# Shank3 Modulates Sleep and Expression of Circadian Transcription Factors differential expression

February 28, 2019

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# Description

This is the report of the analysis made for the paper Shank3 Modulates Sleep and Expression of Circadian Transcription Factors by Ashley M. Ingiosi, Taylor Wintler, Hannah Schoch, Kristan G. Singletary, Dario Righelli, Leandro G. Roser, Davide Risso, Marcos G. Frank and Lucia Peixoto.

Autism Spectrum Disorder (ASD) is the most prevalent neurodevelopmental disorder in the US that often co-presents with sleep problems. Sleep impairments in ASD predict the severity of ASD core diagnostic symptoms and have a considerable impact on the quality of life of caregivers. However, little is known about the underlying molecular mechanism(s) of sleep impairments in ASD. In this study we investigated the role

of Shank3, a high confidence ASD gene candidate, in the regulation of sleep. We show that Shank3 mutant mice have problems falling asleep despite accumulating sleep pressure. Using RNA-seq we show that sleep deprivation doubles the differences in gene expression between mutants and wild types and downregulates circadian transcription factors Per3, Dec2, and Rev-erb $\alpha$ . Shank3 mutants also have trouble regulating locomotor activity in the absence of light. Overall, our study shows that Shank3 is an important modulator of sleep and circadian activity. # Differential Expression Analysis

#### Importing data

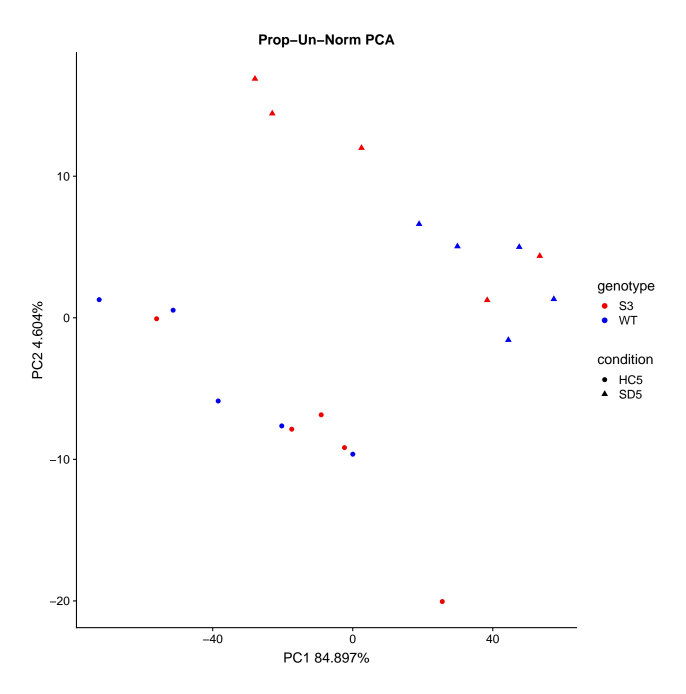
Importing data and filtering out those genes with cpm lesser than 1. We use the *filtered.data* method in *NOISeq* package.

#### Plot PCA of log unnormalized data

PCA Plot of filtered not-normalized data.

```
PlotPCAPlotlyFunction(counts.data.frame=log1p(filteredCountsProp),
    design.matrix=designMatrix,
    shapeColname="condition", colorColname="genotype", xPCA="PC1", yPCA="PC2",
    plotly.flag=FALSE, show.plot.flag=TRUE, prefix.plot="Prop-Un-Norm")
```

## [1] FALSE



## **Control Genes**

#### Negative control genes

Loading Negative Control Genes to normalize data

```
library(readxl)

sd.ctrls <- read_excel(path="./data/controls/Additional File 4 full list of BMC genomics SD&RS2.xlsx",
sd.ctrls <- sd.ctrls[order(sd.ctrls$adj.P.Val),]

sd.neg.ctrls <- sd.ctrls[sd.ctrls$adj.P.Val > 0.9, ]
```

#### positive control genes

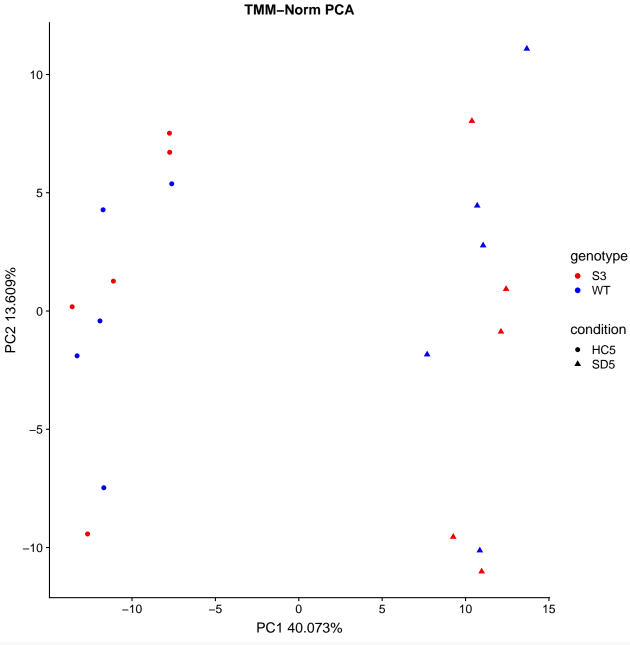
Loading Positive Control Genes to detect them during the differential expression step.

#### **Normalizations**

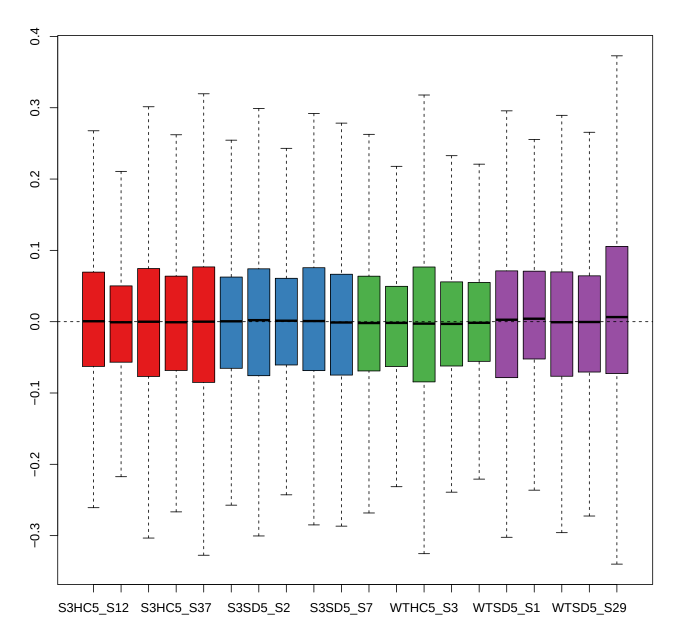
#### TMM Normalization

Normalizing data with TMM, as implemented in edgeR package, and plotting a PCA and an RLE plot of them.

