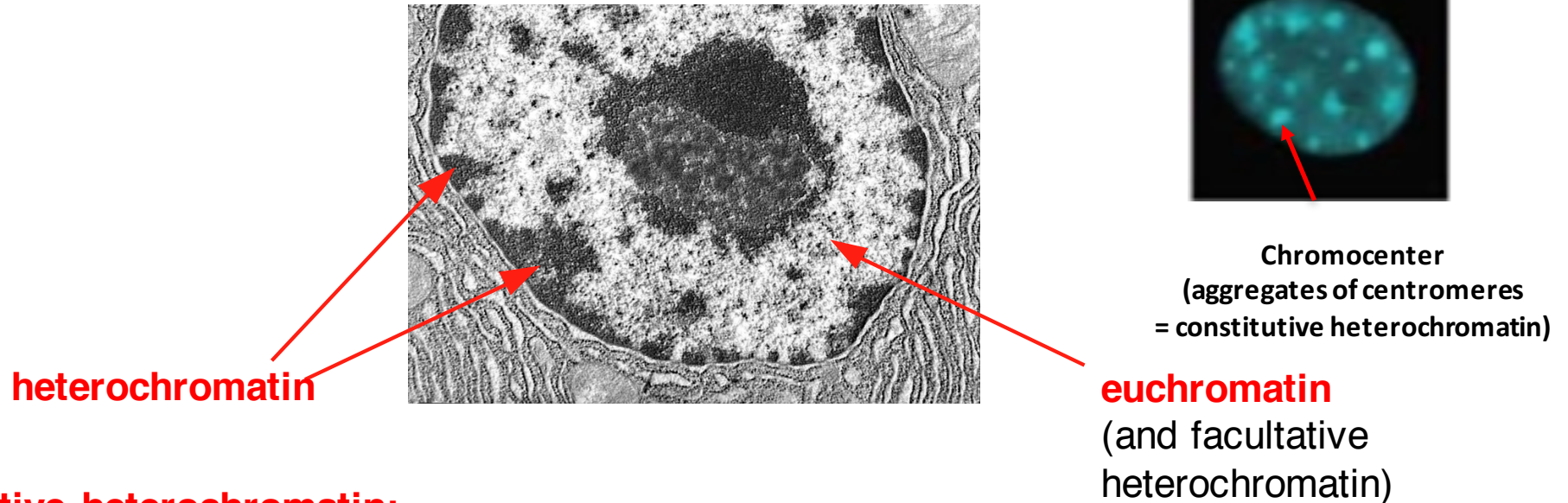


HOW TO STUDY EPIGENETIC MODIFICATIONS

**-- STRATEGIES USING SUV39H1 AS A HALLMARK
MODEL FOR EPIGENETIC REGULATION --**

Chromatin comes in different flavors

Different types of chromatin



Constitutive heterochromatin:

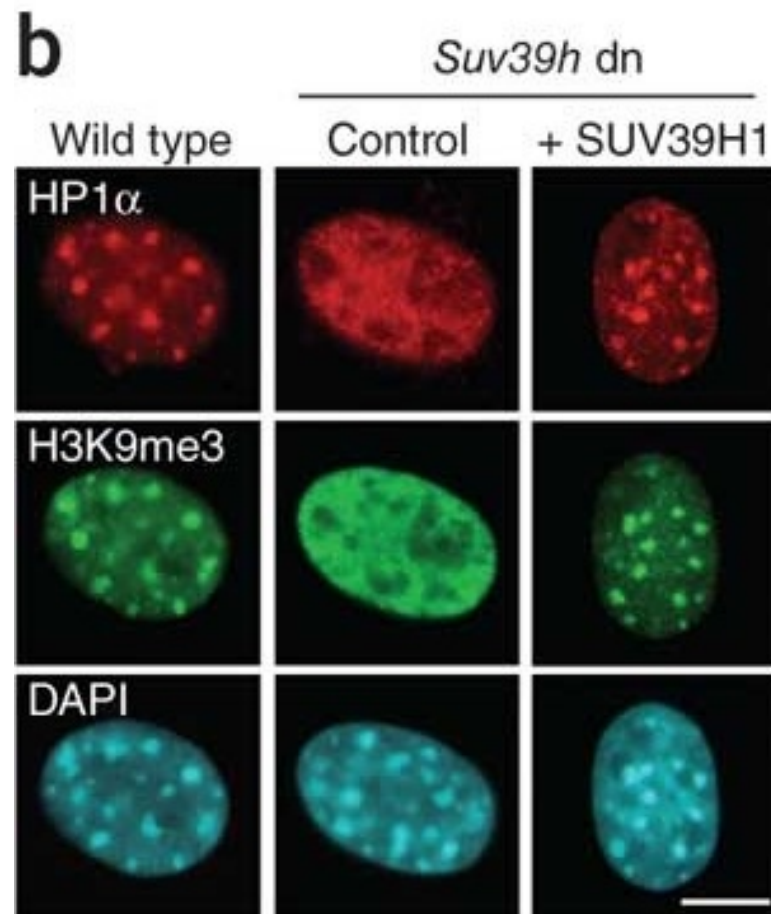
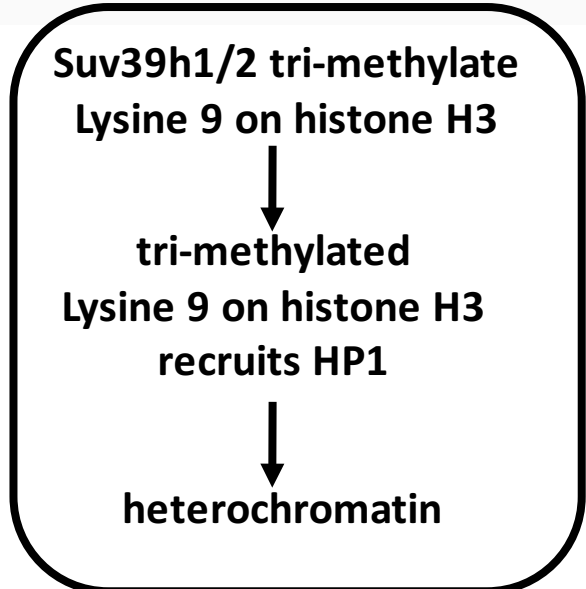
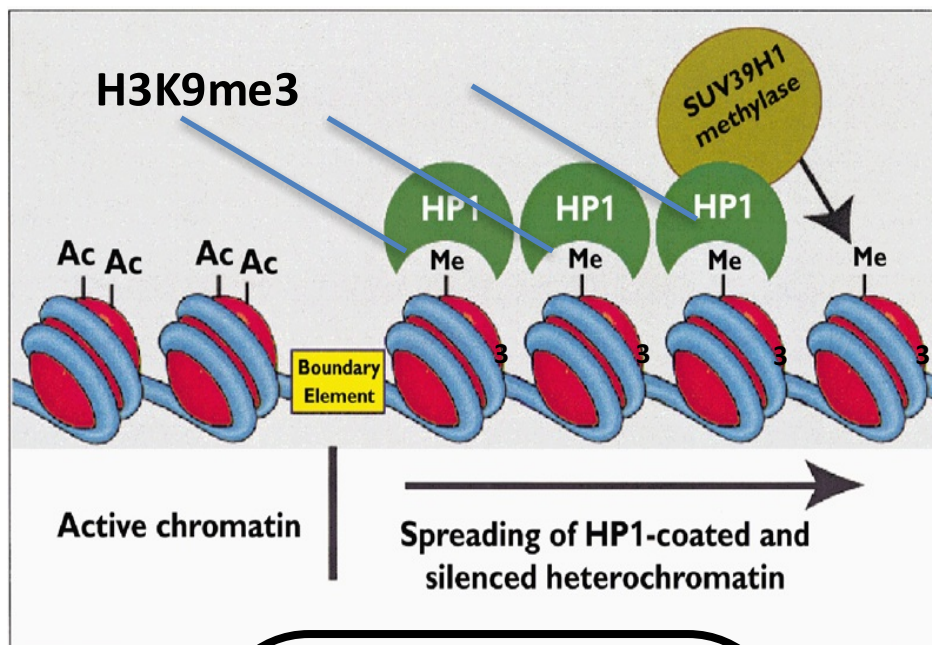
- constitute ~ 10% of nuclear DNA; telomeres, centromeres, and a considerable fraction of repetitive sequences
- highly compacted, replicates late in S phase, (transcriptionally inert)

Euchromatin + facultative heterochromatin:

- constitute ~ 90% of nuclear DNA
- less condensed, rich in genes, replicates early in S phase
- however,**
- only small fraction of euchromatin is transcriptionally active
- the rest is transcriptionally inactive/silenced (but can be activated in certain tissues or developmental stages) → these inactive regions are also known as “facultative heterochromatin”

Post-translational histone modifications can recruit specialized proteins

Example: SUV39H1 and HP1 form heterochromatin at centromeric and telomeric
Heterochromatin in flies and vertebrates and SAHFs

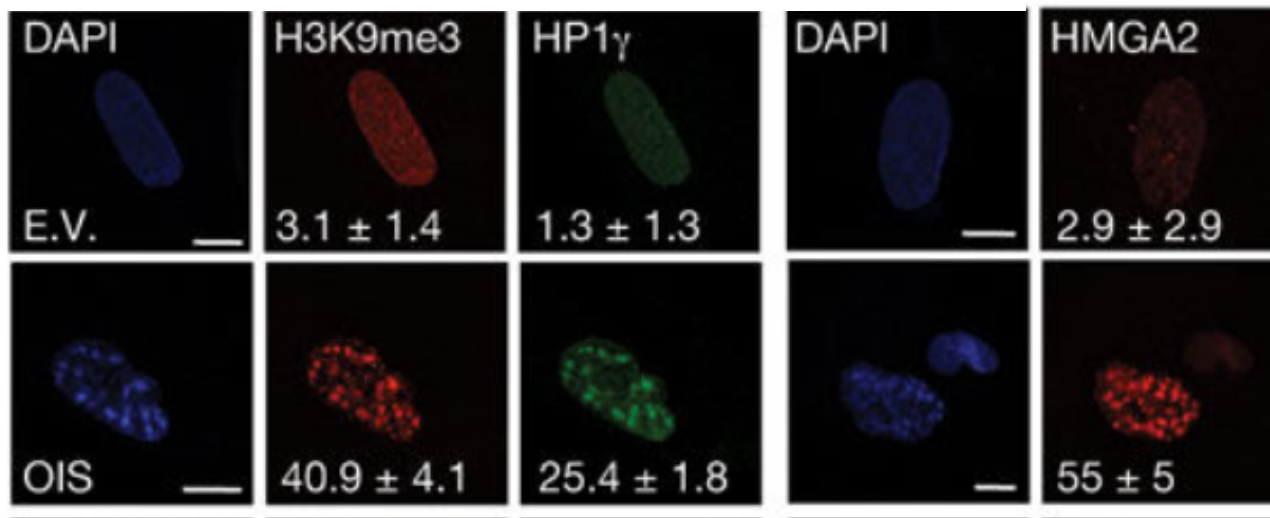


histone modifications can reach high levels in cells and can be visualized by immunofluorescence

Post-translational histone modifications can recruit specialized proteins

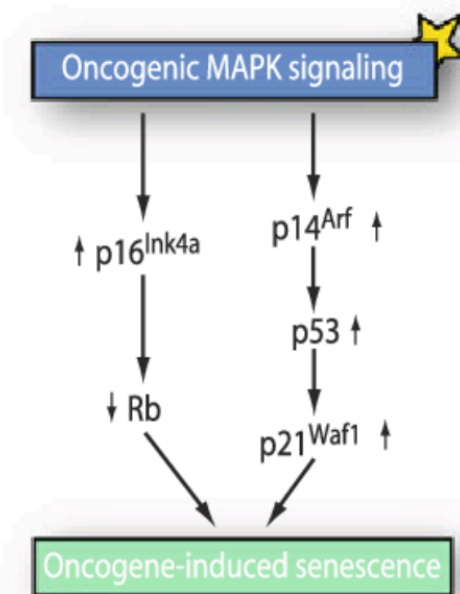
Example: SUV39H1 and HP1 form heterochromatin at centromeric and telomeric
Heterochromatin in flies and vertebrates and SAHFs

CELLULAR STIMULUS IS TRANSLATED IN TO CHROMATIN REGULATION



Senescence activated heterochromatin foci (SAHF)

OIS: Oncogene induced senescence



Oncogene induced
Replication
stress
→ DNA
damage

OIS markers:
e.g. SAHF, SA-β-Gal, p21^{Waf1}, p16^{Ink4a}

Identification of H3 Lys9 methyltransferase activity

Experiment:

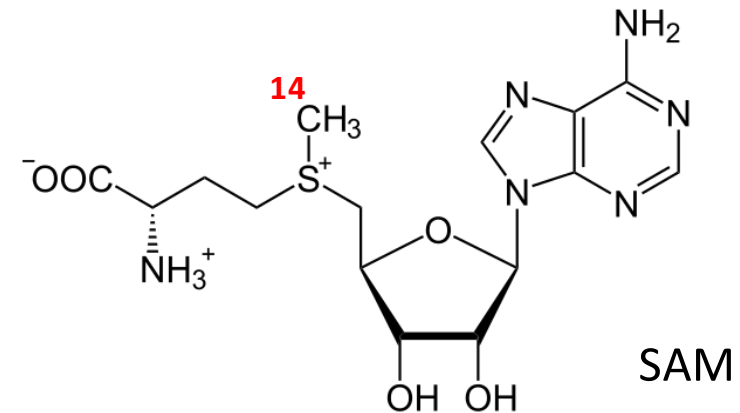
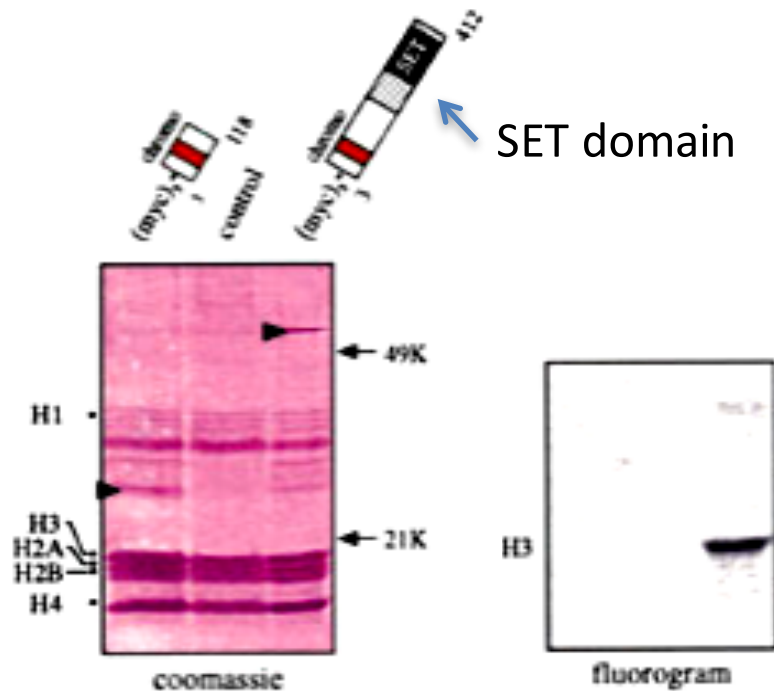
Overexpression of myc-tagged-SUV39H1 in Hela cells

Use an antibody to immunoprecipitate mycSUV39H1 → high concentration of SUV39H1

Incubate Immunoprecipitate with purified histones and S-adenosyl-[methyl-¹⁴C]-L-methionin as methyl donor



■ SUV39H1



SET – domain is required for histone methyl transferases activity

- The SET domain of the SUV39H1 is required for histone methyltransferase activity and this enzyme methylates H3 at Lys9

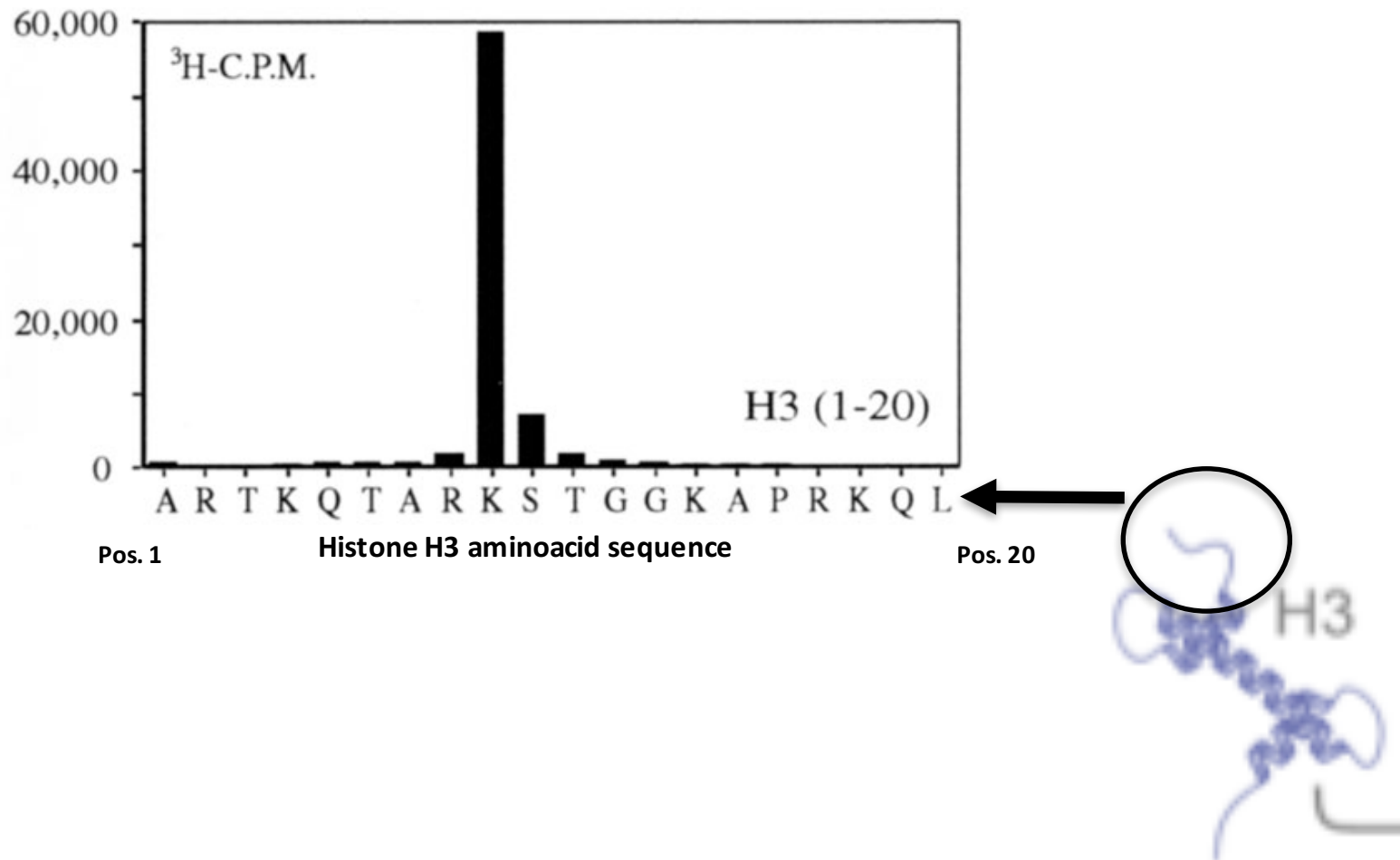
Identification of H3 Lys9 methyltransferase activity

Experiment:

Purify histone H3 by reverse-phase-liquid chromatography after HMT assay

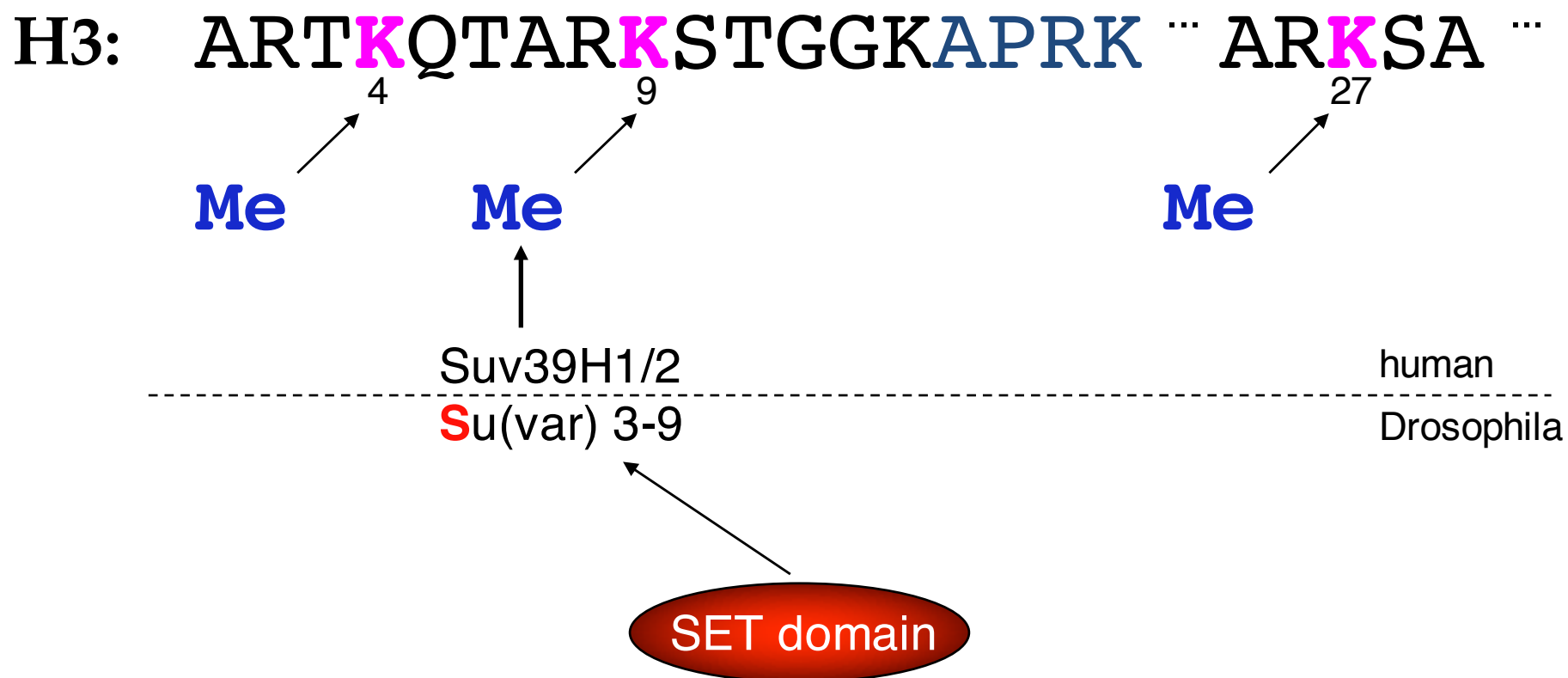
Sequence histone H3 and quantify the amount of radioactivity per amino acid

Suv39H1 methylates histone H3 at lysine 9



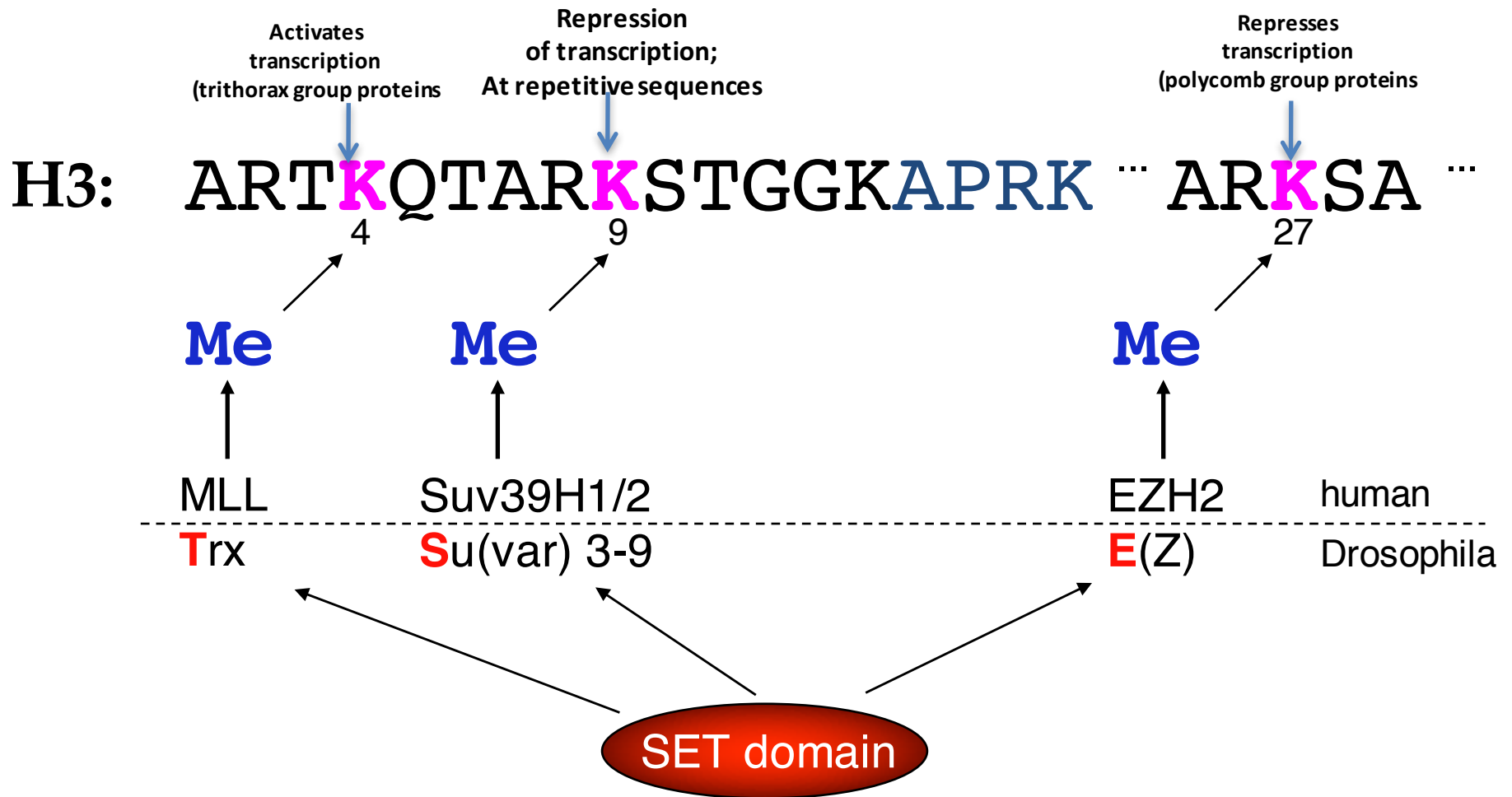
Identification of other Histone H3 Lys9 methyltransferases

- The SET domain is the conserved catalytic core of histone methyltransferases
- The histone H3 tail has 3 sites for methylation at lysines



Identification of other Histone H3 Lys9 methyltransferases

- The SET domain is the conserved catalytic core of histone methyltransferases



Enzymatic domain is conserved

SET domain protein have sequence specificity for peptide sequence around target K

- Mutations of some histone methyltransferases → cancer

Post-translational histone modifications can recruit specialized proteins

Problem:

1. How can we detect epigenetic modifications?

- Generation of antibodies that specifically recognize modified histones (western blotting/Immunofluorescence)
- Mass spectroscopy

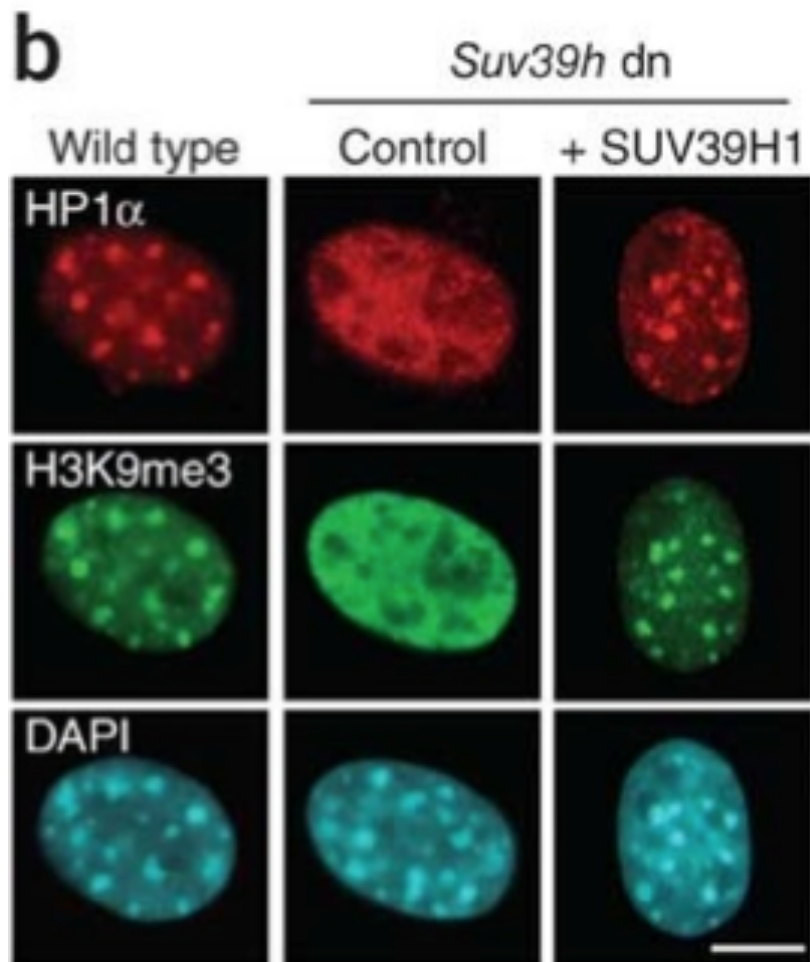
Problem:

2. How can we locate epigenetic information at defined regions or genes or promoters, etc?

- DAPI intense regions; DNA-FISH probes*

- Chromatin immunoprecipitation*

- > Detect histone modifications on specific genomic site of interest (i.e promoter of p53)
- > Detect histone modifications at multiple sites or at the entire genome level



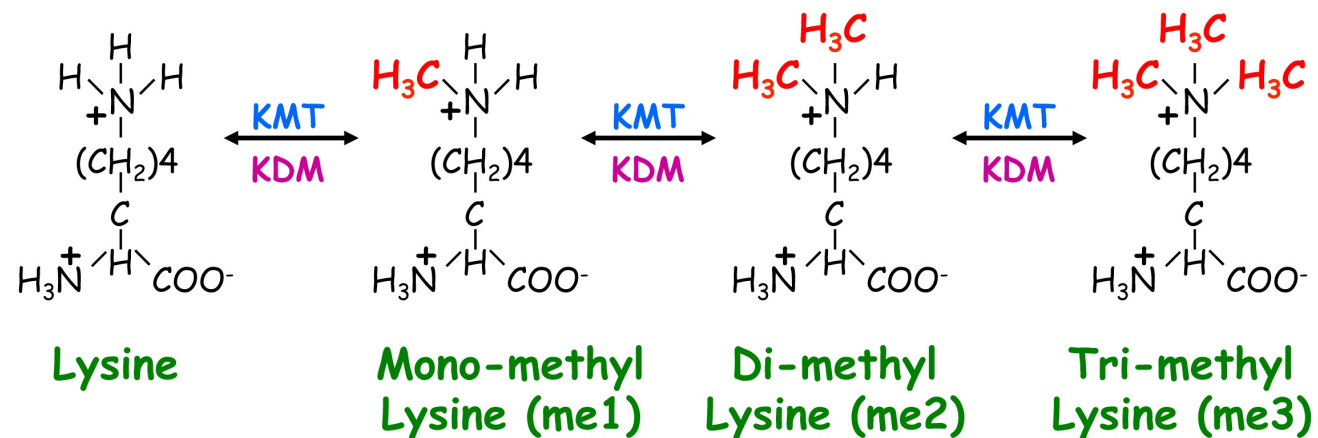
1. Generation of antibodies that specifically recognize modified histones



CAN WE GENERATE MODIFICATION SPECIFIC ANTIBODIES??

