

# Quality control in ChIP-seq data

Using the ChIPQC package

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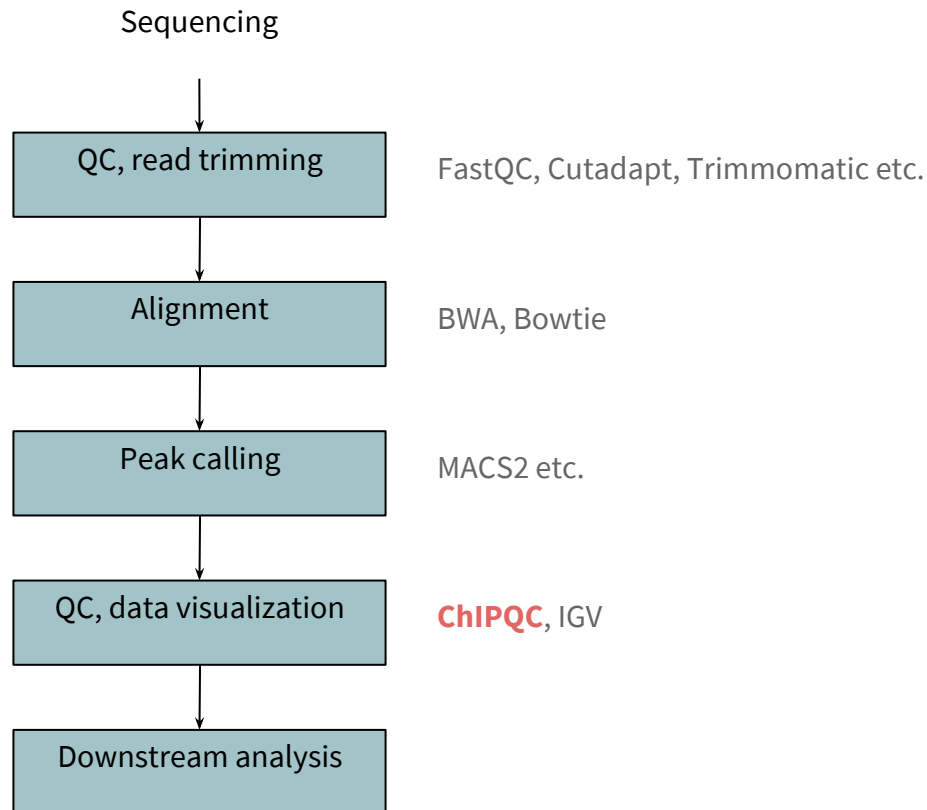
MRC Cancer Unit, University of Cambridge

CRUK Functional Genomics Workshop  
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# Overview

- Introduction
- Distribution of signal
  - Coverage profiles
  - FRiP: fraction of reads in peaks
  - REGI: relative enrichment in genomic intervals
  - FRiBl: fraction of reads in blacklisted regions
- Clustering of Watson/Crick reads
- Other factors affecting site discovery
  - Sequencing depth
  - Duplication rate, library complexity
  - Controls

