6. PCA FACS -lab

Fay

2022-11-03

Always change the knitting directory to the working directory! # Load libraries

```
library(tidyverse)
library(dplyr)
library(stringr)
library(FactoMineR)
library(reshape2)
library(corrplot)
library(factoextra)
library(janitor)
library(janitor)
library(janitor)
library(pheatmap)
library(visdat)
```

Load data

```
hm <- read.csv("output_data/2.imputed_MICE_data_set.csv")</pre>
```

vectors for selecting

```
<- c("IFNy", "CXCR3", "IL.6", "IL.13", "IL.10",
Gene_lab
                "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_lab_imp <- paste(Gene_lab, "imp", sep = "_")</pre>
           <- c("IFNy", "CXCR3", "IL.6", "IL.13", "IL.10",
Genes wild
                  "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("CD4", "Treg", "Div_Treg", "Treg17", "Th1",</pre>
                    "Div_Th1", "Th17", "Div_Th17", "CD8", "Act_CD8",
                    "Div_Act_CD8", "IFNy_CD4", "IFNy_CD8") #"Treg_prop", removed due to many missing va
                     #"IL17A_CD4"
```

FACS

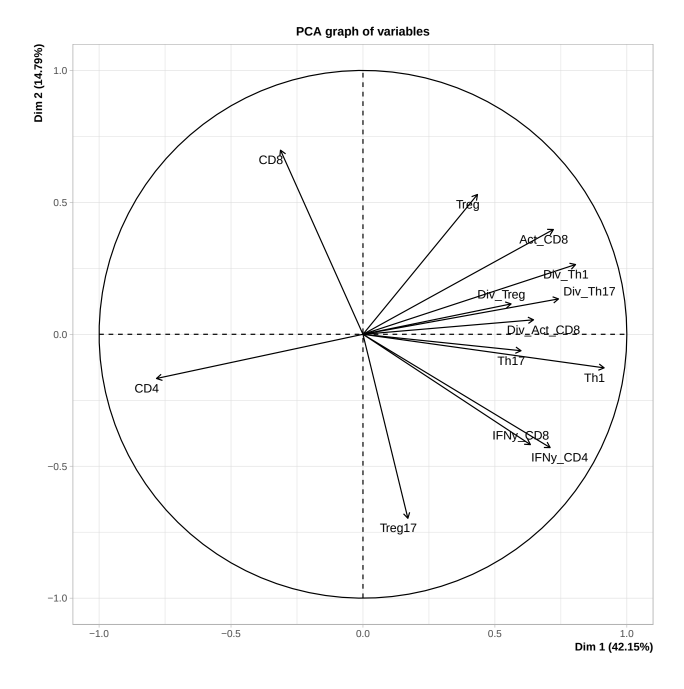
Lab

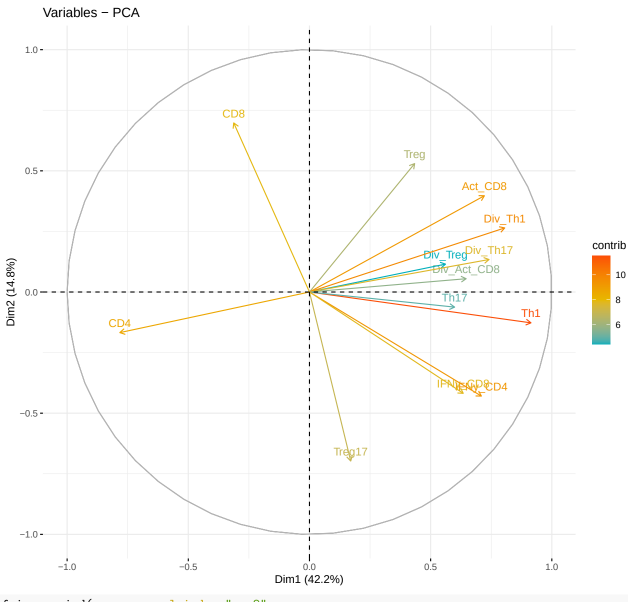
PCA on the lab genes -imputed

