RNA-seq analysis: SLBP

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This R Markdown document contains a walk-through for analyzing pooled (single-end) RNA-seq on single *D.melanogaster* histone overexpression mutant embryos caused by a knockdown of *Slbp* (Stem loop binding protein) gene and corresponding wild-type/ control. Raw transcript count data obtained by mapping to a reference transcriptome using Salmon (slurm/bash scripts calling the relevant functions to preprocess and map are in another repository).

This document includes

- performing differential gene expression (DE) analysis of single stage and time-series RNA-seq
- Gene enrichment analysis with other independent datasets from
- Gene ontology analysis
- custom plotting/ visualization in base R,ggplot2 and pheatmap
- "Desktop/RNAseq_output/" has been used as a local directory for this project, but that can be changed to any desired local or cloud based directory.

```
# Libraries for RNA-seq count data analysis
library(tximport)
library(DESeq2)
library(apeglm)
library(vsn)
library(sva)
library(magrittr)
library(matrixStats)
library(metaRNASeq)
# Libraries for data handling and visualization
library(ggplot2)
library(dplyr)
library(pheatmap)
library(RColorBrewer)
library(genefilter)
library(dendsort)
library(reshape2)
#Libraries for database and GO analysis
library(goseq)
library(GO.db)
library(TxDb.Dmelanogaster.UCSC.dm6.ensGene)
library(org.Dm.eg.db)
```

Loading Gene annotation datasets transcript to gene ID to gene name mappings, downloaded from flybase

```
setwd("~/Desktop/RNAseq_output/Abo_Wt_StagedNC_flybase_transcriptome/")
ttg.slbp <- read.table ("scripts_samples_ttg/flybase_transcript_to_gene.txt", h=T)
head(ttg.slbp)</pre>
```

txId Gene

```
## 1 FBtr0081624 dmel_7SLRNA:CR32864
## 2 FBtr0100521
                              dmel a
## 3 FBtr0071764
                              dmel a
## 4 FBtr0342981
                              dmel_a
## 5 FBtr0071763
                              dmel a
## 6 FBtr0083388
                          dmel abd-A
fbgn.gene.conv <- read.table ("scripts_samples_ttg/flybase_fbgn_to_gene.tsv", h=T)
head(fbgn.gene.conv)
##
                         Gene
          geneId
## 1 FBgn0262029
                       dmel d
## 2 FBgn0052532 dmel_CG32532
## 3 FBgn0023536
                  dmel_CG3156
## 4 FBgn0029718
                  dmel_mRpL30
## 5 FBgn0031101
                  dmel_CG1631
## 6 FBgn0030952 dmel_CG12609
```

Differential Gene expression analysis

The differential gene expression analysis is split into the following phases

- exploratory analysis of the whole data to understand general trends
- comparisons of specific developmental stages to understand stage specific

Exploratory Analysis

5 SLBP_NCO_Pre9_R5_B2

SLBP_NC13_R1_B1

6

```
setwd("~/Desktop/RNAseq_output/Abo_Wt_StagedNC_flybase_transcriptome/data/SLBP/main/")
# Reading in the sample information file
sample.slbp.all <- read.csv("salmon_data_slbp_main_pilot_inc/samples_slbp_wt_main_pilot_inc.csv", h=T)</pre>
str(sample.slbp.all)
## 'data.frame':
                    28 obs. of 7 variables:
                   : Factor w/ 28 levels "SLBP_NCO_Pre9_R1_B2",..: 1 2 3 4 5 6 7 8 9 10 ...
##
   $ Sample
## $ Genotype
                   : Factor w/ 2 levels "SLBP", "WT": 1 1 1 1 1 1 1 1 1 1 ...
## $ Cycle
                   : Factor w/ 5 levels "NCO_Pre9", "NC13",..: 1 1 1 1 1 2 2 2 2 2 ...
                   : Factor w/ 6 levels "R1", "R2", "R3", ...: 1 2 3 4 5 1 1 2 2 3 ....
## $ Replicate
                   : Factor w/ 25 levels "SLBP_NCO_Pre9_R1",..: 1 2 3 4 5 6 6 7 7 8 ...
##
   $ Gen_Rep
## $ Gen_Rep_Batch: Factor w/ 28 levels "SLBP_NCO_Pre9_R1_B2",..: 1 2 3 4 5 6 7 8 9 10 ...
                   : Factor w/ 5 levels "SLBP NCO Pre9",..: 1 1 1 1 1 2 2 2 2 2 ...
   $ Gen Cyc
head(sample.slbp.all)
##
                  Sample Genotype
                                     Cycle Replicate
                                                               Gen_Rep
## 1 SLBP_NCO_Pre9_R1_B2
                             SLBP NCO_Pre9
                                                   R1 SLBP_NCO_Pre9_R1
## 2 SLBP_NCO_Pre9_R2_B2
                             SLBP NCO_Pre9
                                                   R2 SLBP_NCO_Pre9_R2
## 3 SLBP_NCO_Pre9_R3_B2
                             SLBP NCO_Pre9
                                                   R3 SLBP_NCO_Pre9_R3
## 4 SLBP_NCO_Pre9_R4_B2
                             SLBP NCO_Pre9
                                                   R4 SLBP_NCO_Pre9_R4
```