# Workflow of ChIP-seq data processing



# Looking at ChIP-seq data

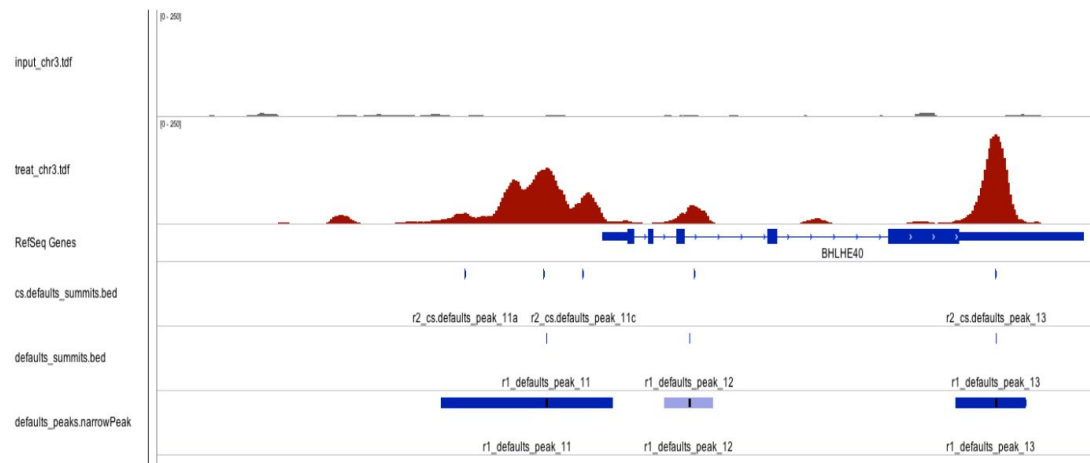
- A good quality ChIP-seq experiment will have high enrichment over background

- Ways to quantify the quality:

- Number of reads in peaks
- High peaks, low background
- Sequencing depth
- Diverse library (duplications)
- Low enrichment in control
- Similarity of replicates
- Genes closeby

- Tools to quantify quality:

- ChIPQC (T Carroll, *Front Genet*, 2014.)
- SPP package - Unix/Linux (PV Karchenko, *Nature Biotechnol*, 2008.)
- ChIP-seq guidelines and practices of the ENCODE and modENCODE consortia (Landt et al, *Genome Research*, 2012.)



# Things that can go wrong

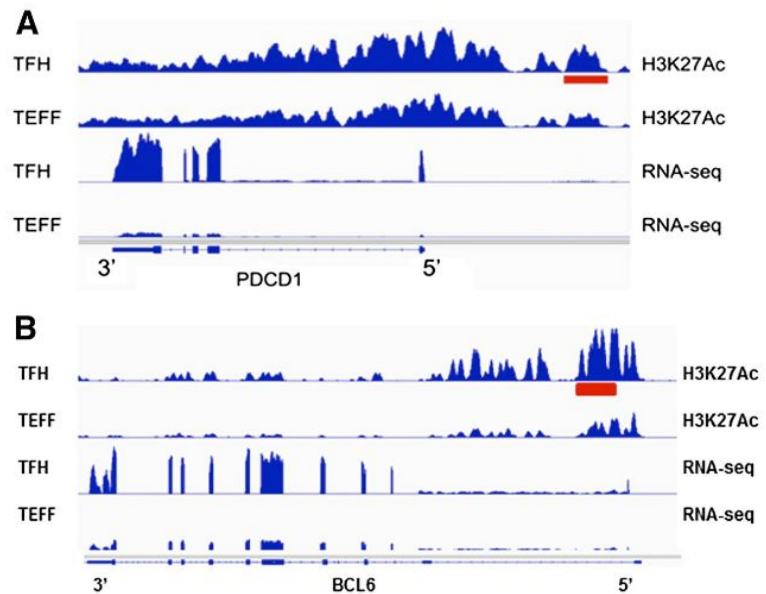
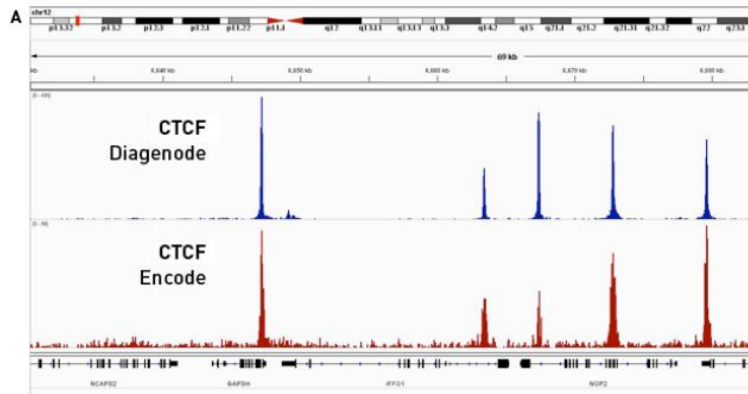
- The specificity of the antibody
  - Poor reactivity against the target of the experiment
  - High cross-reactivity with other proteins
- Degree of enrichment
- Biases during library preparation
  - PCR amplification bias
  - Fragmentation bias
- These can all affect the quality of the data and the number of sites detected
- Identification and removal of technical noise from the data is important

