## Introduction to Bulk RNAseq data analysis

QC of raw reads with FastQC - Solutions

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### 1. Using FastQC to QC raw reads

#### Exercise 1

- 1. a) Check the location of the current directory using the command pwd
  - b) If the current directory is not Course\_Materials, then navigate to the Course\_Materials directory using the cd (change directory) command:

#### cd ~/Course\_Materials

- 2. a) Use 1s to list the contents of the directory. There should be directory called fastq
  - b) Use 1s to list the contents of the fastq directory:

#### ls fastq

```
SRR7657883.sra_1.fastq.gz SRR7657883.subset_2M.sra_1.fastq.gz SRR7657883.sra_2.fastq.gz Test_adapter_contamination.gq.gz. SRR7657883.subset 2M.sra 2.fastq.gz
```

You should see two fastq files called  $SRR7657883.sra\_1.fastq.gz$  and  $SRR7657883.sra\_1.fastq.gz$ . These are the files for read 1 and read 2 of one of the samples we will be working with.

3. Run fastqc on one of the fastq files:

#### fastqc fastq/SRR7657883.sra\_1.fastq.gz

This creates two files in the fastq directory. The first is the QC report in html format and the second is a zip file containing the data summary data used to generate the report.  $\Rightarrow$  SRR7657883.sra\_1\_fastqc.html  $\Rightarrow$  SRR7657883.sra\_1\_fastqc.zip

4. Open the html report in a browser and see if you can answer these questions:

# Basic Statistics

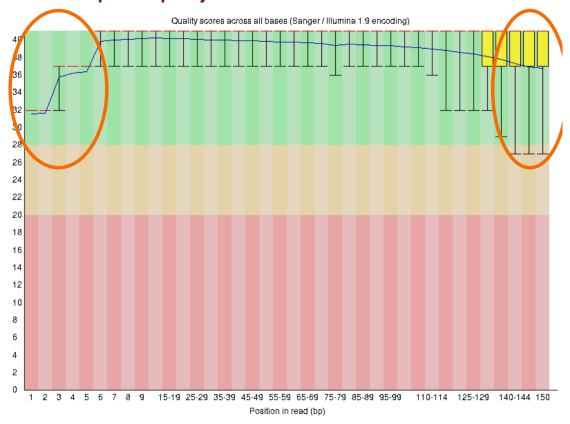
Measure	Value
Filename	SRR7657883_1.fastq.gz
File type	Conventional base calls
Encoding	Sanger / Illumina 1.9
Total Sequences	30120695
Total Bases	4.5 Gbp
Sequences flagged as poor quality	0
Sequence length	150
%GC	49

A) What is the read length?

150

B) Does the quality score vary through the read length?

## Per base sequence quality



Yes, the first few bases and the last few bases are typically of lower quality.

C) How is the data's quality?

Overall, pretty good.