Lysines can be mono-, di- and tri-methylated
 An example: H3K9methylation by Suv39h1

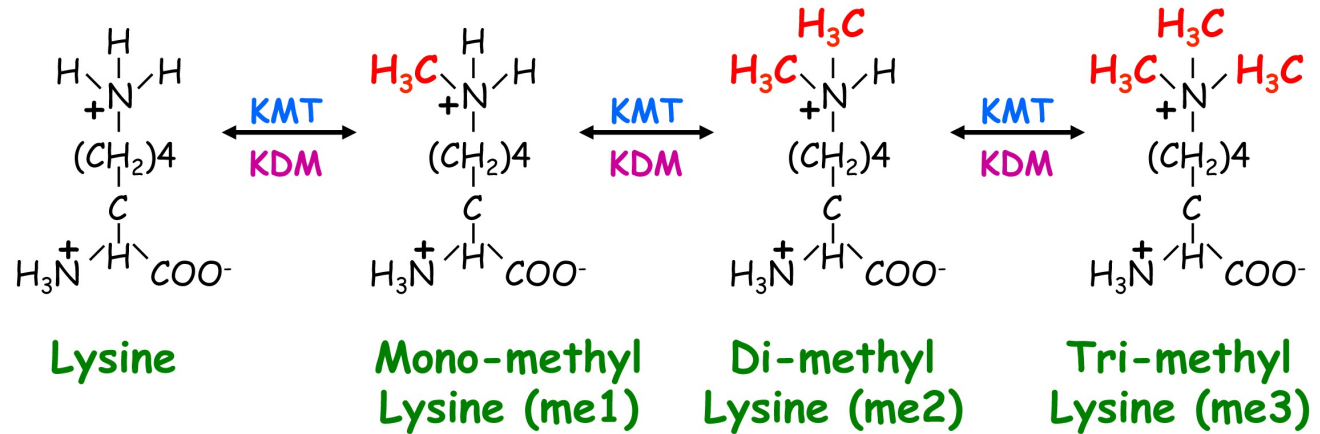
H3K9-specific Lysine-HMTases (KMTases) can mediate mono-, di-, tri-methylation



1. Generation of antibodies that specifically recognize modified histones

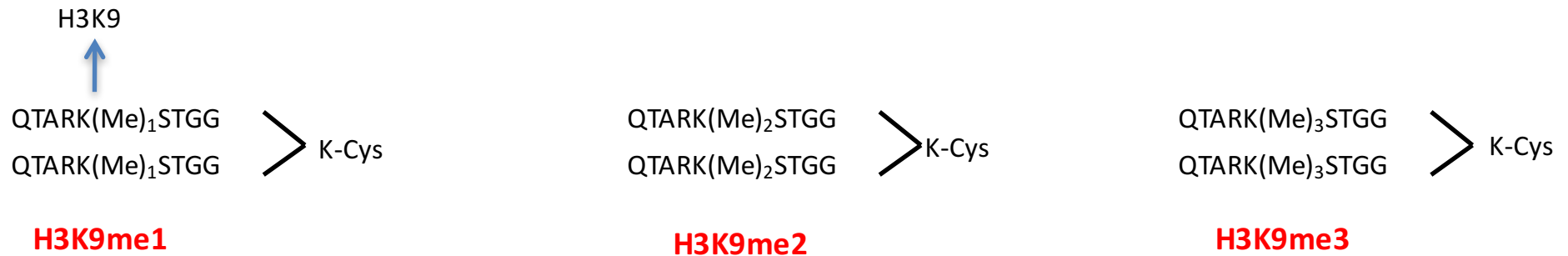
Lysines can be mono-, di- and tri-methylated

H3K9-specific
Lysine-HMTases
(KMTases) can
mediate mono-,
di-, tri-methylation



SYNTHESIS OF BRANCHED PEPTIDES FOR ANTIBODY GENERATION

H3: ARTKQTARK₉STGGKAPRK

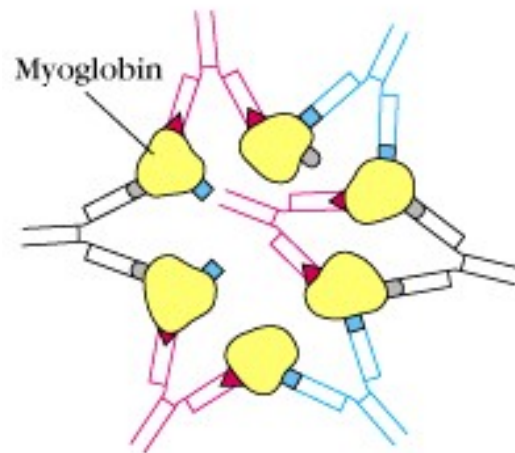


→ Branching allows to place histone modifications in close vicinity
 → → resembles high concentration of chromatin modification in the nucleus

Generation of polyclonal/monoclonal antibodies

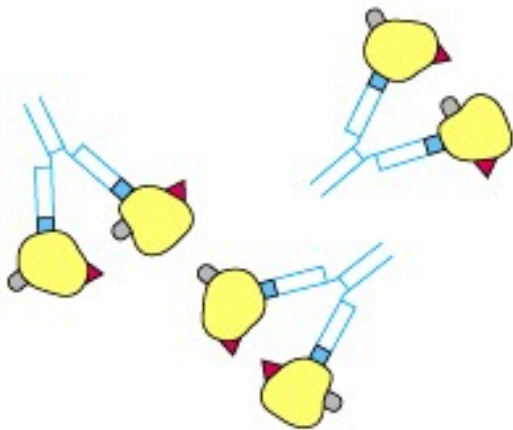
(a)

POLYCLONAL ANTISERUM



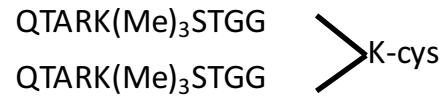
Polyclonal antibodies can form lattices with homogeneous, monomeric protein antigens because each antibody can interact with a different epitope on the antigen.

MONOCLONAL ANTIBODY



Monoclonal antibodies do not form lattices with homogeneous, monomeric proteins, because only they can bind to only one epitope on the antigen.

Generation of polyclonal antibodies



H3K9me3

1 Antigen synthesis



Non-immunogenic
small molecule

Immunogenic
protein carrier

Antigenic
conjugate

2 12-week antibody production



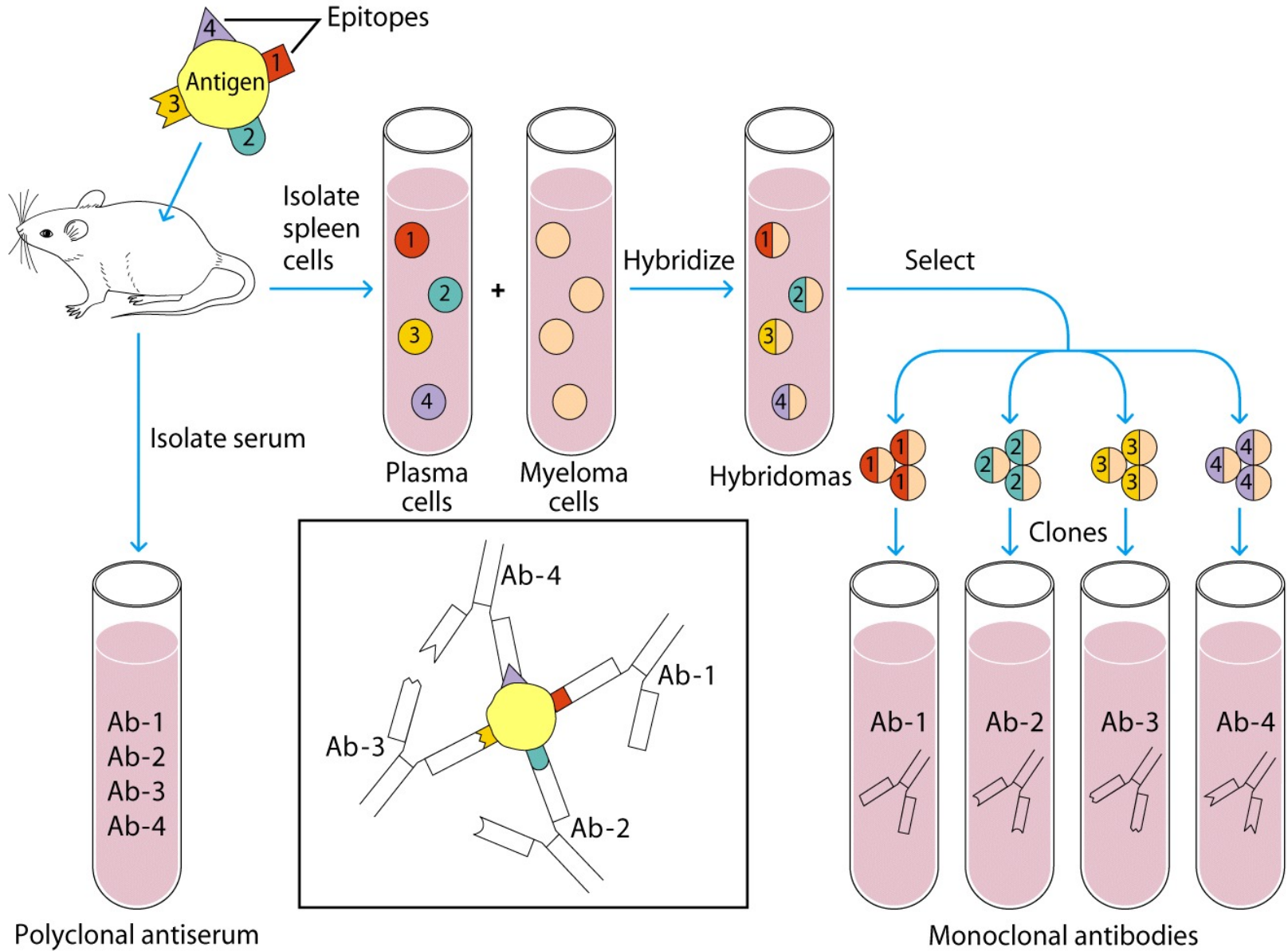
2 rabbits, 4 immunizations, 2 bleeds per rabbit

3 Polyclonal antibody characterization & purification



ELISA evaluation (titer, affinity, specificity)
& ammonium sulfate purification

Generation of monoclonal antibodies



Generation of monoclonal antibodies

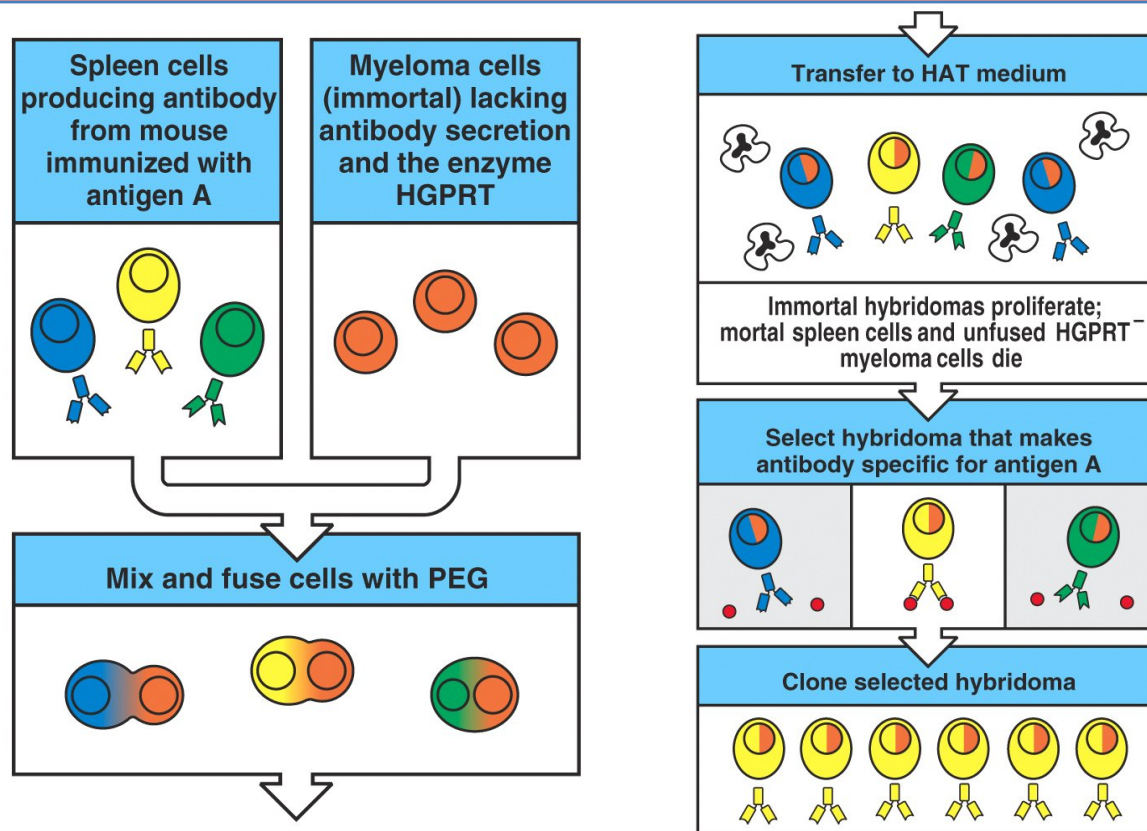


Figure A-14 part 1 of 2 Immunobiology, 6/e. (© Garland Science 2005)

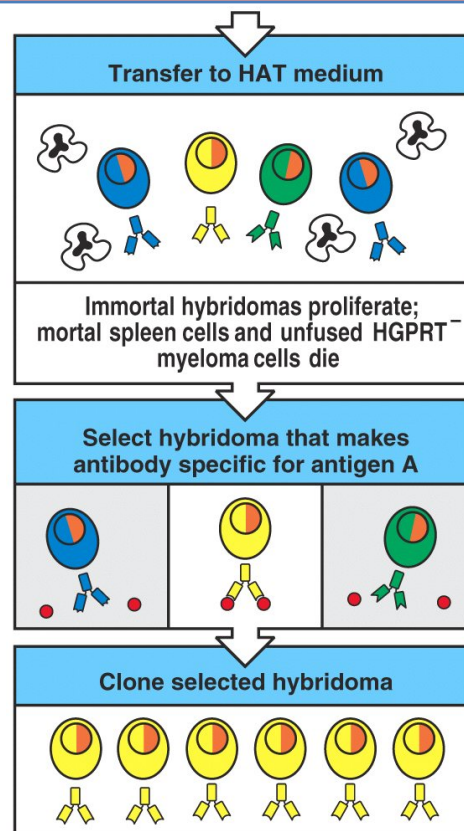
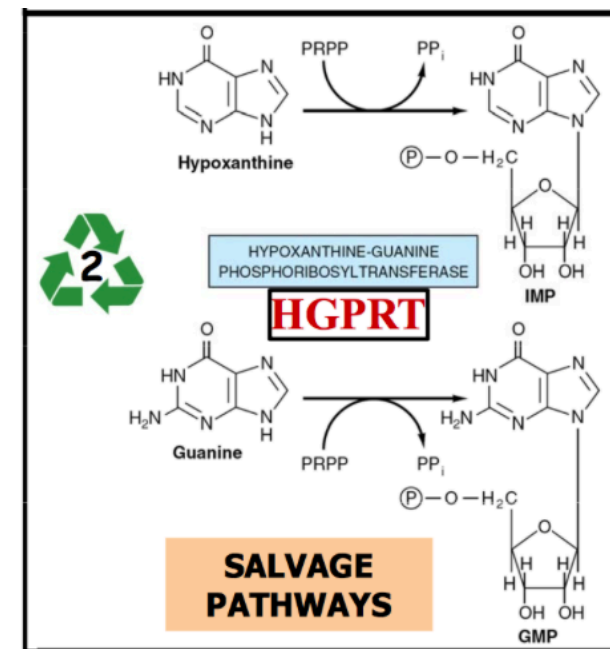


Figure A-14 part 2 of 2 Immunobiology, 6/e. (© Garland Science 2005)



Most purines are recycled rather than degraded. HGPRT can recycle hypoxanthine and guanine to be used in DNA replication

HAT Medium (hypoxanthine-aminopterin-thymidine medium) is a selection medium for mammalian cell culture, which relies on the combination of **aminopterin**, a drug that acts as a powerful folate metabolism inhibitor by inhibiting dihydrofolate reductase, with **hypoxanthine** (a purine derivative) and **thymidine** (a deoxynucleoside) which are intermediates in DNA synthesis. Its binding affinity for dihydrofolate reductase effectively blocks tetrahydrofolate synthesis. This results in the **depletion of nucleotide precursors** and inhibition of DNA, RNA, and protein synthesis. **The trick is that aminopterin blocks DNA de novo synthesis, which is absolutely required for cell division to proceed, but hypoxanthine and thymidine provide cells with the raw material to evade the blockage (the "salvage pathway"), provided that they have the HGPRT gene → rescue of DNA replication**

HAT medium is often used for preparation of monoclonal antibodies. This process is called Hybridoma technology. Laboratory animals (e.g., mice) are first exposed to an antigen against which we are interested in isolating an antibody. Once splenocytes are isolated from the mammal, the B cells are fused with HGPRT negative, immortalized myeloma cells using polyethylene glycol or the Sendai virus. Fused cells are incubated in the HAT medium. Aminopterin in the medium blocks the de novo pathway. Hence, unfused myeloma cells die, as they cannot produce nucleotides by de novo or salvage pathway. Unfused B cells die as they have a short lifespan. In this way, only the B cell-myeloma hybrids survive. These cells produce antibodies (a property of B cells) and are immortal (a property of myeloma cells). The incubated medium is then diluted into multiwell plates to such an extent that each well contains only 1 cell. Then the supernatant in each well can be checked for desired antibody. Since the antibodies in a well are produced by the same B cell, they will be directed towards the same epitope, and are known as monoclonal antibodies.

Generation of monoclonal antibodies

Cloning hybridomas from fusion event

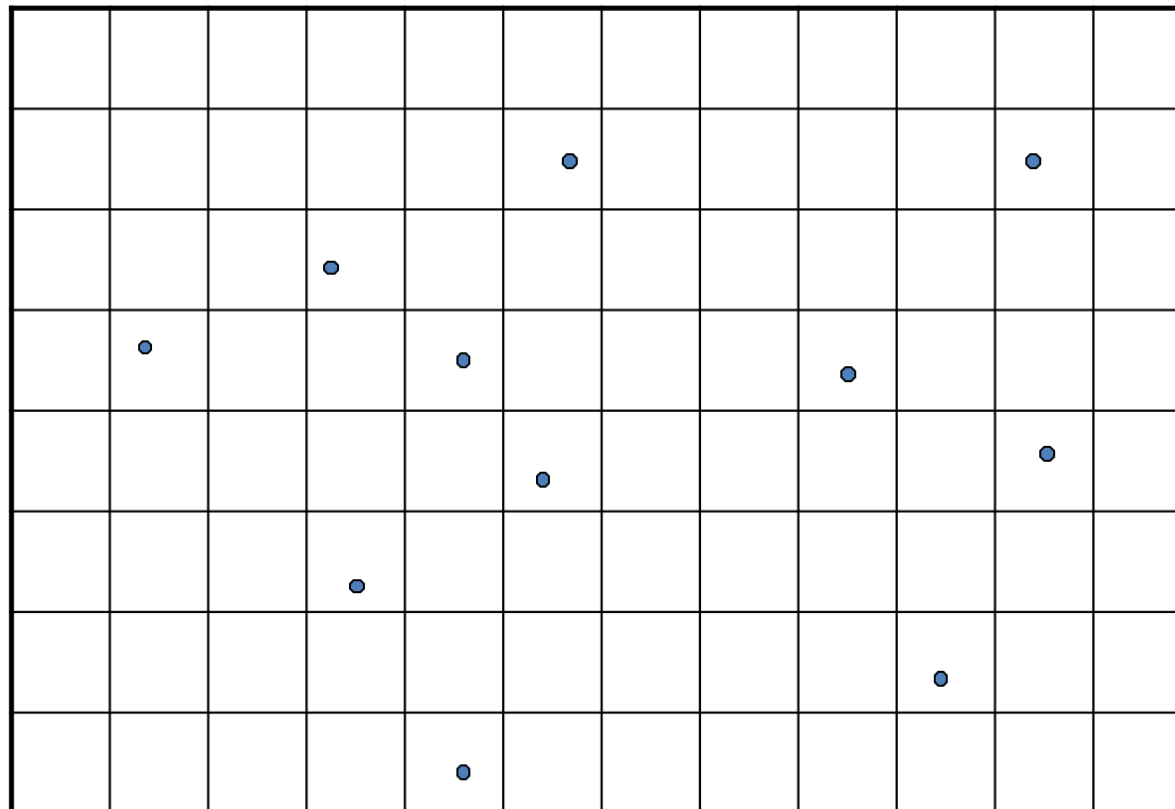
1. Plate at limiting dilution (<1 cell/well) in 96 well plates.

- for example: volume per well: 100 microliter

- use fused cells and prepare cell suspension of ca. 1 cell/ml (0,1 cell per 100ul well → 1 cell every 10 wells)

2. Allow clones to expand in 96 well = cell clone or cell line (a population of hybridoma cells derived from a single cell = genetically identical) → Hybridoma line

3. Further expand positive well and test for production of antibody of desired specificity in culture supernatant



Generation of monoclonal antibodies

Cloning hybridomas from fusion event

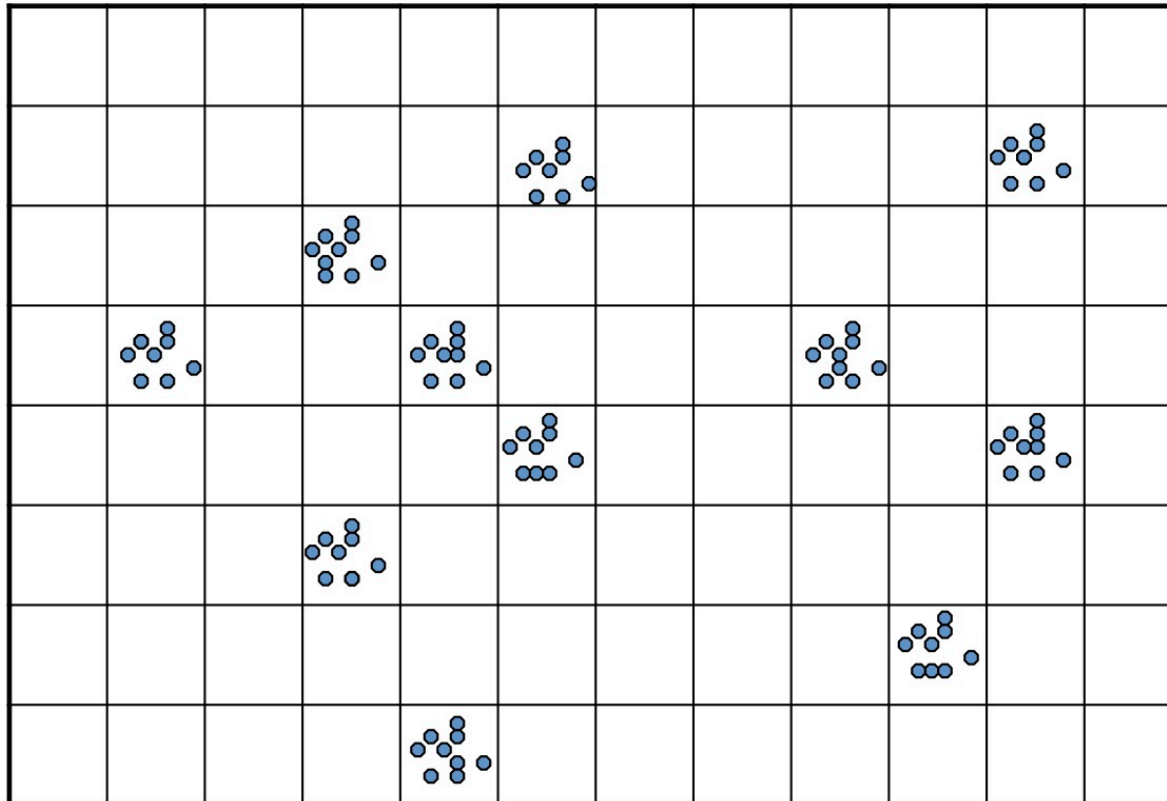
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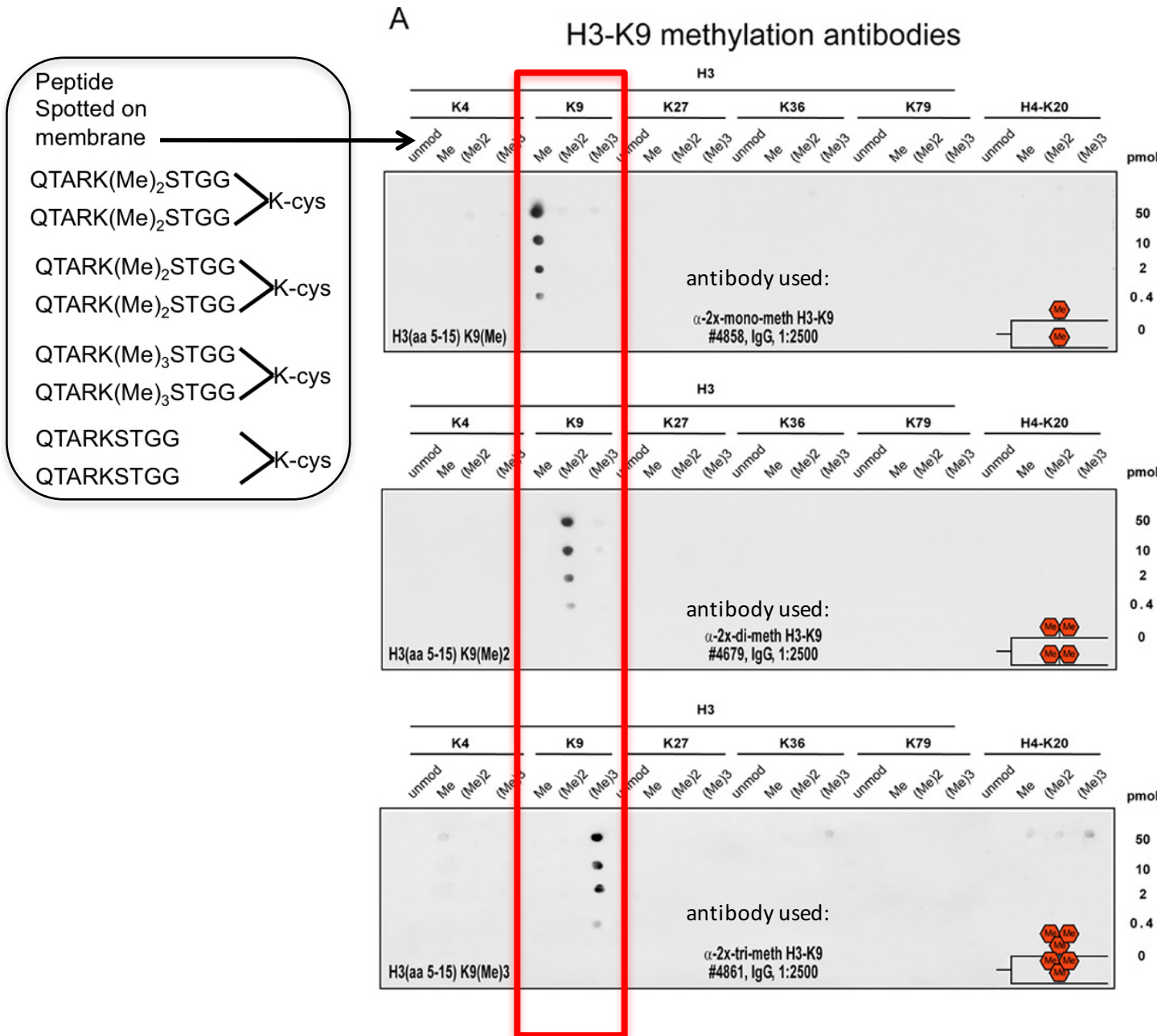


-Hybridoma cells secrete antibody into cell culture medium

-Purification of antibody

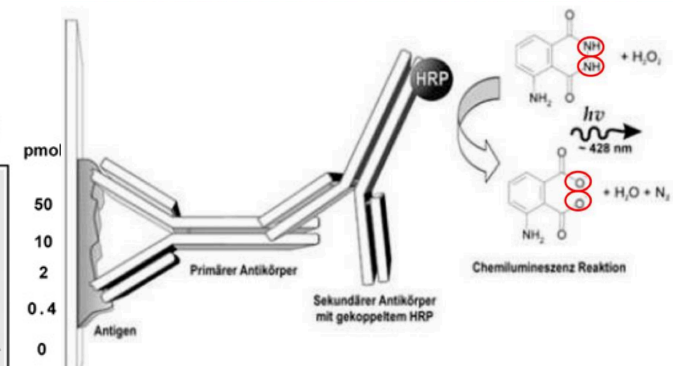
-Validation of antibody

Validation of antibodies

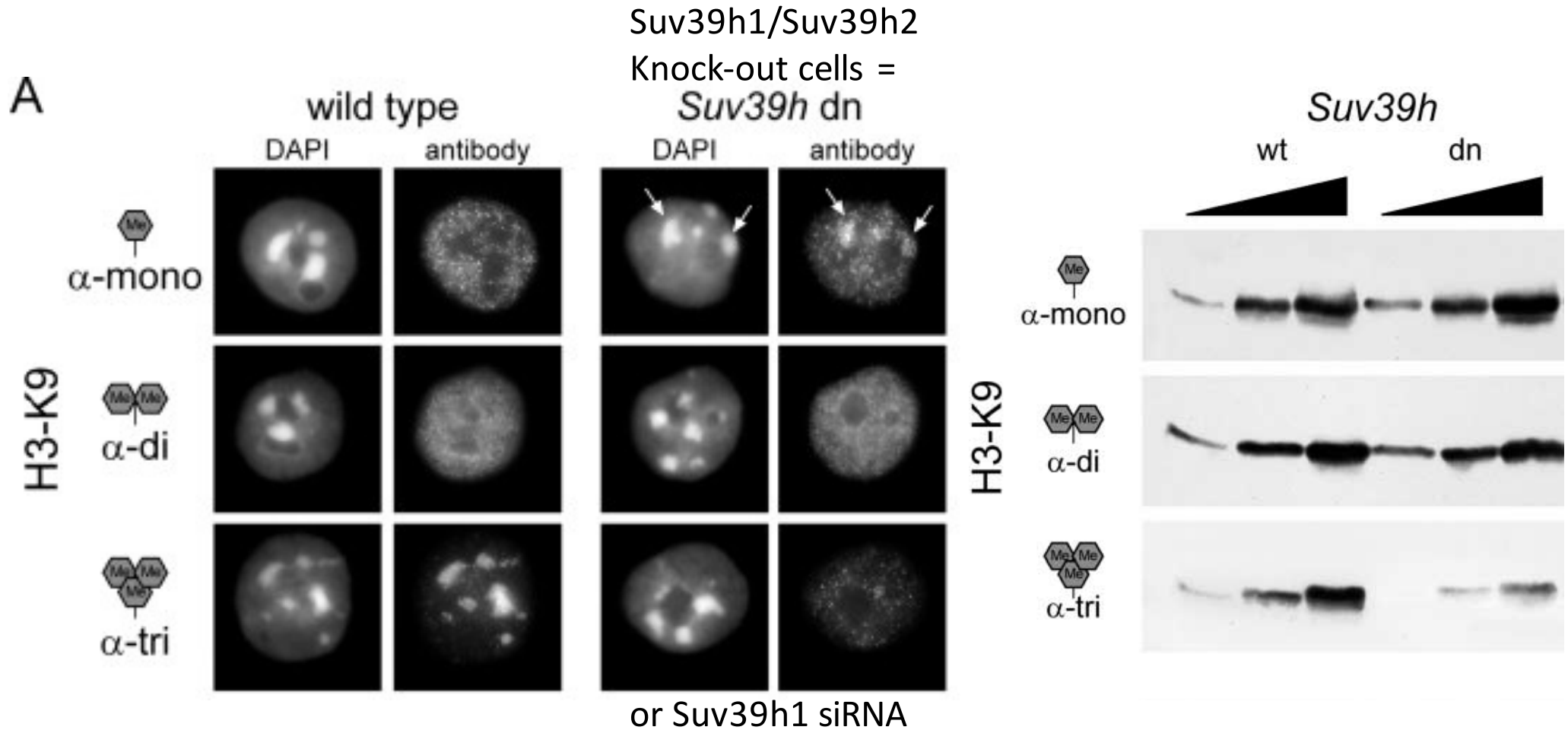


Validation by dot-blot
(pure antigen spotted on membrane)

1. Spot branched peptides at different concentration on membrane
2. Incubate with respective Antibody
3. Incubate with secondary Antibody that is coupled with horseradish peroxidase
4. Add substrate for Horseradish peroxidase
5. Develop



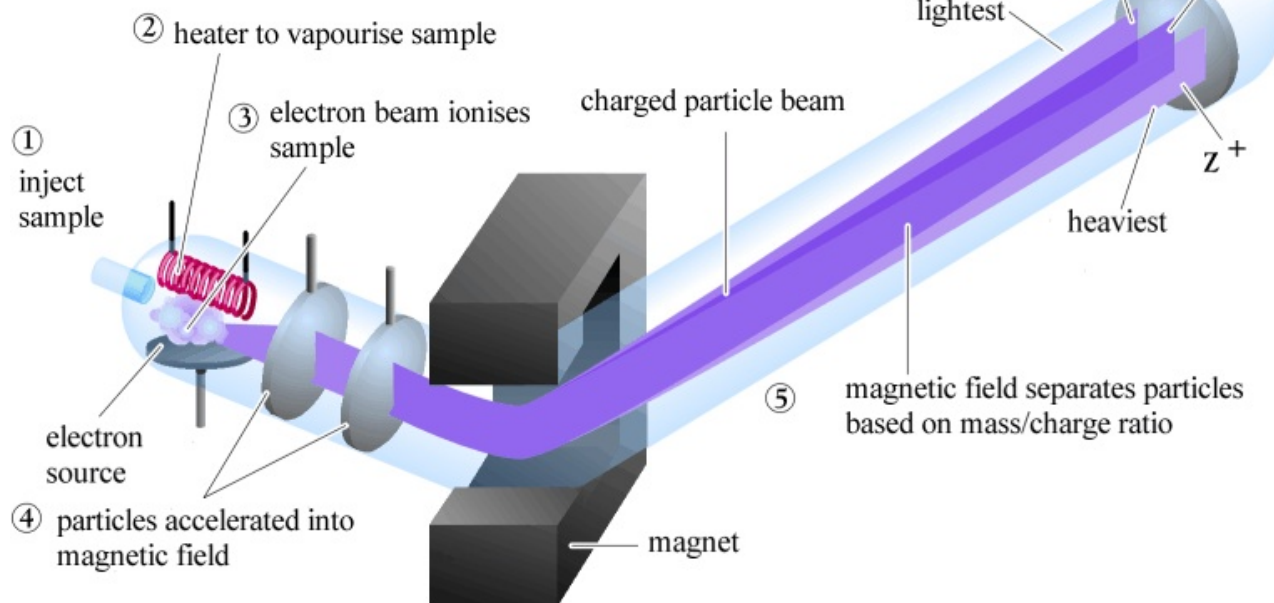
Obtaining evidence for the specificity of histone modifying enzymes → in cells



In *Suv39h* dn cells, H3K9me2 and H3K9me2 are still present.
***Suv39h1* is a H3K9 specific HMTase that is required to establish the TRI-METHYLATION of H3K9 in mammalian cells**

Note, that in the absence of H3K9me3, H3K9me1 is localized at DAPI rich regions

2. Studying histone modifications by mass spectrometry



In a typical MS procedure proteins are ionized, for example by bombarding it with electrons. This may cause some of the sample's molecules to break into charged fragments.