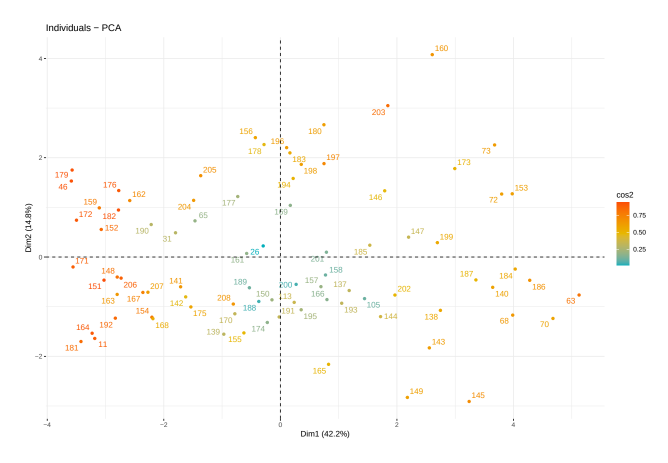
```
#select the genes and lab muce
lab <- hm %>%
  dplyr::filter(origin == "Lab", Position == "mLN") #selecting for mln to avoid
# duplicates
lab <- unique(lab)</pre>
facs_mouse <- lab %>%
  dplyr::select(c(Mouse_ID, all_of(Facs_lab)))
facs <- facs_mouse[, -1]</pre>
#remove rows with only nas
facs <- facs[,colSums(is.na(facs))<nrow(facs)]</pre>
#remove colums with only nas
facs <- facs[rowSums(is.na(facs)) != ncol(facs), ]</pre>
vis_dat(facs)
## Warning: `gather_()` was deprecated in tidyr 1.2.0.
## i Please use `gather()` instead.
## i The deprecated feature was likely used in the visdat package.
## Please report the issue at <a href="https://github.com/ropensci/visdat/issues">https://github.com/ropensci/visdat/issues</a>.
```



res.pca <- PCA(facs)







Dimensions of the pca

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

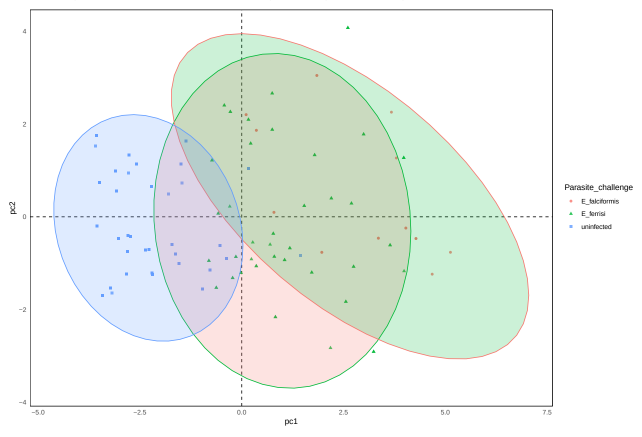
We here visualize the variability, that is uncertainty on the plane defined by two pca axes.

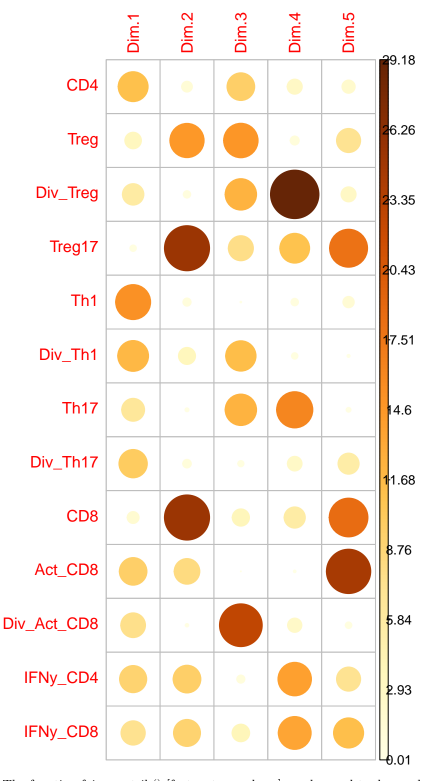
Biplot of the imputed facs pca

```
theme(panel.grid = element_blank(),
    panel.border = element_rect(fill= "transparent"))
```

