# ATAC-seq analysis

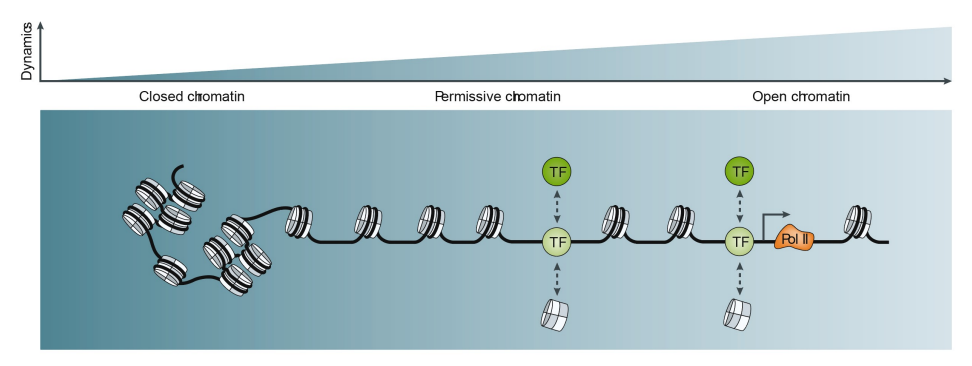
Epigenomics Data Analysis
Jacques Serizay
Physalia 2023



# Beyond textbook statements: a spectrum of regulatory capacity

Chromatin accessibility continuum that ranges from closed chromatin to highly dynamic, accessible or permissive chromatin

This landscape of chromatin accessibility reflects the spectrum of regulatory capacity — rather than a bistate organization

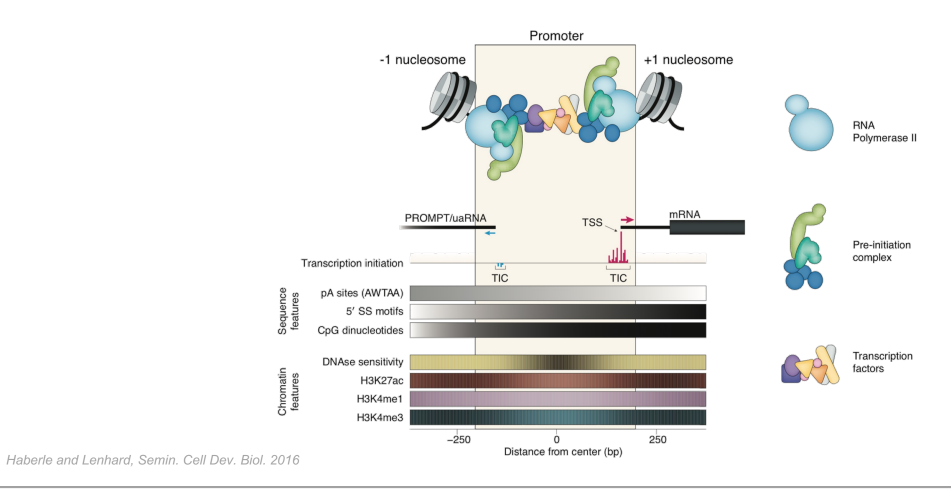


Klemm et al., Nat .Rev. Genet 2019



# **Promoter organization**

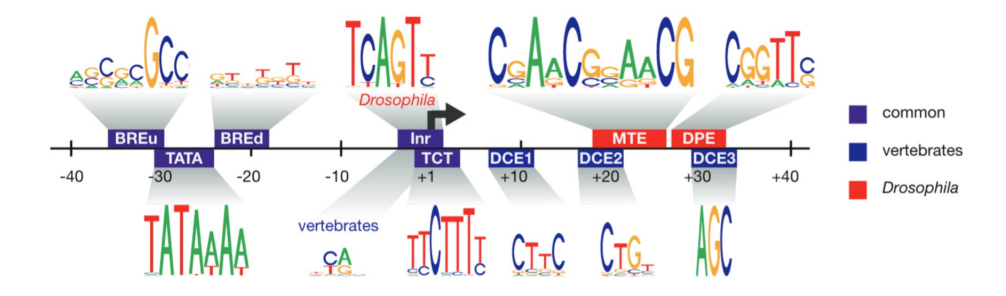
#### Promoters are crowded environments!