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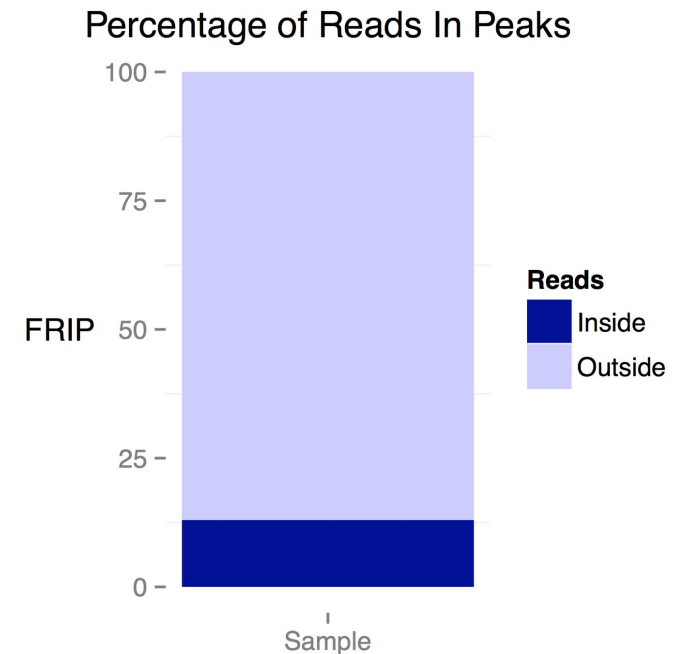
# Visualisation of coverage profiles

- Using IGV or USCS genome browser



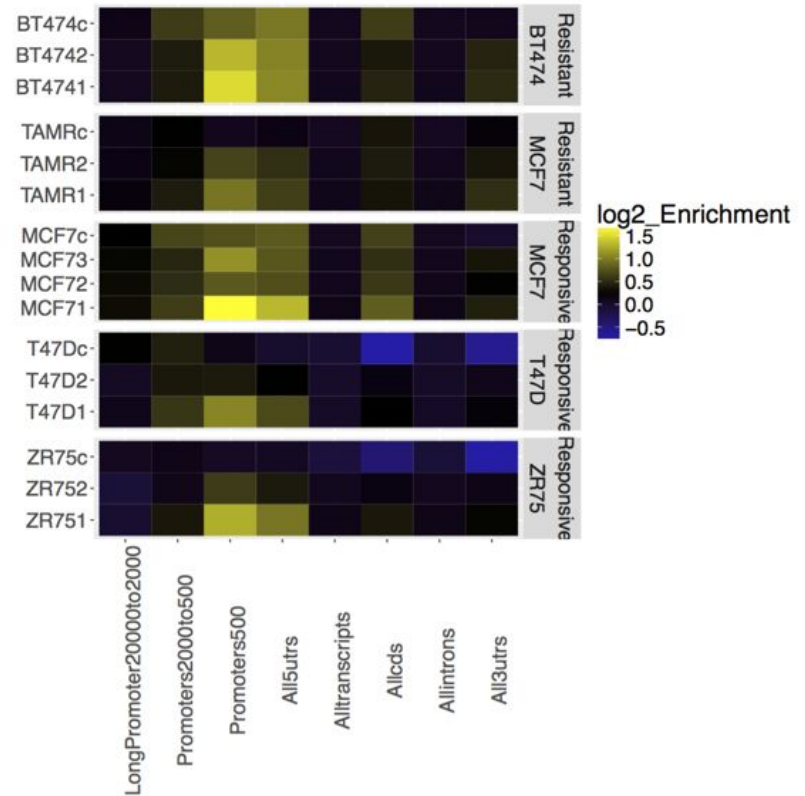
# FRiP - fragment of reads in peaks

- A useful metric to measure global ChIP enrichment
- Gives a quick understanding of the success of immunoprecipitation
- Guideline: in case of good quality FRiP is  $> 5\%$ 
  - But there are known examples of good quality data with FRiP  $< 1\%$