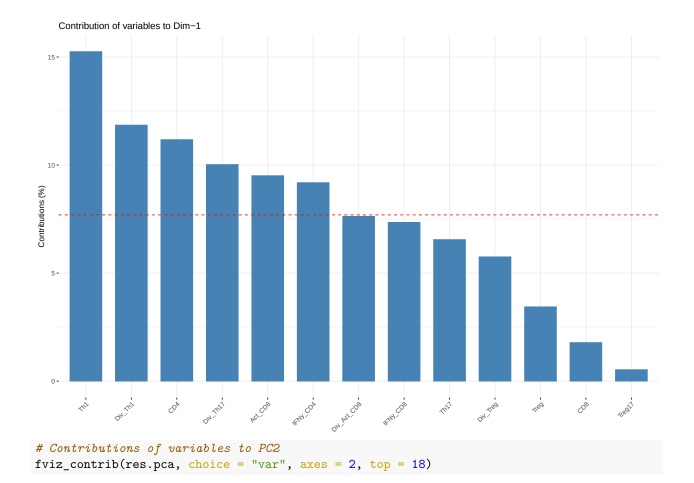
Warning: Removed 56 rows containing non-finite values (`stat_ellipse()`).

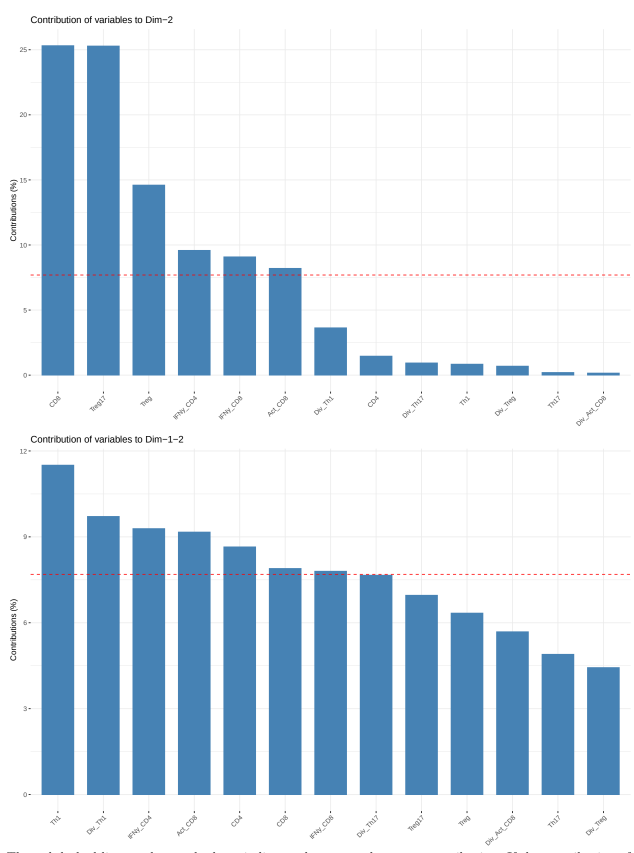
Warning: Removed 56 rows containing missing values (`geom_point()`).





