# Clinical and Research Genomics Assignment #2 From Lecture\_04-06 (March 28th):

### RNA-Sequencing, Single-cell Expression, Epitranscriptomes, and Gene Fusions

Assignment: Answer questions about RNA-Sequencing and their methods Due Date: 5:00PM on April 5<sup>th</sup>

This week has two sections: short-answer questions and a data exercise.

## **Reading and Problem Sets:**

Chapters 5 and 6 from "Computational Genomics With R."

Chapter 5: <a href="http://compgenomr.github.io/book/processingReads.html">http://compgenomr.github.io/book/processingReads.html</a>

Chapter 6: <a href="http://compgenomr.github.io/book/rna-seq-analysis-overview.html">http://compgenomr.github.io/book/rna-seq-analysis-overview.html</a>

For more background information on genomics and R, see Chapters 1-4.

#### **Questions:**

State whether the statement is true or false, then explain with one sentence:

- 1) There are 20 types of RNA present in cells.
- 2) Once spliced and 5' capped, an RNA is no longer modified.
- 3) Gene fusions identified by RNA sequencing always correspond to rearrangements in the DNA as well.

#### **Essay Questions:**

- 1) If you were designing an experiment with RNA-Sequencing for each sample:
  - a.) How much sequencing is required to assay each sample?
  - b.) What are the parameters that affect this depth of sequencing?
  - c.) How would you calculate an expression value?

Please hand the assignment on the day of the lecture, or email if you cannot attend.

For any questions, please contact Chandrima Bhattacharya (<a href="mailto:chb4004@med.cornell.edu">chb4004@med.cornell.edu</a>), Ebrahim Afshinnekoo (<a href="mailto:eba2001@med.cornell.edu">eba2001@med.cornell.edu</a>), or Professor Mason (<a href="mailto:chm2042@med.cornell.edu">chm2042@med.cornell.edu</a>)