## Introduction to Bulk RNAseq data analysis

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

print version string

produce help message

quasi index.

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 ]
 -h [ --help ]
  -t [ --transcripts ] arg Transcript fasta file.
 -k [ --kmerLen ] arg (=31)
```

salmon index. This flag will expect the input transcript

-d [ --decoys ] arg

-i [ --index ] arg

--gencode

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

The size of k-mers that should be used for the

```
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.subset\_2M.sra\_1.fastq.gz
  - File containing the #2 mates  $fastq/SRR7657883.subset\_2M.sra\_2.fastq.gz$
  - --writeMappingssalmon\_output/SRR7657883.salmon.sam Instructs Salmon to output the read alignments in SAM format to the file salmon\_output/SRR7657883.salmon.sam.
  - 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.subset_2M.sra_1.fastq.gz \
    -2 fastq/SRR7657883.subset_2M.sra_2.fastq.gz \
    -writeMappings salmon_output/SRR7657883/SRR7657883.salmon.sam \
    -o salmon_output/SRR7657883
```

## Exercise 3

1. Transform your aligned SAM file in to a BAM file called SRR7657883.salmon.bam. Use the option -@ 7 to use 7 cores. This vastly speeds up the compression.

```
samtools view -b -@ 7 salmon_output/SRR7657883/SRR7657883.salmon.sam \
> salmon_output/SRR7657883/SRR7657883.salmon.bam
```

 $\Rightarrow$  salmon\_output/SRR7657883/SRR7657883.salmon.bam