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:



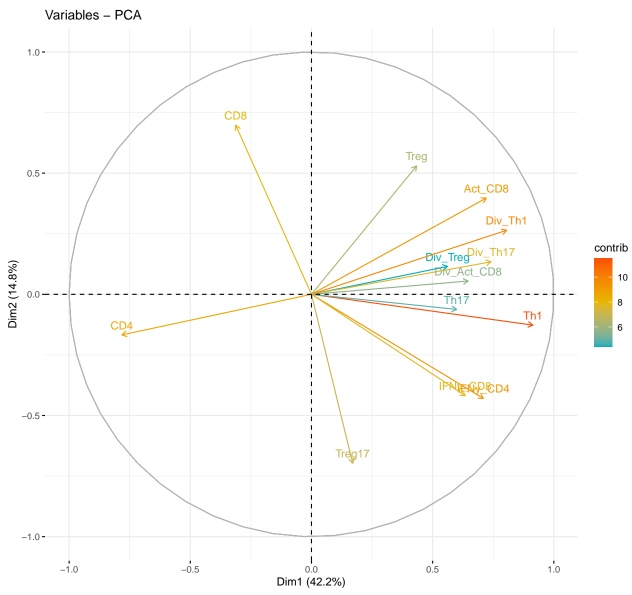


The red dashed line on the graph above indicates the expected average contribution. If the contribution of the variables were uniform, the expected value would be 1/length(variables) = 1/10 = 10%. For a given

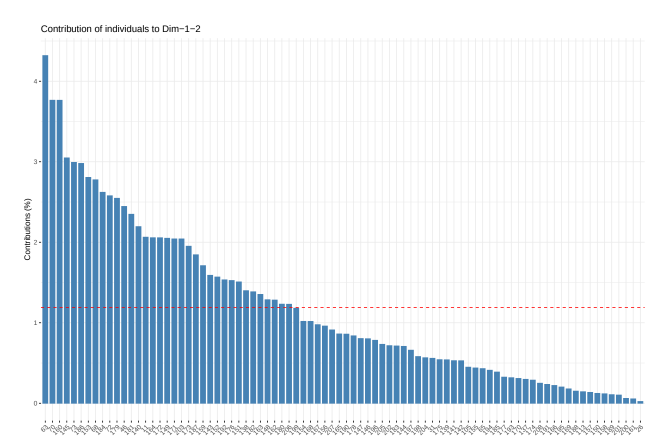
component, a variable with a contribution larger than this cutoff could be considered as important in contributing to the component.

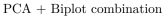
Note that, the total contribution of a given variable, on explaining the variations retained by two principal components, say PC1 and PC2, is calculated as contrib = [(C1 * Eig1) + (C2 * Eig2)]/(Eig1 + Eig2), where

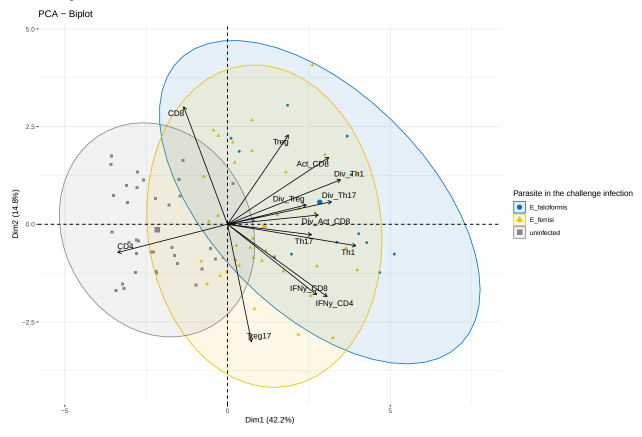
C1 and C2 are the contributions of the variable on PC1 and PC2, respectively Eig1 and Eig2 are the eigenvalues of PC1 and PC2, respectively. Recall that eigenvalues measure the amount of variation retained by each PC. In this case, the expected average contribution (cutoff) is calculated as follow: As 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:







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.

