# Bioinformatic approaches to regulatory genomics and epigenomics

376-1347-00L | week 05

Pierre-Luc Germain



### Plan

Debriefing on the assignment

• The 'histone code' & functional elements

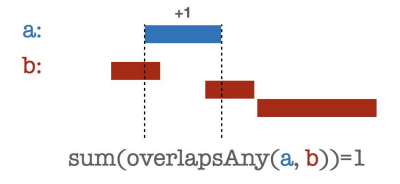
More on overlaps and comparing signals

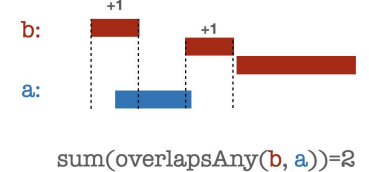
### Debriefing on the assignments

Symmetry of overlaps:

```
a: queryb: subjectoverlapsAny(a, b)
```

overlapsAny(a, b) findoverlaps(a, b)





### Debriefing on the assignments

### Symmetry of overlaps:

```
gr1 <- GRanges(seqnames=c(1), IRanges(start=c(1), end=c(10)))
gr2 <- GRanges(seqnames=c(1,1,1), IRanges(start=c(1,6,15), end=c(5,9,20)))

# Symmetry
ov1 <- overlapsAny(gr1, gr2)
sum(ov1)</pre>
```

```
## [1] 1
```

```
ov2 <- overlapsAny(gr2, gr1)
sum(ov2)</pre>
```

```
## [1] 2
```

### Debriefing on the assignments

Difference findOverlaps() vs overlapsAny():

```
# findOverlaps vs overlapsAny
gr1 \leftarrow GRanges(segnames=c(1,1,1), IRanges(start=c(1,1,15), end=c(8,9,20)))
gr2 \leftarrow GRanges(segnames=c(1,1), IRanges(start=c(1), end=c(10,9)))
ov1 <- overlapsAny(gr1, gr2)
## [1] TRUE TRUE FALSE
fo <- findOverlaps(gr1, gr2)
## Hits object with 4 hits and 0 metadata columns:
         queryHits subjectHits
         <integer> <integer>
     [1]
     [2]
    queryLength: 3 / subjectLength: 2
# If there are multiple overlapping ranges in the subject these are not the same
length(fo)
## [1] 4
sum(ov1)
## [1] 2
```

p300:

Biosample summary:

Mus musculus strain Bruce4 ES-Bruce4

H3K4me1:

Biosample summary:

Mus musculus strain 129/Ola ES-E14

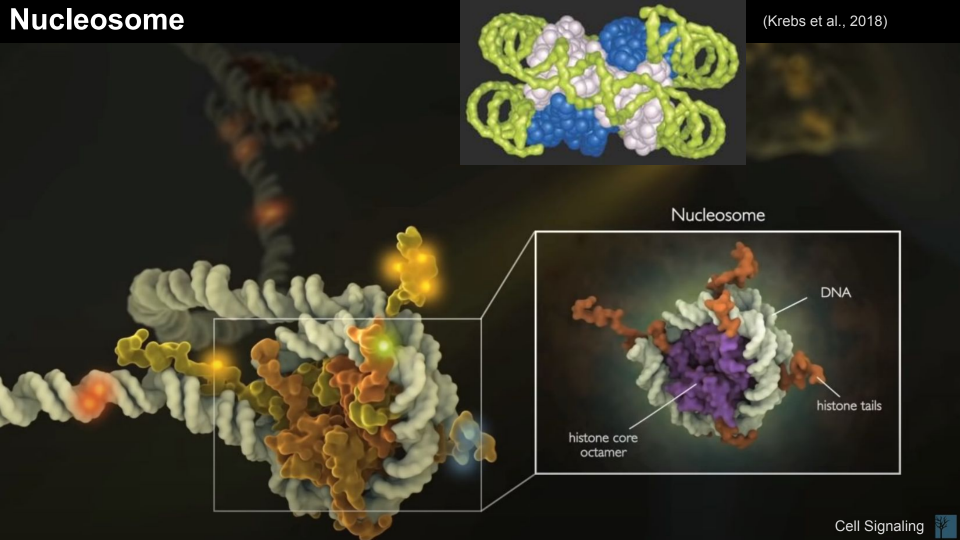
Assay:	ChIP-seq (TF ChIP-seq)		
Target:	EP300		
Biosample summary:	Mus musculus strain B6NCrl liver tissue embryo (14.5 days)		
Biosample Type:	tissue		
Replication type:	isogenic		
Description:	Chip-Seq on e14.5 liver		
		Assay:	ChIP-seq (Histone ChIP-seq)
		Target:	H3K4me3
		Biosample summary:	Mus musculus strain Bruce4 ES-Bruce4
		Biosample Type:	cell line
		Replication type:	isogenic
		Description:	H3K4me3 ChIP-seq on E0 mouse ES-Bruce4

