

# Introduction to ATAC-seq

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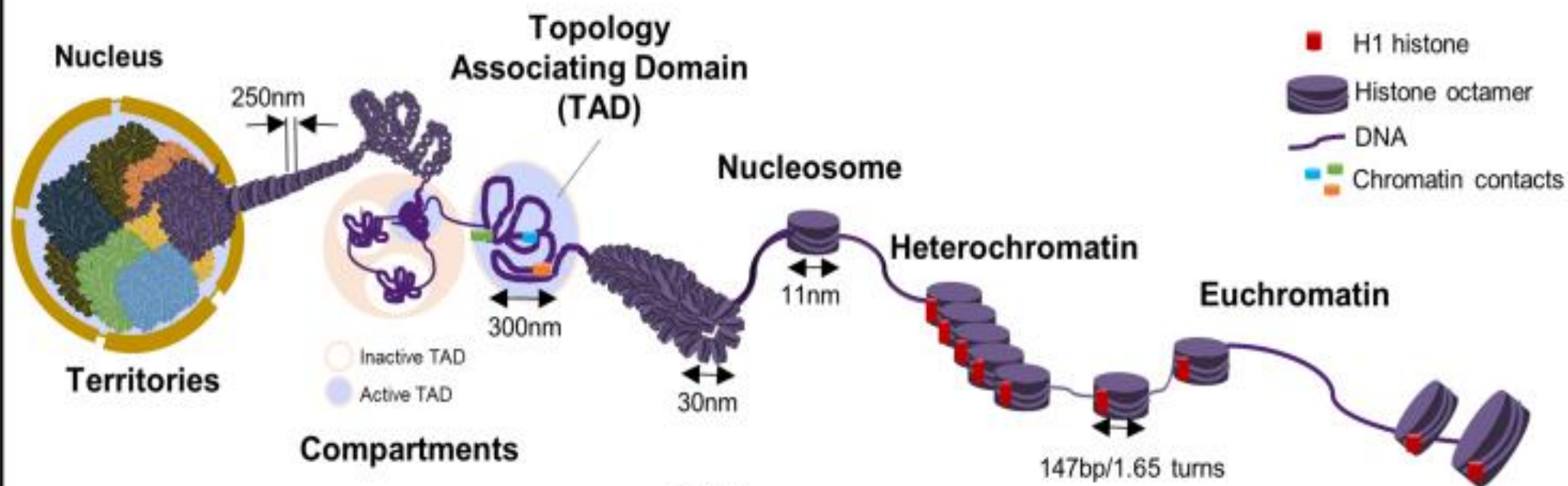
