Statistical Analysis of Pathobiochemical Signatures in Bile Duct Ligated Mice

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Introduction

This document contains the statistical analysis performed in the publication *Pathobiochemical signatures of cholestatic liver disease in bile duct liquted mice*.

A comprehensive data set of serum markers, histological parameters and transcript profiles was compiled at 8 time points after bile duct ligation (BDL) in mice, comprising different stages of the disease. The data set consists of $N_r = 5$ repeats ($N_r = 3$ for the measured antibodies) for $N_t = 8$ time points denoted by $t_1, ..., t_{N_t}$ consisting of a total $N_f = 154$ measured parameters in the following referred to as factors (Fluidigm gene expression, antibodies, serum markers, histological measurements).

The main steps of the analysis comprise

- Explorative data anaysis
- Dimension reduction via ANOVA
- Correlation analysis based on time course correlation measure
- · Hierarchical clustering
- Decision trees for prediction

The complete data set, source code and documentation of this analysis is available from https://github.com/matthiaskoenig/bdl-analysis .

The following naming conventions are used

- factor: one of the measured quantities/parameters over time, i.e. either
 - gene expression of a single gene (e.g. Actb);
 - one of the biomarkers (e.g. ALT, albumin, bilirubin)
 - one of the histological markers (e.g. BrdU-positive Kupffer cells)
 - one of the antibodies (e.g. CTGF, S100A4)
- time point: a single value t_i from the measured time points 0h (control), 6h, 12h, 18h, 30h, 2d, 5d, 14d
- sample: one of the $N_t \cdot N_r = 40$ mice, i.e. a one of the repeats for a given time point

The results of the analysis are written to the results directory defined via the BDL_RESULTS environment variable. To reproduce this analysis create the respective variable.

```
# read results directory from environment variable
resultsPath <- Sys.getenv("BDL_RESULTS")
if (identical(resultsPath, "")){
   stop("No results directory defined, set the BDL_RESULTS environment variable")
}
# create directory to store data sets of analysis
dir.create(file.path(resultsPath, 'data'), showWarnings=FALSE)</pre>
```

Explorative data analysis

Data import

In a first step the processed data set is loaded from the data folder. The data consists of the time course data for all factors (BDLdata), additional information for the factors (BDLfactors), the sample definition, i.e. the assignment of sample ids to respective time point and repeat (BDLsamples), and a mapping of the Fluidigm (gene) probe ids to UniProt identifiers (BDLprobes).

```
suppressPackageStartupMessages(library(BDLanalysis))
suppressPackageStartupMessages(library(calibrate))
suppressPackageStartupMessages(library(pander))
dir.create(file.path(resultsPath, 'control'), showWarnings=FALSE)
# path definition
baseLoc <- system.file(package="BDLanalysis")</pre>
extPath <- file.path(baseLoc, "extdata")</pre>
# load data
data(BDLdata)
data(BDLsamples)
data(BDLfactors)
data(BDLprobes)
# counters
Nr <- 5 # repeats
Nt <- length(levels(BDLsamples$time_fac)) # time points</pre>
# store all data sets in the results folder
save(BDLdata, file=file.path(resultsPath, "data", "BDLdata.Rdata"))
save(BDLsamples, file=file.path(resultsPath, "data", "BDLsamples.Rdata"))
save(BDLfactors, file=file.path(resultsPath, "data", "BDLfactors.Rdata"))
save(BDLprobes, file=file.path(resultsPath, "data", "BDLprobes.Rdata"))
```

In addition to the individual data points per time point, the mean data averaged over the N_r repeats per time points is used in parts of the correlation analysis. This mean factor data set is calculated once at the beginning via

```
BDLmean <- bdl_mean_data(BDLdata, BDLsamples)
BDLmean.time <- as.numeric(levels(as.factor(BDLsamples$time)))</pre>
```

In total 153 factors were measured in the this BDL study falling in the categories: Antibodies, Biochemistry, GE_ADME, GE_Cytokines, GE_Fibrosis, Histology. The majority of factors belongs herby to the 3 fluidigm chips with 47 probes per chip.

An overview of the number of factors per category is provided in the following table

```
cat_table <- as.data.frame(table(BDLfactors$ftype))
colnames(cat_table) <- c("Category", "Freq")
set.caption(sub(".", " ", "Factors per category", fixed = TRUE))
pander(cat_table)</pre>
```

Table 1: Factors per category

Category	Freq
Antibodies	3

Category	Freq
Biochemistry	4
GE_ADME	47
$GE_Cytokines$	47
$GE_Fibrosis$	46
Histology	6

```
rm(cat_table)
```

Data visualization

Time course of single factors

In a first step overview plots of the raw and mean data for all individual factors are generated. These are available in the resultsPath/factors folder

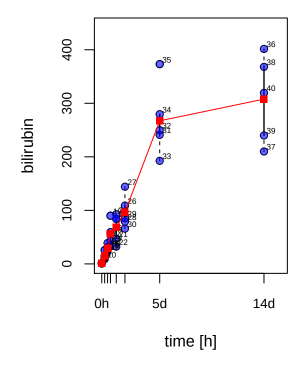
```
# Create figures for all factors
factors_path <- file.path(resultsPath, 'factors')
dir.create(factors_path, showWarnings=FALSE)
plot_all_factors(path=factors_path)
rm(factors_path)</pre>
```

The example for a single factor is depicted below, here for the factor bilirubin.

```
plot_single_factor('bilirubin', path=NULL)
```



bilirubin (Biochemistry)



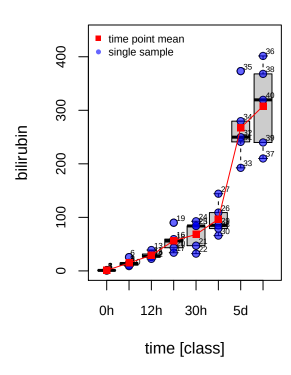


Figure Single factor: Plot of the raw time course data for a single factor, here for bilirubin. On the left the data is plotted against the time [h], on the right against the different time classes. Individual data points are depecticed in blue with the respective sample number shown next to the data points. The mean averaged of the repeats per time point are depicted in red. Box-and-whisker plots were added with default R parameters of boxwex=0.8, staplewex=0.5, outwex=0.5.

Time course of all factors (Heatmap)

In a next step the heatmap of the full data set was generated, i.e. of all time points and repeats. This provides a first overview over the complete data set. Rows correspond to the individual factors (factor order corresponding to the original data set: GE_ADME, GE_Cytokines, GE_Fibrosis, Biochemistry, Histology, Antibodies). Columns correspond to the 40 samples with 5 subsequent samples belonging to one of the 8 time points (with from left to right: 0h, 6h, 12h, 18h, 30h, 2d, 5d, 14d). The data is row scaled, i.e. every individual factor is scaled to have mean zero and standard deviation one.

```
suppressPackageStartupMessages(library(gplots))
suppressPackageStartupMessages(library("RColorBrewer"))
# define horizontal and vertical helper lines
v_{lines} <- ((1:Nt)*Nr+0.5)
factor_types <- c("Antibodies", "Histology", "Biochemistry", "GE_Fibrosis", "GE_Cytokines", "GE_ADME")
factor_table <- table(BDLfactors$ftype)</pre>
h_lines <- 0.5 + cumsum(factor_table[factor_types])</pre>
timecourse_heatmap <- function(){</pre>
  # create better row names
  dtmp <- BDLdata
  rownames(dtmp) <- paste(rownames(BDLsamples), BDLsamples$time_fac, sep=" ")
  # heatmap colors
  hmap_colors <- HeatmapColors()</pre>
  # colors for factor groups
  colorset <- brewer.pal(length(factor_types), "Set2")</pre>
  color.map <- function(factor_id) {return(colorset[ which(factor_types==BDLfactors$ftype[which(BDLfact</pre>
  factorColors <- unlist(lapply(BDLfactors$id, color.map))</pre>
  # heatmap
  heatmap.2(t(as.matrix(dtmp)), col=hmap_colors(100), scale="row", dendrogram="none", Rowv=NULL, Colv=N
            key=TRUE, trace="none", cexRow=0.5, keysize=0.8, density.info="none",
            RowSideColors=factorColors,
            add.expr=abline(v=v_lines, h=h_lines, col="black", lwd=0.5),
            main="Heatmap of BDL time course data")
            # xlab="sample", ylab="factor")
  # legend
  legend("left",
      inset=c(-0.03,0),
      legend = rev(factor_types), # category labels
      col = rev(colorset), # color key
      lty= 1, lwd = 10, cex = 0.7, bty="n"
  )
}
# plot to file
pdf(file.path(resultsPath, "control", "timecourse_heatmap.pdf"), width=10, height=10, pointsize=12)
timecourse_heatmap()
invisible(dev.off())
```

```
timecourse_heatmap()
```

```
# cleanup
rm(factor_table, h_lines, v_lines)
```

Figure All factors: Heatmap of the complete data set, i.e. all factors and repeats over time. The data is row scaled, i.e. every individual factor is scaled to have mean zero and standard deviation one, with positive Z-score in blue, negative Z-score in red. The row order is according to the factor categories, the order within the fluidigm chips according to the order of the probes on the chip. the respective categories are depicted on the left.

Results: Various patterns are visible in the plotted raw data:

- Two main classes of response can be observed. One class with an increase in the early phase up to 6h after BDL (many of the ADME genes fall into this class) and a second class increasing in the later stage after 2-5 days after BDL. Many of the genes on the Cytokines and Fibrosis chips as well as some of the biochemical, histological and antibody fall in this second class.
- The individual animals show heterogeneous responses to BDL. Within one time point the 5 repeats can show very different patterns. For instance at time 6h after BDL 3/5 of the mice show a marked increase in the ADME genes, whereas 2/5 do not show such a marked increase. Another example is the mice sample 27 at time 2d, with a high increase in the genes on the Fibrosis chip, which is not observed in the other 4 samples at time 2d.

Actb quality control

Actb (Actin, cytoplasmic 1) probes were included on all Fluidigm chips (GE_ADME, GE_Cytokines, GE_Fibrosis) and not used in the normalization of the gene expression data. Hence, ActB can serve as quality control for the technical reproducibility of the Fluidigm chips. If the data is reproducible between chips the pairwise correlation between all individual Actb measurements should have high correlation coefficients close to 1. Plotting the data of the Actb measurements of two chips against each other should lie on a straight line

```
# Actb control figure
plot_actb_control <- function(){</pre>
  par(mfrow=c(2,3))
  plot_single("Actb")
  plot_single("Actb.x")
  plot_single("Actb.y")
  plot_cor_pair("Actb", "Actb.x", single_plots=FALSE)
  plot_cor_pair("Actb", "Actb.y", single_plots=FALSE)
  plot_cor_pair("Actb.x", "Actb.y", single_plots=FALSE)
  par(mfrow=c(1,1))
}
# plot to file
pdf(file.path(resultsPath, "control", "Actb_control.pdf"), width=10, height=6, pointsize=12)
plot_actb_control()
invisible(dev.off())
# calculate Spearman and Pearson correlation coefficients on N=8*5=40 data points
actb.spearman <- cor(data.frame(Actb=BDLdata$Actb,</pre>
```


Heatmap of BDL time course data

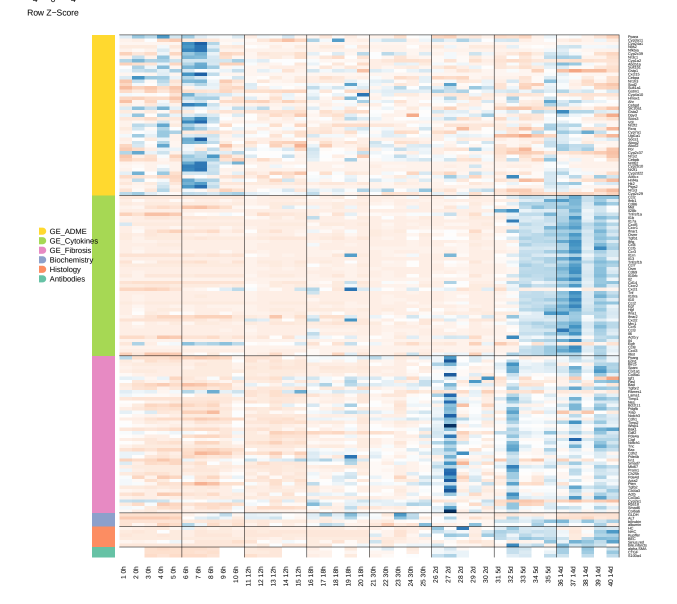


Figure 1:

Table 2: Spearman correlation of Actb controls

	Actb	Actb.x	Actb.y
Actb	1	0.908	0.944
Actb.x	0.908	1	0.917
$\mathbf{Actb.y}$	0.944	0.917	1

```
set.caption(sub(".", " ", "Pearson correlation of Actb controls", fixed = TRUE))
pander(round(actb.pearson, digits=3))
```

Table 3: Pearson correlation of Actb controls

	Actb	Actb.x	Actb.y
Actb	1	0.945	0.938
Actb.x	0.945	1	0.935
Actb.y	0.938	0.935	1

```
plot_actb_control()
```

```
rm(actb.pearson, actb.spearman)
```

Figure Actb Control: Correlation plot of the Actb probes from the 3 Fluidigm chips: Actb (fibrosis), Actb.x (ADME), Actb.y (Cytokines). The top row shows the individual time courses, the bottom row the pair wise plot of individual data points.

Results: The Actb Fluidigm gene expression measurements are highly reproducible for the measured chips, with Spearman as well as Pearson correlation coefficients all > 0.9 for pairwise Actb comparison.

Dimension reduction

Introduction

A one-way analysis of variance (ANOVA) was applied to reduce the factors to the subset showing significant $(p_{adjusted} < 0.05)$ changes during the time course. In its simplest form, ANOVA provides a statistical test of whether or not the means of several groups are equal, and therefore generalizes the t-test to more than two groups, with the groups being the sampled time points. The Holm's procedure was used to correct the p-values for any artificial p-value inflation due to multiple testing.

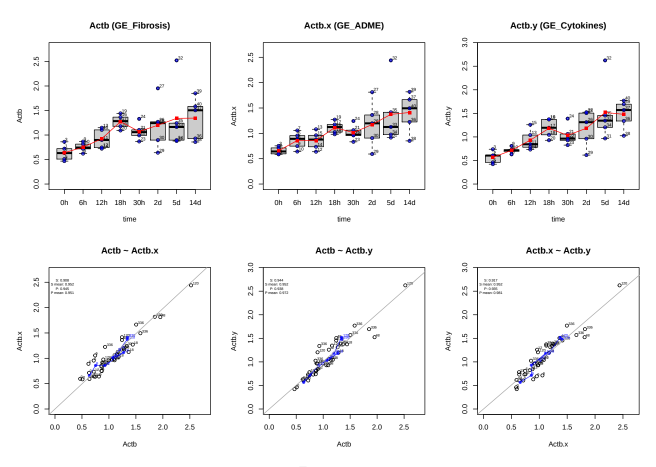


Figure 2:

For every of the individual factors in the BDL data set an ANOVA was calculated. Dimension reduction of the BDL data set was than performed by filtering out factors which did not significantly changing over time.

The BDLdata data set is reshaped into matrix format for the ANOVA calculation, with time points in rows and repeats as columns for every factor.

```
BDLmatrices <- bdl_matrix_data(BDLdata, BDLsamples)
```

ANOVA for single factor

The following shows the ANOVA calculation for a single factor, here for bilirubin.

```
# example ANOVA for one factor
  mat.anova <- t(BDLmatrices[['bilirubin']])</pre>
  colnames(mat.anova) <- levels(BDLsamples$time_fac)</pre>
  # concatenate the data rows of df1 into a single vector r .
  r = c(t(as.matrix(mat.anova))) # response data
  # assign new variables for the treatment levels and number of observations.
  f = levels(BDLsamples$time_fac) # treatment levels
  k = 8
                                  # number of treatment levels
 n = 5
                                  # observations per treatment
  # create a vector of treatment factors that corresponds to each element of r in step 3 with the gl fu
  tm <- gl(k, 1, n*k, factor(f)) # matching treatments</pre>
  # apply the function aov to a formula that describes the response r by the treatment factor tm.
  # fit an analysis of variance model
  av \leftarrow aov(r \sim tm)
  # print out the ANOVA table with the summary function.
  summary(av)
               Df Sum Sq Mean Sq F value
##
                                            Pr(>F)
## tm
                7 479470
                           68496
                                       41 3.27e-14 ***
               32 53463
                             1671
## Residuals
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
 # print the corresponding p-value
 p.value <- summary(av)[[1]][["Pr(>F)"]][[1]]
  # show data matrix
 print(mat.anova)
##
        0h
              6h
                   12h
                         18h
                                30h
                                        2d
                                              5d
                                                    14d
## R1 1.49 25.80 31.51 59.40 47.11 108.95 241.0 401.5
## R2 0.93 13.87 22.51 33.95 32.26 144.20 249.8 210.0
## R3 1.65 10.70 38.99 55.80 83.98 79.10 192.5 368.0
```

```
# cleanup
rm(r, f, k, n, tm, av, p.value, mat.anova)
```