Linear models:

```
##
## Call:
## lm(formula = max_WL ~ pc1 + pc2, data = lab)
##
## Residuals:
##
      Min
                1Q
                   Median
                                3Q
                                       Max
##
  -16.865
           -2.911
                     1.123
                             4.404
                                     9.741
##
## Coefficients:
##
               Estimate Std. Error t value Pr(>|t|)
               92.9127
                            0.6719 138.277
                                           < 2e-16 ***
## (Intercept)
                -0.8623
                            0.2870
                                   -3.004
                                           0.00354 **
                -0.3550
                            0.4845 -0.733 0.46583
## pc2
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 6.158 on 81 degrees of freedom
## Multiple R-squared: 0.1056, Adjusted R-squared: 0.08351
## F-statistic: 4.781 on 2 and 81 DF, p-value: 0.01089
## [1] 548.7185
##
## Call:
## lm(formula = max_WL ~ pc1 + pc2 + hybrid_status, data = lab)
##
## Residuals:
##
        Min
                  1Q
                       Median
                                    3Q
                                            Max
                       0.5969
                                4.0292
##
  -14.7941
            -3.9487
                                        13.2617
##
## Coefficients:
##
                                    Estimate Std. Error t value Pr(>|t|)
## (Intercept)
                                     89.8897
                                                 1.5060 59.687
                                                                 < 2e-16 ***
## pc1
                                     -0.7531
                                                         -2.410
                                                                 0.01837 *
                                                 0.3125
## pc2
                                     -1.9165
                                                 0.8753
                                                         -2.189
                                                                 0.03163 *
## hybrid statusFO M. m. musculus
                                      5.5814
                                                 3.3176
                                                           1.682
                                                                 0.09660 .
## hybrid statusF1 hybrid
                                                 2.0974
                                                           2.880
                                                                 0.00517 **
                                      6.0398
## hybrid statusF1 M. m. domesticus
                                     -0.7613
                                                 2.4155
                                                         -0.315
                                                                 0.75349
## hybrid_statusF1 M. m. musculus
                                      8.3437
                                                 3.6664
                                                                 0.02568 *
                                                           2.276
## hybrid_statusother
                                      2.2420
                                                 2.3853
                                                          0.940 0.35023
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 5.955 on 76 degrees of freedom
## Multiple R-squared: 0.2153, Adjusted R-squared: 0.1431
## F-statistic: 2.979 on 7 and 76 DF, p-value: 0.008133
## [1] 547.7233
```

```
weight_lm3 <- lm(max_WL ~ pc1 + hybrid_status, data = lab)</pre>
summary(weight_lm3)
##
## Call:
## lm(formula = max_WL ~ pc1 + hybrid_status, data = lab)
##
## Residuals:
##
                       Median
        Min
                  1Q
                                     3Q
                                             Max
## -16.9566 -3.7143
                       0.9613
                                 4.5793
                                          9.0997
##
## Coefficients:
##
                                     Estimate Std. Error t value Pr(>|t|)
## (Intercept)
                                      91.9512
                                                   1.2040 76.371
                                                                    <2e-16 ***
                                                   0.3195 -2.231
## pc1
                                      -0.7129
                                                                    0.0286 *
## hybrid_statusF0 M. m. musculus
                                      -0.2341
                                                   2.0362 -0.115
                                                                    0.9088
## hybrid_statusF1 hybrid
                                       3.9198
                                                   1.9058
                                                            2.057
                                                                    0.0431 *
## hybrid_statusF1 M. m. domesticus -0.3417
                                                   2.4665 -0.139
                                                                    0.8902
## hybrid_statusF1 M. m. musculus
                                       2.9391