These ions are then separated according to their mass-to-charge ratio, typically by accelerating them and subjecting them to an electric or magnetic field:

Ions of the same mass-to-charge ratio will undergo the same amount of deflection. Ions with different mass-to-charge ratio will show different deflection

→ **mono-methylated H3K9 has different deflection than di- or tri-methylated H3K9**

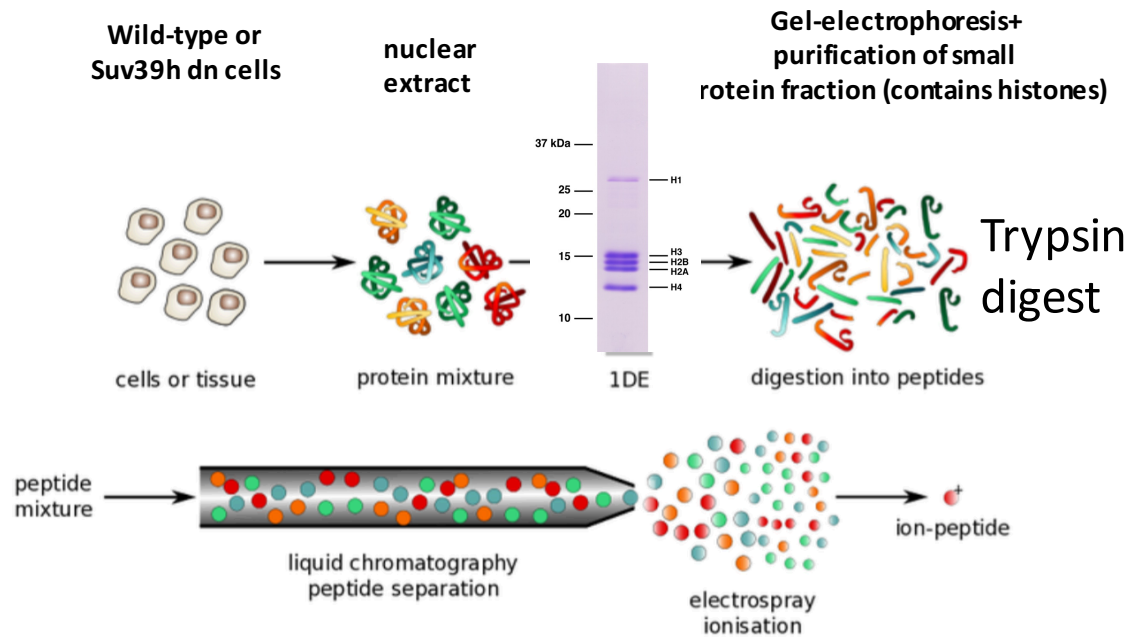
The ions are detected by a mechanism capable of detecting charged particles, such as an electron multiplier. Results are displayed as spectra of the relative abundance of detected ions as a function of the mass-to-charge ratio.

The molecules in the sample can be identified by correlating known masses to the identified masses or through a characteristic fragmentation pattern.

MW of all amino acids and all their possible modifications are known = identifiable by mass-to-charge ratio: also when present in a series of amino acids

MOST IMPORTANT: provide a sample with defined peptide fragments → achieved by digest with proteases that cut peptides at defined positions (comparable with restriction enzymes)

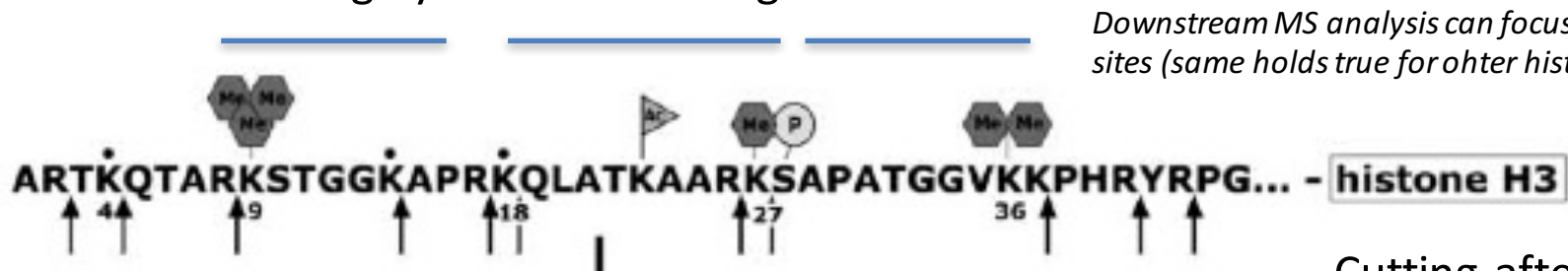
2. Studying histone modifications by mass spectrometry



Digestion of fraction containing histones with proteases creates a “library” of small peptides that are derived from histones:
Protease= Trypsin
Trypsin cleaves after every Lysine (K) and Arginine (R)

IMPORTANT: we know already All proteins and the amino acid sequence of all proteins of a cell!!!
That means we can predict all possible small peptide sequences that result from a trypsin cleavage

Highly informative fragments



Cutting after K and R

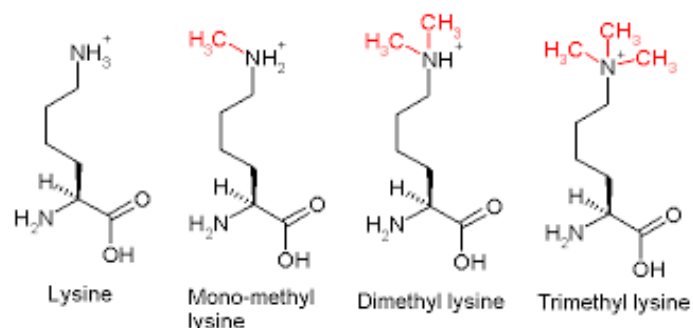
Arrows indicate trypsin target sites for cleavage

Histone modifications change m/z ratio in mass spec experiments



Table of amino acid residues .

Symbol	Structure	Mass (Da)
Ala A	-NH.CH.(CH ₃).CO-	71.0
Arg R	-NH.CH.[(CH ₂) ₃ .NH.C(NH).NH ₂].CO-	156.1
Asn N	-NH.CH.(CH ₂ CONH ₂).CO-	114.0
Asp D	-NH.CH.(CH ₂ COOH).CO-	115.0
Cys C	-NH.CH.(CH ₂ SH).CO-	103.0
Gln Q	-NH.CH.(CH ₂ CH ₂ CONH ₂).CO-	128.1
Glu E	-NH.CH.(CH ₂ CH ₂ COOH).CO-	129.0
Gly G	-NH.CH ₂ .CO-	57.0
His H	-NH.CH.(CH ₂ C ₃ H ₃ N ₂).CO-	137.1
Ile I	-NH.CH.[CH.(CH ₃)CH ₂ .CH ₃].CO-	113.1
Leu	-NH.CH.[CH ₂ CH(CH ₃) ₂].CO-	113.1
Lys K	-NH.CH.[(CH ₂) ₄ NH ₂].CO-	128.1
Met M	-NH.CH.[(CH ₂) ₂ .SCH ₃].CO-	131.0
Phe F	-NH.CH.(CH ₂ Ph).CO-	147.1
Pro P	-NH.(CH ₂) ₃ .CH.CO-	97.1
Ser S	-NH.CH.(CH ₂ OH).CO-	87.0
Thr T	-NH.CH.[CH(OH)CH ₃].CO-	101.0
Trp W	-NH.CH.[CH ₂ .C ₈ H ₆ N].CO-	186.1
Tyr Y	-NH.CH.[(CH ₂) ₂ .C ₆ H ₄ .OH].CO-	163.1
Val V	-NH.CH.[CH(CH ₃) ₂].CO-	99.1



KSTGGK: MW= 577 g/mol

CH₃ : MW= 15 g/mol

Kme₃STGGK: MW=577 – 3 + 45 = 619 g/mol

Kme₂STGGK: MW= 577 – 2 + 30 = 605 g/mol

Kme₁STGGK: MW= 577 – 1 + 15 = 591g/mol

Protonation
changes MW
in a defined
manner

Molecular weight

ionization

For details: An Introduction to Mass Spectrometry

<http://www.astbury.leeds.ac.uk/facil/MStut/mstutorial.htm>

2. Studying histone modifications by mass spectrometry

PROBLEM:

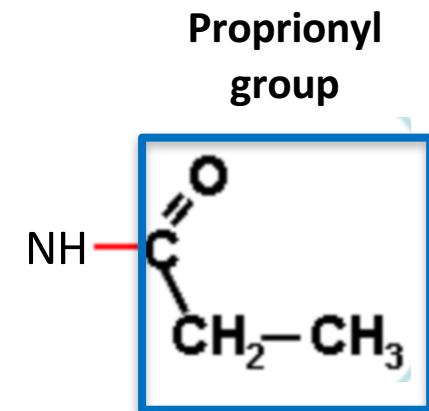
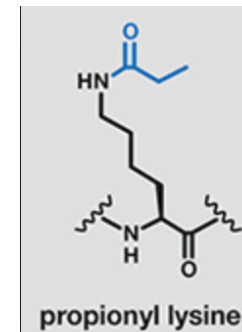
1. Trypsin cleavage after K is **not complete** because me2 or me3 on **some lysines** block block cutting by Trypsin
2. K is frequent in histone tails --> result many cleavages (mostly at me0, me1) → many small heterogeneous peptides → difficult to analyze



Trypsin can cleave after Arg (R)

Trypsin can cleave after K-unmethylated; K-me1

Trypsin cannot cleave after K-me2; K-me3



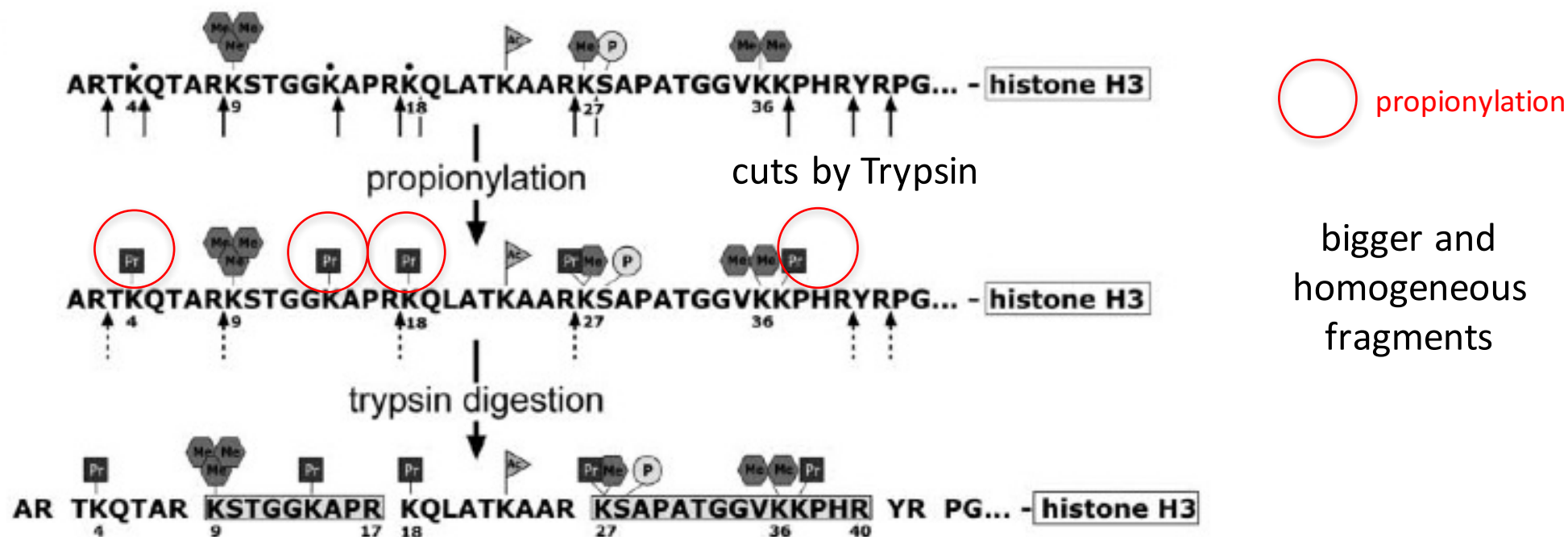
SOLUTION: In vitro Propionylation of **unmodified or mono-methylated lysine prevents trypsin cleavage**

→ now Trypsin can only cut after Arginine.

→ this allows a uniform cleavage of histone tails

→ creates a mass:charge ratio that **allows to differentiate between fragments carrying me0, me1, me2 or me3 marks** (me0 + 1x propionyl group; me1 + 1x propionyl group; me2 + 0 propionyl group; me3 + 0x propionyl groups)

2. Studying histone modifications by mass spectrometry



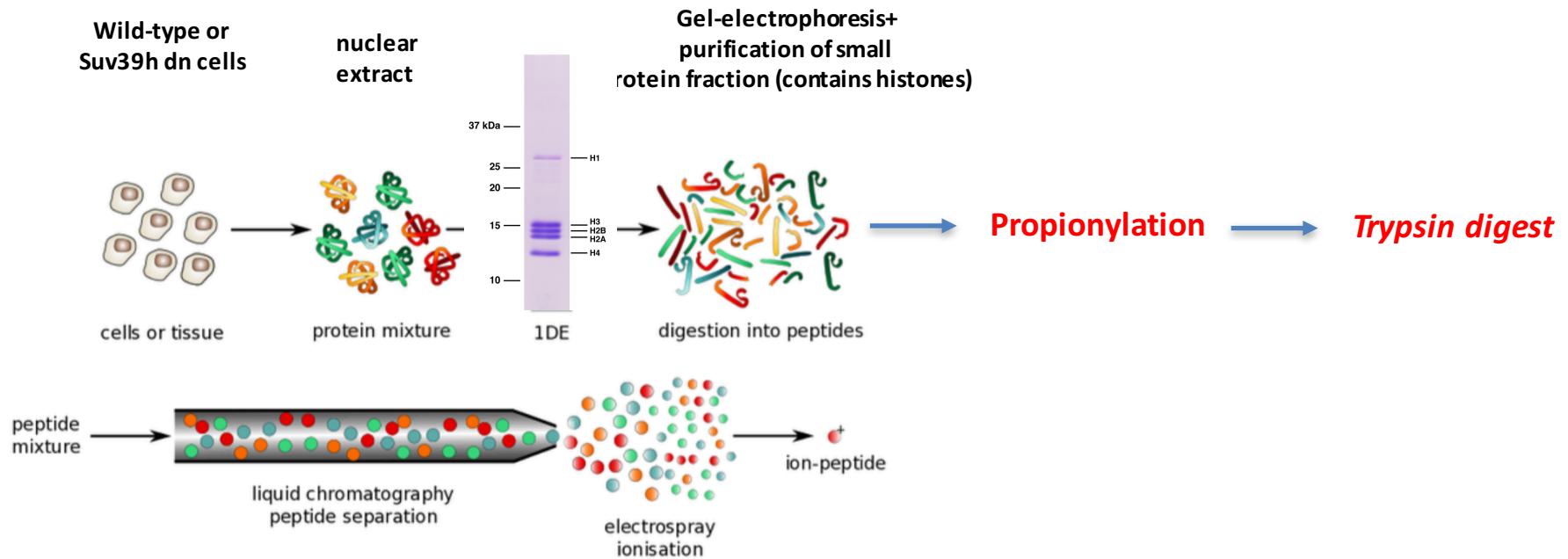
TRYPSIN cuts ONLY at Arg by Trypsin after propionylation

Now Trypsin can only cut after Arginine. This allows a uniform cleavage of histone tails

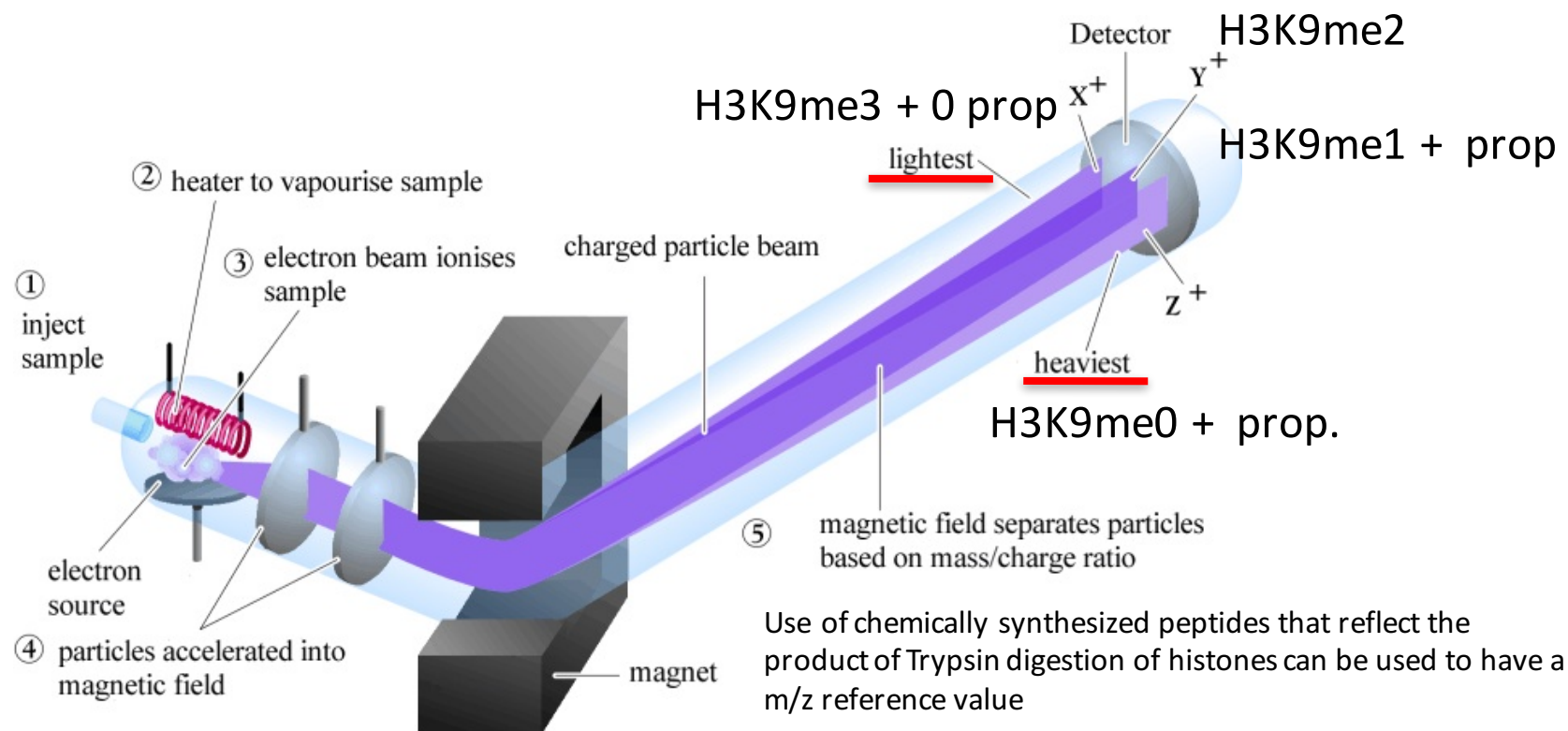
The number of methyl-groups/propinyl groups at the H3 peptide K9 – R17 decides on the deflection of this specific peptide in mass spectrometry.

Important: All peptides cut at Arg by Trypsin are analyzed contemporarily!!!!

2. Studying histone modifications by mass spectrometry



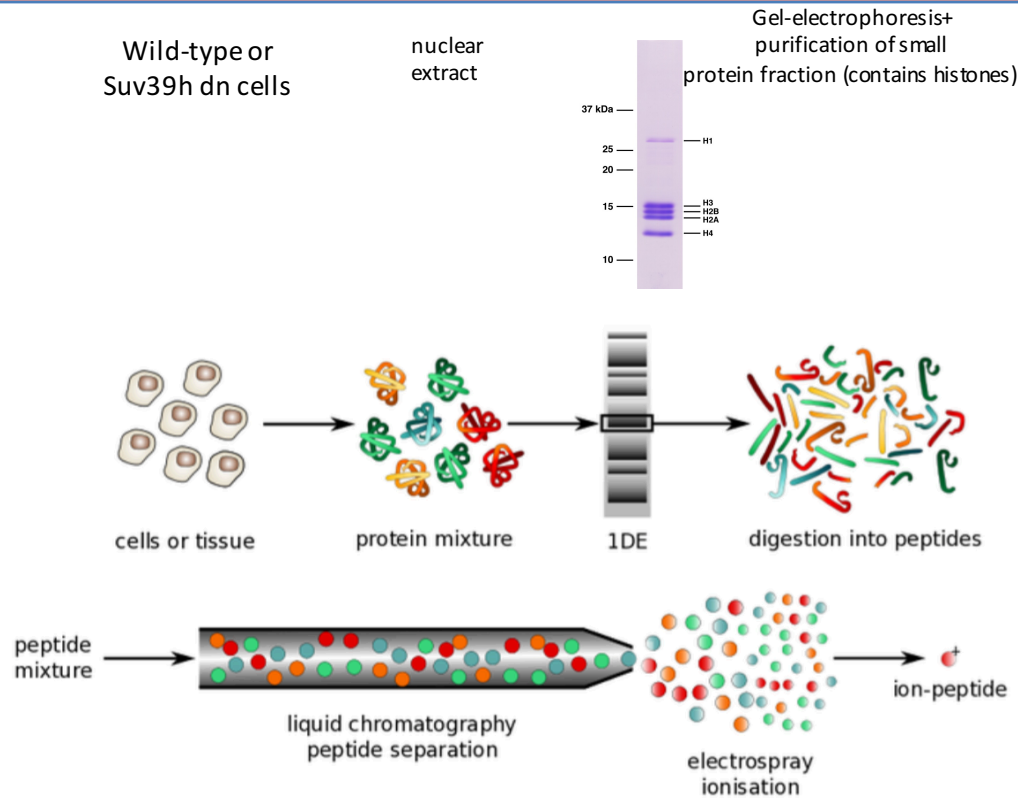
2. Studying histone modifications by mass spectrometry



The method allows to quantify the numbers of H3K9me0, H3K9me1 H3K9me2 and H3K9me3 in a sample → We can calculate the % of each histone modification in the given sample.

Important: The analysis is not limited to H3K9 → other histone modifications can be quantified in the same analysis

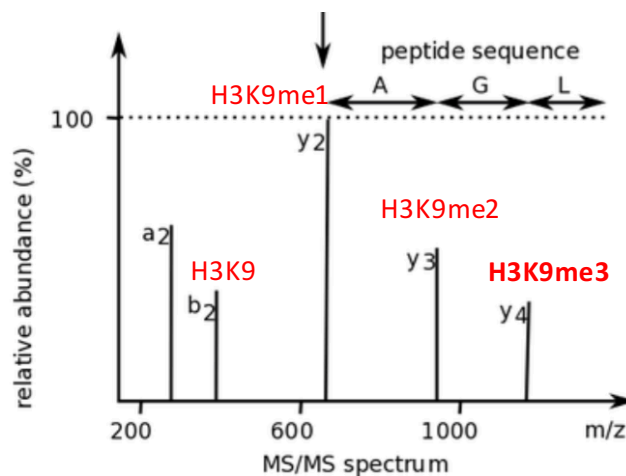
2. Studying histone modifications by mass spectroscopy



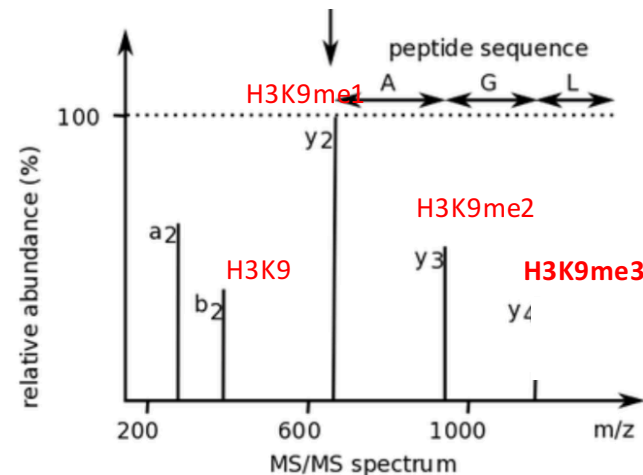
Digestion of fraction containing histones with proteases creates a “library” of small peptides that are derived from histones:
Protease= Trypsin
Trypsin cleaves after every Lysine (K) and Arginine (R)

IMPORTANT: we know already All proteins and the amino acid sequence of all proteins of a cell!!!
That means we can predict all possible small peptide sequences that result from a trypsin cleavage

MS

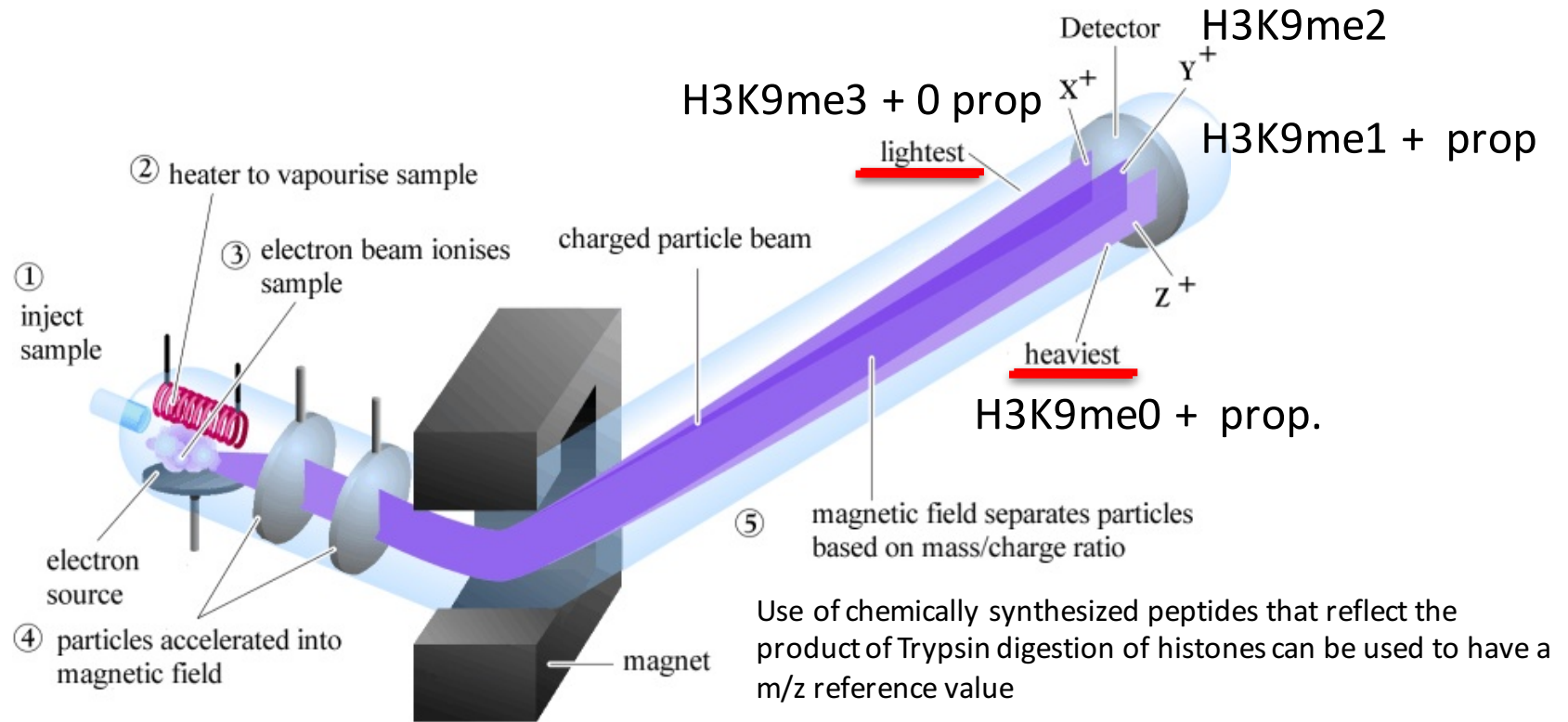


WT profile: Mass/charge ratio



Suv39DN profile: Mass/charge ratio

2. Studying histone modifications by mass spectrometry

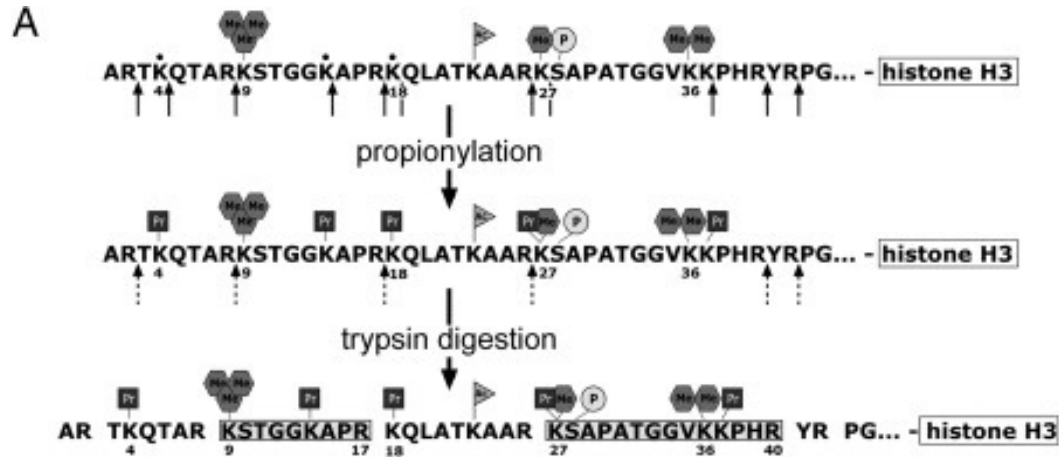


The method allows to quantify the numbers of H3K9me0, H3K9me1 H3K9me2 and H3K9me3 in a sample → We can calculate the % of each histone modification in the given sample.

Important MS is a proteomics approach: The analysis is not limited to H3K9 → all other histone modifications can be quantified in the same analysis. Remember: the histone proteins were isolated prior to MS.