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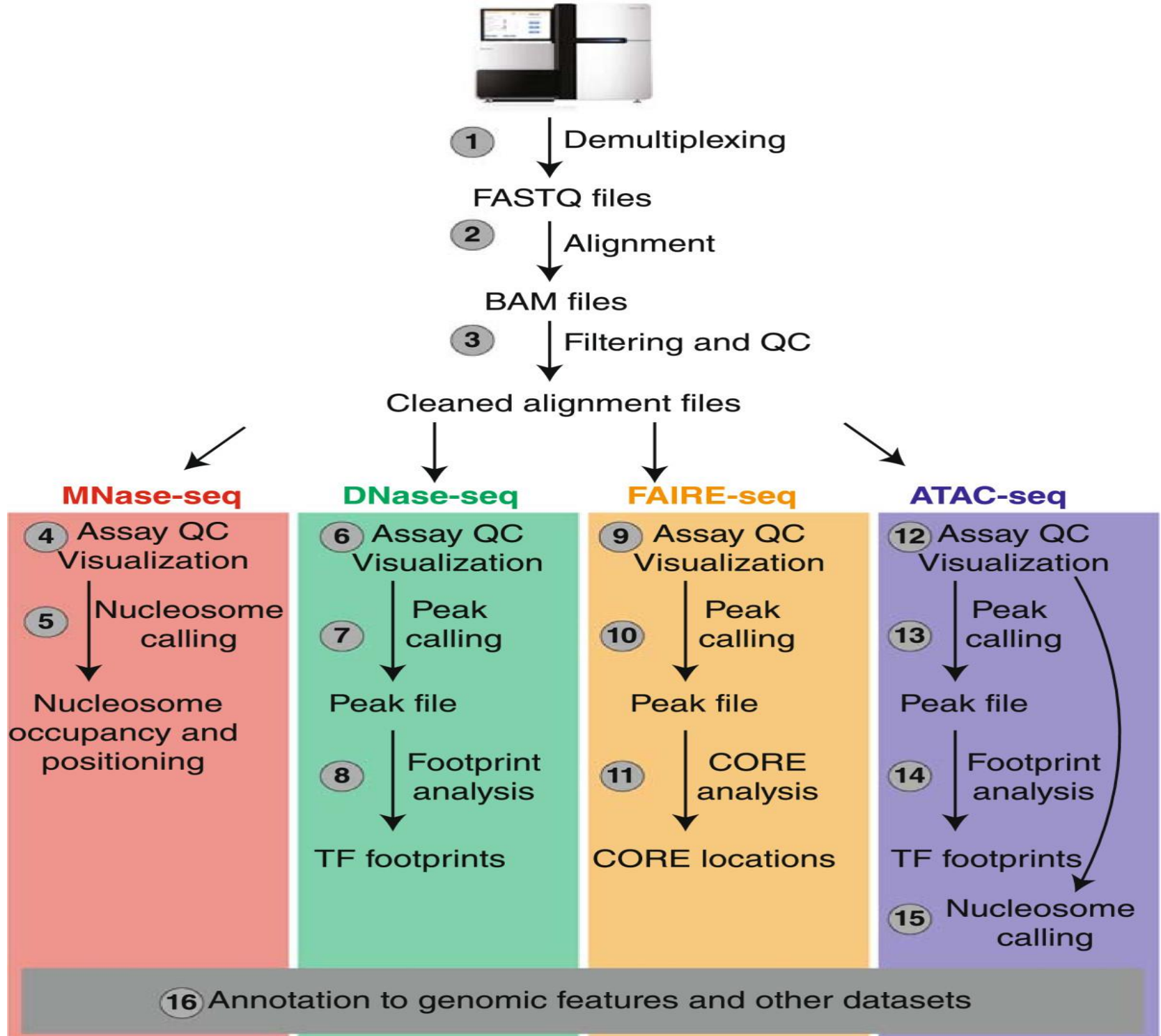
## Higher-order

