Read counts to DGE, Part I

This script will show you how to:

- Read in `featureCounts` results into R.
- Use `DESeq2` to:
  - normalize read counts for sequencing depth
  - transform reads to the log2 scale
- Accompany each step by exploratory plots.

You can generate an `html` document out of this entire script by clicking the Knit HTML button in RStudio.

```r
options(stringsAsFactors = FALSE) # this will change a global setting, but just for this session

library(knitr)
opts_chunk$set(echo = TRUE, message = FALSE, cache=FALSE) # tuning knitr output

featureCounts

We aligned five samples for the WT and SNF2 condition, respectively.

How can you check which command was used to generate those `BAM` files?

Let’s read the result file into R (you’ll have to download it to your laptop first).

Loading additional libraries:

```r
library(ggplot2) # for making plots
library(magrittr) # for "pipe"-like coding in R
```

First, make sure you set the path to your working directory which should contain the count table.

```r
code <- "~/Documents/Teaching/ANGSD/RNA/" # download count table!
sfwd(code)
```

We will use the `DESeq2` package to normalize the samples for differences in their sequencing depth.

```
# not available via install.packages(), but through bioconductor
source("http://bioconductor.org/biocLite.R")
biocLite("DESeq2")
```

```r
library(DESeq2)
```

We will have to generate a `DESeqDataSet`, which is a specific R class that combines `data.frames` and `matrices` into one object. The `data.frames` typically contain metadata about the samples and genes (e.g. gene IDs, sample conditions), while the matrices contain the expression values.

Find out via `?DESeqDataSetFromMatrix` how to generate a `DESeqDataSet`!

We need two tables: `countData` and `colData`.

- `colData`: `data.frame` with all the variables you know about your samples, e.g., experimental condition, the type, and date of sequencing and so on. Its `row.names` should correspond to the unique sample names.
- `countData`: should contain a matrix of the actual values associated with the genes and samples. Is equivalent to `assay()`. Conveniently, this is almost exactly the format of the `featureCounts` output.
```r
##folder <- "~/Documents/Teaching/ANGSD/RNA/"
folder <- "./"
# reading in featureCounts output
readcounts <- read.table(paste0(folder, "featCounts_Gierlinski_genes.txt"), header=TRUE)
head(readcounts)
```

```
## Geneid   Chr    Start   End   Strand Length
## 1  YAL012W chrI 130799 131983  +   1185
## 2  YAL069W chrI  335   649    +   315
## 3  YAL068W-A chrI  538   792    +   255
## 4  YAL068C chrI 1807  2169    -   363
## 5  YAL067W-A chrI 2480  2707    +   228
## 6  YAL067C chrI 7235  9016    -  1782
## ...alignment.SNF2_1_Aligned.sortedByCoord.out.bam
## 1  7351
## 2   0
## 3   0
## 4   2
## 5   0
## 6  103
## ...alignment.SNF2_2_Aligned.sortedByCoord.out.bam
## 1  7180
## 2   0
## 3   0
## 4   2
## 5   0
## 6   51
## ...alignment.SNF2_3_Aligned.sortedByCoord.out.bam
## 1  7648
## 2   0
## 3   0
## 4   2
## 5   0
## 6   44
## ...alignment.SNF2_4_Aligned.sortedByCoord.out.bam
## 1  8119
## 2   0
## 3   0
## 4   1
## 5   0
## 6   90
## ...alignment.SNF2_5_Aligned.sortedByCoord.out.bam
## 1  5944
## 2   0
## 3   0
## 4   0
## 5   0
## 6   53
## ...alignment.WT_1_Aligned.sortedByCoord.out.bam
## 1  4312
## 2   0
## 3   0
## 4   0
```

2
In principle, this is the format that we’ll need (columns = Samples, rows = genes), but particularly the sample names are a bit unwielding and we’re completely missing row.names.

Preparing the count matrix for DESeq2:

```r
# gene IDs should be stored as row.names
row.names(readcounts) <- gsub("-", ".", readcounts$Geneid)

# exclude the columns without read counts (columns 1 to 6 contain additional info such as genomic coordinates)
readcounts <- readcounts[, -c(1:6)]

# give meaningful sample names - there are many ways to achieve this
orig_names <- names(readcounts)
names(readcounts) <- c("SNF2_1", "SNF2_2", "SNF2_3", "SNF2_4", "SNF2_5", "WT_1", "WT_2", "WT_3", "WT_4", "WT_5") # most error-prone way!