## [1] FALSE



pal <- RColorBrewer::brewer.pal(9, "Set1")
plotRLE(as.matrix(normPropCountsUqua), outline=FALSE, col=pal[designMatrix\$gcondition])</pre>

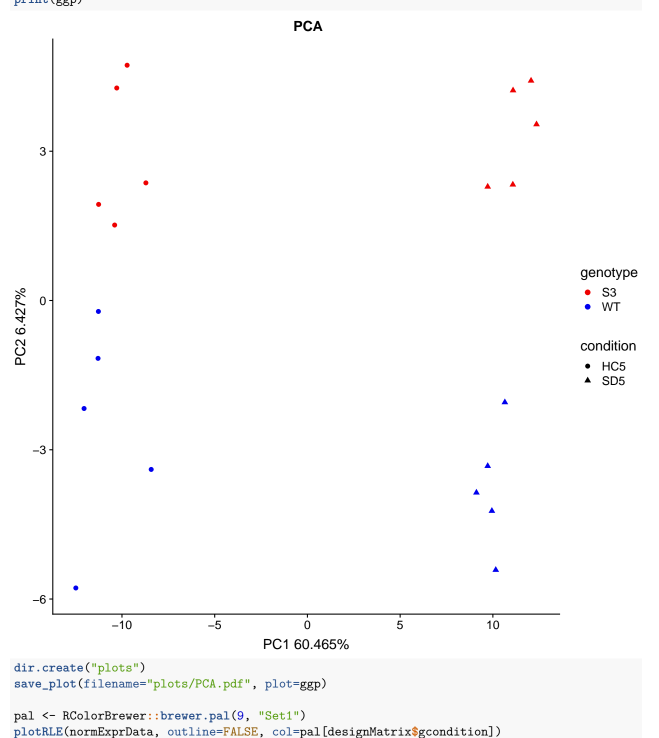


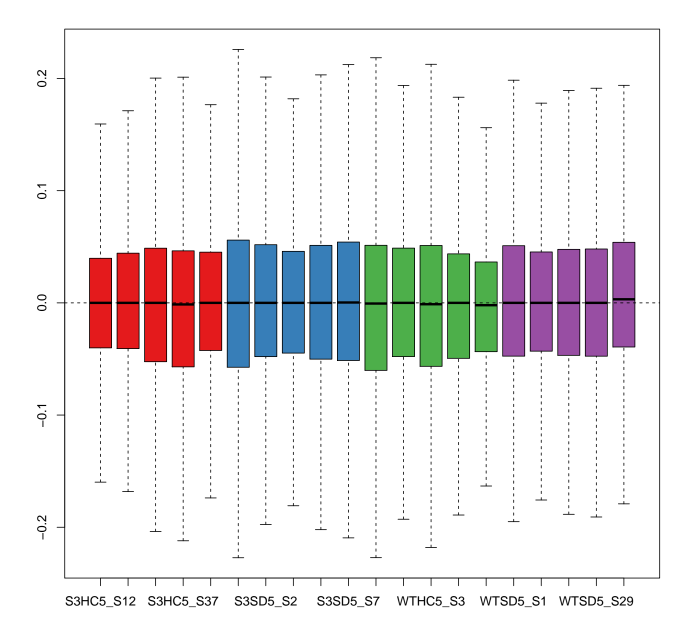
#### TMM + RUVs Normalization

Applying a RUVs method of RUVSeq package on normalized data, in order to adjust the counts for the unwanted variation. And of corse we plot a PCA and an RLE plot on these data.

```
plotly.flag=FALSE, show.plot.flag=FALSE, save.plot=FALSE,
prefix.plot=NULL)
```

## [1] FALSE
print(ggp)





## edgeR Differential Expression Analysis

Making differential expression analysis with edgeR package on four different contrasts.

Here is a brief legend:

- WTSD5: Wild Type Sleep Deprivation 5 days.
- KOHC5: Knock Out Home Cage Control 5 days.
- KOSD5: Knock Out Sleep Deprivation 5 days.

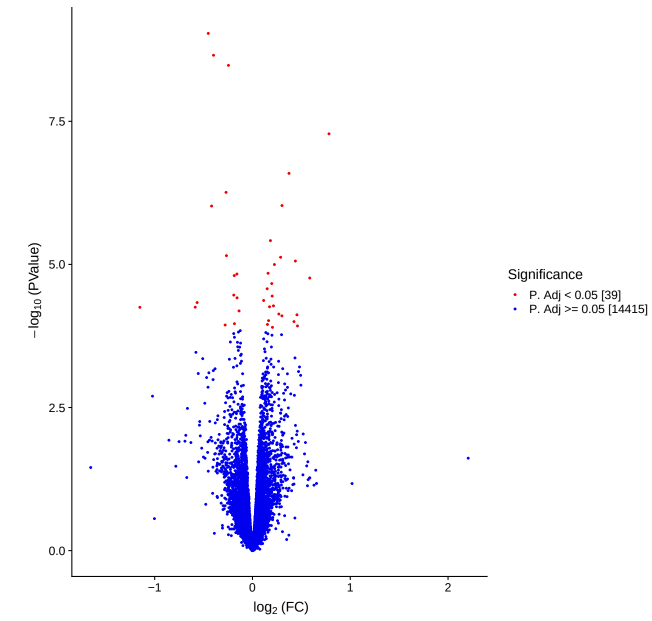
```
padj.thr <- 0.05
venn.padgj.thr <- 0.1
desMat <- cbind(designMatrix, ruvedSExprData$W)
colnames(desMat) <- c(colnames(designMatrix), colnames(ruvedSExprData$W))
cc <- c("S3HC5 - WTHC5", "S3SD5 - WTSD5", "WTSD5 - WTHC5")</pre>
```

#### Shank3 Home Cage control VS Wild Type Home Cage Controls

#### volcano plot

A volcano plot of differential expressed genes.





```
de <- sum(res.o$FDR < padj.thr)
nde <- sum(res.o$FDR >= padj.thr)
detable <- cbind(de,nde)
rownames(detable) <- names(rescList1)[1]
ddetable <- detable

tot.ctrls <- dim(sd.pos.ctrls)[1]
idx.pc <- which(tolower(res.o$gene) %in% tolower(sd.pos.ctrls[,1]))
tot.pc.de <- sum(res.o$FDR[idx.pc] < padj.thr)
tot.pc.nde <- length(idx.pc) - tot.pc.de

wt <- res.o[which(res.o$FDR < padj.thr),]
wt.sign.genes.entrez <- rownames(res.o)[which(res.o$FDR < venn.padgj.thr)]</pre>
```

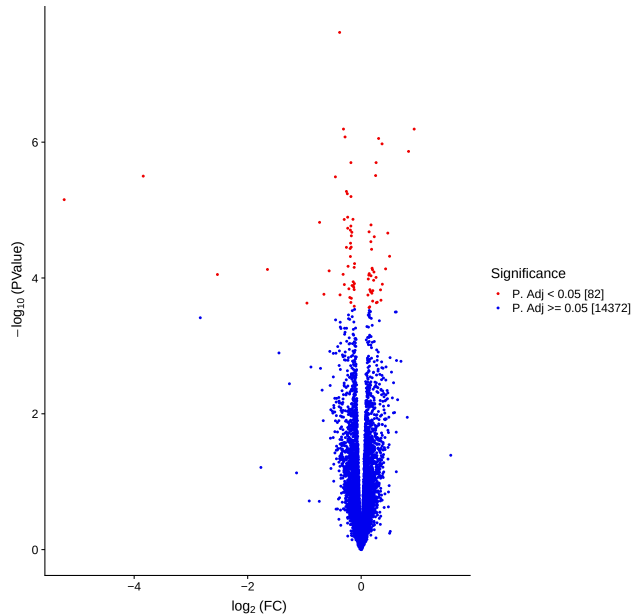
```
kowthc5 <- res.o[which(res.o$FDR < padj.thr),]
kowthc5.sign.genes.entrez <- rownames(res.o)[which(res.o$FDR < venn.padgj.thr)]</pre>
```

#### Shank3 Sleed Deprivation VS Wild Type Sleep Deprivation

#### volcano plot

A volcano plot of differential expressed genes.