## **Promoter organization**

Transcription machinery and general transcription factors need access to DNA to recognize their binding motif



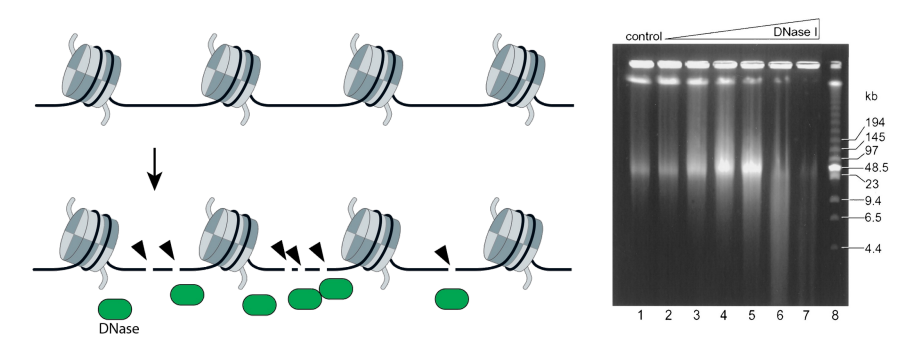
Haberle and Lenhard, Semin. Cell Dev. Biol. 2016



# How to measure chromatin accessibility: originally with nucleases

Nuclease enzymes were historically used to profile chromatin accessibility

o Dnase I (deoxyribonuclease I): a non-specific double-strand endonuclease



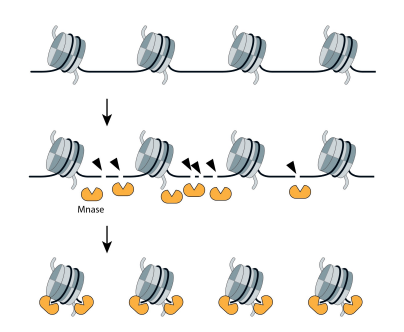


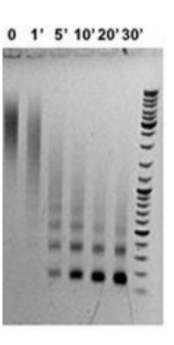


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- Mnase (Micrococcal nuclease): a endo-exonuclease with preference to single-strand DNA but with also double-strand nuclease activity





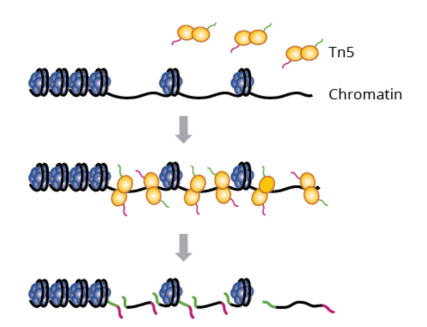
Rodríguez-Campos & Azorín, PLoS One 2007



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- o ATAC: a Tn5 transposase, integrating transposons wherever it is possible (i.e. accessible)



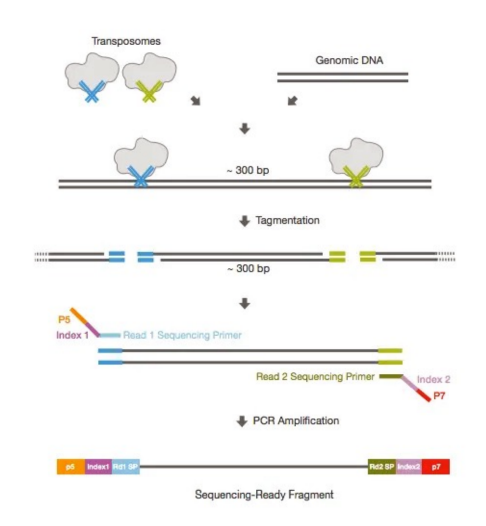
Buenrostro et al., 2013



## ATAC-seq: from chromatin to NGS library

Since the sequence of the <u>transposons</u> loaded on the Tn5 <u>transposome</u> is known, one can use them to in a PCR

- → "Tagmented" (i.e. DNA with inserted transposons) will be amplified.
- → Each end of a fragment corresponds to a transposition event.



