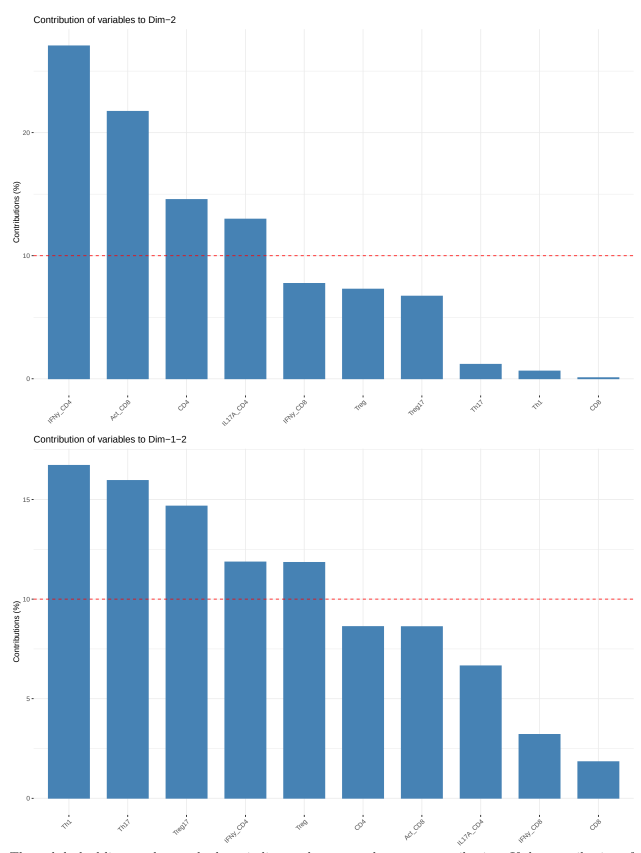
-> mulrimple imputation: generate several plausible values for each missing data point

We here visualize the variability, that is uncertainty on the plane defined by two pca axes.



The function fviz\_contrib() [factoextra package] can be used to draw a bar plot of variable contributions. If your data contains many variables, you can decide to show only the top contributing variables. The R code below shows the top 10 variables contributing to the principal components:



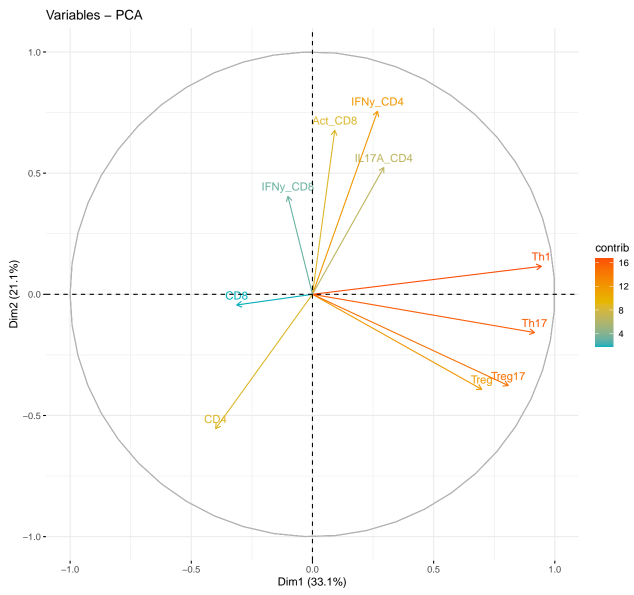


The red dashed line on the graph above indicates the expected average contribution. If the contribution of the variables were uniform, the expected value would be 1/length(variables) = 1/10 = 10%. For a given

component, a variable with a contribution larger than this cutoff could be considered as important in contributing to the component.

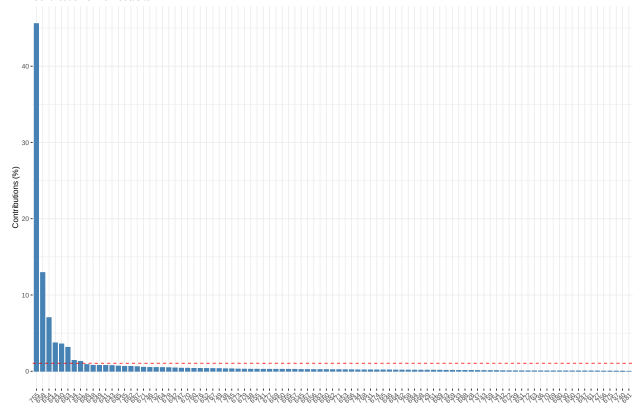
Note that, the total contribution of a given variable, on explaining the variations retained by two principal components, say PC1 and PC2, is calculated as contrib = [(C1 \* Eig1) + (C2 \* Eig2)]/(Eig1 + Eig2), where

C1 and C2 are the contributions of the variable on PC1 and PC2, respectively Eig1 and Eig2 are the eigenvalues of PC1 and PC2, respectively. Recall that eigenvalues measure the amount of variation retained by each PC. In this case, the expected average contribution (cutoff) is calculated as follow: As mentioned above, if the contributions of the 10 variables were uniform, the expected average contribution on a given PC would be 1/10 = 10%. The expected average contribution of a variable for PC1 and PC2 is : [(10\* Eig1) + (10\* Eig2)]/(Eig1 + Eig2)



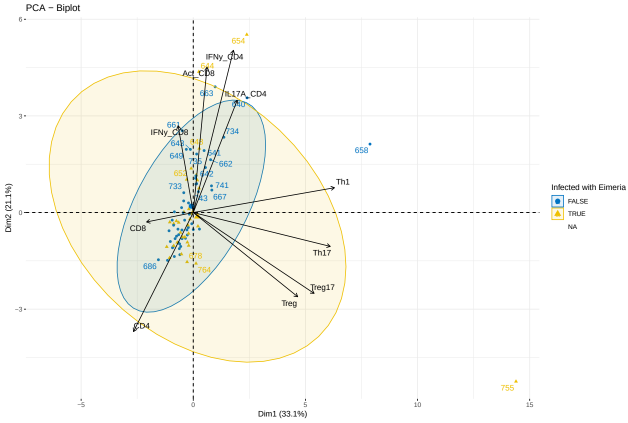
To visualize the contribution of individuals to the first two principal components:





### PCA + Biplot combination

- ## Warning: Removed 8 rows containing missing values (geom\_point).
- ## Warning: Removed 1 rows containing missing values (geom\_point).
- ## Warning: ggrepel: 71 unlabeled data points (too many overlaps). Consider
- ## increasing max.overlaps



In the following example, we want to color both individuals and variables by groups. The trick is to use pointshape = 21 for individual points. This particular point shape can be filled by a color using the argument fill.ind. The border line color of individual points is set to "black" using col.ind. To color variable by groups, the argument col.var will be used.

#### repeating the heatmap o

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

# turn the first row into column names
facs_mouse %>%
    row_to_names(row_number = 1) -> heatmap_data
heatmap_data <- as.data.frame(heatmap_data)

table(rowSums(is.na(heatmap_data)) == nrow(heatmap_data))

##
## FALSE
## 10

# 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>
```

Heatmap on gene expression data:

