

NGS: technologies and challenges

Olga Vinnere Pettersson, PhD
Project Coordinator
NGI/SciLifeLab, Uppsala Node



@OlgaVPettersson

Outline

INTRO

- Sequencing service at NGL-SciLifeLab

NGS general knowledge:

- History of NGS
- Current technologies

NGS Challenges:

- Sequencing artefacts
- NGS sample quality requirements

- Philosophical reflection upon NGS analysis



Operational principles of SciLifeLab

User community

- Open to all Swedish academic scientists on equal terms.
- Consultation and introduction of new protocols.
- Workshops, courses and seminars.

Cost basis

- Academic users of NGI only cover reagent cost.
- Staff salaries at NGI covered by SciLifeLab, VR, and host universities.
- Pre- and service contracts covered by SciLifeLab, VR, KAW and host universities.
- Capital equipment covered by KAW, VR, SciLifeLab.

Quality

- Emphasis on data quality and needs of the users.
- Illumina sequencing and genotyping processes accredited by SWEDAC, ISO/IEC 17025
- Ion and PacBio: accreditation due 2017

We are non-profit

We have technology and knowledge

We want to help you to do GREAT research

We do not want co-authorship

Let us help YOU



NGI Support

Pre-sequencing

- **Project design** via discussions with users
- **Advise** in sample collection and preparation
- Case-to-case **DNA extraction service**

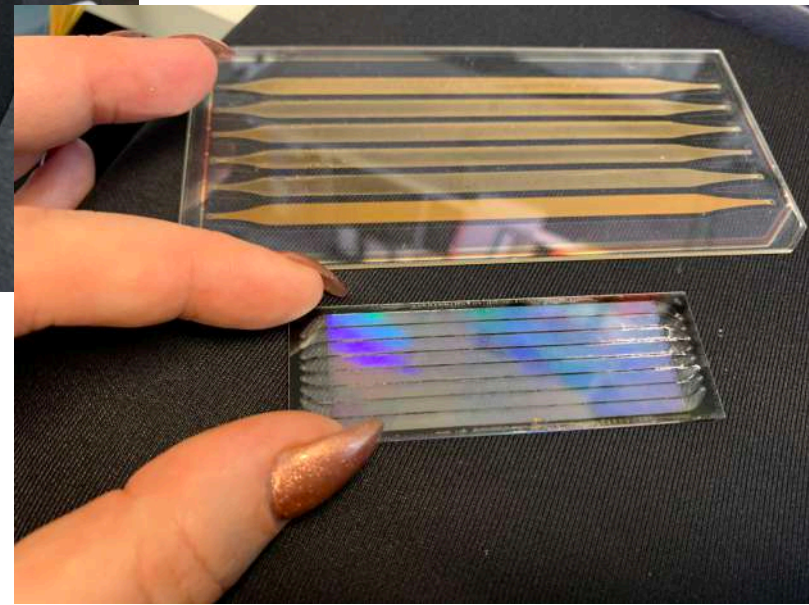
Post-sequencing:

- Control over produced data: making sure data meet our **high standards** in terms of quality and yield.
- Primary **analysis of human genomes** is enabled
- **Genome assembly** of PacBio data is offered as a service
- Data is delivered to **UPPMAX** (Uppsala Multidisciplinary Center for Advanced Computational Science)

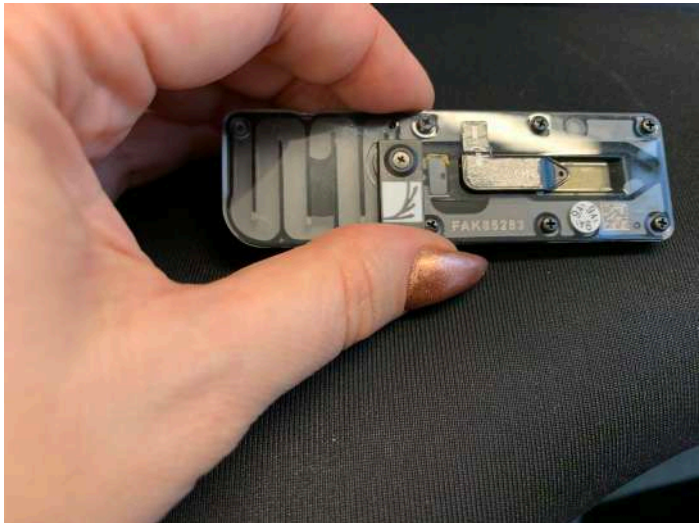
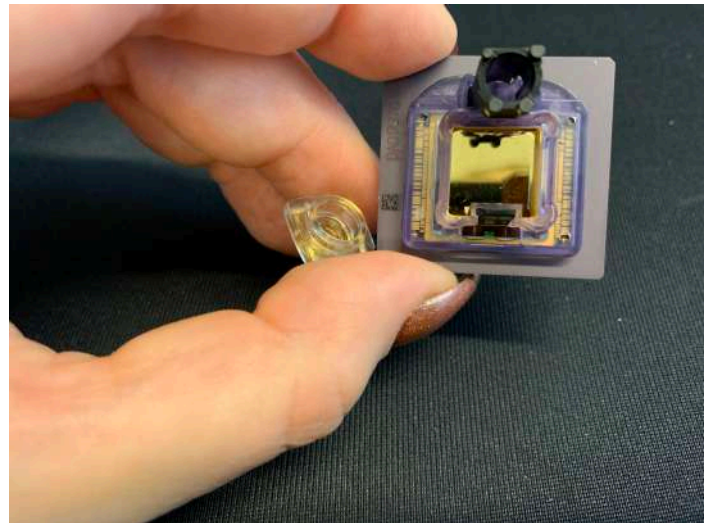
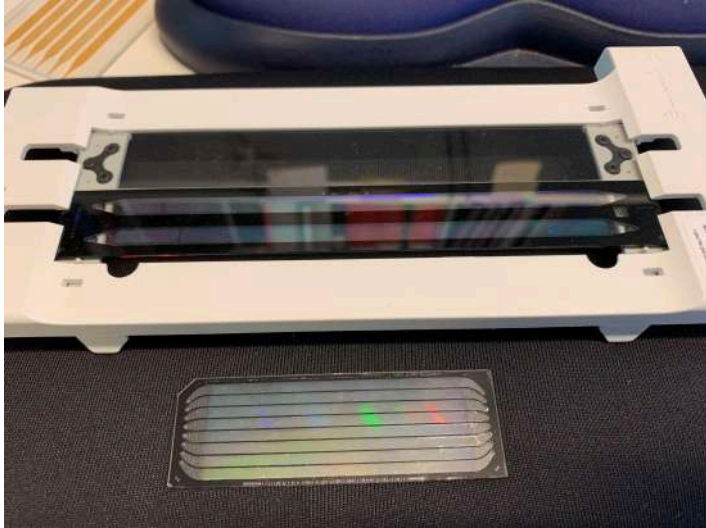
Collaborative projects for technology and method development

