

Homework 5
BIOS-7659/CPBS-7659
Due 10/27/2020, 9AM

1. Next Generation Sequencing: Sample Size Estimates

- Install the **RNASeqPower** package from BioConductor.
- Install the **edgeR** and **cqn** packages from BioConductor.
- From **cqn**, use **data(montgomery.subset)** to load human RNA-seq data from lymphoblastoid cells from 10 subjects (see Montgomery et al., *Nature* 464:773-777). Use **data(uCovar)** to load the GC content and length of the genes in this data set.

- (a) Using **rnapower()**, recreate Figure 3 from the journal club paper using : Hart et al., (2013) “Calculating sample size estimates for RNA sequencing data.” *J Comput Biol.* 20:970-8. What does this figure show?
- (b) For the Montgomery data, create a row in Table 1 in the Hart et al. paper. Note that genes that have zero counts in all 10 samples were already excluded (see **help(montgomery.subset)**), so “% mapped”, will be 100%. How does the Montgomery data compare to the other data sets in Table 1?
- (c) Calculate the biological coefficient of variations (CV) from the Montgomery human lymphoblastoid data (use the function **estimateTagwiseDisp()** in **edgeR**, see problem 3). Plot the histogram and empirical cdf (use **ecdf()** function) and report the median and 90% percentile. How do the CVs compare with the examples in the Hart et al., paper in Figure 2?

Hint: Note that Hart et al., “estimated the coefficient of variation (CV) in expression across samples in the data set using a negative binomial model (**edgeR**).” See the section “Biological coefficient of variation” in <http://www.ncbi.nlm.nih.gov/pmc/articles/PMC3378882/> for a description on how the CVs are obtained and the **edgeR** User’s Guide, see section 2.8.2-2.8.3 and 2.9.1

<http://bioconductor.org/packages/release/bioc/vignettes/edgeR/inst/doc/edgeRUsersGuide.pdf>

- (d) Using **rnapower()**, recreate Figure 3 from Hart et al. again but with two curves using the median and the 90% percentile CV across genes for the Montgomery data. What sample size do you recommend?
- (e) Using **rnapower()**, recreate the curve (not the histogram) in the top Figure 4 from the Hart et al., paper. If you cannot recreate the figure, please explain any differences.

2. Next Generation Sequencing: Pre-Processing

- Install the `yeastRNASeq` and `EDASeq` packages from BioConductor. Familiarize yourself with these packages by looking at the manuals and the reference Risso et al. (2011) “GC-Content Normalization for RNA-Seq Data.” *BMC Bioinformatics* 12(1), pp. 480.
 - Use `data(geneLevelData)` to load the `geneLevelData` data, which is a yeast RNA-seq data set on two mutant and two wildtype strains (see Lee A et al. (2008) *PLoS Genet* 4(12): e1000299). Use `data(yeastGC)` to also load the GC content and length (`data(yeastLength)`) of annotated yeast genes.
- (a) Within `geneLevelData`, how many genes have all zeros as counts? How many have at least one sample with a zero?
- Save a new object `geneLevelDataFilter` only containing genes with ≥ 10 counts (summed over all samples).
 - Based on `geneLevelDataFilter`, run the following code to create a `SeqExpressionSet` object for the `EDASeq` functions.

```
exprs = as.matrix(geneLevelDataFilter) # matrix of counts
sub = intersect(rownames(geneLevelDataFilter), names(yeastGC))
exprs = exprs[sub,] #only examine genes with annotated GC content/length
row.names(exprs) = NULL #remove row and column names
colnames(exprs) = NULL
#Create SeqExpressionSet, which contains counts, labels for the
#samples and GC content/length
counts = newSeqExpressionSet(counts=exprs,
phenoData=data.frame(conditions = colnames(geneLevelDataFilter)),
featureData=AnnotatedDataFrame(data.frame(gc=yeastGC[sub],
length = yeastLength[sub])))
```
- (b) For the following plots, use the log scale (look at the manual/help for options to use the log scale).
- To plot the counts by sample, use the `boxplot()` function from the `EDASeq` package on `counts`. Do you see a need for normalization?
 - To plot the mean by variance plot, use `meanVarPlot()`. What trends do you see?
 - To assess any biases by GC content, use `biasPlot()`. What trends do you see?
 - To assess any biases by length, use `biasPlot()`. What trends do you see?
- (c) Apply `withinLaneNormalization()` to normalize by GC content. See help file and try different methods for normalization with the “which” option. Describe the different methods. Use `biasPlot()` before and after normalization. How do the methods compare? Save the normalized results in a new object to use in the next part.

- (d) Using the within-lane normalized data from the previous part, apply `betweenLaneNormalization()` to normalize across samples. Try different methods for normalization with the “which” option. Describe the different methods. Use `boxplot()` before and after normalization. How do the methods compare?