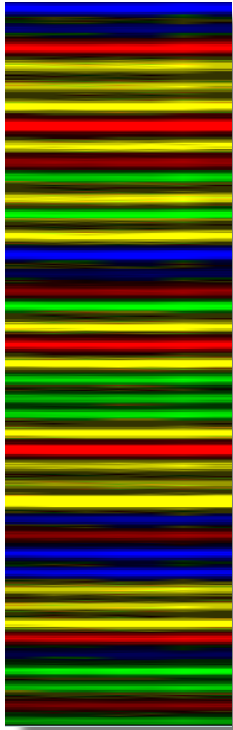


WS2017/2018

„Genomforschung und Sequenzanalyse

- Einführung in Methoden der Bioinformatik- “

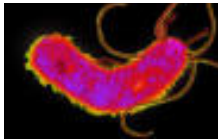
Thomas Hankeln



Methodik der Genomsequenzierung

NGS-Technologie

Meilensteine

	Phi X 174	1977	5.386 bp
	λ- Phage	1982	48.502 bp
	M. genitalium	1995	580.000 bp
	H. influenzae	1995	1.830.000 bp
	M. jannaschii	1996	1.660.000 bp
	S. cerevisiae	1997	12.500.000 bp
	E. coli	1997	4.654.000 bp
	C. elegans	1998	97.000.000 bp
	D. melanog.	1999	116.000.000 bp
	A. thaliana	2000	115.000.000 bp
	H. sapiens	2001	2.693.000.000 bp
	2012 > 1000genomes project (human) soon > Genome 10K (vertebrates)		

05.11.13 | **Erbgutanalyse**

Liegt das Talent für Mathematik in den Genen?

Ist ein Talent für Mathematik angeboren oder liegt es an der Erziehung?
Ein US-Unternehmer will die Frage nun klären: Beim "Projekt Einstein"
wird das Erbgut von 400 Mathematik-Genies analysiert. *Von Norbert Lossau*

ARTI

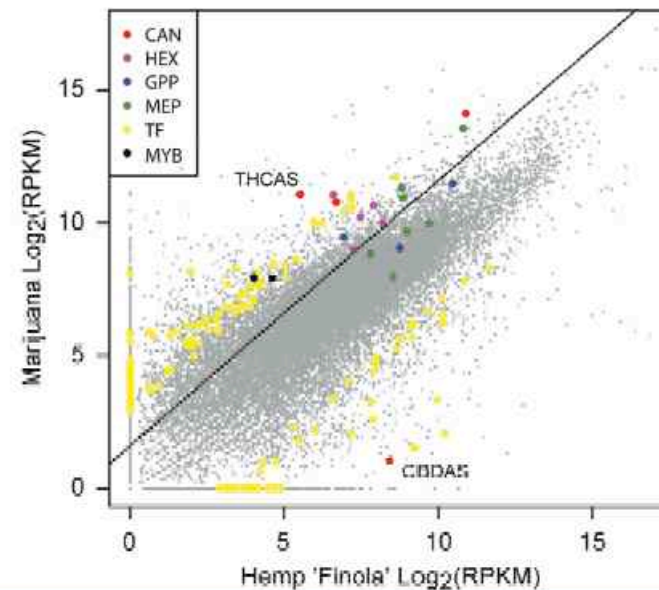
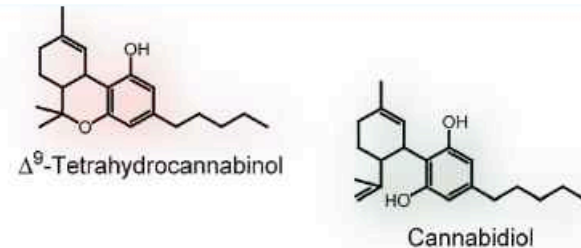
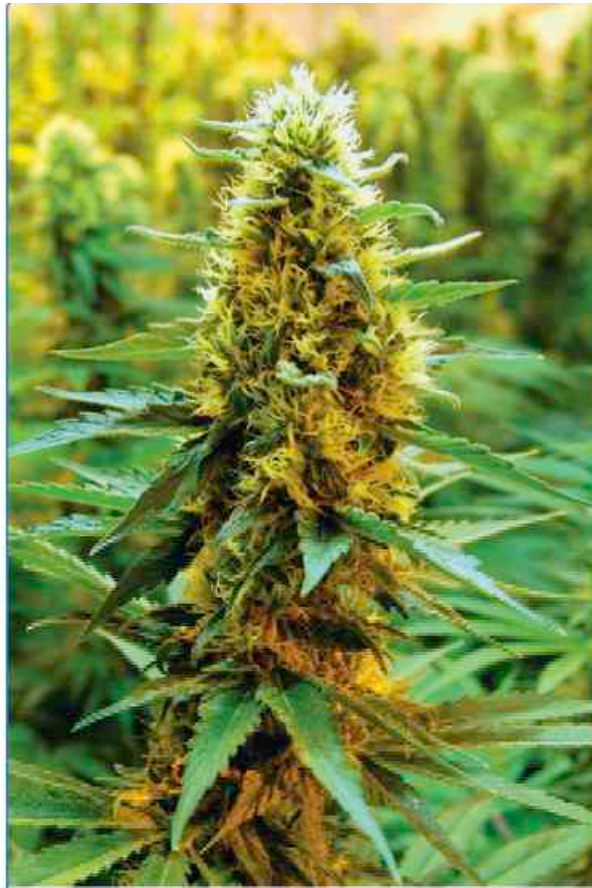
E-M

Koi

ANZEIGE



How hemp got high: Cannabis genome sequenced



van Bakel et al., Genome Biology 2011

Celebrity genomics



Science confirms the Neanderthal in Ozzy Osbourne

Among the findings that will be presented Friday: Osbourne's genes reveal a probability of alcohol dependency "six times higher than the average person."

"All this is big news for blokes everywhere, I think: If the Neanderthals could get laid, there's hope for us all."

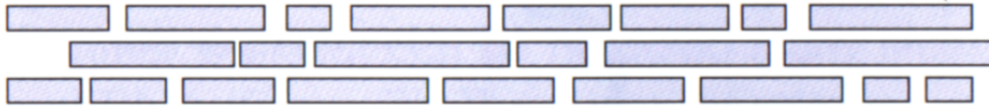
Methodik der Genomsequenzierung

- zu sequenzierende DNA muss zunächst in „handliche“ Abschnitte **zerlegt** und dann **vermehrt** werden (beim Sanger-Verfahren durch Klonierung oder bei **Next-Generation Sequencing** (NGS) meist durch PCR)
- DNA kann mit dem klassischen **Sanger**-Verfahren in Teilabschnitten von **600-1000 Bp** („long reads“) sequenziert werden.
- NGS-Verfahren lesen derzeit meist kurz (z.B. 50 bis 250 Bp; = **short reads**“), aber auch mittlerweile sehr lang (> 5000 Bp; long reads)

Target DNA



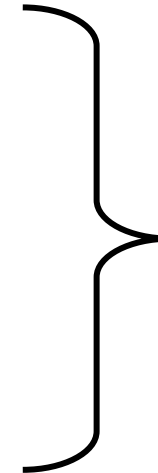
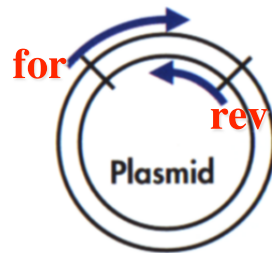
shear



Clone fragments in vector (Sanger)



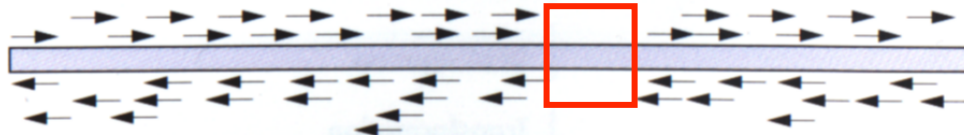
Sequence



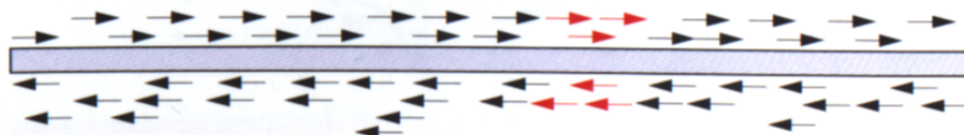
no cloning,
but PCR and
sequencing
in NGS



Assemble sequences
into contigs



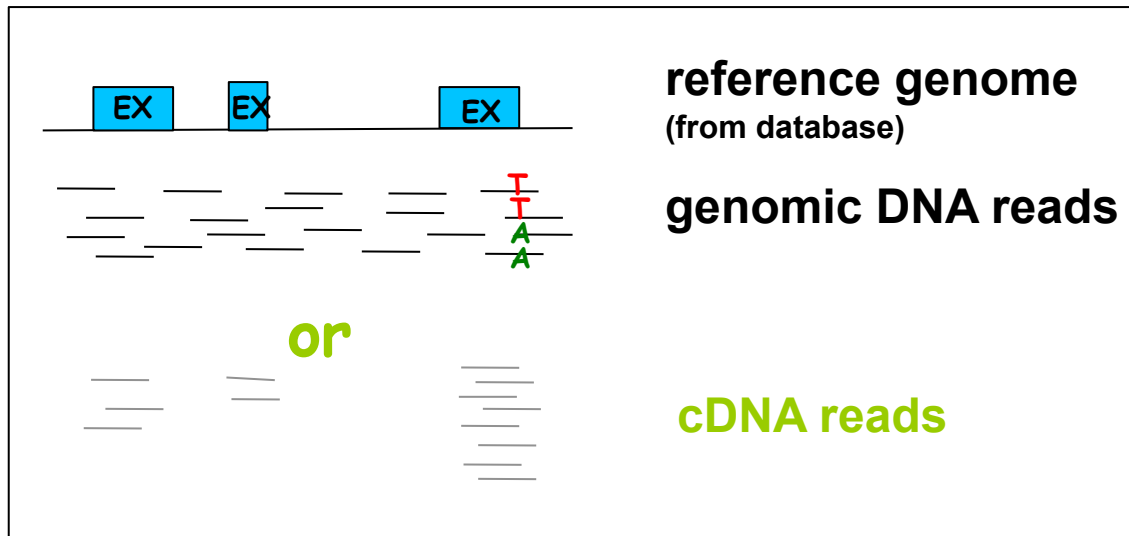
Close gaps with primer
walking or PCR (→)



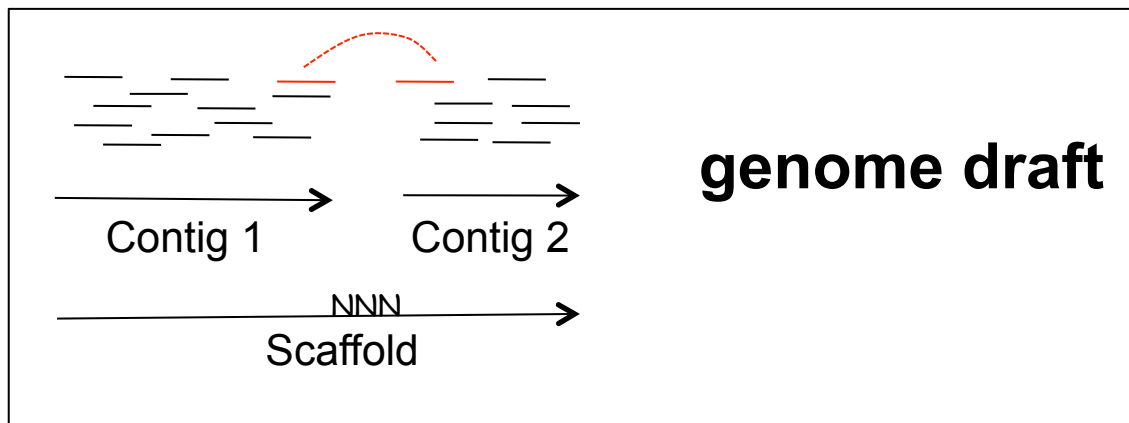
**Shotgun-
Strategie**

Genome sequencing:

Re- or *de novo*



- variant detection
- differential gene expression (RNA-Seq)



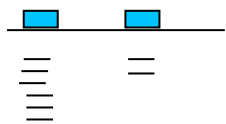
genome draft

- de novo genome
- gene discovery
- genome evolution

Re- vs. de novo-sequencing

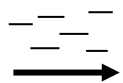
...require different bioinformatics algorithms

▪ Re-Seq



- > short reads ok
- > very high redundancy (SNP detection!)
- > **no assembly**, but **alignment to existing reference sequence („Mapping“)**
- > possible on normal PCs (64bit, quadcore, 16 GB RAM)

▪ de novo



- > longer reads better (due to repeats)
- > **classic assembly** (OLC, de Bruijn graph)
- > extremely RAM intensive (HPC, 512 GB RAM)

NGS-Bioinformatik

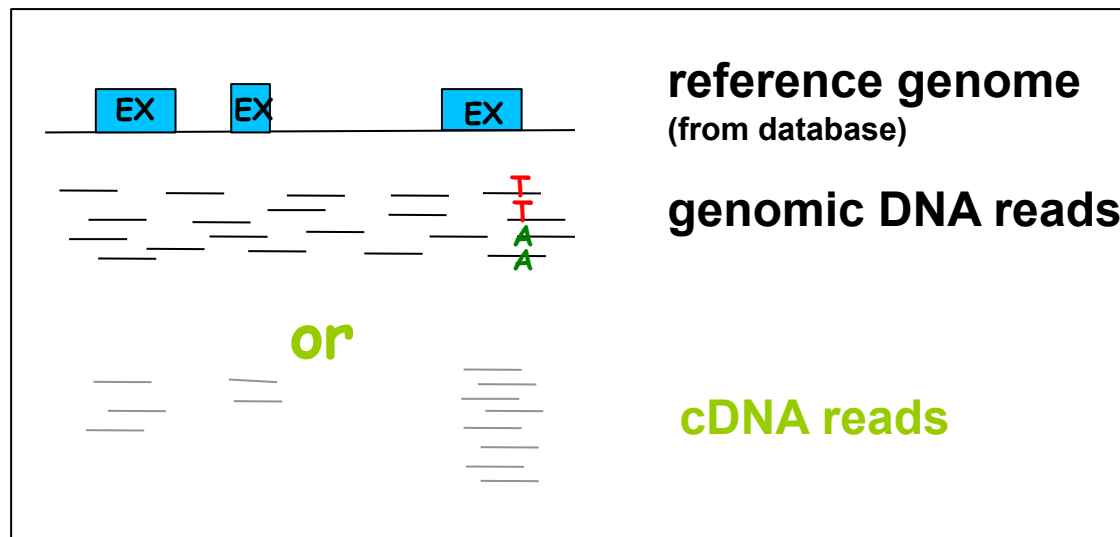


Applikationen von Re-Sequencing

- Wo ist unser Genom variabel? Wo sind Mutationen?
> **Gesamtgenom**-Resequenzierung
- Wo sind unsere Exons variabel? Mutationen?
> **Exome-Seq**
- Welche Gene werden transkribiert? Wie stark?
> **RNA-Seq**
- Welchen Chromatin-Status hat das Genom?
> **ChIP-Seq (u.a.)**
- Welchen Methylierungsstatus hat das Genom?
> **Methyl-Seq, Bisulfite-Seq (u.a.)**

...entdecke die Möglichkeiten!

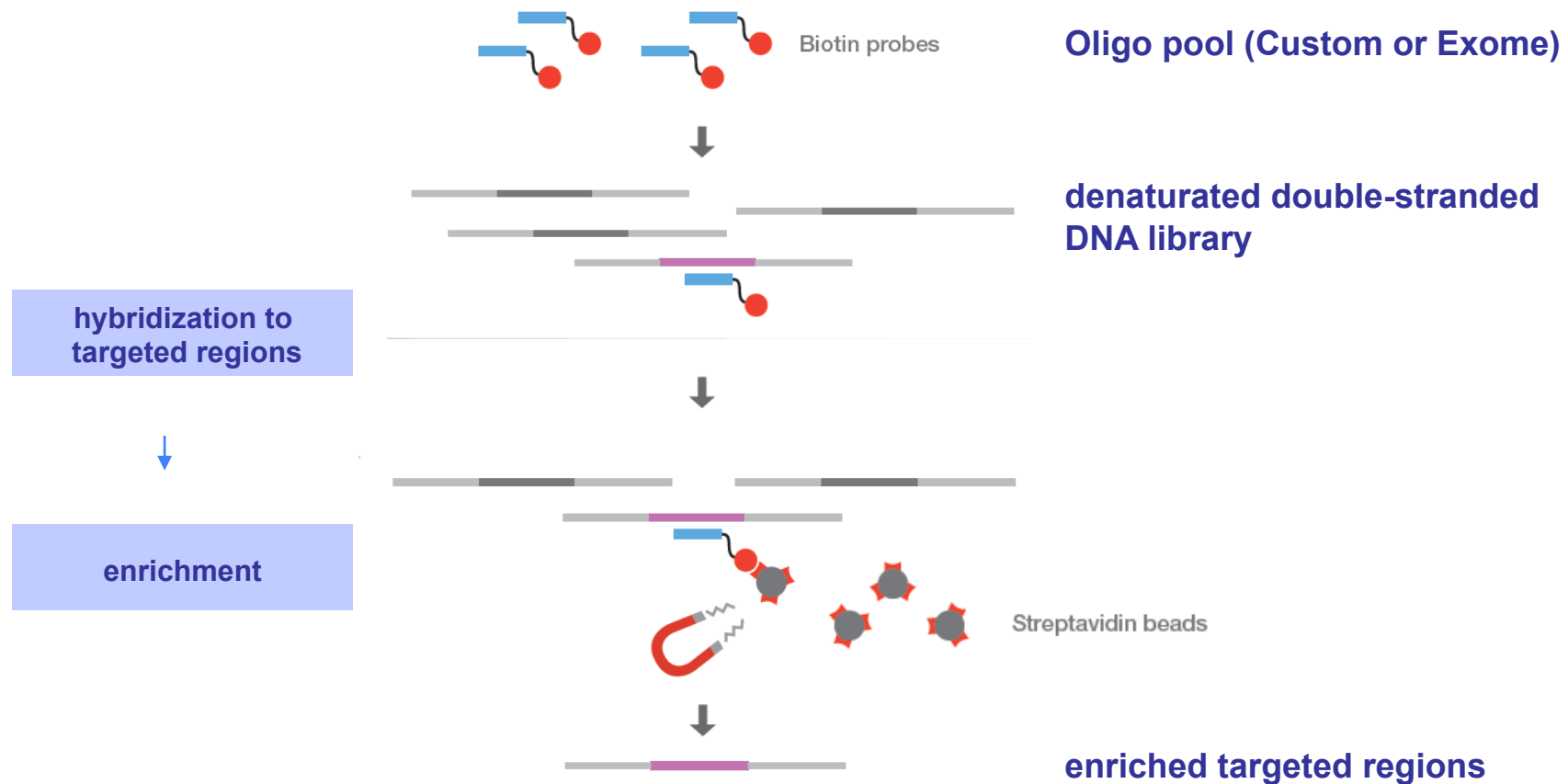
Gemeinsames Prinzip bei Re-Sequencing



Wichtigster Schritt: Zuordnung der Reads zu den passenden Regionen im Genom („Mapping“)

Exome-Seq

- requires ***a priori* knowledge of target** to be sequenced (e.g. exons)
- target enrichment by **hybridization**

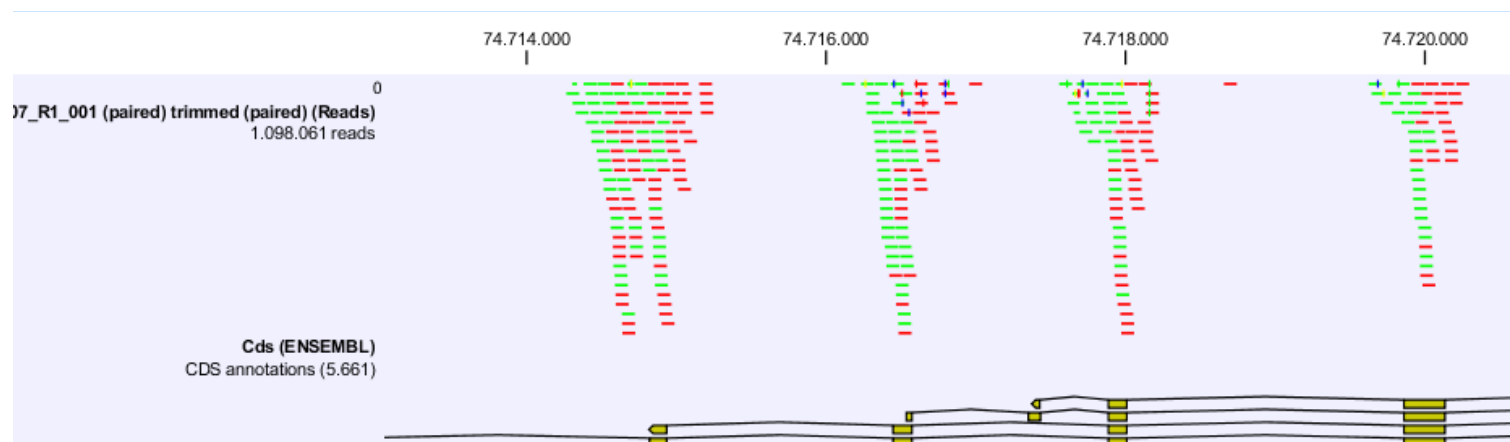




Exome-Seq

Exome Enrichment:

Target region size	62 Mb
Number of target genes	20,794
Number of target exons	201,121
Number of probes	340,427



Exome-Seq

Ergebnis der Exome-Seq ist eine Liste von Nt-Veränderungen, durch die sich das sequenzierte Exom von einem Referenzgenom unterscheidet.