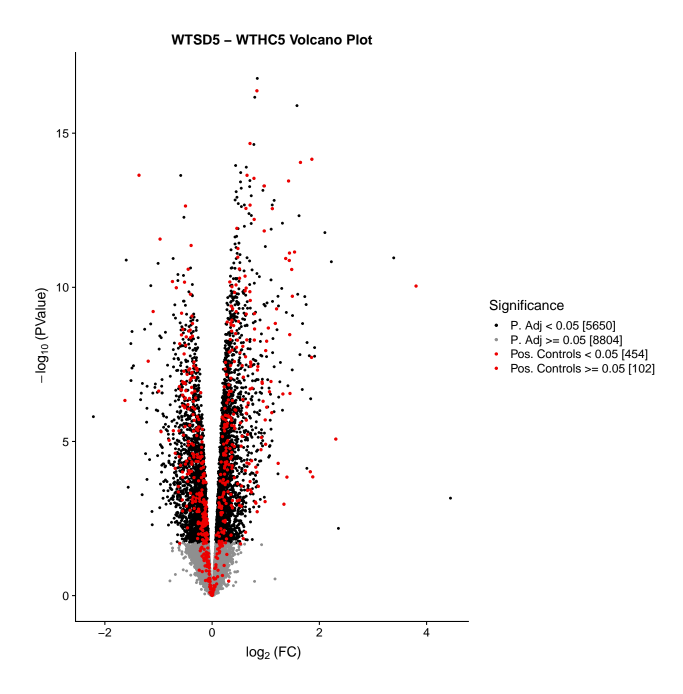




#### DE TABLE + Positive Controls table

We present a summarization of the results. The first table is a summarization on how many genes are Differentially Expressed. The second table explains on the first column how many positive controls we have, on the second column how many positive controls have been identified over the differentially expressed genes, and, finally, on the third column how many positive controls have beed identified on the NOT differentially expressed genes.

#### WTSD5 - WTHC5 positive controls

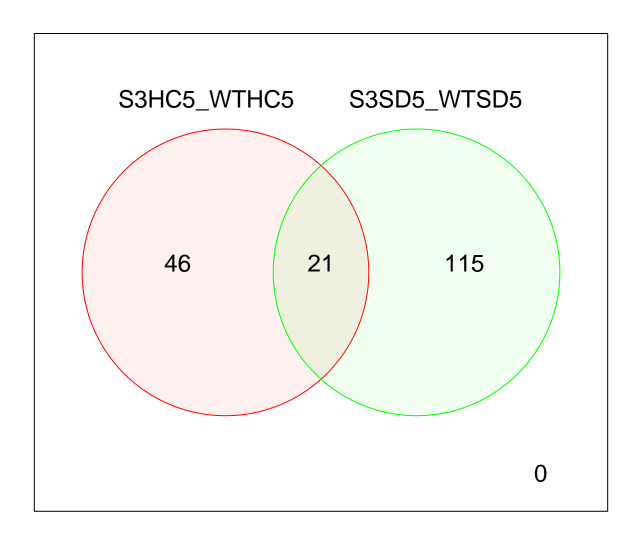


## Venn Diagram

#### KOHC5-WTHC5 vs KOSD5-WTSD5

We take the results of the two contrasts. Knock Out Sleed Deprivation VS Wild Type Sleep Deprivation and Knock Out Home Cage control VS Wild Type Home Cage Controls . And plot the results in a Venn Diagram

# S3HC5\_WTHC5 venn S3SD5\_WTSD5



## Heatmaps

Setting up the data structures for the heatmps.

```
source("./R/heatmapFunctions.R")
de.genes.entr <- union(rownames(venn$int), rownames(venn$XnoY))
de.genes.entr <- union(de.genes.entr, rownames(venn$YnoX))</pre>
```

```
gene.map <- convertGenesViaMouseDb(gene.list=de.genes.entr,</pre>
                             fromType="ENTREZID")
de.genes.symb <- attachGeneColumnToDf(as.data.frame(de.genes.entr,</pre>
                                                      row.names=de.genes.entr),
                                     genesMap=gene.map,
                                     rowNamesIdentifier="ENTREZID",
                                     mapFromIdentifier="ENTREZID",
                                     mapToIdentifier="SYMBOL")
# de.qenes.symb[which(is.na(de.qenes.symb$qene)),]
de.genes.symb$gene[which(de.genes.symb$de.genes.entr=="100039826")] <- "Gm2444" ## not annotated in no
de.genes.symb$gene[which(de.genes.symb$de.genes.entr=="210541")] <- "Entrez:210541" ## not annotated in
de.genes.counts <- normExprData[match(de.genes.symb$de.genes.entr, rownames(normExprData)),]</pre>
rownames(de.genes.counts) <- de.genes.symb$gene</pre>
de.gene.means <- computeGeneMeansOverGroups(counts=de.genes.counts,</pre>
                             design=designMatrix, groupColumn="gcondition")
library(gplots)
library(clusterExperiment)
color.palette = clusterExperiment::seqPal3#c("black", "yellow")
pal <- colorRampPalette(color.palette)(n = 1000)</pre>
library(pheatmap)
filter2 <- rowMeans(de.gene.means)>0
filter <- apply(de.gene.means, 1, function(x) log(x[4]/x[3]) * log(x[2]/x[1]) < 0)
filter[is.na(filter)] <- FALSE</pre>
```