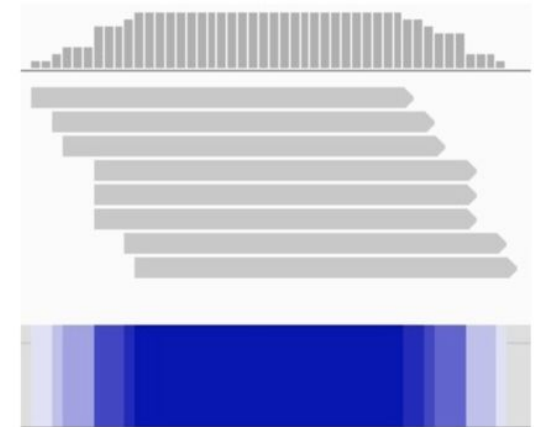
# REGI - relative enrichment in genomic intervals

- Proteins might have a high expected enrichment in certain genomic regions, like promoters, UTRs, introns, etc.
- This plot helps to identify whether our experiment worked as expected and/or to reveal interesting behaviours



# Dispersion of coverage

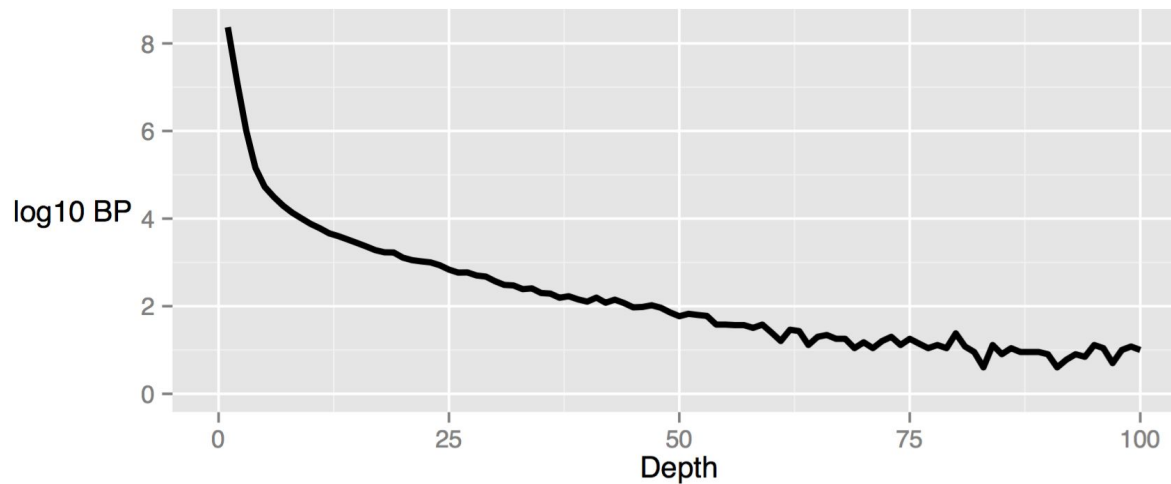
- The depth of coverage is the number of fragments at a specific genomic region
- To build a coverage profile
  - Measure the number of base pairs with a given depth of coverage
  - Normalise to the number of reads to compare samples
- We expect the depth to have large diversity in an enriched ChIP dataset