SNVs : single nucleotide variants

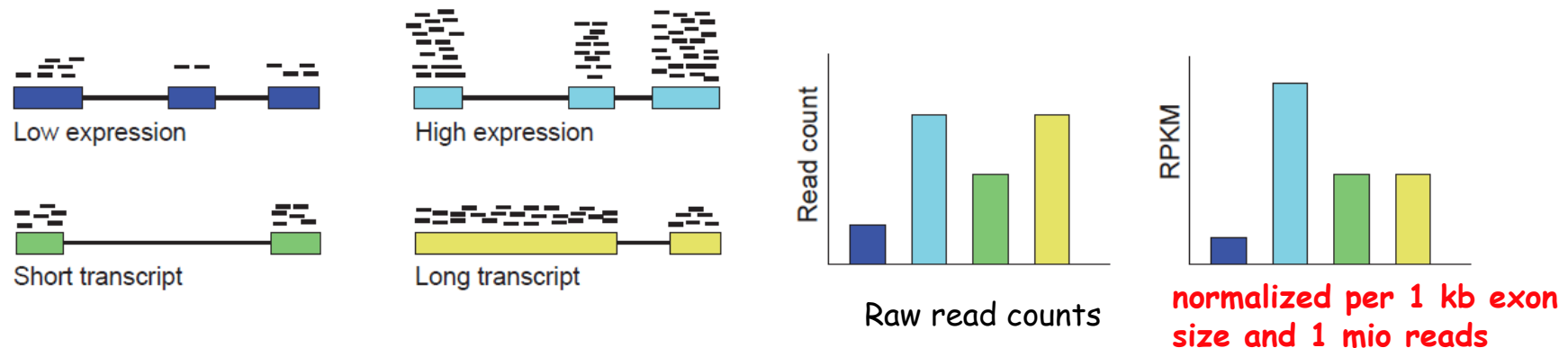
SNPs : single nucleotide polymorphisms*

cSNPs : coding SNPs

* mind. zu 1% als Allel in Human-Population vorhanden

Transcriptome analysis (RNA-Seq)

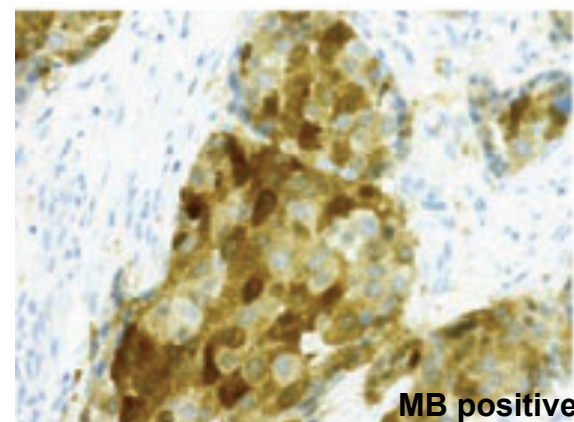
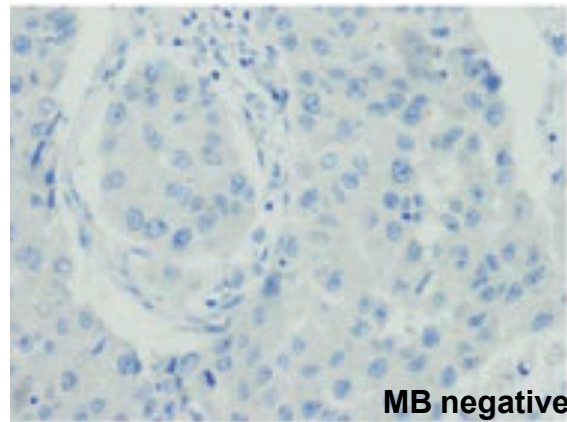
- RNA isolation > cDNA production & fragmentation
- high-throughput sequencing
- mapping of fragments to reference genome
- read-counting as measure of gene transcription



A typical measure is “RPKM”
= reads per Kb of exon sequence and 1 mio reads in the dataset

Beispiel: Was macht Myoglobin in Brustkrebs-Zellen?

MB immunostaining on breast tumors



RNA-Seq

**Differenziell exprimierte Gene
= Hinweis auf molekulare Veränderungen**

Transcriptome analysis (RNA-Seq)

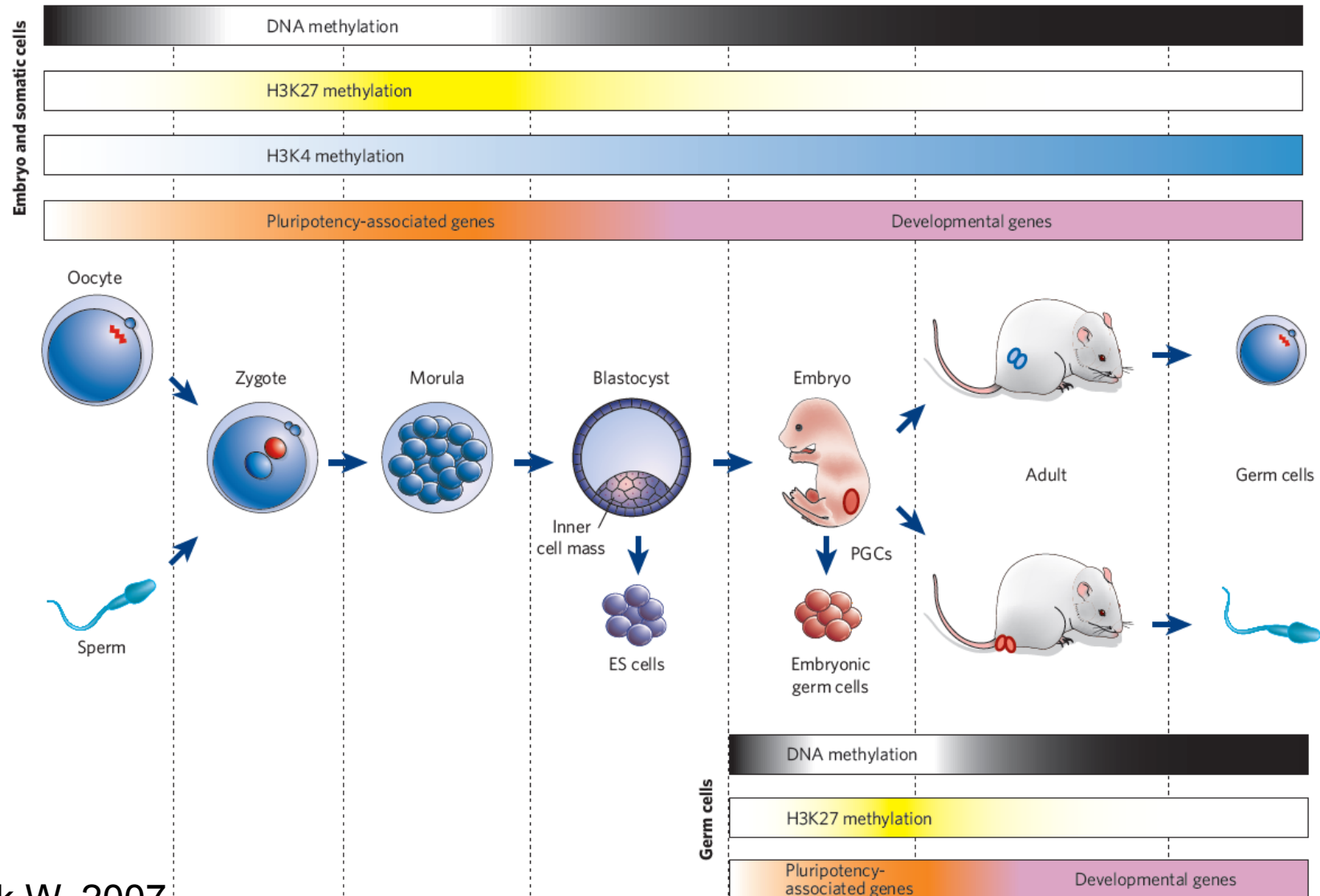
Ergebnis der RNA-Seq ist eine **Liste von Genen**, die zwischen den betrachteten Datensätzen statistisch signifikant **differenziell reguliert** sind.

Manchmal enthalten diese Listen Hunderte oder Tausende von Genen.

Dies macht es erforderlich, die Gene funktionell zu kategorisieren (GeneOntology-Vokabular, KEGG Pathways).

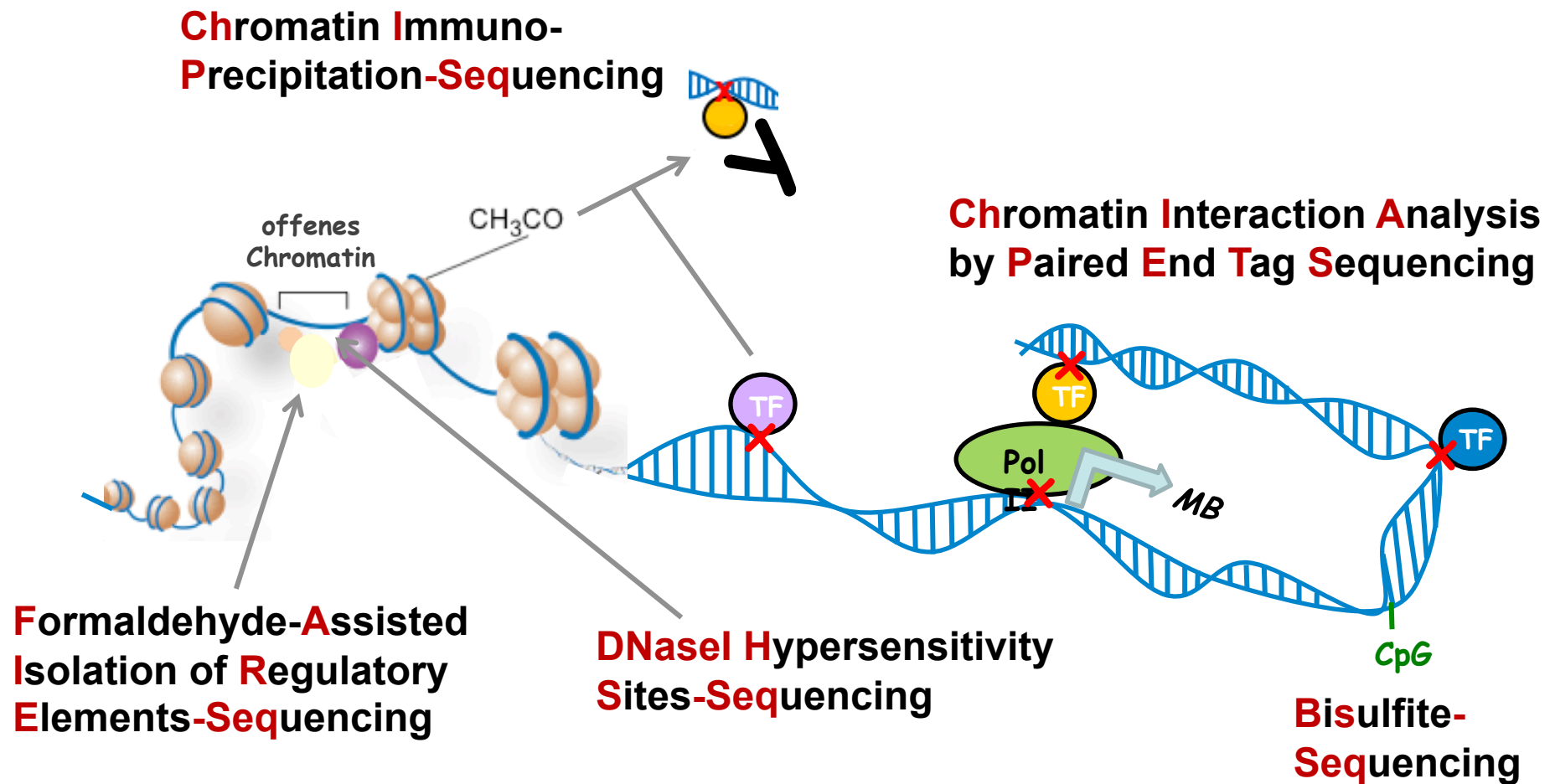
Sind bestimmte funktionelle Kategorien in den differenziell regulierten Genen über- oder unterrepräsentiert?
Was sagt das über die „Biologie“ der verglichenen Proben?

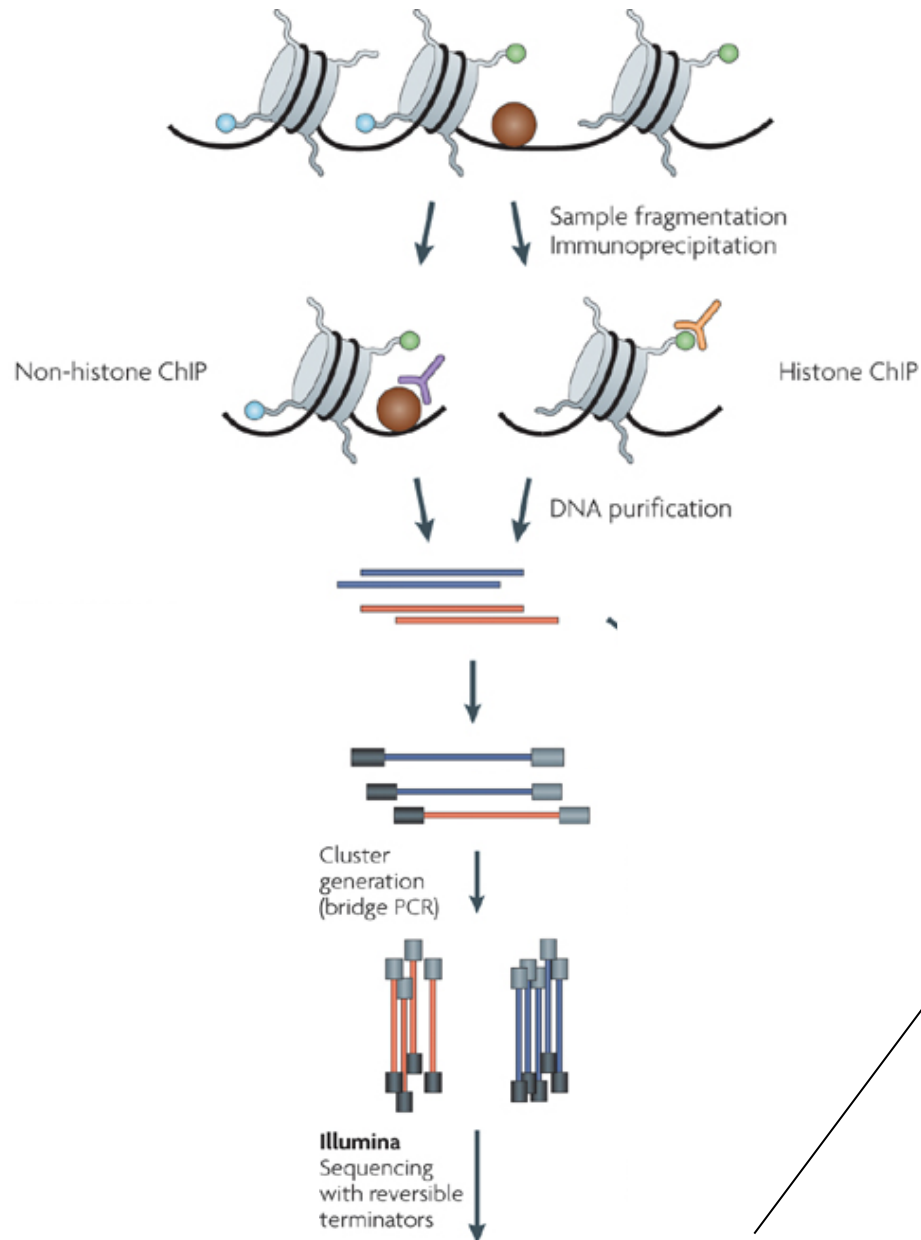
Epigenetische Genommodifikationen



Reik W, 2007

Genomweite Detektion von Chromatin-Modifikationen durch NGS-Methoden

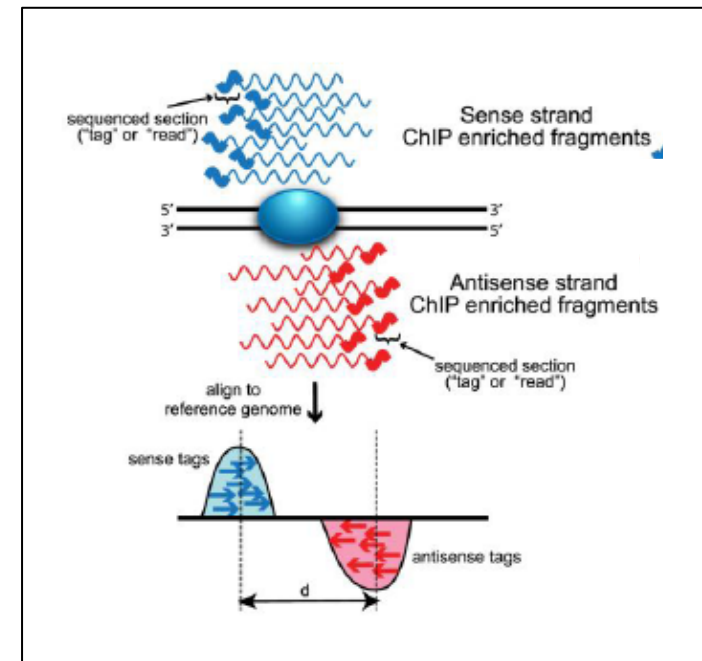




Reads > Mapping to reference genome

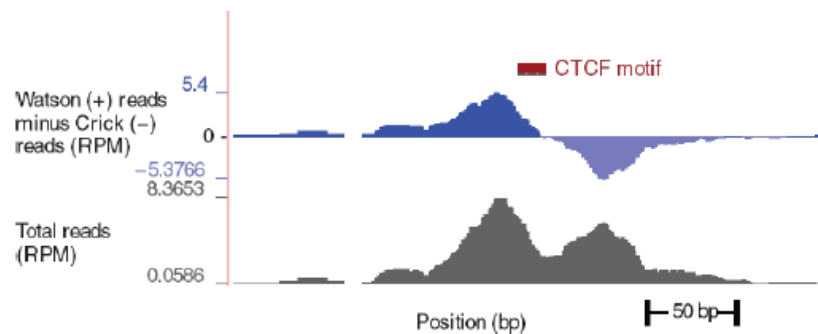
ChIP-Seq

„peak calling“

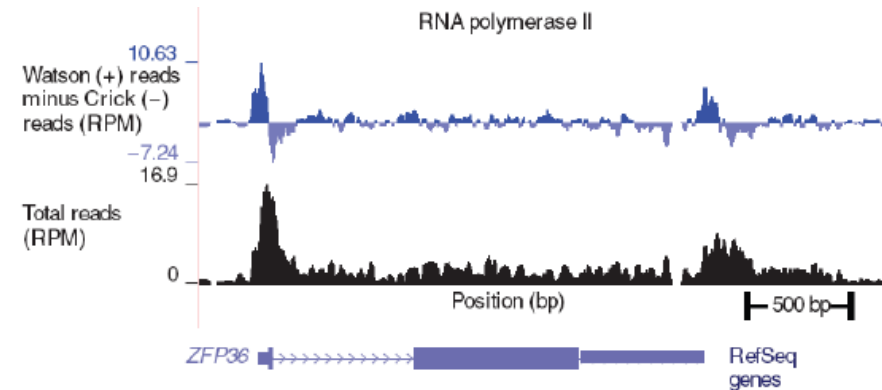


➤ peaks zeigen Orte, wo TF gebunden hat

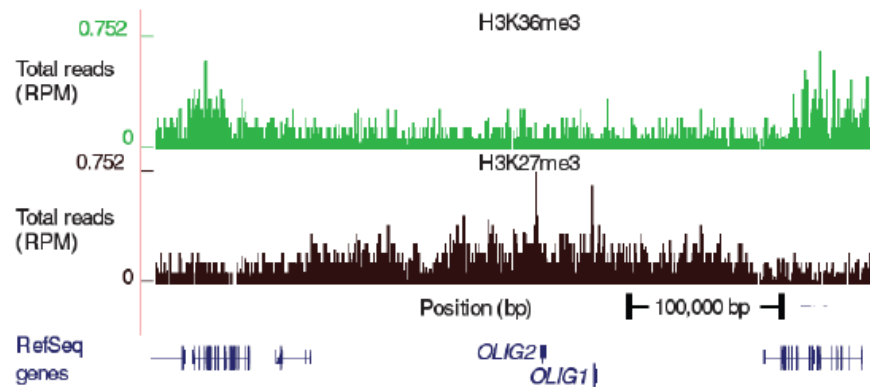
Chip-Seq: Ergebnisbeispiele



Transcriptional regulation protein (CTCF)