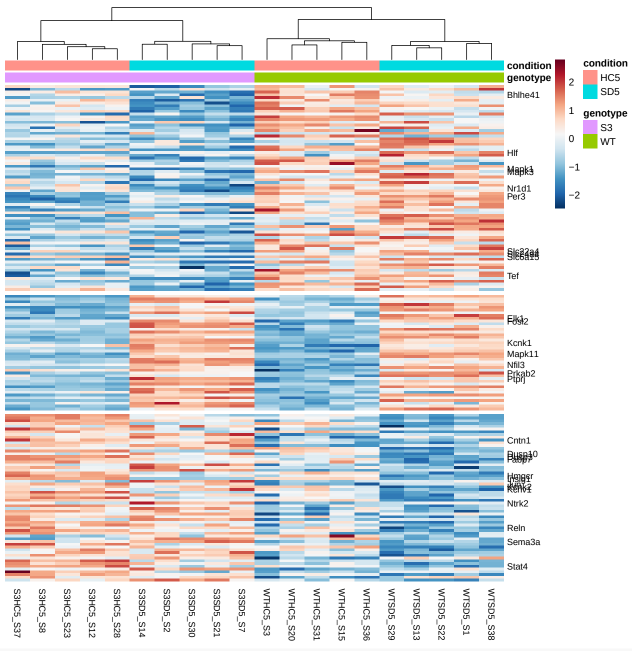
#### Heatmap gene by bene

```
0.15
                                                                                                      HC5
                                                                             genotype
                                                                                                      SD5
                                                                                              0.1
                                                                                                    genotype
                                                                                              0.05
                                                                                                      S3
                                                                                                      WT
                                                                                               -0.05
                                                                             Cluster: 2 Size: 42
                                                                                              -0.1
                                                                             Cluster: 1 Size: 75
                                                                             Cluster: 3 Size: 61
                      S3HC5_S8
                                 WTHC5_S3
                                    WTHC5_S3
                                       WTHC5_S36
                                                        WTSD5_S22
                                                            S3SD5_S2
                          WTHC5_S20
                                           WTSD5_S1
                                              WTSD5_S38
                                                  WTSD5_S29
                                                     WTSD5_S13
                                                               S3SD5_S7
                                                                   S3SD5_S1
                             WTHC5_S16
clusterized.genes <- as.data.frame(ph1$kmeans$cluster)</pre>
gene.map <- convertGenesViaMouseDb(gene.list=rownames(clusterized.genes), fromType="SYMBOL")</pre>
converted.clusterized.gens <- attachGeneColumnToDf(mainDf=clusterized.genes, genesMap=gene.map,</pre>
                        rowNamesIdentifier="SYMBOL", mapFromIdentifier="SYMBOL", mapToIdentifier="ENTREZID"
converted.clusterized.gens$gene[which(rownames(converted.clusterized.gens)=="Gm2444")] <- "100039826"
converted.clusterized.gens$gene[which(rownames(converted.clusterized.gens)=="Entrez:210541")] <- "2105"
converted.clusterized.gens <- converted.clusterized.gens[order(converted.clusterized.gens$`ph1$kmeans$c</pre>
save_pheatmap_pdf(filename="plots/heatmap_kmeans_k3.pdf", plot=ph1, width=20, height=20)
## pdf
##
      2
```

condition

condition

```
WriteDataFrameAsTsv(data.frame.to.save=converted.clusterized.gens, file.name.path="plots/clustered_gene"
ord.de.genes.counts <- de.heatmap[match(rownames(converted.clusterized.gens), rownames(de.heatmap)),]
idx <- which(!(rownames(ord.de.genes.counts) %in% gene_names))</pre>
rownames(ord.de.genes.counts)[idx] <- ""</pre>
gaps.row <- c()</pre>
for(i in c(1:3))
    li <- length(which(converted.clusterized.gens$`ph1$kmeans$cluster`==i))</pre>
    1 <- ifelse(i!=1, gaps.row[i-1]+li, li)</pre>
    gaps.row <- c(gaps.row, 1)</pre>
}
heatmap_data_scaled <- t(scale(t(log(ord.de.genes.counts+1)), center = TRUE, scale = TRUE))
library(dendextend)
column_dend <- as.dendrogram(hclust(dist(t(heatmap_data_scaled))))</pre>
ord <- labels(column_dend)</pre>
ord[11:15] <- labels(column_dend)[16:20]
ord[16:20] <- labels(column_dend)[11:15]
column_dend <- rotate(column_dend, ord)</pre>
ph1 <- pheatmap(heatmap_data_scaled, cluster_cols=as.hclust(column_dend), scale="none",</pre>
             color=pal, border_color=NA, fontsize_row=9, fontsize_col=9, cluster_rows=FALSE,
             annotation_col=ann.col, gaps_row=gaps.row)
```



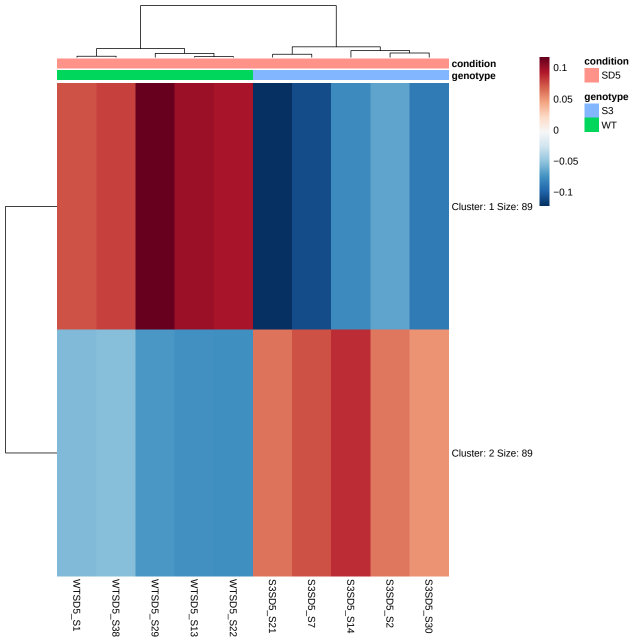
save\_pheatmap\_pdf(filename="plots/heatmap\_gg\_k3.pdf", plot=ph1, width=20, height=20)

```
## pdf
## 2
```

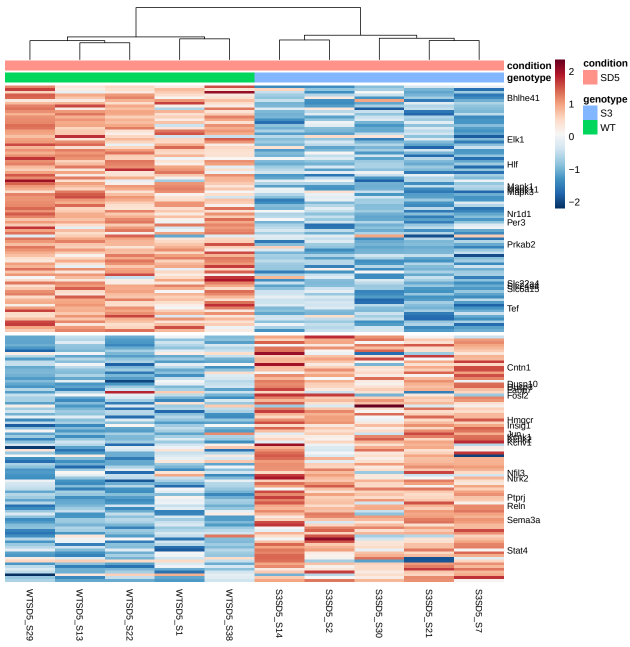
#### other heatmaps

condition condition HC5 genotype SD5 0.5 Bhlhe41 genotype S3 WT HIf Mapk3 -1.5 Nr1d1 Per3 -2 \$168394 Tef Fbs12 Kcnk1 Mapk11 Nfil3 Brkab2 Ptprj Cntn1 Pusp<sub>2</sub>0 Higher Kenka Ntrk2 Reln Sema3a Stat4 WTSD5\_S38 S3HC5\_S37 S3HC5\_S8 S3HC5\_S12 S3HC5\_S28 S3SD5\_S14 S3SD5\_S30 S3SD5\_S21 S3SD5\_S7 WTHC5\_S3 WTHC5\_S31 WTHC5\_S15 WTHC5\_S36 WTSD5\_S29 WTSD5\_S13 WTSD5\_S22 WTSD5\_S1 S3HC5\_S23 S3SD5\_S2 WTHC5\_S20 save\_pheatmap\_pdf(filename="plots/heatmap\_gg\_k3\_no\_scale.pdf", plot=ph1, width=20, height=20) ## pdf ## ## Only SD samples heatmap\_data <- t(scale(t(log(de.heatmap[, grep("^SD", desMat[,2])]+1)), center = TRUE, scale = FALSE))</pre> ph1 <- pheatmap(heatmap\_data, cluster\_cols=TRUE, scale="none",</pre> color=pal, border\_color=NA, fontsize\_row=10, kmeans\_k=2, annotation\_col=ann.col)

breaks = c(min(heatmap\_data), seq(quantile(as.vector(heatmap\_data), .01), quantile(as.vector



idx <- which(!(rownames(ord.de.genes.counts) %in% gene\_names))</pre>



save\_pheatmap\_pdf(filename="plots/heatmap\_gg\_sd\_only.pdf", plot=ph1, width=20, height=20)

