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 your 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  $\geq 10$  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?