RNA polymerase II



Epigenetics: Repressive mark for H3K27me

ChIP Seq peak types from different experiments

Computation for ChIP-seq and RNA-seq studies
Shirley Pepke, Barbara Wold & Ali Mortazavi
Nature Methods 6, S22 - S32 (2009)

NGS-Kursprogramm Modul 7A

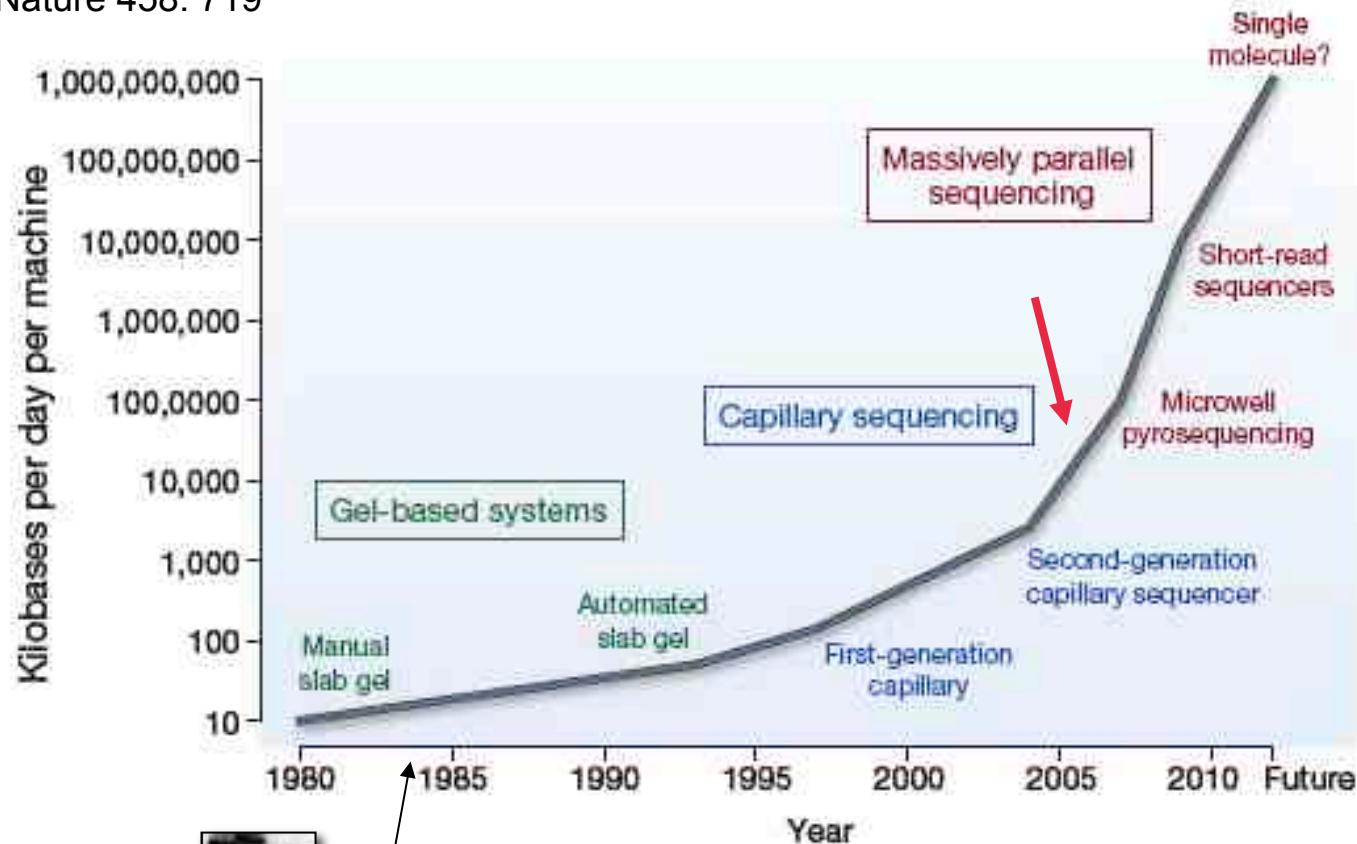
Trio-Analyse: Identifizierung von Krankheitsgenen durch Exome-Seq von Familien (Tag 1 & 2)

Transkriptomanalyse: Der Einfluss der Myoglobinexpression auf das Gesamt-Transkriptom von Krebszellen (RNA-Seq; Tag 3 & 4)

Q: Wie funktioniert NGS technisch?

Sequencing technology: A million-fold improvement!

Nature 458: 719



my diploma thesis: 1kb Maxam-Gilbert, 4 weeks (day & night in the isotope lab)

NGS technology: How to...

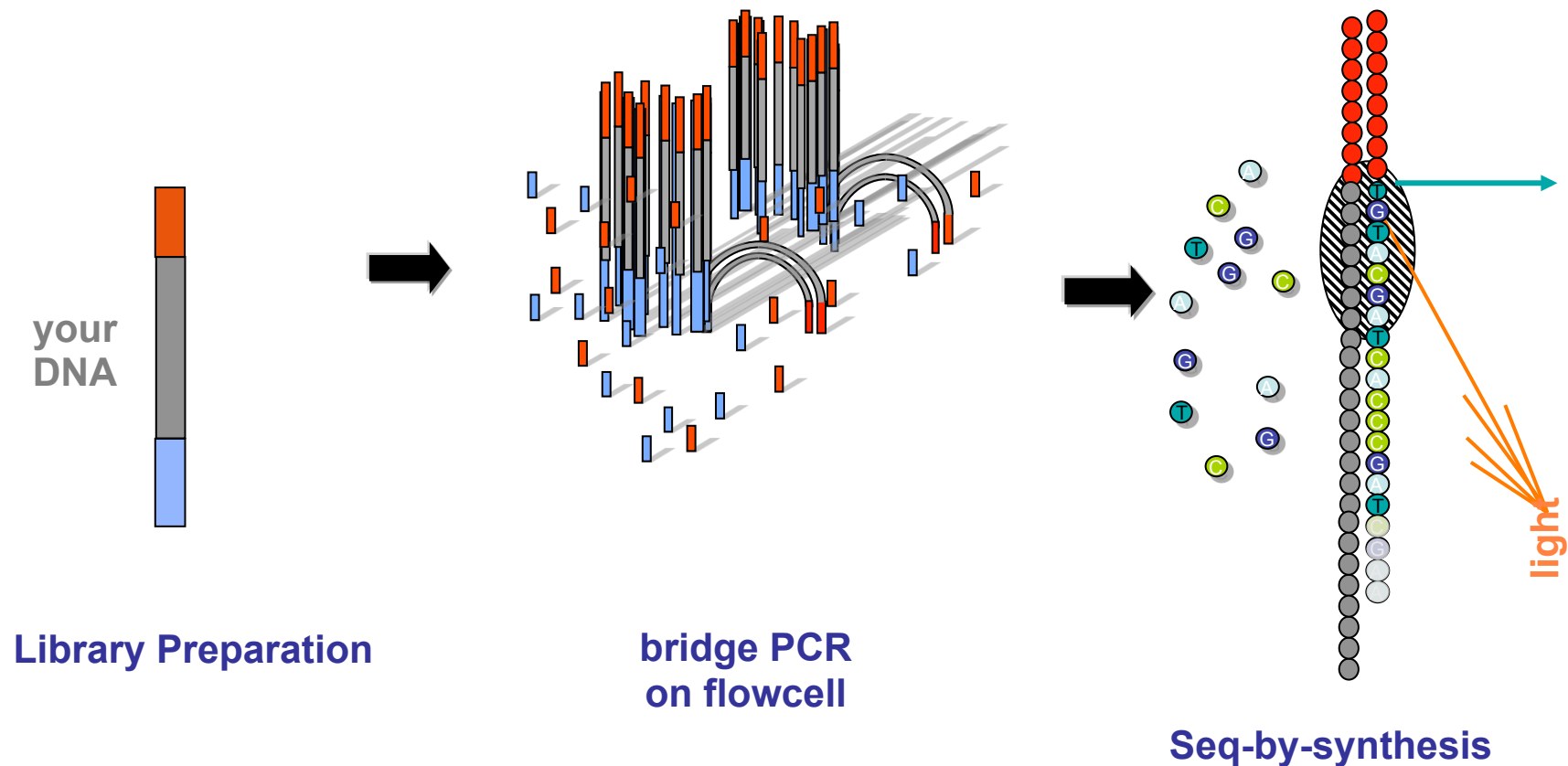


tedious cloning
high chemical costs
slow electrophoresis



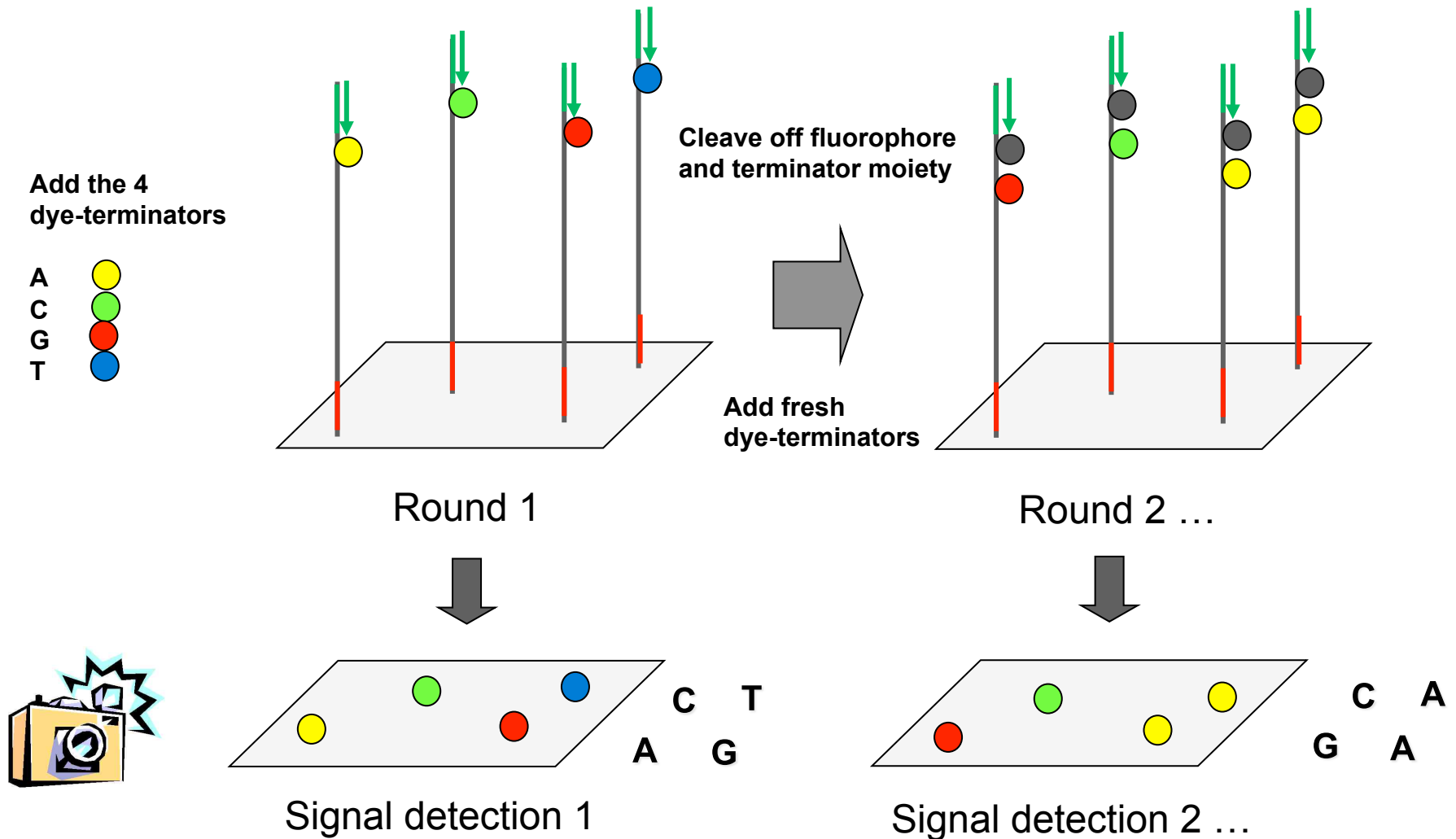
PCR or even single molecules
extreme miniaturisation
massively-parallel read-out

Illumina Sequencing - principle

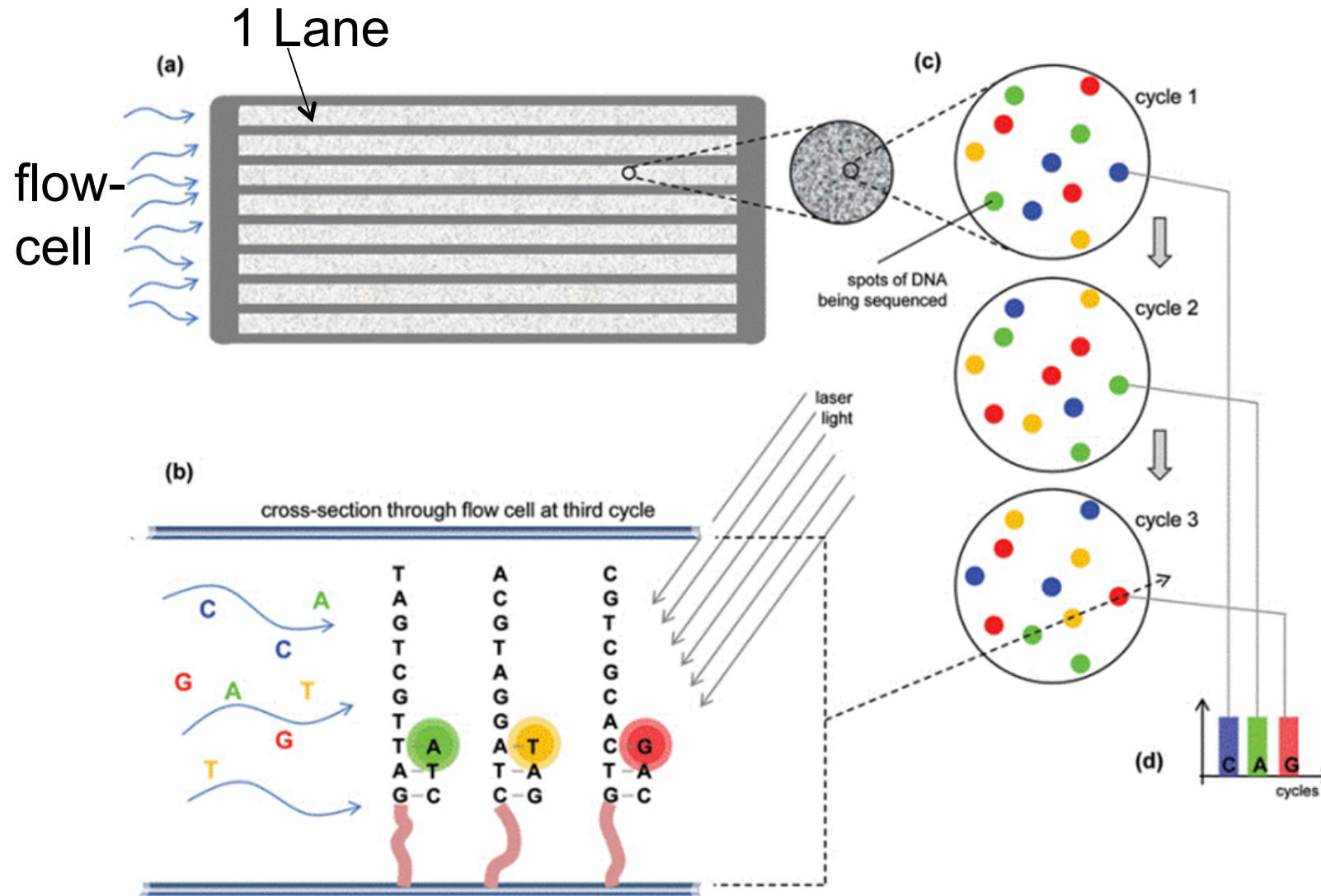


Illumina Sequencing

- „Sequencing-by-synthesis“ using reversible dye-terminators

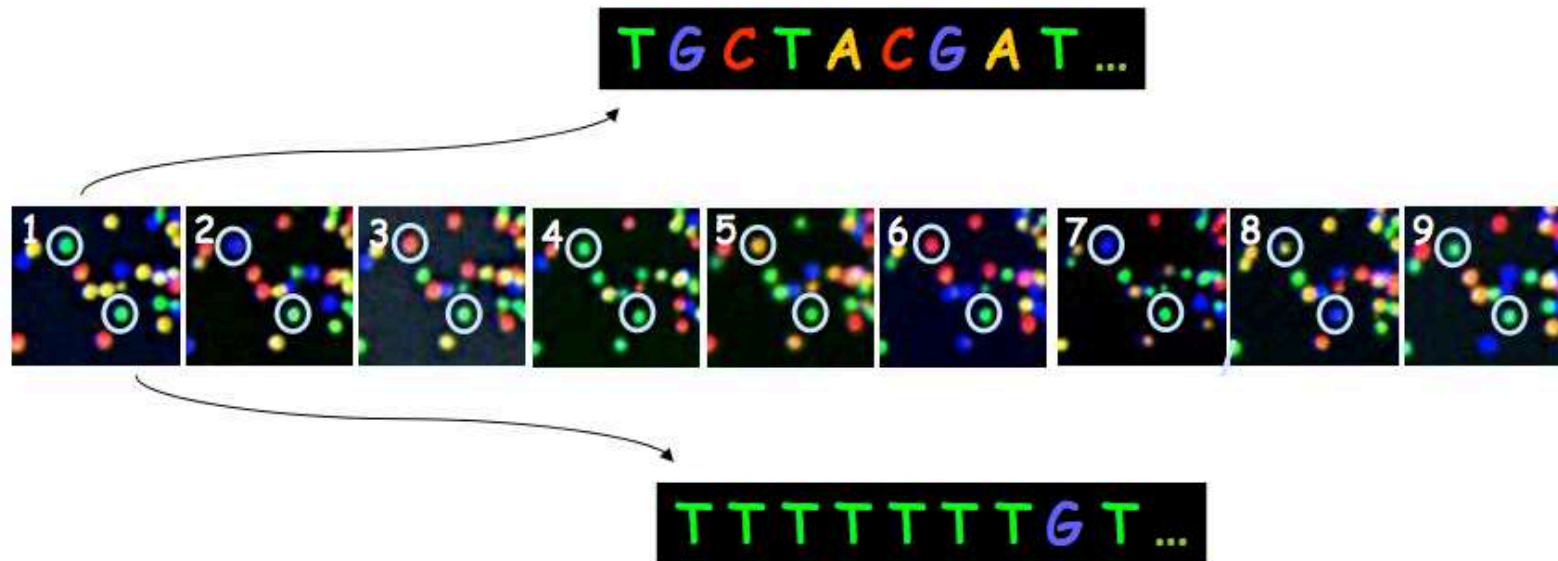


Illumina-Sequenzierung



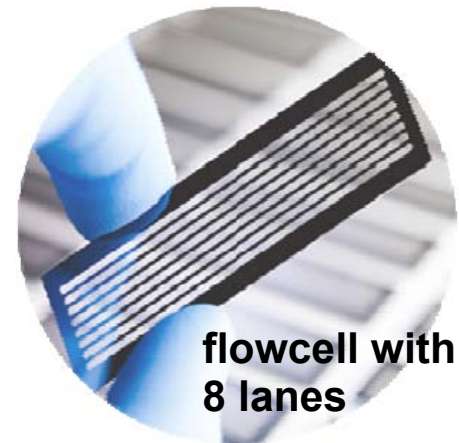
Illumina-Sequencing

9 cycles shown, two cluster position marked...



The sequence of each cluster is determined by consecutive images.

Illumina HiSeq2000



- 600 Gbp / run = **6 Billion Reads** x 100 Bp!
- 600 Gbp / 3 Gbp / 30 x coverage = 6 human genomes
- run time 11 days (high-throughput mode; rapid mode possible)

Illumina Library-Preparation

gDNA, cDNA, plasmids,
amplicons ...