Depth	Base Pairs
1	3
2	4
3	3
5	3
6	4
7	3
8	26

# Dispersion of coverage

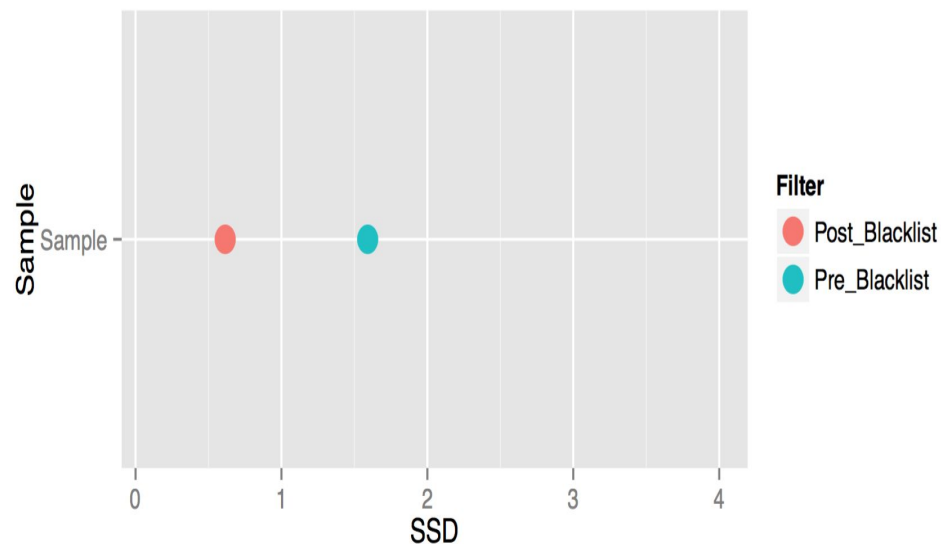
- Dispersion coverage profile plotted with CHIPQC
- More enriched libraries have higher number of bases at greater depths
- Profile of control samples usually drops more quickly
- The gap between samples and controls indicates enrichment



# Dispersion of coverage

- **SSD**: standardised standard deviation
- Metric to assess dispersion coverage developed in htSeqTools package
- Provides measure of pile-up across the genome, it is expected to be:
  - High for samples with enriched regions
  - Low for controls with uniform coverage
- This measure is highly influenced by regions, where the coverage is high because of some mapping error, like blacklisted regions