This allows also to discover possible regulatory loops between different histone modifications!!! (for this you always need 2 biologically different samples: genetic model; differentiation, treatment, etc...)

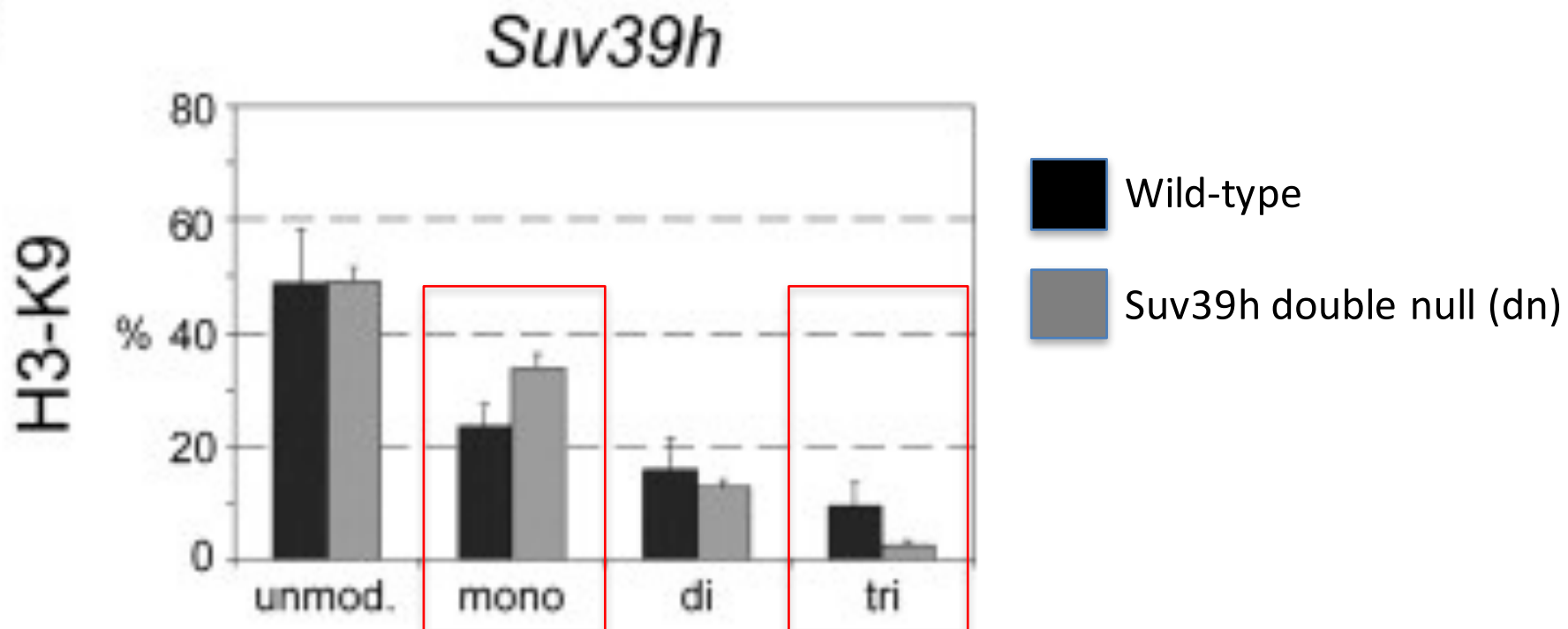
2. Quantifying histone modifications by mass spectroscopy



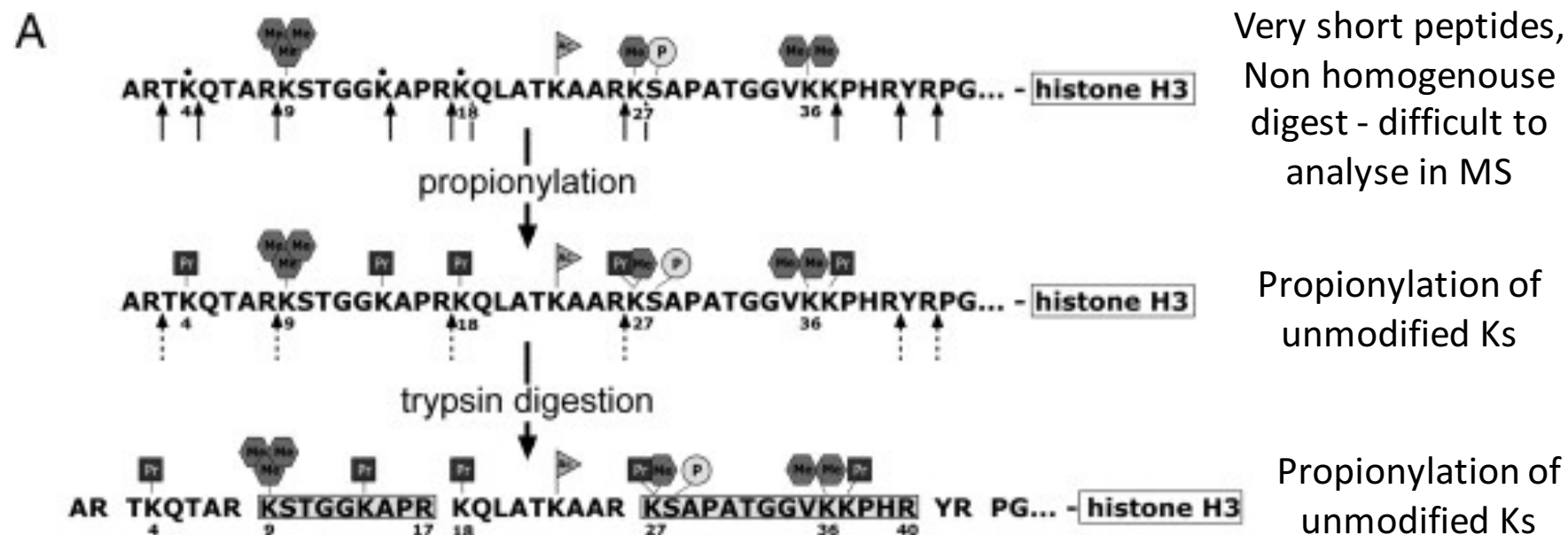
Suv39h1 and Suv39h2 ensure H3K9me3.

H3K9me1 is a preferred substrate for Suv39h1/h2

Suv39h1/h2 methylate H3K9me1 until reaching the tri-methylated state



2. Quantifying histone modifications by mass spectroscopy



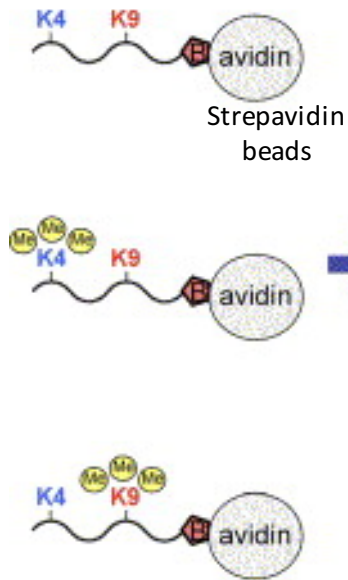
1. Identifying the enzymatic activity of a histone modifying enzymes (LOF of epigenetic writer)
2. Identification of “modifiable” aminoacids in histone tails
3. Functional link between histone modifications
4. Defining the ENTIRE epigenetic status of a cell type (for example: differentiated/stem cell)
5. Quantitative information on histone modifications (%)

HOW CAN EPIGENETIC READERS BE IDENTIFIED?

Identifying methyl-H3 binding proteins

- histone peptide pulldown assay: protein identification

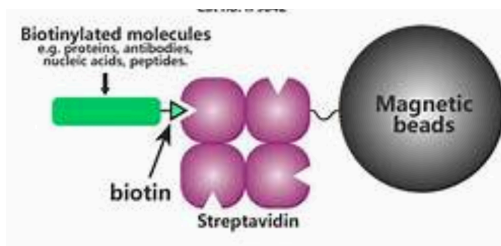
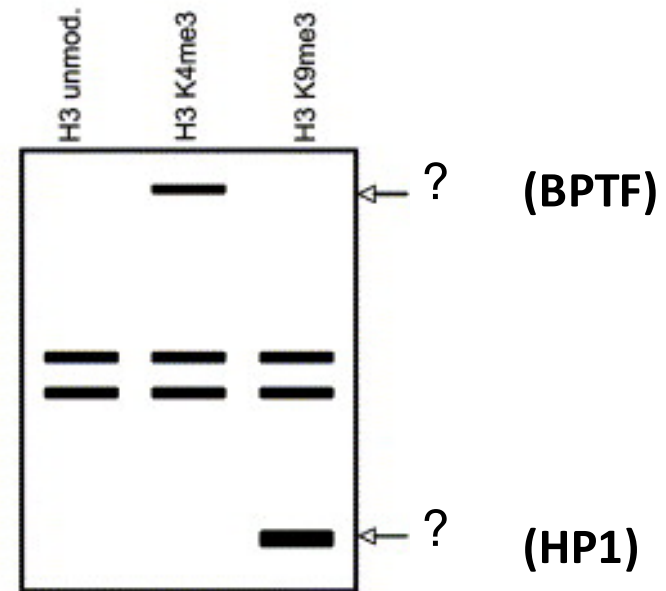
Chemically synthesized peptides, fused to biotin



nuclear extract



elution and SDS PAGE



- stain proteins in gel;
- cut out specific band,
- identify protein by Mass Spec

What are the target sites for Suv39h1 and H3K9me???

- histone peptide pull-down assay: validation of interaction and functional analysis
- Using the peptide pull-down assay, it was found that Lys9-methylated H3 binds to heterochromatin protein 1 (HP1)



Pull-down assay:

- Couple biotinylated histone tail-peptides, carrying specific modifications (methylated, or unmethylated) to streptavidine coated resin

- Incubate with recombinant ^{35}S -labelled HP1, produced in E-coli

- Wash resin

- Elute bound proteins, run gel and make radiography

- HP1 is a protein previously identified to be enriched in and important for heterochromatin assembly
- Lys9-methylated H3 binds to HP1 via the chromodomain motif in HP1

Lys9-methylated H3 binds to the conserved motif called chromodomain

What are the target sites for Suv39h1 and H3K9me???

- histone peptide pulldown assay: validation of interaction and functional analysis



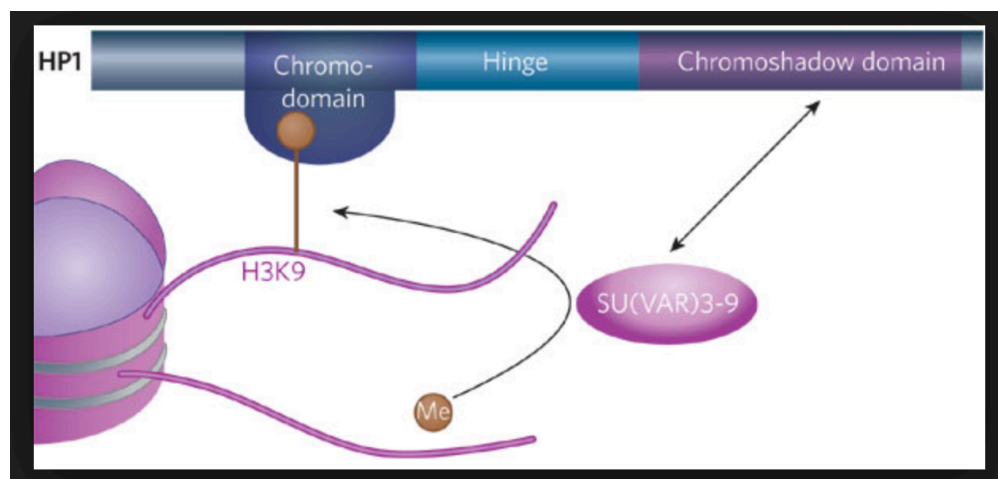
Pull-down assay:

- Couple biotinylated histone tail-peptides, carrying specific modifications (methylated, or unmethylated) to streptavidine coated resin

- ³⁵S-labelled mutant versions of HP1: ΔC; ΔCS; ΔH

- Wash resin

- Elute bound proteins, run gel and make radiography



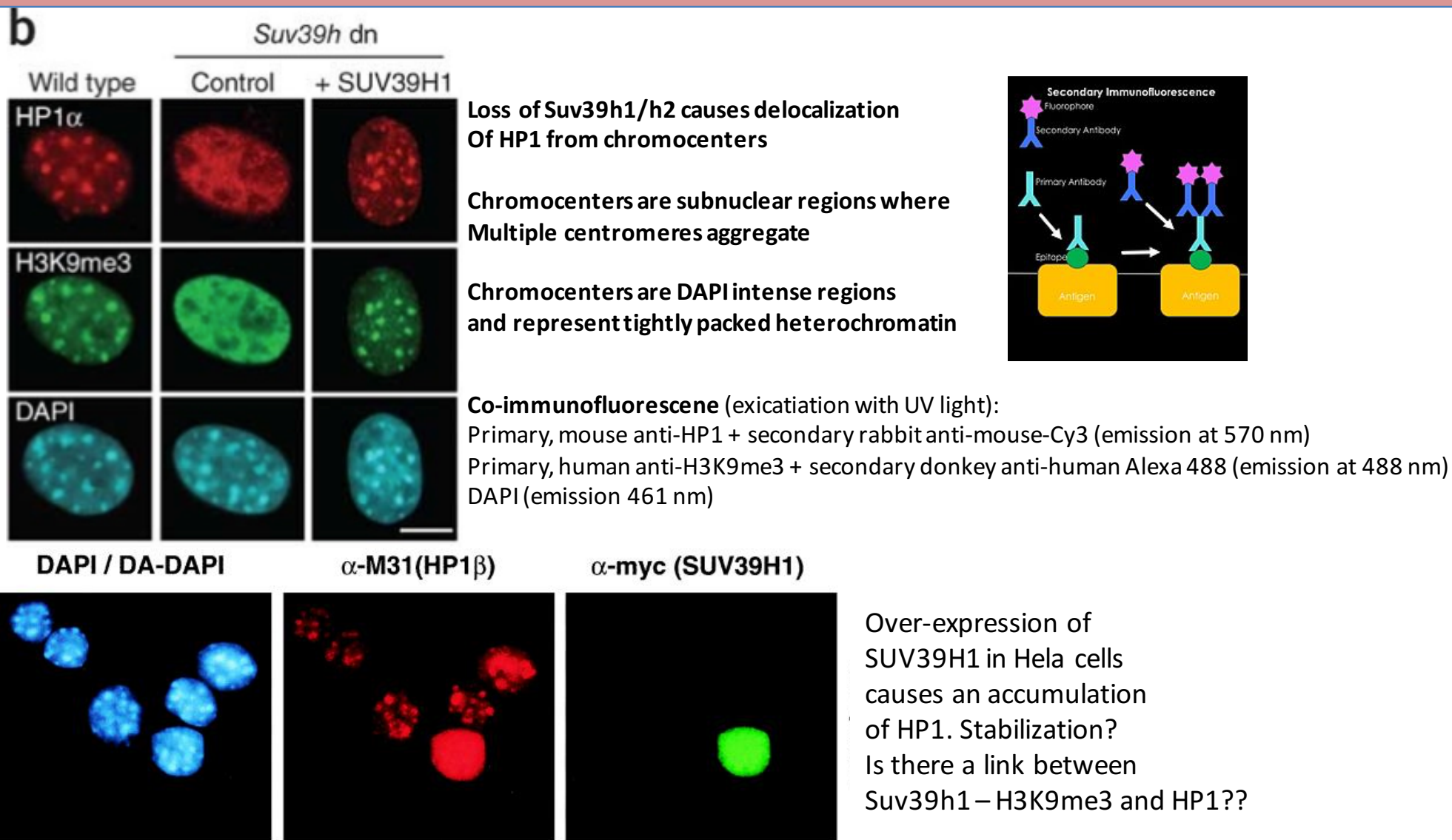
Delta C domain (chromodomain) doesn't bind:
= binds modified histone tail

Lys9-methylated H3 binds to the conserved motif called chromodomain

Where does Suv39h1 act in a cell ???

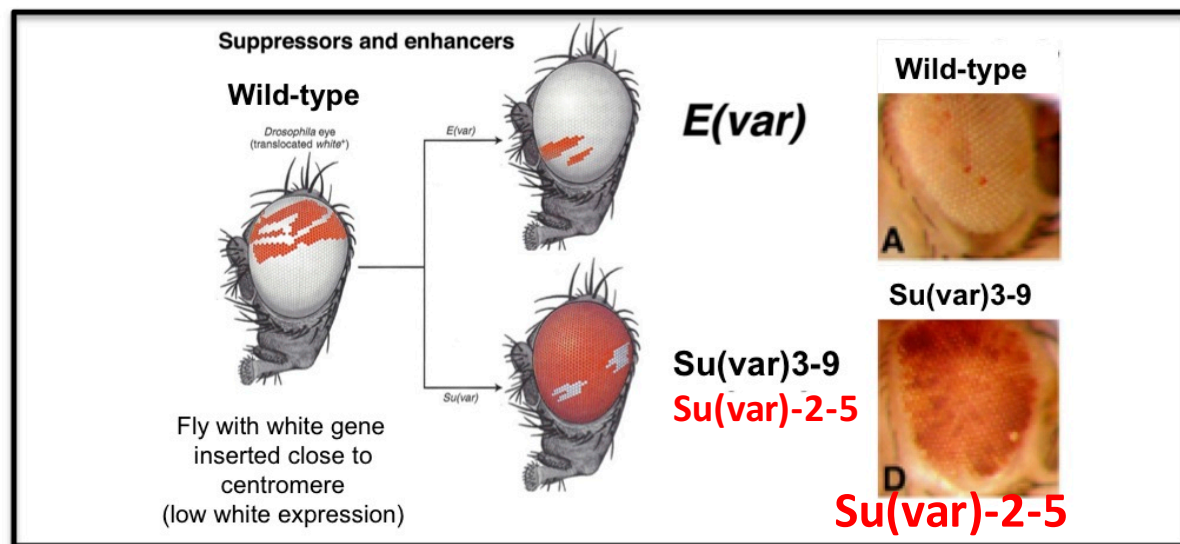
IMMUNOFLUORESCENCE → MACROSCOPIC ANALYSIS

→ Localization of protein across large regions of DNA



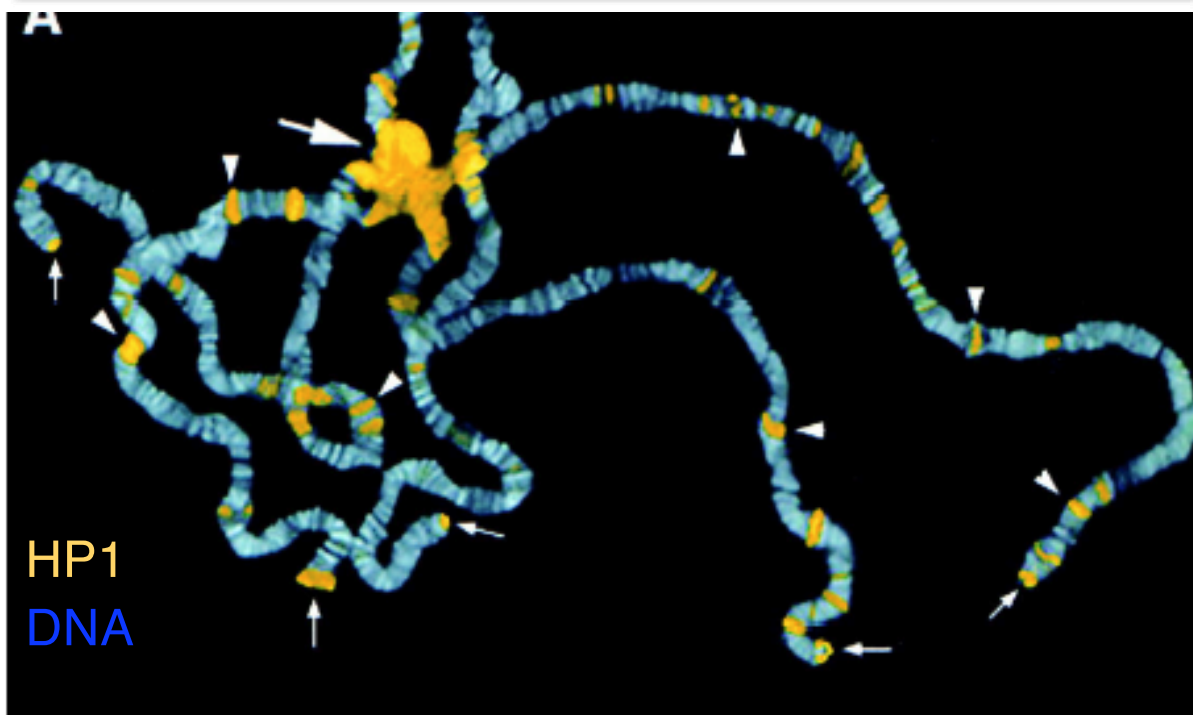
myc-SUV39H1: myc-epitope tagged Suv39H1

What are the target sites for Suv39h1 and H3K9me???

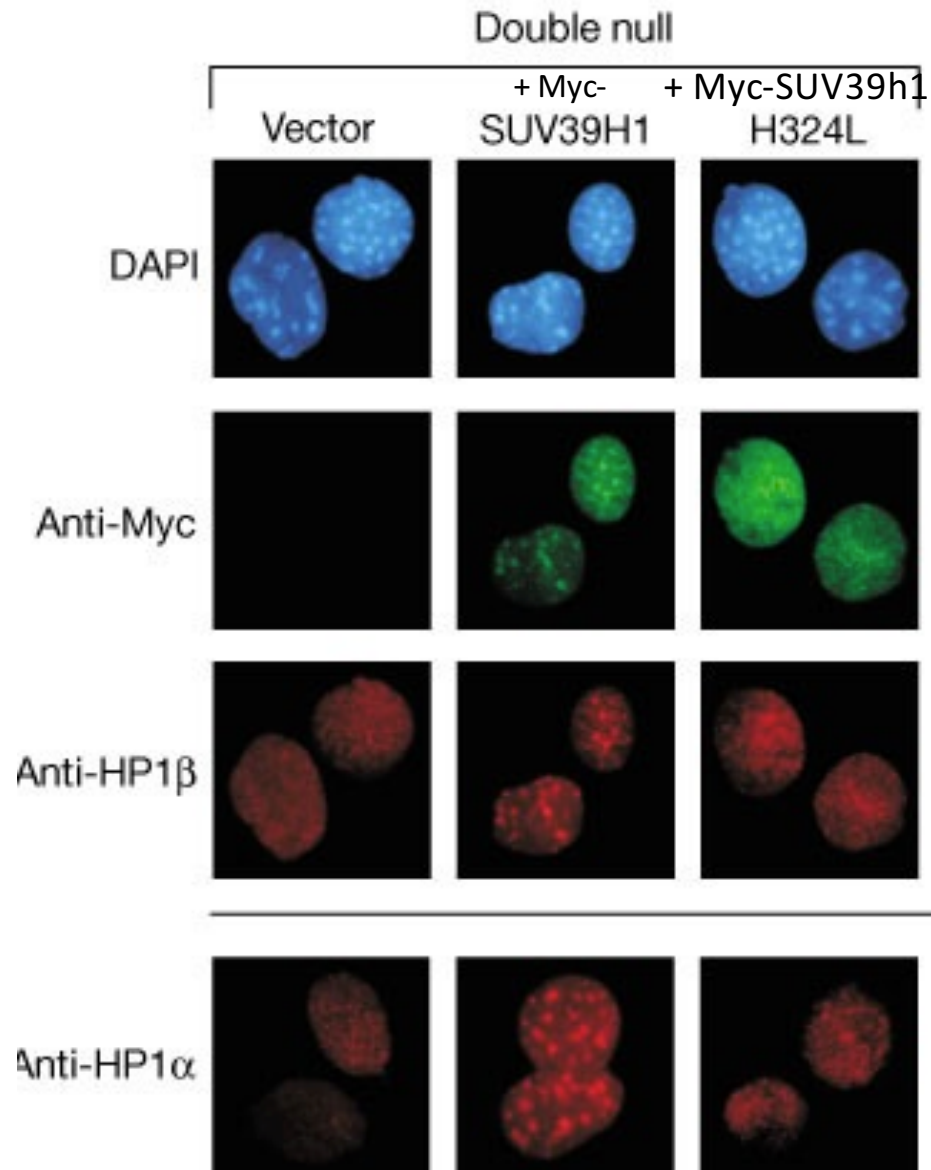


Another evidence for HP1 and Suv39h interaction came from *Drosophila*

- back to early genetics studies in *Drosophila*:
- *Su(var) 2-5* (gene) codes for heterochromatin protein 1
- *Su(var) 2-5* shows similar phenotype like *Su(var)3-9*
- HP1 in *Drosophila* is localized to the chromocenter



SUV39h HMTase activity is important to build constitutive heterochromatin



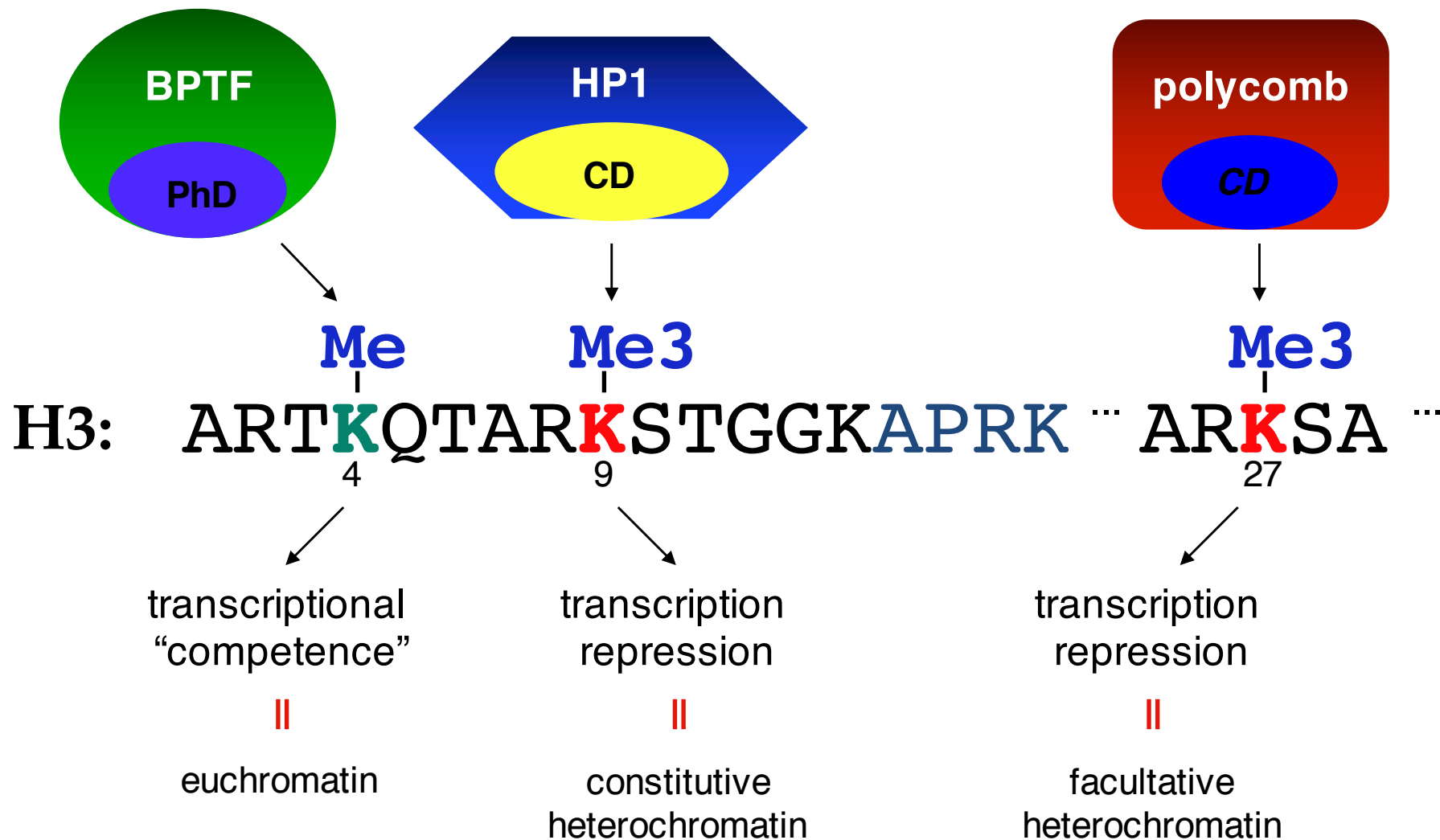
Loss of Suv39h1 expression results in delocalization of HP1 from chromocenters

Re-expression of Suv39h1 rescues Localization of HP1 to chromocenters

Re-expression of Suv39h1 that Contains an enzymatic dead mutation (H324L) in SET domain does not rescue the localization of HP1 to chromocenters

→ The enzymatic activity of Suv39h1 Is required for recruitment of HP1 to chromocenters

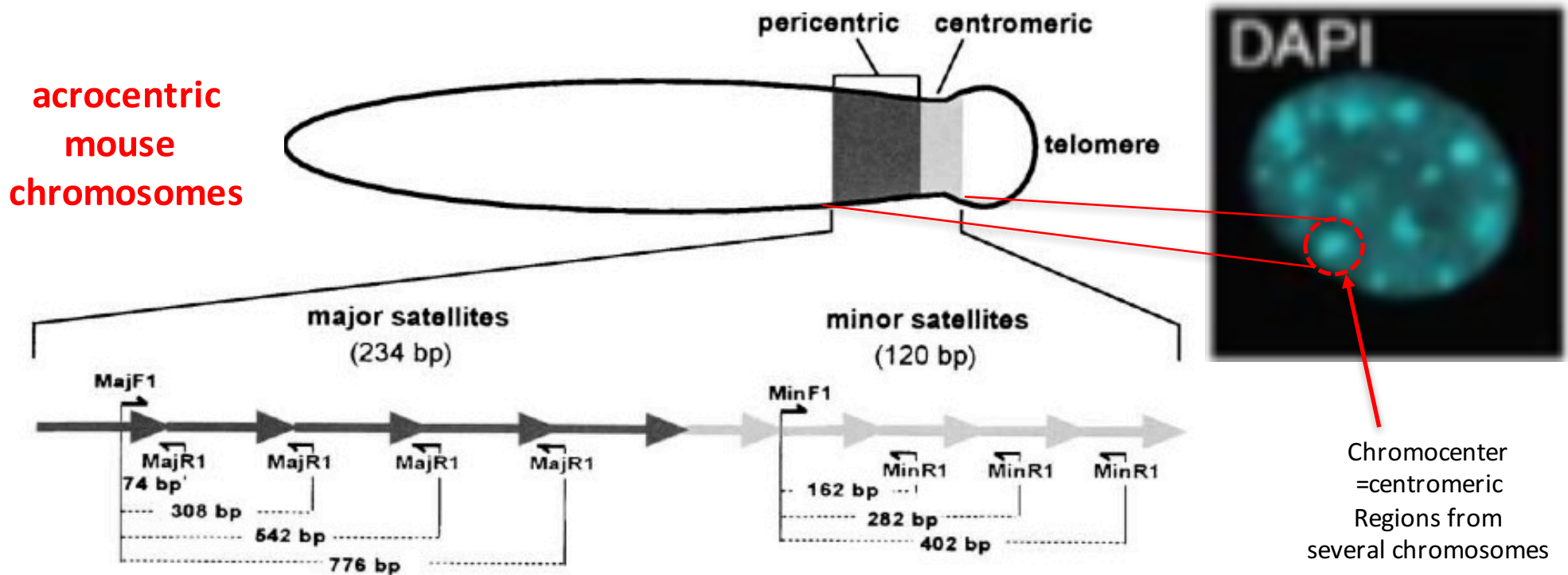
Site specific methylation of the H3 tail has different functions



What are the target sites for Suv39h1 and H3K9me???