# alternatives:
names(readcounts) <- c( paste("SNF2", c(1:5), sep = "_"),
                        paste("WT", c(1:5), sep = "_")) # less potential for typos

# even safer
names(readcounts) <- gsub(".*\([0-9]+\).", \"\1\", orig_names)
```

Always check your data set after you manipulated it!
str(readcounts)

## 'data.frame': 6692 obs. of 10 variables:
## $ SNF2_1: int 7351 0 0 2 0 103 2 5 13 46 ... 
## $ SNF2_2: int 7180 0 0 2 0 51 0 9 8 58 ... 
## $ SNF2_3: int 7648 0 0 2 0 44 0 6 10 45 ... 
## $ SNF2_4: int 8119 0 0 1 0 90 0 3 9 61 ... 
## $ SNF2_5: int 5944 0 0 0 0 53 0 1 6 45 ... 
## $ WT_1 : int 4312 0 0 0 0 12 0 10 9 33 ... 
## $ WT_2 : int 3767 0 0 0 0 23 0 5 12 41 ... 
## $ WT_3 : int 3040 0 0 0 0 21 0 2 4 31 ... 
## $ WT_4 : int 5604 0 0 2 0 30 0 4 4 45 ... 
## $ WT_5 : int 4167 0 0 2 0 29 0 3 8 25 ... 

head(readcounts)

## SNF2_1 SNF2_2 SNF2_3 SNF2_4 SNF2_5 WT_1 WT_2 WT_3 WT_4 WT_5
## YAL012W 7351 7180 7648 8119 5944 4312 3767 3040 5604 4167
## YAL069W 0 0 0 0 0 0 0 0 0 0
## YAL068W.A 0 0 0 0 0 0 0 0 0 0
## YAL067W.A 2 2 2 1 0 0 0 0 2 2
## YAL067C 103 51 44 90 53 12 23 12 21 30 29

In addition to the read counts, we need some more information about the samples. According to \texttt{?colData}, this should be a \texttt{data.frame}, where the \texttt{rows} directly match the \texttt{columns} of the count data.

Here’s how this could be generated in \texttt{R} matching the \texttt{readcounts data.frame} we already have:

\begin{verbatim}
sample_info <- DataFrame(condition = gsub("_\[0-9]+", ",", names(readcounts)),
                          row.names = names(readcounts))
\end{verbatim}

```
sample_info
```

## DataFrame with 10 rows and 1 column
## <character>
## SNF2_1 SNF2
## SNF2_2 SNF2
## SNF2_3 SNF2
## SNF2_4 SNF2
## SNF2_5 SNF2
## WT_1 WT
## WT_2 WT
## WT_3 WT
## WT_4 WT
## WT_5 WT

str(sample_info)

## Formal class 'DataFrame' [package "S4Vectors"] with 6 slots
## ..@ rownames : chr [1:10] "SNF2_1" "SNF2_2" "SNF2_3" "SNF2_4" ... 
## ..@ nrows : int 10
## ..@ listData :List of 1
## ..$ condition: chr [1:10] "SNF2" "SNF2" "SNF2" "SNF2" ...
## ..@ elementType : chr "ANY"
## ..@ elementMetadata: NULL
## ..@ metadata : list()
Let’s generate the DESeqDataSet:

```r
DESeq.ds <- DESeqDataSetFromMatrix(countData = readcounts,
                                   colData = sample_info,
                                   design = ~ condition)
```