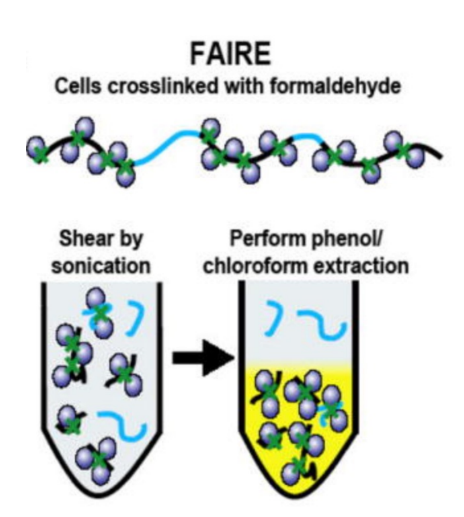


## **Emergence of other enzymatic or mechanical approaches**

#### Mechanical approaches

 FAIRE-seq: Formaldehyde-Assisted Isolation of Regulatory Elements

→ Uses a crosslinking + sonication + phenol extraction to isolate nucleosome-depleted chromatin

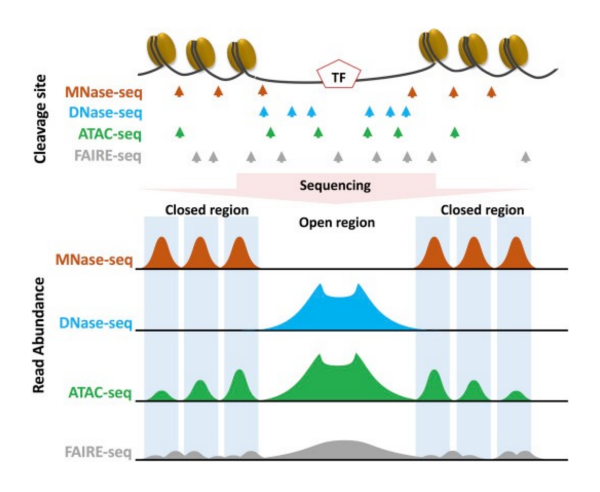


Giresi & Lieb, Methods 2009



# Comparison of the main experimental approaches

Each assay generates a specific type of profile.



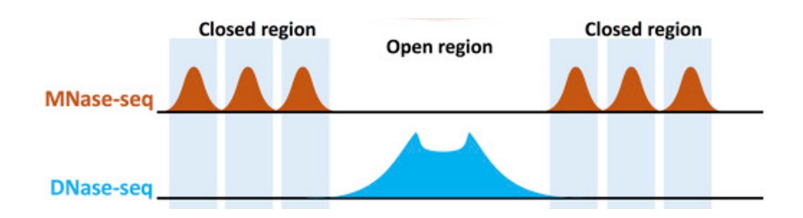
Hsu et al., Epigenetics in Human Diseases 2018



## Positive vs. negative measurements

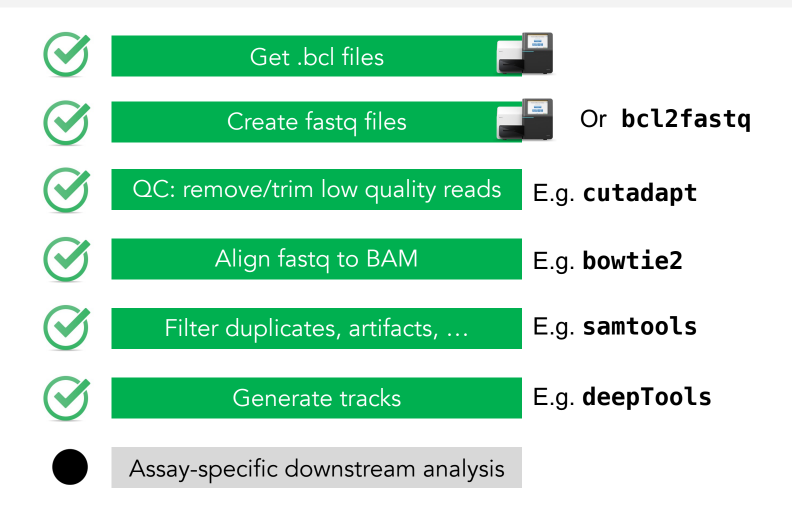
DNAse-seq relies on presence of signal (**positive measurements**) to map accessible regulatory elements