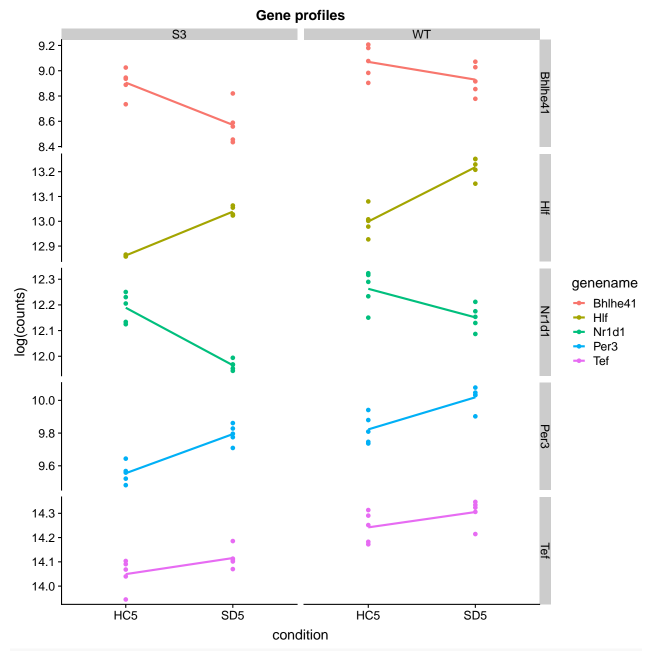
```
## pdf
## 2
```

## Group gene profiles

#### Group gene profiles by genotype

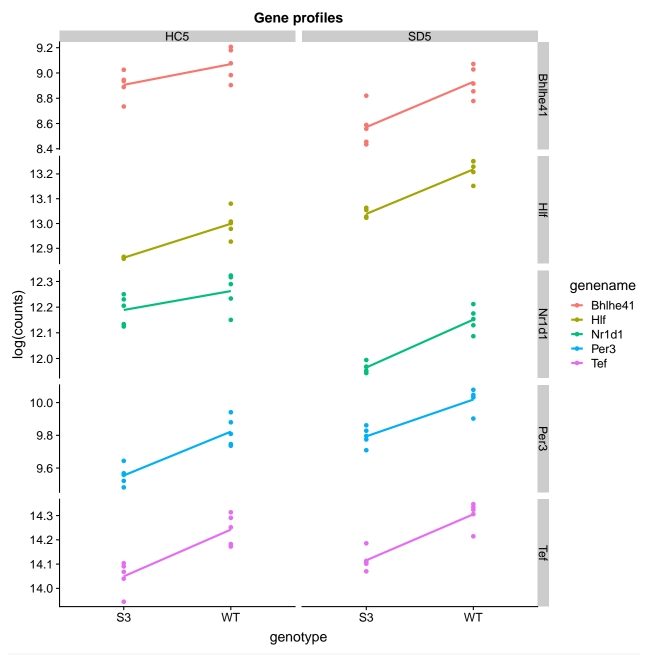
```
g <- geneGroupProfileRows(normalized.counts=normExprData, design.matrix=designMatrix, gene.names=c("Nr1d1", "Hlf", "Per3", "Bhlhe41", "Tef"), res.o=de.genes.symb, show.plot=TRUE, plotly.flag=FALSE, log.flag=TRUE)
```





save\_plot(filename=paste0("plots/", "Nr1d1\_Hlf\_Per3\_Bhlhe41\_Tef", "\_log\_gene\_profile\_genotype.pdf"), pl
base\_height=15, base\_width=15)

#### Group gene profiles by condition



save\_plot(filename=paste0("plots/", "Nr1d1\_Hlf\_Per3\_Bhlhe41\_Tef", "\_log\_gene\_profile\_condition.pdf"), p
 base\_height=15, base\_width=15)

# Circadian Analysis LD-LD

# Analysis for activity

```
wt <- read_xlsx("data/LD_LD_Activity_analysis_4_R.xlsx", sheet = 1)
mut <- read_xlsx("data/LD_LD_Activity_analysis_4_R.xlsx", sheet = 2)
wt <- wt %>%
```

```
bind_cols(WT.M=rep("WT", nrow(wt)), time = decimal_date(ymd(wt$`Total_revolutions/day`)), .) %>%
  gather(mice, activity, -c(1:3)) %>%
  mutate(time = time-min(time)) %>%
  dplyr::select(-`Total_revolutions/day`)
mut <- mut %>%
  bind_cols(WT.M=rep("M", nrow(mut)), time = decimal_date(ymd(mut$`Total_revolutions/day`)), .) %>%
  gather(mice, activity, -c(1:3)) %>%
  mutate(time = time-min(time)) %>%
  dplyr::select(-`Total_revolutions/day`)
data <- wt %>% bind_rows(mut)
#data <- data %>% filter(week>=3)
data$mice <- factor(data$mice, levels= unique(data$mice))</pre>
data$time_scaled <- scale(data$time, scale=FALSE)</pre>
#data$period <- factor(data$period, levels= unique(data$period))
data$WT.M <-factor(data$WT.M, levels=c("WT", "M"))</pre>
mod3 <- lme(activity ~ time_scaled * WT.M, random=~1 | mice, data = data)</pre>
cat("Estimates, errors and the significance")
## Estimates, errors and the significance
summary(mod3)
## Linear mixed-effects model fit by REML
  Data: data
          ATC
##
                   BIC
                          logLik
##
     11661.69 11687.61 -5824.844
##
## Random effects:
## Formula: ~1 | mice
##
           (Intercept) Residual
             14529.36 7940.975
## StdDev:
##
## Fixed effects: activity ~ time_scaled * WT.M
                         Value Std.Error DF
                                              t-value p-value
## (Intercept)
                      40118.57 5158.779 542 7.776758 0.0000
## time scaled
                       2089.01 1016.983 542 2.054126 0.0404
## WT.MM
                     -19109.48 7295.615 14 -2.619311 0.0202
## time_scaled:WT.MM -13336.51 1438.232 542 -9.272853 0.0000
## Correlation:
##
                     (Intr) tm_scl WT.MM
                      0.000
## time scaled
## WT.MM
                     -0.707 0.000
## time_scaled:WT.MM 0.000 -0.707 0.000
##
## Standardized Within-Group Residuals:
                                   Med
           Min
                        Q1
                                                 QЗ
                                                            Max
## -3.44738653 -0.64119095 0.03137236 0.52237000 3.77912348
##
```

```
## Number of Observations: 560
## Number of Groups: 16
cat("Bootstrap confidence intervals for the estimates")
## Bootstrap confidence intervals for the estimates
mod_lmer3 <- lmer(activity ~ time_scaled * WT.M + (1 mice), data = data)
suppressMessages(confint.merMod(mod_lmer3, method = "boot", nsim = 999))
                                   97.5 %
##
                         2.5 %
## .sig01
                      9338.623 19997.721
## .sigma
                      7455.925
                                8419.105
## (Intercept)
                     30323.250 49785.615
## time scaled
                       135.387
                                4004.274
## WT.MM
                    -34122.484 -4141.786
## time_scaled:WT.MM -16139.161 -10536.146
cat("ANOVA table")
## ANOVA table
anova.lme(mod3, type = "marginal", adjustSigma = F)
##
                   numDF denDF F-value p-value
## (Intercept)
                       1 542 60.47796 <.0001
                       1 542 4.21943 0.0404
## time scaled
## WT.M
                       1 14 6.86079 0.0202
## time scaled:WT.M
                      1 542 85.98580 <.0001
```