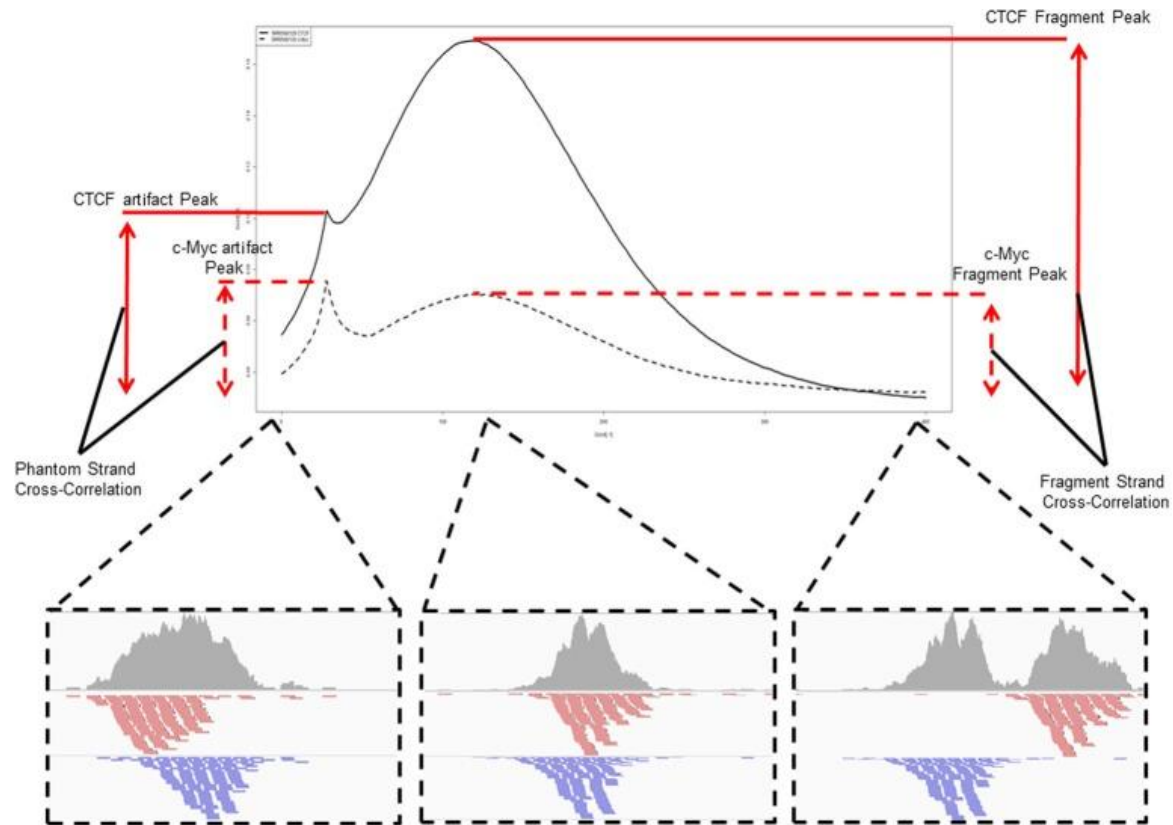
$$SSD = \frac{SD}{\sqrt{n}}$$



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# Clustering of Watson/Crick reads



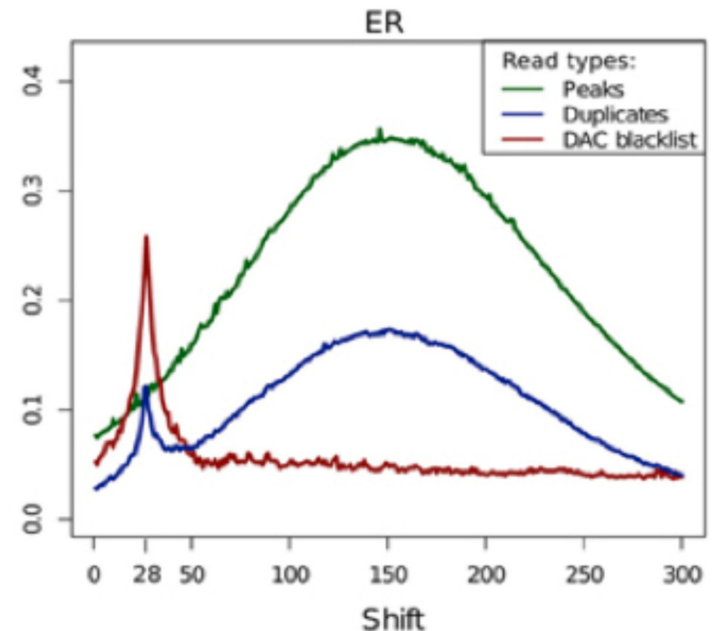
# Clustering of Watson/Crick reads

- Fragment length can be estimated from the data
  - Cross-correlation: correlation of reads on positive and negative strand after successive read shifts
  - Cross-coverage: coverage of reads on both strands after successive shifts of reads on one strand; the area covered by reads will be reduced after the shifting
- These metrics are computed in ChIPQC:

$$FragCC = CC_{fragmentLength}$$

$$RelCC = \frac{FragCC}{CC_{readLength}}$$

- Blacklisted regions have a large contribution to read-length cross-coverage peaks