## Primary-order



| Techniques | Validation |              |            |                    | MNase-seq               | DNase-seq      | FAIRE-seq      | ATAC-seq              |
|------------|------------|--------------|------------|--------------------|-------------------------|----------------|----------------|-----------------------|
|            | 4C         | 5C           | Hi-C       | ChIP, Imaging etc. |                         |                |                |                       |
|            | 1-vs-Many  | Many-vs-Many | All-vs-All |                    | Inferred closed regions | Open chromatin | Open chromatin | Open/Closed chromatin |

| Procedure | Higher-order     |                             | Primary-order     |                   |
|-----------|------------------|-----------------------------|-------------------|-------------------|
|           | Experimental     | Computational               | Experimental      | Computational     |
|           | 1. Crosslinking  | 1. Alignment                | 1. Crosslinking*  | 1. Size-selection |
|           | 2. Fragmentation | 2. Filtering                | 2. Fragmentation  | 2. Alignment      |
|           | 3. Ligation      | 3. Binning                  | 3. Size-selection | 3. Peak calling   |
|           | 4. Detection     | 4. Normalization            | 4. Sequencing     | 4. Normalization  |
|           |                  | 5. Identifying interactions |                   | 5. Visualization  |
|           |                  | 6. Visualization            |                   |                   |