Mnase-seq relies on absence of signal (**negative measurements**) to map accessible regulatory elements





### **ATAC-seq downstream analysis**

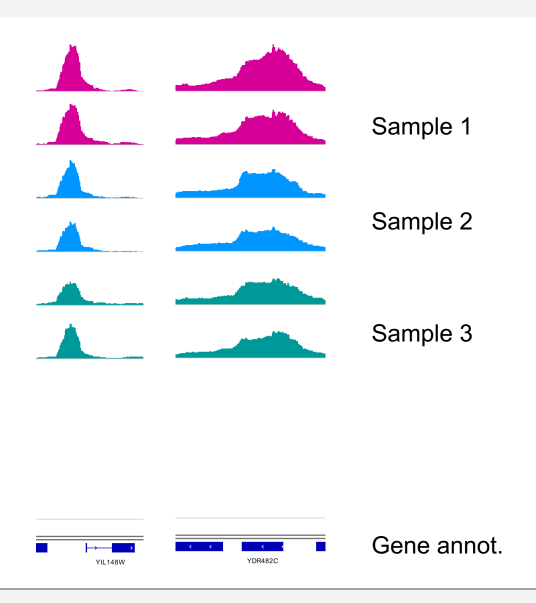




- Most peak callers were designed before the emergence of ATAC-seq
- Few of them directly aim at identifying peaks in chromatin accessibility signals

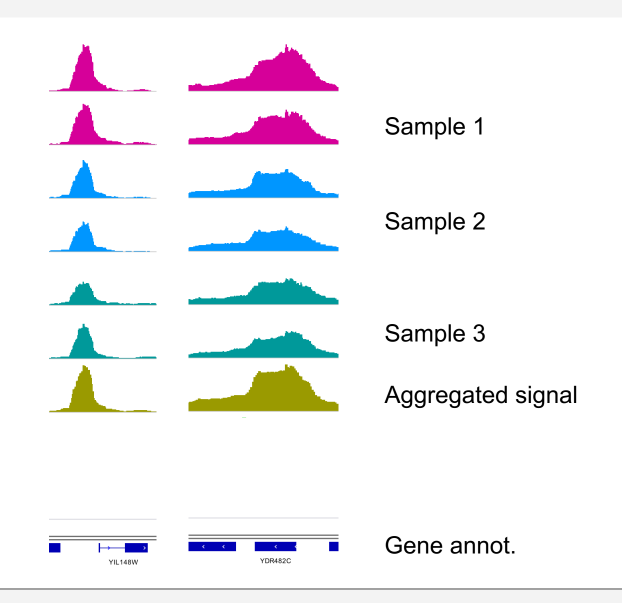


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- Since the emergence of ATAC-seq, new peak callers were designed with the identification of peak "shape" in mind.



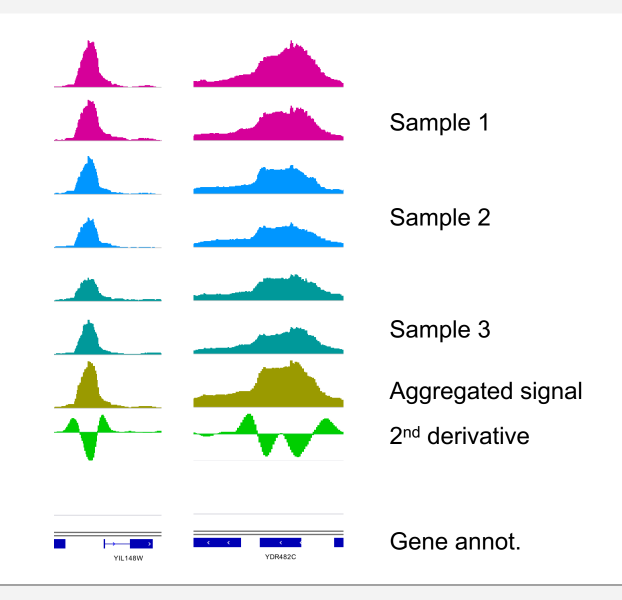


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