#### Analysis for alpha

```
wt <- read_xlsx("data/LD_LD_Alpha_Activity_analysis_4_R.xlsx", sheet = 1, na = "NA")
mut <- read_xlsx("data/LD_LD_Alpha_Activity_analysis_4_R.xlsx", sheet = 2, na = "NA")</pre>
wt <- wt %>%
  bind_cols(WT.M = rep("WT", nrow(wt)), time = decimal_date(ymd(wt$`Total_revolutions/day`)), .)%>%
  gather(mice, alpha, -c(1:3)) %>%
  mutate(time = time-min(time)) %>%
  dplyr::select(-`Total_revolutions/day`)
mut <- mut %>%
  bind_cols(WT.M=rep("M", nrow(mut)), time = decimal_date(ymd(mut$`Total_revolutions/day`)), .)%%
  gather(mice, alpha, -c(1:3)) %>%
  mutate(time = time-min(time)) %>%
  dplyr::select(-`Total_revolutions/day`)
alpha data <- wt %>% bind rows(mut)
#alpha_data <- alpha_data %>% filter(week>=3)
alpha_data<- na.omit(alpha_data)</pre>
alpha_data$mice <- factor(alpha_data$mice, levels= unique(alpha_data$mice))
alpha_data$time_scaled <- scale(alpha_data$time, scale=FALSE)</pre>
#alpha_data$period <- factor(alpha_data$period, levels= unique(alpha_data$period))
alpha_data$WT.M <- factor(alpha_data$WT.M, levels=c("WT", "M"))</pre>
```

```
alpha_data$alpha <- as.numeric(alpha_data$alpha)</pre>
mod4 <- lme(alpha ~ time_scaled * WT.M, random=~1 mice, data = alpha_data, na.action = na.omit)
cat("Estimates, errors and the significance")
## Estimates, errors and the significance
summary(mod4)
## Linear mixed-effects model fit by REML
   Data: alpha_data
##
         AIC
                 BIC
                        logLik
##
    3188.158 3214.58 -1588.079
##
## Random effects:
## Formula: ~1 | mice
##
           (Intercept) Residual
## StdDev:
             1.301648 3.255454
##
## Fixed effects: alpha ~ time_scaled * WT.M
##
                         Value Std.Error DF
                                             t-value p-value
                     ## (Intercept)
## time scaled
                    -18.117603 6.214771 590 -2.915249 0.0037
## WT.MM
                     -1.255362 0.702350 14 -1.787374 0.0955
## time scaled:WT.MM 6.997015 8.789013 590 0.796109 0.4263
## Correlation:
                    (Intr) tm_scl WT.MM
##
## time_scaled
                     0.000
## WT.MM
                    -0.707 0.000
## time_scaled:WT.MM 0.000 -0.707 0.000
## Standardized Within-Group Residuals:
          Min
                       Q1
                                  Med
                                               Q3
                                                          Max
## -2.87208043 -0.48234294 -0.01803405 0.36708937 4.80926449
##
## Number of Observations: 608
## Number of Groups: 16
cat("Bootstrap confidence intervals for the estimates")
## Bootstrap confidence intervals for the estimates
mod4_lmer <- lmer(alpha ~ time_scaled * WT.M + (1|mice), data = alpha_data)</pre>
suppressMessages(confint.merMod(mod4_lmer, method = "boot", nsim = 999))
##
                          2.5 %
                                    97.5 %
## .sig01
                      0.6842351 1.8257756
                      3.0622418 3.4435839
## .sigma
## (Intercept)
                      9.0826202 11.0532511
## time_scaled
                    -30.3568760 -5.9398413
## WT.MM
                     -2.6549706 0.1080982
## time_scaled:WT.MM -10.2458584 25.1892030
cat("ANOVA table")
```

## ANOVA table

```
anova.lme(mod4, type = "marginal", adjustSigma = F)
##
                    numDF denDF F-value p-value
                            590 412.7941 < .0001
## (Intercept)
## time_scaled
                        1
                            590
                                  8.4987 0.0037
## WT.M
                        1
                            14
                                  3.1947 0.0955
## time_scaled:WT.M
                            590
                                  0.6338 0.4263
                        1
Analysis for period
wt <- read xlsx("data/LD LD Period analysis 4 R.xlsx", sheet = 1) %>% gather(mice, value, -1)
wt <- data.frame(WT.M=rep("WT", nrow(wt))) %>% bind_cols(wt)
mut <- read_xlsx("data/LD_LD_Period_analysis_4_R.xlsx", sheet = 2) %>% gather(mice, value, -1)
mut <- data.frame(WT.M=rep("M", nrow(mut))) %>% bind_cols(mut)
period_data <- wt %>% bind_rows(mut)
period_data$value <- as.numeric(period_data$value)</pre>
mod5 <- lme(value ~ week * WT.M, random = ~1 | mice, data = period_data)
cat("Estimates, errors and the significance")
## Estimates, errors and the significance
summary(mod5)
## Linear mixed-effects model fit by REML
   Data: period_data
##
         AIC
                  BIC
                          logLik
##
     373.0096 399.9916 -174.5048
##
## Random effects:
## Formula: ~1 | mice
           (Intercept) Residual
##
## StdDev:
           0.3722568 2.496563
## Fixed effects: value ~ week * WT.M
##
                           Value Std.Error DF t-value p-value
## (Intercept)
                        21.26375 0.8924266 56 23.826890 0.0000
                         2.59000 1.2482815 56 2.074853
## weekLD_Week_2
                                                         0.0426
## weekLD_Week_3
                         2.98500 1.2482815 56
                                               2.391288
                                                         0.0202
## weekLD_Week_4
                         2.75500 1.2482815 56 2.207034
                                                         0.0314
## weekLD_Week_5
                         2.81000 1.2482815 56 2.251095
                                                         0.0283
## WT.MWT
                         2.75375 1.2620818 14 2.181911 0.0467
## weekLD_Week_2:WT.MWT -2.62000 1.7653366 56 -1.484136 0.1434
## weekLD_Week_3:WT.MWT -3.02375 1.7653366 56 -1.712846 0.0923
## weekLD_Week_4:WT.MWT -2.76125 1.7653366 56 -1.564149 0.1234
## weekLD_Week_5:WT.MWT -2.81625 1.7653366 56 -1.595305 0.1163
## Correlation:
##
                        (Intr) wkLD W 2 wkLD W 3 wkLD W 4 wkLD W 5 WT.MWT
                        -0.699
## weekLD_Week_2
## weekLD_Week_3
                        -0.699 0.500
## weekLD_Week_4
```

