Homework 1

Homework can be submitted via the github link which will create a repository for you with basic template of files you can edit to solve the homework. See the the table for homework submission links which will help you create a github repository in the class team.cd https://piazza.com/ucr/fall2020/gen220/resources

- 1. Write a script called download_count.sh which does the following.
 - Download the data file https://ftp.ncbi.nlm.nih.gov/pub/UniVec/Un iVec_Core from NCBI
 - Print out the count of the number of FASTA format sequences in this file - see Wikipedia FASTA format - each record starts with a >
- 2. Write a script called summary_exons.sh which summarizes the total length of exons in the file data/rice_random_exons.bed. These data are in the BED file format. The columns are "Chromosome", "Start position", "Stop position". The length of a feature (or exon in this case) is computed by doing the computation: STOP START
 - read in the file
 - use a loop structure to read each line
 - add up the length of each exon by summing this into a variable
 - Print out the total length of exon features at the end.
 - You do not need to save this for each chromosome, just print out the total length for this example - however if this is too easy for you, go ahead and make a more sophisticated report which presents, per chromosome, the total length of exons as well as the total number of exons, and the average length of exons.
- 3. Write a script called strand_gene_count.sh to calculate the number of genes that are on the positive (+) and negative (-) strand in the file. This file format is called GFF the strand of the gene is encoded in 7th column.
- https://fungidb.org/common/downloads/release-48/ScerevisiaeS288c/gf f/data/FungiDB-48_ScerevisiaeS288c.gff