R1

R5 SLBP_NCO_Pre9_R5

SLBP_NC13_R1

SLBP NCO_Pre9

NC13

SLBP

```
##
           Gen_Rep_Batch
                                Gen_Cyc
## 1 SLBP_NCO_Pre9_R1_B2 SLBP_NCO_Pre9
## 2 SLBP_NCO_Pre9_R2_B2 SLBP_NCO_Pre9
## 3 SLBP_NCO_Pre9_R3_B2 SLBP_NCO_Pre9
## 4 SLBP_NCO_Pre9_R4_B2 SLBP_NCO_Pre9
## 5 SLBP NCO Pre9 R5 B2 SLBP NCO Pre9
         SLBP NC13 R1 B1
                             SLBP NC13
levels(sample.slbp.all$Gen_Cyc)
## [1] "SLBP_NCO_Pre9" "SLBP_NC13"
                                        "WT_NCO_Pre9"
                                                         "WT_NC13"
## [5] "WT_NC14"
sample.slbp.all$Gen_Cyc <- relevel(sample.slbp.all$Gen_Cyc, ref="WT_NCO_Pre9")</pre>
# Ordering the dataset such that WT NCO Pre9
# i.e. the most initial developmental stages is the reference for all comparisons
# Reading in the raw count matrix files and coverting it into a DESeq object
files.slbp.all <- file.path("salmon_data_slbp_main_pilot_inc", sample.slbp.all$Sample, "quant.sf")
names(files.slbp.all) <- paste0("Sample", 1:28)</pre>
all(file.exists(files.slbp.all))
## [1] TRUE
txi.slbp.all <- tximport(files.slbp.all, type = "salmon", tx2gene = ttg.slbp)</pre>
## reading in files with read_tsv
## 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
## summarizing abundance
## summarizing counts
## summarizing length
names(txi.slbp.all)
## [1] "abundance"
                              "counts"
                                                    "length"
## [4] "countsFromAbundance"
rownames(sample.slbp.all) <- colnames(txi.slbp.all$counts)</pre>
ddsTxi.slbp.all <- DESeqDataSetFromTximport(txi.slbp.all, colData = sample.slbp.all, design = ~ Gen_Cyc
## using counts and average transcript lengths from tximport
# The design here is Gen Cyc == Genotype*Cycle == Genotype + Cycle + Genotype:Cycle
```

There are a couple of other manipulations that can be performed before performing the actual analysis - The same libraries were sequenced on multiple lanes. These form technical replicates and can be combined into the appropriate biological replicate by summing the counts, using the collapseReplicates() function - The genes that have not received any reads or say below a threshold level of counts can be eliminated before analysis