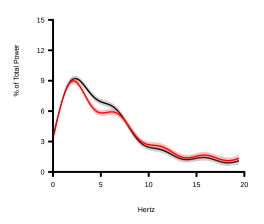
0.500

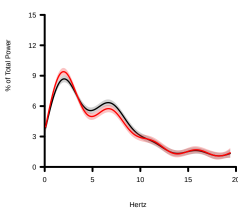
-0.699 0.500

```
## weekLD_Week_5
                        -0.699 0.500
                                         0.500
                                                  0.500
## WT.MWT
                        -0.707 0.495
                                         0.495
                                                  0.495
                                                           0.495
## weekLD Week 2:WT.MWT 0.495 -0.707
                                        -0.354
                                                 -0.354
                                                          -0.354
                                                                    -0.699
## weekLD_Week_3:WT.MWT 0.495 -0.354
                                        -0.707
                                                 -0.354
                                                           -0.354
                                                                    -0.699
## weekLD_Week_4:WT.MWT 0.495 -0.354
                                        -0.354
                                                 -0.707
                                                          -0.354
                                                                    -0.699
## weekLD Week 5:WT.MWT 0.495 -0.354
                                                          -0.707
                                                                    -0.699
                                        -0.354
                                                 -0.354
                        wLD_W_2: wLD_W_3: wLD_W_4:
## weekLD Week 2
## weekLD_Week_3
## weekLD_Week_4
## weekLD_Week_5
## WT.MWT
## weekLD_Week_2:WT.MWT
## weekLD_Week_3:WT.MWT
                         0.500
## weekLD_Week_4:WT.MWT
                         0.500
                                  0.500
## weekLD_Week_5:WT.MWT
                         0.500
                                  0.500
                                           0.500
##
## Standardized Within-Group Residuals:
           Min
                          Q1
                                      Med
                                                    03
## -7.277846353 -0.022845832 -0.004456104 0.031513974 2.241303977
##
## Number of Observations: 80
## Number of Groups: 16
cat("Bootstrap confidence intervals for the estimates")
## Bootstrap confidence intervals for the estimates
mod5_lmer <- lmer(value ~ week * WT.M + (1|mice), data = period_data)</pre>
suppressMessages(confint.merMod(mod5 lmer, method = "boot", nsim = 999))
##
                              2.5 %
                                        97.5 %
## .sig01
                         0.00000000 1.3513179
## .sigma
                         2.00418701 2.8931814
## (Intercept)
                        19.32598071 23.1408941
## weekLD_Week_2
                         0.15814999 5.0850499
## weekLD_Week_3
                         0.26043155 5.4351006
## weekLD_Week_4
                         0.08875410 5.1606052
## weekLD_Week_5
                         0.30404482 5.3296867
## WT.MWT
                         0.02750816
                                    5.2315235
## weekLD_Week_2:WT.MWT -6.14831649 0.6980356
## weekLD Week 3:WT.MWT -6.37762320 0.8897963
## weekLD_Week_4:WT.MWT -6.25747849 0.9877018
## weekLD_Week_5:WT.MWT -6.23504956 0.9621325
cat("ANOVA table")
## ANOVA table
anova.lme(mod5, type = "marginal", adjustSigma = F)
               numDF denDF F-value p-value
## (Intercept)
                        56 567.7207 < .0001
                   1
## week
                   4
                        56
                             2.0166 0.1045
## WT.M
                        14
                   1
                             4.7607 0.0467
## week:WT.M
                        56
                             1.0236 0.4031
```

## Analysis for Spectral Data

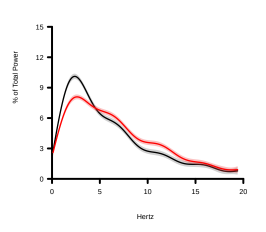
```
# GAM plots
library(mgcv)
data<-read xlsx("data/BL spectral.xlsx")</pre>
data <- data %>% gather(Hertz, value, -c(1:3))
data <- data %>%
 mutate(GT=replace(GT,GT == 1, "WT")) %>%
 mutate(GT=replace(GT,GT == 2, "MT")) %>%
 mutate(LD=replace(LD,LD == 1, "LIGHT")) %>%
 mutate(LD=replace(LD,LD == 2, "DARK")) %>%
 mutate(hz= as.numeric(Hertz)) %>%
  mutate(STATE = factor(STATE, levels = unique(STATE))) %>%
  mutate(GT = factor(GT, levels = unique(GT))) %>%
  mutate(LD = factor(LD, levels = unique(LD))) %>%
  mutate(value = replace(value, value == -99, NA))
temp<-data %>% filter(STATE == "WAKEFULNESS" & LD == "LIGHT")
index <-paste(data$STATE, data$LD, sep = "")</pre>
index_lev <- unique(index)</pre>
layout(matrix(seq_len(6), nrow = 3, ncol = 2, byrow = TRUE))
shadow_col \leftarrow c(rgb(109, 109, 109, max = 255, alpha = 80),
                 rgb(244, 66, 66, max = 255, alpha = 80))
for(this index in index lev) {
  state <- unique(data[index == this_index, ][, 1])</pre>
  state <- as.character(unlist(state))</pre>
  light <- unique(data[index == this_index, ][, 3])</pre>
  light <- as.character(unlist(light))</pre>
  temp2 <- data[index == this_index, ]</pre>
  plot(x = temp2\$hz, y = temp2\$value, type = "n",
       ylab = "% of Total Power", ylim = c(0,20),
       xlab = 'Hertz', lwd = 3, cex = 1.2,
       main = paste0(state, "-", light), axes = FALSE)
  axis(1, at = seq(0, 20, by = 5), las = 1, pos = 0, lwd = 3)
  axis(2, at = seq(0, 15, by = 3), las = 2, pos = 0, lwd = 3)
  mod <- list(wt = gam(value~s(hz), data = temp2[temp2$GT == "WT",]),</pre>
              mt = gam(value~s(hz), data = temp2[temp2$GT == "MT",]))
  for(i in seq along(mod)) {
    ss \leftarrow seq(min(temp2\$hz) + 0.1, max(temp2\$hz) - 0.1, 0.1)
    pred <- predict(mod[[i]], data.frame(hz = ss), se = TRUE)</pre>
    fit <- pred$fit</pre>
    se <- pred$se.fit
    lower <- fit - 1.96 * se
    upper <- fit + 1.96 * se
    to_plot <- data.frame(hz = ss, fit, lower, upper)</pre>
```