ANOVA for all factors

Analog to the single factor ANOVA, the ANOVA is performed on all the factors. Hereby, a multitude of tests are performed, namely an ANOVA for every single factor. Consequently, the reported p-values of the ANOVA have to be adjusted via multiple testing procedures. Using the p.adjust function which given a set of p-values, returns p-values adjusted using one of several methods. The Bonferroni, Holm, Hochberg, Hommel are designed to give strong control of the family-wise error rate. We used the Holm's method for adjustment (Holm, S. (1979). A simple sequentially rejective multiple test procedure. Scandinavian Journal of Statistics 6, 65-70.).

```
# Calculation of ANOVA for all factors
df.anova <- all_factor_anova()
df.anova$sig <- sapply(df.anova$p.value, significant_code)  # add significant codes

df.anova$p.holm <- p.adjust(df.anova$p.value, method ="holm", n = length(df.anova$p.value))
df.anova$sig.holm <- sapply(df.anova$p.holm, significant_code)
df.anova$ftype <- BDLfactors$ftype
df.anova$fshort <- BDLfactors$ftype.short

# order factors by adjusted p-values, and cleanup for printing
df.anova.ordered <- df.anova[with(df.anova, order(p.holm)), ]
rownames(df.anova.ordered) <- df.anova.ordered$factors

df.anova.ordered <- df.anova.ordered[c("p.holm", "sig.holm", "ftype", "fshort")]
df.anova.ordered$p.holm <- as.numeric(sprintf("%1.2E", df.anova.ordered$p.holm))
print(df.anova.ordered)</pre>
```

```
##
                   p.holm sig.holm
                                            ftype fshort
                 2.93e-14
## Cyp1a2
                                ***
                                         GE ADME
## bilirubin
                 4.98e-12
                                *** Biochemistry
                                                       В
## Il10rb
                                *** GE_Cytokines
                 1.15e-11
## Tgfb1
                 3.30e-11
                                *** GE_Cytokines
## Cc12
                 3.46e-11
                                *** GE Cytokines
                                *** GE Cytokines
## Cd86
                 6.56e-11
                                *** GE_Cytokines
## Ccr2
                 6.94e-11
                                *** GE_Cytokines
## Mrc1
                 6.95e-11
## Tnfrsf1b
                                *** GE_Cytokines
                 7.89e-11
## Cxc15
                 6.26e-10
                                *** GE_Cytokines
## CTGF
                 7.82e-10
                                      Antibodies
                                                       Α
## Il10ra
                 1.64e-09
                                *** GE_Cytokines
## Gstm1
                 9.18e-09
                                          GE\_ADME
## Cc17
                 3.40e-08
                                *** GE_Cytokines
## Ccr5
                 4.36e-08
                                *** GE_Cytokines
## Hgf
                 5.80e-08
                                *** GE_Cytokines
## Osmr
                 1.01e-07
                                *** GE Cytokines
## Cc14
                 1.04e-07
                                *** GE_Cytokines
## NrOb2
                 1.30e-07
                                          GE ADME
                                     GE_Fibrosis
## Tgfbr2
                 1.71e-07
                                ***
## BEC
                 2.09e-07
                                       Histology
                                                       Η
```

```
## Cc15
                  2.78e-07
                                 *** GE_Cytokines
## Col1a1
                  4.40e-07
                                      GE_Fibrosis
## Ifnar1
                  7.71e-07
                                     GE_Cytokines
## S100a4
                  9.19e-07
                                        Antibodies
                                                         Α
## Sparc
                  1.07e-06
                                      GE_Fibrosis
                                      GE Fibrosis
## Cyp2e1
                  1.50e-06
## Cxcr2
                  1.75e-06
                                     GE_Cytokines
## Ccr3
                  1.91e-06
                                     GE_Cytokines
## Cd69
                  2.73e-06
                                 *** GE_Cytokines
## Cyp2c29
                  2.93e-06
                                 ***
                                           GE_ADME
## Gsta2
                  3.88e-06
                                           GE_ADME
## Tnf
                  4.36e-06
                                 ***
                                     GE_Cytokines
## Gdf2
                                      GE_Fibrosis
                  5.95e-06
                                 ***
                                 *** GE_Cytokines
## Il1b
                  7.29e-06
## Ifng
                  7.51e-06
                                 *** GE_Cytokines
## Osm
                  7.51e-06
                                 *** GE_Cytokines
## Cc13
                  8.53e-06
                                 *** GE_Cytokines
## Il13
                  9.29e-06
                                     GE_Cytokines
## Cxcr1
                  1.07e-05
                                     GE_Cytokines
## Cyp2c37
                  1.09e-05
                                           GE_ADME
## Cd14
                  1.15e-05
                                 *** GE_Cytokines
## Col3a1
                  2.02e-05
                                      GE_Fibrosis
## Tnfrsf1a
                  3.33e-05
                                 *** GE_Cytokines
## I12
                  4.84e-05
                                 *** GE_Cytokines
## Ifnb1
                  4.86e-05
                                 *** GE_Cytokines
## Egf
                  4.88e-05
                                 *** GE_Cytokines
## I128b
                  4.88e-05
                                 *** GE_Cytokines
## Il10
                  4.88e-05
                                 *** GE_Cytokines
## I14
                  4.88e-05
                                     GE_Cytokines
## Slc10a1
                                           GE_ADME
                  5.26e-05
                                 ***
## Timp2
                  6.24e-05
                                      GE_Fibrosis
## Cxcl3
                  6.82e-05
                                 *** GE_Cytokines
## Cc18
                  1.23e-04
                                     GE_Cytokines
## Ctgf
                                      GE_Fibrosis
                  1.32e-04
                                 ***
                  1.38e-04
                                           GE\_ADME
## Gstp1
## Ppara
                                           GE_ADME
                  1.63e-04
                                 ***
## Ifnar2
                  1.82e-04
                                     GE Cytokines
## I16
                  2.23e-04
                                     GE_Cytokines
## Il17a
                                     GE_Cytokines
                  2.43e-04
## Bad
                  3.84e-04
                                      GE_Fibrosis
                                 ***
## Timp1
                  4.30e-04
                                      GE_Fibrosis
## Cdh1
                  4.48e-04
                                 ***
                                      GE_Fibrosis
## Cebpa
                  4.93e-04
                                 ***
                                           GE_ADME
## alpha.SMA
                  5.02e-04
                                 ***
                                        Antibodies
                                                         A
## NHC
                  5.28e-04
                                         Histology
                                                         Η
                                 ***
## Cdh2
                  5.38e-04
                                 ***
                                      GE_Fibrosis
## Sirius.red
                  7.59e-04
                                 ***
                                         Histology
                                                         Η
## Pdgfb
                  7.68e-04
                                 ***
                                      GE_Fibrosis
## Il6st
                  8.71e-04
                                     GE_Cytokines
                                 ***
## Fn1
                  1.21e-03
                                      GE_Fibrosis
## Mki67
                  1.46e-03
                                      GE_Fibrosis
## Ifna1
                  1.46e-03
                                     GE Cytokines
## Egfr
                                     GE_Cytokines
                  1.59e-03
## Kupffer
                  1.95e-03
                                         Histology
                                                         Η
```

```
## Tnc
                  2.04e-03
                                      GE_Fibrosis
## Ugt1a1
                  2.75e-03
                                          GE_ADME
                                  **
## Sult1a1
                  2.98e-03
                                          GE ADME
## GLDH
                                                        В
                  4.28e-03
                                     Biochemistry
## Notch1
                  4.45e-03
                                      GE_Fibrosis
## Met
                  4.97e-03
                                     GE_Cytokines
                                          GE ADME
## Cyp7a1
                  9.83e-03
## Cyp24a1
                  9.88e-03
                                          GE_ADME
## Tgfb2
                  1.08e-02
                                      GE_Fibrosis
## Birc5
                  1.69e-02
                                      GE_Fibrosis
## Actb.y
                  1.91e-02
                                     GE_Cytokines
## Bak1
                  2.73e-02
                                      GE_Fibrosis
## Bax
                  2.89e-02
                                      GE_Fibrosis
## Rarres1
                  3.57e-02
                                      GE_Fibrosis
## bile.infarcts 3.78e-02
                                                         Н
                                        Histology
## Cyp3a11
                  5.30e-02
                                           GE_ADME
                  5.74e-02
                                           GE\_ADME
## Sult1b1
## Cyp4a10
                  6.79e-02
                                           GE\_ADME
                  6.79e-02
                                      GE_Fibrosis
## Pparg
## Hk2
                  7.50e-02
                                           GE ADME
## ALT
                  8.79e-02
                                   . Biochemistry
                                                         В
## Smad6
                                      GE_Fibrosis
                  9.49e-02
                                           GE_ADME
## Cyp2b10
                  1.17e-01
## HC
                  1.27e-01
                                        Histology
                                                         Η
## Il1rn
                  1.35e-01
                                     GE_Cytokines
## Nes
                  1.37e-01
                                      GE_Fibrosis
## Nfkbia
                                           GE_ADME
                  1.57e-01
## Rps18
                  1.68e-01
                                      GE_Fibrosis
## Cxcl15
                  2.18e-01
                                           GE_ADME
                  2.43e-01
## Socs3
                                           GE_ADME
## Vdr
                  2.43e-01
                                           GE_ADME
## Edn1
                  2.43e-01
                                      GE_Fibrosis
## Nr2f1
                  2.63e-01
                                           GE_ADME
                                           GE_ADME
## Abcg2
                  2.82e-01
## Nr1i3
                  2.82e-01
                                           GE\_ADME
## Hnf4a
                  2.88e-01
                                           GE_ADME
## Ptgs2
                  3.43e-01
                                           GE ADME
## Bcl2l11
                  3.43e-01
                                      GE_Fibrosis
## Socs1
                  3.46e-01
                                           GE_ADME
## Pten
                  3.66e-01
                                      GE_Fibrosis
                                          GE_ADME
## Actb.x
                  3.90e-01
## Cxc12
                                     GE_Cytokines
                  4.06e-01
## Xiap
                  4.06e-01
                                      GE_Fibrosis
## Pde4a
                  4.42e-01
                                      GE_Fibrosis
## Dpyd
                  4.43e-01
                                           GE_ADME
                                     GE_Cytokines
## Cxcl1
                  4.43e-01
## Lama1
                  4.57e-01
                                      GE_Fibrosis
## Col8a1
                  5.07e-01
                                      GE_Fibrosis
## Prom1
                  5.54e-01
                                      GE_Fibrosis
## Actb
                  5.54e-01
                                      GE_Fibrosis
## Ahr
                  5.57e-01
                                          GE_ADME
## Nr2f2
                  5.57e-01
                                           GE_ADME
## Nos2
                  5.82e-01
                                           GE_ADME
## Notch3
                  7.26e-01
                                      GE_Fibrosis
```

```
## Cebpd
                  7.29e-01
                                          GE_ADME
## Hmox1
                  7.69e-01
                                          GE_ADME
## Cyp2d22
                  8.36e-01
                                          GE ADME
## Igf1
                  9.78e-01
                                      GE_Fibrosis
## Fasl
                  9.78e-01
                                      GE_Fibrosis
## Cyp2c39
                                          GE ADME
                  1.00e+00
## Nr3c1
                                          GE ADME
                  1.00e+00
                                          GE_ADME
## Abcb1a
                  1.00e+00
## Nr1h3
                  1.00e+00
                                          GE_ADME
## Sod2
                  1.00e+00
                                          GE_ADME
## Rxra
                  1.00e+00
                                          GE_ADME
                                          GE\_ADME
## Abcc2
                  1.00e+00
## Por
                  1.00e+00
                                          GE\_ADME
## Nr1i2
                  1.00e+00
                                          GE_ADME
## Cebpb
                                          GE_ADME
                  1.00e+00
## Wisp1
                  1.00e+00
                                      GE_Fibrosis
## Pde4b
                  1.00e+00
                                      GE_Fibrosis
## Smad7
                  1.00e+00
                                      GE Fibrosis
## Ch25h
                  1.00e+00
                                      GE_Fibrosis
## Pde4d
                  1.00e+00
                                      GE Fibrosis
## Acta2
                  1.00e+00
                                      GE_Fibrosis
## Col4a3
                  1.00e+00
                                      GE_Fibrosis
## Col6a6
                                      GE_Fibrosis
                  1.00e+00
## albumin
                  1.00e+00
                                     Biochemistry
                                                        В
```

```
# save results
write.table(df.anova.ordered, file=file.path(resultsPath, "data", 'BDLanova.csv'), sep="\t", quote=FALS.
save(df.anova, file=file.path(resultsPath, "data", "BDLanova.Rdata"))
# rm(df.anova.ordered)
```

Filter factors

The factors are filtered based on the respective acceptance level, with the cutoff for the adjusted p-value being p_{accept} , i.e. all factors with a ANOVA with $p_{adjusted} \ge p_{accept}$ are filtered out. The filtered raw data is available as BDLdata.fil, the filtered mean data set as BDLmean.fil. All subsequent analyses are performed on the filtered data set, which is depicted in the following heatmap

```
p.accept = 0.05  # acceptance level
anova.accept = (df.anova$p.holm < p.accept)  # accepted subset
# subset of filtered data
BDLdata.fil <- BDLdata[, anova.accept]
BDLmean.fil <- BDLdata[, anova.accept]

# accepted
table(anova.accept)  # 64 rejected / 90 accepted (adjusted)

## anova.accept
## FALSE TRUE
## 63 90</pre>
```

```
# which factors were accepted in the various categories
fil_tab <- data.frame(</pre>
  table(BDLfactors$ftype[anova.accept]),
 table(BDLfactors$ftype),
  round(table(BDLfactors$ftype[anova.accept])/table(BDLfactors$ftype), 2)
fil_tab <- fil_tab[, c('Var1', 'Freq', 'Freq.1', 'Freq.2')]</pre>
names(fil_tab) <- c('Category', 'Accepted', 'All', 'Percent')</pre>
# overview of filtered factors
print(fil_tab)
##
         Category Accepted All Percent
## 1
       Antibodies
                        3
                             3
                                  1.00
                        2
## 2 Biochemistry
                            4
                                  0.50
## 3
          GE_ADME
                        14 47
                                  0.30
## 4 GE_Cytokines
                        44 47
                                  0.94
## 5 GE_Fibrosis
                        22 46
                                  0.48
## 6
       Histology
                        5 6
                                  0.83
rm(fil_tab)
```

Based on the adjusted p-values the data set was reduced from original {r} ncol(BDLdata.fil) factors to 90. Almost all Cytokines genes (inflammation panel) were retained in the data set whereas many of the ADME and Fibrosis genes are filtered.

Heatmap of filtered time course data

The heatmap of the filtered raw data is depicted below

```
v_{lines} \leftarrow ((1:Nt)*Nr+0.5)
timecourse_heatmap_filtered <- function(){</pre>
  # prepare data with row names
  dtmp <- BDLdata.fil</pre>
  rownames(dtmp) <- paste(rownames(BDLsamples), BDLsamples$time_fac, sep=" ")</pre>
  # color definitions
  hmap_colors <- HeatmapColors()</pre>
  colorset <- brewer.pal(length(factor_types), "Set2")</pre>
  color.map <- function(factor_id) {return(colorset[ which(factor_types==BDLfactors$ftype[which(BDLfact</pre>
  factorColors <- unlist(lapply(colnames(BDLdata.fil), color.map))</pre>
  # heatmap
  heatmap.2(t(as.matrix(dtmp)), col=hmap_colors(100), scale="row", dendrogram="none",
             Rowv=NULL, Colv=NULL,
            key=TRUE, trace="none", cexRow=0.5, keysize=0.8, density.info="none",
             RowSideColors=factorColors,
             add.expr=abline(v=v_lines, col="black", lwd=0.5),
             main="ANOVA Filtered BDL factors")
             # xlab="sample", ylab="factor")
  # legend
  legend("left",
      inset=c(-0.03,0),
```

```
legend = rev(factor_types), # category labels
col = rev(colorset), # color key
lty= 1, # line style
lwd = 10, # line width
cex = 0.7,
bty="n"
)
}

# plot to file
pdf(file.path(resultsPath, "control", "timecourse_heatmap_filtered.pdf"),
    width=10, height=10, pointsize=12)
timecourse_heatmap_filtered()
invisible(dev.off())
```

```
# plot to report
timecourse_heatmap_filtered()
```

```
rm(v_lines)
```

Figure Timcourse Filtered: Plot of ANOVA filtered data set.

t-test for initial phase

A t-test was performed to find significantly changed factors in the initial phase after BDL.

```
# t-test for the initial phase changes
calculate_inital_phase_changes <- function(){</pre>
  # empty vectors
  p.t_test <- rep(NA, ncol(BDLdata))</pre>
  names(p.t_test) <- colnames(BDLdata)</pre>
  up_down <- rep(NA, ncol(BDLdata))
  names(up_down) <- colnames(BDLdata)</pre>
  for (name in colnames(BDLdata)){
    # data for control (0h) and initial response (6h)
    d0 <- BDLdata[BDLsamples$time_fac == "Oh", name]</pre>
    d6 <- BDLdata[BDLsamples$time_fac == "6h", name]</pre>
    # remove NA
    d0 <- d0[!is.na(d0)]</pre>
    d6 <- d6[!is.na(d6)]
    # unpaired two.sided t-test
    t.test.res <- t.test(d0, d6, alternative="two.sided", var.equal=FALSE)</pre>
    p.t_test[name] <- t.test.res$p.value</pre>
    # going up or down
    up_down[name] <- "-"</pre>
    if (mean(d6) > mean(d0)){
      up_down[name] <- "up"</pre>
    } else if (mean(d6) < mean(d0)){</pre>
      up_down[name] <- "down"</pre>
```

Color Key

ANOVA Filtered BDL factors



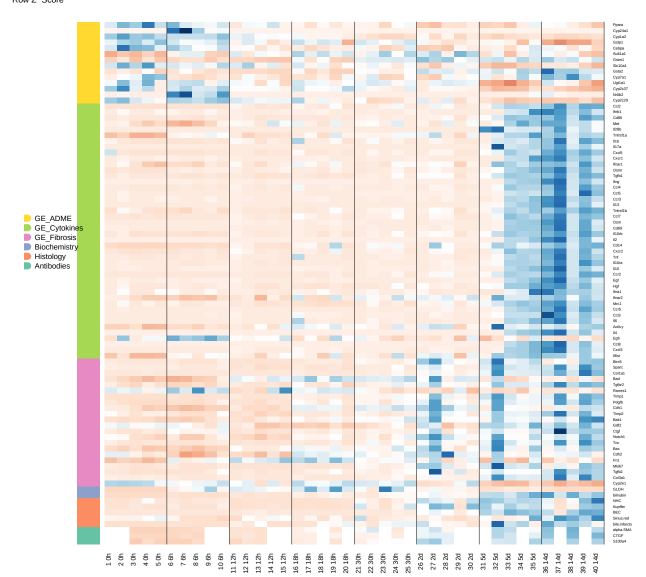


Figure 3:

```
}
  # results data.frame
  p.df <- data.frame(p.value=p.t_test, up_down=up_down)</pre>
  p.df$sig <- sapply(p.df$p.value, significant_code)</pre>
  rownames(p.df) <- colnames(BDLdata)</pre>
  # sort by p.value
  p.df.ordered <- p.df[order(p.df$p.value),]</pre>
  return(p.df.ordered)
}
# top up and down regulated in initial phase
p.df.ordered <- calculate_inital_phase_changes()</pre>
p.df.up <- p.df.ordered[p.df.ordered$p.value<0.05 & p.df.ordered$up_down=="up", ]
p.df.down <- p.df.ordered[p.df.ordered$p.value<0.05 & p.df.ordered$up_down=="down", ]
# top up
print(p.df.up)
##
                  p.value up_down sig
## Tnfrsf1a 2.418075e-05
                                up ***
## Il6st
            4.777944e-04
                                up ***
## Osmr
             5.635117e-04
                                up ***
             2.057930e-03
## Fn1
                                up
                                    **
            4.500421e-03
## Cd14
                                    **
                                up
## ALT
             5.488831e-03
                                    **
## bilirubin 9.221295e-03
                                up
                                    **
## NrOb2
           9.693272e-03
                                up
                                    **
## Ctgf
             1.026407e-02
                                up
           1.159292e-02
## Cxcl1
## Timp1
             1.196340e-02
                                up
## Egfr
             1.444607e-02
                                up
## Cyp4a10 2.288889e-02
                                up
## Cxcl2
             2.507493e-02
                                up
## GLDH
             2.896681e-02
                                up
## Hmox1
             2.913971e-02
                                up
## Socs3
             3.604221e-02
                                up
## Sult1a1
             3.632014e-02
                                up
# top down
print(p.df.down)
##
                  p.value up_down sig
## Cdh2
              0.005994558
                              down
## Pde4a
              0.010806348
                              down
## Sirius.red 0.014252420
                              down
## Pten
              0.014853870
                              down
              0.017104924
## Col3a1
                              down
## Nes
              0.026669560
                              down
## I128b
              0.027571476
                              down
## Cdh1
              0.027687939
                              down
```

```
## Tgfb1 0.032921400 down *
## Il10ra 0.035845273 down *
## Cyp2e1 0.038398167 down *
## Cyp7a1 0.038785457 down *
## Xiap 0.041507273 down *
```

Correlation analysis

Introduction

For the correlation analysis between factors and the subsequent cluster analysis a correlation measure for time series data {Son2008} in combination with Complete-Linkage hierarchical clustering was used. This combination of methods provided the best enrichments on gene-expression time-series in a recent comparisons of methods {Jaskowiak2014, Jaskowiak2013} testing various correlation measures and clustering algorithms.