main and interaction effects of Genotype and Cell cycle

```
ddsColl.slbp.all <- collapseReplicates (ddsTxi.slbp.all, ddsTxi.slbp.all$Gen_Rep, ddsTxi.slbp.all$Gen_R
colData(ddsColl.slbp.all)
## DataFrame with 25 rows and 8 columns
##
                                 Sample Genotype
                                                     Cycle Replicate
##
                               <factor> <factor> <factor> <factor>
## SLBP_NCO_Pre9_R1 SLBP_NCO_Pre9_R1_B2
                                            SLBP NCO_Pre9
## SLBP_NCO_Pre9_R2 SLBP_NCO_Pre9_R2_B2
                                            SLBP NCO Pre9
                                                                  R2
## SLBP_NCO_Pre9_R3 SLBP_NCO_Pre9_R3_B2
                                            SLBP NCO_Pre9
                                                                  R3
## SLBP_NCO_Pre9_R4 SLBP_NCO_Pre9_R4_B2
                                            SLBP NCO Pre9
## SLBP_NCO_Pre9_R5 SLBP_NCO_Pre9_R5_B2
                                            SLBP NCO_Pre9
                                                                  R5
## ...
                                    . . .
                                              . . .
                                                      . . .
                                                                 . . .
## WT NC14 R1
                          WT NC14 R1 B2
                                              WT
                                                      NC14
                                                                  R1
## WT NC14 R2
                          WT_NC14_R2_B2
                                               WT
                                                      NC14
                                                                  R2
                                                                  R3
## WT_NC14_R3
                          WT_NC14_R3_B2
                                               WT
                                                      NC14
## WT_NC14_R4
                          WT_NC14_R4_B2
                                               WT
                                                      NC14
                                                                  R4
## WT_NC14_R5
                                               WT
                                                      NC15
                                                                  R5
                          WT_NC14_R5_B2
                                           Gen_Rep_Batch
                             Gen_Rep
                                                                Gen_Cyc
                            <factor>
                                                <factor>
## SLBP_NCO_Pre9_R1 SLBP_NCO_Pre9_R1 SLBP_NCO_Pre9_R1_B2 SLBP_NCO_Pre9
## SLBP_NCO_Pre9_R2 SLBP_NCO_Pre9_R2 SLBP_NCO_Pre9_R2_B2 SLBP_NCO_Pre9
## SLBP_NCO_Pre9_R3 SLBP_NCO_Pre9_R3 SLBP_NCO_Pre9_R3_B2 SLBP_NCO_Pre9
## SLBP_NCO_Pre9_R4 SLBP_NCO_Pre9_R4 SLBP_NCO_Pre9_R4_B2 SLBP_NCO_Pre9
## SLBP_NCO_Pre9_R5 SLBP_NCO_Pre9_R5 SLBP_NCO_Pre9_R5_B2 SLBP_NCO_Pre9
## ...
## WT_NC14_R1
                          WT_NC14_R1
                                           WT_NC14_R1_B2
                                                                WT_NC14
## WT_NC14_R2
                          WT_NC14_R2
                                           WT_NC14_R2_B2
                                                                WT NC14
## WT_NC14_R3
                          WT_NC14_R3
                                           WT_NC14_R3_B2
                                                                WT_NC14
## WT NC14 R4
                          WT NC14 R4
                                           WT NC14 R4 B2
                                                                WT NC14
## WT NC14 R5
                          WT_NC14_R5
                                           WT_NC14_R5_B2
                                                                WT NC14
##
                          runsCollapsed
##
                            <character>
## SLBP_NCO_Pre9_R1 SLBP_NCO_Pre9_R1_B2
## SLBP_NCO_Pre9_R2 SLBP_NCO_Pre9_R2_B2
## SLBP_NCO_Pre9_R3 SLBP_NCO_Pre9_R3_B2
## SLBP_NCO_Pre9_R4 SLBP_NCO_Pre9_R4_B2
## SLBP_NCO_Pre9_R5 SLBP_NCO_Pre9_R5_B2
## ...
## WT_NC14_R1
                          WT_NC14_R1_B2
## WT_NC14_R2
                          WT_NC14_R2_B2
## WT_NC14_R3
                          WT_NC14_R3_B2
## WT NC14 R4
                          WT NC14 R4 B2
## WT_NC14_R5
                          WT_NC14_R5_B2
# Retaining a gene if it has over 1 reads in 2 or more samples
# This is a very permissive threshold to eliminate any unintentional filtration bias
ddsColl.slbp.all
## class: DESeqDataSet
## dim: 13728 25
```