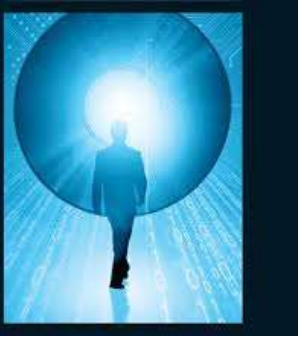
Education

How it looked yesterday



How it looks like now





... paradigm changes

- From single genes to complete genomes
- From single transcripts to whole transcriptomes
- From single organisms to complex metagenomic pools
- From model organisms to the species you are studying
- Personal genome = personalized medicine



Current Technologies



Current leader on the NGS market

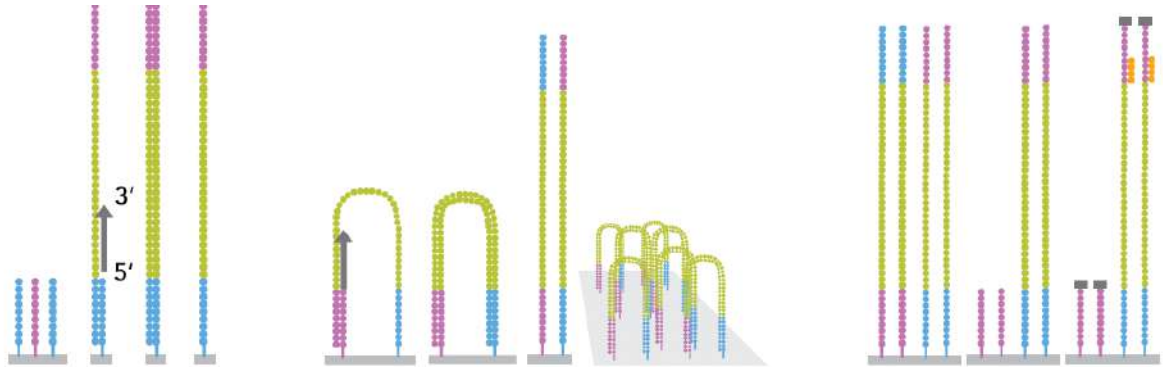
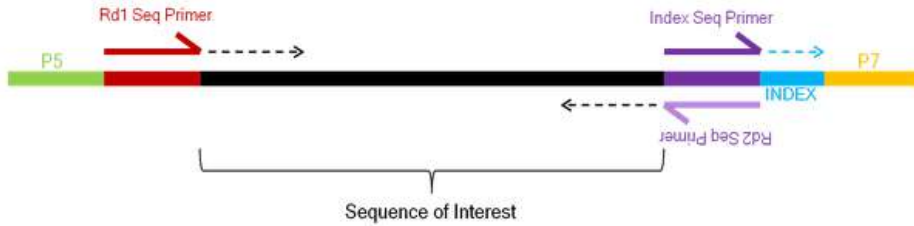
Instrument	Run time	Max output	Max reads/run	Max read length
iSeq	9.5 – 19 hrs	1.2 Gb	4 mln	PE 150
MiniSeq	4-24 hrs	7.5 Gb	25 mln	PE 150
MiSeq	4-55 hours	15 Gb	25 mln	PE 300
NextSeq series	12-48 hours	120-300 Gb	0.4 – 1 bln	PE 150
NovaSeq 6000	13-44 hours	6 Tb	20 bln	PE 250

RIP: HiSeq 2500 & HiSeq X

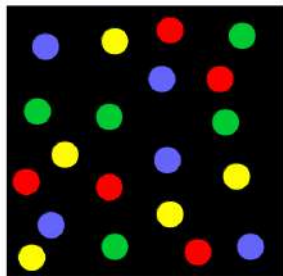
Used for everything



Illumina: bridge amplification



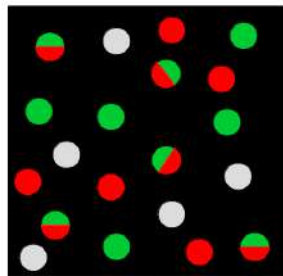
4-Channel system (4 dyes)



4 Filter channels



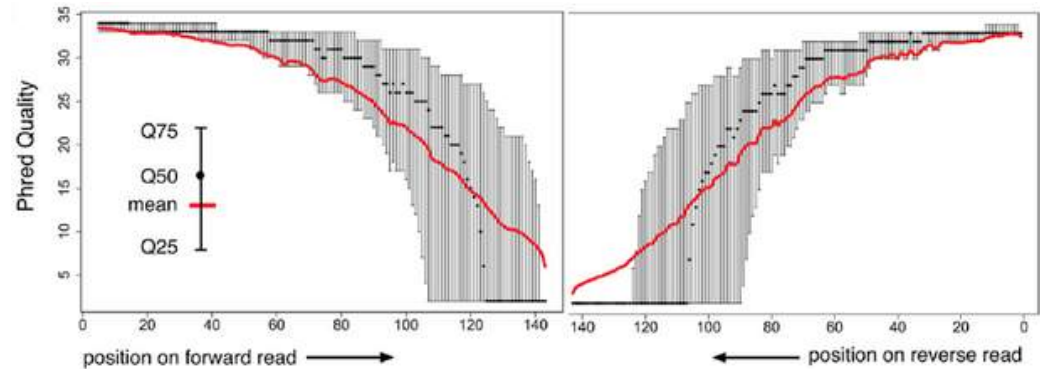
2-Channel system (2 dyes)



2 Filter channels

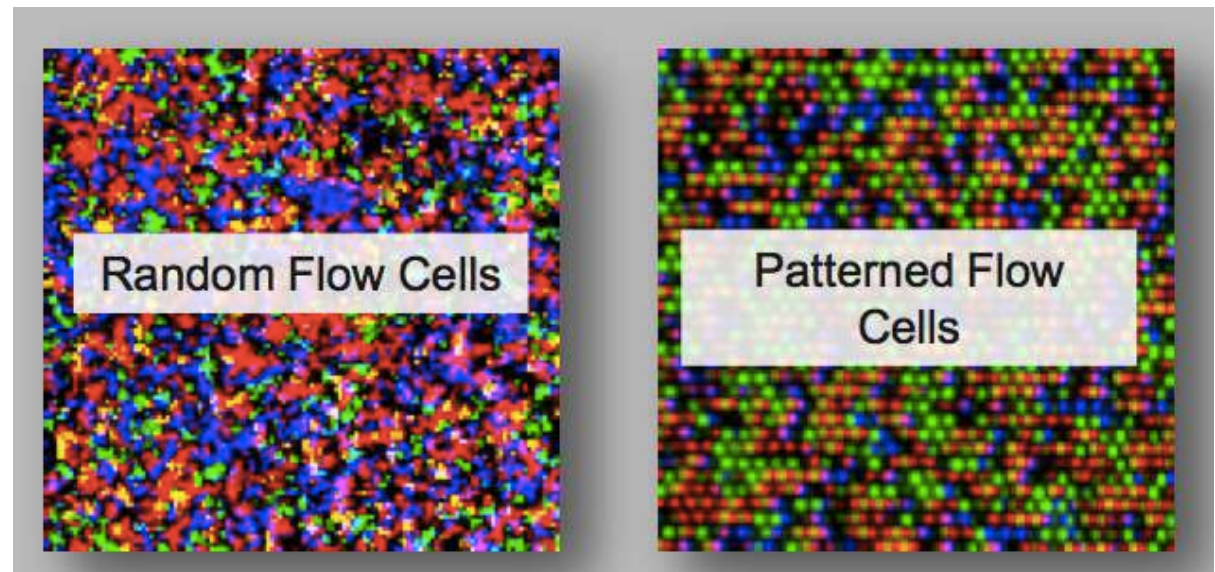
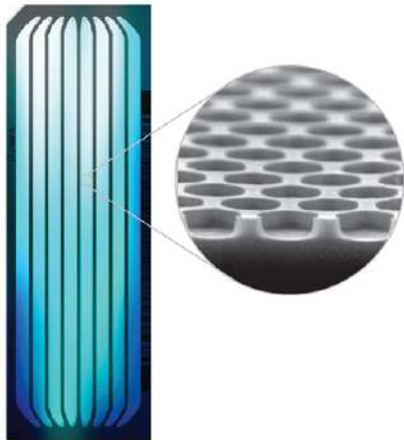


*No detected dye



Illumina sequencing: before vs now

Patterned flow cells introduced on HiSeq X and NextSeq systems



MiSeq flow cells still do not have a pattern



Ion S5 XL

Chip:	Run time	Output	Max reads/ run	Max read length
510	2.5-4 hrs	0.3 - 0.5 Gb	2-3 mln	SE 400 bp
520	2.5-4 hrs	0.6-2 Gb	3-6 mln	SE 600 bp
530	2.5-4 hrs	3-8 Gb	15-20 mln	SE 600 bp
540	2.5-4 hrs	10-15 Gb	60-80 mln	SE 400 bp
550	2.5-4 hrs	18-20 Gb	100-130 mln	SE 200 bp

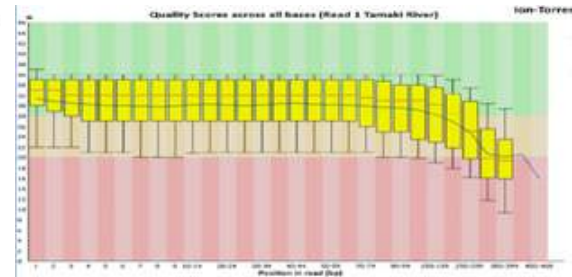
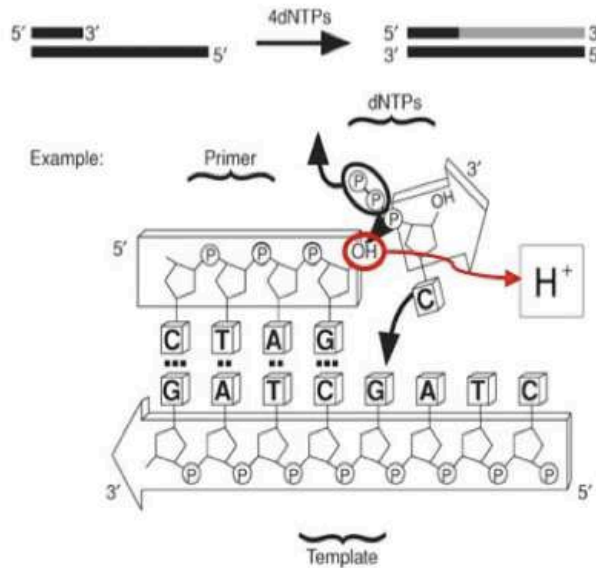
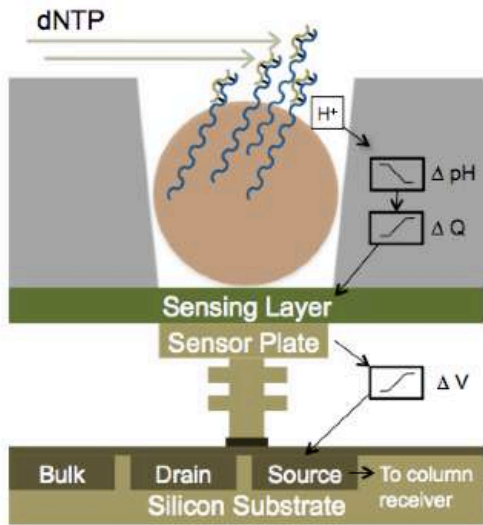
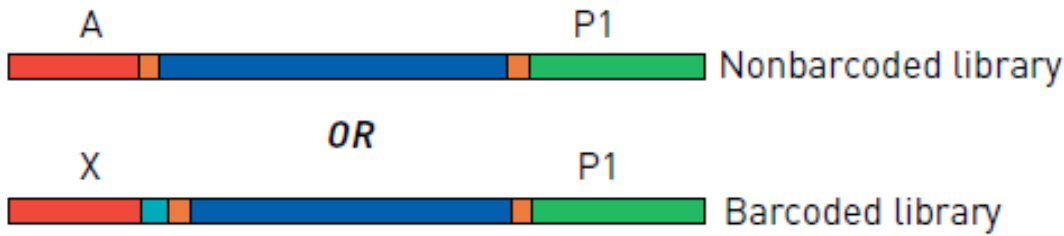
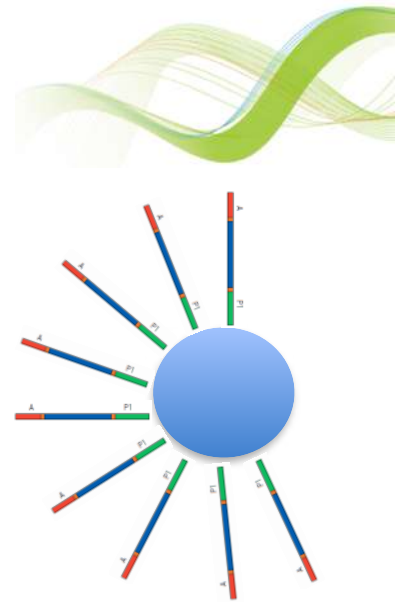
RIP: IonTorrent PGM, IonProton

Clinical applications mainly
Standard analysis directly on the instrument

Multiplex-PCR panels



Ion Torrent: H⁺ ion-sensitive field effect transistors





PACBIO®



Instrument	Run time /SMRT	Output /SMRT	Max reads / SMRT	Max read length*
RSII	30 min – 6 hrs	500 Mb – 2 Gb	50 000	40 kb
Sequel	30 min – 20 hrs	2 – 35 Gb	200 000	60 kb
Sequel II				
HiFi	30 hrs	<i>320 Gb</i>	<i>4 mln</i>	25 kb
CLR	15 hrs	<i>300 Gb</i>	<i>3 mln</i>	120 kb

