

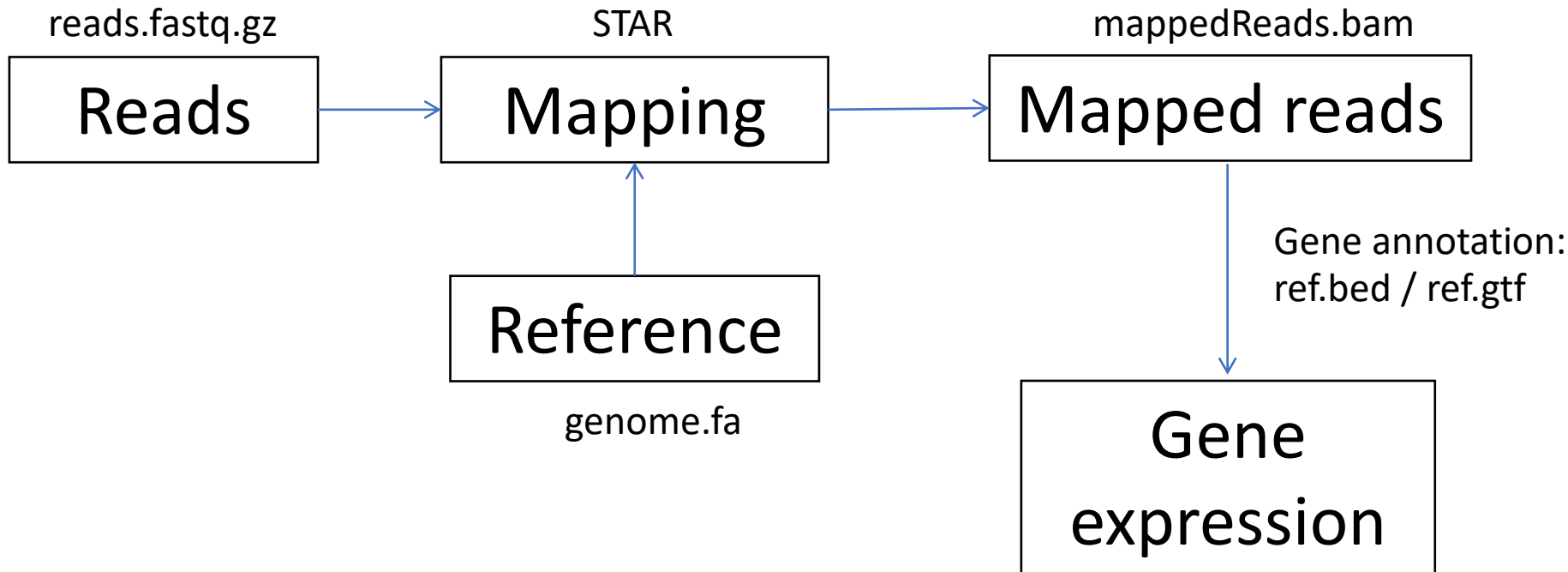
# RNA-seq QC analysis

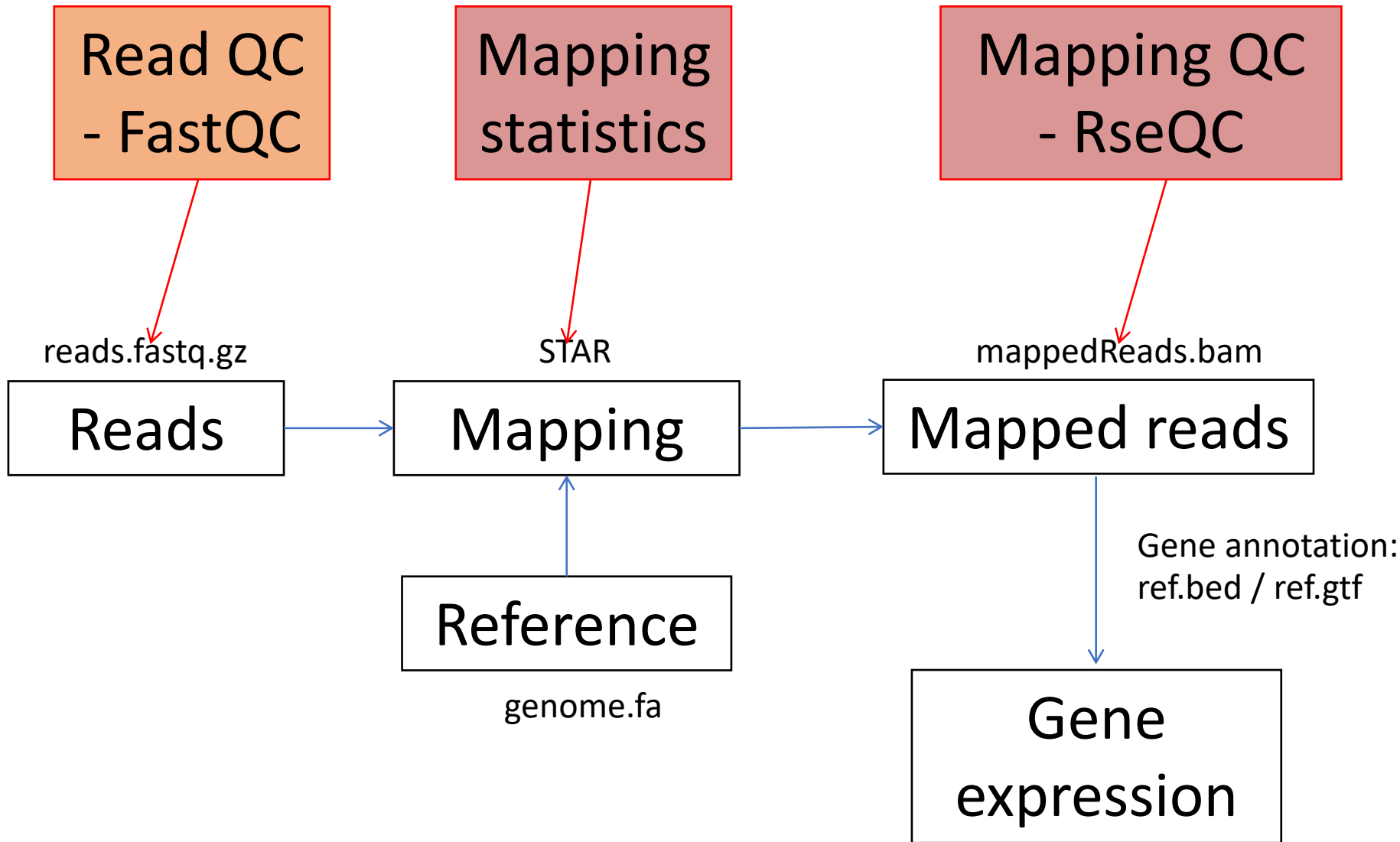
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RNA-seq data analysis

**Johan Reimegård**

# RNA-seq analysis workflow





# What could go wrong?

- RNA quality:
  - Degradation
  - Contaminations (pathogens or other sources)
  - GC-bias
  - Nuclear vs organelle reads
- Library prep:
  - Failed reactions
  - RNA / Adapter ratios – primer dimers
  - Clonal duplicates
  - Chimeric reads
  - Contaminations
- Sequencing:
  - Base calling errors
  - Uncalled bases
  - Low quality bases (3' end)
  - Contaminations
  - Sequence complexity

# From samples to reads

- may not be what you think they are

- Mixing samples
  - 30 samples with 5 steps from samples to reads has 24 300 000 potential mix ups of samples
  - Error rate 1/ 100 with 5 steps suggest that one of every 20 sample is mislabeled
- Experiments go wrong
  - 30 samples with 5 steps from samples to reads has 150 potential steps for errors
  - Error rate 1/100 with 5 steps suggest that one of every 20 samples the reads does not represent the sample
- Combine the two error sources and approximately one in every 10 samples is wrong

# From samples to reads

- may not be what you think they are

- Mixing samples
- Experiments go wrong
- How do we understand what went wrong?