Article | Published: 06 October 2013

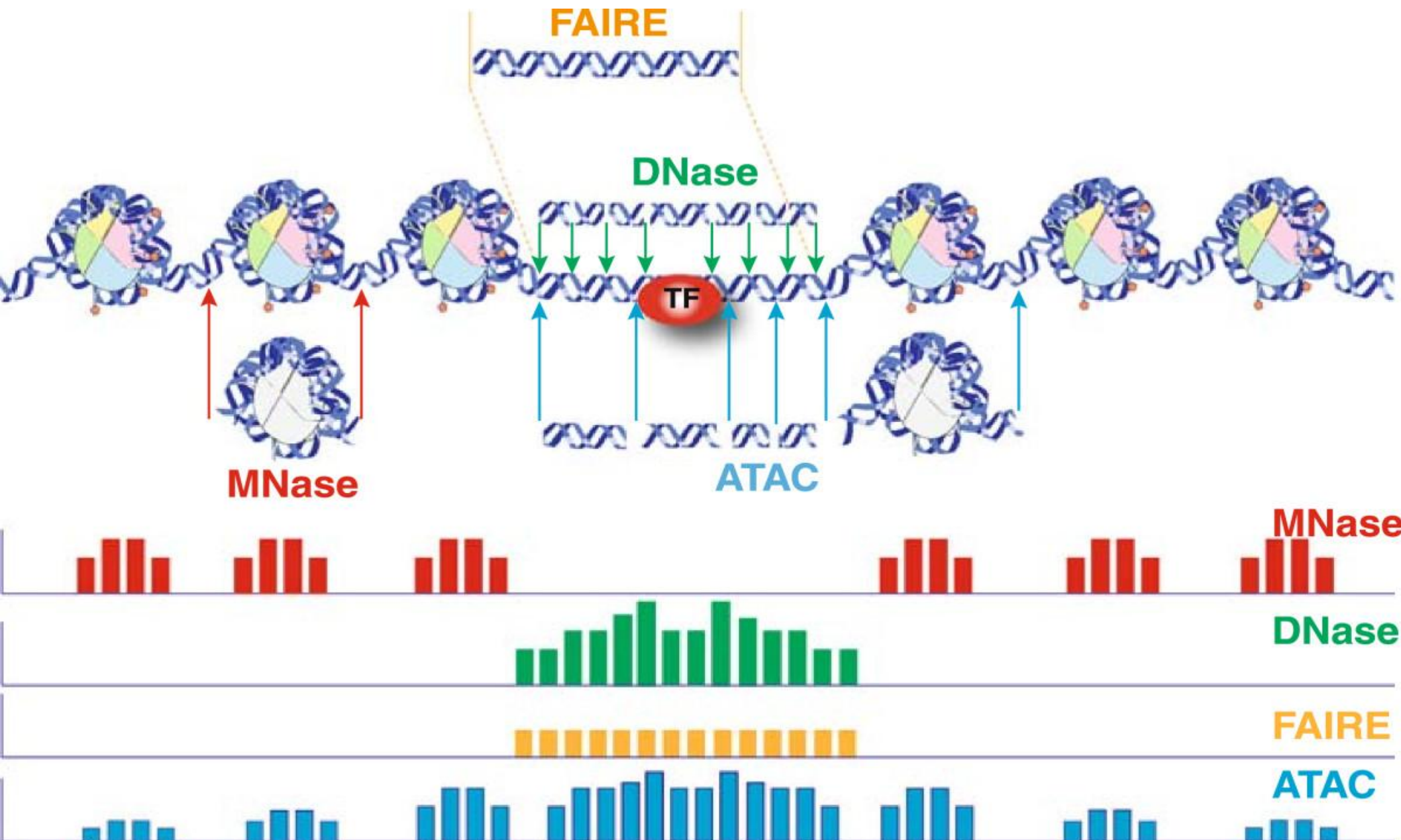
# Transposition of native chromatin for fast and sensitive epigenomic profiling of open chromatin, DNA-binding proteins and nucleosome position

