

# 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
```

```
ls
GCF_009858895.2_ASM985889v3_genomic.fna
GCF_009858895.2_ASM985889v3_protein.fa
```