Single Molecule Real Time sequencing: SMRT

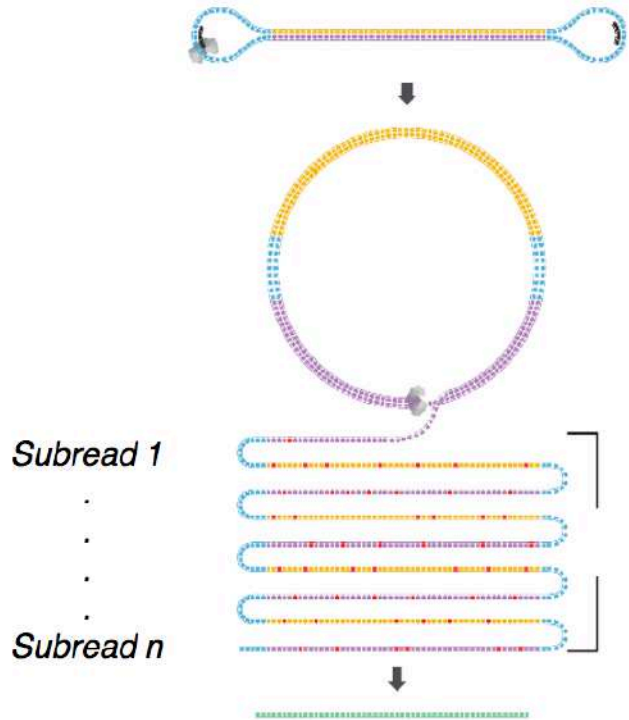




TWO MODES OF SMRT SEQUENCING

Circular Consensus Sequencing (CCS) Mode

Inserts 10-20 kb



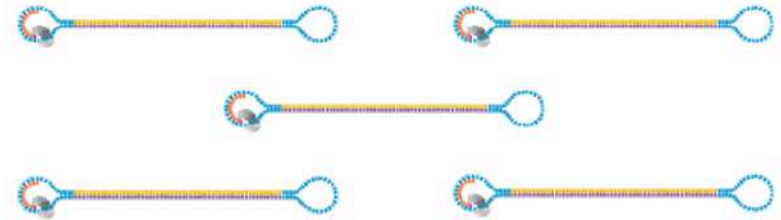
Qv 20+



Single-molecule consensus sequence

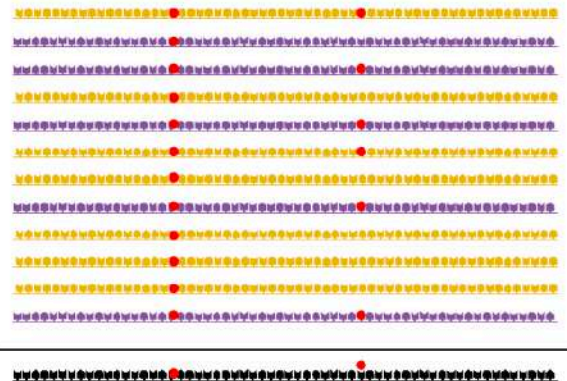
Continuous Long Read (CLR) Sequencing Mode

Inserts >25 kb, up to 175 kb



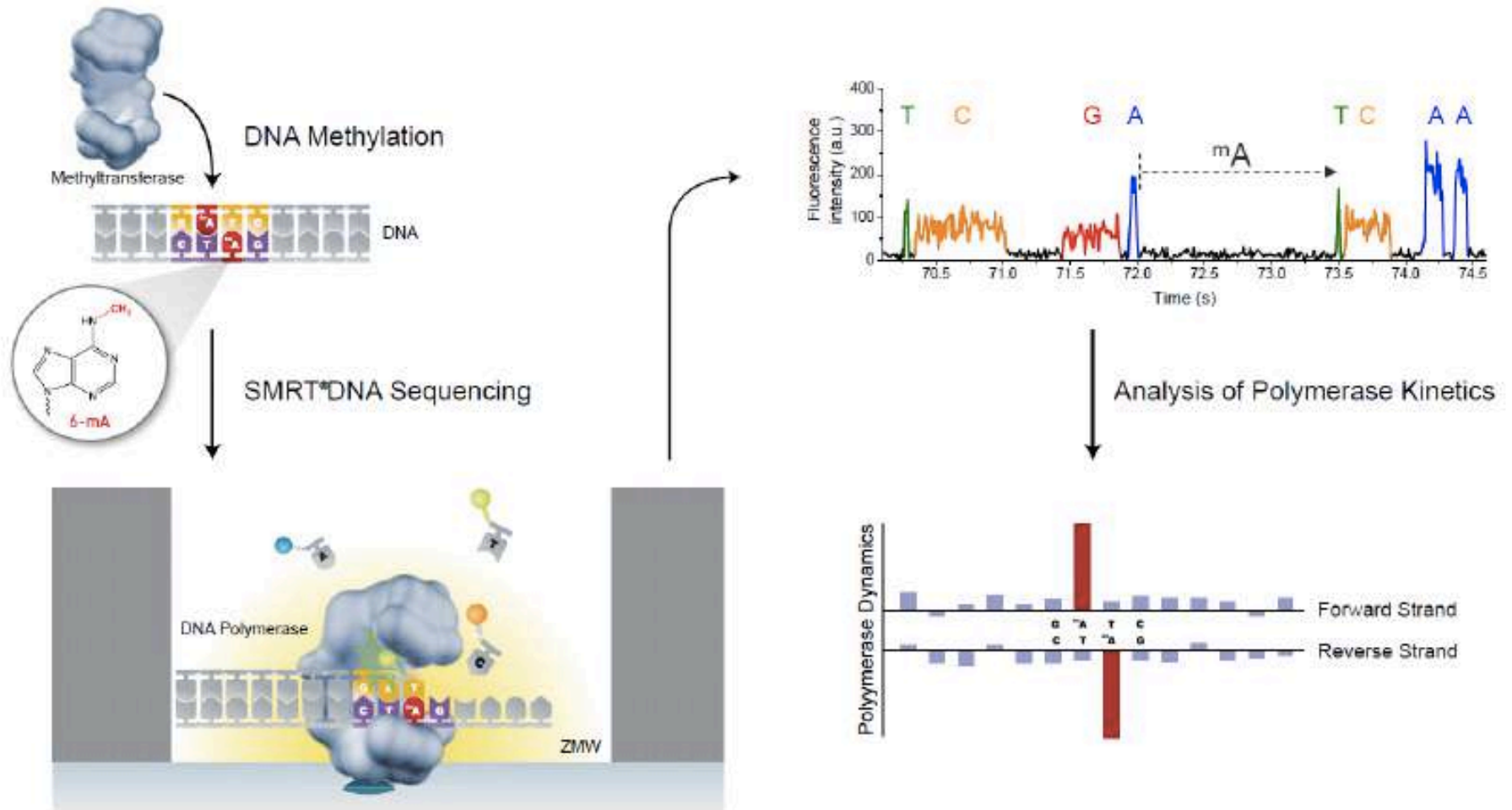
CLR 1

CLR n



Multi-molecule consensus sequence

Base Modification: Discover the Epigenome



Detect base modifications using the kinetics of the polymerization reaction during normal sequencing