CHROMATIN IMMUNOPRECIPITATION → DETAILED ANALYSIS
→ Localization of protein at a defined region - sequence

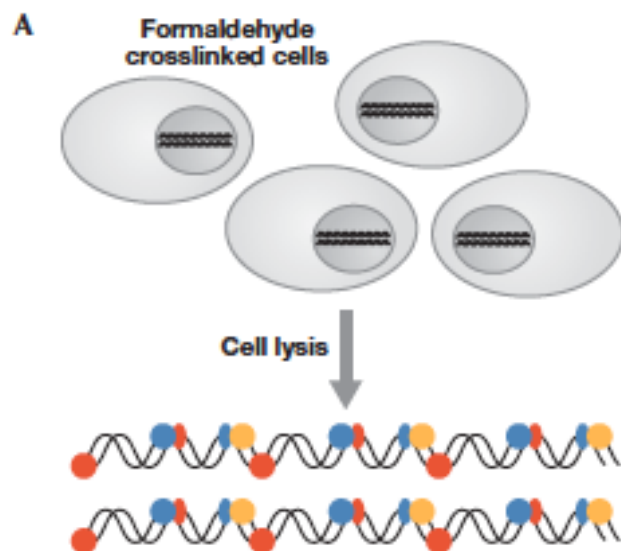
The combination of Immunoprecipitation methods and PCR analysis allows to define the histone code at defined sequences.
PCR primers define the site of analysis in the genome



EXAMPLE: Pericentric heterochromatin in mouse cells

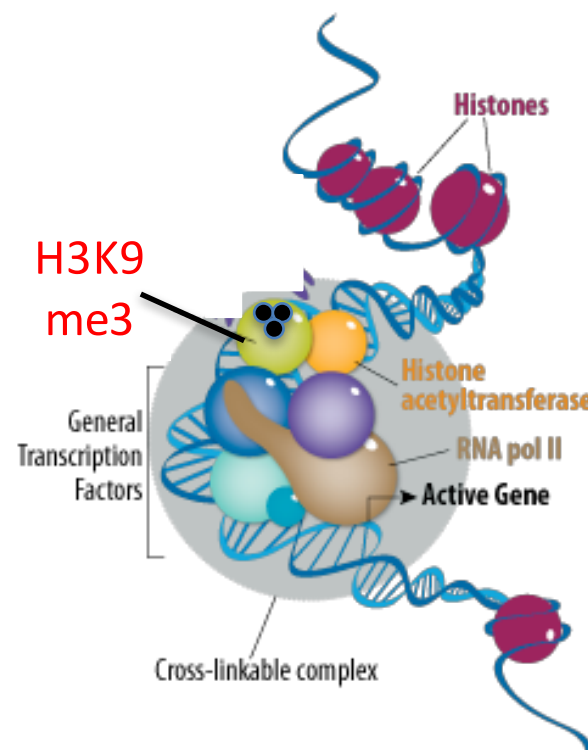
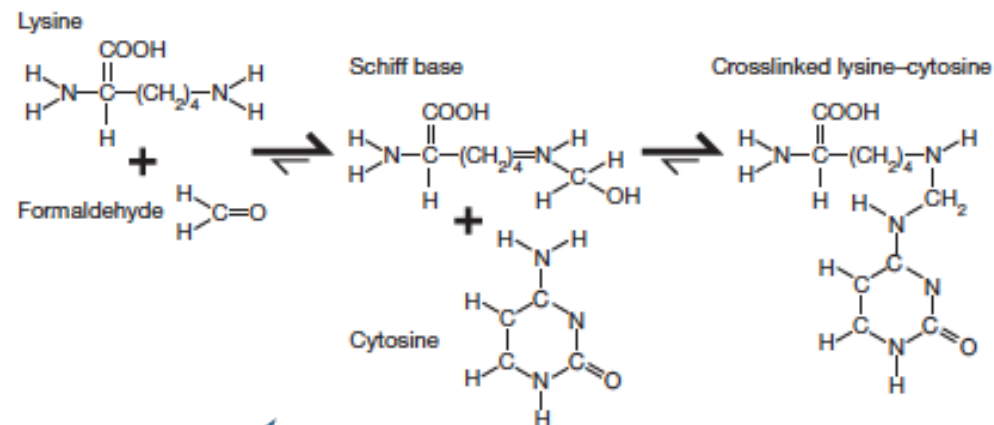
CHROMATIN IMMUNOPRECIPITATION

1. Cross linking with FA



Optimization is crucial.

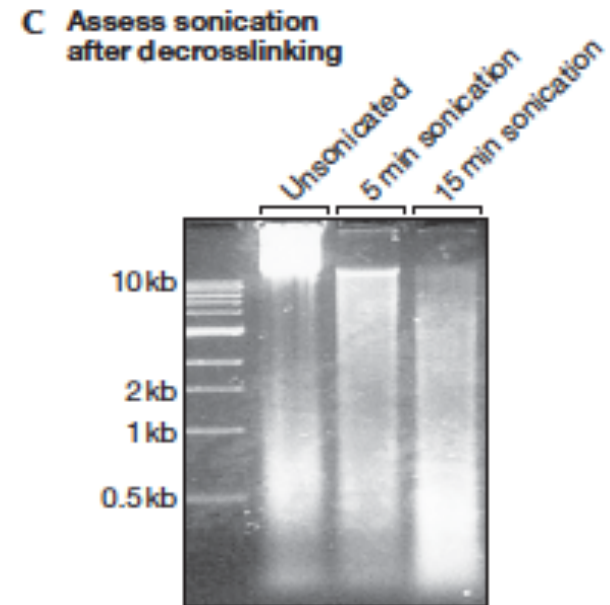
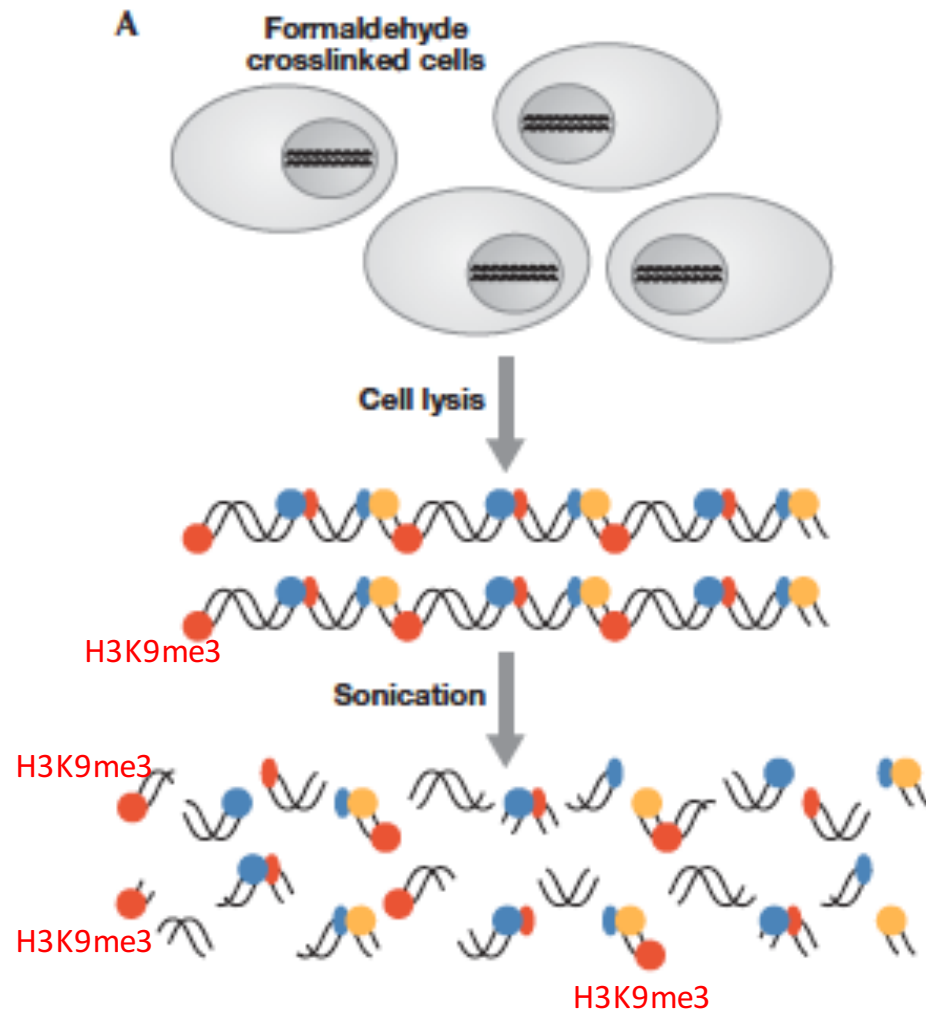
B Formaldehyde will crosslink amino or imino groups within 2Å, for example:



Covalently crosslinked chromatin

CHROMATIN IMMUNOPRECIPITATION

2. Cross linking followed by sonication (fragmentation of chromatin)

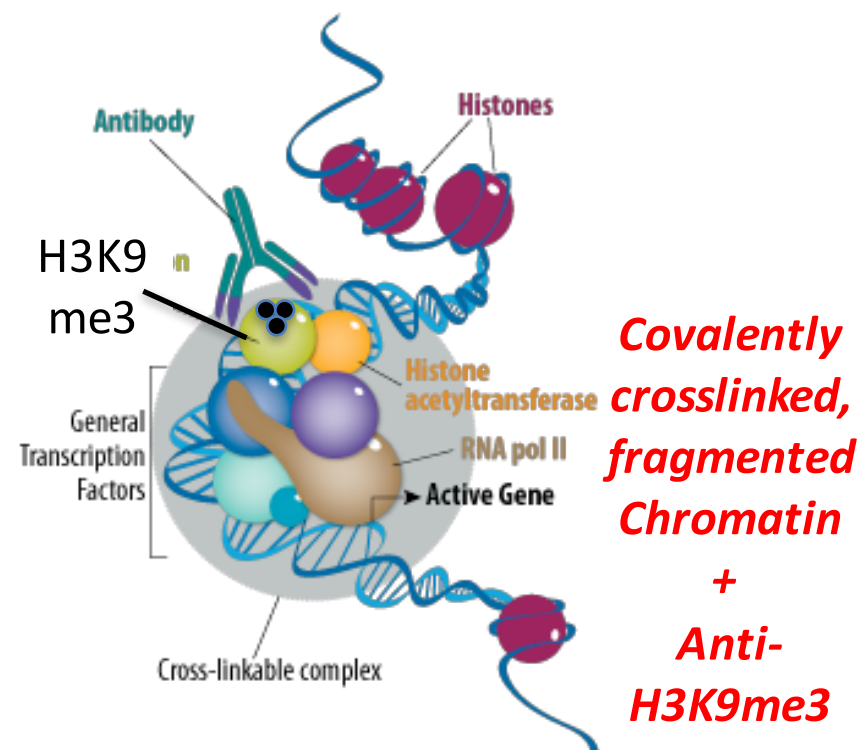
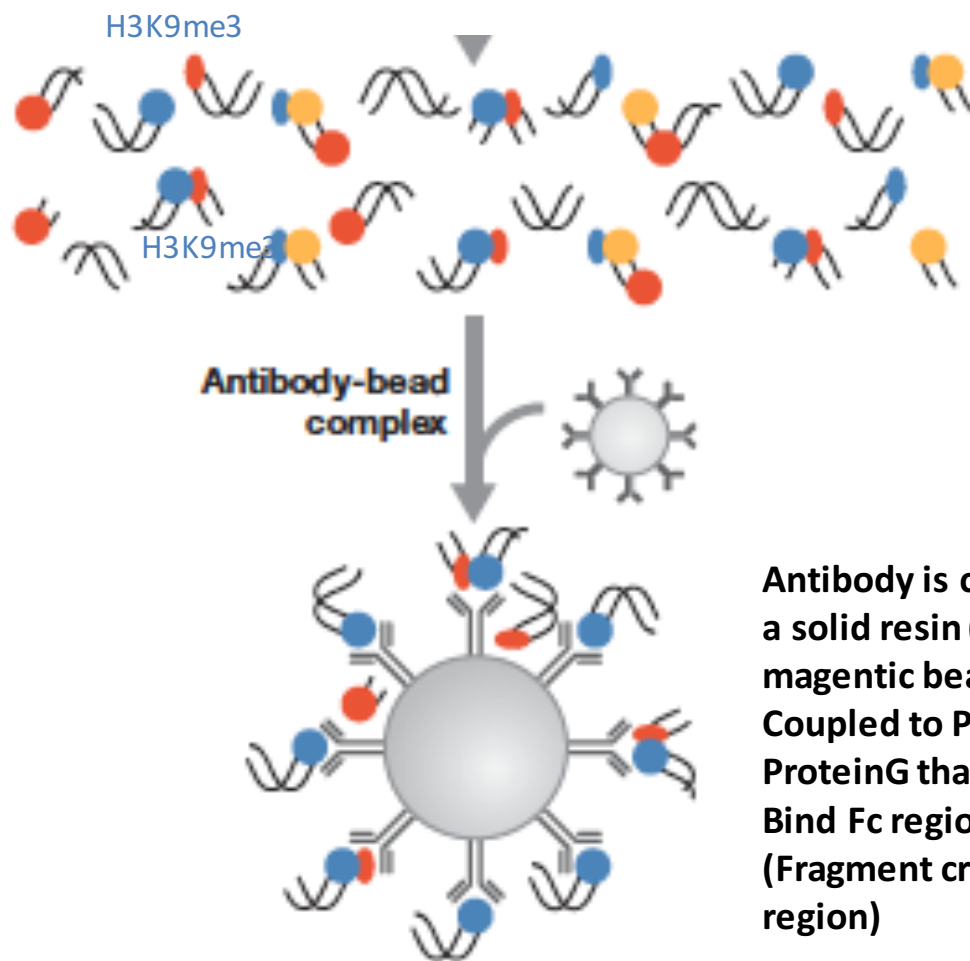


Chromatin is bulky → needs to be cut into small pieces to become soluble

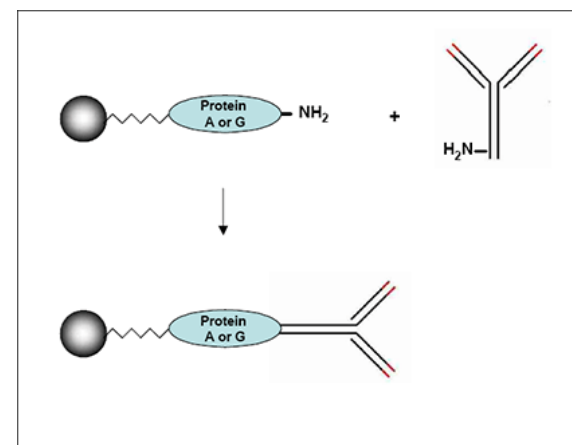
CHROMATIN IMMUNOPRECIPITATION

3. Immunoprecipitation (IP)

The protein of interest is immunoprecipitated together with the crosslinked DNA: Modified histones; epigenetic writers, epigenetic readers



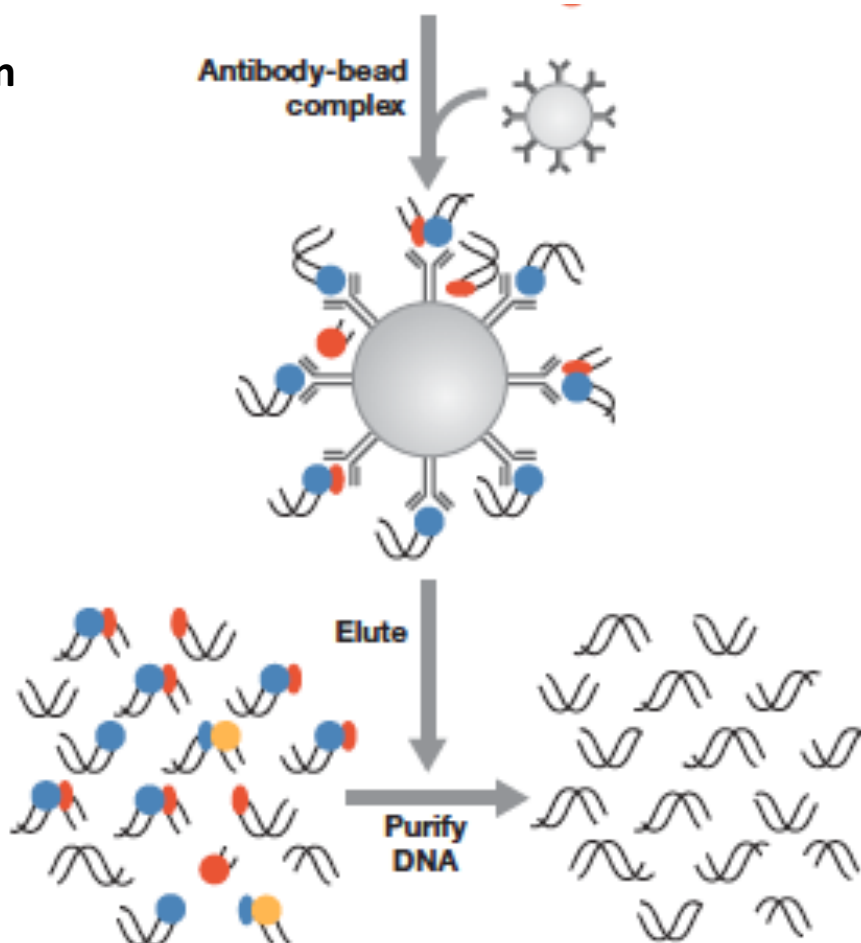
Antibody is coupled to a solid resin (agarose, magnetic beads) Coupled to Protein A or Protein G that Bind Fc region of antibody (Fragment crystallizable region)



CHROMATIN IMMUNOPRECIPITATION

4. Decrosslinking of PFA crosslinked chromatin and purification of the DNA

- Recover beads by centrifugation
- Wash beads
- Reverse the FA: 65°C, high salt concentrations: crosslink break
- RNase and Protease treatment
- Purification of DNA

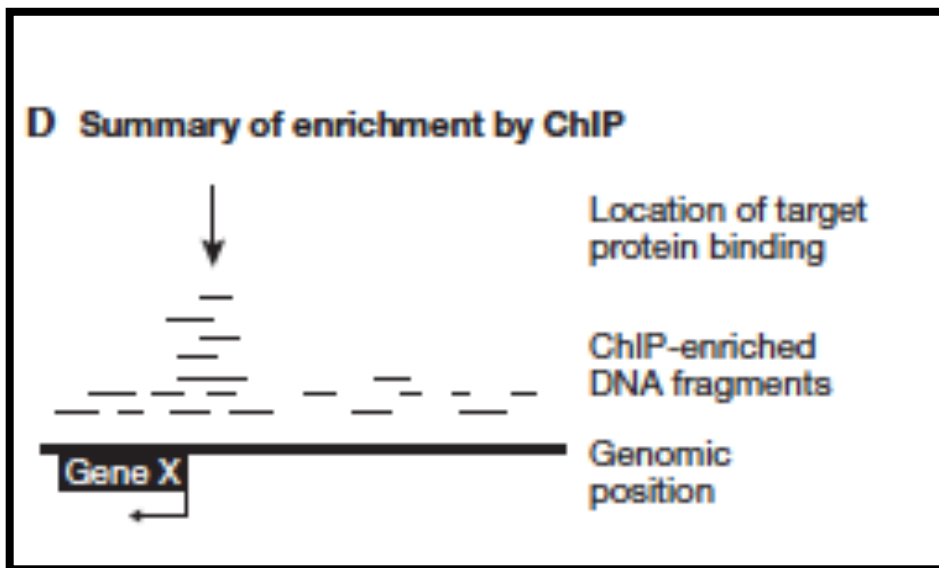


ANALYZE DNA TO IDENTIFY REGIONS WITH HIGH H3K9me3
(sequencing, hybridization, PCR with gene specific primers)

CHROMATIN IMMUNOPRECIPITATION

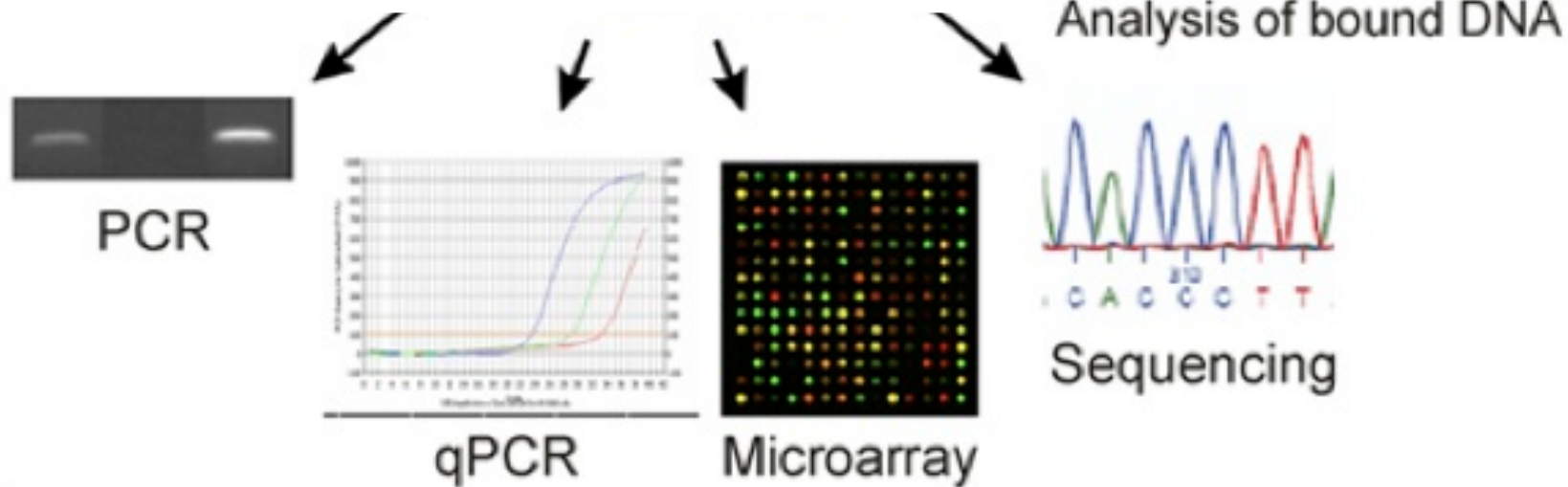
5. Analysis of ChIP DNA

Identification of DNA regions associated with the protein/modification of interest

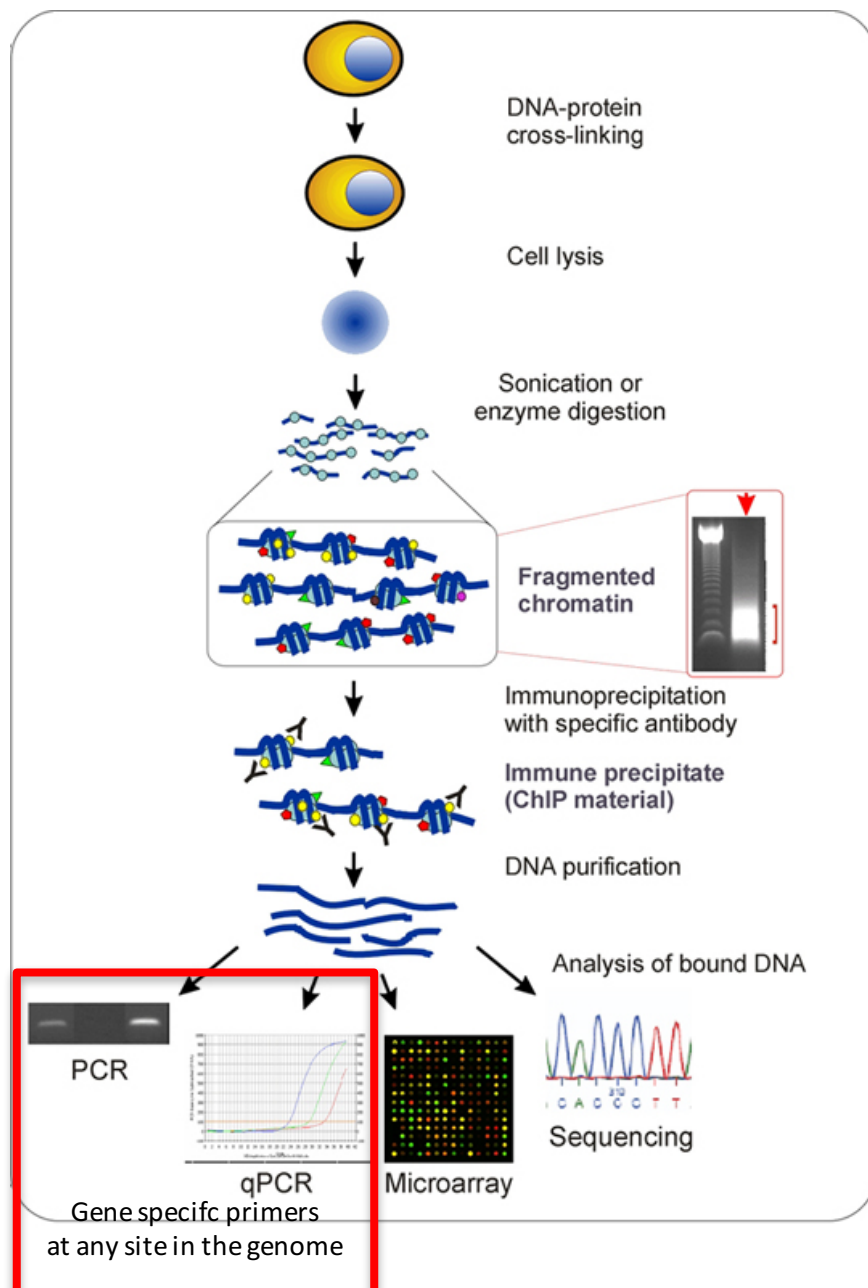


DNA from chromatin enriched in H3K9me
→ Correspond to precise sites in the genome

Methods of ChIP analysis



CHROMATIN IMMUNOPRECIPITATION (ChIP) → DETAILED ANALYSIS → Localization of protein at a defined region - sequence



Cell model system:

i.e. **Wild-type or Suv39 dn cells**
that grow in cell culture dish

1. Crosslink chromatin
(treatment of cells with Paraformaldehyde)
2. Sonicate crosslinked cells
3. Incubate chromatin fragments with antibodies raised against **H3K9me3**
4. Recover antibody-chromatin complexes using a resin that binds to the constant region of antibodies
5. Elute chromatin at high salt concentration and revert crosslinks at high temperature
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7. Purify DNA and precipitate DNA
8. Measure the amount of immunoprecipitated DNA
In control versus Suv39h dn cells

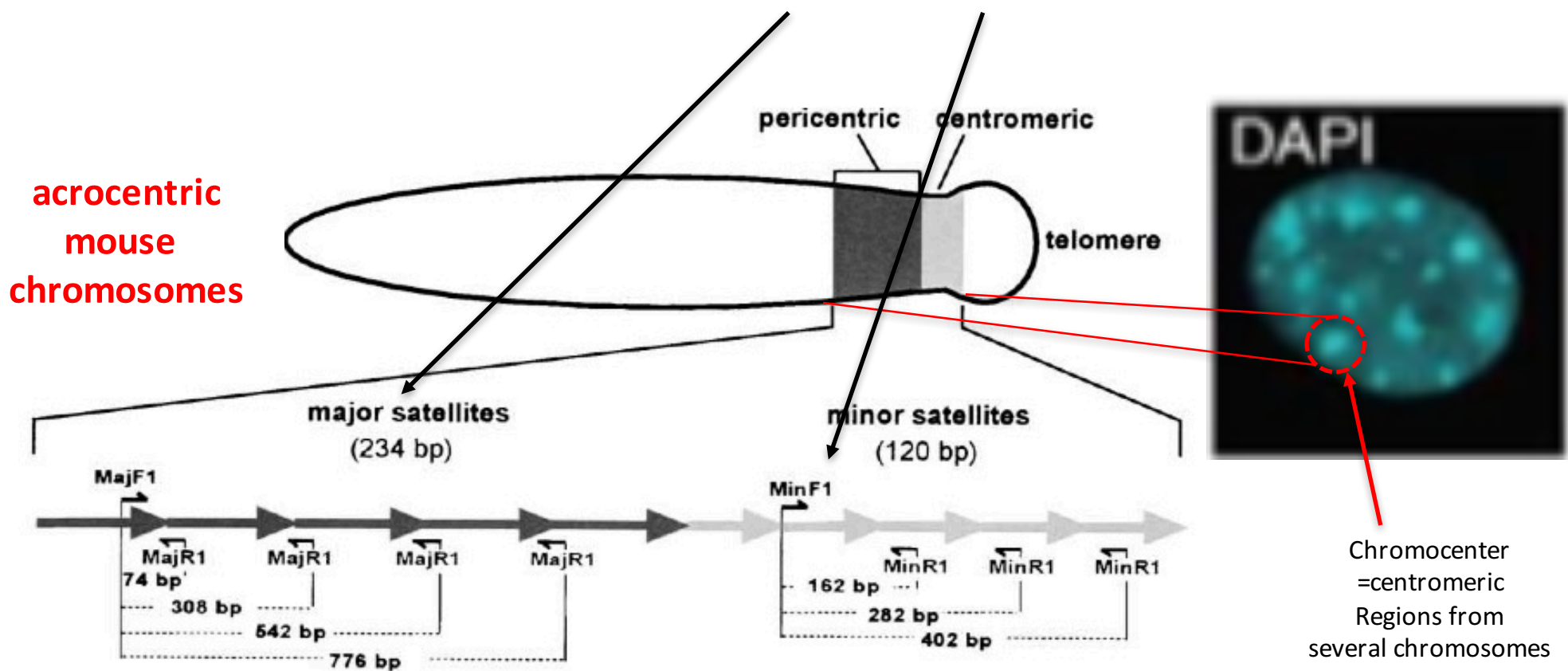
METHODS:

Quantitative PCR: design PCR oligos to amplify defined sequences in immunoprecipitates. Per PCR reaction only one Locus can be examined by real-time PCR

What are the target sites for Suv39h1 and H3K9me???