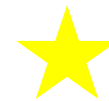
# From samples to reads

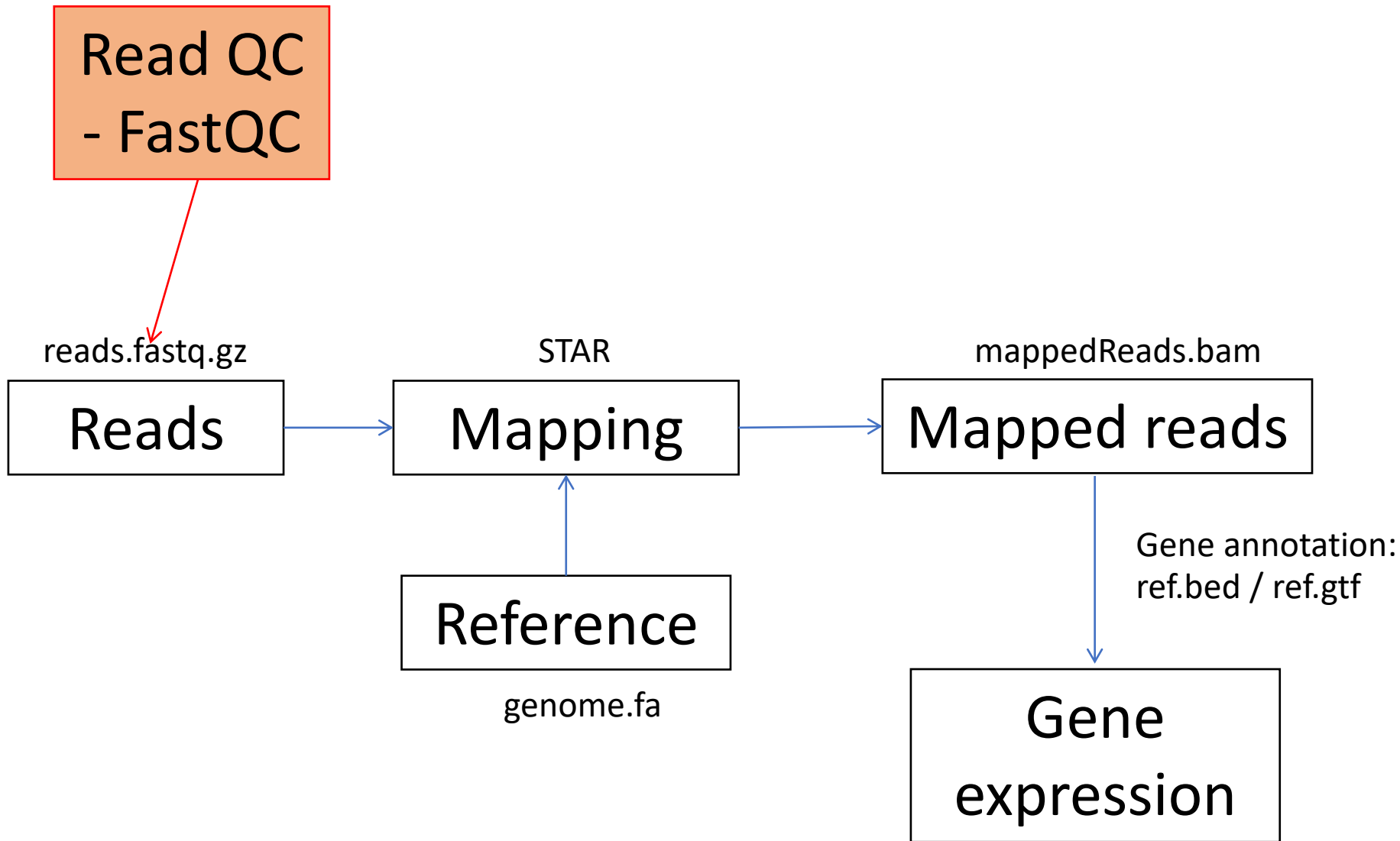
- may not be what you think they are

- Mixing samples
- Experiments go wrong
- How do we understand what went wrong?



**BIOINFORMATICS!**







# Fastq – read file format

Unique identifier

Sequence

```
@SEQ_ID  
GATTTGGGGTTCAAAGCAGTATCGATCAAATAGTAAATCCATTTGTTCAACTCACAGTTT  
+  
!''*(((((***+))%%%++) (%%%) .1***-+*'')) **55CCF>>>>>CCCCCCC65
```

Sequence quality

Paired end data usually in format sampleX\_1.fastq and sampleX\_2.fastq with same SEQ\_ID for both mate pairs, followed by /1 and /2 (or \_f and \_r)

# Basic read metrics with FastQC

A program that analyses some of the basic metrics on fastq raw read files.

- Quality
- Length
- Sequence bias
- GC content
- Repeated sequences
- Adapter contamination

```
$ fastqc -o outdir seqfile.fastq
# multiple files:
$ fastqc -o outdir seqfile_*.fastq
```

<http://www.bioinformatics.babraham.ac.uk/projects/fastqc/>



**Thank you. Questions?**

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**Johan Reimegård**