DNA fragmentation

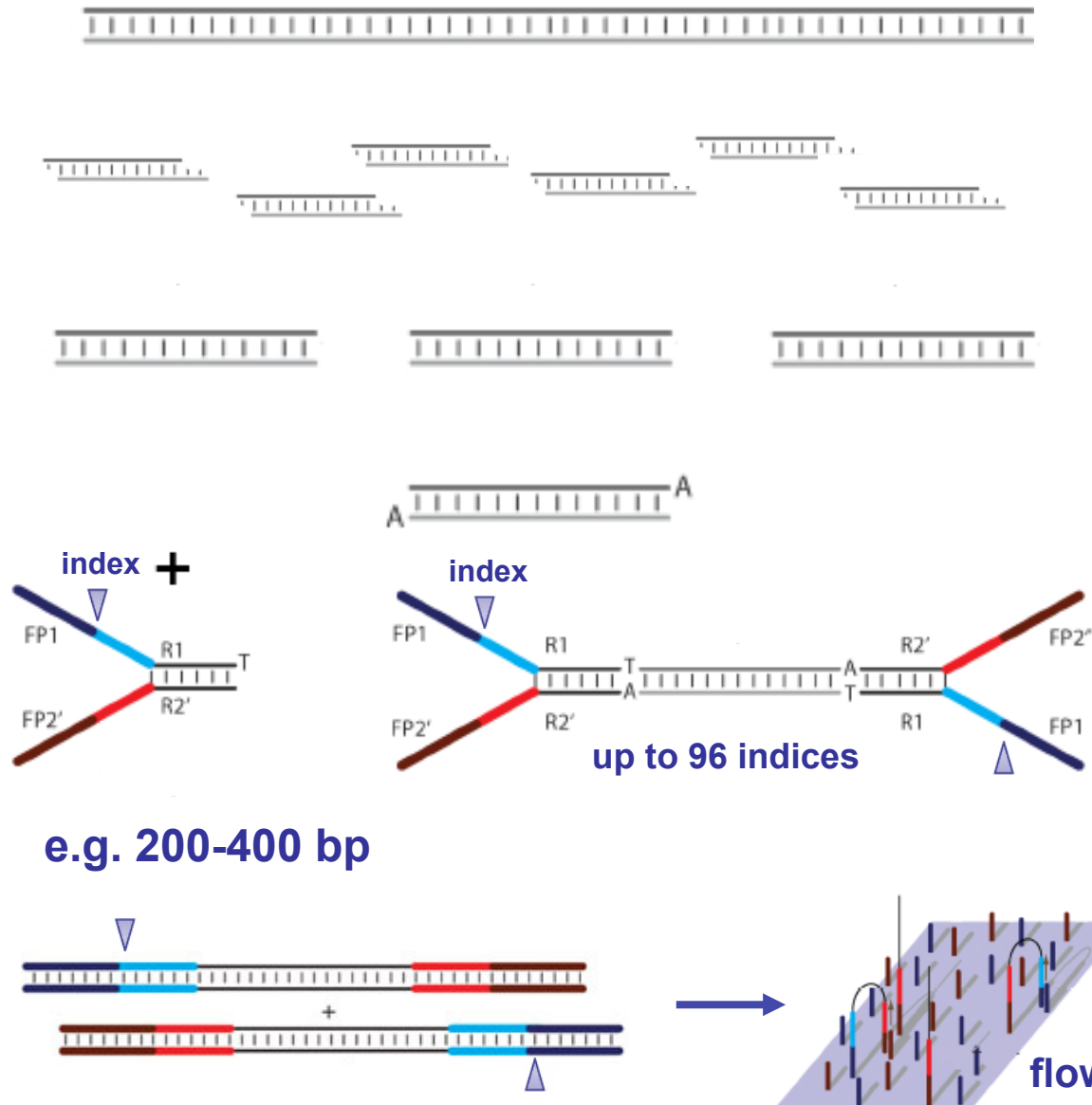
end-repair

A-tailing

adapter-ligation

size selection

enrichment



Illumina Library-Preparation

Genom-DNA oder cDNA

ATTGCGTAGCATCGCGATACGACGTGCTAGATGACTGATCGTACGACGATGATGATCGAGTAGCATGCTCATTGCGTAGCATCGCGATACGACGTGCTAGATGACTGATCGT
TAACGCATCGTAGCGCTATGCTGCACGATCTAGTGACTAGCTAGCTGCTACTACTAGCTCATCGTACGAGTAACGCATCGTAGCGCTATGCTGCACGATCTAGTGACTAGCA

Fragmentierung



ATTGCGTAGCATCGCGATACGA
TAACGCATCGTAGCGCTATG

CTGATCGTACGAC
AGTGACTAGCTAGCTG

AGTAGCATGCT
TCATCGTACGAGTAACGCA

CGTGCTAGATGA
CTGCACGATCT

GATGATGATCG
CTACTACTAGC

CATTGCGTAGCATCGCGATACGACGTGCTAGATGACTGATCGT
TCGTAGCGCTATGCTGCACGATCTAGTGACTAGCA

Nach Größe sortieren



CATTGCGTAGCATCGCGATACGACGTGCTAGATGACTGATCGT
TCGTAGCGCTATGCTGCACGATCTAGTGACTAGCA

ATTGCGTAGCATCGCGATACGA
TAACGCATCGTAGCGCTATG

AGTAGCATGCT
TCATCGTACGAGTAACGCA

CTGATCGTACGAC
AGTGACTAGCTAGCTG

CGTGCTAGATGA
CTGCACGATCT

GATGATGATCG
CTACTACTAGC

gewünschte Fraktion isolieren
(oft 200 bis 500 bp, richtet sich nach
Leselänge)

Illumina Library-Preparation

Extrahierte Fraktion

AGTAGCATGCT CTGATCGTACGAC CGTGCTAGATGA
TCATCGTACGAGTAACGCA AGTGACTAGCTAGCTG CTGCACGATCT

Enden „behandeln“



AGTAGCATGCTA	CTGATCGTACGACA	CGTGCTAGAA
ATCATCGTACGA	AGACTAGCTAGCTG	AGCACGATCT

Fragmente mit Y-Adaptern ligieren



3' - TCTAGCCTTCTCGAGCATACGGCAGAAGACGAAC - 5'
 - A C A C T C T T T C C C T A C A C G A C
 G C T C T T C G A T C T C T G A T C G T A C G A C A G A T C G G A A G A G C T C G T A T G C C G T C T T C G T T G - 3'
 - G T T C G T C T T C T G C C G T A T G C T C G A G A A G G C T A G A G A C T A G C T A G C T G T C T A G C C T T C T C G
 C A G C A C A T C C C T T T C T C A C A - 5'

PCR: Zyklus 1



ACACTCTTTCCCTACACGACGCTCTTCCGATCTCTGATCGTACGACA**GATCGGAAGAGCTC**GTATGCCGTCTTCTGCTTG
TGTGAGAAAGGGATGTGCTGCGAGAAGGCTAGAGACTAGCTAGCTG**TCTAGCCTTCTCGAG**CATACGGCAGAAGACGAAC

DNA-Strang mit 2 unterschiedlichen Enden
(das reziproke Produkt des anderen Strangs ist nicht gezeigt)

Illumina Library-Preparation

DNA-Strang mit 2 unterschiedlichen Enden

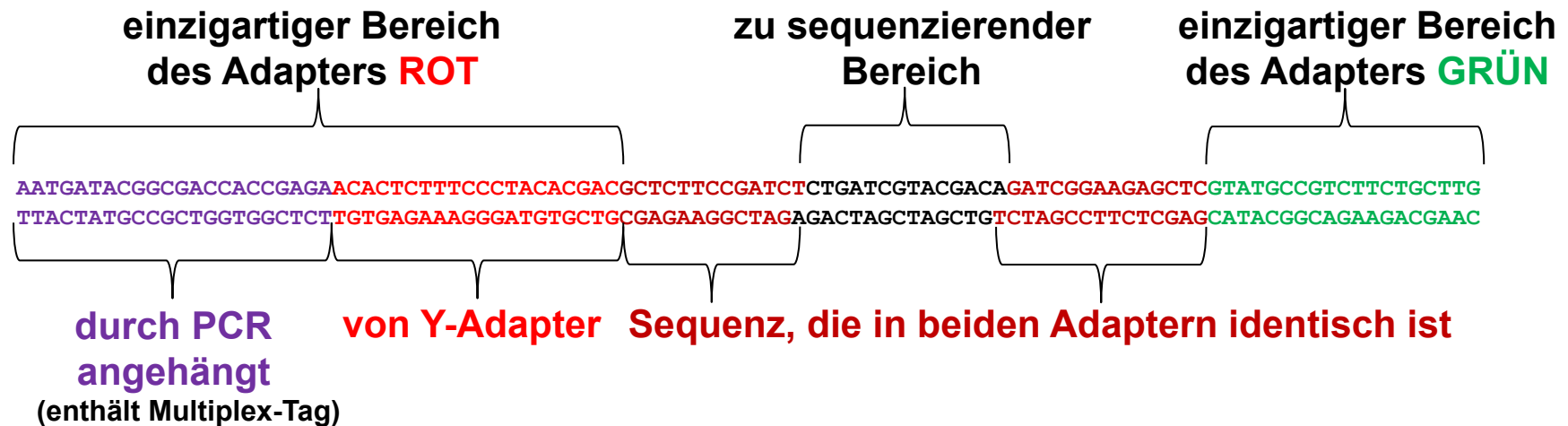
ACACTCTTTCCCTACACGACGCTCTTCCGATCTCTGATCGTACGACAGATCGGAAGAGCTCGTATGCCGTCTTCTGCTTG
TGTGAGAAAGGGATGTGCTGCGAGAAGGCTAGAGACTAGCTAGCTGTCTAGCCTTCTCGAGCATACGGCAGAAGACGAAC

PCR: Zyklen 2+



AATGATACGGCGACCACCGAGAACACTCTTTCCCTACACGACGCTCTTCCGATCT >
AATGATACGGCGACCACCGAGAACACTCTTTCCCTACACGACGCTCTTCCGATCTCTGATCGTACGACAGATCGGAAGAGCTCGTATGCCGTCTTCTGCTTG
TTACTATGCCGTGGTGGCTCTTGTGAGAAAGGGATGTGCTGCGAGAAGGCTAGAGACTAGCTAGCTGTCTAGCCTTCTCGAGCATACGGCAGAAGACGAAC
< TCTAGCCTTCTCGAGCATACGGCAGAAGACGAAC

PCR: Zyklen 2+



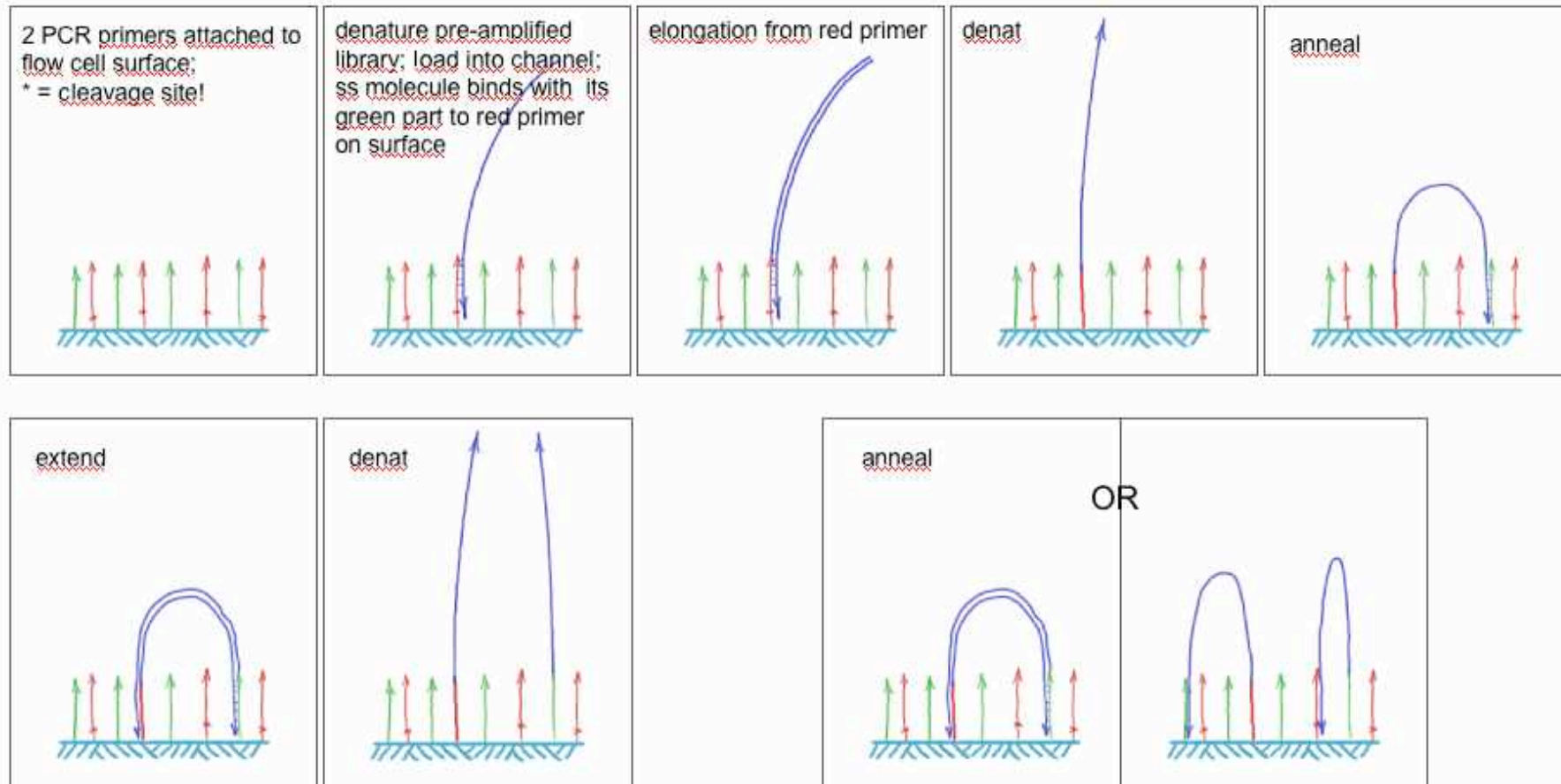
Illumina Library-Preparation



- Multiplex-**Tag** ist also nicht in den Sequenzen enthalten, die von den Primern der Runde 1 und 2 ausgehen
- Für **Tags** gibt es daher einen separaten kleinen Sequenzierlauf ausgehend vom Primer **DUNKELROT**
- die Sequenzierungen von Primer 1 und Primer 2 aus nennt man **PAIRED-ENDS!**

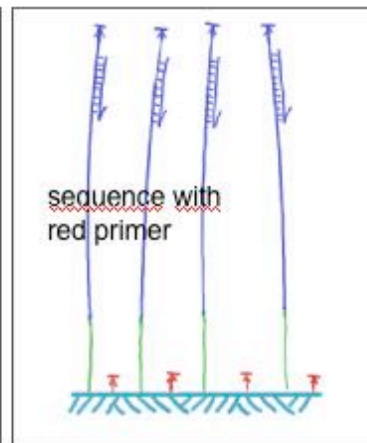
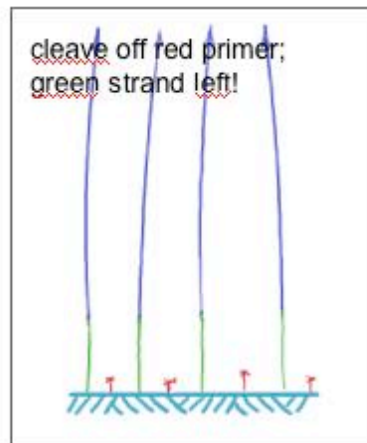
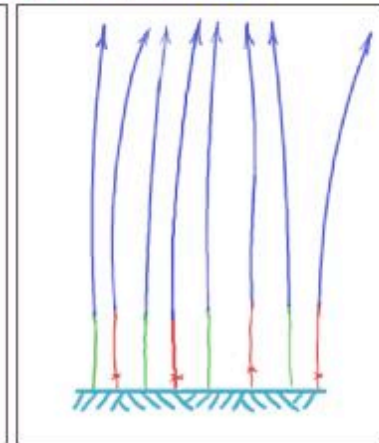
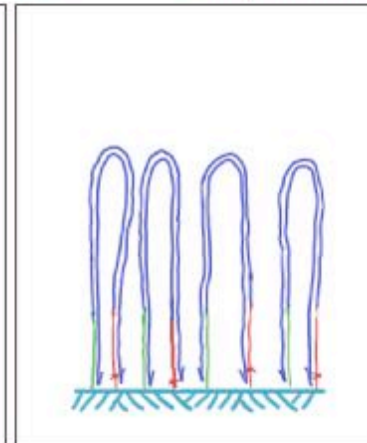
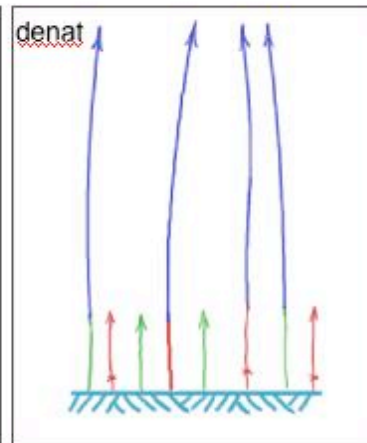
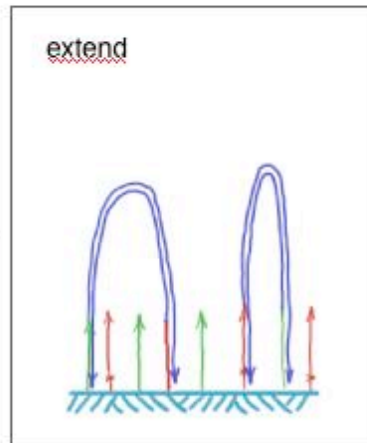
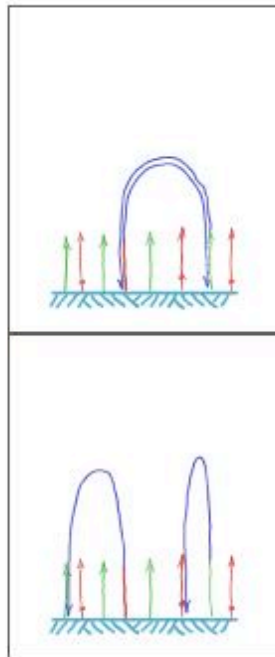
Illumina template prep: bridge PCR

Shear DNA > size fractionate > attach adapters > pre-amplify by PCR with red and green primers > then: bridge PCR on flowcell



Illumina template prep: bridge PCR

35 cycles
bridge formation
ca 1000 sequencing templates

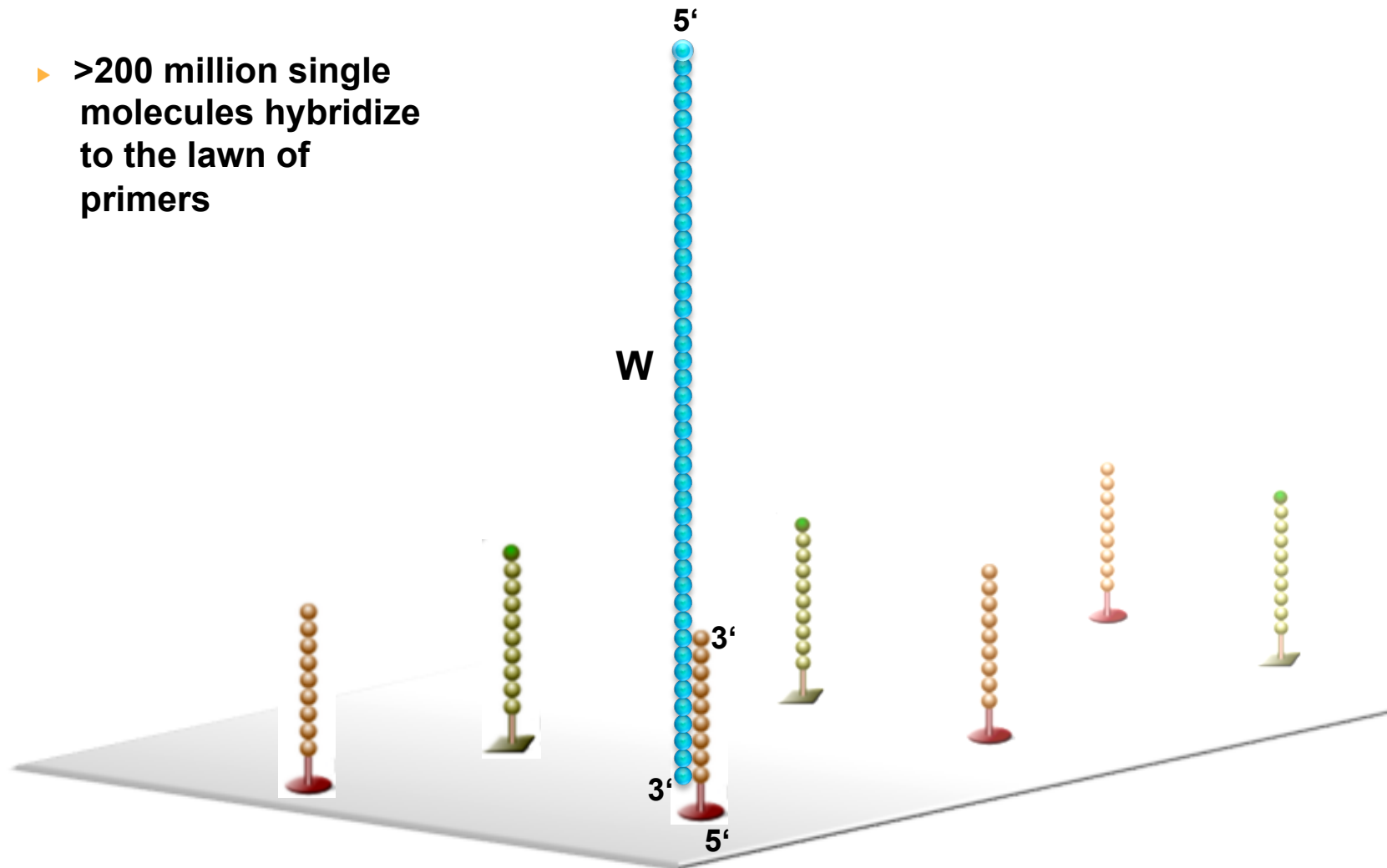


Achtung:

Cleavage-Punkt des roten Primers liegt „weiter oben“ und lässt Großteil der roten Primerseq in der flowcell intakt. Nach Seq-Runde1 kann man also erneut eine Bridge-PCR machen, dann den Strang grünen Strang durch Cleavage entfernen und den verbleibenden roten Strang mit dem grünen Primer sequenzieren (= „paired end“-Sequenzierung; Runde 2)

Illumina Cluster Generation

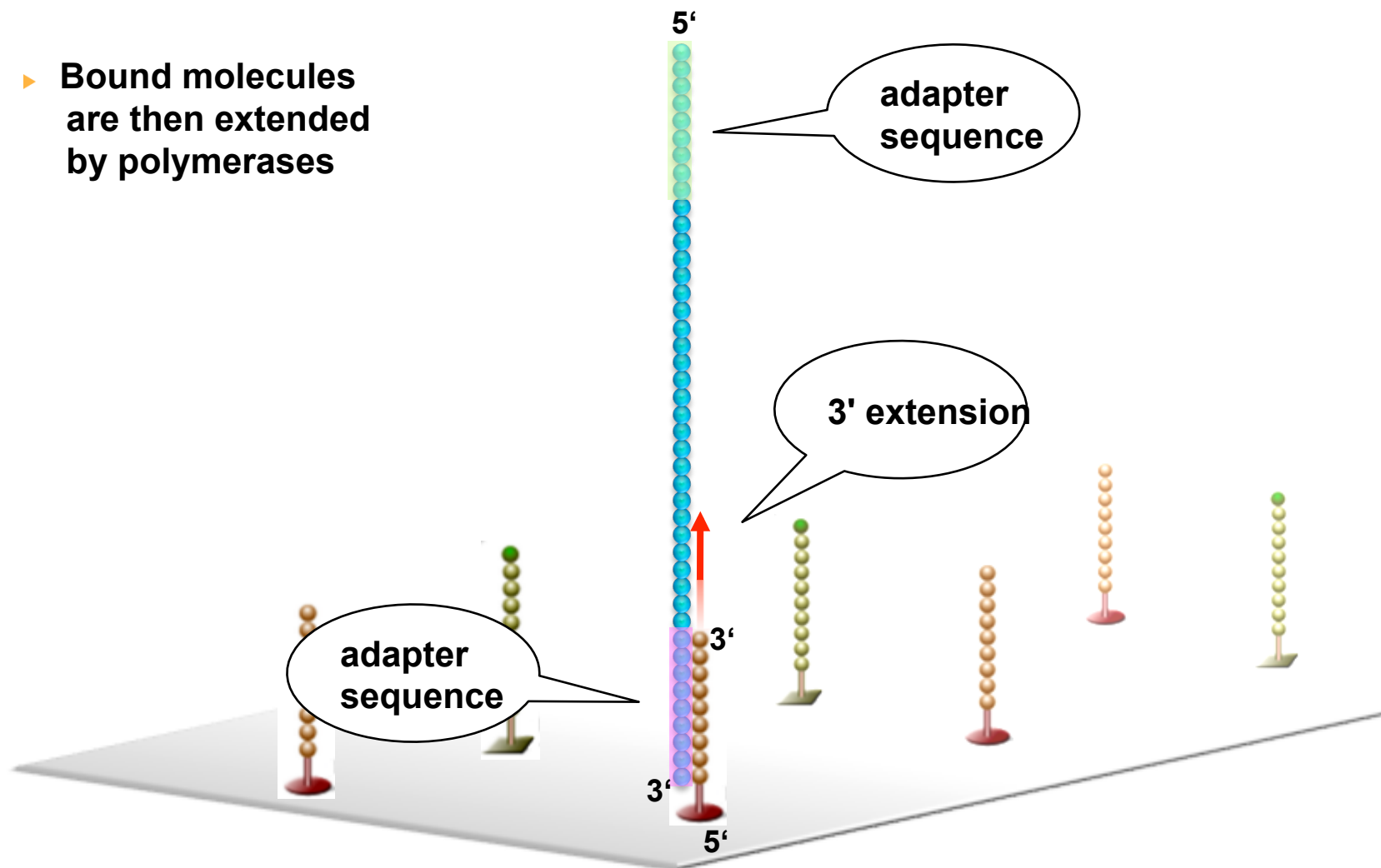
- ▶ >200 million single molecules hybridize to the lawn of primers





Illumina Cluster Generation

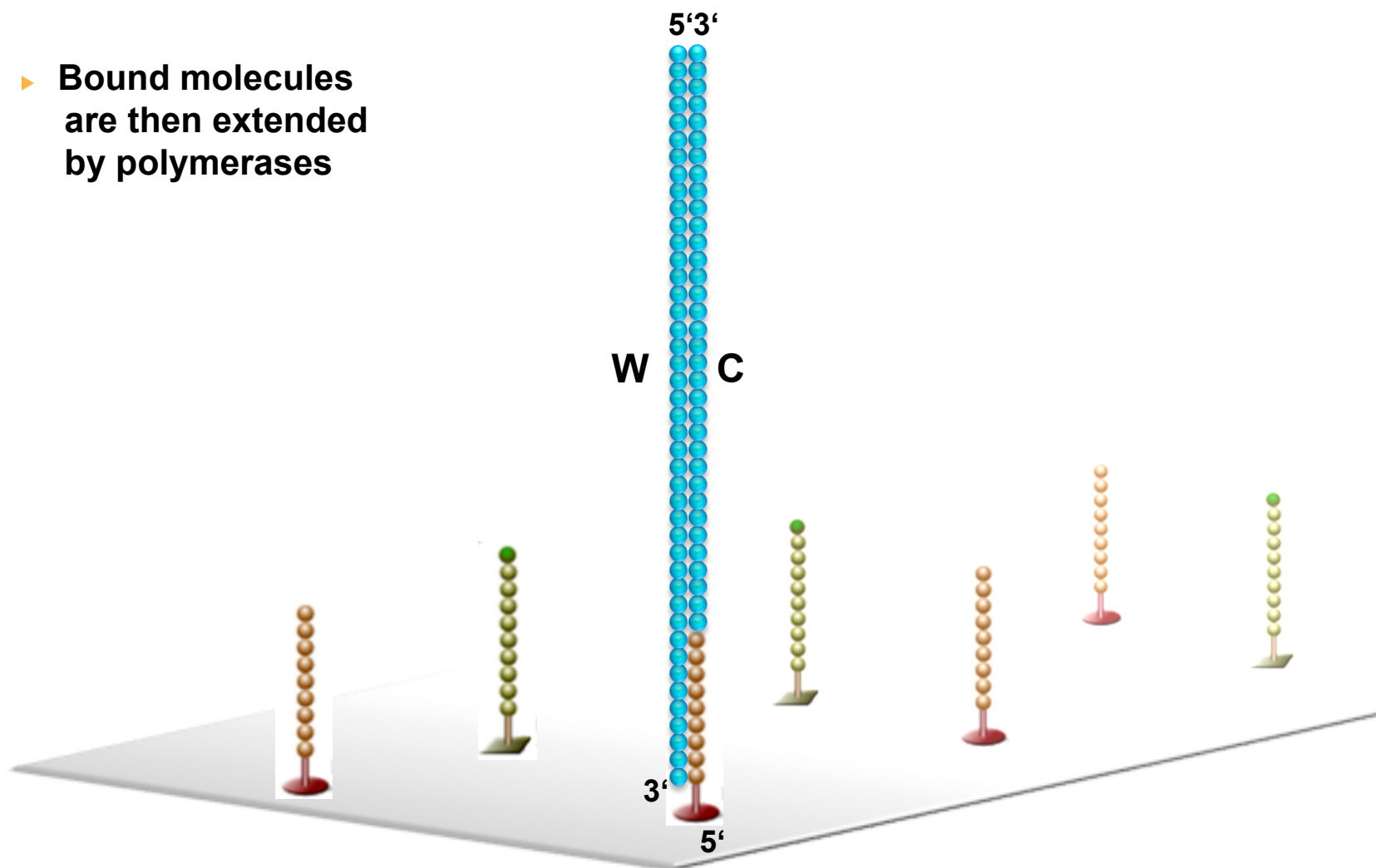
- ▶ Bound molecules are then extended by polymerases





Illumina Cluster Generation

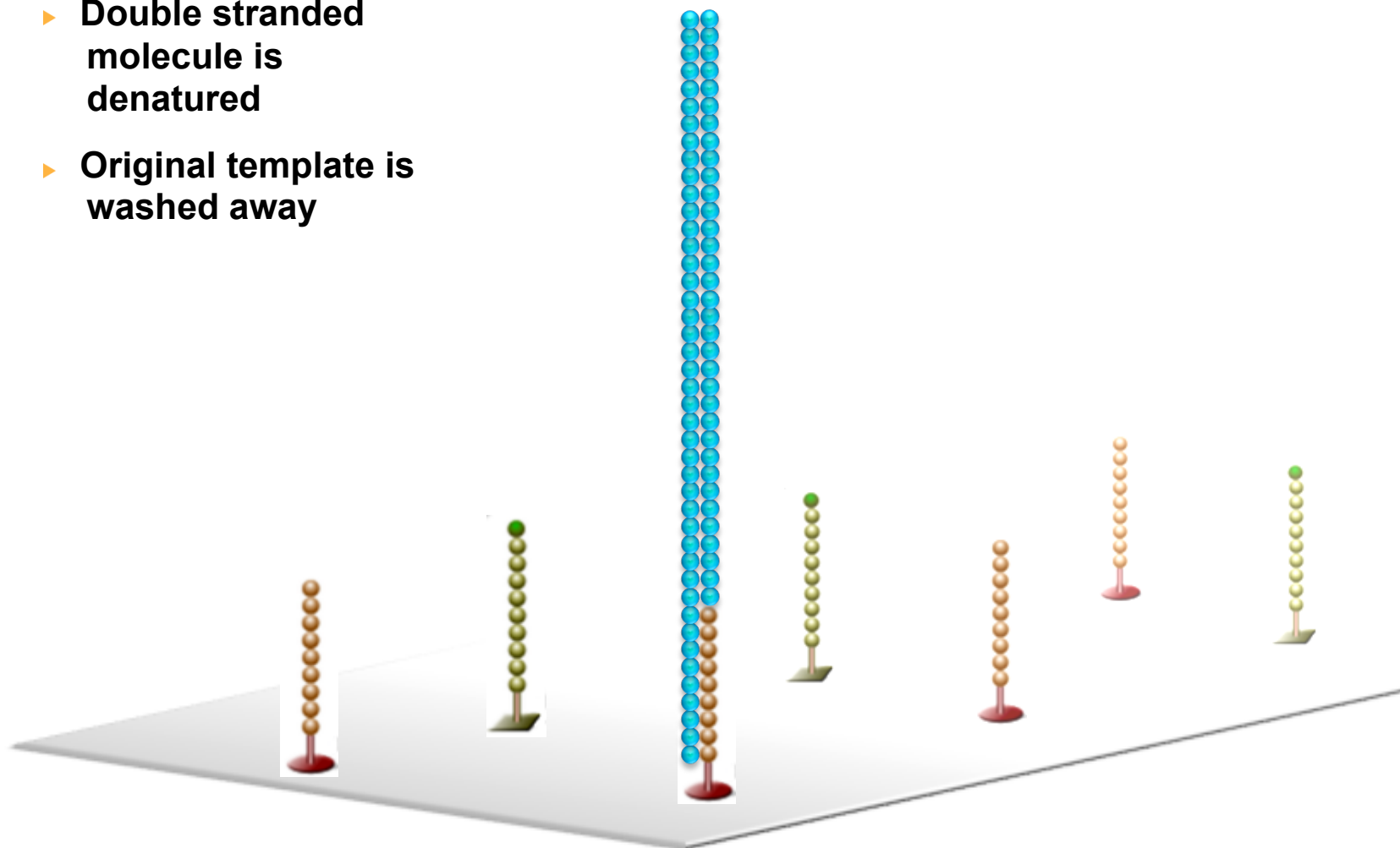
- ▶ Bound molecules are then extended by polymerases





Illumina Cluster Generation

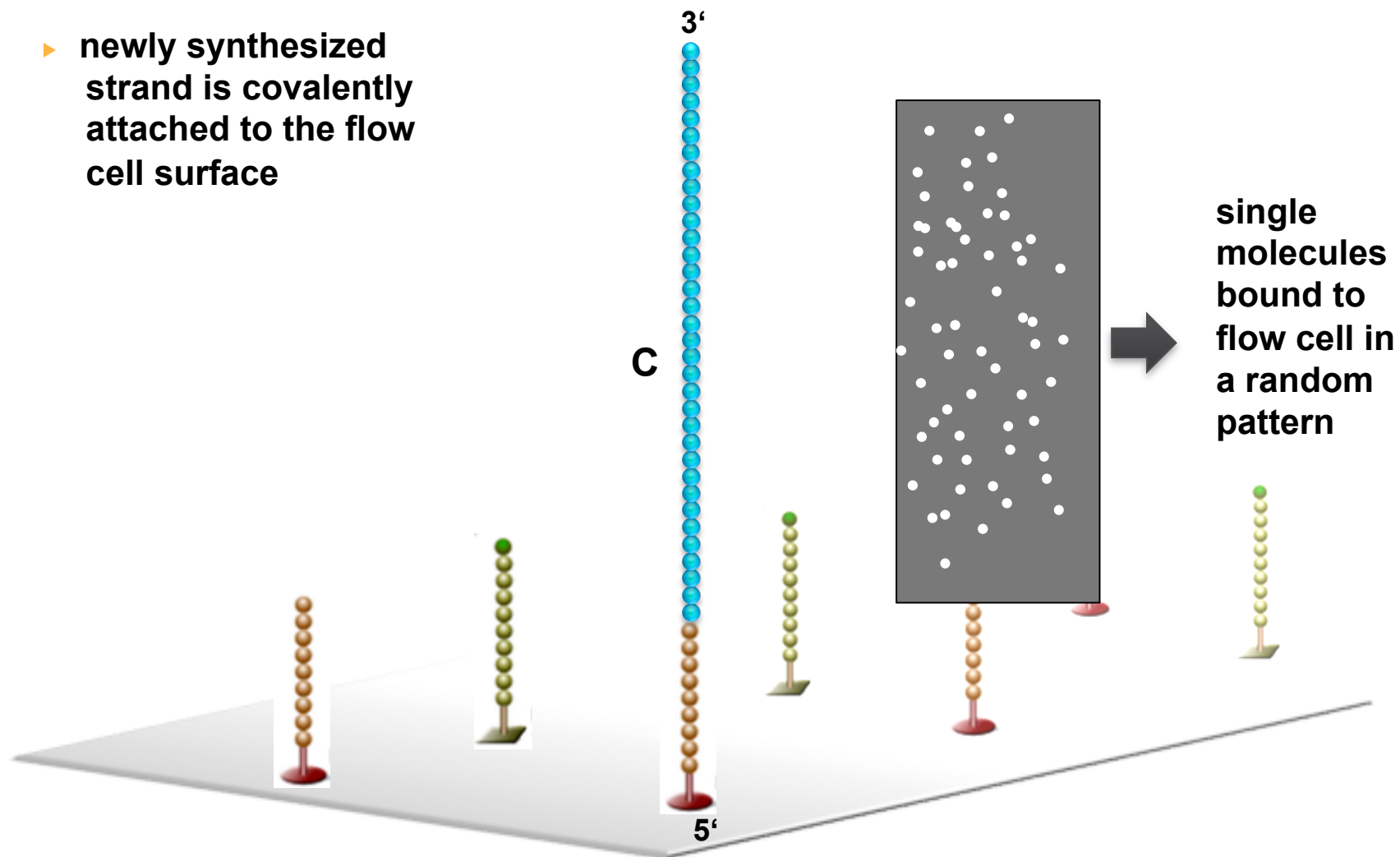
- ▶ Double stranded molecule is denatured
- ▶ Original template is washed away





Illumina Cluster Generation

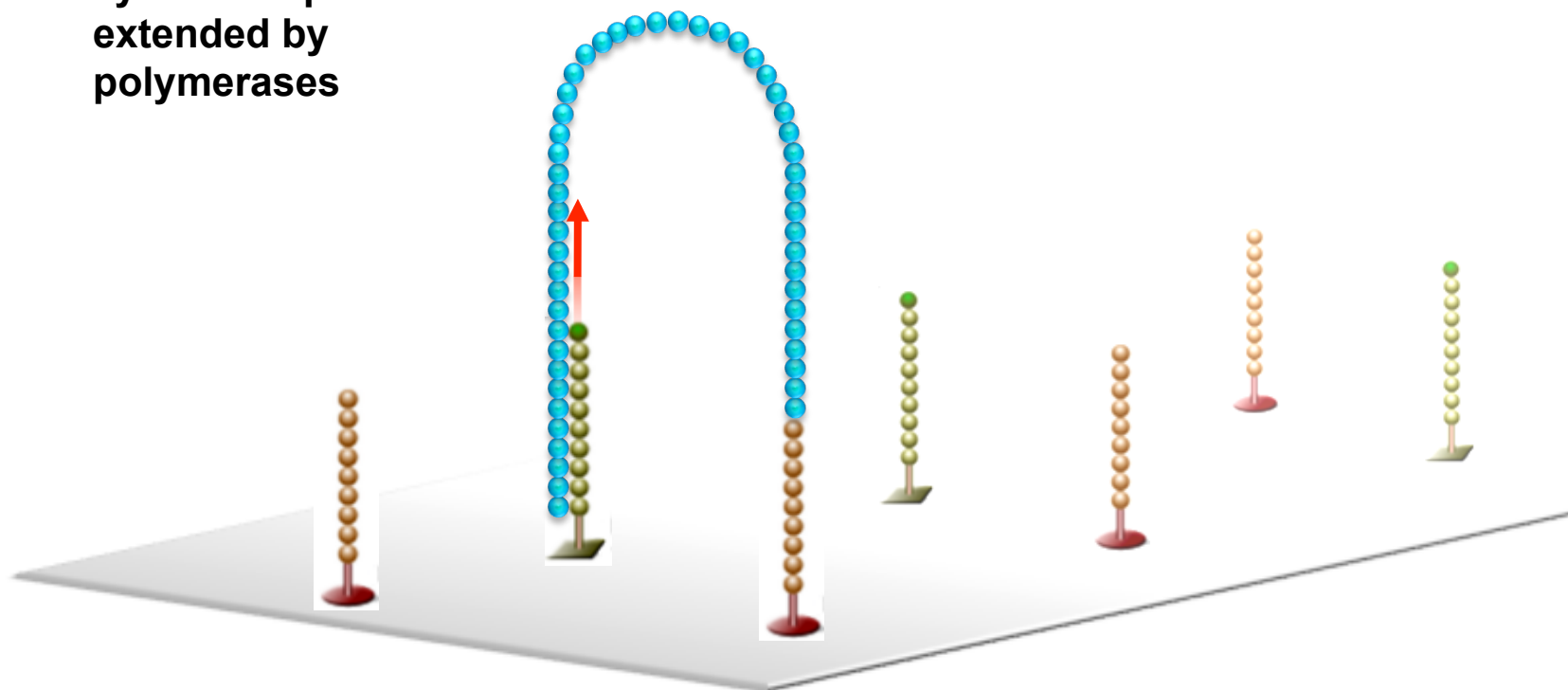
- ▶ newly synthesized strand is covalently attached to the flow cell surface