The calculation of correlation coefficients between factors i and j $(i, j = 1, ..., N_p)$ was performed using the slightly modified correlation coefficient based similarity measure developed for clustering of time-course data $(Y_{i,j}^{S2} \text{ and } Y_{i,j}^{R2})$ {Son2008}. $Y_{i,j}^{S2}$ and $Y_{i,j}^{R2}$ are linear combinations of (i) a classical correlation part based on Spearman correlation $S_{i,j}^*$ in case of $Y_{i,j}^{S2}$ or Pearson $R_{i,j}^*$ in case of $Y_{i,j}^{R2}$, (ii) a component $A_{i,j}^*$ accounting for the similarity in changes between two time courses, (iii) a component $M_{i,j}^*$ comparing the location of minimum and maximum values of the time course (see {Son2008} for definitions)

$$Y_{i,j}^{S2} = w_1 S_{i,j}^* + w_2 A_{i,j}^* + w_3 M_{i,j}^*$$

$$Y_{i,j}^{R2} = w_1 R_{i,j}^* + w_2 A_{i,j}^* + w_3 M_{i,j}^*$$

 $R_{i,j}^*$ and $S_{i,j}^*$ are hereby calculated on the individual data points for the factors i and j, $A_{i,j}^*$ and $M_{i,j}^*$ on the mean time courses averaged over the N_r repeated measurements. Throughout the analysis the following weights were used $w_1 = 0.5$, $w_2 = 0.3$, $w_3 = 0.2$.

In the calculation of the change component we used a Spearman correlation based measure (A^{**}) instead of the originally proposed Pearson measure (A^*) resulting in the correlation scores $Y_{i,j}^{S3}$ and $Y_{i,j}^{S3}$

$$Y_{i,j}^{S3} = w_1 S_{i,j}^* + w_2 A_{i,j}^{**} + w_3 M_{i,j}^*$$

$$Y_{i,j}^{R3} = w_1 R_{i,j}^* + w_2 A_{i,j}^{**} + w_3 M_{i,j}^*$$

Herein, $A_{i,j}^{**}$ calculates the correlation of changes between factors i and j based on Spearman correlation analog $A_{i,j}^{*}$ as

$$A_{i,j}^{**} = (S(d_i, d_j) + 1)/2$$

$$A_{i,j}^* = (R(d_i, d_j) + 1)/2$$

The reason for this adaption was that initial analysis showed a strong dependency of the change components on outliers.

All calculated correlation scores Y^S and Y^R are transformed from [0, 1] to [-1, 1] via

$$Y_{norm}^S = 2(Y^S - 0.5)$$

$$Y_{norm}^{R} = 2(Y^{R} - 0.5)$$

In addition to the used Y^S and Y^R correlation scores Pearson (R) and Spearman (S) correlations were calculated for comparison.

```
suppressPackageStartupMessages(library(corrplot))
dir.create(file.path(resultsPath, 'correlation'), showWarnings=FALSE)

# list of calculated correlation methods
correlation_methods <- c("pearson", "spearman", "ys1", "ys2", "ys3", "yr1", "yr2", "yr3")

# The calculated correlation matrices are stored in the `cor.matrices
# Provides simple access to the respective correlation matrix.
cor.matrices <- vector("list", length=length(correlation_methods))
names(cor.matrices) <- correlation_methods</pre>
```

Pearson & Spearman correlation

In a first step Pearson $R_{i,j}$ and Spearman $S_{i,j}$ correlation were calculated for the subset of filtered factors.

```
# Spearman and Pearson on individual data points
cor.matrices$pearson <- cor(BDLdata.fil, method="pearson", use="pairwise.complete.obs")
cor.matrices$spearman <- cor(BDLdata.fil, method="spearman", use="pairwise.complete.obs")</pre>
```

Heatmap plot of the correlation matrices with factors in original order

```
# Heatmap of correlation matrix
correlation_heatmap <- function(method){</pre>
  data <- cor.matrices[[method]]</pre>
  if(is.null(data)){
    stop("Correlation matrix does not exist for method: ", method)
  cor_colors <- HeatmapColors() # color palette for correlation (red - white - blue)
  heatmap.2(data, col=cor_colors(10), scale="none",
          key=TRUE, symkey=FALSE, trace="none", cexRow=0.8, cexCol=0.8,
          main=method,
          density.info="none", dendrogram="none",
          Rowv=NULL, Colv=NULL,
          keysize=0.8, key.xlab = method,
          #revC=TRUE,
          sepwidth=c(0.01,0.01),
          sepcolor="black",
          colsep=1:ncol(data),
          rowsep=1:nrow(data))
```

```
# Pearson correlation (no clustering)
correlation_heatmap(method="pearson")
```



pearson

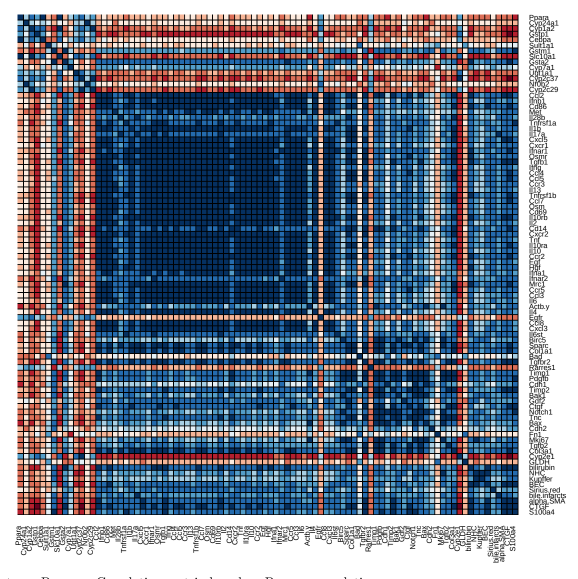


Figure Heatmap Pearson: Correlation matrix based on Pearson correlation.

```
# Spearman correlation (no clustering)
correlation_heatmap(method="spearman")
```



spearman

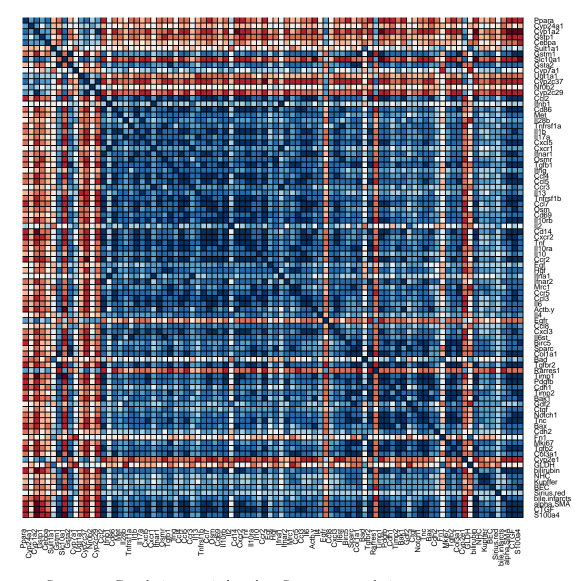


Figure Heatmap Spearman: Correlation matrix based on Spearman correlation.

The pearson correlation is highly sensitive to outliers (correlation between factors, but also in the changes between time points). Prelimary analysis of A^* showed similar problems, so we used A^{**} in Y^{S3} instead.

YS & YR correlation

Here, the time-course based correlation measurements Y^{S1} , Y^{S2} , Y^{S3} , Y^{R1} , Y^{R2} and Y^{R3} are calculated for the factors in the filtered BDL data set. To create the correlation matrix all pairwise correlations between all factors are calculated.

```
# weighting factors
w \leftarrow list(w1=0.5, w2=0.3, w3=0.2)
# calculate the YSR component matrices on the filtered data set (A, A*, A**, M, M*)
# all components are calculated on the mean data
ysr.res <- ysr.matrices(BDLmean.fil, BDLmean.time, use="pairwise.complete.obs")
# S* and R* (Pearson & Spearman correlation on individual data points)
cor.S star <- (cor.matrices$spearman + 1)/2</pre>
cor.R_star <- (cor.matrices$pearson + 1)/2</pre>
# Calculate YS and YR scores based on the components
ysr_methods <- c("ys1", "ys2", "ys3", "yr1", "yr2", "yr3")
cor.ysr <- vector("list", length(ysr_methods))</pre>
names(cor.ysr) <- ysr_methods</pre>
# unnormalized correlations in [0, 1] as combination of weighted (S/R, A/A*/A**, M/M*)
cor.ysr$ys1 <- w$w1*cor.S_star + w$w2*ysr.res$A + w$w3*ysr.res$M
cor.ysr$ys2 <- w$w1*cor.S_star + w$w2*ysr.res$A_star + w$w3*ysr.res$M_star</pre>
cor.ysr$ys3 <- w$w1*cor.S_star + w$w2*ysr.res$A_star2 + w$w3*ysr.res$M_star
cor.ysr$yr1 <- w$w1*cor.R star + w$w2*ysr.res$A</pre>
                                                       + w$w3*ysr.res$M
cor.ysr$yr2 <- w$w1*cor.R_star + w$w2*ysr.res$A_star + w$w3*ysr.res$M_star
cor.ysr$yr3 <- w$w1*cor.R_star + w$w2*ysr.res$A_star2 + w$w3*ysr.res$M_star</pre>
# scaling of ysr correlation coefficient to interval [-1,1]
for (method in ysr_methods){
  cor.matrices[[method]] <- 2*(cor.ysr[[method]]-0.5)</pre>
}
# Save correlation matrices
save(cor.matrices, file=file.path(resultsPath, "data", "cor.matrices.Rdata"))
rm(cor.ysr, cor.R_star, cor.S_star, ysr.res, ysr_methods)
```

Create heatmap plots for all calculated correlation matrices



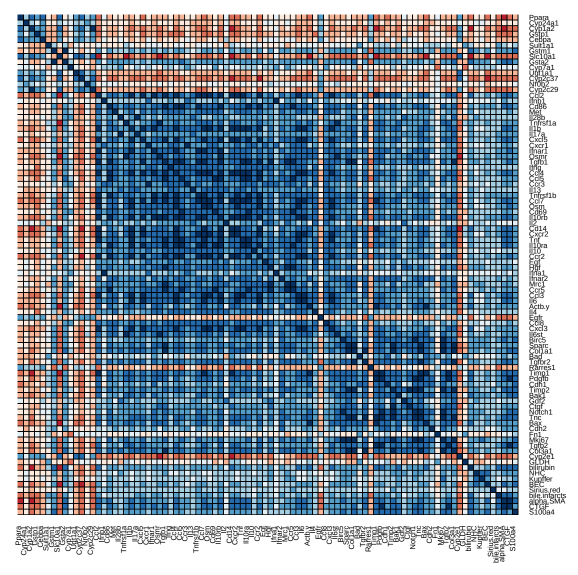


Figure Heatmap YS3: Correlation matrix based on YS3 correlation.

Hierarchical clustering

Clustering

In the next step clusters are calculated on the correlation matrices using hierarchical clustering with complete linkage. Hereby, the factors are grouped into a number of clusters with the correlation within each cluster being larger than between clusters. This effectively finds groups of factors which have similar time courses.

The complete linkage method defines the cluster distance between two clusters to be the maximum distance between their individual components.

The hierarchical clustering is now calculated based on the respective correlation matrices. Hierarchical clustering based on the given correlation matrix is applied with Ngroups=6 clusters. For every correlation measure the cluster plots are created. The number of clusters was choosen, so that at least >1 members exist per cluster.

```
hclust.res <- vector("list", length=length(correlation_methods))
names(hclust.res) <- correlation_methods</pre>
# Calculate the hierarical clustering: N clusters for correlation based on method
calculate_clusters <- function(cor_method, N){</pre>
  # get correlation matrix
  cor.cluster <- cor.matrices[[cor_method]]</pre>
  # perform hierarchical clustering (complete linkage & Euclidion distance measure)
  hc <- hclust(dist(cor.cluster, method="euclidian"), method="complete")</pre>
  # cut into the clusters
  groups <- cutree(hc, k=Ngroups)</pre>
  groups.hc.order <- groups[hc$order]</pre>
  # store results
 return(list(hc=hc,
              groups=groups,
              groups.hc.order=groups.hc.order))
}
Ngroups <- 6 # number of clusters
for (method in correlation_methods){
 hclust.res[[method]] <- calculate_clusters(cor_method=method, N=Ngroups)
}
# save clustering
save(hclust.res, file=file.path(resultsPath, "data", "hclust.res.Rdata"))
rm(method)
```

In the next step the correlation matrices are plotted in combination with the clustering results

```
# Create correlation heatmap with hierachical clustering results
correlation_heatmap_cluster <- function(method){</pre>
  # matrix and cluster results
  cor.cluster <- cor.matrices[[method]]</pre>
  hc.res <- hclust.res[[method]]</pre>
  hc <- hc.res$hc
  groups <- hc.res$groups</pre>
  # colors
  hmap_colors <- HeatmapColors()</pre>
  colorset <- brewer.pal(Ngroups, "Set1")</pre>
  color.map <- function(cluster_id) {return(colorset[cluster_id])}</pre>
  clusterColors <- unlist(lapply(groups, color.map))</pre>
  heatmap.2(cor.cluster, col=hmap_colors(10), scale="none",
          key=TRUE, symkey=FALSE, trace="none", cexRow=0.8, cexCol=0.8,
          main=method,
          density.info="none", dendrogram="column", Rowv=as.dendrogram(hc), Colv=as.dendrogram(hc), key
          key.xlab=method,
```

```
ColSideColors=clusterColors, revC=TRUE,
          sepwidth=c(0.01,0.01),
          sepcolor="black",
          colsep=1:ncol(cor.cluster),
          rowsep=1:nrow(cor.cluster),
          margins=c(12,8))
  legend("left", legend=paste("c", 1:6, sep=""), col= unlist(lapply(1:6, color.map)), pch=15, bty="n")
}
# plot to files
for(method in correlation_methods){
  pdf(file.path(resultsPath, "correlation", sprintf("cor.%s.hclust.pdf", method)),
      width=10, height=10, pointsize=12)
  correlation_heatmap_cluster(method=method)
  invisible(dev.off())
}
rm(method)
```

Display the clusters with the heatmap

```
# plot ys3 to report
correlation_heatmap_cluster(method="ys3")
```

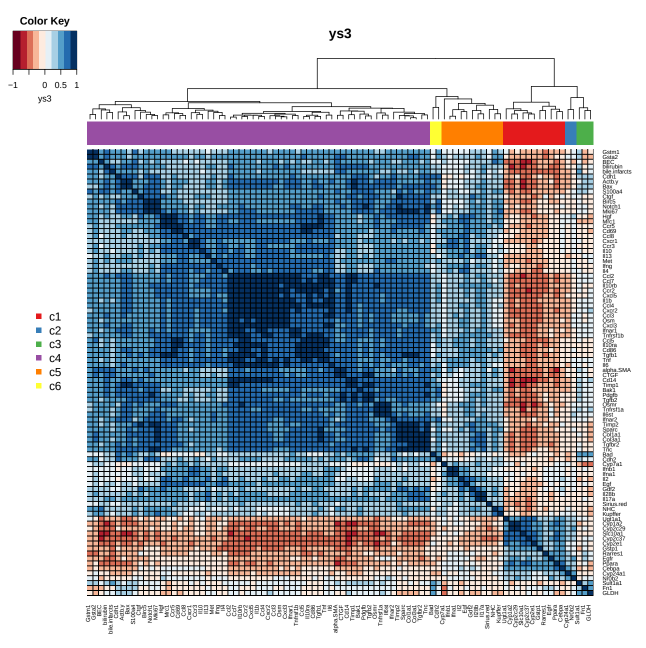


Figure Heatmap Cluster: Correlation matrix based on YS3 correlation with hierarchical clustering infromation. The following factors are in the ys3 time course clusters

```
# print(paste(names(g), sep=", ", collapse =", "))
}
list_cluster_members(method="ys3")
```

```
## Correlation method: *** ys3 ***
## Cluster 1 (N=11)
                                         "Cebpa"
                                                   "Slc10a1" "Ugt1a1"
    [1] "Ppara"
                   "Cyp1a2"
                              "Gstp1"
                                                                         "Cyp2c37"
##
    [8] "Cyp2c29" "Egfr"
                              "Rarres1" "Cyp2e1"
##
##
## Cluster 2 (N=2)
## [1] "Cyp24a1" "Nr0b2"
##
## Cluster 3 (N=3)
## [1] "Sult1a1" "Fn1"
                             "GLDH"
## Cluster 4 (N=61)
                          "Gsta2"
                                                            "Cd86"
##
   [1] "Gstm1"
                                           "Cc12"
    [5] "Met"
                          "Tnfrsf1a"
                                           "Il1b"
                                                            "Cxcl5"
##
##
    [9] "Cxcr1"
                          "Ifnar1"
                                           "Osmr"
                                                            "Tgfb1"
                          "Cc14"
##
  [13] "Ifng"
                                           "Cc15"
                                                            "Ccr3"
  [17] "Il13"
                          "Tnfrsf1b"
                                           "Cc17"
                                                            "Osm"
   [21] "Cd69"
                          "Il10rb"
                                           "Cd14"
                                                            "Cxcr2"
##
        "Tnf"
                          "Il10ra"
                                                            "Ccr2"
##
   [25]
                                           "Il10"
## [29] "Hgf"
                          "Ifnar2"
                                           "Mrc1"
                                                            "Ccr5"
## [33] "Cc13"
                          "I16"
                                                            "I14"
                                           "Actb.y"
## [37] "Cc18"
                          "Cxcl3"
                                           "I16st"
                                                            "Birc5"
## [41]
        "Sparc"
                          "Col1a1"
                                           "Tgfbr2"
                                                            "Timp1"
  [45]
                          "Cdh1"
                                           "Timp2"
        "Pdgfb"
                                                            "Bak1"
  [49] "Ctgf"
                          "Notch1"
                                           "Tnc"
                                                            "Bax"
##
                          "Tgfb2"
##
   [53] "Mki67"
                                           "Col3a1"
                                                            "bilirubin"
##
   [57] "BEC"
                          "bile.infarcts" "alpha.SMA"
                                                            "CTGF"
   [61] "S100a4"
##
##
## Cluster 5 (N=11)
    [1] "Cyp7a1"
                                    "I128b"
                                                  "Il17a"
                                                                "I12"
##
                      "Ifnb1"
    [6] "Egf"
                      "Ifna1"
                                    "Gdf2"
                                                  "NHC"
                                                                 "Kupffer"
  [11] "Sirius.red"
##
## Cluster 6 (N=2)
## [1] "Bad"
               "Cdh2"
```

Correlations for non-RNA factors

Plot correlation matrix for histological, biochemical and antibody markers. Calculation of the largest correlation between the non-transcript factors and other factors. The columns in which no correlation coefficient has abs(value) >= cor.cutoff are filtered out, i.e. only factors are retained with a absolute correlation coefficient above this threshold.