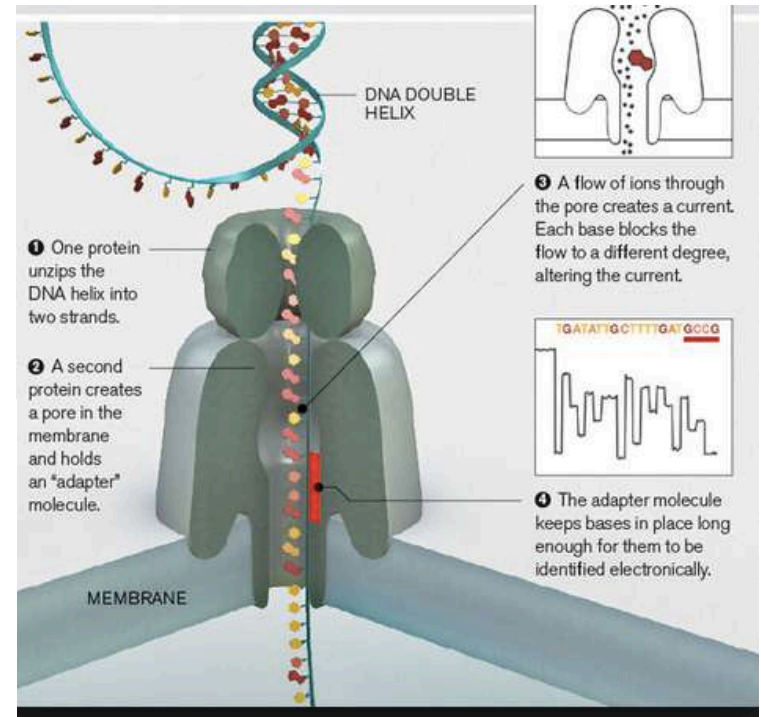
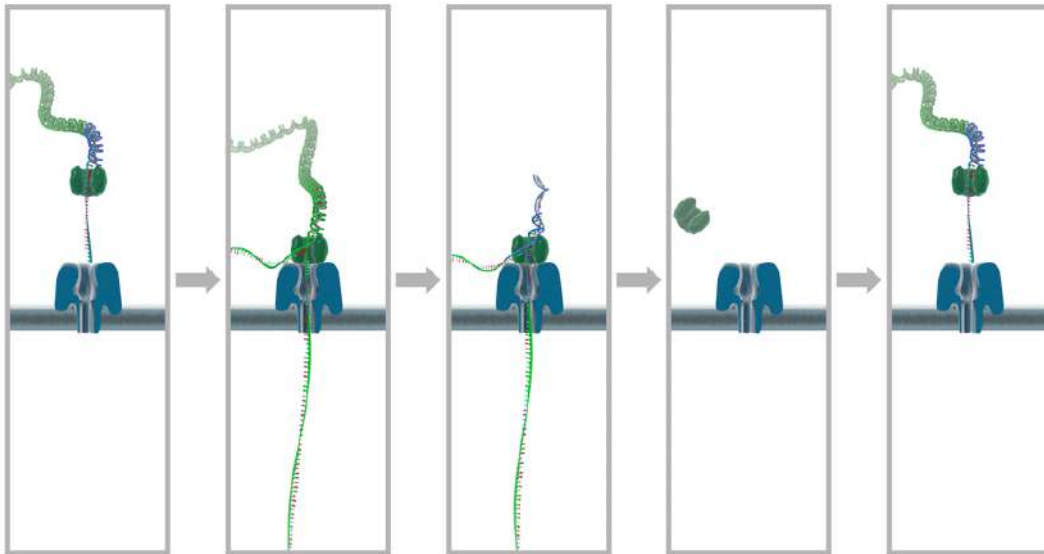
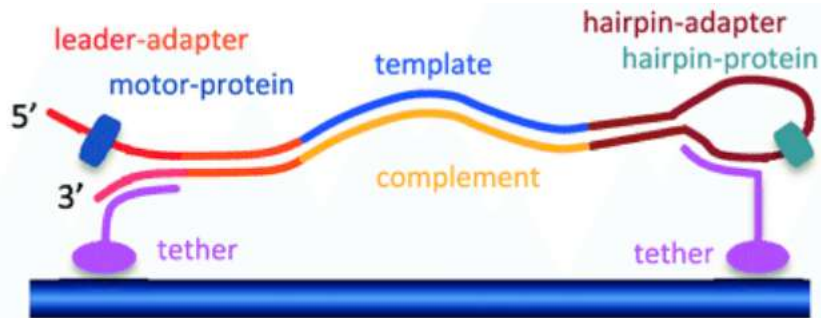
Instrument	Run time /FC	Output / FC	Nr of pores	Max read length
Flongle	16 hrs	1 Gb	126	1 Mb
MinION	24 hrs	2-15 Gb	512	1 Mb
GridION	24 hrs	2-15 Gb	512	1 Mb
PromethION	72 hrs	10 – 150 Gb	3 000	2 Mb

Q&A: "It depends"...





ONT: DNA + Motor + Pore



Base modification info is retained

Main advantages of ONT: SPEED and PORTABILITY

Rapid Confirmation of the Zaire Ebola Virus in the Outbreak of the Equateur Province in the Democratic Republic of Congo: Implications for Public Health Interventions

Placide Mbala-Kingebeni, Christian-Julian Villabona-Arenas, Nicole Vidal, Jacques Likofata, Justus Nsio-Mbeta, Sheila Makiala-Mandanda, Daniel Mukadi, Patrick Mukadi, Charles Kumakamba, Bathe Djokolo ... [Show more](#)

Clinical Infectious Diseases, Volume 68, Issue 2, 15 January 2019, Pages 330–333, <https://doi.org/10.1093/cid/ciy527>

Published: 29 June 2018 **Article history** ▼

ORIGINAL ARTICLE BRIEF REPORT

A Novel Coronavirus from Patients with Pneumonia in China, 2019

Na Zhu, Ph.D., Dingyu Zhang, M.D., Wenling Wang, Ph.D., Xinwang Li, M.D., Bo Yang, M.S., Jingdong Song, Ph.D., Xiang Zhao, Ph.D., Baoying Huang, Ph.D., Weifeng Shi, Ph.D., Roujian Lu, M.D., Peihua Niu, Ph.D., Faxian Zhan, Ph.D., et al., for the China Novel Coronavirus Investigating and Research Team

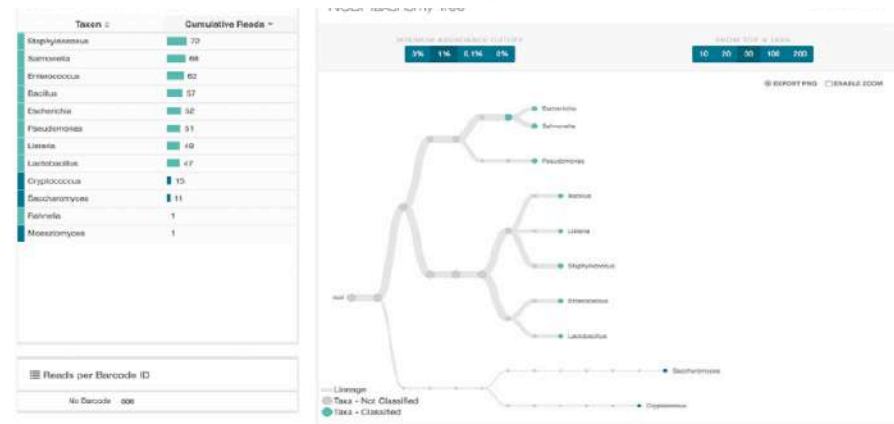


RESEARCH ARTICLE | [Full Access](#)