```
DESeq.ds
## class: DESeqDataSet
## dim: 6692 10
## metadata(1): version
## assays(1): counts
## rownames(6692): YAL012W YAL069W ... YMR325W YMR326C
## rowData names(0):
## colnames(10): SNF2_1 SNF2_2 ... WT_4 WT_5
## colData names(1): condition
```

```
head(counts(DESeq.ds))
##     SNF2_1 SNF2_2 SNF2_3 SNF2_4 SNF2_5 WT_1 WT_2 WT_3 WT_4 WT_5
## YAL012W   7351   7180   7648   8119   5944  4312  3767  3040  5604  4167
## YAL069W    0     0     0     0     0    0    0    0    0    0
## YAL068W.A  0     0     0     0     0    0    0    0    0    0
## YAL068C    2     2     2     1     0    0    0    0    2    2
## YAL067W.A  0     0     0     0     0    0    0    0    0    0
## YAL067C   103    51    44    90    53    12   23   21   30   29
```

How many reads were counted for each sample (= library sizes)?

```r
colSums(counts(DESeq.ds))
```

```
##     SNF2_1 SNF2_2 SNF2_3 SNF2_4 SNF2_5 WT_1 WT_2 WT_3 WT_4 WT_5
## 9518261  8025575  8099295   9933479  59444312 37673040 56044167
```

```r
colSums(counts(DESeq.ds)) %>% barplot
```

Remove genes with no reads.
keep_genes <- rowSums(counts(DESeq.ds)) > 0
dim(DESeq.ds)

## [1] 6692 10
DESeq.ds <- DESeq.ds[ keep_genes, ]
dim(DESeq.ds)

## [1] 6394 10
counts(DESeq.ds) %>% str

## int [1:6394, 1:10] 7351 2 103 2 5 13 46 17 20 249 ...
## - attr(*, "dimnames")=List of 2
## ..$: chr [1:6394] "YAL012W" "YAL068C" "YAL067C" "YAL066W" ...
## ..$: chr [1:10] "SNF2_1" "SNF2_2" "SNF2_3" "SNF2_4" ...
assay(DESeq.ds) %>% str

## int [1:6394, 1:10] 7351 2 103 2 5 13 46 17 20 249 ...
## - attr(*, "dimnames")=List of 2
## ..$: chr [1:6394] "YAL012W" "YAL068C" "YAL067C" "YAL066W" ...
## ..$: chr [1:10] "SNF2_1" "SNF2_2" "SNF2_3" "SNF2_4" ...
Now that we have the data, we can start using DESeq's functions, e.g. estimateSizeFactors() for sequencing depth normalization.

DESeq.ds <- estimateSizeFactors(DESeq.ds)
sizeFactors(DESeq.ds)

## SNF2_1 SNF2_2 SNF2_3 SNF2_4 SNF2_5 WT_1 WT_2
## 1.4257612 1.1080380 1.1007930 1.4800919 0.8917712 0.6003659 0.9428913
## WT_3 WT_4 WT_5
## 0.7674773 1.1383612 0.9003437
plot(sizeFactors(DESeq.ds), colSums(counts(DESeq.ds)))
The read counts normalized for sequencing depth can be accessed via `counts(..., normalized = TRUE)`. Let’s check whether the normalization helped adjust global differences between the samples.

```r
# setting up the plotting layout
par(mfrow=c(1,2))

counts.sf_normalized <- counts(DESeq.ds, normalized=TRUE)

# adding the boxplots
boxplot(counts.sf_normalized, main = "SF normalized")
boxplot(counts(DESeq.ds), main = "read counts only")
```

We can’t really see anything. It is usually helpful to transform the normalized read counts to bring them onto more similar scales.
To see the influence of the sequencing depth normalization, make two box plots of log2(read counts) - one for unnormalized counts, the other one for normalized counts (exclude genes with zero reads in all samples).