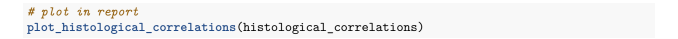
```
get_histological_correlations <- function(method="ys3", cor.cutoff=0.6){</pre>
  # correlation matrix
  cor_mat <- cor.matrices[[method]]</pre>
  # non RNA factors which were accepted by ANOVA
  hist facs <- BDLfactors$id[anova.accept &
                                (BDLfactors$ftype %in% c("Antibodies", "Biochemistry", "Histology"))]
  # print(hist facs)
  # get the indices of these factors in the correlation matrix
  hist_idx <- rep(NA, length(hist_facs))</pre>
  for (k in 1:length(hist_facs)){
    hist_idx[k] <- which(colnames(cor_mat) == hist_facs[k])
  # subset of correlation matrix for histological markers (corresponding rows)
  hist_data <- cor_mat[hist_idx, ]</pre>
  # filter columns by correlation cutoff
  col.accept <- rep(NA, ncol(hist_data))</pre>
  for (k in 1:ncol(hist_data)){
    # if there is any correlation >= cutoff in the column, keep it
    col.accept[k] <- any( abs(hist_data[,k])>=cor.cutoff )
  # print(table(col.accept)) # filtered factors based on correlation threshold
  hist_accept <- hist_data[, col.accept]</pre>
  # sort by the hierarchical cluster ordering
  hist_gene_names <- colnames(hist_accept)[1:(ncol(hist_accept)-nrow(hist_accept))]
  # correlation based cluster for ordering
  hc.res <- hclust.res[[method]]</pre>
  # create sort index for gene names based on clustering
  sort_idx <- rep(NA, length(hist_gene_names))</pre>
  for (k in 1:length(hist_gene_names)){
    sort_idx[k] <- which(names(hc.res$groups.hc.order) == hist_gene_names[k])</pre>
  # first sorted genes than the histological factors
  hist_sorted <- hist_accept[, c(hist_gene_names[order(sort_idx)], rownames(hist_data))]</pre>
  return(hist sorted)
histological_correlations <- get_histological_correlations(method="ys3")
# plot the subset of the correlation matrix to file
plot_histological_correlations <- function(hist_data){</pre>
  hmap_colors <- HeatmapColors()</pre>
  corrplot(hist_data, method="circle", type="full",
           tl.cex=0.7, tl.col="black", col=hmap_colors(10))
}
# plot to file
pdf(file.path(resultsPath, "correlation", "histological_correlations.pdf"),
```

```
width=10, height=4, pointsize=12)
plot_histological_correlations(histological_correlations)
invisible(dev.off())

# print correlation values
print(round(t(histological_correlations), digits=2))
```

##		GLDH	bilirubin	NHC	Kupffer	BEC	Sirius.red	bile.infarcts
##	Gstm1	-0.02	0.61	0.42	0.40	0.43	0.10	0.59
	Gsta2	-0.21	0.61	0.51	0.41	0.46	0.19	0.59
##	Cdh1	0.24	0.46	0.25	0.34	0.35	0.18	0.42
##	Actb.y	0.22	0.61	0.29	0.33	0.47	0.29	0.50
	Bax	0.23	0.61	0.31	0.35	0.42	0.38	0.44
##	Birc5	-0.04	0.57	0.53	0.50	0.42	0.36	0.41
##	Notch1	-0.04	0.52	0.43	0.50	0.31	0.26	0.39
##	Mki67	-0.15	0.50	0.59	0.61	0.47	0.35	0.39
##	Cc12	0.17	0.66	0.25	0.25	0.52	0.36	0.54
##	Cc17	0.15	0.60	0.18	0.22	0.49	0.32	0.48
##	Il10rb	0.00	0.57	0.31	0.33	0.50	0.40	0.46
	Ccr2	0.03	0.56	0.29	0.29	0.53	0.33	0.49
##	Cxc15	0.00	0.54	0.26	0.23	0.49	0.34	0.43
##	Il1b	0.07	0.54	0.18	0.17	0.38	0.37	0.37
##	Ccl4	0.13	0.55	0.16	0.22	0.45	0.33	0.42
##	Cxcr2	0.22	0.55	0.15	0.15	0.44	0.33	0.42
##	Cc13	0.18	0.62	0.16	0.17	0.45	0.23	0.53
##	Osm	0.19	0.59	0.15	0.14	0.38	0.37	0.45
##	Cxcl3	0.18	0.61	0.12	0.18	0.41	0.36	0.44
##	Ifnar1	0.11	0.56	0.21	0.23	0.36	0.30	0.44
##	Il10ra	-0.15	0.46	0.31	0.30	0.39	0.33	0.38
##	Cd86	-0.07	0.45	0.39	0.41	0.46	0.40	0.32
##	Tgfb1	-0.14	0.48	0.41	0.40	0.46	0.38	0.39
##	Tnf	0.00	0.49	0.25	0.25	0.34	0.31	0.40
##	I16	0.03	0.51	0.13	0.20	0.40	0.25	0.37
##	Cd14	0.21	0.67	0.29	0.21	0.50	0.37	0.47
##	Timp1	0.12	0.67	0.32	0.34	0.50	0.27	0.57
##	Bak1	0.08	0.60	0.43	0.40	0.40	0.35	0.41
##	Pdgfb	0.08	0.62	0.33	0.37	0.41	0.30	0.48
##	Tgfb2	0.11	0.58	0.30	0.33	0.38	0.28	0.51
##	Timp2	-0.24	0.51	0.47	0.39	0.37	0.42	0.38
##	Col1a1	-0.22	0.55	0.53	0.38	0.48	0.38	0.42
##	Tgfbr2	-0.10	0.59	0.47	0.31	0.37	0.31	0.46
##	Tnc	-0.10	0.54	0.43	0.37	0.40	0.33	0.42
##	Cyp1a2	-0.15		-0.11		-0.44	-0.20	-0.56
	Slc10a1	-0.17		-0.38		-0.66	-0.37	-0.50
	Cyp2c37	0.03		-0.41		-0.66	-0.41	-0.49
	Ppara	-0.22		-0.18		-0.49	-0.05	-0.41
	GLDH	1.00		-0.30		-0.02	-0.10	0.14
	bilirubin	0.18		0.39	0.26		0.35	0.71
	NHC	-0.30	0.39	1.00	0.75		0.32	0.25
	Kupffer	-0.20	0.26		1.00	0.57	0.30	0.10
	BEC	-0.02	0.49		0.57		0.36	0.39
	Sirius.red	-0.10	0.35	0.32	0.30	0.36	1.00	0.06
##	bile.infarcts	0.14	0.71	0.25	0.10	0.39	0.06	1.00

##	alpha.SMA	-0.04	0.65	0.41	0.27	0.60	0.20	0.61
##	CTGF	-0.06	0.69	0.46	0.25	0.56	0.21	0.63
##	S100a4	0.08	0.62	0.54	0.47	0.62	0.38	0.44
##		${\tt alpha.SMA}$	CTGF	S100a4				
##	Gstm1	0.55	0.57	0.70				
##	Gsta2	0.64	0.62	0.59				
##	Cdh1	0.54	0.54	0.61				
##	Actb.y	0.46	0.60	0.52				
##	Bax	0.48	0.55	0.59				
##	Birc5	0.61	0.59	0.72				
	Notch1	0.53	0.61	0.71				
	Mki67	0.55	0.57	0.68				
	Cc12	0.61	0.72	0.46				
	Cc17	0.62	0.67	0.48				
	Il10rb	0.64	0.70	0.50				
	Ccr2	0.59	0.74	0.46				
	Cxcl5	0.73	0.75	0.53				
	Il1b	0.63	0.71	0.48				
	Cc14	0.61	0.68	0.46				
	Cxcr2	0.61	0.71	0.44				
	Cc13	0.62	0.69	0.50				
	Osm	0.59	0.63	0.51				
	Cxcl3	0.53	0.57	0.47				
	Ifnar1	0.56	0.67	0.49				
	Il10ra Cd86	0.62	0.68 0.72	0.44				
	Tgfb1	0.62 0.60	0.72	0.44 0.48				
	Tnf	0.60	0.69	0.48				
	III	0.65	0.61	0.49				
	Cd14	0.67	0.77	0.50				
	Timp1	0.73	0.73	0.64				
	Bak1	0.56	0.67	0.60				
	Pdgfb	0.69	0.80	0.72				
	Tgfb2	0.73	0.82	0.72				
	Timp2	0.63	0.62	0.68				
	Col1a1	0.59	0.61	0.66				
##	Tgfbr2	0.69	0.78	0.59				
	Tnc	0.70	0.73	0.67				
##	Cyp1a2	-0.70	-0.60	-0.45				
##	Slc10a1	-0.54	-0.55	-0.44				
##	Cyp2c37	-0.59	-0.55	-0.44				
##	Ppara	-0.61	-0.59	-0.37				
##	GLDH	-0.04	-0.06	0.08				
##	bilirubin	0.65	0.69	0.62				
##	NHC	0.41	0.46	0.54				
##	Kupffer	0.27	0.25	0.47				
##	BEC	0.60	0.56	0.62				
	Sirius.red	0.20	0.21	0.38				
##	bile.infarcts	0.61	0.63	0.44				
	alpha.SMA	1.00	0.84	0.69				
	CTGF	0.84	1.00	0.60				
##	S100a4	0.69	0.60	1.00				



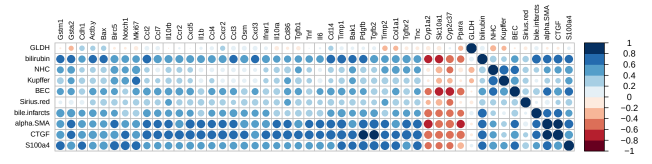


Figure Histological Correlations: Correlations between non-RNA factors and all other factors.

Top correlations for non-RNA factors

List the top correlations of every non-RNA factor, i.e. histological (H), biochemical (B) and immunohistochemical (A). The top correlations are sorted by absolute correlation values.

```
plot_hist_topcors <- function(method="ys3", labels=TRUE, mfrow=c(10,1),</pre>
                                only_RNA=FALSE, Ntop=10, printResults=FALSE){
  hist_facs <- rownames(histological_correlations)</pre>
  cor.mat <- cor.matrices[[method]]</pre>
  par(mfrow=mfrow)
  hmap colors <- HeatmapColors()</pre>
  for (name in hist_facs){
    if (only RNA==TRUE){
      # get the elements which are not histological factors
      v <- cor.mat[!(colnames(cor.mat) %in% hist_facs), name]</pre>
    } else {
      # get all elements (including other non-RNA factors)
      v <- cor.mat[, name]</pre>
    # sort by absolute correlation
    v.sorted <- rev(v[order(abs(v))])</pre>
    # and get the Ntop values without the self-correlation (idx=1)
    mv <- t(as.matrix(v.sorted[2:(Ntop+1)]))</pre>
    rownames(mv) <- c(name)</pre>
    # overview
    if (printResults==TRUE){
      print(round(mv, digits=2))
    # plot without labels to have identical size of figure
    if (labels==FALSE){
      rownames(mv) <- NULL
      colnames(mv) <- NULL</pre>
```

```
corrplot(mv, method="pie", type="full",
            tl.cex=1.0, tl.col="black", col=hmap_colors(100), insig="p-value", sig.level=-1,
            p.mat=mv, cl.pos="n")
 }
 par(mfrow=c(1,1))
# plot to file
# necessary to plot once with and without labels so that all the plots are scaled equally
pdf(file.path(resultsPath, "correlation", "histological_topcors_1.pdf"),
   width=10, height=10, pointsize=12)
plot hist topcors(method="ys3", labels=TRUE)
invisible(dev.off())
pdf(file.path(resultsPath, "correlation", "histological_topcors_2.pdf"),
   width=10, height=10, pointsize=12)
plot_hist_topcors(method="ys3", labels=FALSE)
invisible(dev.off())
# plot in report
plot_hist_topcors(method="ys3", labels=TRUE, mfrow=c(5,2), printResults=TRUE)
       Sult1a1 Bad Fn1 Cyp7a1 Mrc1 Il28b Gdf2 Il2 NHC Cebpa
          0.55 0.54 0.51 -0.47 -0.37 -0.35 -0.32 -0.31 -0.3 -0.3
## GLDH
            bile.infarcts CTGF Timp1 Cd14 Cc12 alpha.SMA Cc13 Slc10a1 S100a4
                     0.71 0.69 0.67 0.67 0.66 0.65 0.62 -0.62 0.62
## bilirubin
            Pdgfb
## bilirubin 0.62
##
      Kupffer BEC Mki67 S100a4 Birc5 Col1a1 Sparc Gsta2 Mrc1 Col3a1
## NHC
         0.75 0.62 0.59 0.54 0.53 0.53 0.53 0.51 0.5
           NHC Mki67 BEC Birc5 Notch1 S100a4 Ccr5 Mrc1 Sparc Hgf
## Kupffer 0.75 0.61 0.57 0.5
                                  0.5 0.47 0.45 0.44 0.43 0.43
      Cyp2c37 Slc10a1 NHC S100a4 alpha.SMA Kupffer CTGF Cyp2e1 Ccr2 Cyp2c29
## BEC -0.66 -0.66 0.62
                            0.62
                                      0.6
                                              0.57 0.56 -0.56 0.53 -0.52
             Timp2 Cyp2c29 Cyp2c37 Tnfrsf1b Il10rb Ccr5 Cd86 Ifng Sparc
## Sirius.red 0.42
                   -0.41 -0.41
                                     0.41
                                            0.4 0.4 0.4 0.39 0.38
             Col3a1
## Sirius.red 0.38
                bilirubin CTGF alpha.SMA Gsta2 Gstm1 Timp1 Cyp1a2 Ccl2
                     0.71 0.63
                                  0.61 0.59 0.59 0.57 -0.56 0.54
## bile.infarcts
                Cyp2e1 Ccl3
## bile.infarcts -0.53 0.53
            CTGF Tgfb2 Cxcl5 Timp1 Tnc Cyp1a2 S100a4 Pdgfb Tgfbr2 Cd14
## alpha.SMA 0.84 0.73 0.73 0.73 0.7 -0.7 0.69 0.69 0.69 0.67
```

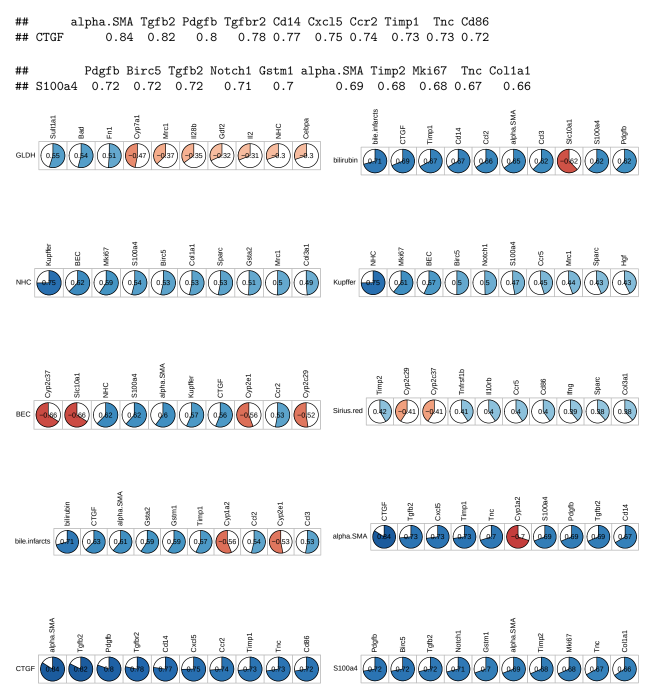


Figure Histological Top Correlations: Top correlations between non-RNA factors and all other factors.

Mean cluster time course

In the next step, we were interested in the time courses of the time course clusters: What are the typical profiles found in the different clusters and which factors are in these clusters. For the comparison of individual factors against each other and against the mean time course of the cluster the factors were normalized. The normalization was hereby performed for every factor separately based on

$$f_k^{norm}(t_{i,r}) = \frac{f_k(t_{i,r}) - \langle f_k \rangle}{max(f_x) - min(f_x)}$$

```
suppressPackageStartupMessages(library('matrixStats'))
dir.create(file.path(resultsPath, 'cluster'), showWarnings=FALSE)
# Normalize the individual factors
normalize_factor <- function(a, min.a, max.a, mean.a){</pre>
  res <- (a - mean.a)/(max.a - min.a)
# Calculate min, max and mean for all single factors (normalization constants)
factor.norm <- data.frame(min=apply(BDLdata, 2, min, na.rm=TRUE),</pre>
                             max=apply(BDLdata, 2, max, na.rm=TRUE),
                             mean=apply(BDLdata, 2, mean, na.rm=TRUE))
# Function for normalizing subset of BDL data with factor normalization constants.
normalize_BDLdata <- function(data, factor.norm){</pre>
  dnorm <- data
  for (name in colnames(data)){
    dnorm[, name] <- normalize_factor(a=dnorm[, name],</pre>
                                       min.a=factor.norm[name, "min"],
                                       max.a=factor.norm[name, "max"],
                                       mean.a=factor.norm[name, "mean"])
 }
 return(dnorm)
# Normalize the full data set
BDLdata.norm <- normalize BDLdata(data=BDLdata, factor.norm=factor.norm)
# Calculate the mean of the normalized data set
BDLmean.norm <- bdl_mean_data(BDLdata.norm, BDLsamples)</pre>
```

Based on the normalized factor data the mean time course of the clusters were calculated, i.e. the time course due to averaging over all factors in the individual clusters. In addition the mean time course averaged over the repeats for all factors are plotted.

```
# Plot mean cluster with SD range and the individual representatives in the cluster.
plot_clusters_mean <- function(method){</pre>
  # clusters for correlation method
  hc.res <- hclust.res[[method]]</pre>
  groups.hc.order <- hc.res$groups.hc.order</pre>
  # par(mfrow=c(ceiling(sqrt(Ngroups))), ceiling(sqrt(Ngroups))))
  par(mfrow=c(2,3))
  steps <- 1:Nt # time points
  for (k in 1:Ngroups){
    # get representatives of cluster
    g <- groups.hc.order[groups.hc.order==k]</pre>
    dgroup <- BDLmean.norm[names(g)] # normalized mean data for group (normalized within factor)
    # mean and sd for timepoints (i.e. over all factors in the cluster)
    g.mean <- rowMeans(as.matrix(dgroup), na.rm=TRUE)</pre>
    g.sd <- rowSds(as.matrix(dgroup), na.rm=TRUE)</pre>
    # plot sd range
    plot(factor(levels(BDLsamples$time_fac), levels=levels(BDLsamples$time_fac)), rep(-2, 8), type="n",
         xlim=c(1, Nt), ylim=1.1*c( min(min(dgroup, na.rm=TRUE), na.rm=TRUE),
```

```
max(max(dgroup, na.rm=TRUE), na.rm=TRUE) ),
         main=sprintf("%s : Cluster %s (N=%s)", method, k, ncol(dgroup)))
    polygon(c(steps, rev(steps)), c(g.mean+g.sd, rev(g.mean-g.sd)),
            col = rgb(0,0,1,0.2), border = NA)
    # individual data
    for (name in names(g)){
      points(steps, dgroup[, name], pch=16, col="black")
      lines(steps, dgroup[, name], col=rgb(0.5,0.5,0.5,1.0), lwd=1)
    # mean over factors in cluster
    lines(steps, g.mean, col="blue", lwd=2)
  }
 par(mfrow=c(1,1))
# plot mean clusters to file
for (method in correlation_methods){
  pdf(file.path(resultsPath, 'cluster', sprintf("%s_cluster_mean.pdf", method)),
        width=10, height=7.5, pointsize=12)
  plot_clusters_mean(method=method)
  invisible(dev.off())
```

```
# plot ys3 clusters to report
plot_clusters_mean(method="ys3")
```

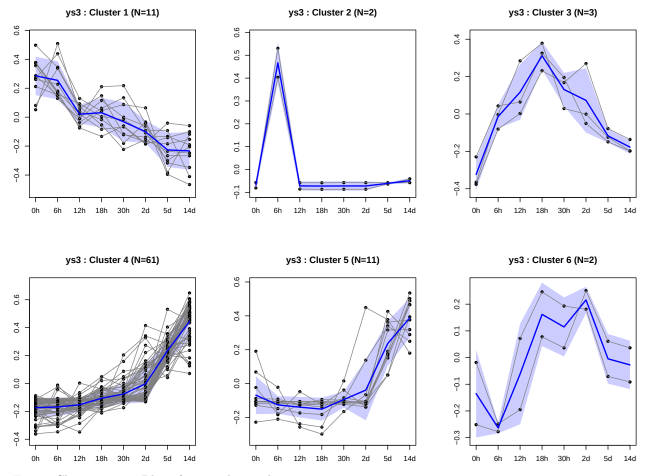


Figure Cluster mean: Plot of mean cluster data.