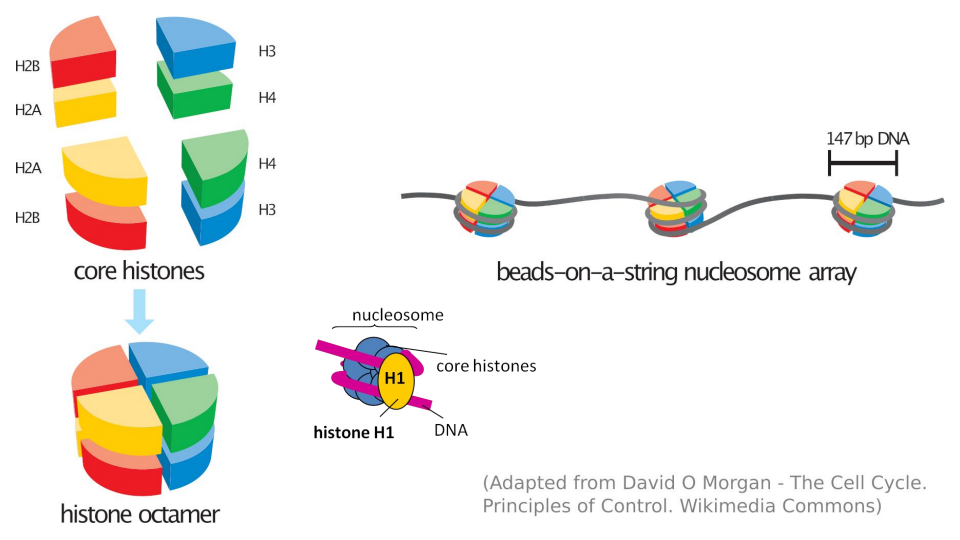
Overlapping proportion:

```
```{r, fraction overlap}
# overlap proportion gr1, gr2
gr1 <- GRanges(seqnames=c(1,1,1), IRanges(start=c(1,1,15), end=c(8,9,20)))
gr2 <- GRanges(seqnames=c(1), IRanges(start=c(1), end=c(10)))