```r
par(mfrow=c(1, 2))  # to plot the two box plots next to each other
boxplot(log2(counts(DESeq.ds)), notch=TRUE,
        main = "Non-normalized read counts\n(log-transformed)",
        ylab="read counts")

boxplot(log2(counts(DESeq.ds, normalize=TRUE)), notch=TRUE,
        main = "Size-factor-normalized read counts\n(log-transformed)",
        ylab="read counts")
```
Understanding more properties of read count data

Characteristics we've touched upon so far:

- zeros can mean two things: no expression or no detection
- fairly large dynamic range

Make a scatterplot of log normalized counts against each other to see how well the actual values correlate which each other per sample and gene.

```r
# non-normalized read counts plus pseudocount
log.counts <- log2(counts(DESeq.ds, normalized = FALSE) + 1)
# instead of creating a new object, we could assign the values to a distinct matrix
# within the DESeq.ds object
assay(DESeq.ds, "log.counts") <- log.counts

# normalized read counts
log.norm.counts <- log2(counts(DESeq.ds, normalized=TRUE) + 1)
assay(DESeq.ds, "log.norm.counts") <- log.norm.counts
```

```r
par(mfrow=c(2,1))
DESeq.ds[, c("WT_1","WT_2")]
#> assay(. , "log.norm.counts") %>% plot(. , cex=.1, main = "WT_1 vs. WT_2")
DESeq.ds[, c("SNF2_1","SNF2_2")]
#> assay(. , "log.norm.counts") %>% plot(. , cex=.1, main = "SNF2_1 vs.
```
Every dot = one gene.

The fanning out of the points in the lower left corner (points below $2^5 = 32$) indicates that read counts correlate less well between replicates when they are low.

This observation indicates that the standard deviation of the expression levels may depend on the mean: the lower the mean read counts per gene, the higher the standard deviation.

This can be assessed visually; the package vsn offers a simple function for this.

```r
par(mfrow=c(1,1))
# generate the base meanSdPlot using sequencing depth normalized log2(read counts)
log.norm.counts <- log2(counts(DESeq.ds, normalized=TRUE) + 1)
msd_plot <- vsn::meanSdPlot(log.norm.counts,
   ranks=FALSE, # show the data on the original scale
   plot = FALSE)
msd_plot$gg +
ggtitle("Sequencing depth normalized log2(read counts)") +
ylab("standard deviation")
```

From the help for `meanSdPlot`: The red dots depict the running median estimator (window-width 10 percent). If there is no variance-mean dependence, then the line formed by the red dots should be approximately horizontal.

The plot here shows that there is some variance-mean dependence for genes with low read counts. This means that the data shows signs of heteroskedasticity.

Many tools expect data to be homoskedastic, i.e., all variables should have similar variances.
DESeq offers two ways to shrink the log-transformed counts for genes with very low counts: \texttt{rlog} and \texttt{varianceStabilizingTransformation (vst)}.

We’ll use \texttt{rlog} here as it is an optimized method for RNA-seq read counts: it transforms the read counts to the log2 scale while simultaneously minimizing the difference between samples for rows with small counts and taking differences between library sizes of the samples into account. \texttt{vst} tends to depend a bit more on the size factors, but generally, both methods should return similar results.

```
DESeq.rlog <- rlog(DESeq.ds, blind = TRUE) # this actually generates a different type of object
# set blind = FALSE if the conditions
# are expected to introduce strong differences in a large proportion of the genes
```

```
par(mfrow=c(1,2))
plot(log.norm.counts[,1:2], cex=.1,
     main = "size factor and log2-transformed")

# the rlog-transformed counts are stored in the accessor "assay"
plot(assay(DESeq.rlog)[,1],
     assay(DESeq.rlog)[,2],
     cex=.1, main = "rlog transformed",
     xlab = colnames(assay(DESeq.rlog[,1])),
     ylab = colnames(assay(DESeq.rlog[,2]) )
```

As you can see in the left plot the variance - that is higher for small read counts - is tightened significantly using \texttt{rlog}. What does the mean-sd-plot show?

```
# rlog-transformed read counts
msd_plot <- vsn::meanSdPlot( rlog.norm.counts, ranks=FALSE, plot = FALSE)
msd_plot$gg + ggtitle("rlog transformation")
```
save.image(file = "RNAseqGierlinski.RData")