In addition the plots of the individual factors in the clusters are generated. Despite being clustered together still quit large variance exists between the different factors in the clusters.

```
# Plot individual time courses in cluster
plot_clusters_items <- function(method, toFile=FALSE){</pre>
  # get the cluster assignment for the given method
 hc.res <- hclust.res[[method]]</pre>
  groups.hc.order <- hc.res$groups.hc.order</pre>
 for (k in 1:Ngroups){
    if (toFile==TRUE){
      pdf(file.path(resultsPath, "cluster", sprintf("%s_cluster_%s.pdf", method, k)),
          width=10, height=10, pointsize=8)
    }
    g <- groups.hc.order[groups.hc.order==k]</pre>
    N <- ceiling(sqrt(length(g)))</pre>
    par(mfrow=c(N,N))
    for (name in names(g)){
      plot_single(name_A=name)
    par(mfrow=c(1,1))
    if (toFile==TRUE){
```

```
invisible(dev.off())
}

}

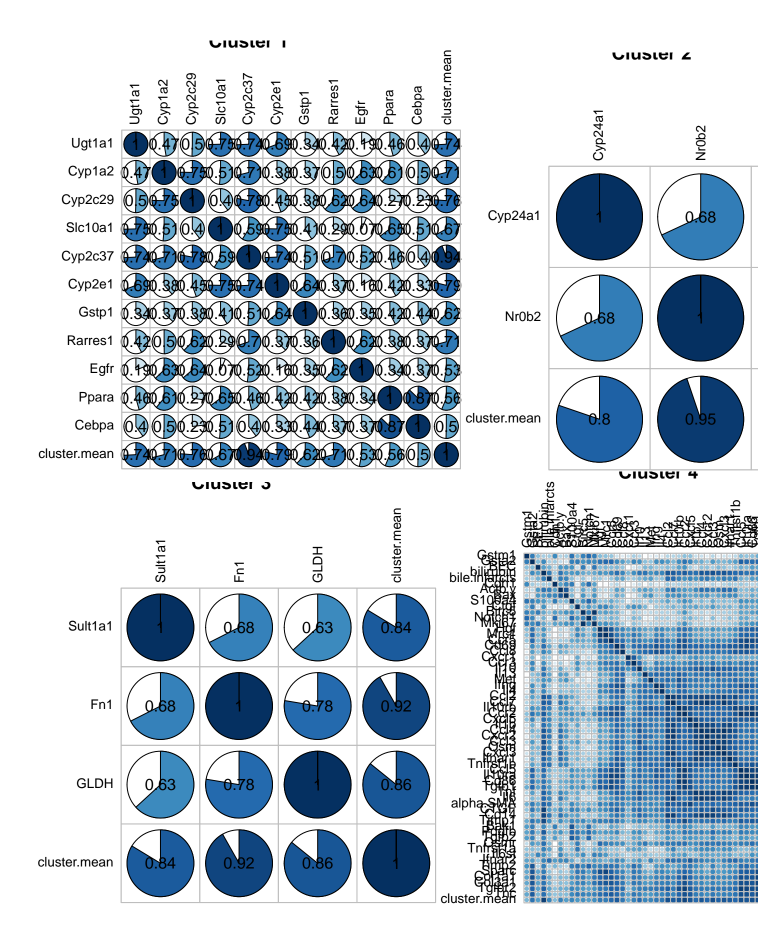
# plot individual factors in clusters on disk
for (method in correlation_methods){
   plot_clusters_items(method=method, toFile=TRUE)
}
rm(method)
```

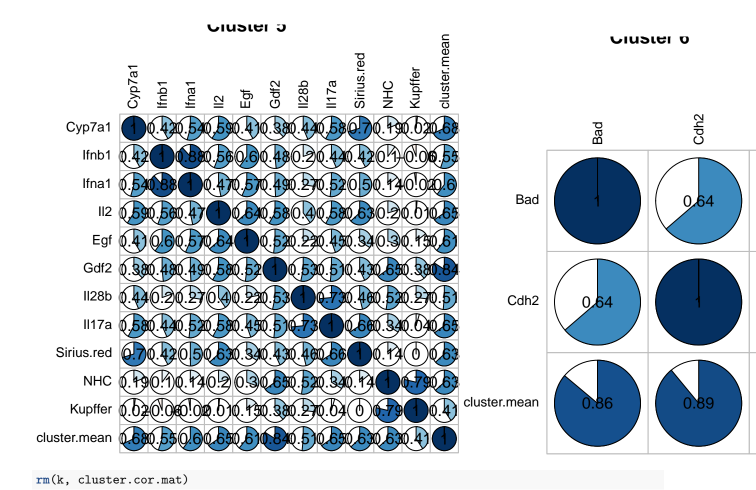
Top cluster representatives

In this step, the top cluster representatives are calculated for all clusters. The higher the correlation between the factor and the cluster mean, the higher the respective factor is scored. The used correlation measure is YS3.

```
# Calculate and plot the top cluster representatives.
# Currently only ys3 based calculation is supported. This function should be refactored
# to support all correlation measures.
hmap_colors <- HeatmapColors()</pre>
# calculates the correlation between the cluster mean and all other members in the cluster
calculate_cluster_correlations <- function(method="ys3"){</pre>
  if (!identical(method, "ys3")){
    stop("Only ys3 top correlation is currently supported")
  }
  # get the clusters
 hc.res <- hclust.res[[method]]</pre>
  groups.hc.order <- hc.res$groups.hc.order</pre>
  # mean cluster time course as additional factor
  name_cluster <- sprintf("cluster.mean")</pre>
  # calculate and plot the top correlations for every cluster
  cluster.cor <- vector("list", length=Ngroups)</pre>
  for (k in 1:Ngroups){
      # get members of cluster
      g <- groups.hc.order[groups.hc.order==k]</pre>
      # get normalized data for members of cluster
      dgroup <- BDLdata.norm[names(g)]</pre>
      # calculate cluster mean and add as factor (mean of all factors in cluster for given time point a
      # (for S*)
      g.mean <- rowMeans(as.matrix(dgroup), na.rm=TRUE)</pre>
      dgroup[[name_cluster]] <- g.mean</pre>
      # calculate mean averaged over repeats (for A** and M*)
      dgroup.mean <- bdl_mean_data(dgroup, BDLsamples)</pre>
      # now calculate the ys3 correlation with the created data frame containing the mean cluster facto
      ysr.res <- ysr.matrices(dgroup.mean, BDLmean.time, use="pairwise.complete.obs")
```

```
# Spearman correlation on individual data points
      cor.S_star <- ( cor(dgroup, method="spearman", use="pairwise.complete.obs") + 1 )/2</pre>
      # YS3 in [-1,1]
      ys3 <- 2*( (w$w1*cor.S_star + w$w2*ysr.res$A_star2 + w$w3*ysr.res$M_star) - 0.5)
      # store the matrix
      cluster.cor[[k]] <- ys3</pre>
  }
 return(cluster.cor)
cluster.cor <- calculate_cluster_correlations(method="ys3")</pre>
# plot correlations for individual clusters
for (k in 1:Ngroups){
  cluster.cor.mat <- cluster.cor[[k]]</pre>
  if (ncol(cluster.cor.mat)<20){
    corrplot(cluster.cor.mat, method="pie", type="full",
           main=sprintf("Cluster %s", k),
             tl.cex=0.8, tl.col="black", col=hmap_colors(100),
             insig="p-value", sig.level=-1, p.mat=cluster.cor.mat,
             cl.pos="n")
  } else {
    corrplot(cluster.cor.mat, method="circle", type="full",
           main=sprintf("Cluster %s", k),
             t1.cex=0.8, t1.col="black", col=hmap_colors(100),
             cl.pos="n")
 }
```





Now the top correlations between the cluster mean and the members of the respective cluster are plotted.

```
plot_top_cluster_representatives <- function(method="ys3", Ntop=11, labels=TRUE){</pre>
  par(mfrow=c(Ngroups,1))
  name_cluster <- sprintf("cluster.mean")</pre>
  for (k in 1:Ngroups){
      # YS3 in [-1,1]
      ys3 <- cluster.cor[[k]]
      # correlation for the cluster mean
      v <- ys3[, which(colnames(ys3)==name_cluster)]</pre>
      # sort by absolute correlation
      v.sorted <- rev(v[order(abs(v))])</pre>
      # reduce to Ntop candidates and fill short clusters with zeros tto Ntop
      if (length(v.sorted)>=(Ntop+1)){
        v.sorted <- v.sorted[2:(Ntop+1)]</pre>
        v.sorted <- c(v.sorted[2:length(v.sorted)], rep(0,Ntop-length(v.sorted)+1))</pre>
      # prepare data for corrplot
      mv <- t(as.matrix(v.sorted))</pre>
      rownames(mv) <- c(name_cluster)</pre>
      if (labels == FALSE){
```

```
colnames(mv) <- NULL</pre>
      }
      corrplot(mv, method="pie", type="full",
             tl.cex=1.0, tl.col="black", col=hmap_colors(100),
             insig="p-value", sig.level=-1, p.mat=mv,
             cl.pos="n")
  }
  par(mfrow=c(1,1))
}
# plot to file (with and without names)
pdf(file.path(resultsPath, "cluster", "cluster_top_representatives_01.pdf"),
    width=5, height=5, pointsize=12)
plot_top_cluster_representatives(labels=FALSE)
invisible(dev.off())
pdf(file.path(resultsPath, "cluster", "cluster_top_representatives_02.pdf"),
    width=10, height=10, pointsize=12)
plot_top_cluster_representatives(labels=TRUE)
invisible(dev.off())
# plot in report
plot_top_cluster_representatives(labels=TRUE)
```

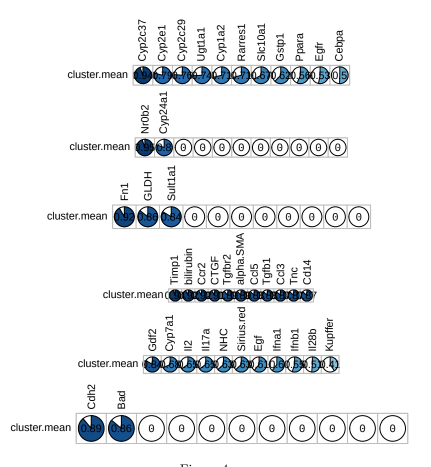


Figure 4:

Overview YS3 clusters

Full overview over the members of the {r} Ngroups clusters with respective YS3 correlation to the cluster mean and ANOVA result.

```
# print representatives of cluster
list_cluster_information <- function(method="ys3"){</pre>
 if(!identical(method, "ys3")){
   stop("Cluster correlation only calculated for ys3")
 }
 groups <- hclust.res[[method]]$groups</pre>
 cat(sprintf("-----\n"))
 cat(sprintf("Correlation method: *** %s ***\n", method))
 cat(sprintf("-----\n"))
 for (k in 1:Ngroups){
   cluster.cor.mat <- cluster.cor[[k]]</pre>
   # create data.frame with ANOVA & cluster correlation
   g <- groups[groups==k]</pre>
   cat(sprintf("Cluster %s (N=%s)\n", k, length(g)))
   Ng <- length(names(g))
   rows <- vector("list", length=Ng)</pre>
   for (kn in 1:Ng){
     name <- names(g)[kn]
     row <- df.anova[which(df.anova$factor==name),]</pre>
     row$cluster.cor <- cluster.cor.mat[name, "cluster.mean"]</pre>
     rows[[kn]] <- row
   }
   df <- do.call("rbind", rows)</pre>
   df <- df[order(-df$cluster.cor), ]</pre>
   # cleanup the data.frame
   rownames(df) <- df$factors</pre>
   df <- df[, c("p.holm", "sig.holm", "ftype", "fshort", "cluster.cor")]</pre>
   df$cluster.cor <- round(df$cluster.cor, digits=2)</pre>
   df$p.holm <- sprintf("%1.2E", df$p.holm)</pre>
   print(df)
   cat("\n")
   # create list for figure legend
    # print(paste(sprintf("%s (%s)", rownames(df), df$cluster.cor),
                sep=", ", collapse =", "))
 }
}
# List information in cluster
list_cluster_information(method="ys3")
## Correlation method: *** ys3 ***
## -----
```

```
## Correlation method: *** ys3 ***

## ------
## Cluster 1 (N=11)

## p.holm sig.holm ftype fshort cluster.cor

## Cyp2c37 1.09E-05 *** GE_ADME 0.94

## Cyp2e1 1.50E-06 *** GE_Fibrosis 0.79

## Cyp2c29 2.93E-06 *** GE_ADME 0.76
```

```
## Ugt1a1 2.75E-03
                                   GE ADME
                                                          0.74
                           **
                                   GE_ADME
## Cyp1a2 2.93E-14
                          ***
                                                          0.71
## Rarres1 3.57E-02
                               GE Fibrosis
                                                          0.71
## Slc10a1 5.26E-05
                                   GE_ADME
                          ***
                                                          0.67
## Gstp1
           1.38E-04
                          ***
                                   GE ADME
                                                          0.62
                                   GE ADME
## Ppara
           1.63E-04
                          ***
                                                          0.56
## Egfr
           1.59E-03
                           ** GE Cytokines
                                                          0.53
## Cebpa
           4.93E-04
                                   GE_ADME
                                                          0.50
##
## Cluster 2 (N=2)
             p.holm sig.holm
                                ftype fshort cluster.cor
## NrOb2
           1.30E-07
                          *** GE_ADME
                                                     0.95
                                                     0.80
## Cyp24a1 9.88E-03
                           ** GE_ADME
##
## Cluster 3 (N=3)
##
             p.holm sig.holm
                                     ftype fshort cluster.cor
## Fn1
           1.21E-03
                           ** GE_Fibrosis
                                                          0.92
## GLDH
           4.28E-03
                           ** Biochemistry
                                                 В
                                                          0.86
## Sult1a1 2.98E-03
                                   GE_ADME
                                                          0.84
                           **
## Cluster 4 (N=61)
                   p.holm sig.holm
                                           ftype fshort cluster.cor
                                *** GE_Fibrosis
## Timp1
                 4.30E-04
                                                                 0.94
## bilirubin
                 4.98E-12
                                                                 0.92
                                *** Biochemistry
## Ccr2
                 6.94E-11
                                *** GE_Cytokines
                                                                 0.92
## CTGF
                 7.82E-10
                                ***
                                      Antibodies
                                                                 0.91
## Tgfbr2
                 1.71E-07
                                ***
                                     GE_Fibrosis
                                                                 0.89
## alpha.SMA
                 5.02E-04
                                ***
                                      Antibodies
                                                                 0.89
## Cc15
                 2.78E-07
                                *** GE_Cytokines
                                                                 0.88
## Tgfb1
                 3.30E-11
                                *** GE_Cytokines
                                                                 0.88
## Cc13
                 8.53E-06
                                *** GE_Cytokines
                                                                 0.87
## Tnc
                 2.04E-03
                                 ** GE_Fibrosis
                                                                 0.87
## Cd14
                 1.15E-05
                                *** GE_Cytokines
                                                                 0.87
## Cc12
                 3.46E-11
                                *** GE_Cytokines
                                                                0.86
## Cd86
                 6.56E-11
                                *** GE_Cytokines
                                                                 0.86
## Pdgfb
                 7.68E-04
                                *** GE_Fibrosis
                                                                0.86
## Col1a1
                 4.40E-07
                                *** GE Fibrosis
                                                                0.86
## Cxcl3
                 6.82E-05
                                *** GE_Cytokines
                                                                0.86
## Cc14
                 1.04E-07
                                *** GE_Cytokines
                                                                 0.85
## Cxc15
                                *** GE_Cytokines
                 6.26E-10
                                                                0.85
## Il10ra
                 1.64E-09
                                *** GE_Cytokines
                                                                0.85
## Col3a1
                 2.02E-05
                                *** GE_Fibrosis
                                                                 0.85
## Il10rb
                 1.15E-11
                                *** GE Cytokines
                                                                 0.84
## Cc17
                                *** GE_Cytokines
                                                                 0.82
                 3.40E-08
## Cd69
                 2.73E-06
                                *** GE_Cytokines
                                                                 0.82
## Ifnar1
                 7.71E-07
                                *** GE_Cytokines
                                                                 0.82
## Tnf
                 4.36E-06
                                *** GE_Cytokines
                                                                0.82
## Osm
                 7.51E-06
                                *** GE_Cytokines
                                                                0.81
## Sparc
                 1.07E-06
                                *** GE_Fibrosis
                                                                 0.80
## I16
                 2.23E-04
                                *** GE_Cytokines
                                                                 0.80
## Tnfrsf1b
                 7.89E-11
                                *** GE_Cytokines
                                                                 0.80
## Cxcr2
                 1.75E-06
                                *** GE_Cytokines
                                                                0.78
## Il1b
                 7.29E-06
                                *** GE_Cytokines
                                                                0.78
## Timp2
                 6.24E-05
                                *** GE_Fibrosis
                                                                 0.77
```

```
## Ifnar2
                  1.82E-04
                                 *** GE_Cytokines
                                                                  0.77
## Ccr5
                  4.36E-08
                                 *** GE_Cytokines
                                                                  0.77
## Il10
                  4.88E-05
                                 *** GE_Cytokines
                                                                  0.76
## Osmr
                                 *** GE_Cytokines
                  1.01E-07
                                                                  0.75
## Gsta2
                  3.88E-06
                                          GE_ADME
                                                                  0.74
## I14
                  4.88E-05
                                 *** GE Cytokines
                                                                  0.71
## Ifng
                  7.51E-06
                                 *** GE_Cytokines
                                                                  0.71
## Cc18
                  1.23E-04
                                 *** GE_Cytokines
                                                                  0.71
## Hgf
                  5.80E-08
                                 *** GE_Cytokines
                                                                  0.70
## Bak1
                  2.73E-02
                                   * GE_Fibrosis
                                                                  0.70
## Mrc1
                  6.95E-11
                                 *** GE_Cytokines
                                                                  0.69
## Tgfb2
                  1.08E-02
                                   * GE_Fibrosis
                                                                  0.69
## Ccr3
                                 *** GE_Cytokines
                                                                  0.68
                  1.91E-06
                  1.91E-02
                                   * GE_Cytokines
                                                                  0.68
## Actb.y
## S100a4
                                                                  0.66
                  9.19E-07
                                       Antibodies
## Il13
                  9.29E-06
                                 *** GE_Cytokines
                                                                  0.66
## Met
                  4.97E-03
                                  ** GE_Cytokines
                                                                  0.66
                                        Histology
## bile.infarcts 3.78E-02
                                                        Η
                                                                  0.65
## Il6st
                  8.71E-04
                                 *** GE_Cytokines
                                                                  0.63
## Tnfrsf1a
                  3.33E-05
                                 *** GE_Cytokines
                                                                  0.63
## Mki67
                  1.46E-03
                                      GE_Fibrosis
                                                                  0.62
## Birc5
                                      GE_Fibrosis
                                                                  0.60
                  1.69E-02
## Ctgf
                  1.32E-04
                                      GE_Fibrosis
                                                                  0.58
                                 ***
## BEC
                  2.09E-07
                                        Histology
                                                        Η
                                                                  0.56
## Bax
                  2.89E-02
                                      GE Fibrosis
                                                                  0.56
## Notch1
                  4.45E-03
                                      GE_Fibrosis
                                                                  0.54
                  1.07E-05
                                 *** GE_Cytokines
                                                                  0.51
## Cxcr1
## Gstm1
                  9.18E-09
                                 ***
                                          GE_ADME
                                                                  0.45
## Cdh1
                  4.48E-04
                                      GE_Fibrosis
                                                                  0.42
##
## Cluster 5 (N=11)
##
                 p.holm sig.holm
                                         ftype fshort cluster.cor
## Gdf2
              5.95E-06
                                   GE_Fibrosis
                                                              0.84
                                       GE_ADME
                                                              0.68
## Cyp7a1
              9.83E-03
                              **
## I12
              4.84E-05
                             *** GE Cytokines
                                                              0.65
## Il17a
              2.43E-04
                             *** GE_Cytokines
                                                              0.65
## NHC
              5.28E-04
                                     Histology
                                                     Η
                                                              0.63
## Sirius.red 7.59E-04
                                     Histology
                                                     Η
                                                              0.63
                             ***
## Egf
              4.88E-05
                             *** GE_Cytokines
                                                              0.61
## Ifna1
              1.46E-03
                              ** GE_Cytokines
                                                              0.60
                             *** GE_Cytokines
## Ifnb1
                                                              0.55
              4.86E-05
## I128b
              4.88E-05
                             *** GE_Cytokines
                                                              0.51
## Kupffer
              1.95E-03
                                     Histology
                                                     Η
                                                              0.41
##
## Cluster 6 (N=2)
                                  ftype fshort cluster.cor
##
          p.holm sig.holm
## Cdh2 5.38E-04
                      *** GE_Fibrosis
                                                       0.89
## Bad 3.84E-04
                       *** GE_Fibrosis
                                                       0.86
```