                                                   2.7769
                                                           1.058
                                                                    0.2932
## hybrid_statusother
                                       0.1930
                                                   2.2475
                                                            0.086
                                                                    0.9318
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 6.1 on 77 degrees of freedom
## Multiple R-squared: 0.1658, Adjusted R-squared: 0.1008
## F-statistic: 2.551 on 6 and 77 DF, p-value: 0.0263
AIC(weight_lm3)
## [1] 550.8613
Try instead: LLR test (likelihood ration) (LM4 package)?
https://www.rdocumentation.org/packages/lmtest/versions/0.9-38/topics/lrtest
In this way you compare each model, with the different variables used to predict.
Another way is to compare the AIC. (function: step)
weight_lm3 <- lm(max_WL ~ pc1 + pc2 + hybrid_status, data = lab)</pre>
weight no pc1 <- lm(max WL ~ pc2 + hybrid status, data = lab)</pre>
weight_no_pc2 <- lm(max_WL ~ pc1 + hybrid_status, data = lab)</pre>
weight_no_hybrid <- lm(max_WL ~ pc1 + pc2, data = lab)</pre>
lrtest(weight_lm2, weight_lm3)
## Likelihood ratio test
## Model 1: max_WL ~ pc1 + pc2 + hybrid_status
## Model 2: max_WL ~ pc1 + pc2 + hybrid_status
     #Df LogLik Df Chisq Pr(>Chisq)
## 1
       9 -264.86
       9 -264.86 0
                                    1
lrtest(weight_lm, weight_lm3)
## Likelihood ratio test
##
## Model 1: max_WL ~ pc1 + pc2
```

```
## Model 2: max_WL ~ pc1 + pc2 + hybrid_status
   #Df LogLik Df Chisq Pr(>Chisq)
## 1 4 -270.36
## 2 9 -264.86 5 10.995
                             0.05147 .
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
lrtest(weight_lm3, weight_no_hybrid)
## Likelihood ratio test
## Model 1: max_WL ~ pc1 + pc2 + hybrid_status
## Model 2: max_WL ~ pc1 + pc2
    #Df LogLik Df Chisq Pr(>Chisq)
      9 -264.86
## 2 4 -270.36 -5 10.995
                             0.05147 .
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## Call:
## lm(formula = max_WL ~ pc1 + pc2 + hybrid_status, data = lab)
## Residuals:
##
                 1Q
                      Median
                                   30
## -14.7941 -3.9487
                      0.5969
                               4.0292 13.2617
## Coefficients:
                                   Estimate Std. Error t value Pr(>|t|)
## (Intercept)
                                               1.5060 59.687 < 2e-16 ***
                                    89.8897
## pc1
                                    -0.7531
                                               0.3125 -2.410 0.01837 *
                                               0.8753 -2.189 0.03163 *
## pc2
                                    -1.9165
## hybrid_statusF0 M. m. musculus
                                     5.5814
                                               3.3176
                                                       1.682 0.09660 .
## hybrid_statusF1 hybrid
                                                        2.880 0.00517 **
                                     6.0398
                                               2.0974
## hybrid_statusF1 M. m. domesticus -0.7613
                                               2.4155 -0.315 0.75349
## hybrid_statusF1 M. m. musculus
                                     8.3437
                                               3.6664
                                                        2.276 0.02568 *
## hybrid_statusother
                                     2.2420
                                               2.3853
                                                        0.940 0.35023
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 5.955 on 76 degrees of freedom
## Multiple R-squared: 0.2153, Adjusted R-squared: 0.1431
## F-statistic: 2.979 on 7 and 76 DF, p-value: 0.008133
## [1] 547.7233
##
## lm(formula = max_WL ~ pc1 + pc2 + infection_history, data = lab)
##
## Residuals:
##
       Min
                 1Q
                     Median
                                   3Q
                                           Max
## -12.6200 -2.7393 -0.1766 3.3119
                                      8.7910
## Coefficients:
##
                                          Estimate Std. Error t value Pr(>|t|)
```

```
## (Intercept)
                                            84.3265
                                                        2.6943 31.298 < 2e-16
                                                        0.4051
                                                                2.655 0.009731
## pc1
                                             1.0756
                                                                0.747 0.457224
## pc2
                                            0.3095
                                                        0.4141
## infection_historyfalciformis_ferrisi
                                            7.3079
                                                        2.6336
                                                                2.775 0.007009
## infection_historyfalciformis_uninfected 15.1902
                                                        3.4750
                                                                4.371 4.03e-05
## infection historyferrisi falciformis
                                            -4.4071