Illumina Cluster Generation

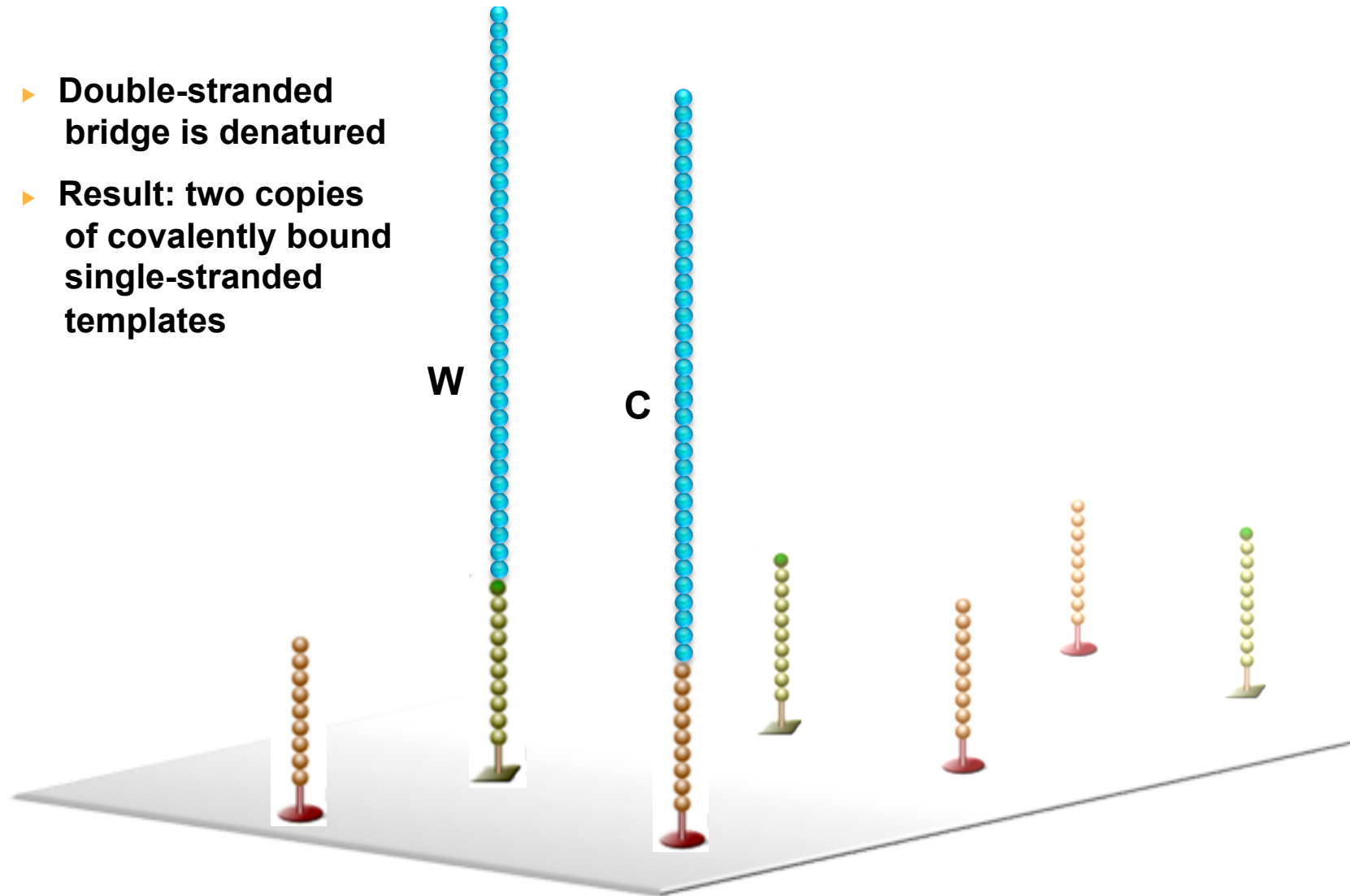
- ▶ **Single-strand flips over to hybridize to adjacent oligos to form a bridge**
- ▶ **Hybridized primer is extended by polymerases**





Illumina Cluster Generation

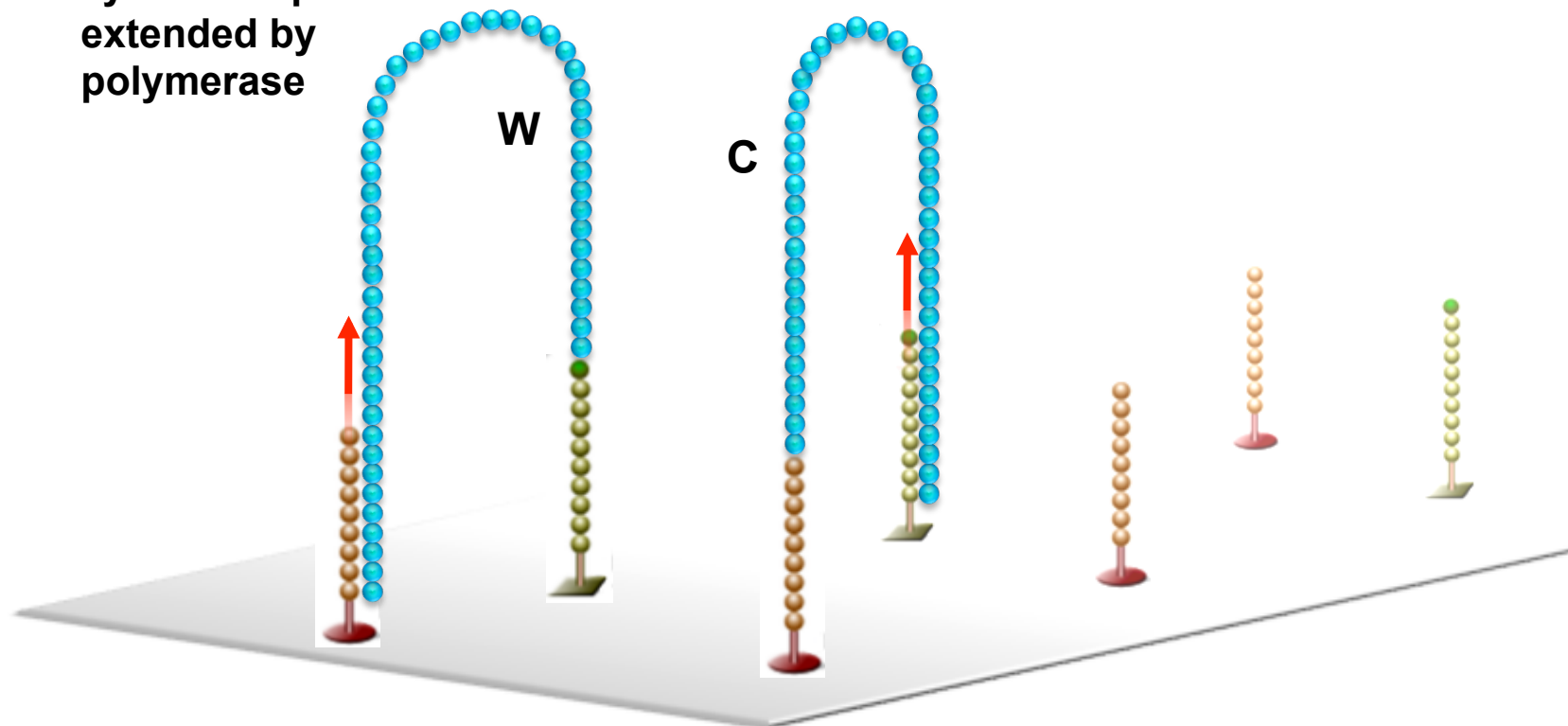
- ▶ **Double-stranded bridge is denatured**
- ▶ **Result: two copies of covalently bound single-stranded templates**





Illumina Cluster Generation

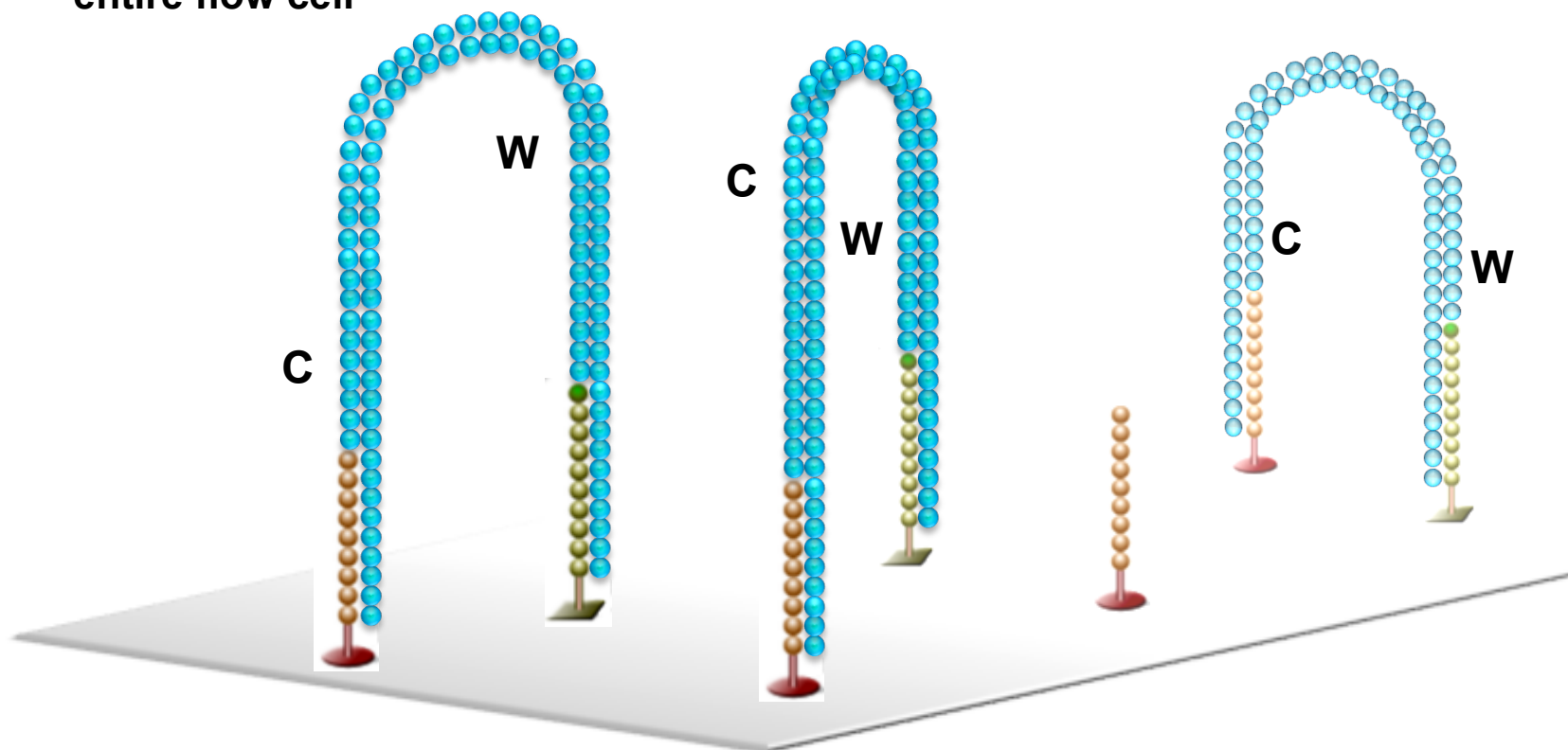
- ▶ Single-strands flip over to hybridize to adjacent oligos to form bridges
- ▶ Hybridized primer is extended by polymerase





Illumina Cluster Generation

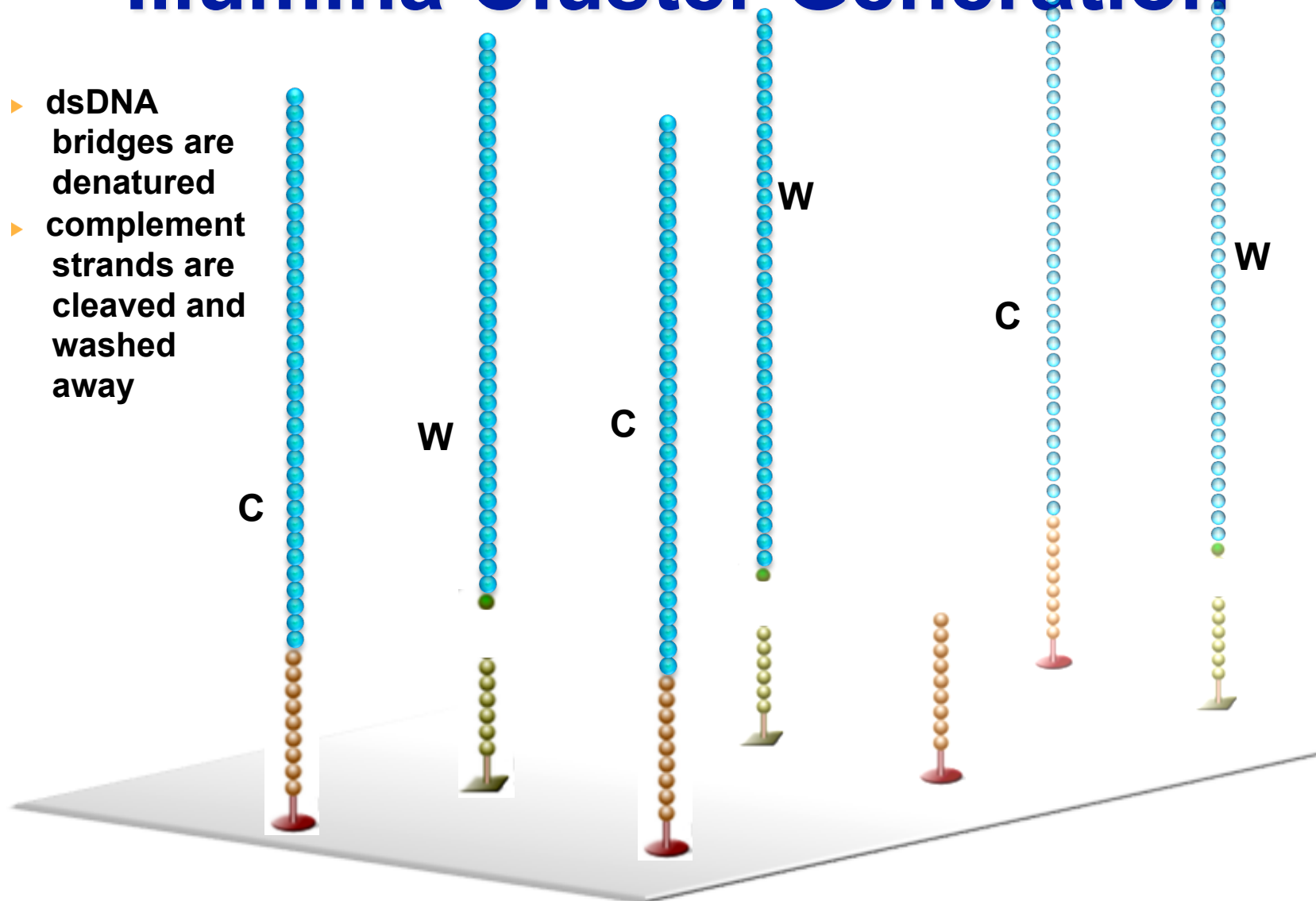
- ▶ Bridge amplification cycle repeated until multiple bridges are formed across the entire flow cell





Illumina Cluster Generation

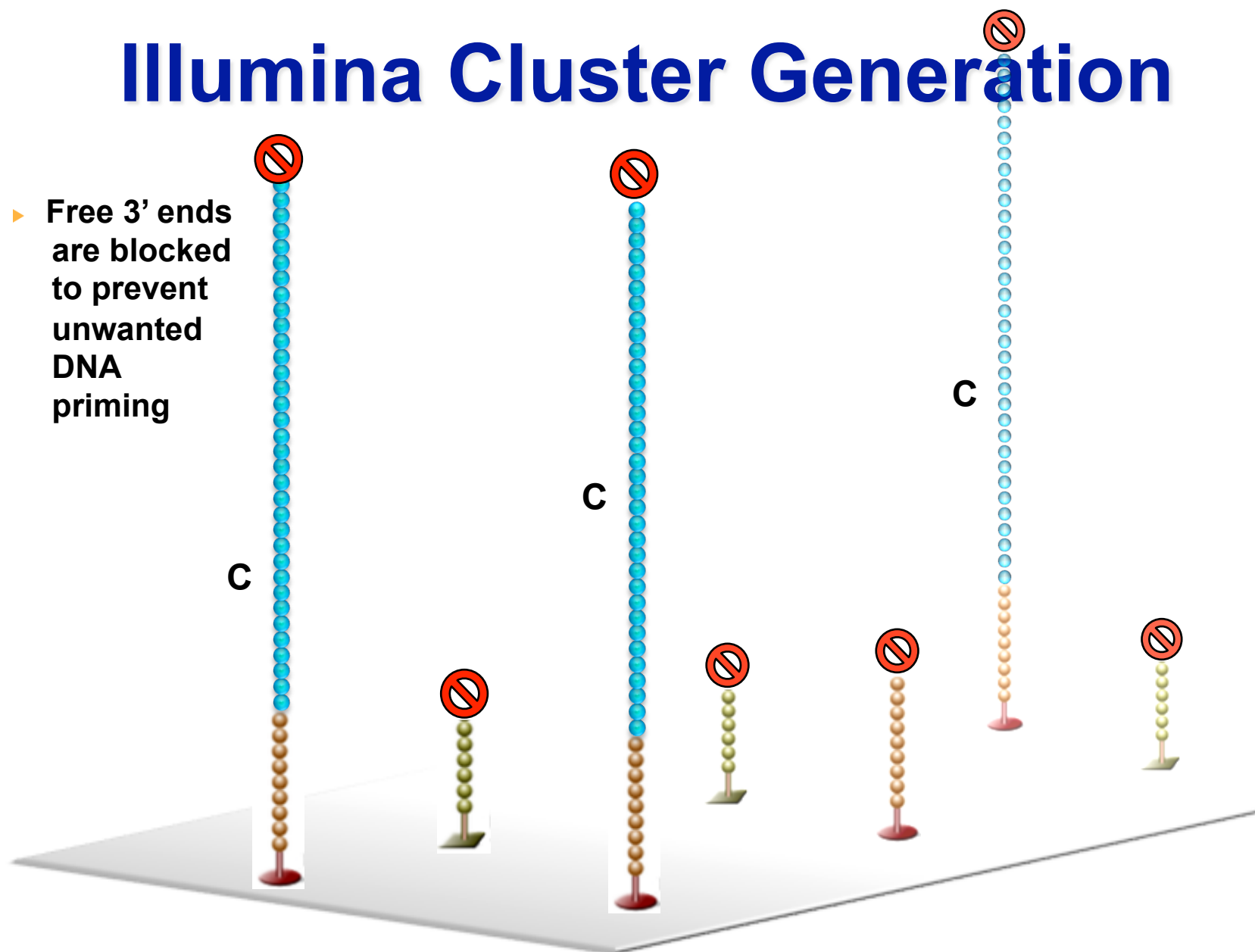
- ▶ dsDNA bridges are denatured
- ▶ complement strands are cleaved and washed away





Illumina Cluster Generation

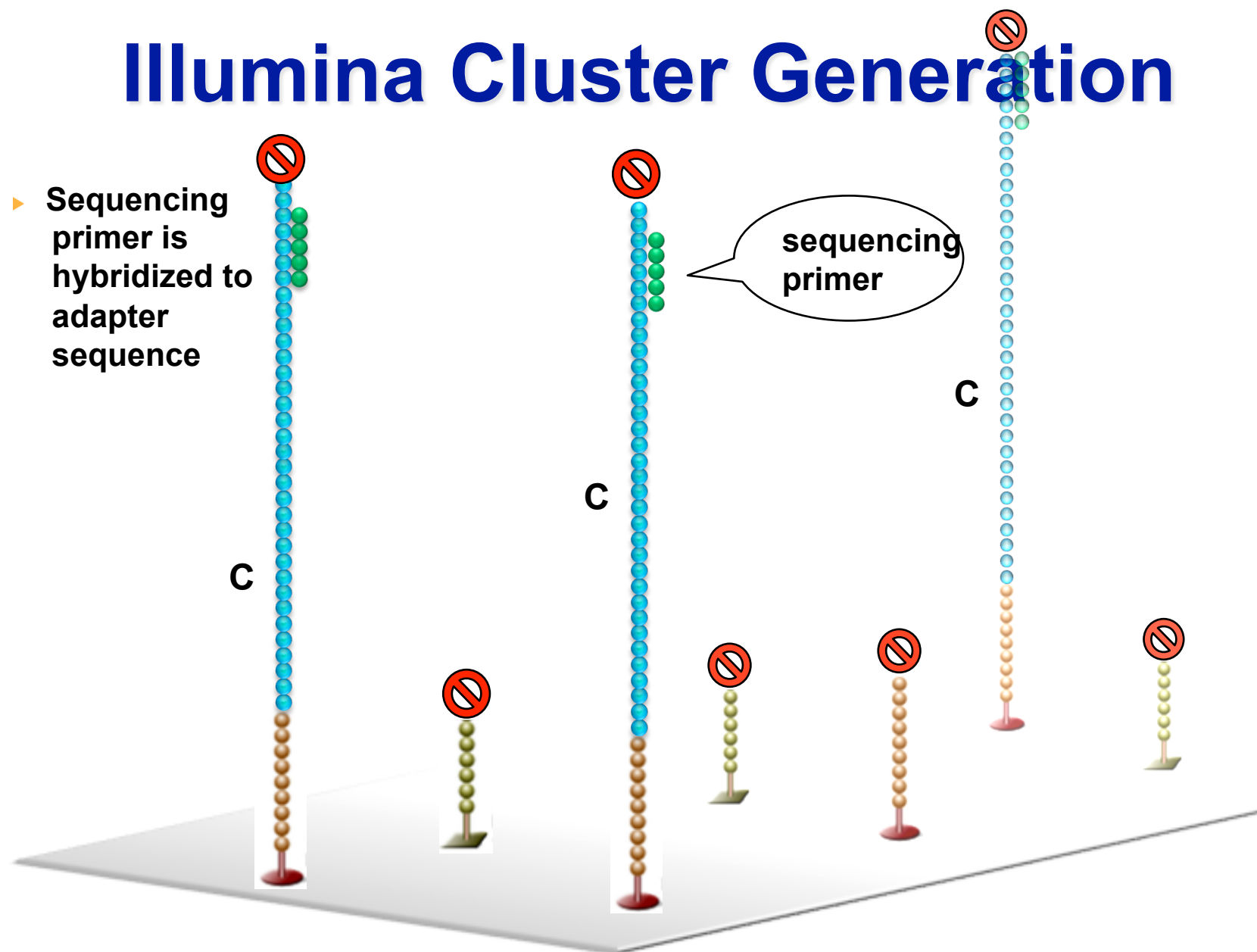
- ▶ Free 3' ends are blocked to prevent unwanted DNA priming