Semi-quantitative characterisation of mixed pollen samples using MinION sequencing and Reverse Metagenomics (RevMet)

Ned Peel, Lynn V. Dicks, Matthew D. Clark, Darren Heavens, Lawrence Percival-Alwyn, Chris Cooper, Richard G. Davies, Richard M. Leggett, Douglas W. Yu

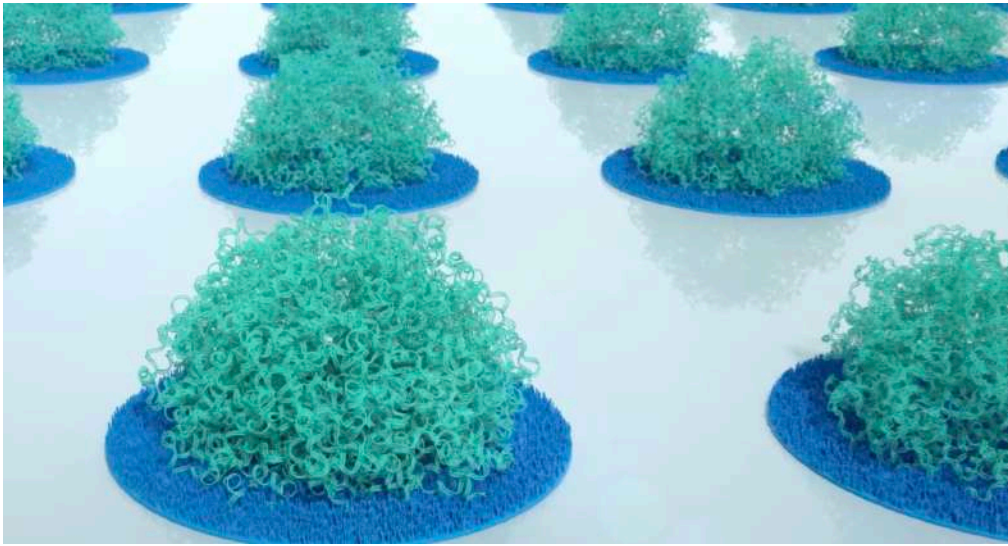
First published: 15 July 2019 | <https://doi.org/10.1111/2041-210X.13265>





MGI
华大智造

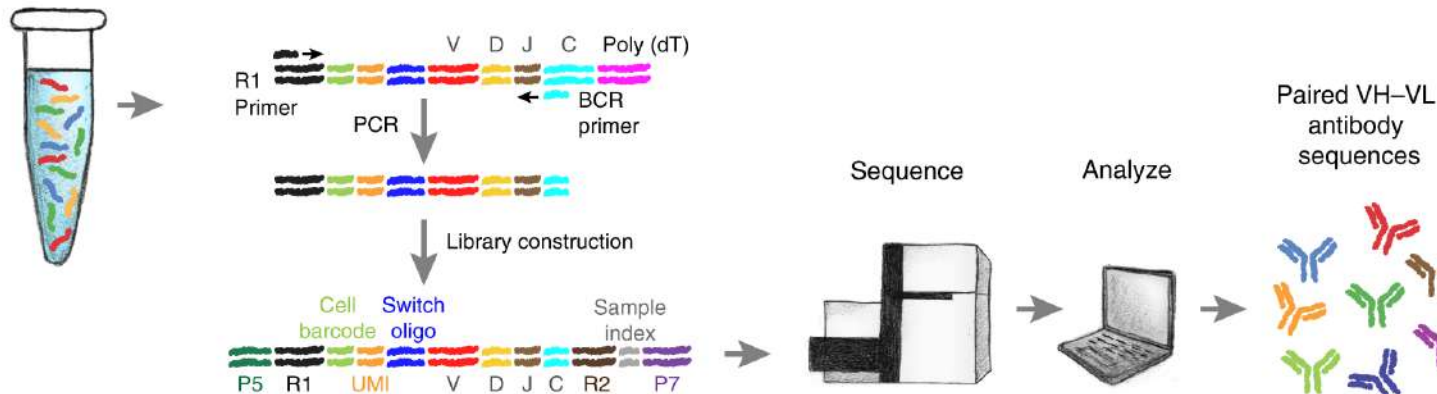
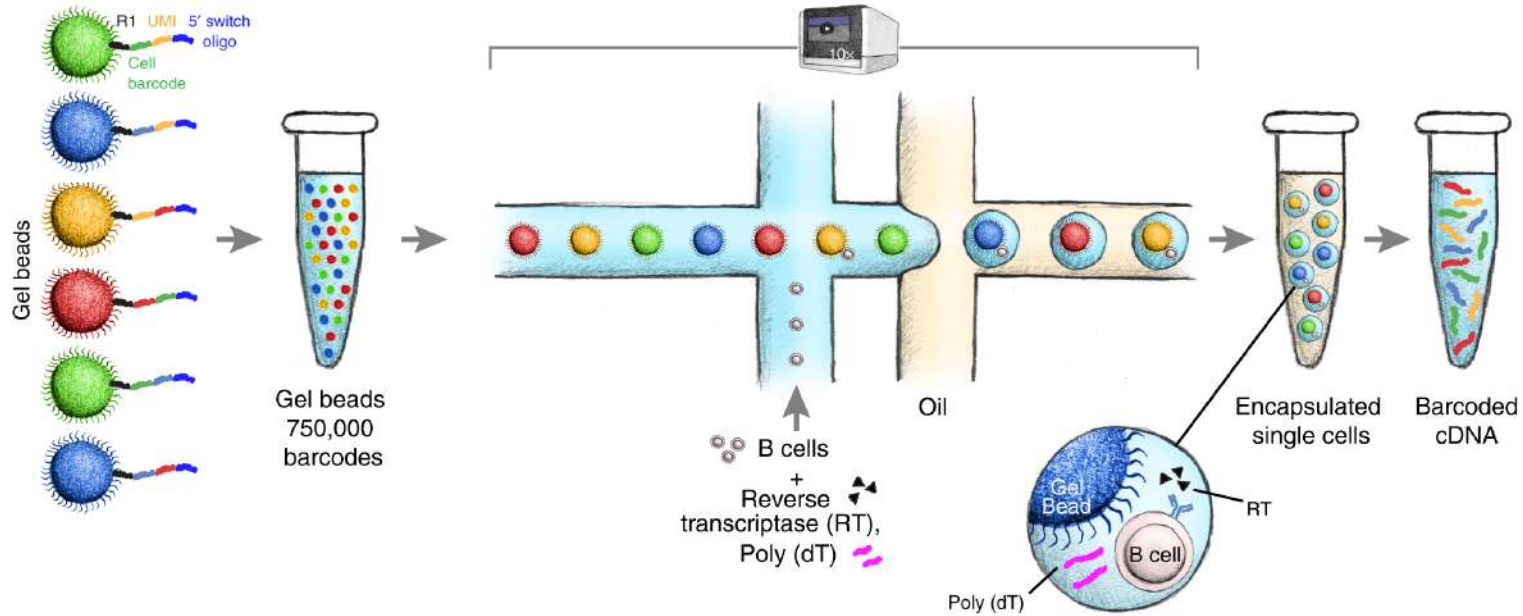
MGI - DNA NanoBall technology