CHROMATIN IMMUNOPRECIPITATION FOLLOWED BY QUANTITATIVE PCR

Design PCR oligos that amplify major and minor satellite repeats



EXAMPLE: Pericentric heterochromatin in mouse cells

Min/Maj F1...: forward primer in unique region

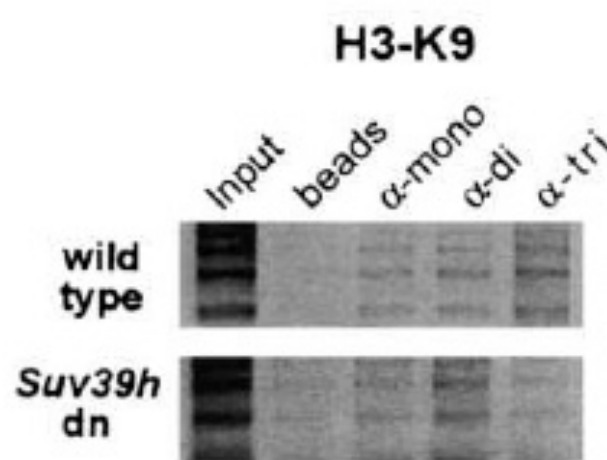
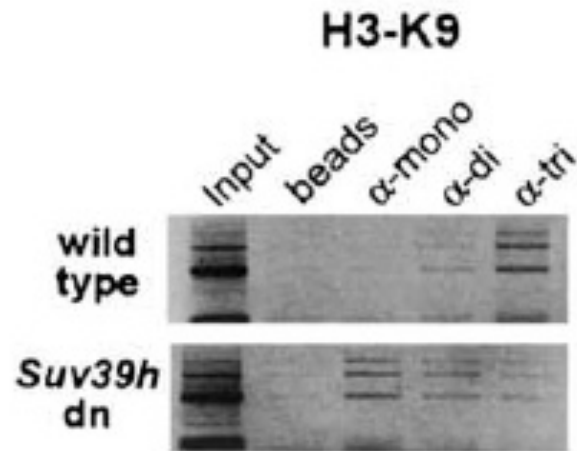
CHROMATIN IMMUNOPRECIPITATION (ChIP) COUPLED WITH PCR

→ H3K9me3 is enriched

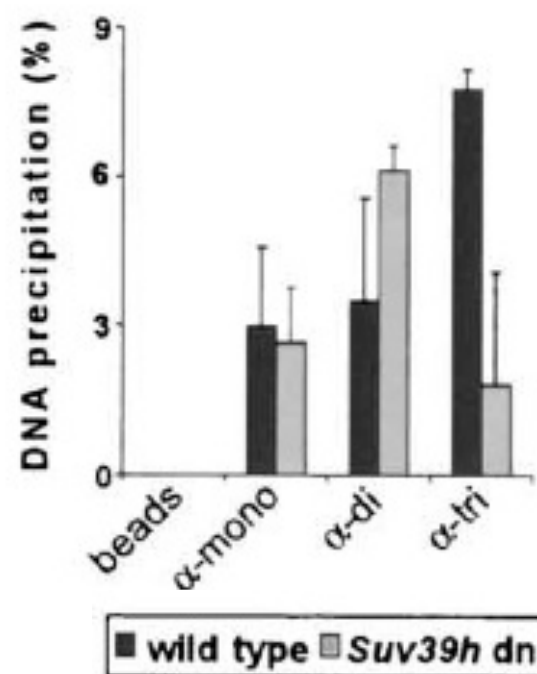
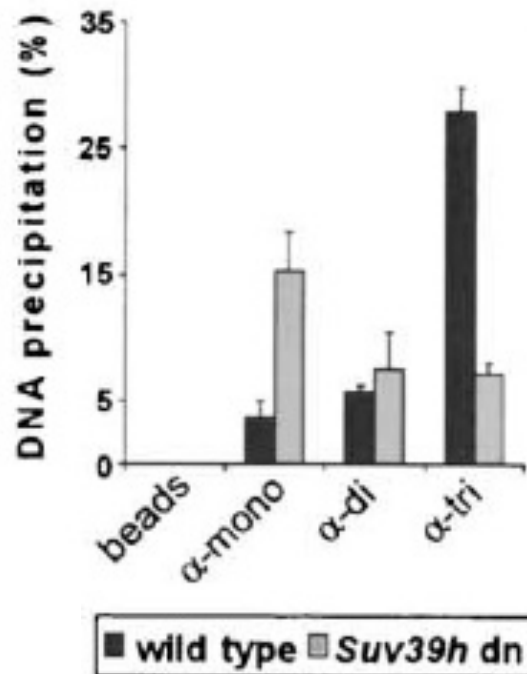
at pericentric (major+minor) repeats in mouse cells

Major satellite repeats

Minor satellite repeats

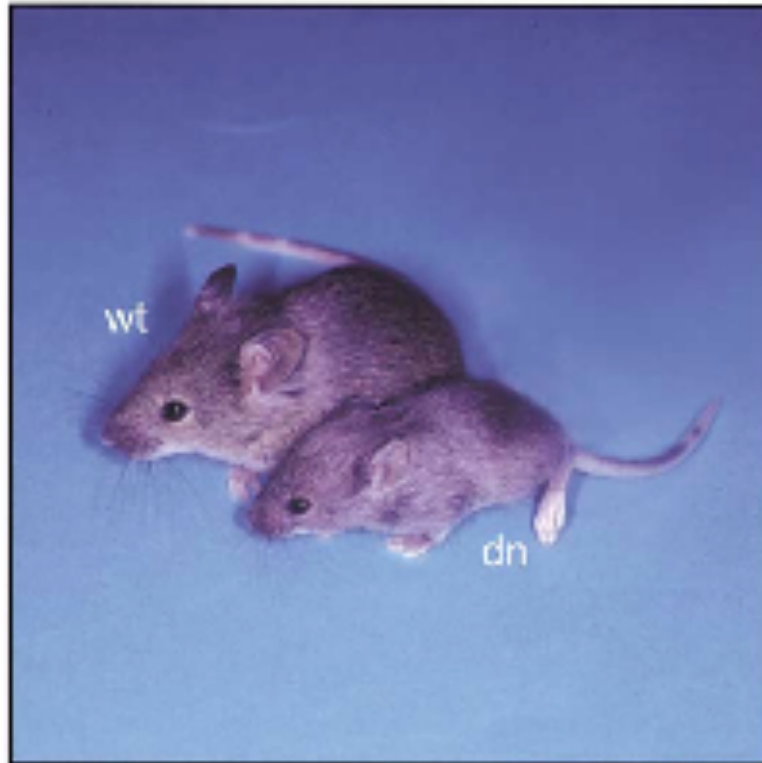


PCR amplification of major/minor satellite Repeats after ChIP using Antibodies that are specific for H3K9me1; H3K9me3; H3K9me3

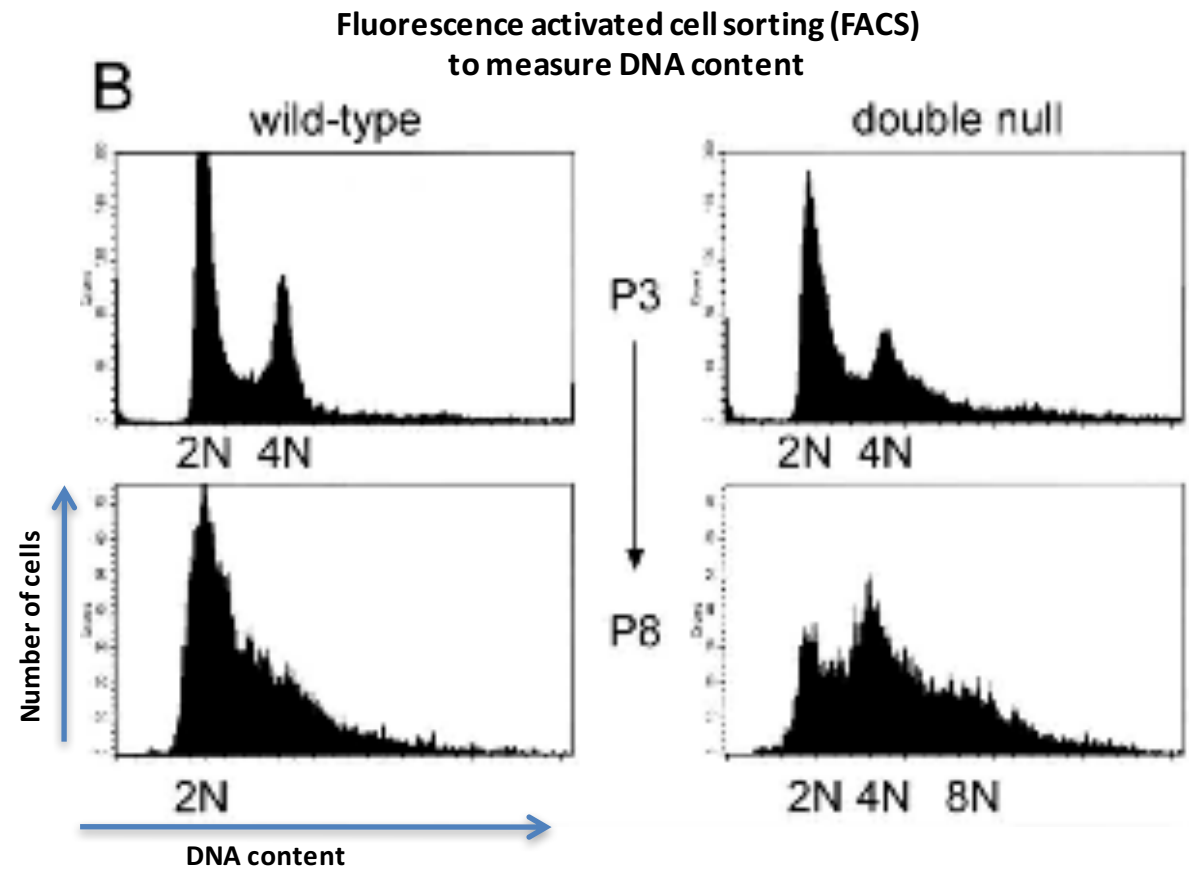


Suv39h1 is required for imposition of H3K9me3 at pericentric repeats

Lack of SUV39h HMTase activity results in genomic instability



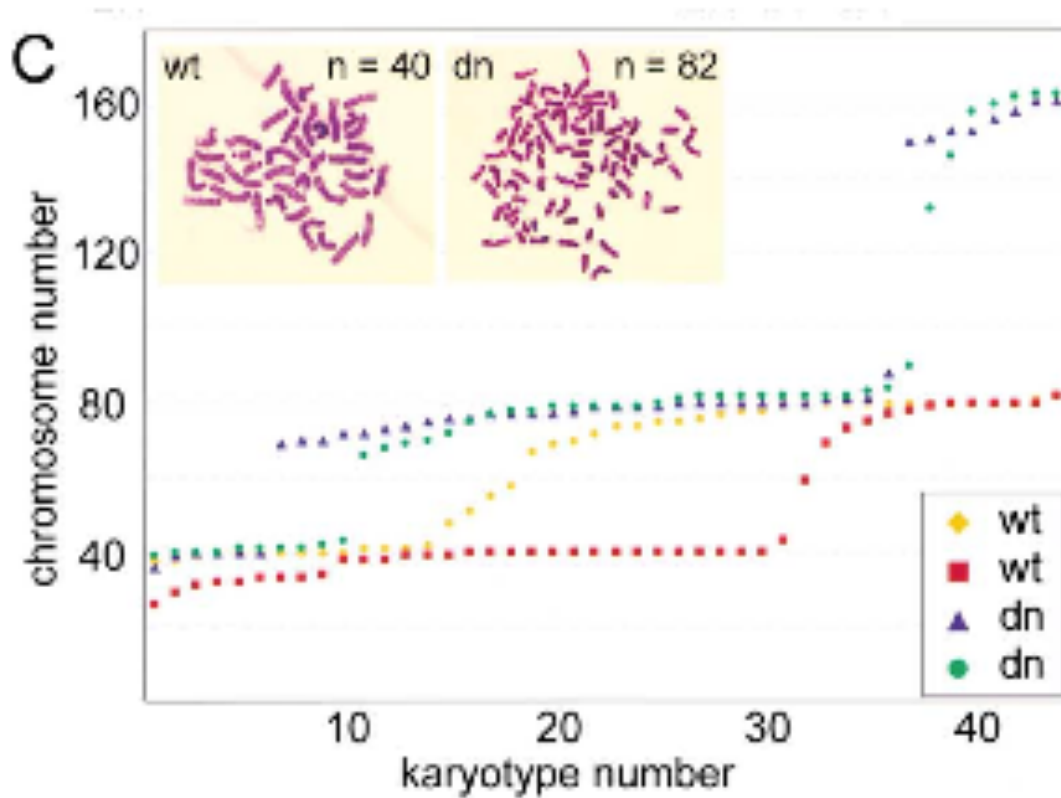
A knock-out model system
for Suv39h1 and Suv39h2
- Loss of Suv39h1/2: smaller body size



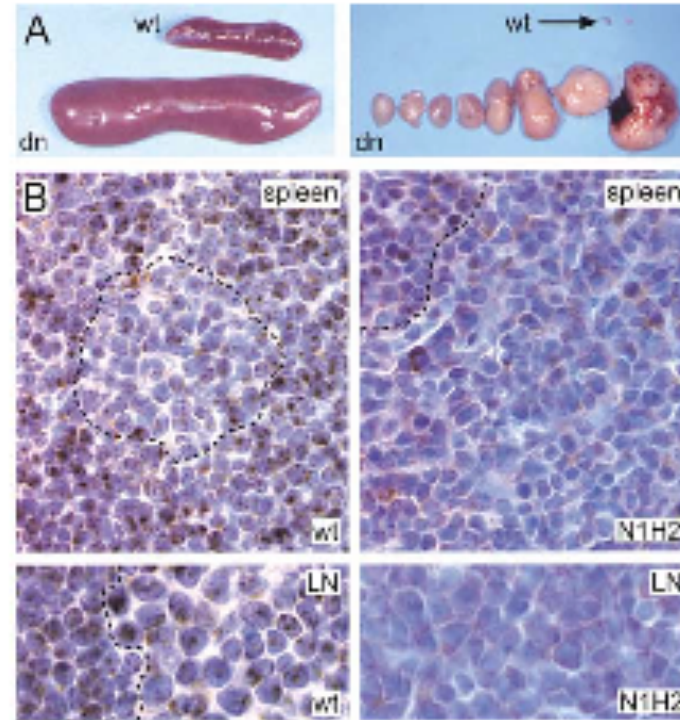
Fibroblasts from Suv39h1/2 null mice
are aneuploidy

Lack of SUV39h HMTase activity results in genomic instability

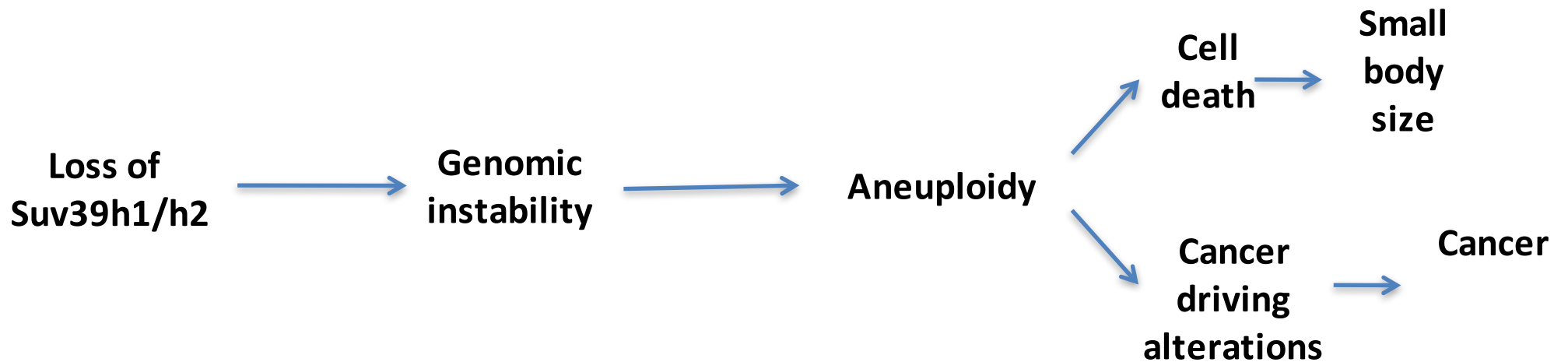
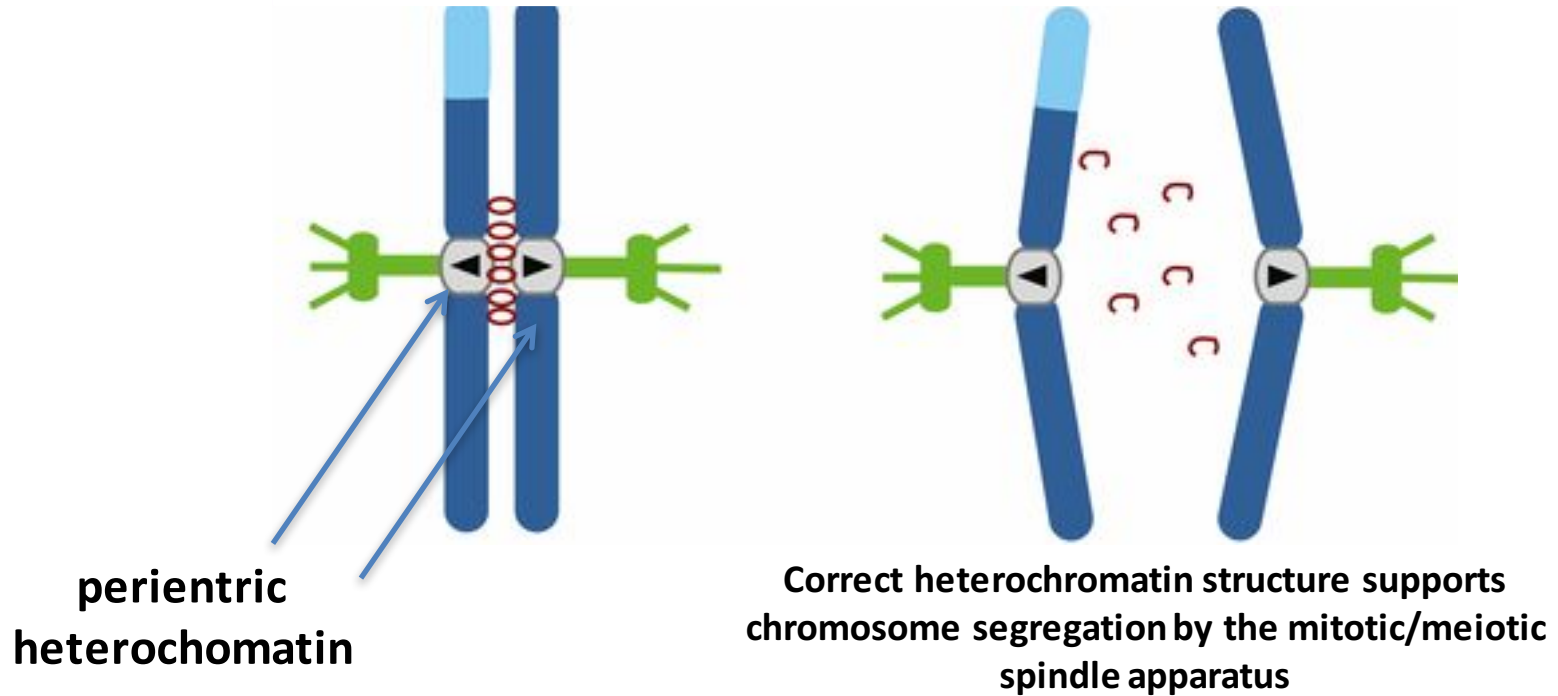
Loss of Suv39h1/2 results in increased chromosome numbers



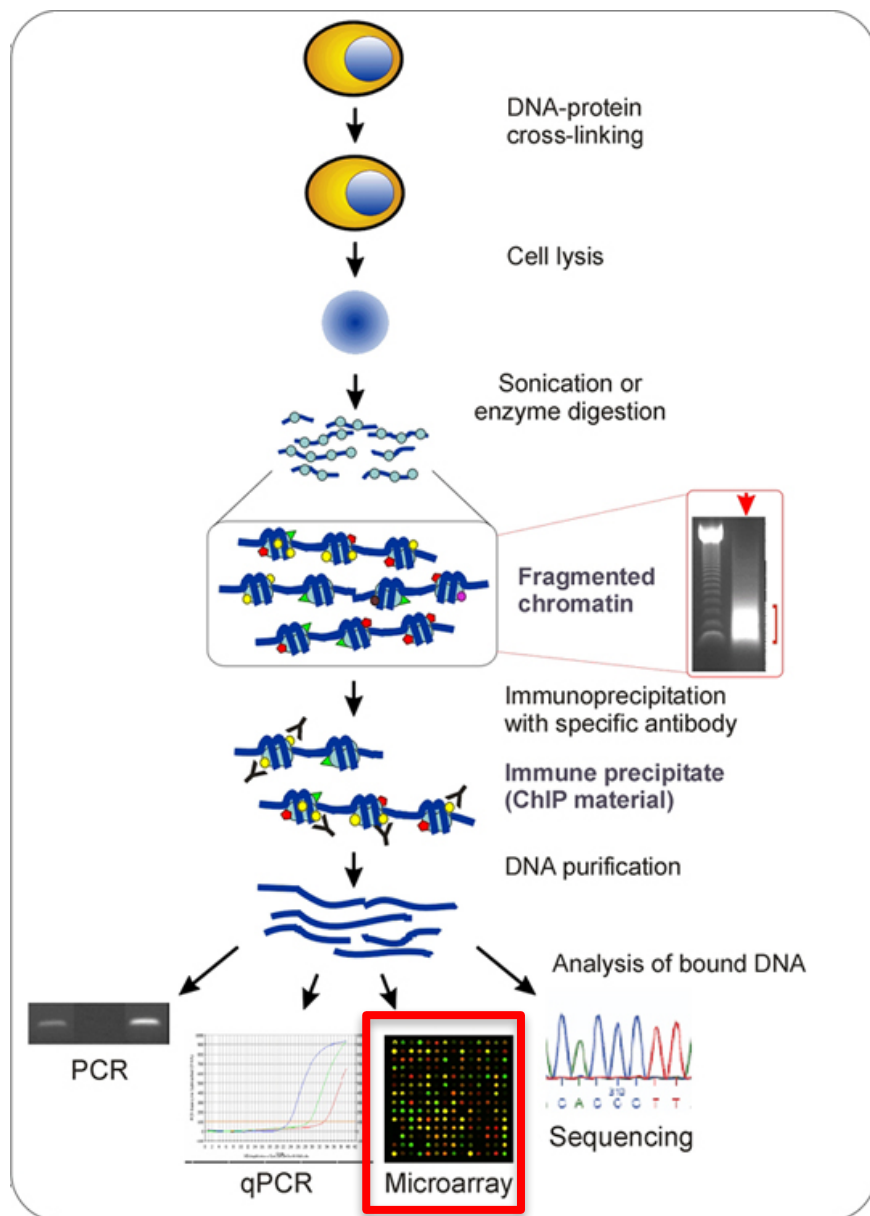
Genomic instability in Suv39h1/2 mice increases lymphomas



SUV39h HMTase activity is essential for fidelity in mitosis/meiosis



CHROMATIN IMMUNOPRECIPITATION (ChIP) → DETAILED ANALYSIS → Localization of protein at a defined region - sequence



Cell model system:

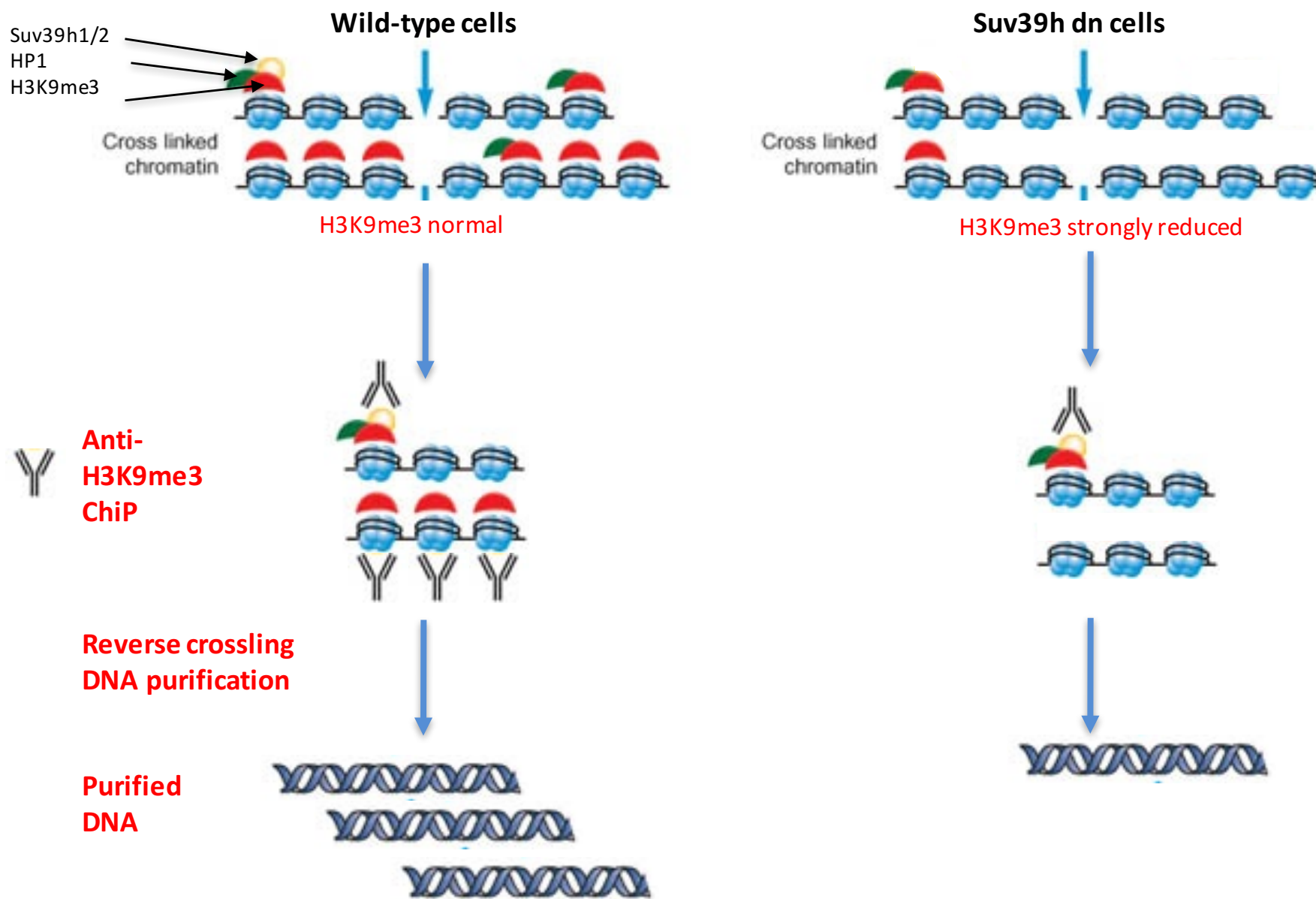
i.e. **Wild-type or Suv39 dn cells**
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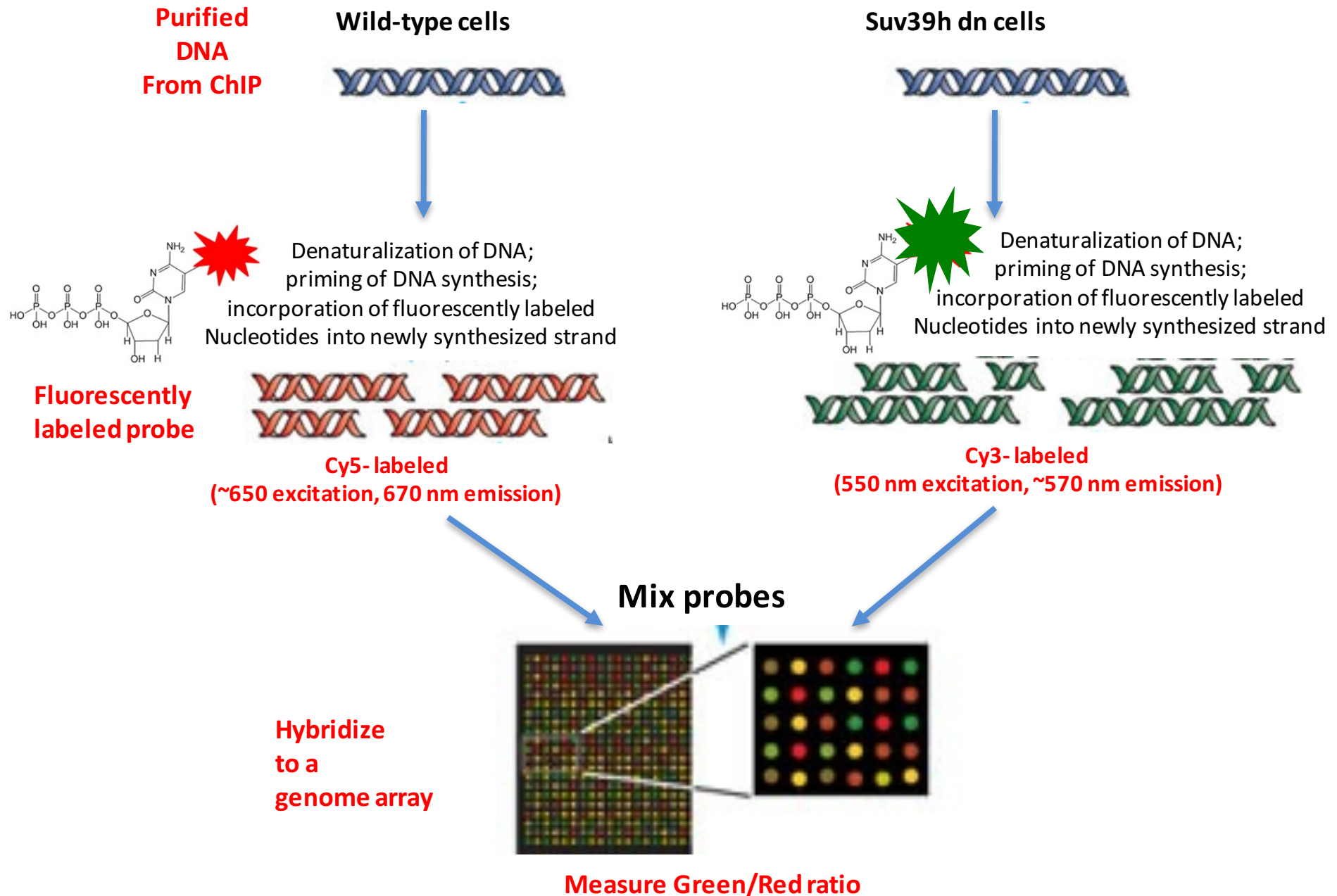
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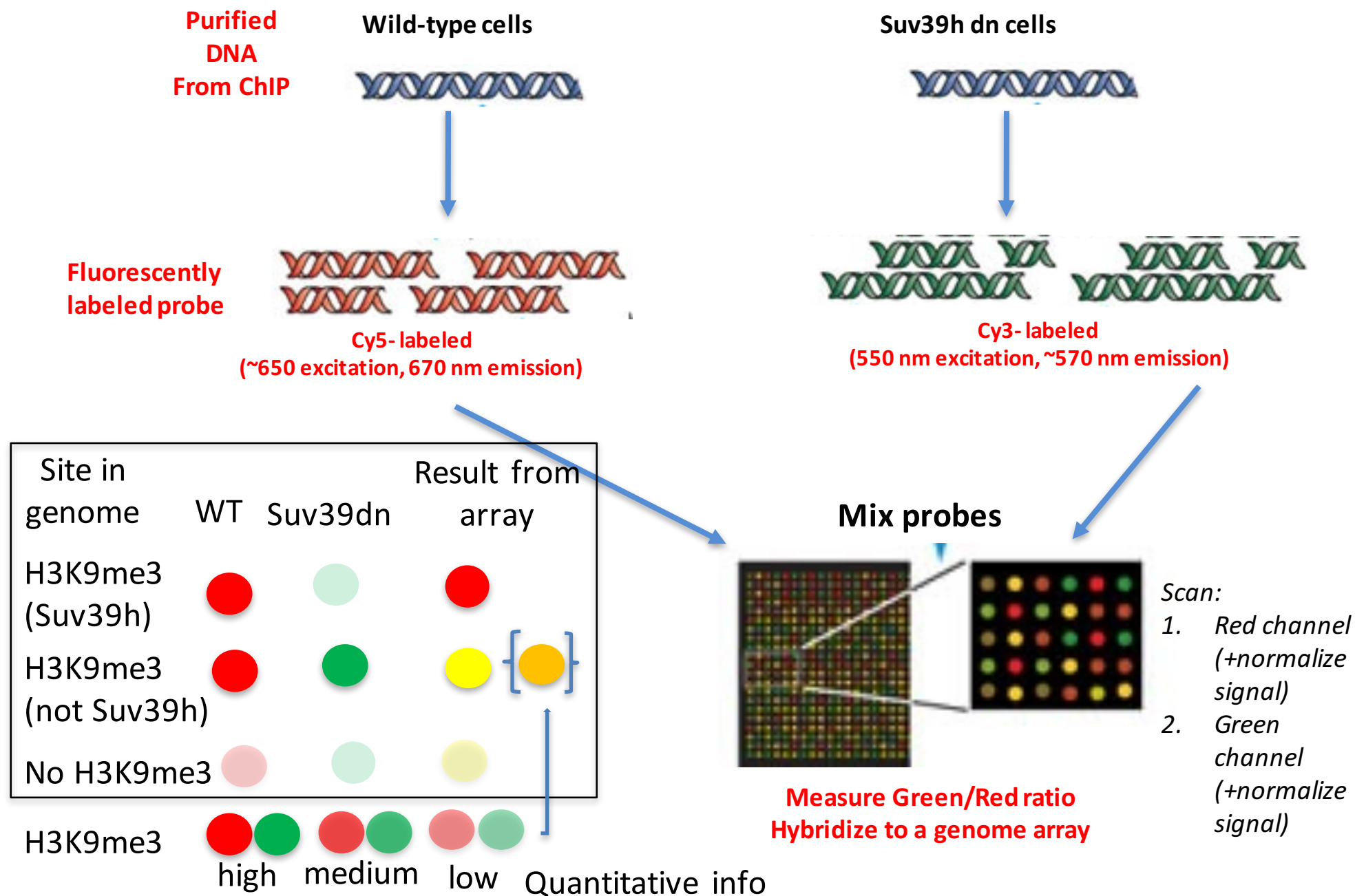
ChIP on ChIP: Analysis of epigenetic information across a high number of genomic sites in the genome



ChIP on ChIP: Analysis of epigenetic information across a high number of genomic sites in the genome