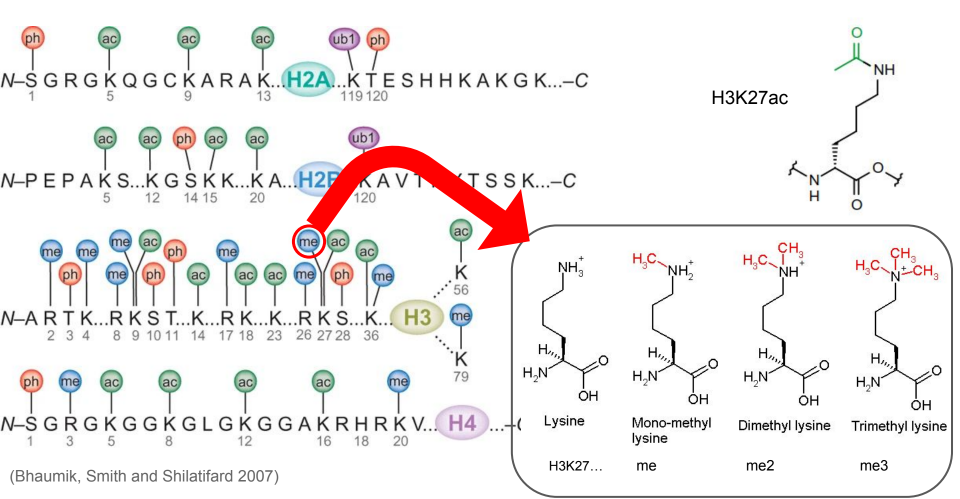
ov1 <- overlapsAny(gr1, gr2)
sum(ov1)/length(ov1)
</pre>
```

[1] 0.6666667

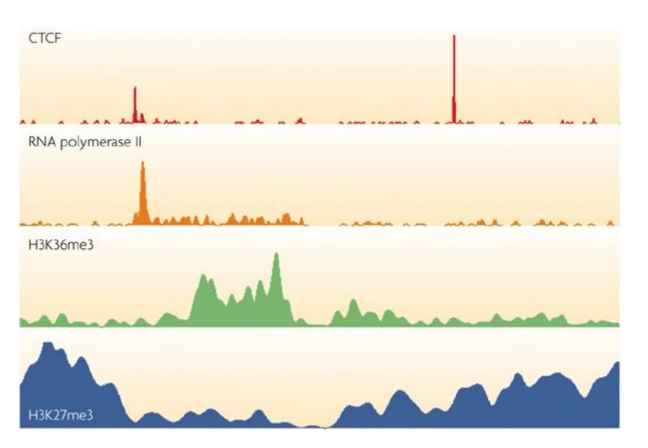




### Many residues on the histone tails can be post-translationally modified

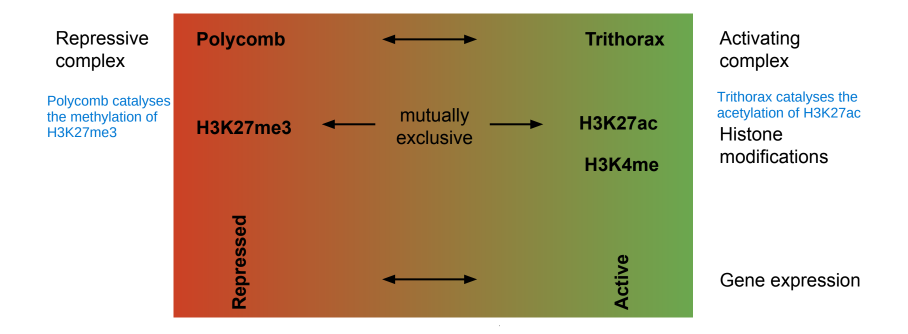


Some histone modifications appear to be very localized, e.g. happening on a specific nucleosome, while most are much more broadly distributed



The strategy of calling 'peaks' must therefore be adapted (e.g. "broad" option of most peak-callers)

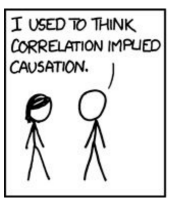
### There is a very strong association of certain histone marks and activation or repression

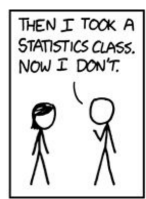


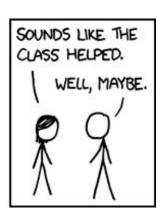
But which comes first?

### Causality or correlation?

Are histone modifications responsible for activation/repression, or are they merely associated side-effects?

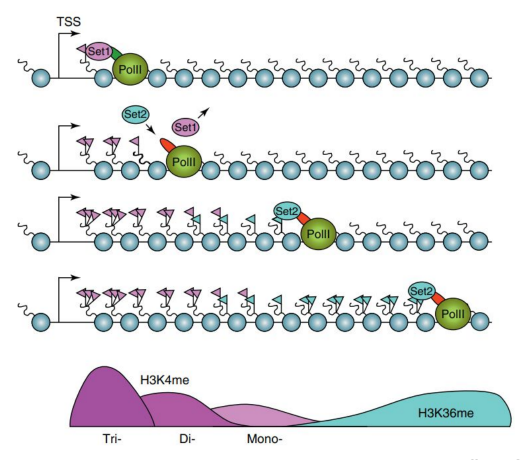




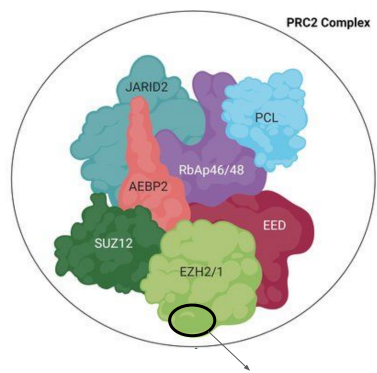


( https://xkcd.com/552 )

### Transcription-mediated histone modification



## The example of H3K27me3, chiefly deposited by the polycomb repressive complex (PRC2)



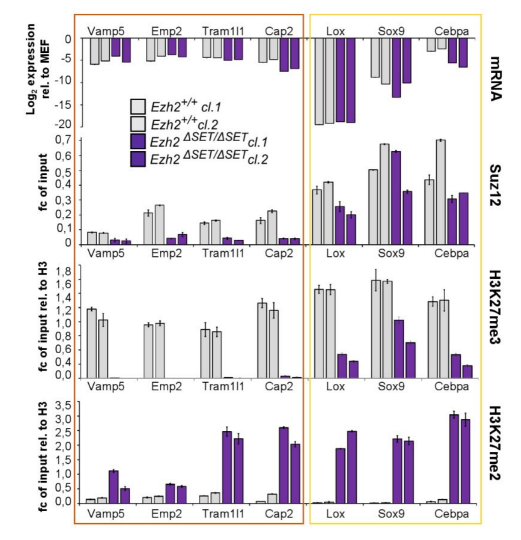
EHZ2's SET domain catalyzes the addition of a 3rd methyl group to H3K27, i.e. H3K27me2 → H3K27me3

Abolishing the enzymatic activity of *Ezh2*, the gene responsible for depositing H3K27me3, abolishes (most of) the mark but does not prevent the repression of the target genes, nor cellular reprogramming

(Fragola et al., PLoS Genetics 2013)

Similarly, the loss of H3K4me3 appears to have no effect on nascent transcription