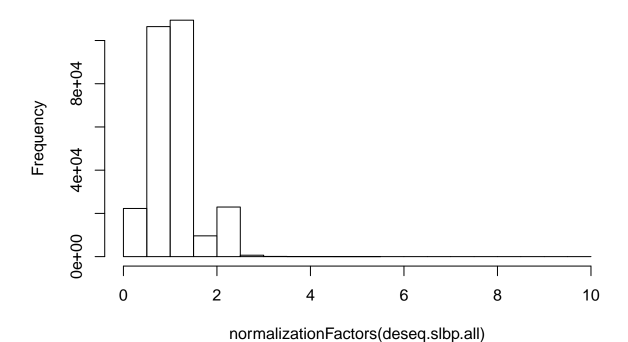
Collapsing the technical replicates

metadata(1): version

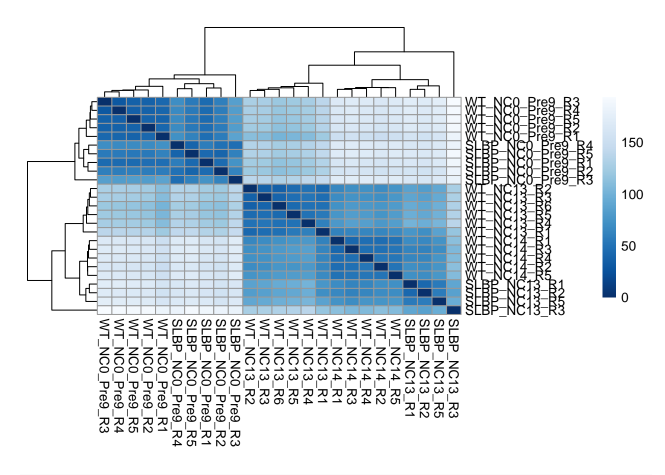
assays(2): counts avgTxLength

```
## rownames(13728): dmel_1-Dec dmel_1-Sep ... dmel_zye dmel_Zyx
## rowData names(0):
## colnames(25): SLBP_NCO_Pre9_R1 SLBP_NCO_Pre9_R2 ... WT_NC14_R4
    WT_NC14_R5
## colData names(8): Sample Genotype ... Gen_Cyc runsCollapsed
keep.slbp.all <- rowSums(counts(ddsColl.slbp.all) >1) >= 2
ddsColl.keep.slbp.all <- ddsColl.slbp.all[keep.slbp.all,]</pre>
ddsColl.keep.slbp.all # 10860 genes
## class: DESeqDataSet
## dim: 10860 25
## metadata(1): version
## assays(2): counts avgTxLength
## rownames(10860): dmel_1-Dec dmel_1-Sep ... dmel_zye dmel_Zyx
## rowData names(0):
## colnames(25): SLBP_NCO_Pre9_R1 SLBP_NCO_Pre9_R2 ... WT_NC14_R4
     WT NC14 R5
## colData names(8): Sample Genotype ... Gen_Cyc runsCollapsed
Perform the DESeq2 normalization followed by
- regularized log transformation
- similarity between samples based on Euclidean distance
- principal components analysis (PCA)
deseq.slbp.all <- DESeq(ddsColl.keep.slbp.all)</pre>
## estimating size factors
## using 'avgTxLength' from assays(dds), correcting for library size
## estimating dispersions
## gene-wise dispersion estimates
## mean-dispersion relationship
## final dispersion estimates
## fitting model and testing
hist(normalizationFactors(deseq.slbp.all))
```

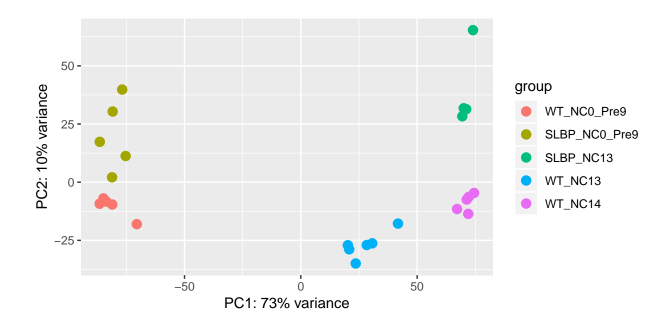
Histogram of normalizationFactors(deseq.slbp.all)



histogram of normalization factors typically distributed with a mean/ median ~1.0 rld.slbp.all <- rlog(deseq.slbp.all, blind=T)</pre> sampleDists.slbp.all <- dist(t(assay(rld.slbp.all)))</pre> hm.mat.slbp.all <- as.matrix(sampleDists.slbp.all)</pre> colors <- colorRampPalette(rev(brewer.pal(9, "Blues")))(255)</pre> # heirarchical clustering of the sample distances hm.col.clust.slbp.all <- hclust(dist(t(hm.mat.slbp.all)))</pre> sort.hclust <- function(...) as.hclust(dendsort(as.dendrogram(...)))</pre> hm.col.clust.slbp.all <- sort.hclust(hclust(dist(t(hm.mat.slbp.all))))</pre> hm.row.clust.slbp.all <- sort.hclust(hclust(dist(hm.mat.slbp.all)))</pre> #visualizing the clustered sample distances pheatmap(= hm.mat.slbp.all, mat color = colors, cluster_cols = hm.col.clust.slbp.all, = hm.row.clust.slbp.all, cluster_rows show_rownames)



#plotting PCA
plotPCA(rld.slbp.all, intgroup=c("Gen_Cyc"),ntop=5000)



The Euclidean distance based similarity of expression data for WT and Slbp embryos at different time-points demonstrate that replicates are more similar to each other than they are to other genotypes and/ or timepoints

Principal Components Analysis of Slbp and WT expression data also shows that global transcriptomic profile is shifted in Slbp NC13 as compared WT NC13 and is more similar to WT NC14 indicating that the onset of transcription is advanced in Slbp embryos