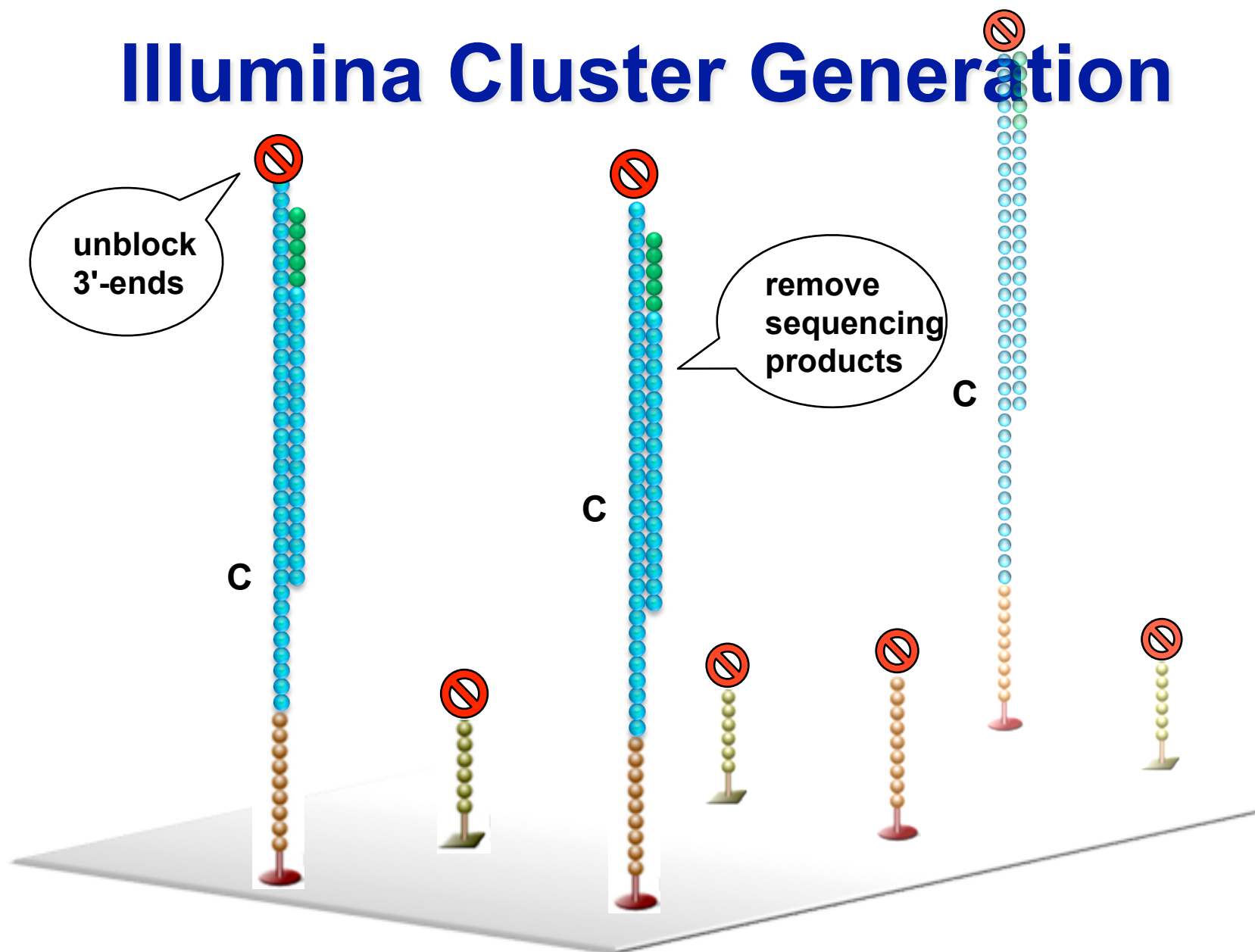
Illumina Cluster Generation

- ▶ Sequencing primer is hybridized to adapter sequence





Illumina Cluster Generation

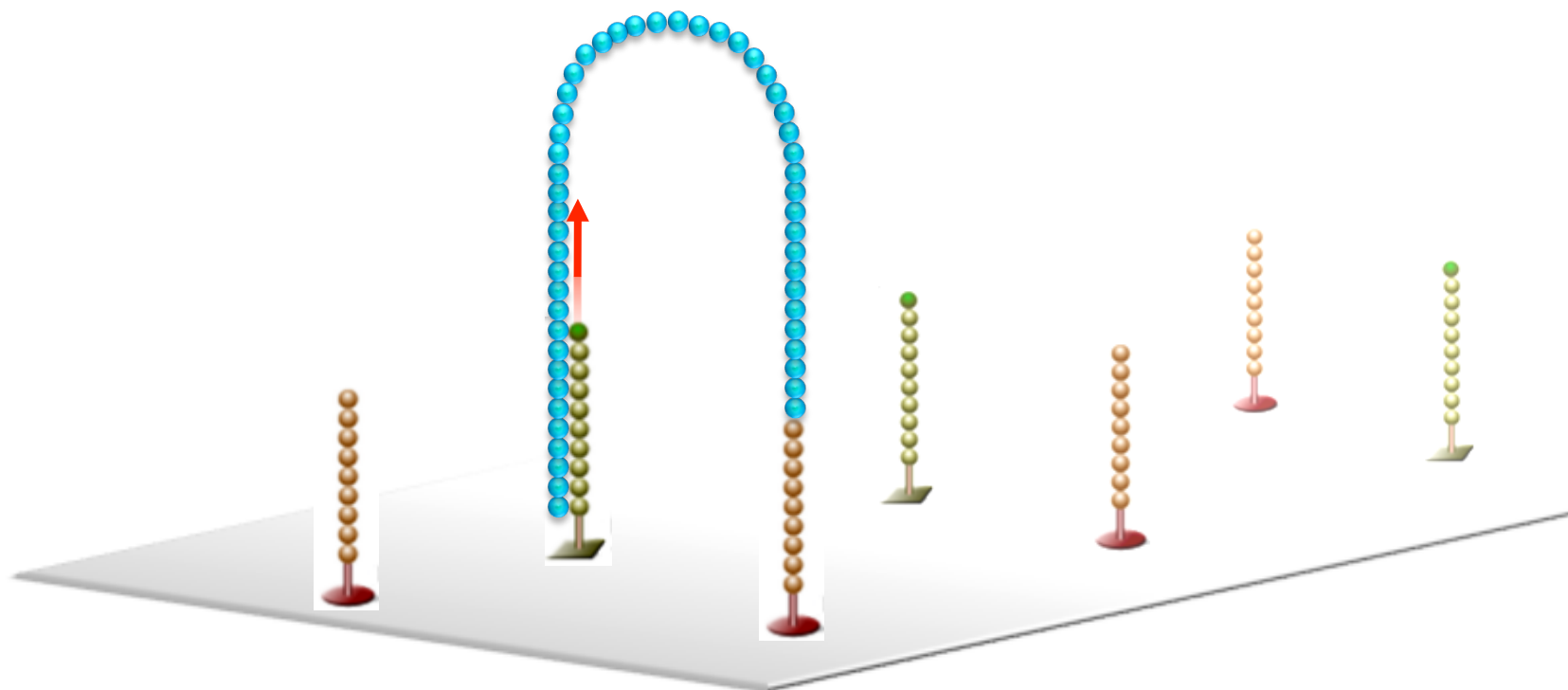


Sequence 2nd strand! (Paired-end sequencing) > W strand needed



Illumina Cluster Generation

- ▶ Bridge formation and 3' extension

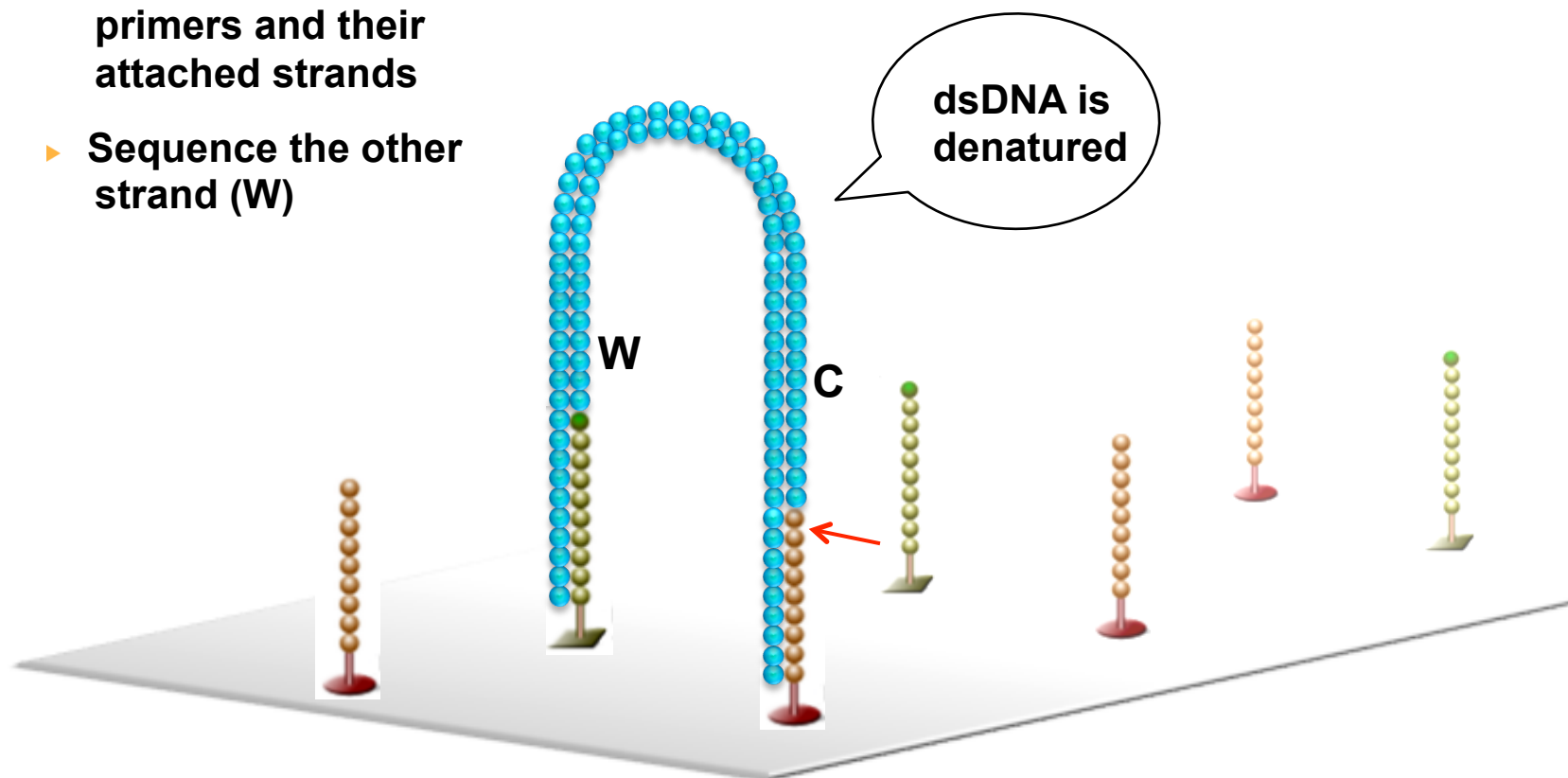


Do it again! New bridge PCR! Then remove C strand...



Illumina Cluster Generation

- ▶ Bridge PCR Round 2
- ▶ **Cleave off** red primers and their attached strands
- ▶ Sequence the other strand (W)



> Paired-end sequencing

Sequenzierstrategien

Illumina/SOLiD



Single end (SE)

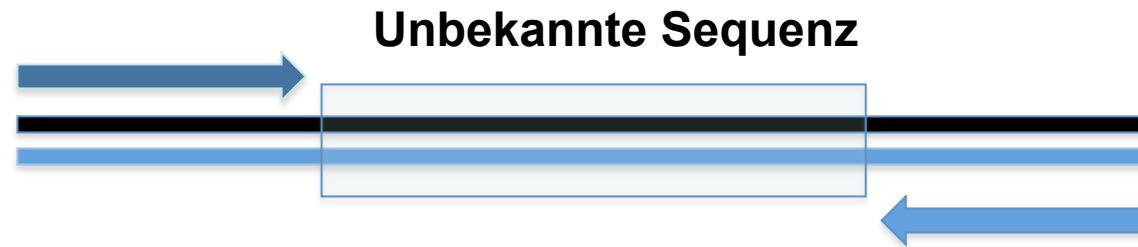


Paired-end (PE), short fragment ends



Mate-pair (MP), another type of paired-end, circularization (cloning)

Paired End Sequencing



„inward-facing reads“, Abstand gering und definiert

**Zusätzlich brauchen wir „long-distance“-Sequenzinformation,
um Contigs relativ zueinander anzuordnen...**

Vergleich von Sequenziermethoden

(2nd generation NGS)



454
(Roche)

Ion
Torrent

Illumina

Complete
Genomics

Sanger

DNA
matrix

Emulsion
PCR

Emulsion
PCR

Bridge
PCR

amplification:
DNA nano balls

Plasmids
Clones
PCR

Sequencing
Method

seq-by-**synth**:
Pyrosequencing

seq-by-**synth**:
Proton release

seq-by-**synth**:
reversible Dye-
Terminators

Seq-by-**ligation**

seq-by-**synth**:
Dye Terminator
96 capillaris

Read
length

av. 600 bp
(up to 1000)

Up to 700

2 x 100 bp
(up to 2 x 300)

70 bp

1000 bp

Data

600 Mbp

1 Gbp

Up to 1.5 Tbp

20-60 Gbp

0,1 Mbp

Runtime

10 hrs

90 min

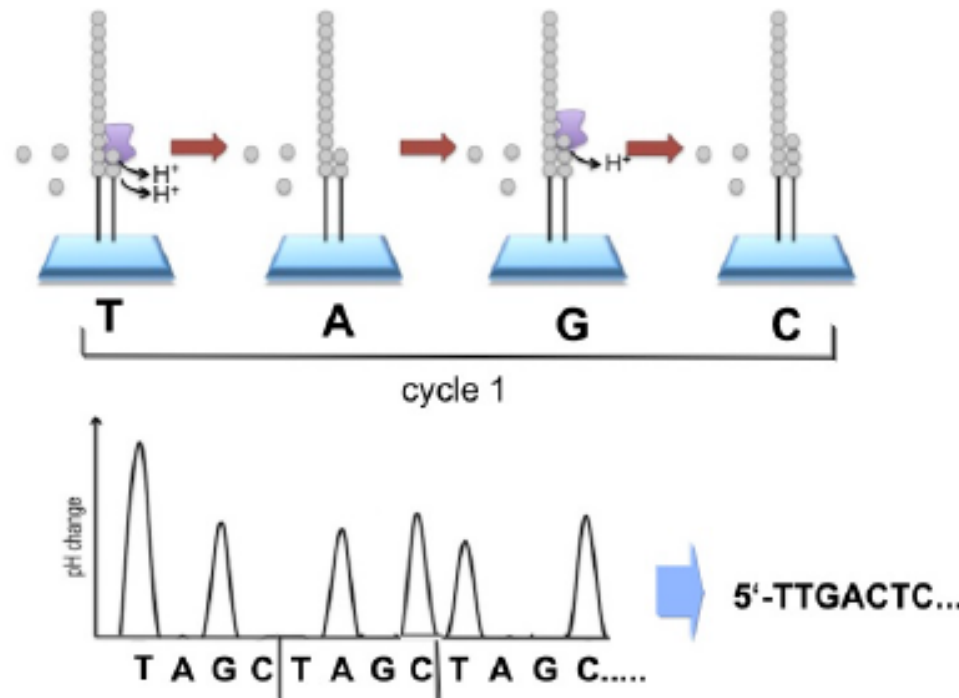
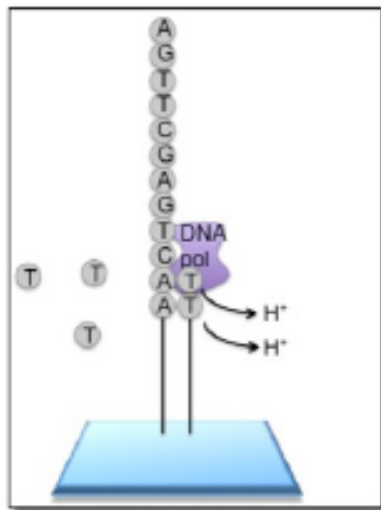
2-10 Days

?

2 hrs

Ion Torrent (Life Tech)

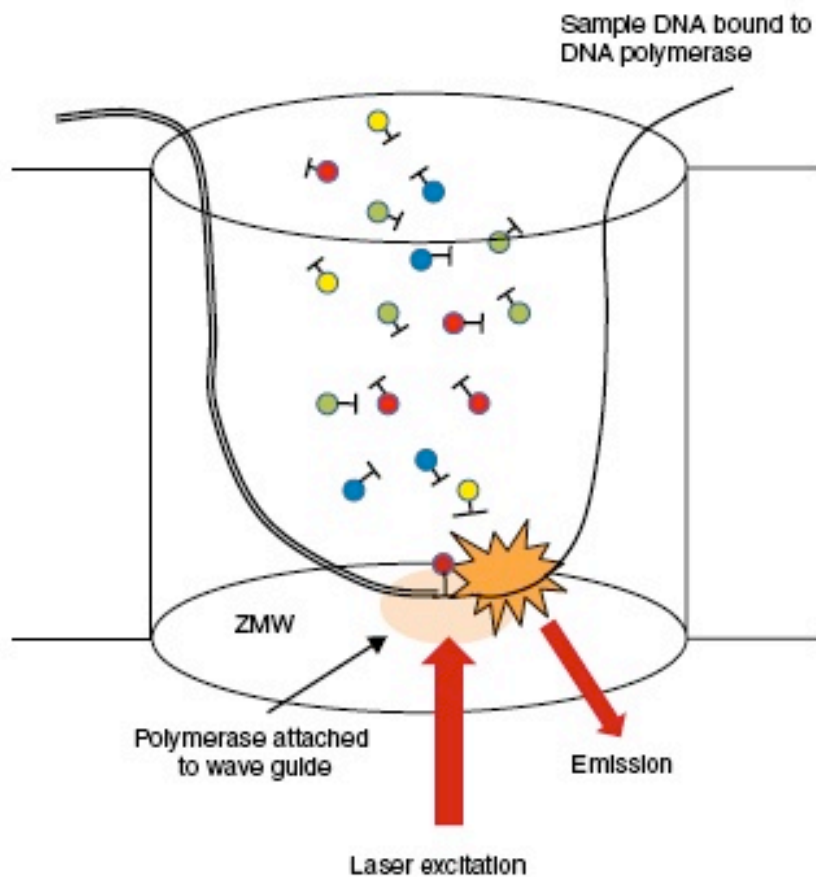
- nukleotide incorporation > proton release > change in pH



Vergleich von Sequenziermethoden (3rd generation NGS)

	Pacific Bioscience	Genia (Roche) Not released	Oxford Nanopore
DNA matrix	Single-mol	?	Single-mol
Sequencing Method	seq-by-synth: labeled hexaphosphate Nt's	seq-by-synth: PEG tagged Nt's + Nanopore	direct sequencing
Read length	20 kbp (mean) 90 kbp max	?	20 kbp (mean) Up to 1.2 Mbp
Data	5-8 Gbp /SMRT Cell (16 Cells)	?	10-20 Gbp / Flow Cell
Runtime	30 min – 10 hrs	?	Realtime (48 hrs max)