(Murray et al., bioRxiv 2019)



### nature

Article Open Access Published: 01 March 2023

# H3K4me3 regulates RNA polymerase II promoter-proximal pause-release

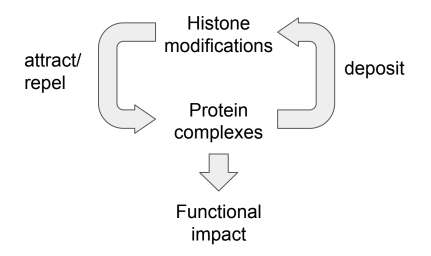
<u>Hua Wang, Zheng Fan, Pavel V. Shliaha, Matthew Miele, Ronald C. Hendrickson, Xuejun Jiang & Kristian Helin</u> ⊠

Nature 615, 339–348 (2023)

"acute loss of H3K4me3 does not have detectable effects on transcriptional initiation but leads to a widespread decrease in transcriptional output, an increase in RNA polymerase II (RNAPII) pausing and slower elongation. We show that H3K4me3 is required for the recruitment of the integrator complex subunit 11 (INTS11), which is essential for the eviction of paused RNAPII and transcriptional elongation."

### Causality or correlation?

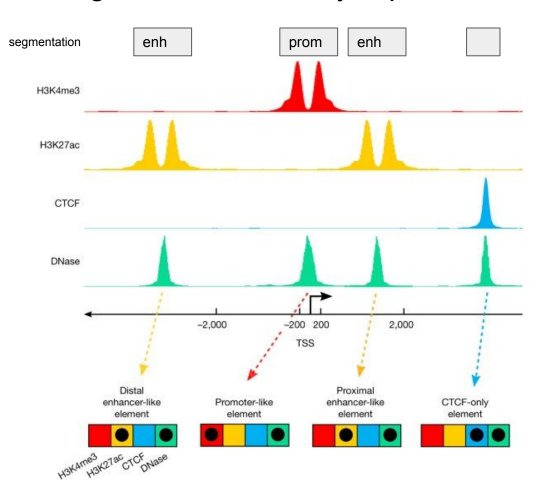
Most likely somewhere in the middle, depending on the modification/context



Whether they're causative or not, they can serve as **proxies** for function.

This means that profiling a few histone modifications gives an overview of the epigenomic landscape of a cellular state which would otherwise require profiling all the potentially-relevant factors/complexes

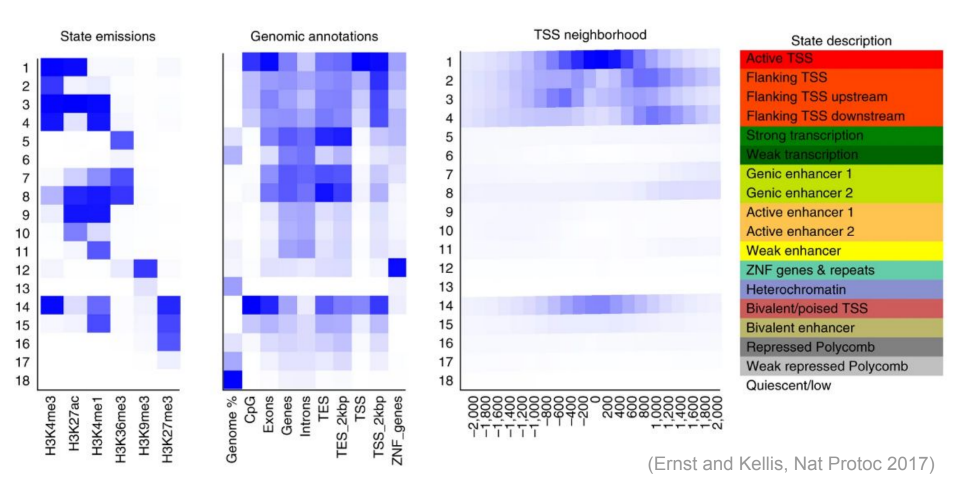
### A signature-based encyclopedia of DNA elements



### ENCODE's "signature strategy":

- Different types of functional genetic elements are associated with different chemical signatures
- We can identify functional elements by identifying these signatures genome-wide

### So how many kinds of functional elements/states are there?



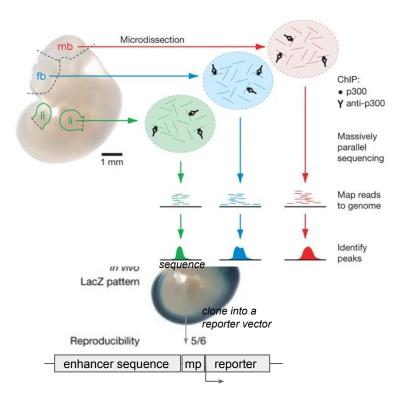