Decision trees

Introduction

For the prediction of disease progression after BDL a decision tree was used. The regression tree was fitted with the R package rpart, being the open-source implementation of CART, implementing algorithms for recursive partitioning for classification following in most details closely Breiman et. al (1984) (Breiman L., Friedman J. H., Olshen R. A., and Stone, C. J. (1984) Classification and Regression Trees. Wadsworth).

The predictor variables are the 6 mean time courses of the clusters, the dependent variable is the log transformed time class, to get approximately equidistant intervals between the trainings classes in the regression models. The tree is hereby built in a two-step process: First the single variable is found which best splits the data into two groups. The data than separated based on the split, and the splitting process is applied separately to each sub-group, and so on recursively until the subgroups either reach a minimum size (minbucket) or until no improvement can be made. The second stage of the procedure consists of using cross-validation to trim back the full tree. The splitting criterion, which is used to decide which variable gives the best split for nodes in the regression trees is $SS_T - (SS_L + SS_R)$, with $SS_T = \sum (y_i - \langle y \rangle)^2$ the sum of squares for the node and SS_R and SS_L the sums of squares for the left and right son. This is equivalent to choosing the split of maximize the between-groups sum-of-squares in a simple analysis of variance (see An Introduction to Recursive Partitioning Using the RPART Routines. T.M. Therneau and E.J. Atkinson, Mayo Foundation, 2015). Two important parameters controlling the resulting tree are

- minsplit: The minimum number of observations in a node for which the routine will even try to compute a split. The default is 20 and is set to 6 in the tree calculation ($N_r = 5$ repeats per time point).
- minbucket: The minimum number of observations in a terminal node. This defaults to minsplit/3.

```
suppressPackageStartupMessages(library(rpart))
suppressPackageStartupMessages(library(rpart.plot))
suppressPackageStartupMessages(library(caret))
dir.create(file.path(resultsPath, 'decision_tree'), showWarnings=FALSE)
```

To generate approximatly equally distant time classes for training the regression tree the log transformed time values are used. The transformation used is

$$\tilde{t_i} = log(t_i + 1)$$

```
# Transform data to log scale (for comparable time intervals)
log_transform <- function(data){
  log(data+1)
}
# Back transformation
log_transform_back <- function(log_data){
  exp(log_data)-1
}
# Resulting time transformations
time_transformation <- data.frame(time=BDLmean.time, log_time=log_transform(BDLmean.time))
print(round(time_transformation,2))</pre>
```

```
## time log_time
## 1 0 0.00
## 2 6 1.95
## 3 12 2.56
```

```
## 4 18 2.94
## 5 30 3.43
## 6 48 3.89
## 7 120 4.80
## 8 336 5.82
```

Trainings data

In a first step the mean cluster trainings data for fitting the decision tree is prepared. The predictor variables are the mean samples of the clusters, the dependent variable is the log transformed time of the respective sample. The trainings set consists of the $N_t * N_r = 40$ samples.

```
prepare_treedata_mean <- function(method="ys3"){</pre>
  # Hierarchical clusters based on ys3 to fit the regression tree
  hc.res <- hclust.res[[method]]</pre>
  groups <- hc.res$groups</pre>
  # Prepare training set for fitting the decision trees (mean cluster data set,
  # i.e. mean over normalized factors in cluster).
  na.vec <- rep(NA, Nt*Nr)</pre>
  treedata.mean <- data.frame(c1=na.vec, c2=na.vec, c3=na.vec, c4=na.vec, c5=na.vec, c6=na.vec)
  # for every sample
  for (ks in 1:(Nt*Nr)){
    # create the mean over the cluster
   for (kgroup in 1:Ngroups){
      # get factors in the cluster
      factors <- names(groups[groups==kgroup])</pre>
      # calculate mean over normalized data
      treedata.mean[ks, kgroup] <- mean(as.numeric(BDLdata.norm[ks, factors]), na.rm=TRUE)
   }
  }
  # add log transformed time [h] for regression
  treedata.mean$logtime <- log_transform(BDLsamples$time)</pre>
  # add experimental time class
  treedata.mean$class <- BDLsamples$time_fac</pre>
  return(treedata.mean)
}
treedata.mean <- prepare_treedata_mean()</pre>
# mean cluster data set for model fitting
print(round(treedata.mean[, -8], digits=2))
##
               c2
                     сЗ
                           c4
                                  с5
                                        c6 logtime
## 1
       0.22 -0.07 -0.36 -0.14 -0.11 -0.03
                                              0.00
## 2
       0.50 -0.06 -0.29 -0.14 -0.07 -0.07
                                              0.00
       0.26 -0.07 -0.32 -0.19 -0.10 -0.17
                                              0.00
## 3
## 4
       0.34 -0.07 -0.31 -0.19 -0.04 -0.23
                                              0.00
       0.12 -0.07 -0.33 -0.19 -0.02 -0.18
## 5
                                              0.00
## 6
       0.31 0.69 -0.01 -0.17 -0.09 -0.29
                                              1.95
## 7
       0.21 0.93 -0.02 -0.18 -0.13 -0.36
                                              1.95
## 8
       0.24 0.27 0.03 -0.18 -0.15 -0.30
                                              1.95
       ## 9
                                              1.95
```

```
1.95
## 11
      0.01 -0.07 0.23 -0.13 -0.15 -0.01
                                             2.56
                  0.01 -0.15 -0.11 -0.07
                                             2.56
      0.08 - 0.07
## 13 -0.07 -0.07
                  0.12 -0.17 -0.15 -0.18
                                             2.56
      0.07 -0.07
                   0.09 -0.15 -0.14
                                             2.56
## 15
      0.01 - 0.07
                  0.13 -0.15 -0.15 -0.08
                                             2.56
      0.13 - 0.07
                  0.30 -0.05 -0.16
                                             2.94
## 17 -0.03 -0.07
                  0.39 -0.12 -0.15
                                     0.24
                                             2.94
## 18
      0.02 -0.07
                   0.22 -0.13 -0.15
                                     0.07
                                             2.94
## 19
      0.02 - 0.07
                  0.43 -0.07 -0.15
                                     0.20
                                             2.94
## 20
      0.01 -0.07 0.22 -0.15 -0.14
                                     0.12
                                             2.94
      0.05 -0.07 -0.01 -0.09 -0.04
## 21
                                     0.14
                                             3.43
## 22 -0.02 -0.07 -0.03 -0.07 -0.10
                                     0.08
                                             3.43
                  0.41 -0.11 -0.16
## 23 -0.11 -0.07
                                     0.03
                                             3.43
## 24 -0.11 -0.07
                  0.13 -0.03 -0.12
                                     0.09
                                             3.43
      0.03 - 0.07
                  0.17 -0.08 -0.05
                                     0.23
                                             3.43
## 26 -0.13 -0.07
                  0.01
                         0.02 - 0.04
                                             3.89
                                     0.11
## 27 -0.17 -0.07
                  0.26
                         0.13 0.00
                                             3.89
## 28 -0.09 -0.07 -0.15
                        0.00 - 0.04
                                    0.28
                                             3.89
## 29 -0.08 -0.07
                  0.12 -0.11 -0.14
                                             3.89
## 30 -0.03 -0.07 0.12 -0.08 -0.02
                                    0.12
                                             3.89
## 31 -0.23 -0.07 -0.27 -0.01
                                             4.80
## 32 -0.26 -0.07 0.11
                         0.27
                               0.17
                                     0.27
                                             4.80
## 33 -0.28 -0.07 -0.27
                         0.29
                               0.19 - 0.13
                                             4.80
## 34 -0.26 -0.07 -0.23
                         0.28
                               0.23 - 0.13
                                             4.80
## 35 -0.10 -0.01 0.07
                         0.33
                               0.44
                                    0.06
                                             4.80
## 36 -0.16 0.03 -0.17
                         0.45
                                             5.82
                               0.44 - 0.06
## 37 -0.22 -0.06 -0.14
                         0.64
                               0.49 - 0.01
                                             5.82
## 38 -0.23 -0.07 -0.13
                         0.23
                               0.26 - 0.07
                                             5.82
## 39 -0.24 -0.07 -0.17
                         0.54
                               0.44 0.12
                                             5.82
## 40 -0.32 -0.07 -0.27
                         0.36
                               0.30 - 0.11
                                             5.82
# save the trainings data
save(treedata.mean, file=file.path(resultsPath, "data", "treedata.mean.Rdata"))
```

Fit regression tree

Fit the log transformed time against the mean cluster data for the time points. Set control parameters for the algorithm are minsplit=6 and minbucket=2 to account for the low number of samples (N=40). In the following the regression tree is fitted and the information for the resulting tree is provided.

```
## n= 40
##
## node), split, n, deviance, yval
         * denotes terminal node
##
##
   1) root 40 1.111047e+02 3.174622
##
      2) c4< -0.1174939 18 2.310447e+01 1.743756
##
        4) c3< -0.2081362 5 0.000000e+00 0.000000 *
##
##
        5) c3>=-0.2081362 13 2.053563e+00 2.414432
         10) c1>=0.1331057 5 2.465190e-31 1.945910 *
##
##
         11) c1< 0.1331057 8 2.700232e-01 2.707258 *
##
      3) c4>=-0.1174939 22 2.099513e+01 4.345331
##
        6) c5< 0.07192204 12 1.384347e+00 3.543160 *
        7) c5>=0.07192204 10 2.622937e+00 5.307937
##
##
         14) c4< 0.3463259 6 8.743124e-01 4.966506 *
##
         15) c4>=0.3463259 4 0.000000e+00 5.820083 *
summary(tree.reg)
## Call:
## rpart(formula = formula.reg, data = treedata.mean, method = "anova",
##
       control = rpart.control(minsplit = 6, minbucket = 2))
##
    n = 40
##
             CP nsplit rel error
                                     xerror
                                                   xstd
## 1 0.60308067
                     0 1.00000000 1.0304620 0.20523641
## 2 0.18946910
                     1 0.39691933 0.4673352 0.06894468
## 3 0.15289945
                     2 0.20745023 0.3681693 0.07373592
                     3 0.05455078 0.1806307 0.06369203
## 4 0.01605279
## 5 0.01573853
                     4 0.03849799 0.1849049 0.06332192
## 6 0.01000000
                     5 0.02275946 0.1348209 0.04821264
## Variable importance
## c4 c5 c1 c3 c6 c2
## 27 19 19 15 12 8
## Node number 1: 40 observations,
                                      complexity param=0.6030807
     mean=3.174622, MSE=2.777617
     left son=2 (18 obs) right son=3 (22 obs)
##
##
    Primary splits:
##
         c4 < -0.1174939
                           to the left, improve=0.6030807, (0 missing)
##
         c1 < -0.07537026 to the right, improve=0.5729499, (0 missing)
         c5 < -0.008508656 to the left, improve=0.5487412, (0 missing)
##
##
         c3 < -0.2829117
                           to the left, improve=0.5183388, (0 missing)
##
         c6 < -0.149531
                           to the left, improve=0.3184694, (0 missing)
##
     Surrogate splits:
##
         c1 < -0.07537026 to the right, agree=0.850, adj=0.667, (0 split)
##
         c5 < -0.0625688
                           to the left, agree=0.800, adj=0.556, (0 split)
##
         c6 < -0.149531
                           to the left, agree=0.750, adj=0.444, (0 split)
                           to the right, agree=0.675, adj=0.278, (0 split)
##
         c2 < 0.07279386
##
         c3 < -0.2829117
                           to the left, agree=0.675, adj=0.278, (0 split)
##
## Node number 2: 18 observations,
                                      complexity param=0.1894691
    mean=1.743756, MSE=1.283582
##
```

```
##
     left son=4 (5 obs) right son=5 (13 obs)
##
    Primary splits:
##
         c3 < -0.2081362
                           to the left, improve=0.9111184, (0 missing)
                          to the right, improve=0.5991729, (0 missing)
##
         c5 < -0.1122942
##
         c1 < 0.09730379
                           to the right, improve=0.5785906, (0 missing)
##
         c4 < -0.1838753
                           to the left, improve=0.4737815, (0 missing)
         c6 < -0.01908181 to the left, improve=0.3296556, (0 missing)
##
     Surrogate splits:
##
         c4 < -0.1838753
                           to the left, agree=0.889, adj=0.6, (0 split)
##
         c5 < -0.1122942
                           to the right, agree=0.889, adj=0.6, (0 split)
##
         c1 < 0.3351088
                           to the right, agree=0.833, adj=0.4, (0 split)
##
## Node number 3: 22 observations,
                                      complexity param=0.1528994
     mean=4.345331, MSE=0.9543241
##
##
     left son=6 (12 obs) right son=7 (10 obs)
##
     Primary splits:
##
         c5 < 0.07192204
                           to the left, improve=0.8091327, (0 missing)
##
         c4 < 0.1840943
                           to the left, improve=0.7540271, (0 missing)
##
         c1 < -0.1437557
                           to the right, improve=0.6642991, (0 missing)
##
         c3 < -0.08194432 to the right, improve=0.6127738, (0 missing)
##
         c6 < 0.007982308 to the right, improve=0.5246106, (0 missing)
##
     Surrogate splits:
##
         c4 < 0.1840943
                           to the left, agree=0.955, adj=0.9, (0 split)
                           to the right, agree=0.909, adj=0.8, (0 split)
##
         c1 < -0.1936876
##
         c3 < -0.08194432 to the right, agree=0.864, adj=0.7, (0 split)
##
         c6 < 0.007982308
                           to the right, agree=0.864, adj=0.7, (0 split)
##
         c2 < -0.07256249 to the right, agree=0.727, adj=0.4, (0 split)
## Node number 4: 5 observations
##
    mean=0, MSE=0
##
## Node number 5: 13 observations,
                                      complexity param=0.01605279
##
     mean=2.414432, MSE=0.1579664
     left son=10 (5 obs) right son=11 (8 obs)
##
##
     Primary splits:
         c1 < 0.1331057
##
                           to the right, improve=0.8685099, (0 missing)
##
         c2 < 0.02364916
                           to the right, improve=0.8685099, (0 missing)
##
         c3 < 0.06868251
                           to the left, improve=0.7242745, (0 missing)
                           to the left, improve=0.7242745, (0 missing)
##
         c6 < -0.09746977
##
         c5 < -0.1383548
                           to the right, improve=0.4701461, (0 missing)
##
     Surrogate splits:
##
         c2 < 0.02364916
                           to the right, agree=1.000, adj=1.0, (0 split)
                           to the left, agree=0.923, adj=0.8, (0 split)
##
         c3 < 0.06868251
##
         c6 < -0.2167742
                           to the left, agree=0.923, adj=0.8, (0 split)
##
         c4 < -0.153395
                           to the left, agree=0.846, adj=0.6, (0 split)
                           to the right, agree=0.846, adj=0.6, (0 split)
##
         c5 < -0.1383548
## Node number 6: 12 observations
##
     mean=3.54316, MSE=0.1153623
##
## Node number 7: 10 observations,
                                      complexity param=0.01573853
    mean=5.307937, MSE=0.2622937
##
##
    left son=14 (6 obs) right son=15 (4 obs)
##
    Primary splits:
```

```
##
         c4 < 0.3463259
                          to the left, improve=0.6666667, (0 missing)
##
        c5 < 0.2491244
                          to the left, improve=0.6666667, (0 missing)
##
        c6 < -0.1217972 to the left, improve=0.2500000, (0 missing)
                          to the right, improve=0.2500000, (0 missing)
##
         c3 < -0.0328633
##
        c1 < -0.247735
                           to the left, improve=0.1666667, (0 missing)
##
    Surrogate splits:
                           to the left, agree=0.9, adj=0.75, (0 split)
##
        c5 < 0.2821139
                           to the left, agree=0.7, adj=0.25, (0 split)
##
        c1 < -0.2249667
##
        c2 < -0.06371928 to the left, agree=0.7, adj=0.25, (0 split)
##
        c3 < -0.1376999
                           to the right, agree=0.7, adj=0.25, (0 split)
##
         c6 < -0.06444851 to the left, agree=0.7, adj=0.25, (0 split)
##
## Node number 10: 5 observations
##
    mean=1.94591, MSE=4.930381e-32
##
## Node number 11: 8 observations
     mean=2.707258, MSE=0.0337529
##
##
## Node number 14: 6 observations
    mean=4.966506, MSE=0.1457187
##
## Node number 15: 4 observations
    mean=5.820083, MSE=0
##
printcp(tree.reg)
##
## Regression tree:
## rpart(formula = formula.reg, data = treedata.mean, method = "anova",
       control = rpart.control(minsplit = 6, minbucket = 2))
##
## Variables actually used in tree construction:
## [1] c1 c3 c4 c5
##
## Root node error: 111.1/40 = 2.7776
##
## n=40
##
          CP nsplit rel error xerror
## 1 0.603081
                  0 1.000000 1.03046 0.205236
## 2 0.189469
                  1 0.396919 0.46734 0.068945
## 3 0.152899
                  2 0.207450 0.36817 0.073736
## 4 0.016053
                  3 0.054551 0.18063 0.063692
                  4 0.038498 0.18490 0.063322
## 5 0.015739
## 6 0.010000
                  5 0.022759 0.13482 0.048213
tree.reg$frame
##
                            dev
                                    yval complexity ncompete nsurrogate
## 1
         c4 40 40 1.111047e+02 3.174622 0.603080672
## 2
         c3 18 18 2.310447e+01 1.743756 0.189469098
                                                            4
                                                                       3
## 4 <leaf> 5 5 0.000000e+00 0.000000 0.010000000
                                                                       0
         c1 13 13 2.053563e+00 2.414432 0.016052788
                                                                       5
## 5
```

```
# pretty plot of tree to file
pdf(file.path(resultsPath, 'decision_tree', "regression_tree.pdf"),
    width=10, height=5, pointsize=12)
prp(tree.reg, type=0, extra=101, yesno=TRUE)
invisible(dev.off())

# and report
prp(tree.reg, type=0, extra=101, yesno=TRUE)
```