Jason D Buenrostro, Paul G Giresi, Lisa C Zaba, Howard Y Chang✉ & William J Greenleaf✉

*Nature Methods* **10**, 1213–1218 (2013) | [Download Citation](#) ⚡

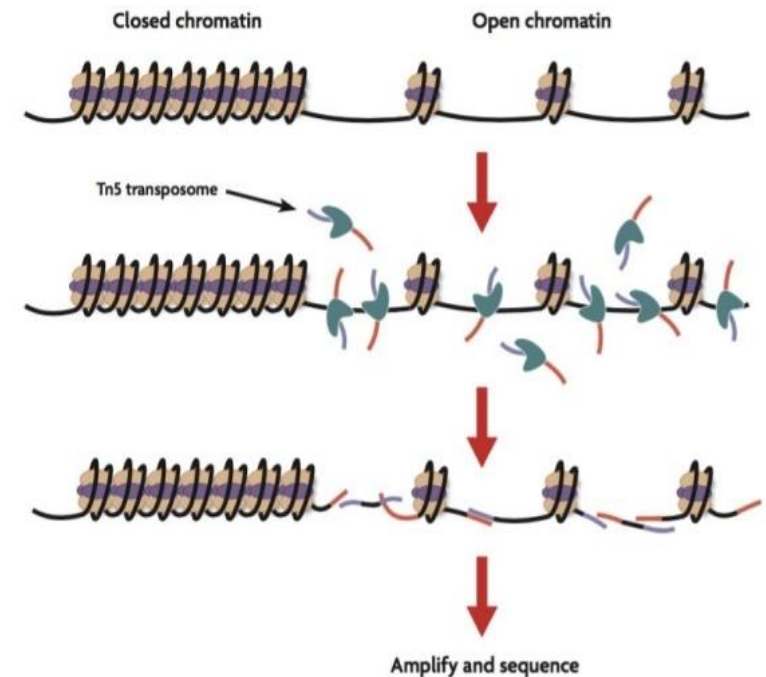


# Assay for Transposase Accessible Chromatin



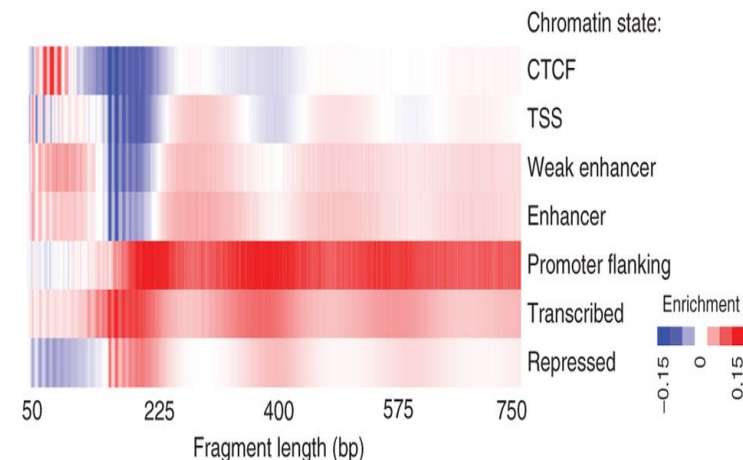
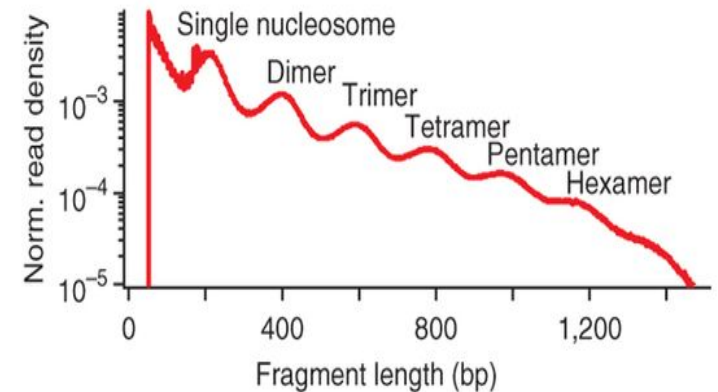
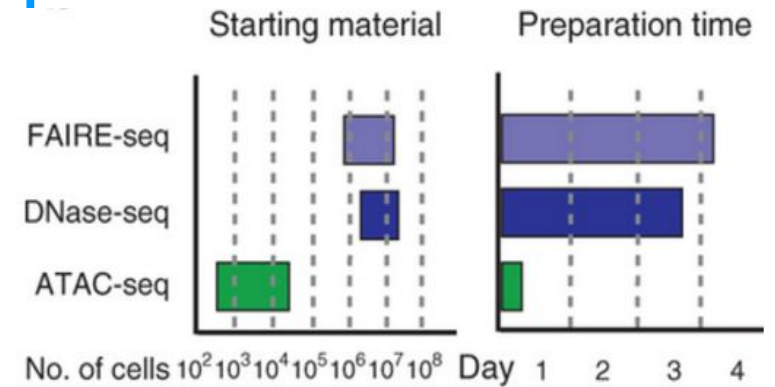
# ATAC-seq