Pacific Biosystems: **single molecule** long-range sequencing

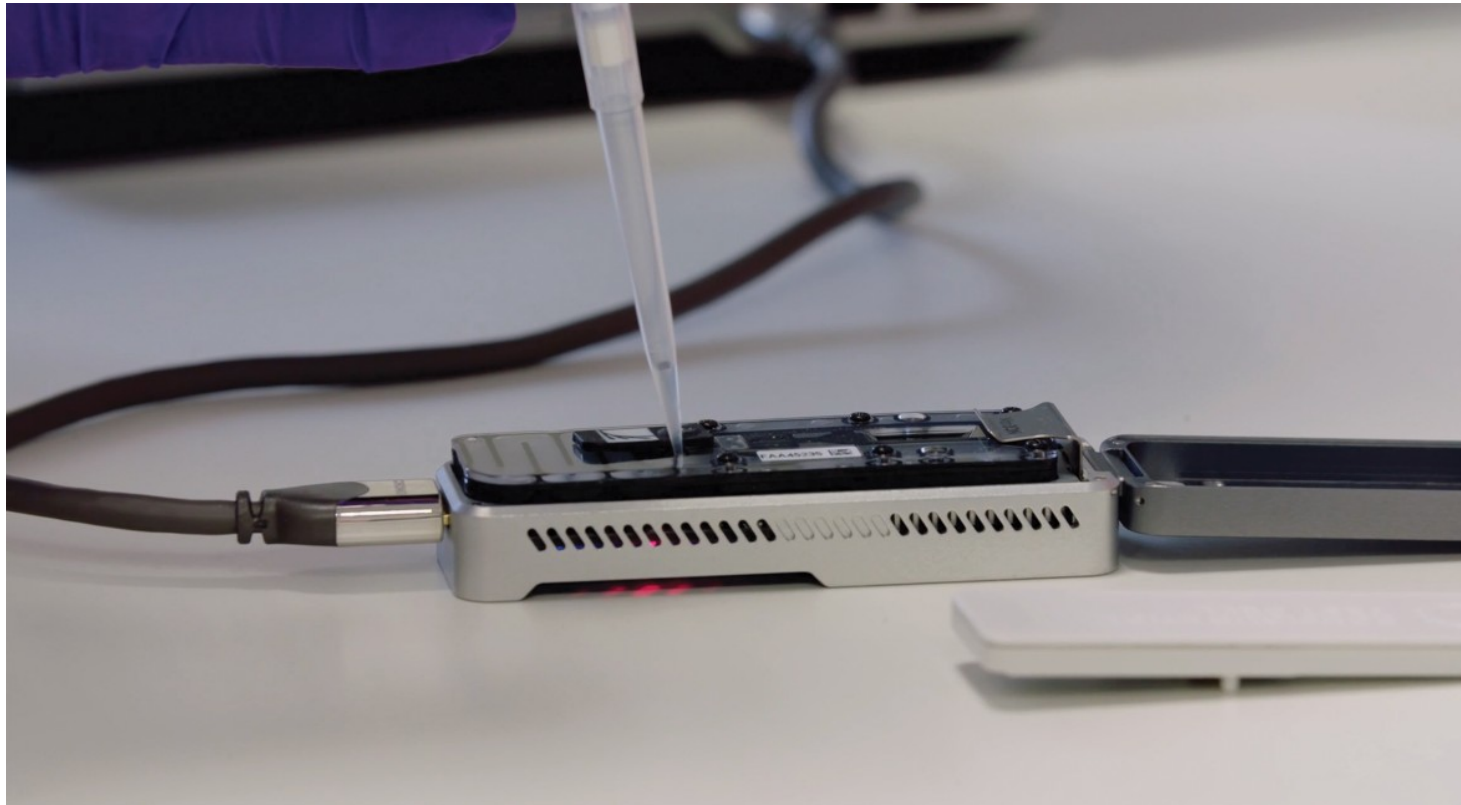


- read length of > 5000 Bp !!
- long-range sequence information !!

but

- expensive machine
- low to medium throughput
- extreme error rate (up to 20%)

Nanopore sequencing: towards **single molecule** detection



**Oxford Nanopore
„MinION“**

Oxford Nanopore MinION

Pro

- Small (90g only)
- Cheap (1000 \$)
- Very long reads (up to 1 MBp)
- Can be used anywhere!
- 1D² Reads (old = 2D)
- direct RNA sequencing

Con

- Low throughput
- High error rate
- New algorithms needed
- Only about 12 % of the pores are atm active

Ebola research in Guinea

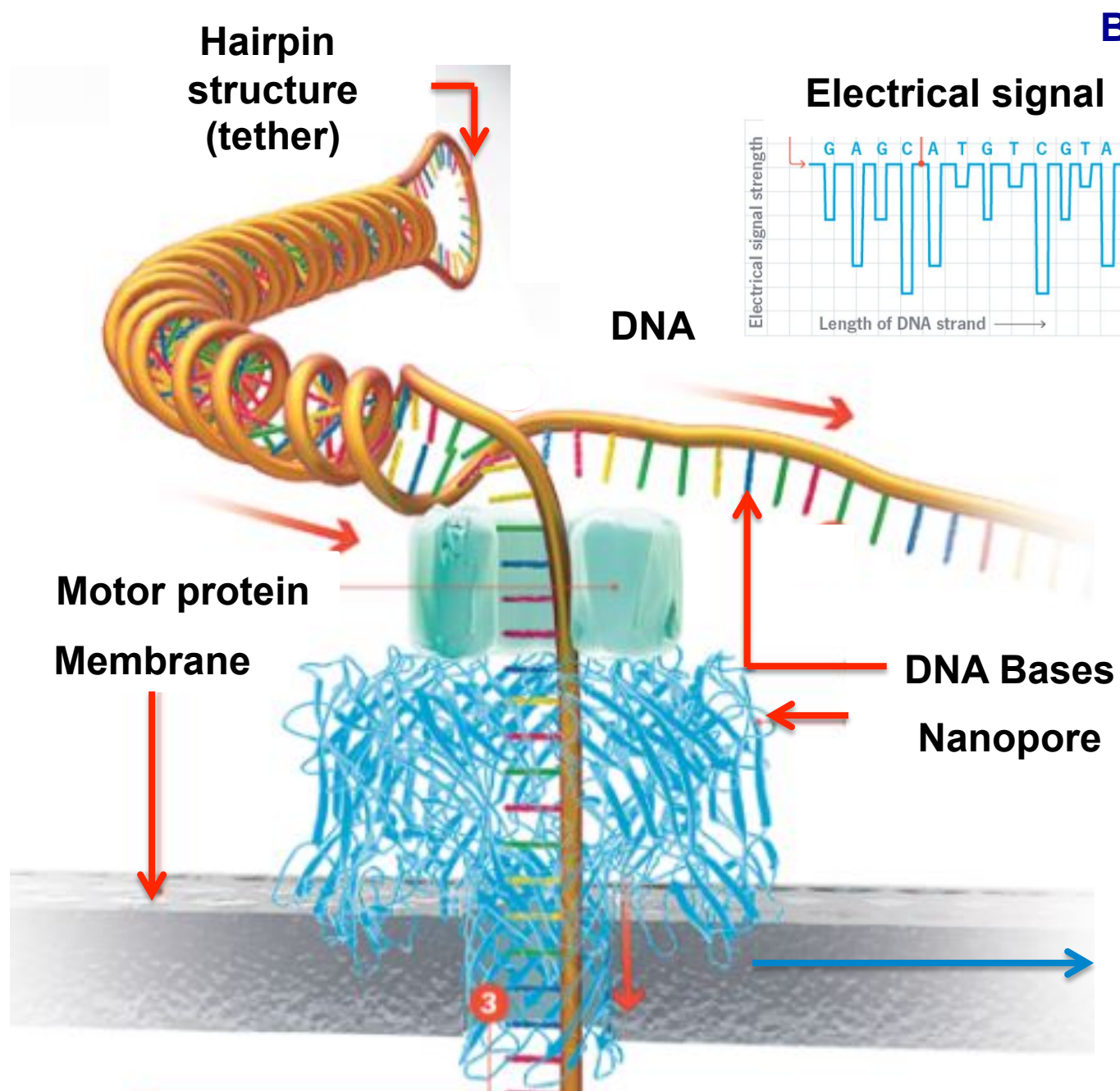


By MUSE/Science Museum of Trento.

First sequencing run in space

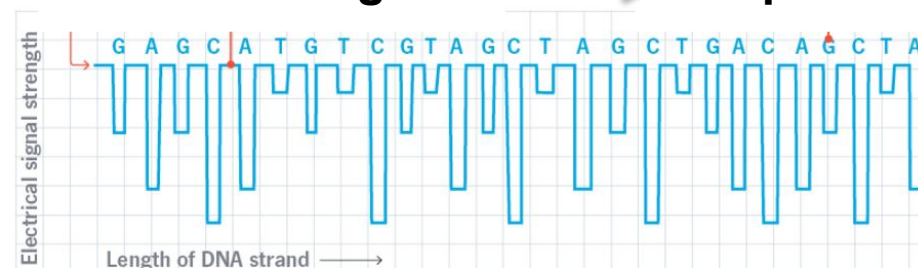


Astronaut Kate Rubins (2016)



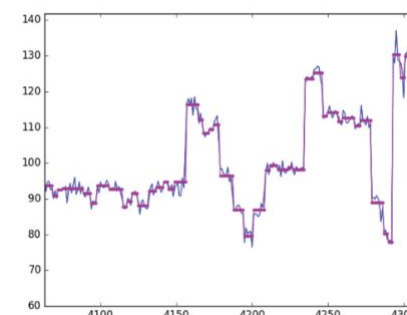
Basecalling

Electrical signal → Sequence



↑
Data processing

Raw current data

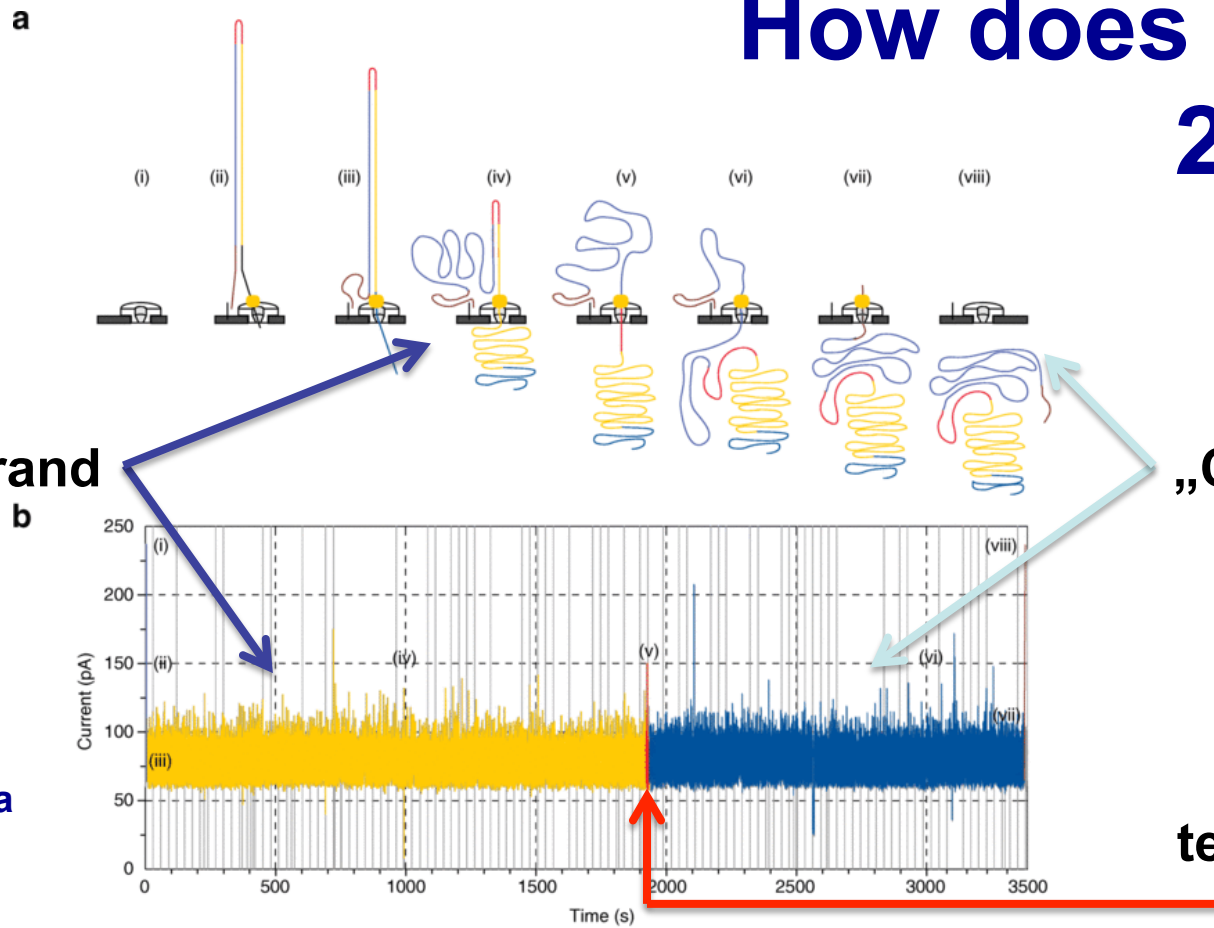


How does it work?

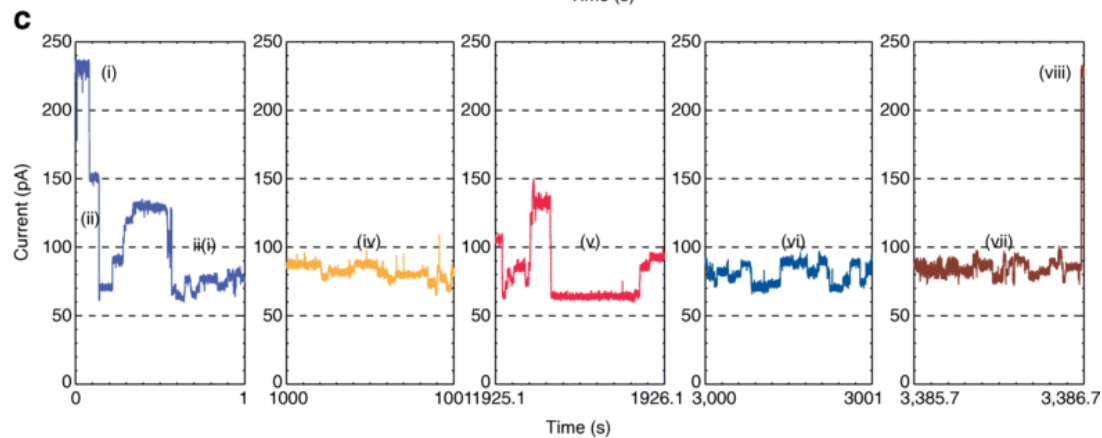
2D-reads

„Watson“ strand

„Crick“ strand



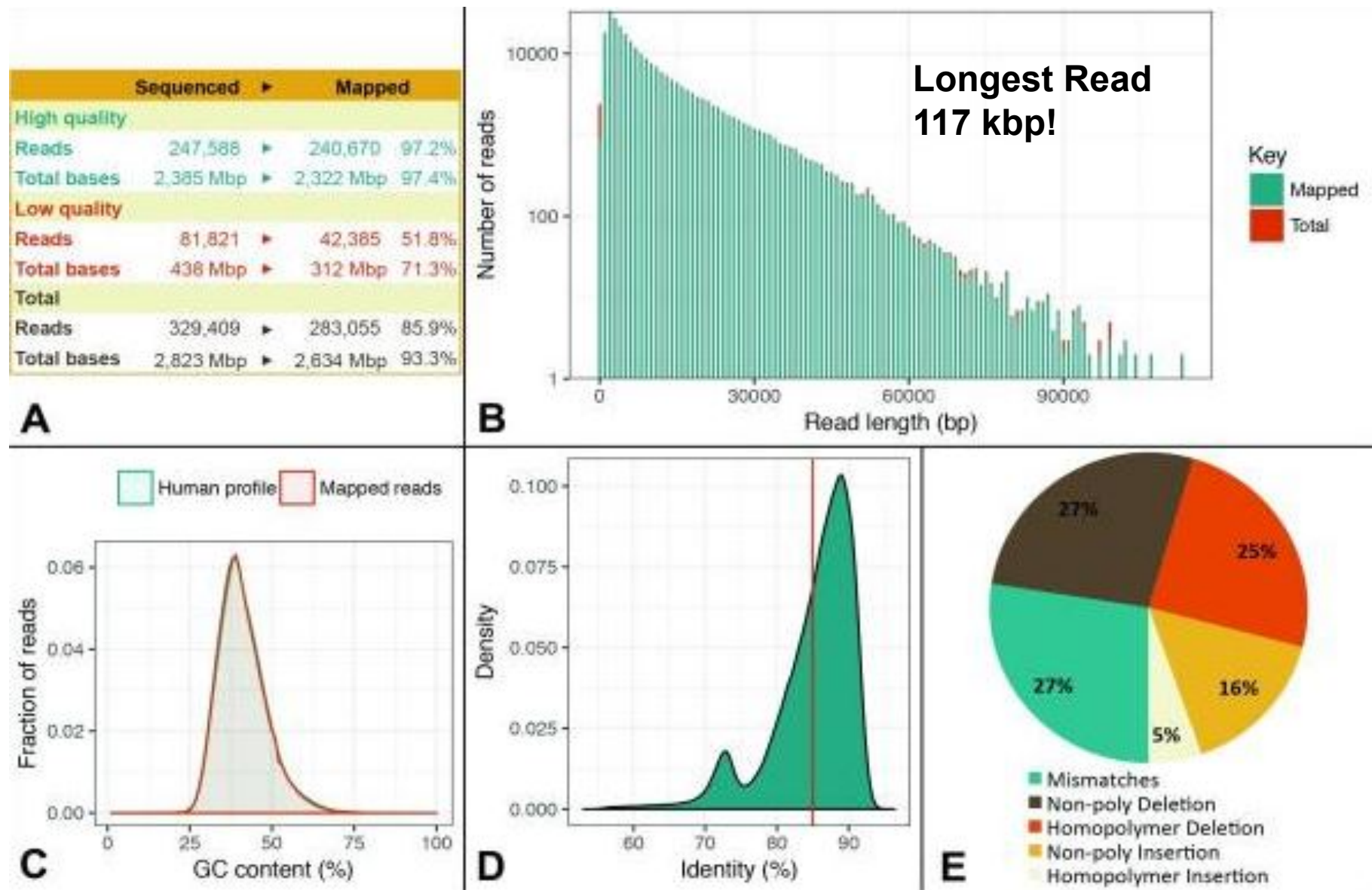
Raw current data
(squiggle data)



processed electrical
signal

Jain et al, 2016

Statistics of a human* genome assembly



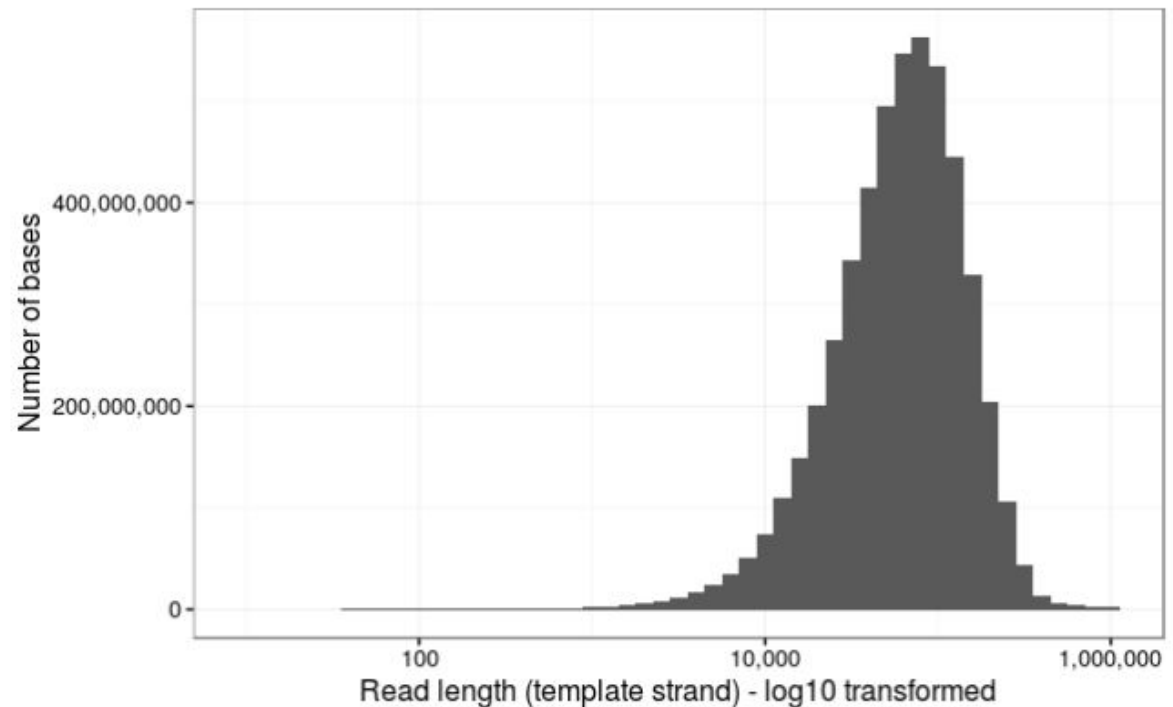
*HAP1 cells (haploid) | Robust long-read native DNA sequencing using the ONT CsgG Nanopore system 2017

Nanopore records

- 1x Coverage of *E.coli* Chromosome (4.6 Mbp) with the 7 longest reads
- Longest read 882 kbp and more
- 1/6 of *E.coli* Genome with one read
- N50 = 63 747 bp



(Loman Labs 2017)



The future...

- Output = ?
- Mobile DNA analysis for everyone and anywhere

PromethION



SmidgION

- 48 Flowcells with 2048 Channels (= 192 MinIONs)
- 6-11 Tbp Output / 24h
- Direct RNA sequencing?
- Direct 5-mC sequencing