## Homework/Mini Project 3

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Now its time to practice what we have learned in class and learn even more! For this homework/mini project you will do a RNA-seq analysis of the TB/HIV dataset. Note that your homework should be written in R Markdown, and turned in by uploading a tarball with your .Rmd, .html (from Rmarkdown, MultiQC, etc) and other outputs on Canvas.

## RNA-sequencing analysis

- 1. To access the data for Homework 3, you will have to download the data from the Sequence Read Archive (SRA) using the sratoolkit. There are 33 fastq files, with the SRR numbers are listed in the homework3\_srr.txt file.
- 2. Align the reads to the human genome reference using your choice of the Rsubread or the STAR aligners.
- 3. Use the featureCounts function in the Rsubread package to generate a counts file for this dataset.
- 4. Generate a SummarizedExperiment object for your counts. The colData for these data are provided in the homework3\_metadata.txt file.
- 5. Preprocess these data by removing TB-HIV-ART samples (should be two of them), removing any genes with 0 expression for all samples, and by generating a log counts per million assay.
- 6. (Extra credit) Create a batch corrected assay in your SummarizedExperiment using ComBat-Seq. You can use the ComBat\_Seq function in the sva package, or simply do it in BatchQC and then extract the SummarizedExperiment. For this example, pretend that disease\_status is the batch variable. Note that this will remove the disease status variability, so don't use this assay in the following analyses! This was merely a practice for cases where you have an actual batch variable.
- 7. Apply PCA and UMAP to your data and generate dimension reduction plots for the results. Color the TB-HIV to the HIV only samples in different colors. Note that you should be using the log CPM values for this analysis.
- 8. Use DESeq2 to do a differential expression analysis (on the counts) comparing the TB-HIV to the HIV only samples. Provide the top 50 most differentially expressed genes.
- 9. Now conduct the same analysis using limma on the log CPM values. How do the DESeq2 results compare to the limma results?
- 10. Give a heatmap plot of either the DESeq2 or the limma results (top 50). Add a colorbar for disease status.
- 11. Conduct a pathway analysis of the top 50 genes usign a tools such as enrichR (through R or online). What are the top scoring pathways?
- 12. (More extra credit) Conduct a TBSignatureProfiler analysis on these data including signature heatmaps, individual boxplots, and AUC boxplots. Interpret your findings.