## Linux Command Line

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For this portfolio I will use the example of bioinformatics to showcase the utility of the Linux command line. The data we will be using is the genome and protein datasets from the SARS-CoV-2 virus, which can be found at https://www.ncbi.nlm.nih.gov/genome/?term=SARS-CoV-2.

## Working with Files and File Systems

We start in a folder containing the files downloaded from the link above:

```
ls
GCF_009858895.2_ASM985889v3_genomic.fna.gz
GCF_009858895.2_ASM985889v3_protein.faa.gz
```

We first create a folder to work within using the mkdir command and use the cd command to move into this new file. Finally we can use the ls command to view the contents of this folder:

```
mkdir bioinf_ex
```

As expected, the directory is empty, but we can move the genome and protein files to our new folders. We use the cd .. to go one 'step back' in the directory and return to our original directory. We use the ls command again and then the mv command to move our two files into the new folder we have created:

```
mv GCF_009858895.2_ASM985889v3_protein.faa.gz -t bioinf_ex/
mv GCF_009858895.2_ASM985889v3_genomic.fna.gz -t bioinf_ex/
cd bioinf_ex/
```

We now check these files have been successfully moved:

```
ls
GCF_009858895.2_ASM985889v3_genomic.fna.gz
GCF_009858895.2_ASM985889v3_protein.faa.gz
```

We notice the .gz file extension, meaning we need to unzip the files. We can do this with the gzip command:

```
gzip -d GCF_009858895.2_ASM985889v3_genomic.fna.gz
gzip -d GCF_009858895.2_ASM985889v3_protein.faa.gz
```

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GCF\_009858895.2\_ASM985889v3\_genomic.fna GCF\_009858895.2\_ASM985889v3\_protein.fa