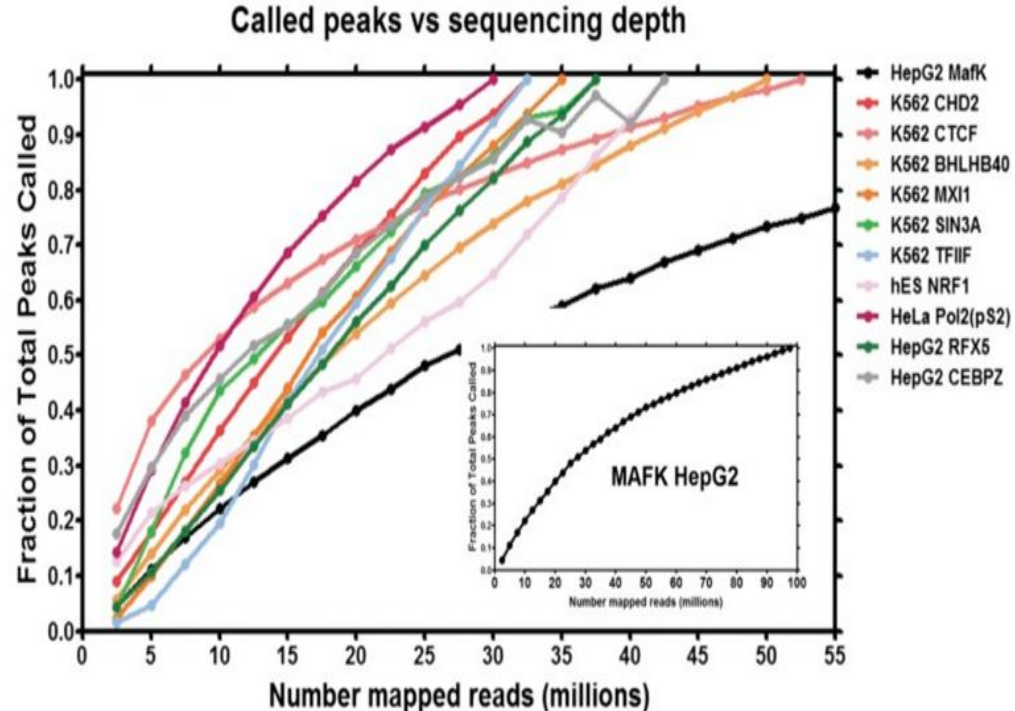
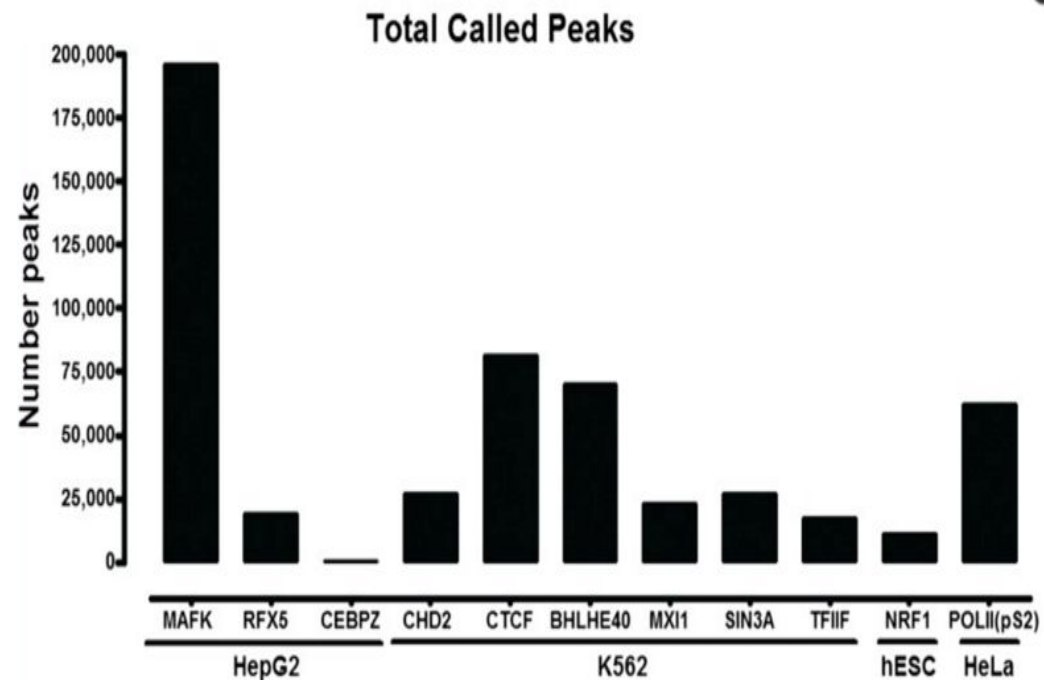
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# Sequencing depth

