## 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. https://classroom.github.com/a/XQDj3DK-

- 1. Write a script called download\_count.sh which does the following.
  - Download the data file https://ftp.ncbi.nlm.nih.gov/pub/UniVec/UniVec\_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.
- https://fungidb.org/common/downloads/release-48/ScerevisiaeS288c/gff/data/FungiDB-48\_ScerevisiaeS288c.gff