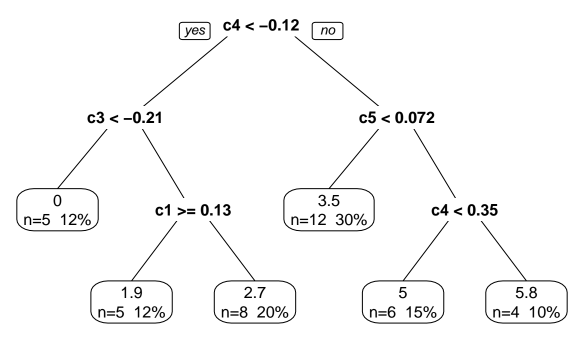


Figure 5:

```
# visualize cross-validation results
# rsq.rpart(tree.reg)
```

Only a subset of all mean clusters is used in the regression tree:

```
# variables used for splitting in the decision tree
tree.nodes <- (tree.reg$frame)$var
tree.nodes <- tree.nodes[tree.nodes != "<leaf>"]
tree.vars <- as.character(sort(unique(tree.nodes)))
rm(tree.nodes)
# variables used for splitting decisions in tree
print(tree.vars)</pre>
```

```
## [1] "c1" "c3" "c4" "c5"
```

Perform leave one out cross validation

```
# fit the 5 repeat regression trees (leave one out)
# in total 40 cross validations ! because mice are not time followed
trees.test <- vector("list", length=Nr*Nt)</pre>
pred.test.log <- rep(NA, length=Nr*Nt)</pre>
for (k in 1:(Nr*Nt)){
 idx.subset <- 1:(Nr*Nt)</pre>
 idx.subset <- idx.subset[-k]</pre>
 print(idx.subset)
 # fit tree with the subset
 t.test <- rpart(formula=formula.reg,</pre>
                 data=treedata.mean[idx.subset, ],
                 method="anova",
                 control=rpart.control(minsplit=6, minbucket=2))
 trees.test[[k]] <- t.test</pre>
 # prediction on left out sample
 pred.test.log[k] <- predict(t.test, newdata=treedata.mean[k,], type="vector")</pre>
}
   [1]
              4 5 6 7 8 9 10 11 12 13 14 15 16 17 18 19 20 21 22 23 24
## [24] 25 26 27 28 29 30 31 32 33 34 35 36 37 38 39 40
   [1]
        1
           3
              4
                 5
                    6 7 8 9 10 11 12 13 14 15 16 17 18 19 20 21 22 23 24
## [24] 25 26 27 28 29 30 31 32 33 34 35 36 37 38 39 40
  [1]
        1 2 4 5 6 7 8 9 10 11 12 13 14 15 16 17 18 19 20 21 22 23 24
## [24] 25 26 27 28 29 30 31 32 33 34 35 36 37 38 39 40
   [1]
        1 2 3 5 6 7 8 9 10 11 12 13 14 15 16 17 18 19 20 21 22 23 24
## [24] 25 26 27 28 29 30 31 32 33 34 35 36 37 38 39 40
   [1]
       1 2 3 4 6 7 8 9 10 11 12 13 14 15 16 17 18 19 20 21 22 23 24
## [24] 25 26 27 28 29 30 31 32 33 34 35 36 37 38 39 40
       1 2 3 4 5 7 8 9 10 11 12 13 14 15 16 17 18 19 20 21 22 23 24
  [1]
## [24] 25 26 27 28 29 30 31 32 33 34 35 36 37 38 39 40
## [1]
        1 2 3 4 5 6 8 9 10 11 12 13 14 15 16 17 18 19 20 21 22 23 24
## [24] 25 26 27 28 29 30 31 32 33 34 35 36 37 38 39 40
       1 2 3 4 5 6 7 9 10 11 12 13 14 15 16 17 18 19 20 21 22 23 24
   [1]
## [24] 25 26 27 28 29 30 31 32 33 34 35 36 37 38 39 40
   [1]
                   5 6 7 8 10 11 12 13 14 15 16 17 18 19 20 21 22 23 24
## [24] 25 26 27 28 29 30 31 32 33 34 35 36 37 38 39 40
        1 2 3 4
                   5 6 7 8 9 11 12 13 14 15 16 17 18 19 20 21 22 23 24
  [1]
## [24] 25 26 27 28 29 30 31 32 33 34 35 36 37 38 39 40
        1 2 3 4 5 6 7 8 9 10 12 13 14 15 16 17 18 19 20 21 22 23 24
## [1]
## [24] 25 26 27 28 29 30 31 32 33 34 35 36 37 38 39 40
       1 2 3 4 5 6 7 8 9 10 11 13 14 15 16 17 18 19 20 21 22 23 24
  [1]
## [24] 25 26 27 28 29 30 31 32 33 34 35 36 37 38 39 40
                   5 6 7 8 9 10 11 12 14 15 16 17 18 19 20 21 22 23 24
   [1]
        1 2 3
                4
## [24] 25 26 27 28 29 30 31 32 33 34 35 36 37 38 39 40
## [1]
        1 2 3 4
                   5 6 7 8 9 10 11 12 13 15 16 17 18 19 20 21 22 23 24
## [24] 25 26 27 28 29 30 31 32 33 34 35 36 37 38 39 40
## [1]
       1 2 3 4 5 6 7 8 9 10 11 12 13 14 16 17 18 19 20 21 22 23 24
```

```
## [24] 25 26 27 28 29 30 31 32 33 34 35 36 37 38 39 40
   [1] 1 2 3 4 5 6 7 8 9 10 11 12 13 14 15 17 18 19 20 21 22 23 24
## [24] 25 26 27 28 29 30 31 32 33 34 35 36 37 38 39 40
                  5 6 7 8 9 10 11 12 13 14 15 16 18 19 20 21 22 23 24
   [1]
## [24] 25 26 27 28 29 30 31 32 33 34 35 36 37 38 39 40
       1 2 3 4 5 6 7 8 9 10 11 12 13 14 15 16 17 19 20 21 22 23 24
##
   [1]
## [24] 25 26 27 28 29 30 31 32 33 34 35 36 37 38 39 40
       1 2 3 4 5 6 7 8 9 10 11 12 13 14 15 16 17 18 20 21 22 23 24
   [1]
## [24] 25 26 27 28 29 30 31 32 33 34 35 36 37 38 39 40
   [1]
       1 2 3 4 5 6 7 8 9 10 11 12 13 14 15 16 17 18 19 21 22 23 24
## [24] 25 26 27 28 29 30 31 32 33 34 35 36 37 38 39 40
          2 3 4 5 6 7 8 9 10 11 12 13 14 15 16 17 18 19 20 22 23 24
   [1]
       1
## [24] 25 26 27 28 29 30 31 32 33 34 35 36 37 38 39 40
   [1]
       1 2 3 4 5 6 7 8 9 10 11 12 13 14 15 16 17 18 19 20 21 23 24
## [24] 25 26 27 28 29 30 31 32 33 34 35 36 37 38 39 40
##
   [1]
       1 2 3 4 5 6 7 8 9 10 11 12 13 14 15 16 17 18 19 20 21 22 24
## [24] 25 26 27 28 29 30 31 32 33 34 35 36 37 38 39 40
   [1]
       1 2 3 4 5 6 7 8 9 10 11 12 13 14 15 16 17 18 19 20 21 22 23
## [24] 25 26 27 28 29 30 31 32 33 34 35 36 37 38 39 40
   [1]
       1
          2 3 4 5 6 7 8 9 10 11 12 13 14 15 16 17 18 19 20 21 22 23
## [24] 24 26 27 28 29 30 31 32 33 34 35 36 37 38 39 40
       1 2 3 4 5 6 7 8 9 10 11 12 13 14 15 16 17 18 19 20 21 22 23
   [1]
## [24] 24 25 27 28 29 30 31 32 33 34 35 36 37 38 39 40
       1 2 3 4 5 6 7 8 9 10 11 12 13 14 15 16 17 18 19 20 21 22 23
##
   [1]
## [24] 24 25 26 28 29 30 31 32 33 34 35 36 37 38 39 40
   [1]
       1 2 3 4 5 6 7 8 9 10 11 12 13 14 15 16 17 18 19 20 21 22 23
## [24] 24 25 26 27 29 30 31 32 33 34 35 36 37 38 39 40
       1 2 3 4 5 6 7 8 9 10 11 12 13 14 15 16 17 18 19 20 21 22 23
   [1]
## [24] 24 25 26 27 28 30 31 32 33 34 35 36 37 38 39 40
   [1] 1 2 3 4 5 6 7 8 9 10 11 12 13 14 15 16 17 18 19 20 21 22 23
## [24] 24 25 26 27 28 29 31 32 33 34 35 36 37 38 39 40
##
   [1]
       1 2 3 4 5 6 7 8 9 10 11 12 13 14 15 16 17 18 19 20 21 22 23
## [24] 24 25 26 27 28 29 30 32 33 34 35 36 37 38 39 40
               4 5 6 7 8 9 10 11 12 13 14 15 16 17 18 19 20 21 22 23
   [1]
       1 2 3
## [24] 24 25 26 27 28 29 30 31 33 34 35 36 37 38 39 40
   [1]
       1 2 3 4 5 6 7 8 9 10 11 12 13 14 15 16 17 18 19 20 21 22 23
##
## [24] 24 25 26 27 28 29 30 31 32 34 35 36 37 38 39 40
       1 2 3 4 5 6 7 8 9 10 11 12 13 14 15 16 17 18 19 20 21 22 23
##
  [1]
## [24] 24 25 26 27 28 29 30 31 32 33 35 36 37 38 39 40
       1 2 3 4 5 6 7 8 9 10 11 12 13 14 15 16 17 18 19 20 21 22 23
##
   [1]
## [24] 24 25 26 27 28 29 30 31 32 33 34 36 37 38 39 40
       1 2 3 4 5 6 7 8 9 10 11 12 13 14 15 16 17 18 19 20 21 22 23
   [1]
## [24] 24 25 26 27 28 29 30 31 32 33 34 35 37 38 39 40
       1 2 3 4 5 6 7 8 9 10 11 12 13 14 15 16 17 18 19 20 21 22 23
   [1]
## [24] 24 25 26 27 28 29 30 31 32 33 34 35 36 38 39 40
       1 2 3 4 5 6 7 8 9 10 11 12 13 14 15 16 17 18 19 20 21 22 23
   [1]
## [24] 24 25 26 27 28 29 30 31 32 33 34 35 36 37 39 40
   [1]
       1 2 3 4 5 6 7 8 9 10 11 12 13 14 15 16 17 18 19 20 21 22 23
## [24] 24 25 26 27 28 29 30 31 32 33 34 35 36 37 38 40
   [1]
        1 2 3 4 5 6 7 8 9 10 11 12 13 14 15 16 17 18 19 20 21 22 23
## [24] 24 25 26 27 28 29 30 31 32 33 34 35 36 37 38 39
# transformation to time in [h]
pred.test <- log_transform_back( pred.test.log )</pre>
```

```
# plot the log prediction and error
par(mfrow=c(1,2))
plot(log_transform(BDLsamples$time), log_transform(pred.test),
     main="Leave one out cross validation")
abline(a=0, b=1, col="gray")
textxy(log_transform(BDLsamples$time), log_transform(pred.test), 1:(Nr*Nt), col="black", cex=0.6)
plot(log transform(BDLsamples$time), log transform(BDLsamples$time)-log transform(pred.test),
     main="Leave one out cross validation error")
abline(a=0, b=0, col="gray")
textxy(log_transform(BDLsamples$time), log_transform(BDLsamples$time)-log_transform(pred.test),
       1:(Nr*Nt), col="black", cex=0.6)
```

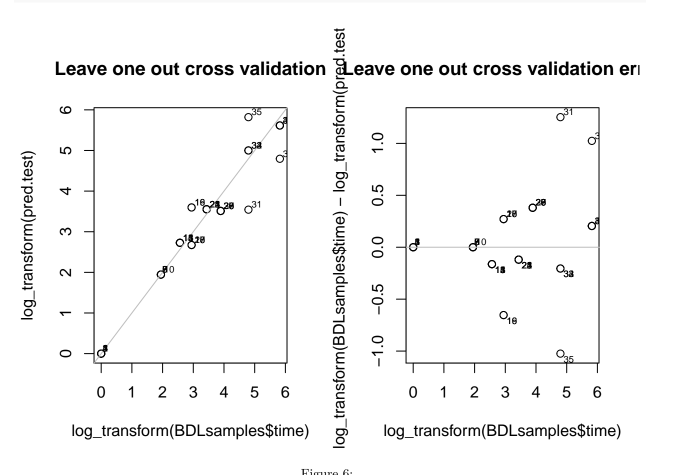


Figure 6:

```
par(mfrow=c(1,1))
```

Prediction on trainings data

Predict data with the tree, here the time class leaves based on the mean cluster data. The regression tree predicts log classes, which are back-transformed to time in [h]. So how good is the tree performing on the trainings data set, i.e. the mean cluster data. Predictons are evaluated based on the distance between the predicted and the experimental time classes based on the following distance measure on log scale

$$d = \frac{1}{N_s} \sqrt{\sum TODO}$$

```
# L2 (euclidian) distance measurement on the log transformed data. Analog to the distance measurement i
# the regression tree.
log_distance <- function(d1, d2){
    # sums over all the distances of the samples in log space
    log_rmsd <- sqrt(sum( (log_transform(d1)-log_transform(d2) )^2 ))/length(d1)
    return(log_rmsd)
}</pre>
```

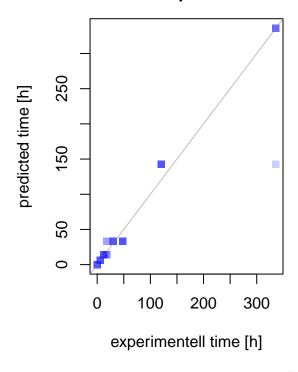
Prediction on trainings data

```
# mean cluster predictions
pred.mean.log <- predict(tree.reg, newdata=treedata.mean, type="vector")</pre>
# transformation to time in [h]
pred.mean <- log_transform_back( pred.mean.log )</pre>
# TODO: create the overview matrix which experimental time classes are predicted in which
# tree time classes (on normal and log scale)
# TODO: heatmap of the matrix
# pred.matrix.mean <- matrix(NA, nrow=nrow(node ranges), ncol=length(levels(as.factor(BDLsamples$time))
# Distance calculation (predicted to experimental)
dist.mean.all <- treedata.mean$logtime - pred.mean.log</pre>
dist.mean <- log_distance(pred.mean, BDLsamples$time)</pre>
# plot predicted ~ experimentell
plot_mean_prediction <- function(){</pre>
  par(mfrow=c(1,2))
  plot(BDLsamples$time, pred.mean, pch=15, col=rgb(0,0,1, 0.2), main="Regression Tree:\nPredicted ~ exp
       xlab="experimentell time [h]", ylab="predicted time [h]")
  abline(a=0, b=1, col=rgb(0.5,0.5,0.5,0.5))
  hist(dist.mean.all, breaks=seq(from=-1.05, to=1.05, by=0.1),
       main="Histogram prediction error:\nmean cluster data",
       col=rgb(0.5,0.5,0.5,0.5),
       xlab="logtime(exp)-logtime(pred)")
  par(mfrow=c(1,1))
}
# plot to file
pdf(file.path(resultsPath, 'decision_tree', "prediction_mean.pdf"),
    width=10, height=5, pointsize=12)
plot_mean_prediction()
invisible(dev.off())
# and report
plot_mean_prediction()
```

The ranges of the predicted classes by the regression trees can be calculated based on the split points on log scale. These provides the information which time points would be classified in which class in the regression tree.