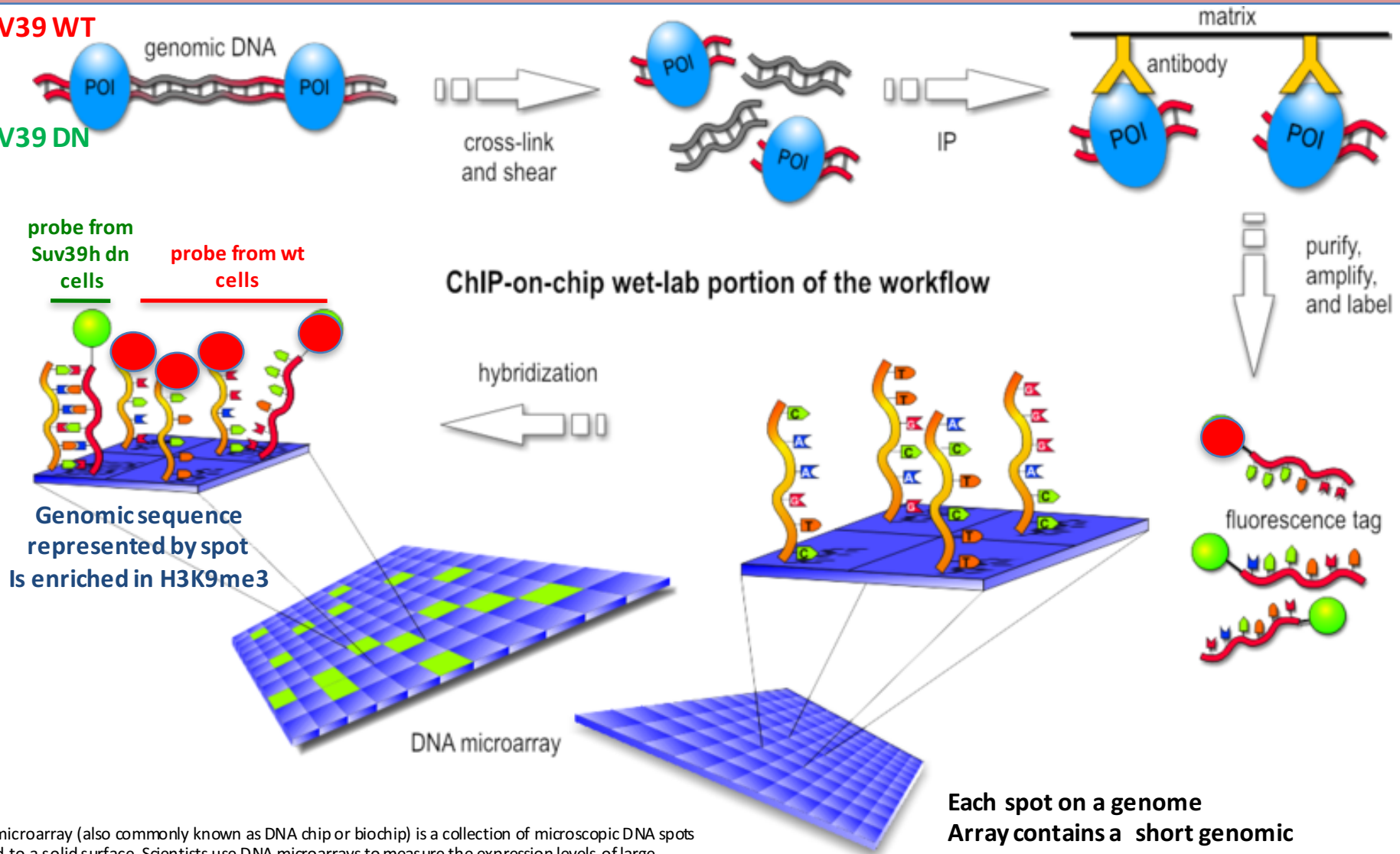
ChIP on ChIP: Analysis of epigenetic information across a high number of genomic sites in the genome



ChIP on ChIP: Analysis of epigenetic information across a high number of genomic sites in the genome

1. SUV39 WT

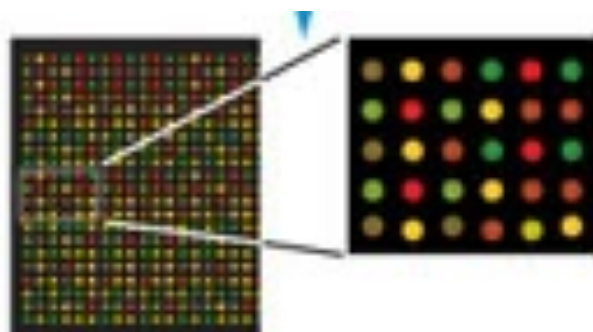
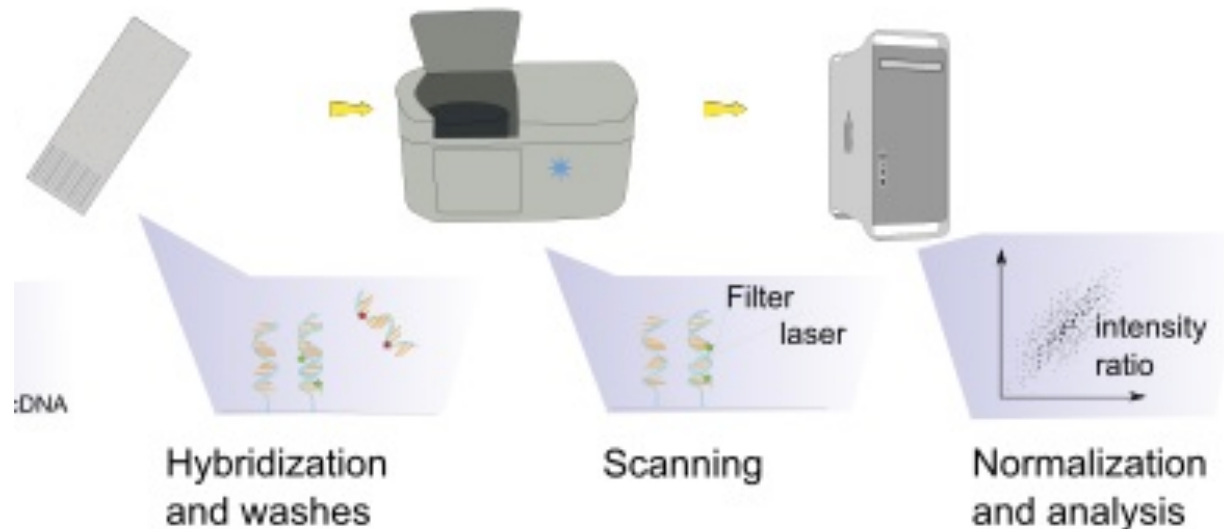
2. SUV39 DN



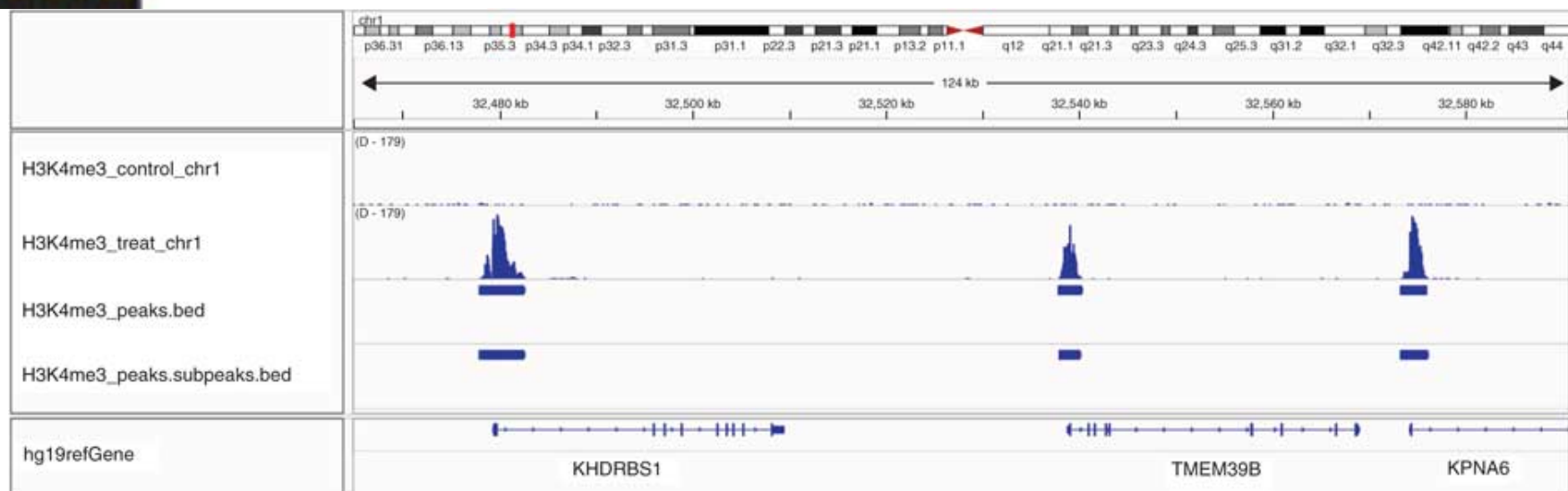
A DNA microarray (also commonly known as DNA chip or biochip) is a collection of microscopic DNA spots attached to a solid surface. Scientists use DNA microarrays to measure the expression levels of large numbers of genes simultaneously or to genotype multiple regions of a genome. Each DNA spot contains picomoles (10⁻¹² moles) of a specific DNA sequence, known as probes (or reporters or oligos). These can be a short section of a gene or other DNA element that are used to hybridize a cDNA or rRNA (also called anti-sense RNA) sample (called target) under high-stringency conditions. Probe-target hybridization is usually detected and quantified by detection of fluorophore-, silver-, or chemiluminescence-labeled targets to determine relative abundance of nucleic acid sequences in the target

Each spot on a genome
Array contains a short genomic
Sequence. Genome arrays cover
a large part of the
Genome
A ChIP holds ca 20000 spots

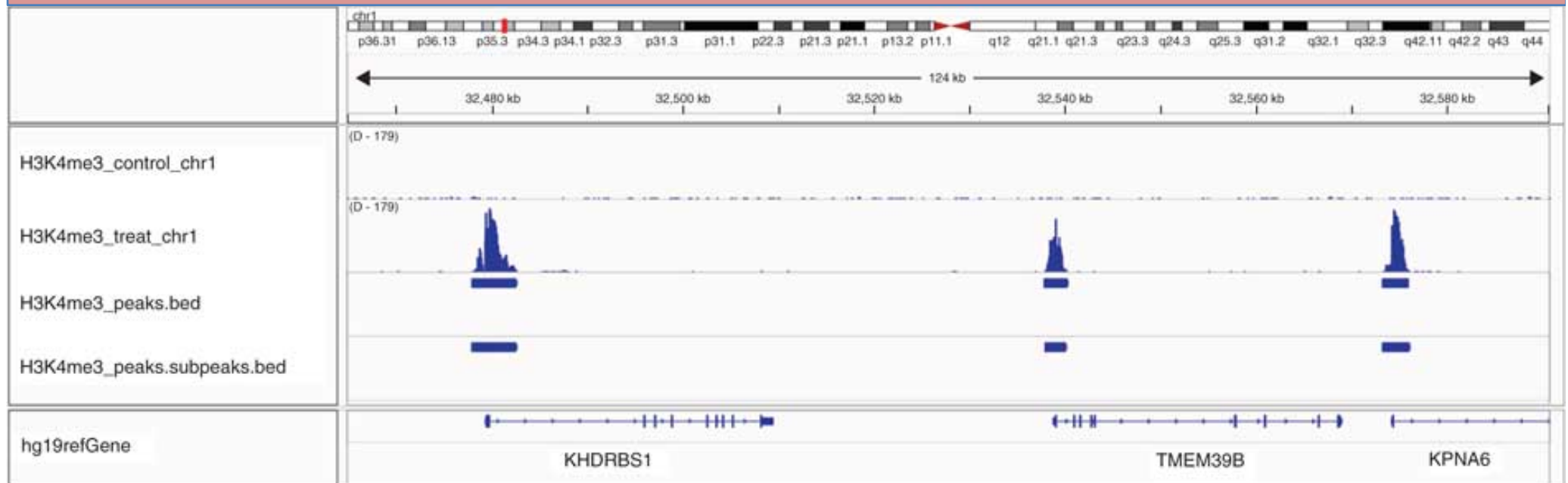
ChIP on CHIP: Analysis of epigenetic information across a high number of genomic sites in the genome



Arrays do not contain the entire human/mouse genome
 Are enriched for particular sequences according to need
 (promoter, enhancer, etc...)



ChIP on ChIP: Analysis of epigenetic information across a high number of genomic sites in the genome

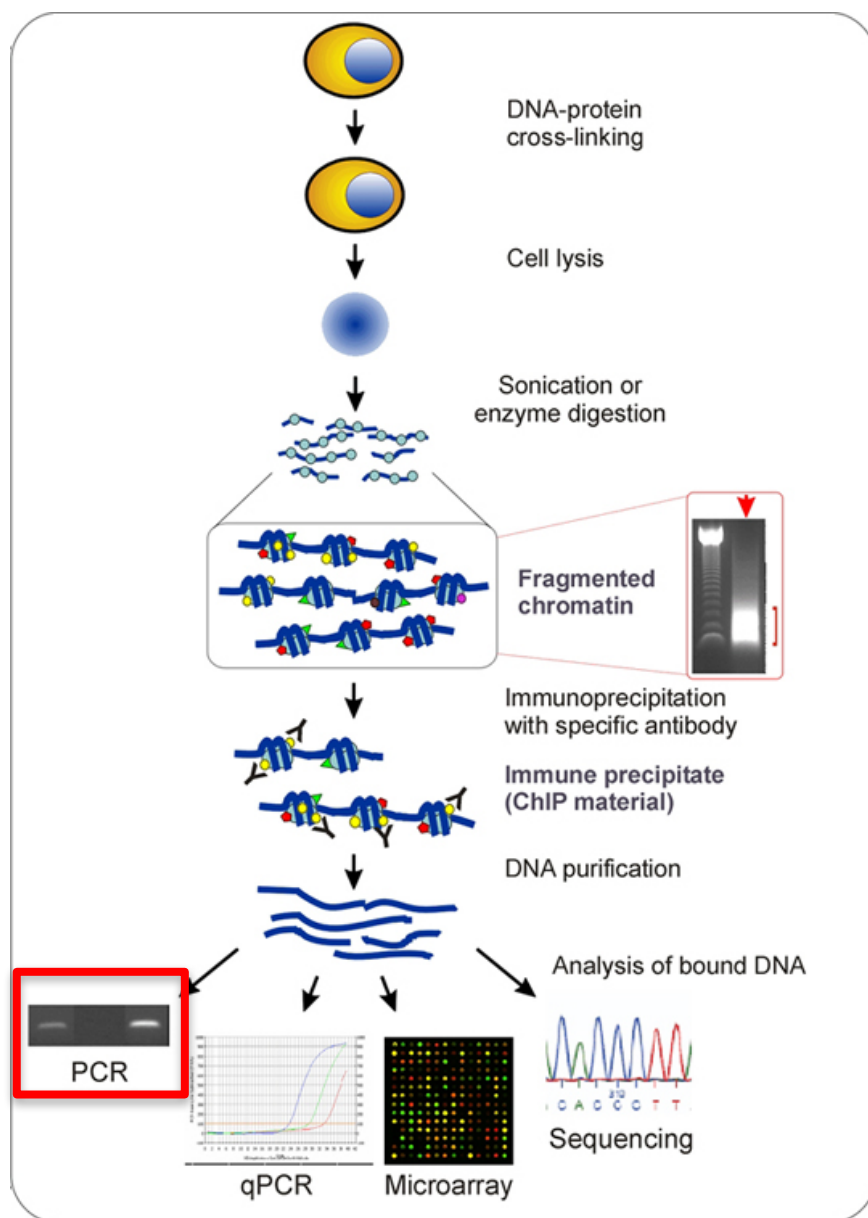


Advantage: low tech, cheap

Disadvantage: low resolution, no data on number of molecules – just proportions;
laborious to reach a good genome coverage

Already outdated → state of the art: **ChIP seq**

ChIP seq: Analysis of epigenetic information on the single nucleotide level → GENERATION OF GENOME WIDE EPIGENETIC MAPS



Cell model system:

i.e. **Wild-type or Suv39 dn cells**
that grow in cell culture dish

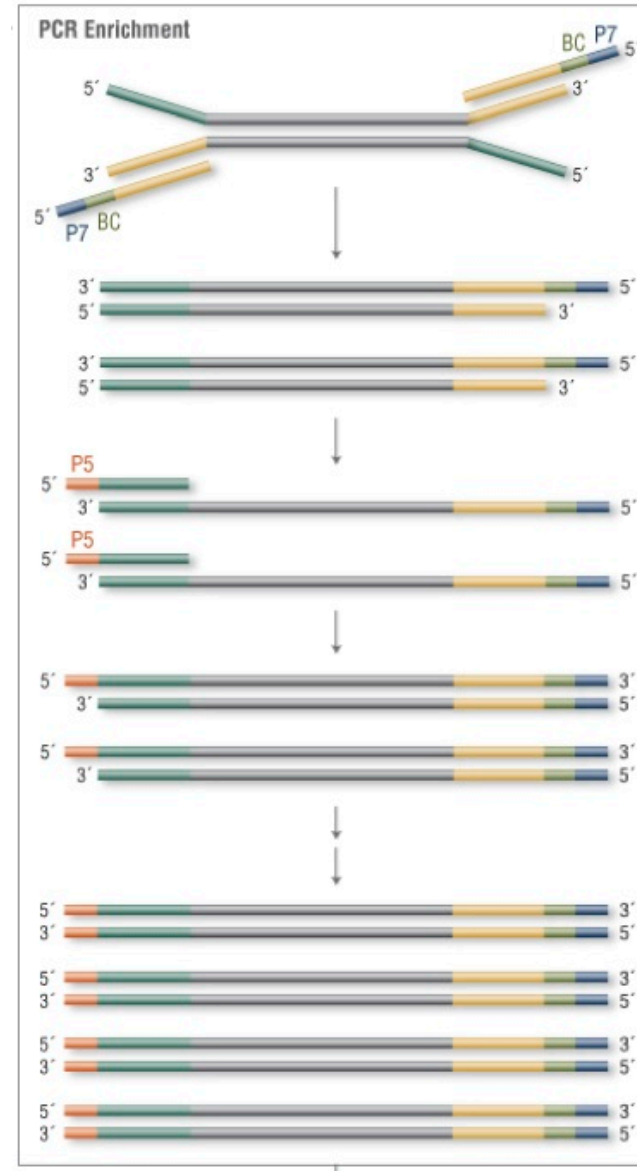
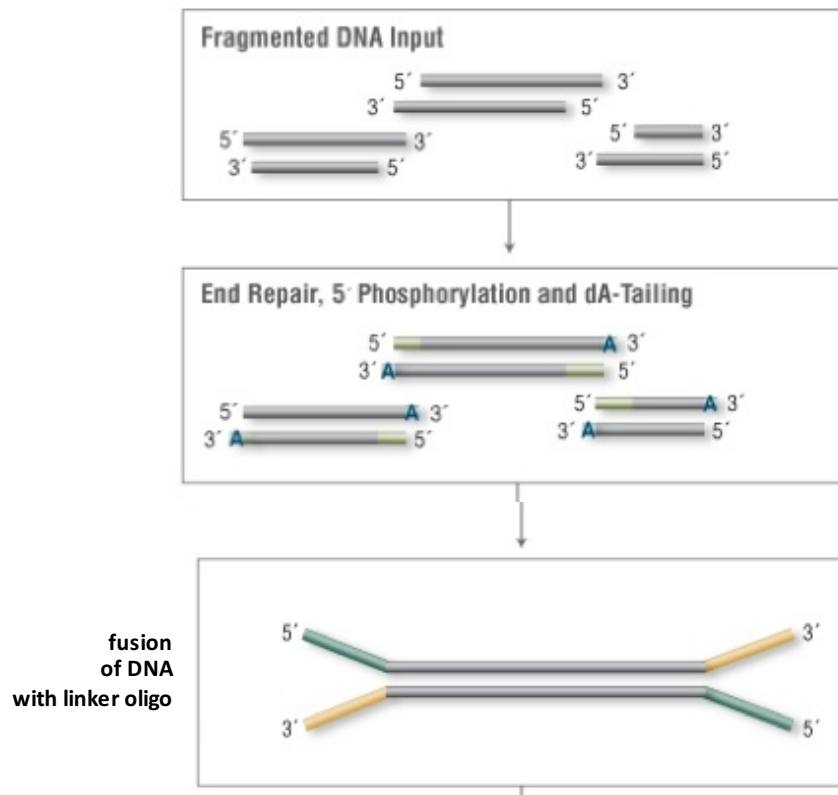
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In control versus Suv39h dn cells

METHODS:

Massive parallel sequencing of immunoprecipitated DNA
Permits to obtain epigenetic information on the single nucleotide level

ChIP seq: Analysis of epigenetic information on the single nucleotide level → GENERATION OF GENOME WIDE EPIGENETIC MAPS

Use immunoprecipitated DNA to generate DNA libraries:



Linkers serve as uniform primer binding sites. This allows the amplification of the entire DNA library using only 2 types of oligonucleotides

Amplified library

READY FOR MASSIVE PARALLEL SEQUENCING

ChIP seq: Analysis of epigenetic information on the single nucleotide level → GENERATION OF GENOME WIDE EPIGENETIC MAPS

Illumina Massively Parallel Sequencing

<https://www.illumina.com/company/video-hub/pfZp5Vgsbw0.html>

HiSeq 2000



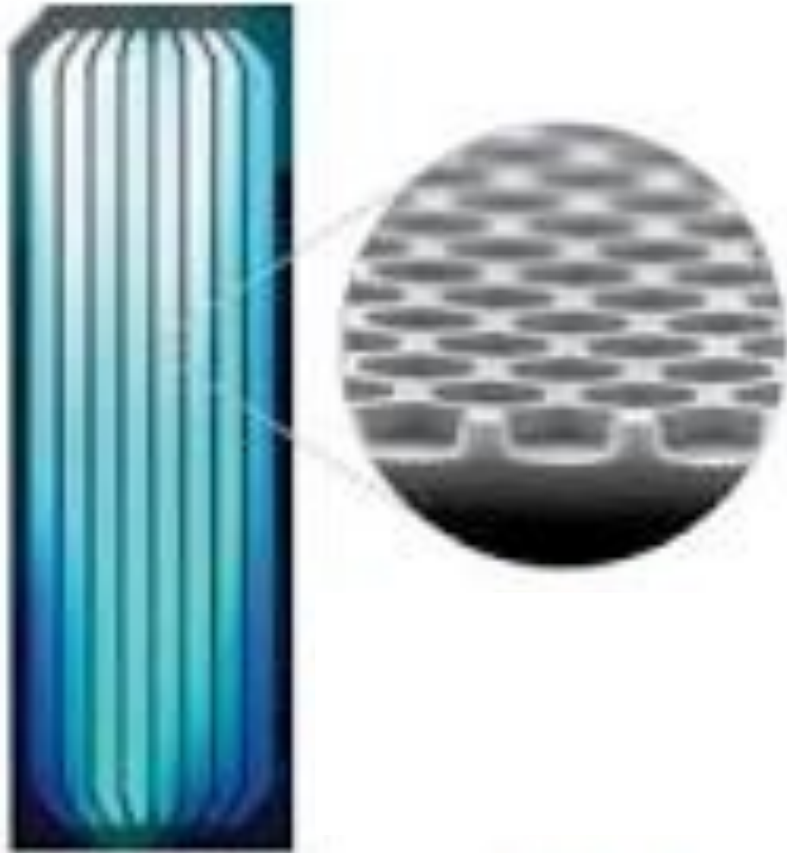
The heart of the Illumina Massive Parallel Sequencer is the “FLOW-CELL”. A surface with millions of small wells that allow thousands of Sanger-sequencing reaction in parallel = “massive parallel sequencing”. In each well a SINGLE MOLECULE of DNA is amplified and sequenced

Illumina offers the most potent massive sequencing instruments – leader on the market

<https://www.youtube.com/watch?v=pfZp5Vgsbw0>

ChIP seq: Analysis of epigenetic information on the single nucleotide level
→ GENERATION OF GENOME WIDE EPIGENETIC MAPS

CLUSTER AMPLIFICATION:



Flow cell contains surface with millions of wells

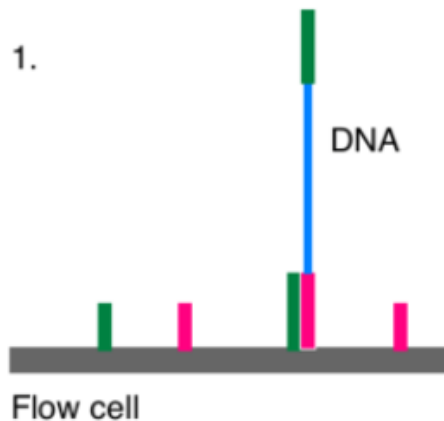
→ Each well contains beads mounted with 2 species of oligonucleotides that hybridize with adaptor oligos of DNA library

→ DNA library will be loaded onto the flow cell in a determined concentration:

ONLY ONE MOLECULE OF DNA WILL BE PROCESSED FOR SEQUENCING IN A SINGLE WELL

CLUSTER AMPLIFICATION:

- making DNA library (~300bp fragments)
- ligation of adapters **A** and **B** to the fragments



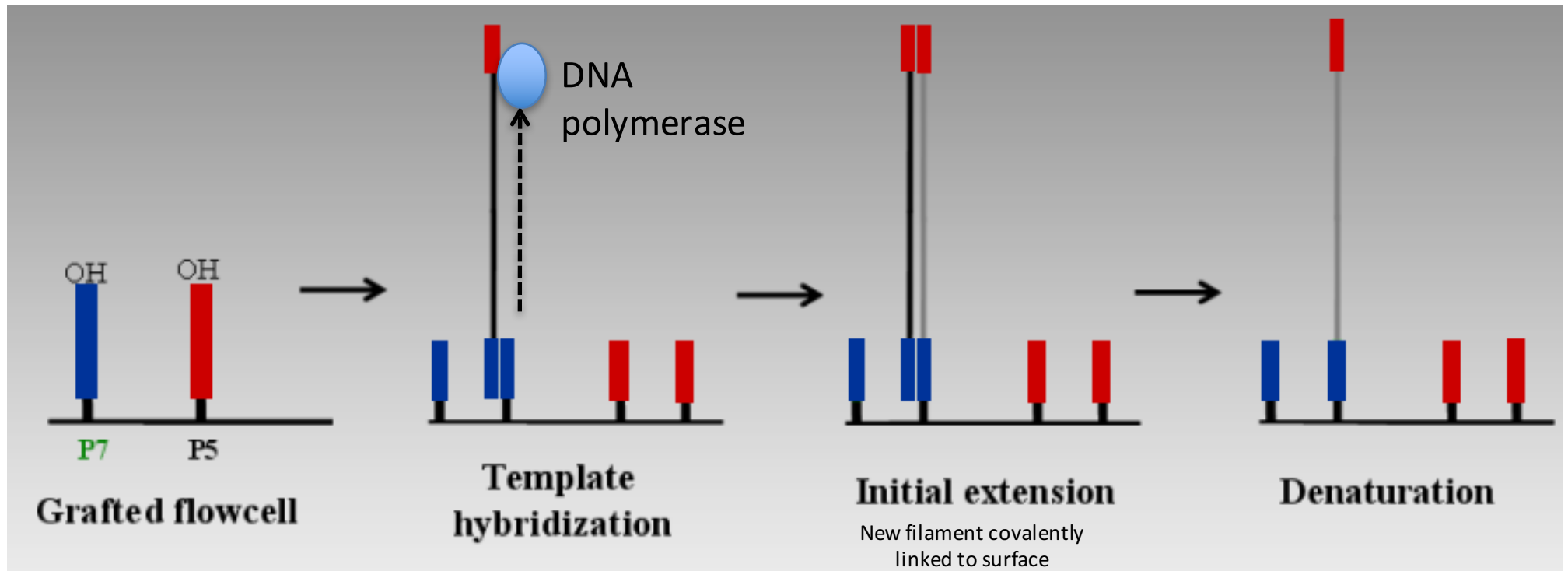
1 well, covered with millions of 2 types of oligos

1 well in a flow-cell with billions of wells

- **complementary** primers are ligated to the surface
- pairing with ChIP ed ssDNA at random position in the well of the flow cell

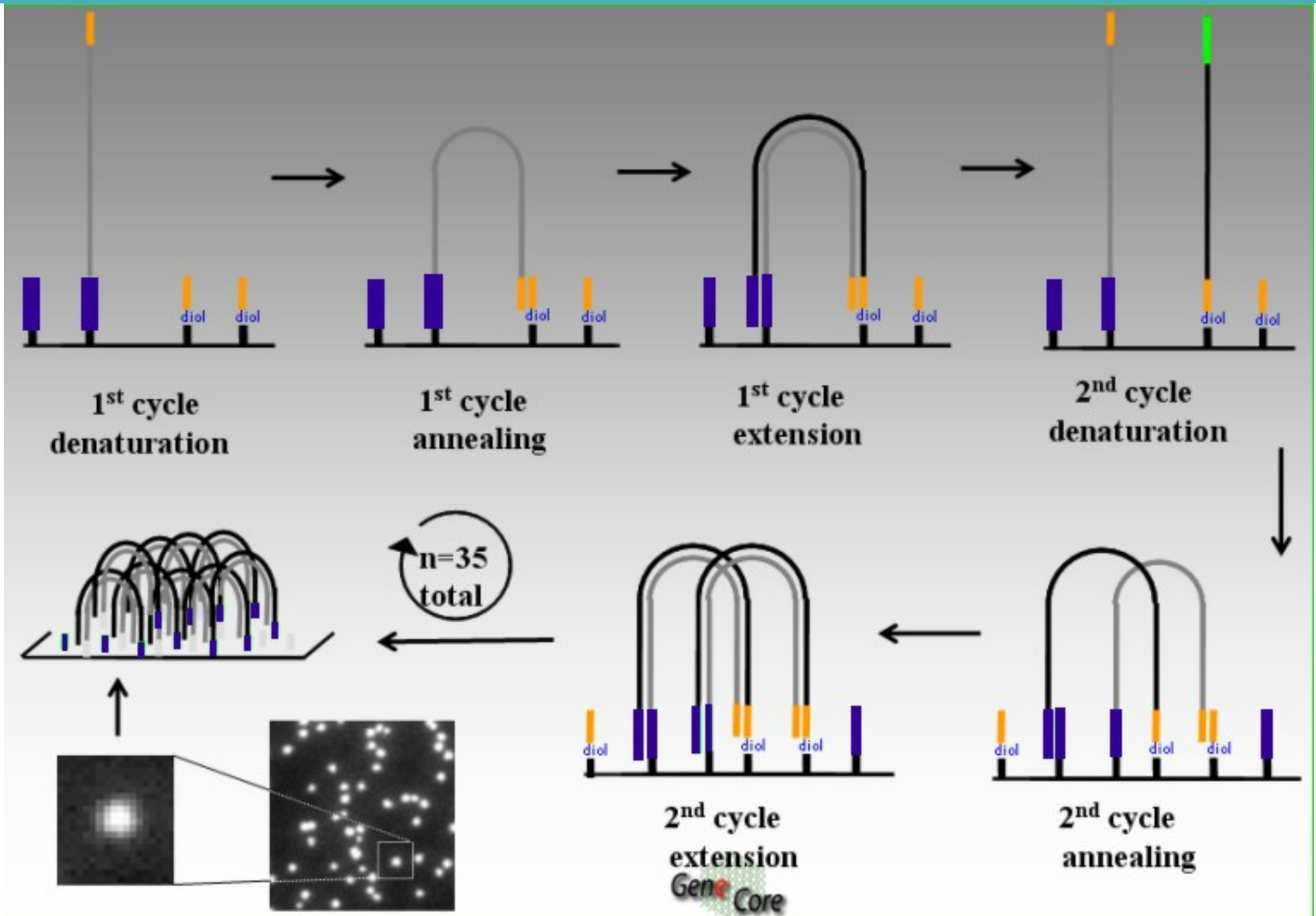
CLUSTER AMPLIFICATION:

Bridge amplification: takes place on surface of beads (each bead is mounted with 2 species of oligos; each oligo can hybridize to a DNA library fragment):
initiation