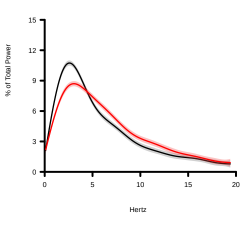




NREM-LIGHT

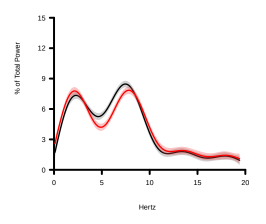
NREM-DARK

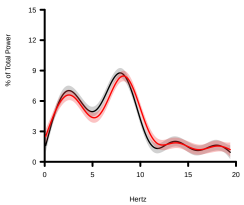




REM-LIGHT

REM-DARK





## Session Info

```
sessionInfo()
```

```
## R version 3.4.3 (2017-11-30)
## Platform: x86_64-pc-linux-gnu (64-bit)
## Running under: Ubuntu 16.04.5 LTS
## Matrix products: default
## BLAS: /usr/local/lib/R/lib/libRblas.so
## LAPACK: /usr/local/lib/R/lib/libRlapack.so
##
## locale:
## [1] LC CTYPE=en US.UTF-8
                                   LC NUMERIC=C
  [3] LC_TIME=en_GB.UTF-8
                                   LC_COLLATE=en_US.UTF-8
   [5] LC_MONETARY=en_GB.UTF-8
                                   LC_MESSAGES=en_US.UTF-8
##
  [7] LC_PAPER=en_GB.UTF-8
                                   LC_NAME=C
##
## [9] LC ADDRESS=C
                                   LC TELEPHONE=C
## [11] LC_MEASUREMENT=en_GB.UTF-8 LC_IDENTIFICATION=C
## attached base packages:
## [1] stats4
                 parallel stats
                                     graphics grDevices utils
                                                                    datasets
## [8] methods
                 base
## other attached packages:
## [1] bindrcpp_0.2.2
                                   mgcv_1.8-24
   [3] dendextend_1.8.0
                                   pheatmap_1.0.10
                                   gplots_3.0.1
## [5] clusterExperiment_1.4.0
## [7] plyr 1.8.4
                                   org.Mm.eg.db 3.5.0
## [9] AnnotationDbi_1.40.0
                                   readxl_1.1.0
## [11] lme4 1.1-18-1
                                   Matrix 1.2-14
## [13] nlme_3.1-137
                                   tidyr_0.8.1
                                   lubridate_1.7.4
## [15] dplyr_0.7.6
## [17] cowplot_0.9.3
                                   RUVSeq_1.12.0
## [19] edgeR 3.20.9
                                   limma 3.34.9
## [21] EDASeq_2.12.0
                                   ShortRead_1.36.1
## [23] GenomicAlignments_1.14.2
                                   SummarizedExperiment_1.8.1
## [25] DelayedArray_0.4.1
                                   matrixStats_0.53.1
## [27] Rsamtools_1.30.0
                                   GenomicRanges_1.30.3
## [29] GenomeInfoDb_1.14.0
                                   Biostrings_2.46.0
## [31] XVector_0.18.0
                                   IRanges_2.12.0
## [33] S4Vectors_0.16.0
                                   BiocParallel_1.12.0
                                   BiocGenerics_0.24.0
## [35] Biobase_2.38.0
## [37] plotly_4.7.1
                                   ggplot2_3.0.0
## loaded via a namespace (and not attached):
     [1] uuid 0.1-2
                                backports_1.1.2
                                                        aroma.light_3.8.0
##
##
     [4] NMF 0.21.0
                                lazyeval 0.2.1
                                                        splines 3.4.3
##
     [7] rncl_0.8.2
                                NOISeq_2.22.1
                                                        gridBase_0.4-7
## [10] digest_0.6.15
                                foreach_1.4.4
                                                        htmltools 0.3.6
## [13] viridis_0.5.1
                                gdata_2.18.0
                                                        magrittr_1.5
                                cluster 2.0.7-1
                                                        doParallel 1.0.11
## [16] memoise 1.1.0
## [19] annotate_1.56.2
                                R.utils_2.6.0
                                                        prettyunits_1.0.2
```

```
[22] colorspace_1.3-2
                                 blob_1.1.1
                                                        crayon 1.3.4
##
    [25] RCurl_1.95-4.11
                                 jsonlite_1.5
                                                        genefilter_1.60.0
                                                        survival 2.42-6
    [28] phylobase 0.8.4
                                 bindr 0.1.1
    [31] iterators_1.0.10
                                 ape_5.1
                                                        glue_1.3.0
##
    [34] registry_0.5
                                 gtable_0.2.0
                                                        zlibbioc_1.24.0
    [37] kernlab 0.9-26
                                 DEoptimR 1.0-8
                                                        prabclus 2.2-6
##
    [40] scales 0.5.0.9000
                                 DESeq 1.30.0
                                                        mvtnorm 1.0-8
                                                        bibtex 0.4.2
    [43] DBI 1.0.0
                                 rngtools_1.3.1
##
##
    [46] Rcpp_1.0.0
                                 viridisLite_0.3.0
                                                        xtable 1.8-2
                                                        mclust_5.4.1
##
    [49] progress_1.2.0
                                 bit_1.1-14
                                                        RColorBrewer_1.1-2
    [52] htmlwidgets_1.2
                                 httr_1.3.1
    [55] fpc_2.1-11
                                 modeltools_0.2-22
                                                        flexmix_2.3-14
##
##
    [58] pkgconfig_2.0.1
                                 XML_3.98-1.12
                                                        R.methodsS3_1.7.1
                                                        howmany_0.3-1
##
    [61] nnet_7.3-12
                                 locfit_1.5-9.1
##
    [64] tidyselect_0.2.4
                                 labeling_0.3
                                                        rlang_0.2.1
##
    [67] reshape2_1.4.3
                                 munsell_0.5.0
                                                        cellranger_1.1.0
##
    [70] tools_3.4.3
                                 RSQLite_2.1.1
                                                        ade4_1.7-11
   [73] evaluate 0.11
                                 stringr 1.3.1
                                                        vaml 2.1.19
   [76] knitr_1.20
                                 bit64_0.9-7
                                                        robustbase_0.93-1.1
##
    [79] caTools 1.17.1
                                 purrr_0.2.5
                                                        whisker 0.3-2
##
   [82] R.oo_1.22.0
                                 xml2_1.2.0
                                                        biomaRt_2.34.2
    [85] compiler 3.4.3
                                 tibble 1.4.2
                                                        statmod_1.4.30
##
                                 RNeXML_2.1.1
    [88] geneplotter_1.56.0
                                                        stringi_1.2.3
##
    [91] GenomicFeatures 1.30.3 RSpectra 0.13-1
                                                        trimcluster 0.1-2
##
   [94] lattice_0.20-35
                                                        pillar 1.3.0
##
                                 nloptr 1.0.4
   [97] data.table_1.11.4
                                                        rtracklayer_1.38.3
                                 bitops_1.0-6
## [100] R6_2.2.2
                                 latticeExtra_0.6-28
                                                        hwriter_1.3.2
## [103] RMySQL_0.10.15
                                                        KernSmooth_2.23-15
                                 gridExtra_2.3
                                                        MASS_7.3-50
## [106] codetools_0.2-15
                                 boot_1.3-20
## [109] gtools_3.8.1
                                 assertthat_0.2.0
                                                        pkgmaker_0.27
                                                        locfdr_1.1-8
## [112] rprojroot_1.3-2
                                 withr_2.1.2
## [115] GenomeInfoDbData_1.0.0 diptest_0.75-7
                                                        hms_0.4.2
## [118] grid_3.4.3
                                 class_7.3-14
                                                        minqa_1.2.4
## [121] rmarkdown_1.10
```