### Some stuff is pretty clear:

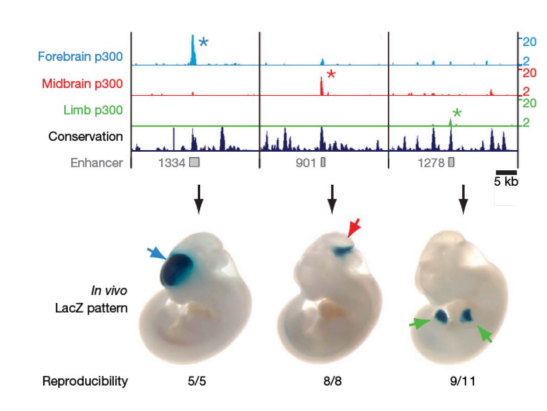
- Transcription start site (TSS):
  - H3K4me3 is almost always associated with active/poised TSS
  - Active TSS are marked by H3K27ac
  - So-called "poised" (or bivalent) TSS are instead marked by both H3K4me3 and H3K27me3

#### Enhancers:

- Most enhancers have H3K4me1
- Active enhancers are marked by H3K27ac
- So-called "poised" (or bivalent) enhancers are marked by H3K4me1 and H3K27me3
- Repressed elements are marked by H3K27me3
- Heterochromatin is marked by H3K9me3
- Insulators: CTCF+cohesin

### p300 and validation of enhancer activity





### Inconsistent seqlevels

```
overlap_H3K27ac <- overlapsAny(peaks_p300, peaks_H3K27ac)

## Warning in .Seqinfo.mergexy(x, y): The 2 combined objects have no sequence levels in common. (Use
## suppressWarnings() to suppress this warning.)

table(overlap_H3K27ac)

## overlap_H3K27ac
## FALSE
## 6394</pre>
```

### Assignment

- Using the peaks you downloaded last week, identify bivalent domains (H3K27me3 + H3K4me3) in mouse embryonic stem cells (mESC)
  - Split those bivalent domains into those that overlap a TSS, and those that don't.

- What happens to those regions upon differentiation?
  - Choose a differentiated cell type (e.g. hepatocytes, neural progenitor, or smooth muscle cells)
  - Download the H3K27me3 and H3K4me3 peaks from this cell type
  - How many of the mESC bivalent domains are, in this differentiated cell type, overlapping either mark or their combination?
    - Provide a separate answer for domains that overlap a TSS and those that don't.