RNA-seq analysis in R

Quantification of Gene Expression with Salmon

Exercise 1 - Create Salmon index

1. Create concatenated trancriptome/genome reference file

```
cat references/Mus_musculus.GRCm38.cdna.chr14.fa.gz \
   references/Mus_musculus.GRCm38.dna_sm.chr14.fa.gz \
        references/gentrome.chr14.fa.gz
```

2. Create decoy sequence list from the genomic fasta

```
echo "14" > references/decoys.txt
```

- 3. Use salmon index to create the index. You will need to provide three pieces of information:
 - the Transcript fasta file references/gentrome.chr14.fa.gz
 - the **decoys** references/decoys.txt
 - the salmon index a directory to write the index to, use references/salmon_index_chr14

Also add -p 7 to the command to instruct salmon to use 7 threads/cores. To find the flags for the other three pieces of information use:

```
salmon index --help
```

Version Info: This is the most recent version of salmon.

```
Index
Creates a salmon index.
Command Line Options:
 -v [ --version ]
                                print version string
 -h [ --help ]
                                produce help message
  -t [ --transcripts ] arg Transcript fasta file.
 -k [ --kmerLen ] arg (=31)
                                The size of k-mers that should be used for the
                                quasi index.
  -i [ --index ] arg
                            salmon index.
  --gencode
                                This flag will expect the input transcript
  . . .
  . . .
 -d [ --decoys ] arg
                            Treat these sequences ids from the reference as
                                the decoys that may have sequence homologous to
                                some known transcript. for example in case of
                                the genome, provide a list of chromosome name
                                --- one per line
```

```
salmon index \
  -t references/gentrome.chr14.fa.gz \
  -d references/decoys.txt \
```

```
-p 7 \
-i references/salmon_index_chr14
```

Exercise 2 - Quantify with Salmon

1. Make directory called salmon_output

mkdir salmon_output

- 2. Use salmon quant to quantify the gene expression from the raw fastq. To see all the options run salmon quant --help-reads. There are lot of possible parameters, we will need to provide the following:
 - salmon index references/salmon_index
 - -1 A Salmon needs to use some information about the library preparation, we could explicitly give this, but it is easy enough for Salmon to Automatically infer this from the data
 - File containing the #1 mates fastq/SRR7657883.sra_1.fastq.gz
 - File containing the #2 mates fastq/SRR7657883.sra_2.fastq.gz
 - Output quantification directory salmon_output/SRR7657883
 - --gcBias salmon can optionally correct for GC content bias, it is recommended to always use this
 - The number of threads to use 7

```
salmon quant \
    -p 7 \
    -i references/salmon_index \
    --gcBias \
    -1 A \
    -1 fastq/SRR7657883.sra_1.fastq.gz \
    -2 fastq/SRR7657883.sra_2.fastq.gz \
    -0 salmon_output/SRR7657883
```

Exercise 3 - Run multique

1. Run multique on the salmon_output directory and create a report called Salmon_quantification_report in the salmon_output directory. Look back to the previous session if you need a reminder on how to do this.

```
multiqc -z -n Salmon_quantification_report -o salmon_output salmon_output
```

- \Rightarrow salmon_output/Salmon_quantification_report.html
- 2. Open the report and determine what percentage of the reads have been *aligned* to the transcriptome. Compare this to the QC report from the previous session.

With Salmon we have $\sim 85\%$ of reads mapped to the transcriptome.

If you look at the RNAseq metrics section in the multiqc report we generated in the previous section, you should see that for SRR7657883 we had $\sim\!85\%$ reads aligned to Coding + UTR.