- Measure **chromatin accessibility** and **nucleosome free regions**
- Can also detect nucleosome packing, positioning and TF footprints
- Does not require sonication and phenol chloroform extractions, antibodies or sensitive enzymatic digestions that can introduce potential bias
- A hyperactive Tn5 transposase is used to fragment DNA and integrate into active regulatory regions
- During ATAC-seq, 500–50,000 unfixed nuclei are tagged *in vitro* with sequencing adapters by Tn5 transposase



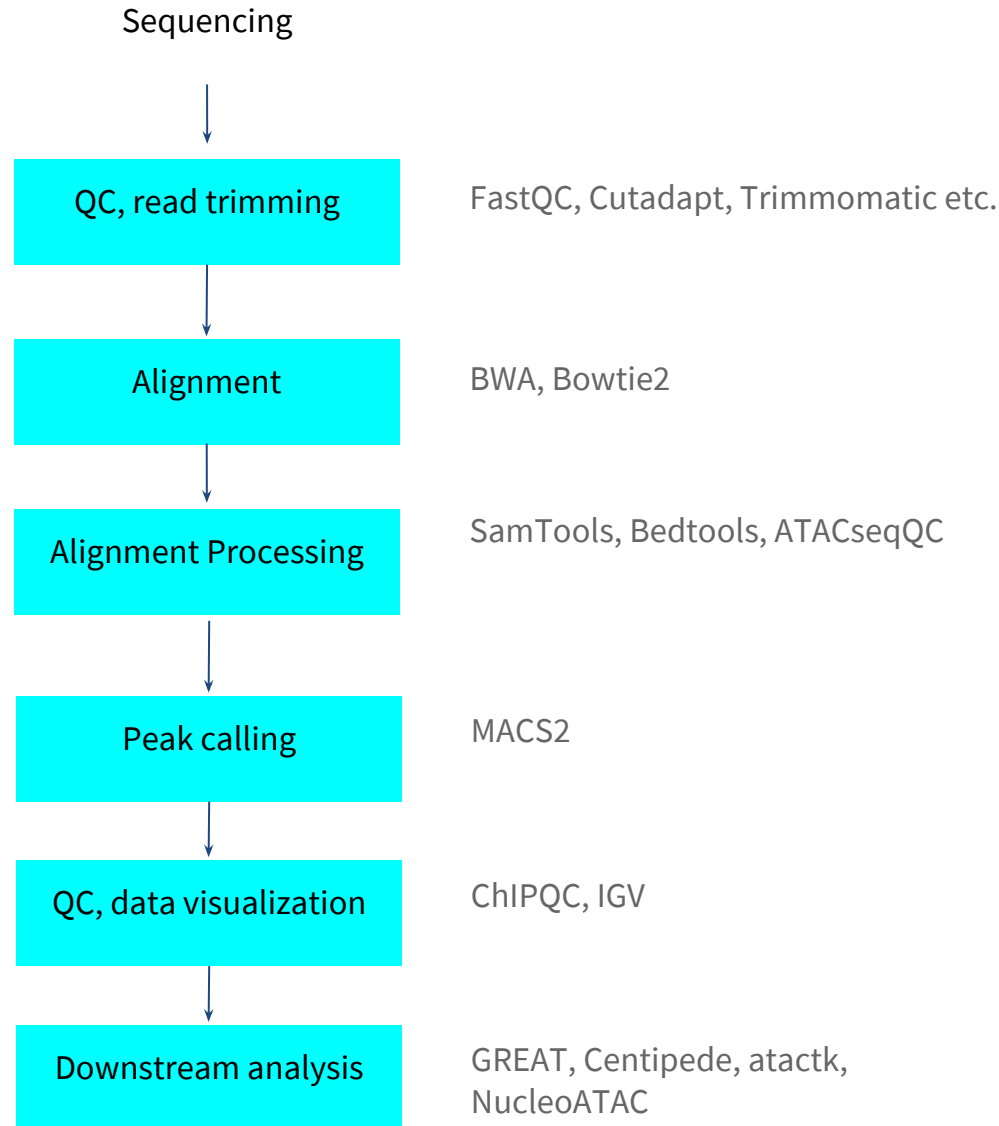
# ATAC-seq

- Two-step protocol
  - Insertion of Tn5 transposase with adaptors
  - PCR amplification
- Needs ~500-50,000 cells
- Paired-end reads produce information about nucleosome positioning
- Insert size distribution of fragments has a periodicity of ~200 bp, suggesting that fragments are protected by multiples of nucleosomes
- Different fragmentation patterns can be associated with different functional states (eg. TSSs are more accessible than promoter flanking or transcribed regions)



[1] JD Buenrostro et al, *Nature Methods*, 2013.

# Workflow of ATAC-seq data processing





# Working with ATAC-seq

- Remove mitochondrial reads
  - A large fraction of ATAC-seq reads map to the mitochondrial genome (up to 40-60%)
- Remove blacklisted regions before peak calling
  - hg19 or hg38 blacklisted regions can be obtained from ENCODE
- The signal (open chromatin where the transposase was inserted) is a mixture of feature types:
  - Short fragments - signal from nucleosome free regions (**NFRs**) and open regions around DNA bound transcription factors. TSS rich.
  - Longer fragments - open regions from around nucleosomes. Includes +1 and -1 nucleosome positions

# Differences from ChIP-seq data processing

- Peak calling: Use fragment size for smoothing when calling peaks with MACS2
- Alternatively, to identify open regions following MACS parameters are used:  
**MACS2 callpeak -t bamfile --nomodel --shift -100 --extsize 200 --format BAM -g hg38**

For nucleosome occupancy shift and extension can centre the signal on nucleosomes (147 bp DNA is wrapped in a nucleosome)