                                                       3.0643 -1.438 0.154652
## infection historyferrisi ferrisi
                                            9.3843
                                                       2.8404
                                                                3.304 0.001479
## infection_historyferrisi_uninfected
                                            13.3682
                                                       3.4287
                                                                3.899 0.000213
## infection historyuninfected
                                            13.7948
                                                        3.9599
                                                                3.484 0.000840
## infection_historyuninfected_falciformis
                                           -8.9692
                                                       4.0447 -2.218 0.029701
## infection_historyuninfected_ferrisi
                                            -1.1569
                                                        3.3462 -0.346 0.730544
## (Intercept)
                                           ***
## pc1
                                           **
## pc2
## infection_historyfalciformis_ferrisi
## infection_historyfalciformis_uninfected ***
## infection historyferrisi falciformis
## infection_historyferrisi_ferrisi
                                           **
## infection historyferrisi uninfected
                                           ***
## infection_historyuninfected
                                           ***
## infection_historyuninfected_falciformis *
## infection_historyuninfected_ferrisi
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 4.514 on 73 degrees of freedom
## Multiple R-squared: 0.5669, Adjusted R-squared: 0.5075
## F-statistic: 9.554 on 10 and 73 DF, p-value: 5.886e-10
## [1] 503.8078
##
## Call:
## lm(formula = max_WL ~ pc1 + pc2, data = lab)
## Residuals:
      Min
                1Q Median
                                30
                                      Max
## -16.865 -2.911
                    1.123
                             4.404
                                     9.741
##
## Coefficients:
              Estimate Std. Error t value Pr(>|t|)
                           0.6719 138.277 < 2e-16 ***
## (Intercept) 92.9127
## pc1
               -0.8623
                            0.2870 -3.004 0.00354 **
## pc2
               -0.3550
                            0.4845 -0.733 0.46583
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 6.158 on 81 degrees of freedom
## Multiple R-squared: 0.1056, Adjusted R-squared: 0.08351
## F-statistic: 4.781 on 2 and 81 DF, p-value: 0.01089
##
                     df
                              AIC
## weight lm
                      4 548.7185
## weight_lm_exp_only 4 548.7185
```

```
weight_lm5 <- lm(max_WL ~ hybrid_status, data = lab)</pre>
summary(weight_lm5)
##
## Call:
## lm(formula = max_WL ~ hybrid_status, data = lab)
##
## Residuals:
##
                                3Q
      Min
                1Q Median
                                       Max
## -18.041 -3.038
                   1.720
                             4.178
                                     9.494
##
## Coefficients:
                                    Estimate Std. Error t value Pr(>|t|)
##
## (Intercept)
                                     92.2550
                                                 1.2264 75.224
                                                                  <2e-16 ***
## hybrid_statusF0 M. m. musculus
                                     -0.7685
                                                 2.0730 -0.371
                                                                  0.7119
## hybrid_statusF1 hybrid
                                      4.1652
                                                 1.9505
                                                         2.135 0.0359 *
## hybrid_statusF1 M. m. domesticus -0.4312
                                                 2.5283 -0.171
                                                                  0.8650
## hybrid_statusF1 M. m. musculus
                                                 2.8323
                                     3.5654
                                                         1.259
                                                                 0.2118
## hybrid_statusother
                                     -1.7492
                                                 2.1242 -0.823
                                                                 0.4128
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 6.253 on 78 degrees of freedom
## Multiple R-squared: 0.1119, Adjusted R-squared: 0.05499
## F-statistic: 1.966 on 5 and 78 DF, p-value: 0.09301
AIC(weight_lm5)
## [1] 554.1228
repeating the heatmap on the now imputed data
 # 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))</pre>
 # turn the first row into column names
 facs mouse %>%
     row_to_names(row_number = 1) -> heatmap_data
heatmap_data <- as.data.frame(heatmap_data)</pre>
table(rowSums(is.na(heatmap_data)) == nrow(heatmap_data))
##
## FALSE
##
# 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>
 # remove columns with only NAs
heatmap_data <- Filter(function(x)!all(is.na(x)), heatmap_data)</pre>
```

#remove rows with only Nas

Heatmap on facs expression data:

