## QCB Workshop W5a Quiz

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## This quiz is due November 30, 2021.

Feel free to use any materials you like to help you answer these questions. Email your responses to me at abtbhatt@g.ucla.edu. There are 20 total points. There are bonus points along the way too, but you are capped at a total of 20 points.

- 1. (1 point) There are two files at this Google Drive link: https://drive.google.com/drive/u/1/folders/1HZUWZBzr9pCUNXFzJcNMCjmvOklZ2VHw. They are named KD2.fastq.gz and SC2.fastq.gz. Download them to your local computer and upload them to hoffman2. What software did you use to upload them to hoffman2? For 1 bonus point, what kind of client did you use (four words)?
- 2. (3 points) Using UNIX commands, how many lines are in KD2.fastq.gz? How many reads does this correspond to? What commands did you use? For this last question, there is no one right answer.
- 3. (3 points) Run quality evaluation on both fastq.gz files using FastQC. What kind of files does FastQC output? What commands did you run? What is the sequence length for SC2.fastq.gz, according to the FastQC report?
- 4. (4 points) Using trimmomatic, trim both FastQC files with the following parameters: minimum leading and trailing base qualities of 15 and a minimum final read length of 60 basepairs. How many reads were trimmed from both fastq files? For 1 bonus point, what command(s) did you use?
- 5. (2 points) You already have a reference genome for Chromosome 18 built. What are the two types of files that are required to build a reference genome sequence database using STAR?
- 6. (3 points) Align the cleaned and trimmed reads for both samples to this reference genome. What commands did you use? What proportion of reads mapped unambiguously to the genome database?
- 7. (3 points) Count the number of reads that map to each gene using htseq-count. At the end of the run, there is some information that the program outputs to the terminal. What commands did you use? For each sample, how many alignments were processed? For 1 bonus point, HTSeq is a software package written in what language?
- 8. (1 point) What R package can you use to read in the counts from htseq-count output?