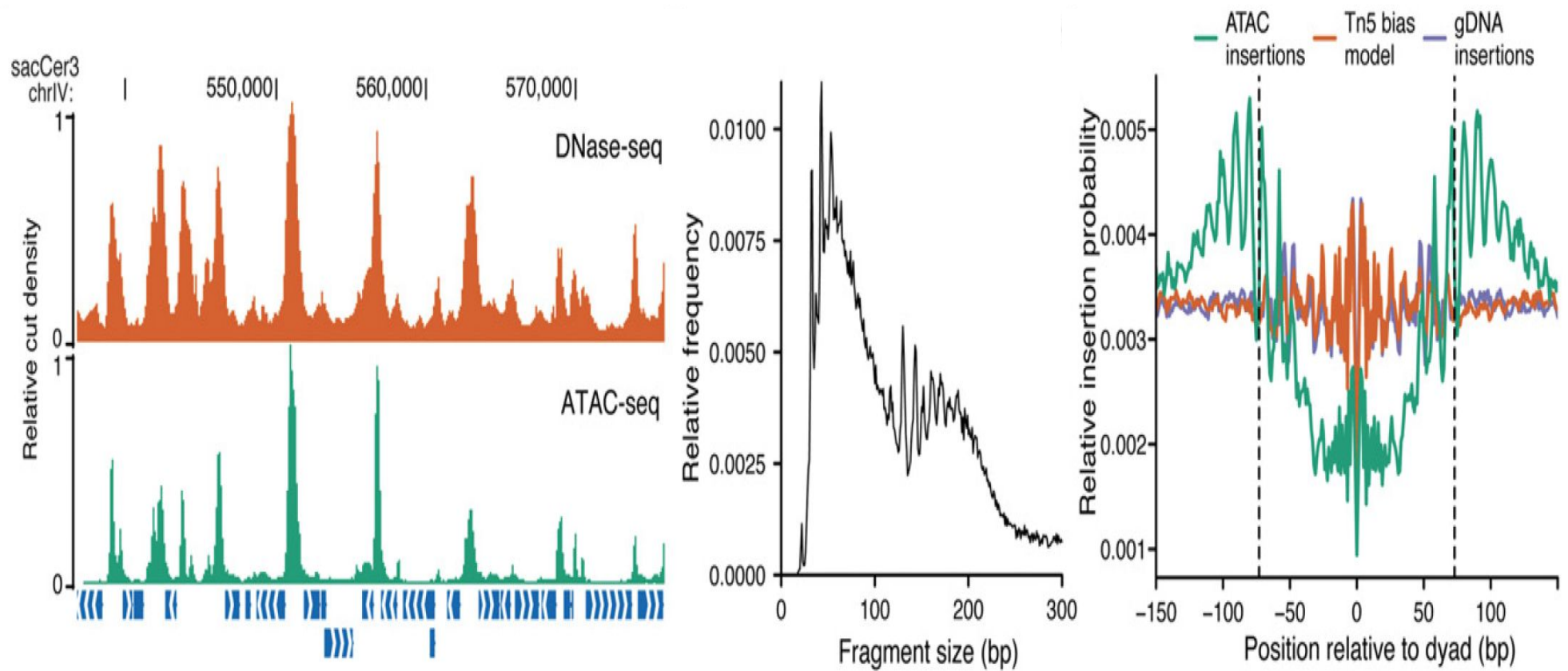
**MACS2 callpeak -t bamfile --nomodel --shift 37 --extsize 73 --format BAM -g hg38**

Optionally, for paired end data

**MACS2 callpeak -t bamfile --format BAMPE -g hg38**

- --nomodel: don't build shifting model
- --shift: when this value is negative, ends will be moved toward 3'→5' direction
- --extsize: extend reads in 5'→3' direction to fix-sized fragments

# ATAC-seq signal



# Normalization and Differential Accessibility of ATAC-seq

- Normalisation across samples might be needed
  - Efficiency of the ATAC-seq protocol in assaying open regions is affected by how many transposons get into the nuclei
  - One normalisation solution is to use signal from ‘essential or housekeeping genes’. see: *Sarah K. Denny et al, Cell, 2016.*
- **THOR** is an HMM-based approach to detect and analyze differential peaks in two sets of ChIP-seq data from distinct biological conditions with replicates
  - Normalizes bam files (given a bed file of housekeeping genes)
  - Can compare two ATAC-seq datasets to perform differential openness analysis

# Cutting site and TF Footprinting analysis

- ATAC-seq produces shorter fragments around smaller protected regions such as TF bound regions.
- Cut sites pileup signal is a good guide to TF binding sites
- Shifting reads +4/-5 nt depending on strand, should adjust for expected shift from Tn5 insertion.
- TF footprinting of these regions enables the detection of motifs for bound TFs
  - Centipede
  - msCentipede
  - PIQ
  - Mocap