On the surface: complementary oligos

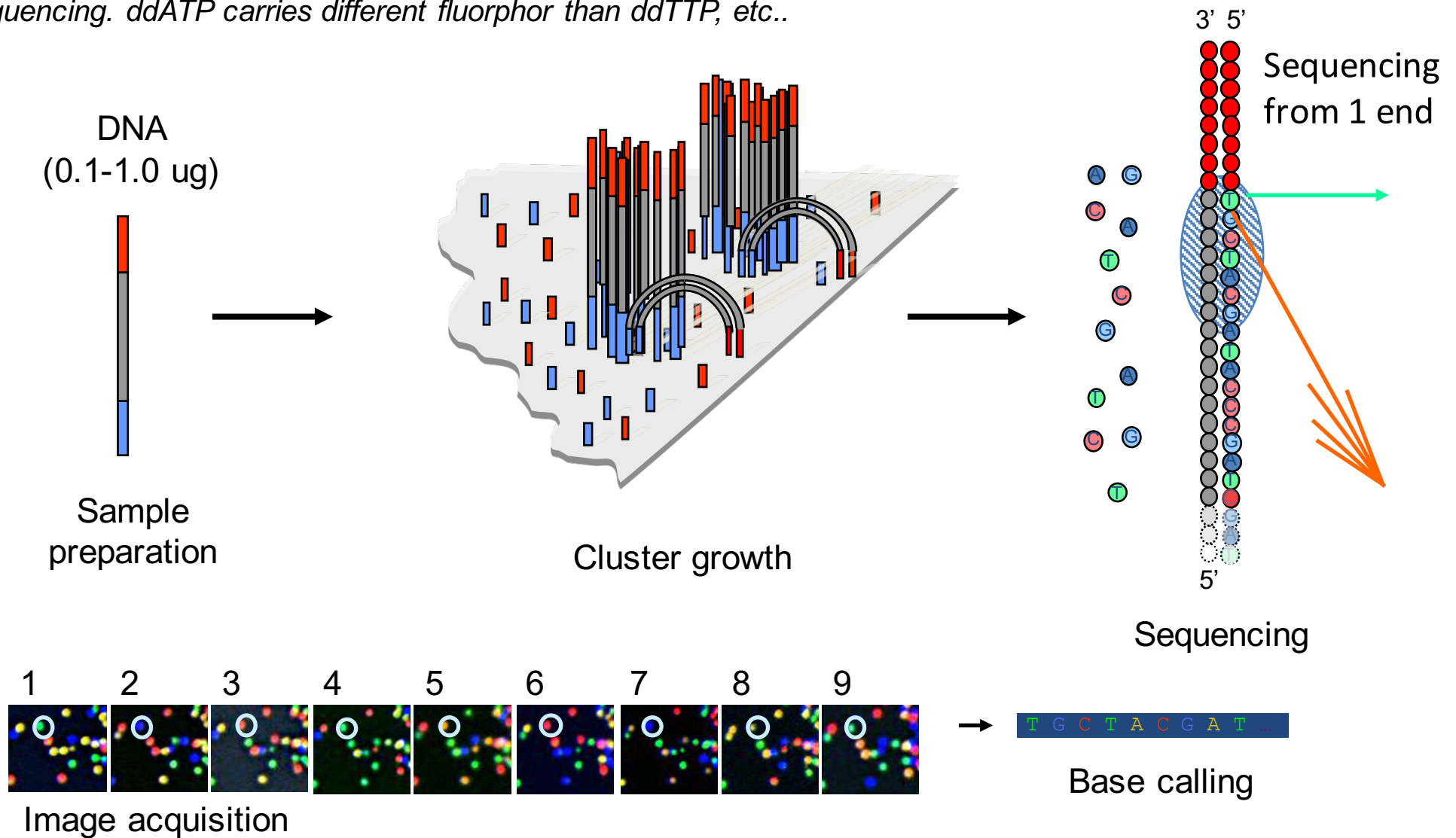
CLUSTER AMPLIFICATION:



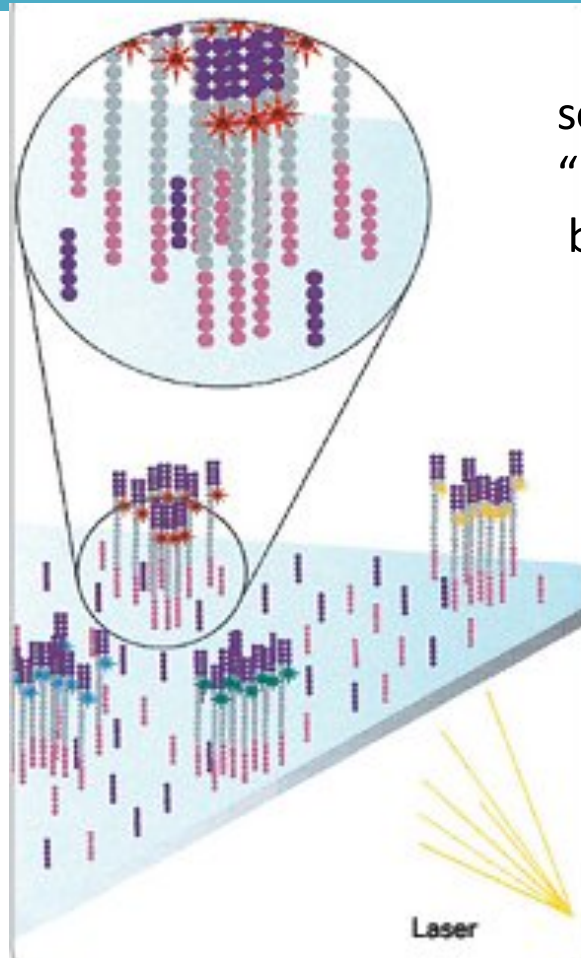
Illumina Sequencing Technology

Robust Reversible Terminator Chemistry Foundation

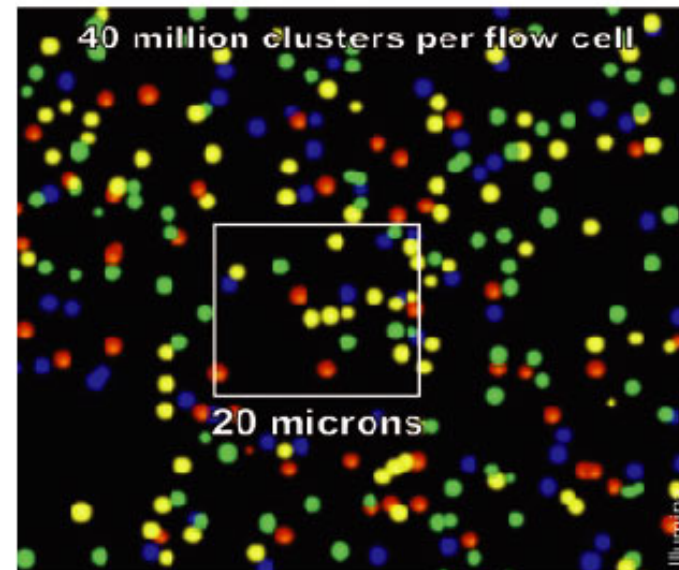
In each round of sequencing a fluorescently labelled ddNTP will be used for sequencing. ddATP carries different fluorophore than ddTTP, etc..



Illumina: massive parallel sequencing:



sequencing by synthesis:
“reversible terminator” nucleotides
blocked + fluorescently labeled

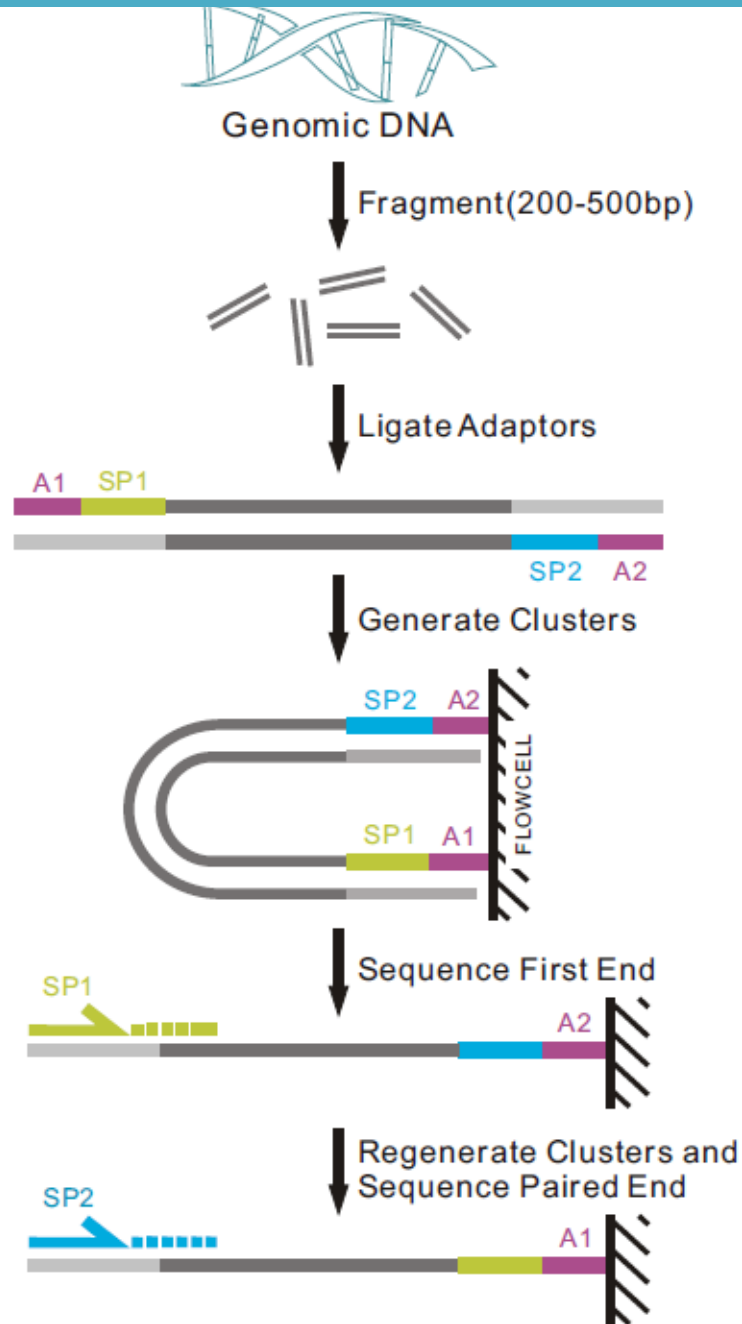


1. Synthesis = incorporation of fluorescent nucleotide: blocking synthesis
2. 4. Scanning of fluorescent signal
3. dye cleavage + elimination
4. wash step

1. Synthesis = incorporation of fluorescent nucleotide: blocking synthesis

READ LENGTH: ca: 150nt from each primer (2x150nt = 300nt)

Illumina: paired end sequencing increases information content



After 1° strand sequencing, A1 anneals to A1 in nanowell-->DNA synthesis --> template strand cleaved off → new strand sequenced

<https://www.youtube.com/watch?v=9YxExTSwgPM>

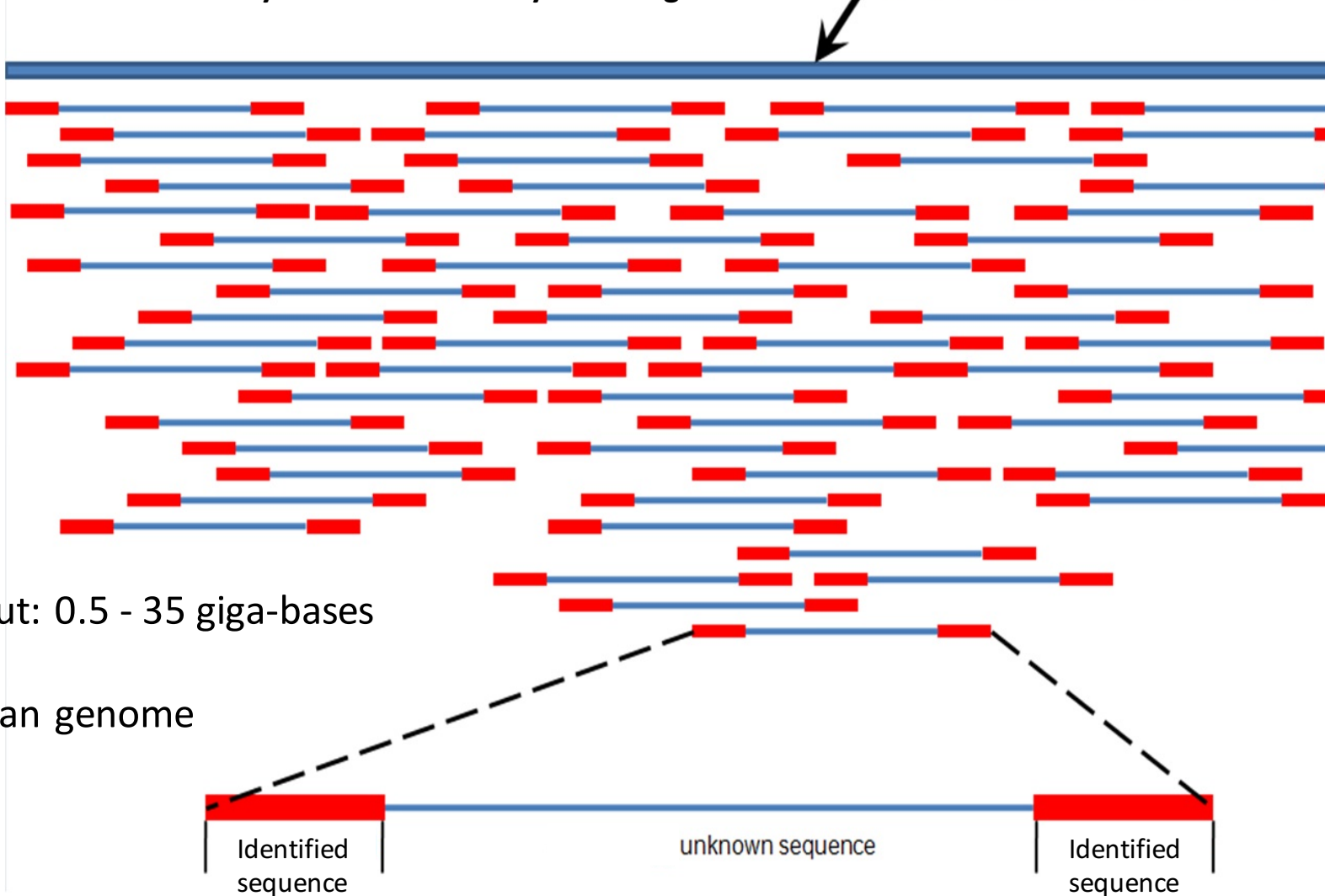
Figure 1-2-1 Pipeline of paired-end sequencing (www.illumina.com)

Data analysis: obtained sequence reads are aligned along genomic DNA sequence → high number of reads necessary to obtain full sequence coverage

Read length: 50 – max. 300 nt

Read does not necessarily cover entire library DNA fragment

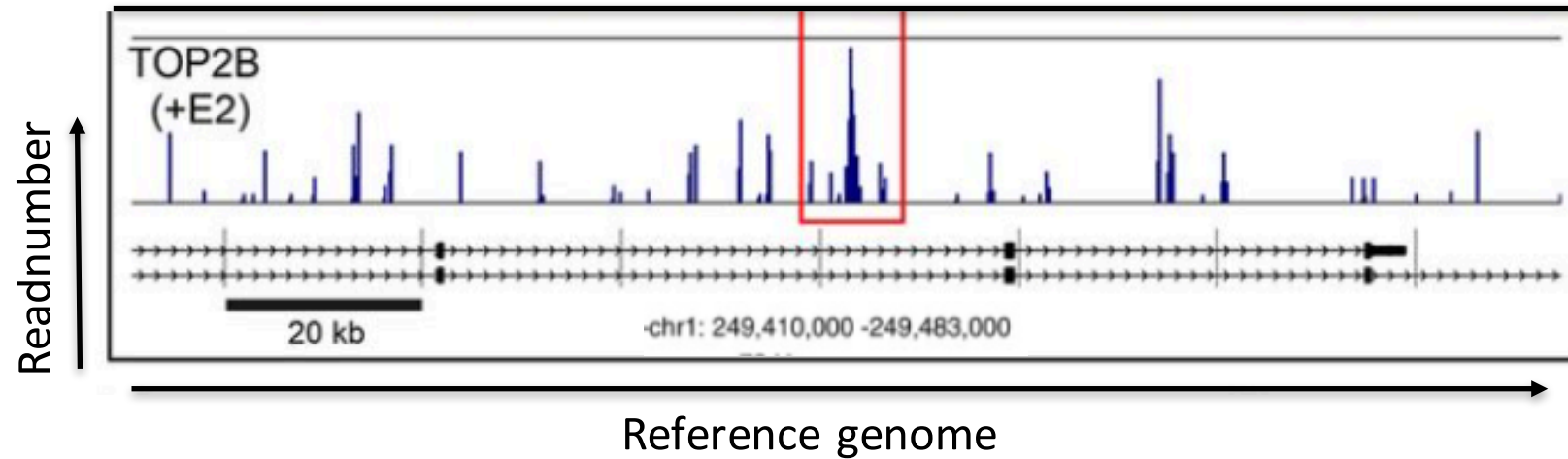
Reference Genome Sequence



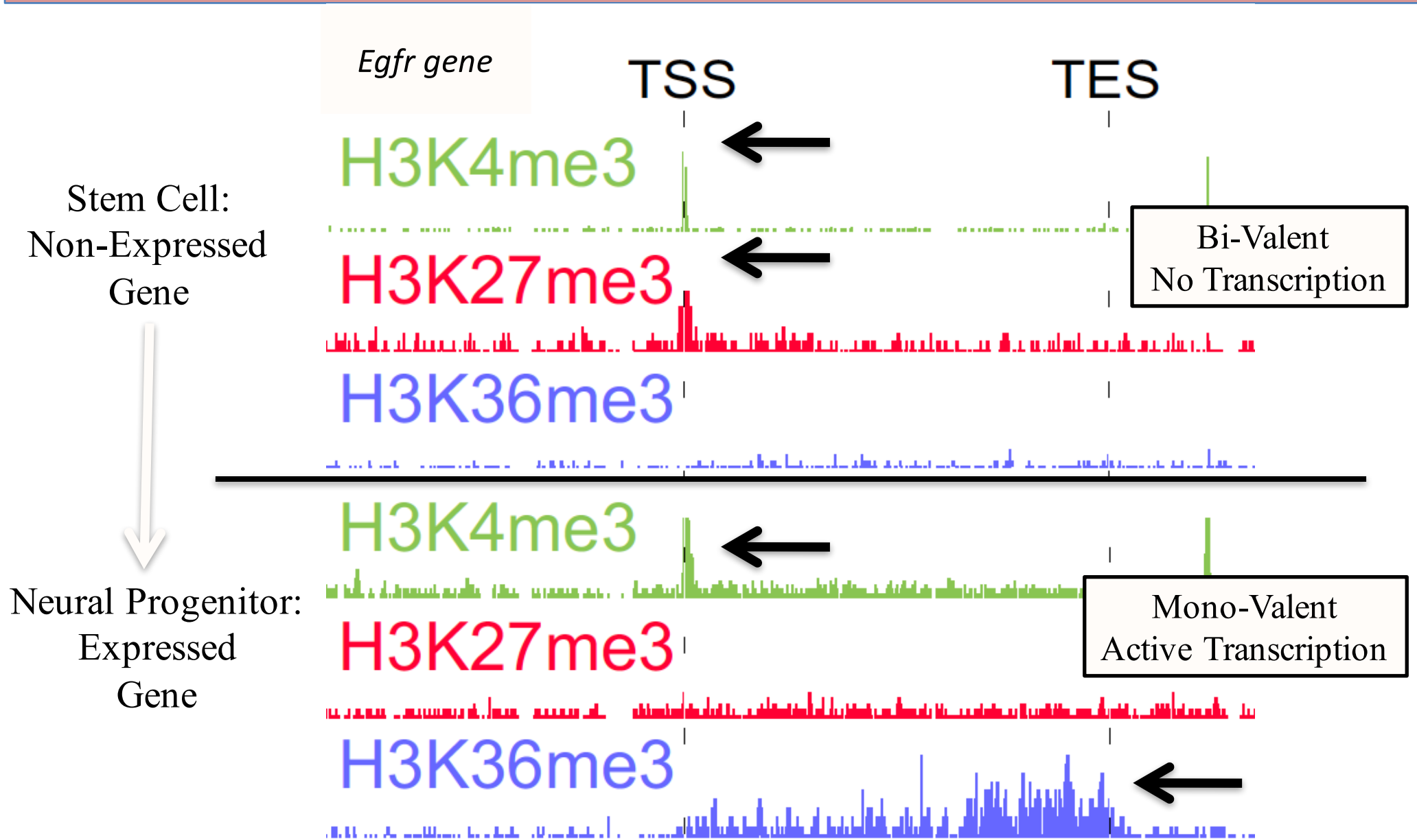
Max. output: 0.5 - 35 giga-bases
= 3.5×10^{10}
= 10x human genome

Sequence derived from one amplified cluster

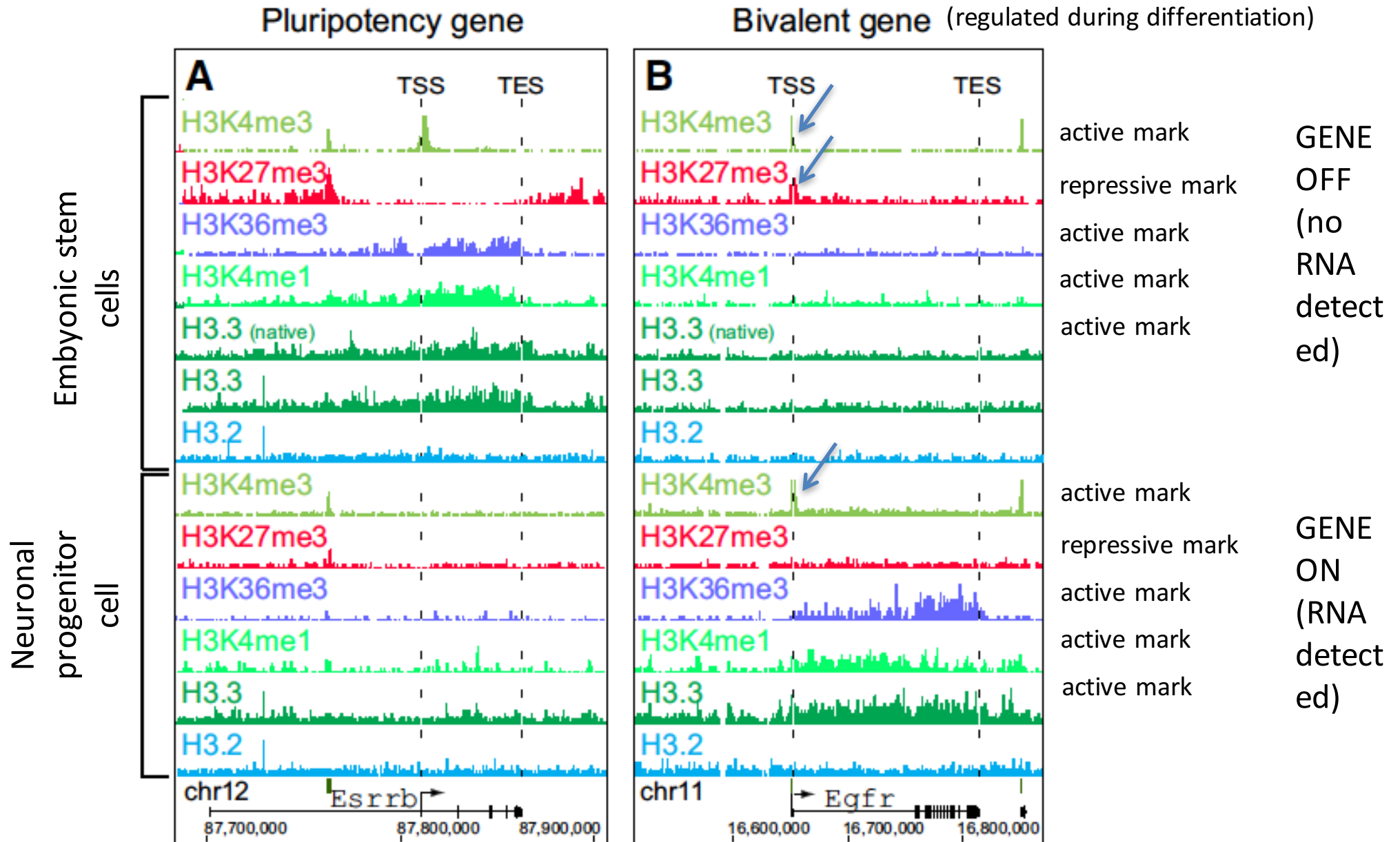
Data analysis: obtained sequence reads are aligned along genomic DNA sequence → high number of reads necessary to obtain full sequence coverage



BIOINFORMATICS ANALYSIS: Mapping ChIP seq reads agins the human genomic sequence

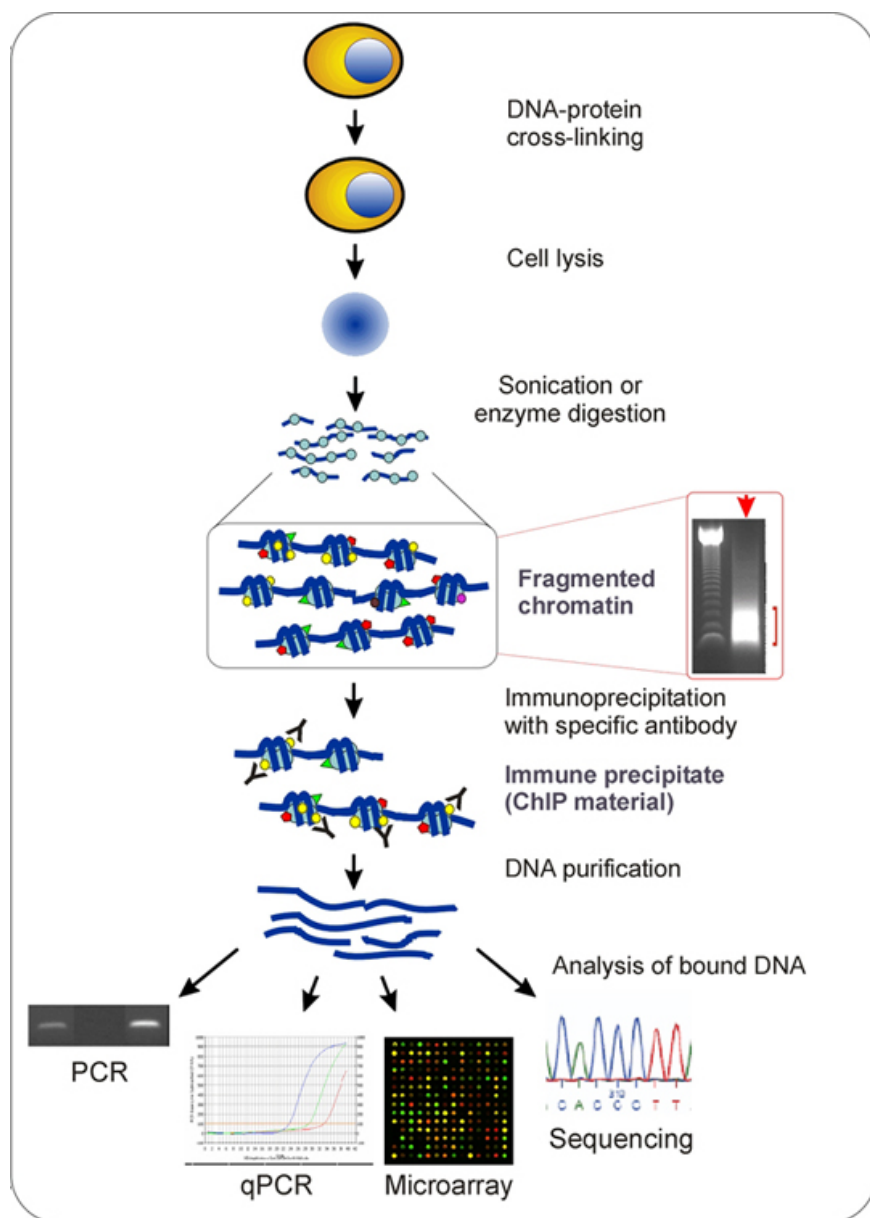


Mapping the epigenetic landscape enables to define “key rules” to define the epigenetic code of active and silent genes



CHROMATIN IMMUNOPRECIPITATION → DETAILED ANALYSIS

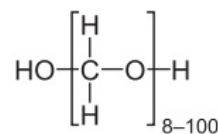
→ Localization of protein at a defined region - sequence



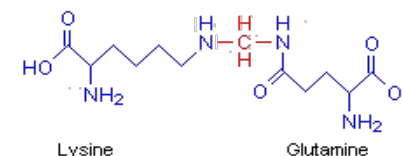
Cell model system:

i.e. Wild-type or Suv39 dn cells that grow in cell culture dish

1. Crosslink chromatin (treatment of cells with Paraformaldehyde)



Paraformaldehyde (PFA) is a polymer of formaldehyde, and covalently links NH₂ groups of biomolecules



Example of crosslinking reaction

2. Sonicate crosslinked cells

→ DNA breaks in a random manner until reaching a size of ca. 250 – 500nt

3. Incubate chromatin fragments with antibodies

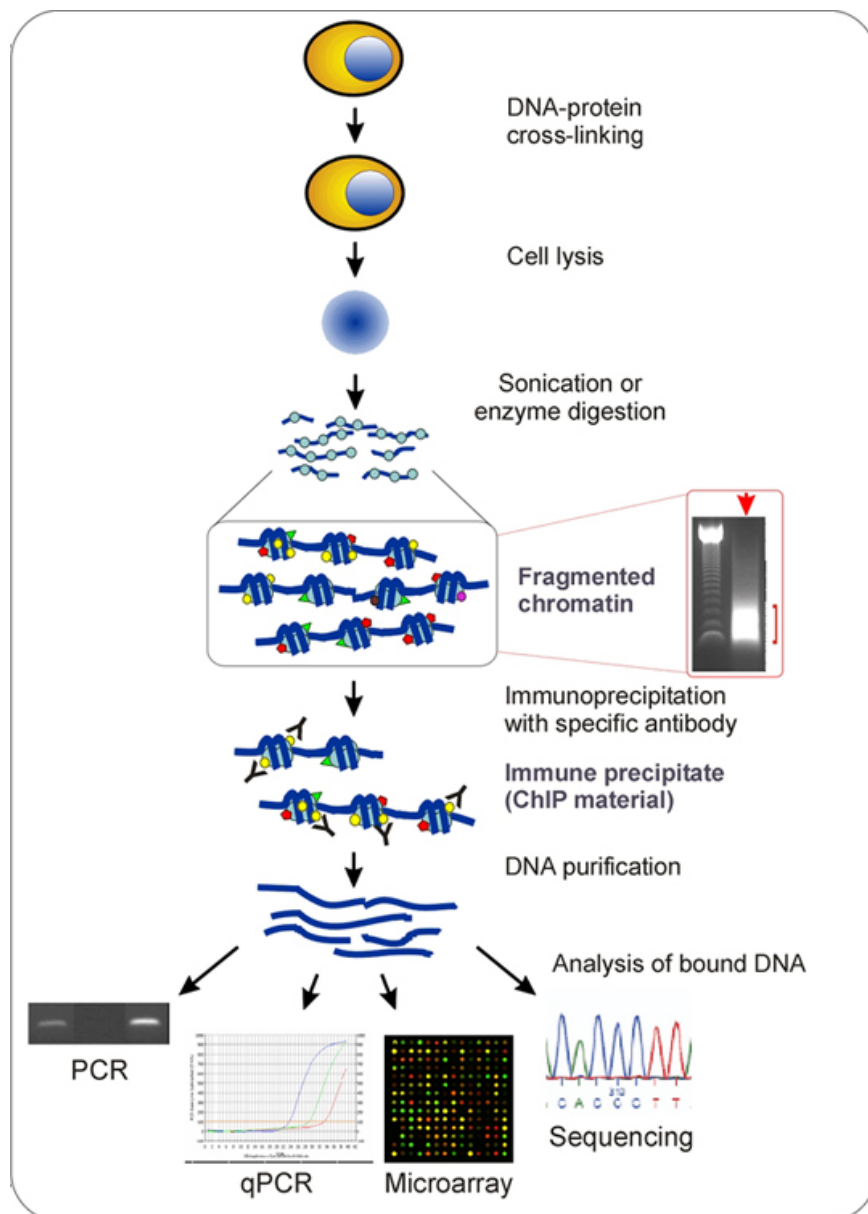
that are specific for modified histones (i.e. H3K9me3) or chromatin modifying enzymes (i.e. Suv39h1); another sample with a non-specific, control antibody

4. Recover antibody-chromatin complexes using a resin that binds to the constant region of antibodies

5. Elute chromatin at high salt concentration and revert crosslinks at high temperature

6. Digest protein with protease K and RNA with RNase

CHROMATIN IMMUNOPRECIPITATION (ChIP) → DETAILED ANALYSIS → Localization of protein at a defined region - sequence



5. Elute chromatin at high salt concentration and revert crosslinks at high temperature

6. Digest protein with protease K and RNA with RNase

7. Purify DNA and precipitate DNA

8. Measure the amount of immunoprecipitated DNA
In control versus Suv39h dn cells

METHODS:

A. PCR: design PCR oligos to amplify defined sequences in immunoprecipitates. Per PCR reaction only one locus can be examined by real-time PCR

B. ChIP on ChIP:

Immunoprecipitated DNA from control and Suv39h dn cells are labeled with nucleotides that are fluorescently labeled (555nm emission fluorophore or 488 nm emission fluorophore). Samples are mixed and hybridized to an array that contains spotted DNA probes that cover a large fraction of the genome. 488/555 nm ratio is measured → relative enrichment of histone modification at thousands of genomic sites in control/Suv39h dn cells

B. ChIP Seq:

Immunoprecipitated DNA is sequenced by massive parallel Sequencing. Control vs. Suv39h dn cells reveals enrichment of H3K9me3 at the ENTIRE genome level. single nucleotide levels