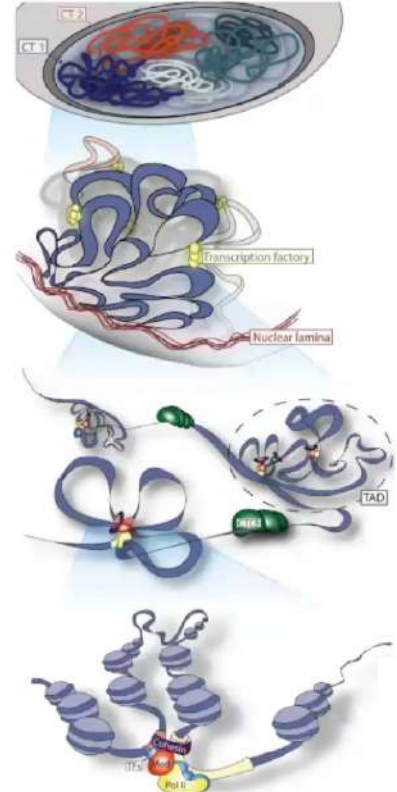
Instrument	Run time /FC	Output / FC	Nr of reads	Read length
DNBSEQ-T7	24 hrs	6Tb	5000M	PE150, PE100
DNBSEQ-G400	14-109 hrs	1440Gb	1500-1800M	SE400/PE200
DNBSEQ-G50	10-66 hrs	150Gb	500M / 100M	PE150
DNBSEQ-G400 FAST	13-37 hrs	330G	550M	PE150



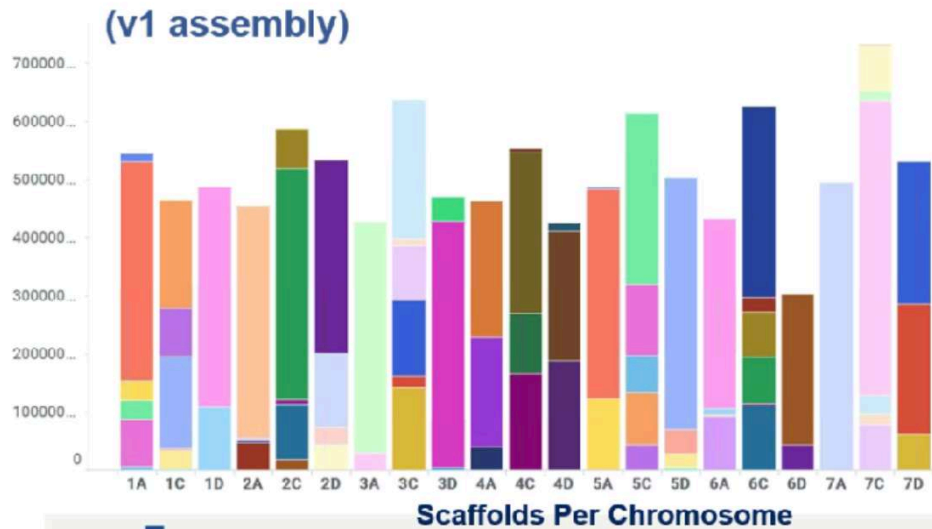
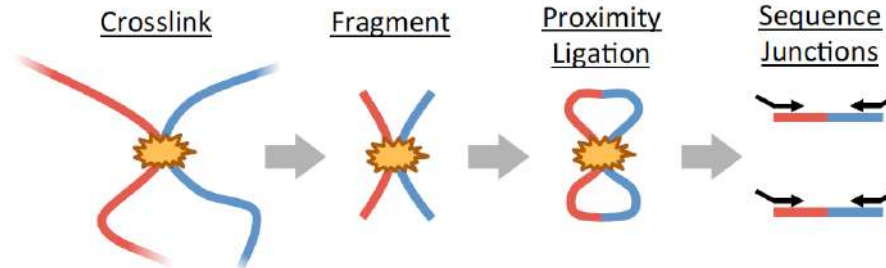
10x Genomics (Chromium)



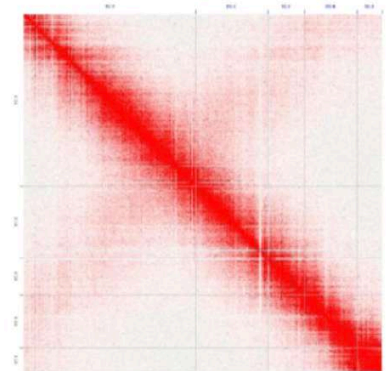
Hi-C / OmniC: linking reads to chromosomes



Sorkin, Leung and Ren, 2014



Chr 5C (612 Mb)



Start with a tissue!

Capture DNA bound to the same nucleosome

Make a library and sequence on Illumina NovaSeq

Technologies and Applications at NGI



NGS technologies

Short read NGS

Long-read NGS



- Whole genome re-sequencing
- RNA-seq
- Exome
- Targeted re-seq
- Panels
- Amplicons up to 600 bp

- De novo* genome sequencing
- Whole-transcript sequencing
- Structural variant resolving
- Allele phasing
- Targeted re-seq
- Amplicons up to 13 kb

Research and development

NGS Technologies: SUMMARY

- Development goes VERY FAST
- All technologies have their PROs and CONs
- One technology does not suit all the applications
- In some projects, several technologies should be combined