Regression Tree: Predicted ~ experimentell time

Histogram prediction error: mean cluster data



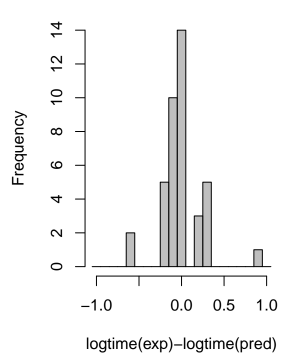


Figure 7:

```
calculate_node_ranges <- function(){</pre>
  # classes via predicted classes for cluster data
  node_levels <- levels(as.factor(pred.mean))</pre>
  # These are the predicted classes
  node_classes <- round(as.numeric(node_levels), digits=1)</pre>
  # get the intervals of the time classes
  node_mean <- as.numeric(levels(as.factor(pred.mean.log)))</pre>
  node_midpoints <- (node_mean[2:length(node_mean)] + node_mean[1:(length(node_mean)-1)])/2
  # minimum of range
  node_min <- node_mean</pre>
  node_min[2:length(node_min)] <- node_midpoints</pre>
  # maximum of range
  node_max <- node_mean
  node_max[1:(length(node_min)-1)] <- node_midpoints</pre>
  # ranges in log scale
  node_ranges.log <- data.frame(node_mean, node_min, node_max)</pre>
  node_ranges <- data.frame(mean=log_transform_back(node_mean),</pre>
                              min=log_transform_back(node_min),
                              max=log_transform_back(node_max))
  rownames(node_ranges) <- paste("class", 1:nrow(node_ranges))</pre>
  return(node_ranges)
}
```

```
# predicted time classes by decision tree
node_ranges <- calculate_node_ranges()
print(round(node_ranges, digits=1))</pre>
```

```
## class 1 0.0 0.0 1.6
## class 2 6.0 1.6 9.2
## class 3 14.0 9.2 21.8
## class 4 33.6 21.8 69.4
## class 5 142.5 69.4 218.9
## class 6 336.0 218.9 336.0
```

Test data for evaluation

Single factor per cluster

Prepare the evaluation data sets for the fitted trees consisting of all single factor combinations from the clusters used in the decision tree. Only the combinations are created which are from clusters used for decisions in the tree

```
# names of factors in the clusters
cluster_names <- paste('c', 1:Ngroups, sep="")
cluster.factors <- vector("list", length=Ngroups)
groups <- (hclust.res$ys3)$groups # get the ys3 groups
for (k in 1:Ngroups){
    cluster.factors[[k]] <- as.character(names(groups[groups==k]))
}
names(cluster.factors) <- cluster_names

# create data.frame of all single combinations from clusters
single.combinations <- expand.grid(cluster.factors, stringsAsFactors=TRUE)
names(single.combinations) <- names(cluster.factors)
# number of single combinations
Nsingle <- nrow(single.combinations)
print(Nsingle) # 88572 combinations</pre>
```

[1] 88572

```
# create all single factor data
print("Calculating single factor data (~ 3min) ... ")
```

[1] "Calculating single factor data (~ 3min) ... "

```
# get factor data
  tmp <- BDLdata.norm[, t(single.combinations[k, ]) ]</pre>
  # add regression values
  tmp[c("class", "logtime")] <- treedata.mean[ c("class", "logtime")]</pre>
  # add factor fields
  tmp[, paste(cluster_names, '.id', sep="")] <- single.combinations[k,]</pre>
  colnames(tmp) <- c(cluster_names, 'class', 'logtime', paste(cluster_names, '.id', sep=""))</pre>
  # store data
  treedata.single[[k]] <- tmp</pre>
  # if (k%%500 == 0){print(k)}
  # print(proc.time()-ptm) # Stop the clock
# Stop the clock
print(proc.time() - ptm)
      user system elapsed
## 114.019
            0.076 114.021
print("... calculated.")
## [1] "... calculated."
rm(tmp,k)
# treedata.single[[1]]
# which factor combinations only use genes
factor_is_gene <- BDLfactors$ftype %in% c("GE_ADME", "GE_Cytokines", "GE_Fibrosis")</pre>
names(factor_is_gene) <- BDLfactors$id</pre>
# vector for lookup if only genes were used
gene_only.single <- vector("logical", Nsingle)</pre>
for (k in 1:Nsingle){
  gene_only.single[k] <- all(factor_is_gene[t(single.combinations[k,])])</pre>
}
rm(k)
Double factor per cluster
```

Create a sample of double combinations from the various clusters.

```
print("Calculating double factor data (~ 1min) ... ")
## [1] "Calculating double factor data (~ 1min) ... "
ptm <- proc.time()</pre>
set.seed(123456)
Ndouble <- 10000
```

```
treedata.double <- vector("list", Ndouble) # list for sampled double combinations
for (k in 1:Ndouble){
  # sample from the 4 clusters without replacement
 n1 = sample(cluster.factors[[1]], 2, replace=FALSE)
 n2 = sample(cluster.factors[[2]], 2, replace=FALSE)
  n3 = sample(cluster.factors[[3]], 2, replace=FALSE)
  n4 = sample(cluster.factors[[4]], 2, replace=FALSE)
 n5 = sample(cluster.factors[[5]], 2, replace=FALSE)
  n6 = sample(cluster.factors[[6]], 2, replace=FALSE)
  # The mean of the combination is used (handle NAs)
  tmp \leftarrow 0.5 * (BDLdata.norm[, c(n1[1], n2[1], n3[1], n4[1], n5[1], n6[1])]
                + BDLdata.norm[, c(n1[2], n2[2], n3[2], n4[2], n5[2], n6[2])])
   # add class and regression values
  tmp[c("class", "logtime")] <- treedata.mean[ c("class", "logtime")]</pre>
  # add factor fields
  tmp[ , paste(cluster_names, '.id', sep="")] <- data.frame(paste(n1, collapse="__"),</pre>
                                               paste(n2, collapse="__"),
                                               paste(n3, collapse="_"),
                                               paste(n4, collapse="__"),
                                               paste(n5, collapse="__"),
                                               paste(n6, collapse="__"))
  colnames(tmp) <- c(cluster_names, 'class', 'regvalue', paste(cluster_names, '.id', sep=""))</pre>
  # store data
  treedata.double[[k]] <- tmp</pre>
print(proc.time() - ptm)
      user system elapsed
##
            0.048 31.967
   31.942
print("... calculated.")
## [1] "... calculated."
rm(k, tmp)
# treedata.double[[5]]
```

Prediction on test data

Single representative predictions

Time class prediction with regression tree for single representative from each cluster

```
print("Predicting single factor data (~ 2min) ... ")
## [1] "Predicting single factor data (~ 2min) ... "
```

```
pred.single.all <- vector("list", length(treedata.single))
for (k in (1:length(treedata.single))){
    # prediction and back transformation
    pred.single.all[[k]] <- log_transform_back( predict(tree.reg, newdata=treedata.single[[k]], method="a")
}
pred.single <- do.call("rbind", pred.single.all)

# distance for all predictions on single factor per cluster
dist.single <- rep(NA, Nsingle)
for (k in 1:Nsingle){
    dist.single[k] <- log_distance(pred.single[k,], BDLsamples$time)
}</pre>
```

Double representative predictions

Time class prediction with regression tree for random selection of two representatives from each clusters

```
print("Predicting double factor data (~ 1min) ... ")

## [1] "Predicting double factor data (~ 1min) ... "

pred.double.all <- vector("list", length(treedata.double))) for (k in (1:length(treedata.double))) {
    pred.double.all[[k]] <- log_transform_back( predict(tree.reg, newdata=treedata.double[[k]], method="a:") }

pred.double <- do.call("rbind", pred.double.all)

# distance for predictions on 2 sampled factors per cluster dist.double <- rep(NA, Ndouble) for (k in 1:Ndouble) {
    dist.double[k] <- log_distance(pred.double[k,], BDLsamples$time) }
}</pre>
```

Best factor combinations

Finding the best regression trees based on i) all single representatives from the clusters; and ii) single representatives only consisting of gene probes. The best is defined by minimal euclidian distance between experimental classes and predicted classes on the log scale. The best tree is not refitted with the respective factors in the tree, but the mean cluster tree uses the respective factor data for prediction.

```
# Best decision tree using all factors
# Find unique combinations with minimal distance
dist.rep.best <- min(dist.single)
# best combination of representatives for the clusters (remove duplicates)
rep.best <- unique(single.combinations[which(dist.single==dist.rep.best), c("c1", "c3", "c4", "c5")])
rep.best.idx <- rownames(rep.best)[1]
# predictions of best representative
pred.rep.best <- pred.single[as.numeric(rep.best.idx),]</pre>
print("Best single representatives for decision tree:")
```

```
## [1] "Best single representatives for decision tree:"
print(rep.best)
                          c4
                  сЗ
                                с5
             c1
## 16062 Cyp1a2 Fn1 S100a4 Il17a
## 16084 Cyp1a2 GLDH S100a4 Il17a
print(dist.rep.best)
## [1] 0.05554854
# Best decision tree using only gene factors, i.e. first reduce to the gene combinations
dist.single.genes <- dist.single[gene_only.single]</pre>
single.combinations.genes <- single.combinations[gene_only.single,]</pre>
dist.gene.best <- min(dist.single.genes)</pre>
# find best gene combination
gene.best <- unique(single.combinations.genes[which(dist.single.genes==dist.gene.best), c("c1", "c3", "</pre>
# predictions with best representative
gene.best.idx <- rownames(gene.best)</pre>
pred.gene.best <- pred.single[as.numeric(gene.best.idx), ]</pre>
pred.gene.best
##
                     2
                                3
                                                               6
                                                                         7
           1
                                                     5
                         0.00000
                                   0.00000
##
     0.00000
               0.00000
                                              0.00000
                                                         6.00000
                                                                   6.00000
##
                               10
                                                    12
                                                              13
                                                        13.98812 13.98812
##
     6.00000
               6.00000
                         6.00000 13.98812
                                              6.00000
##
          15
                               17
                                         18
                                                    19
                                                              20
##
    13.98812
              13.98812 13.98812 13.98812 13.98812
                                                       13.98812
                                                                  13.98812
##
          22
                    23
                               24
                                                    26
                                                              27
                                                                         28
                                         25
   33.57599
              13.98812
                       33.57599
                                  13.98812 33.57599
                                                        33.57599
                                                                  33.57599
##
##
          29
                    30
                               31
                                         32
                                                    33
                                                              34
##
    33.57599 33.57599
                        33.57599 336.00000 142.52453 142.52453 142.52453
          36
                    37
                               38
                                                    40
## 142.52453 336.00000 142.52453 336.00000 336.00000
print("Best single representatives based on genes for decision tree:")
## [1] "Best single representatives based on genes for decision tree:"
print(gene.best)
             c1 c3
                         c4
## 14808 Cyp1a2 Fn1 Col1a1 Il17a
print(dist.gene.best)
```

[1] 0.06672908

```
Nc <- ncol(combination)</pre>
 Nr <- nrow(combination)</pre>
 par(mfrow=c(Nr, Nc))
 for (kr in 1:Nr){
    for (kc in 1:Nc){
      cluster <- colnames(combination)[kc]</pre>
      name <- as.character(combination[kr, kc])</pre>
      plot_single(name)
    }
 }
 par(mfrow=c(1,1))
# plot to file
pdf(file.path(resultsPath, "decision_tree", "rep.best.representatives.pdf"),
    width=10, height=6, pointsize=12)
plot_tree_representatives(rep.best)
invisible(dev.off())
pdf(file.path(resultsPath, "decision_tree", "gene.best.representatives.pdf"),
    width=10, height=3, pointsize=12)
plot_tree_representatives(gene.best)
invisible(dev.off())
# plot to report
plot_tree_representatives(rep.best)
```

Plot time courses of the representatives/factors in provided tree combinations

plot tree representatives <- function(combination){</pre>

Prediction errors for factor combinations

plot tree representatives(gene.best)

Plot the distance distributions of single and double representatives and the best gene and representative trees.

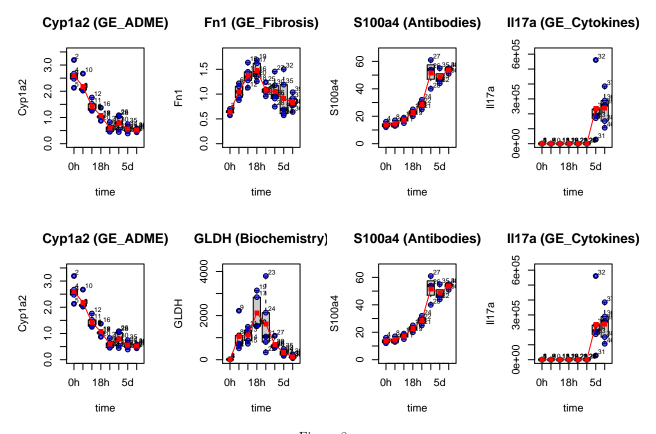


Figure 8:

Predictive performance

Evaluation of the predictions. Which experimental classes were predicted in which time classes of the regression tree for the mean cluster data (trainings data), single representative from each clusters and double representatives from each cluster.

```
# Predicted classes of the regression tree
node_levels <- levels(as.factor(pred.mean))
node_classes <- round(as.numeric(node_levels), digits=1)
node_classes</pre>
```

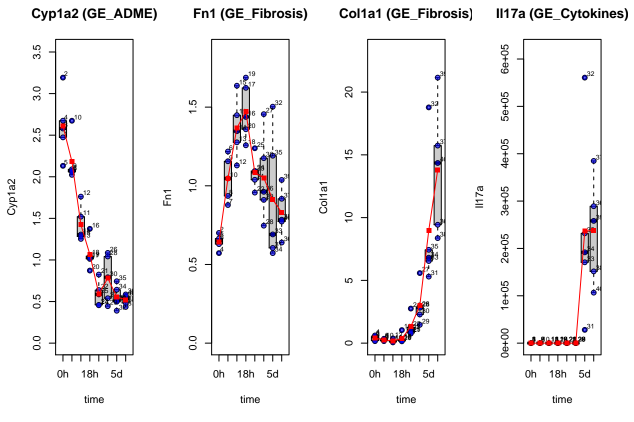


Figure 9:

Distance between predicted and experimentell time

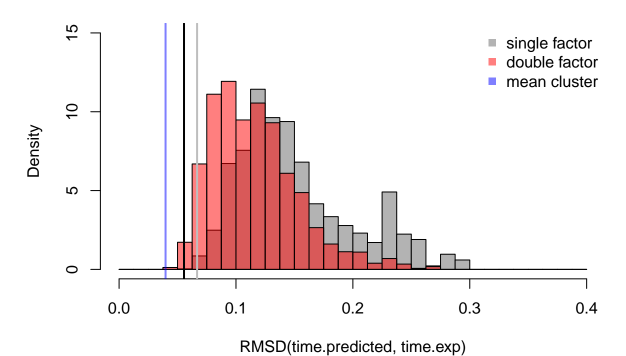


Figure 10:

```
# Plot of the predicted classes with the decision tree
plot_predicted_classes <- function(){</pre>
  # bar_colors <- brewer.pal(4, "Set3")</pre>
  bar_colors <- c(rgb(0.9,0.9,0.9),
                   rgb(0.7,0.7,0.7),
                   rgb(1,0,0, 0.5),
                   rgb(0,0,1, 0.5))
  par(mfrow=c(2,4))
  for (k in 1:Nt){
    # single factor predictions
    data <- as.vector(pred.single[, ((1:Nr)+Nr*(k-1))])</pre>
    tab.single <- table(factor(data, levels=node_levels))/length(data)</pre>
    # two factor predictions
    data <- as.vector(pred.double[, ((1:Nr)+Nr*(k-1))])</pre>
    tab.double <- table(factor(data, levels=node_levels))/length(data)</pre>
    # mean cluster predictions
    data <- as.vector(pred.mean[((1:Nr)+Nr*(k-1))])</pre>
    tab.mean <- table(factor(data, levels=node_levels))/length(data)</pre>
    # best single representative
    # data \leftarrow as.vector(pred.rep.best[((1:Nr)+Nr*(k-1))])
    # tab.rep.best <- table(factor(data, levels=node_levels))/length(data)</pre>
    # best single gene representative
    data <- as.vector(pred.gene.best[((1:Nr)+Nr*(k-1))])</pre>
    tab.gene.best <- table(factor(data, levels=node_levels))/length(data)</pre>
    # combined table
    tab <- rbind(tab.single, tab.double, tab.gene.best, tab.mean)</pre>
    colnames(tab) <- round(as.numeric(colnames(tab)), digits=1)</pre>
    # create the bar plot
    name <- sprintf("Time after BDL: %sh", levels(as.factor(BDLsamples$time))[k])</pre>
    barplot(tab, beside=TRUE,
            main=name,
            xlab="predicted time class [h]", ylab="fraction of predictions",
            ylim=c(0,1), col=bar_colors)
    if (k==1){
      legend("topright", legend=c("single factors", "double factors",
                                    "best single gene", "mean cluster"),
             col=bar_colors,
             bty="n", cex=1.0, pch=15)
    }
  }
  par(mfrow=c(1,1))
# barplot of the predicted classes
plot_predicted_classes()
```

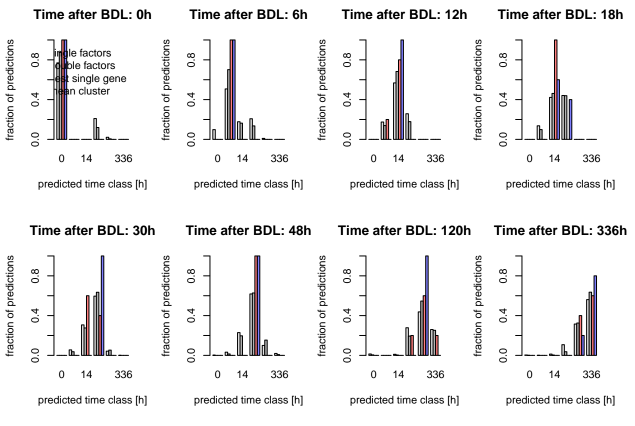


Figure 11:

Reverse bar plots showing the classes of the prediction tree

```
# make the bar plot reverse
library(plyr)
##
## Attaching package: 'plyr'
##
## The following object is masked from 'package:matrixStats':
##
##
       count
library(reshape2)
# which experiments are predicted in which class
plot_predicted_classes2 <- function(){</pre>
  bar_colors \leftarrow c(rgb(1,1,1),
                   rgb(0.8,0.8,0.8),
                   rgb(1,0,0,0.5),
                   rgb(0,0,1,0.5)
  # single
  df <- data.frame(exp=rep(BDLsamples$time, Nsingle),</pre>
                  pre=as.vector(t(round(pred.single, digits=1))) )
  tmp <- count(df, c("exp", "pre"))</pre>
  tab.single <- acast(tmp, exp~pre, value.var="freq", fill=0)</pre>
```

```
# double
  df <- data.frame(exp=rep(BDLsamples$time, Ndouble),</pre>
                  pre=as.vector(t(round(pred.double, digits=1))) )
  tmp <- count(df, c("exp", "pre"))</pre>
  tab.double <- acast(tmp, exp~pre, value.var="freq", fill=0)</pre>
  # best gene
  df <- data.frame(exp=rep(BDLsamples$time, 1),</pre>
                 pre=as.vector(t(round(pred.gene.best, digits=1))) )
  tmp <- count(df, c("exp", "pre"))</pre>
  tab.gene.best <- acast(tmp, exp~pre, value.var="freq", fill=0)</pre>
  # mean cluster
  df <- data.frame(exp=rep(BDLsamples$time, 1),</pre>
                  pre=as.vector(t(round(pred.mean, digits=1))) )
  tmp <- count(df, c("exp", "pre"))</pre>
  tab.mean <- acast(tmp, exp~pre, value.var="freq", fill=0)</pre>
  par(mfrow=c(2,3))
  for (k in 1:length(node classes)){
    # combined table (normalized within each class)
    tab <- rbind(tab.single[,k]/sum(tab.single[,k]),</pre>
                  tab.double[,k]/sum(tab.double[,k]),
                  tab.gene.best[,k]/sum(tab.gene.best[,k]),
                  tab.mean[,k]/sum(tab.mean[,k]))
    # create the bar plot
    name <- sprintf("Predicted: %sh", node_classes[k])</pre>
    barplot(tab, beside=TRUE,
            main=name,
            xlab="time after BDL [h]", ylab="fraction",
            ylim=c(0,1), col=bar_colors)
      legend("topright", legend=c("single factors", "double factors",
                                    "best gene", "mean cluster"),
             col=bar_colors,
             bty="n", cex=1.0, pch=15)
    }
 par(mfrow=c(1,1))
plot_predicted_classes2()
```

```
# barplot to file
pdf(file.path(resultsPath, "decision_tree", "predicted_classes2.pdf"),
    width=12, height=7.5, pointsize=14)
plot_predicted_classes2()
invisible(dev.off())
```