- The number of peaks depends on the depth of sequencing
- Some ENCODE guidelines:
  - Sharp peaks (like transcription factors):
    - Mammalian: 10M reads
    - Worms and flies: 2M reads
  - Broad peaks (some histone marks):
    - Mammalian: 20M reads
    - Worms and flies: 5M reads

<sup>[1]</sup> Landt et al, *Genome Research*, 2012.



# Duplication rate, library complexity

- Duplication rate is also a QC metric:

- expected to be low (<1%) for inputs

$$\frac{\text{DuplicateReads}}{\text{TotalMappedReads}} \times 100$$

- Duplicates can be artefacts:

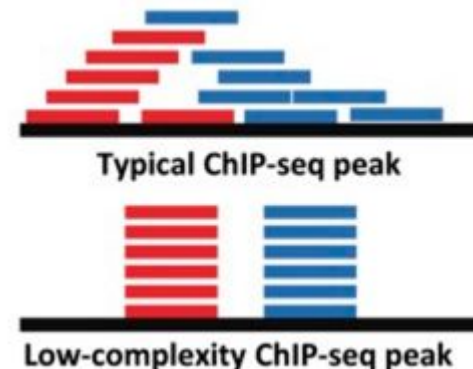
- PCR bias: certain genomic regions are preferentially amplified
- Low initial starting material can introduce artificially enriched regions with overamplification

- Duplicates can also be “legitimate”:

- It is unavoidable in highly enriched experiments and deeply sequenced ChIPs since it is naturally increasing with the sequencing depth

- Removing duplicates limits the dynamic range of ChIP signal:

- Maximum signal/base:  $(2^{\text{readLength}}) - 1$



# Duplication rate, library complexity

- What to do with duplicates?
- Always keep in mind enrichment efficiency and read depth
- Some approaches:
  - Don't remove duplicates as long as it has a reasonable rate
  - Remove duplicates before peak-calling
  - Keep duplicates for differential binding analysis
  - htSeqTools:
    - Estimate duplicate numbers expected taking into account the sequencing depth and using negative binomial model
    - Attempt to identify significantly outstanding duplicate numbers

# Control/input samples

- The use of some kind of a control is always recommended
- You need different controls for:
  - Different cell lines, cell types
  - Different organisms
  - Different treatments/conditions
- Types of controls:
  - Input DNA:
    - Most popularly used
    - Controls for CNVs, sequencing -, fragmentation - and shearing biases
  - IgG:
    - Also controls for non-specific binding
    - Introduces other biases

# Acknowledgement

- Ines de Santiago

[https://github.com/bioinformatics-core-shared-training/ngs-in-bioc/blob/master/Day3/Lect7.ChIP\\_QC\\_presentation.pdf](https://github.com/bioinformatics-core-shared-training/ngs-in-bioc/blob/master/Day3/Lect7.ChIP_QC_presentation.pdf)

- Tom Carroll

[http://bioconductor.org/help/course-materials/2014/BioC2014/ChIPQC\\_Presentation.pdf](http://bioconductor.org/help/course-materials/2014/BioC2014/ChIPQC_Presentation.pdf)

[https://github.com/bioinformatics-core-shared-training/ngs-in-bioc/blob/master/Lectures/Lect6b\\_ChIP---Seq%20Data%20Analysis.pdf](https://github.com/bioinformatics-core-shared-training/ngs-in-bioc/blob/master/Lectures/Lect6b_ChIP---Seq%20Data%20Analysis.pdf)