BREAK

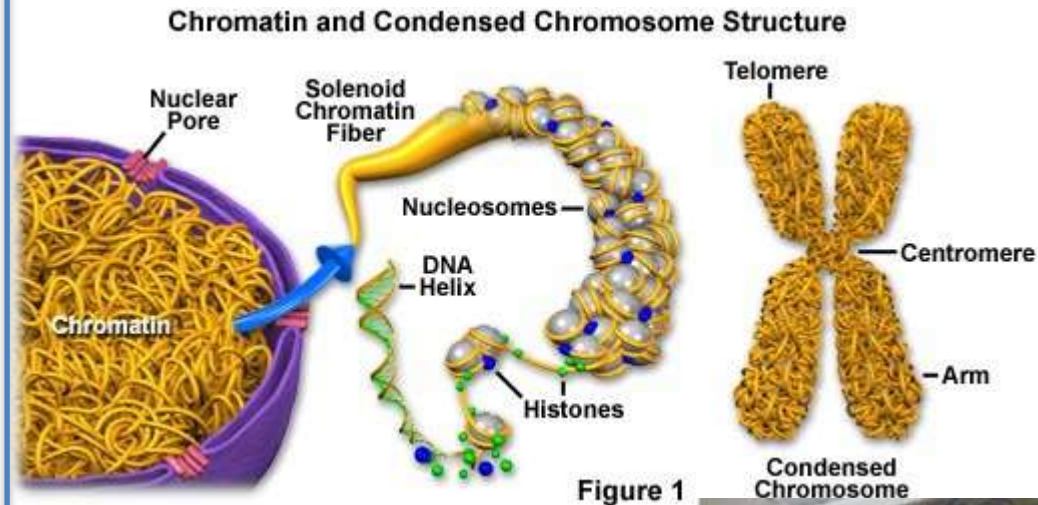
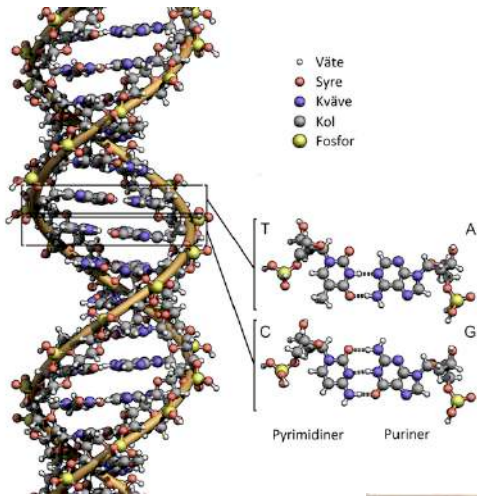
Making sense of genomics data:

Understanding sequencing bias

You do not see them before it is too late

Sequencing artefacts: what are they?

Sequencing a representative, completely randomized subsample:
it starts with input material



What textbook tells you



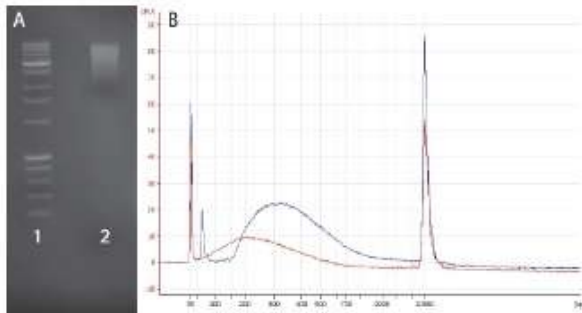
Brutal reality

*Do not forget:
DNA in solution behaves differently*

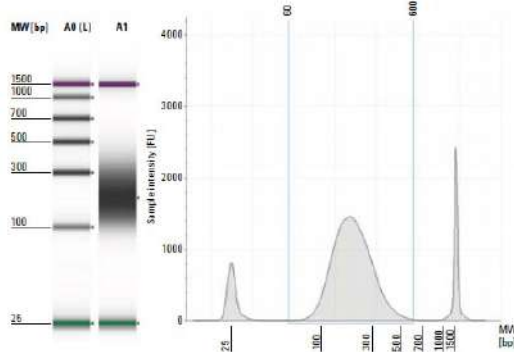


Sequencing artefacts: what are they?

Sequencing a representative, completely randomized subsample:
continues with library preparation

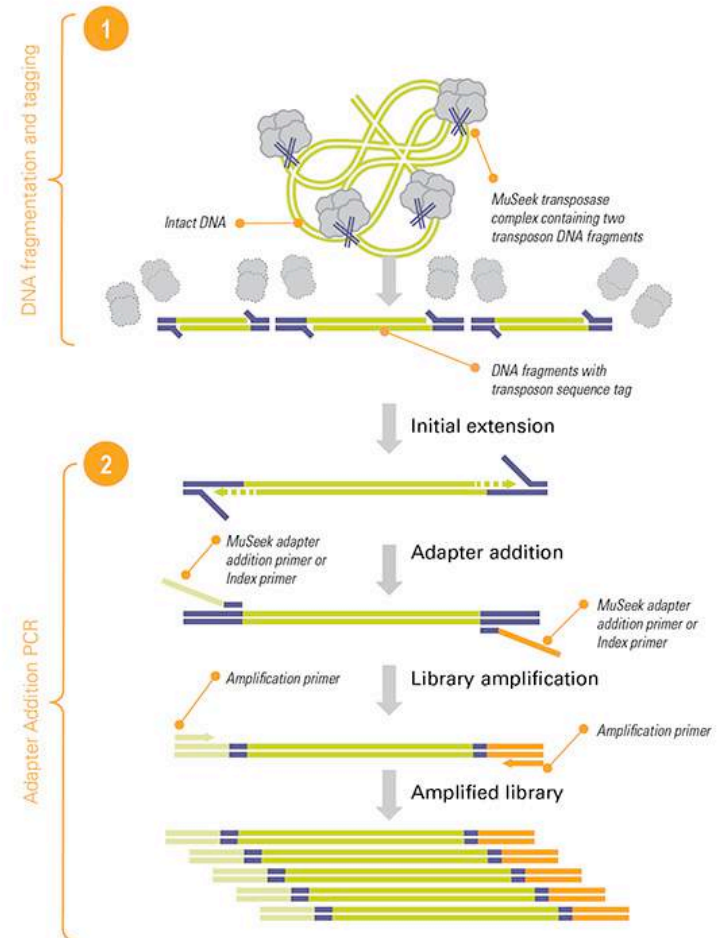


Input sample



Shearing and size-selection

Loosing molecules all the way



Less material -> more amplification cycles

Sequencing artefacts: what are they?

PCR bias – important source of sequencing artefacts

PCR steps involved in any NGS but PacBio and Oxford Nanopore:

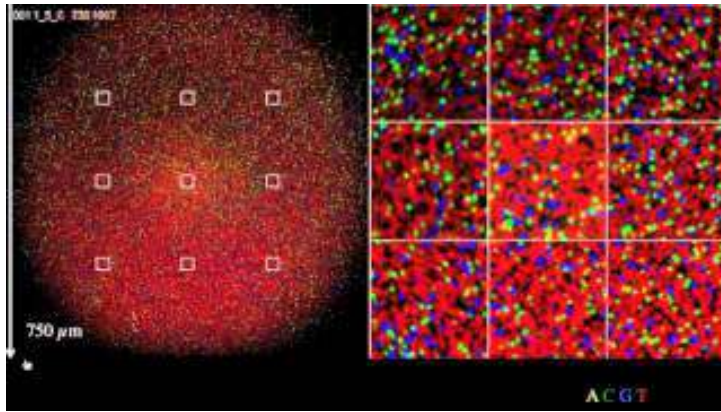
1. Library amplification
2. Amplification during templating (Illumina – on glass; Ion – emPCR)

Main PCR bias:

1. Size: shorter fragments amplify faster -> higher sequencing signal and coverage
2. Polymerase errors
 - slippage in low complexity regions
 - incorporation of erroneous bases & indels
3. GC-bias (fragments with high GC diminish to 1/10th from initial amount)

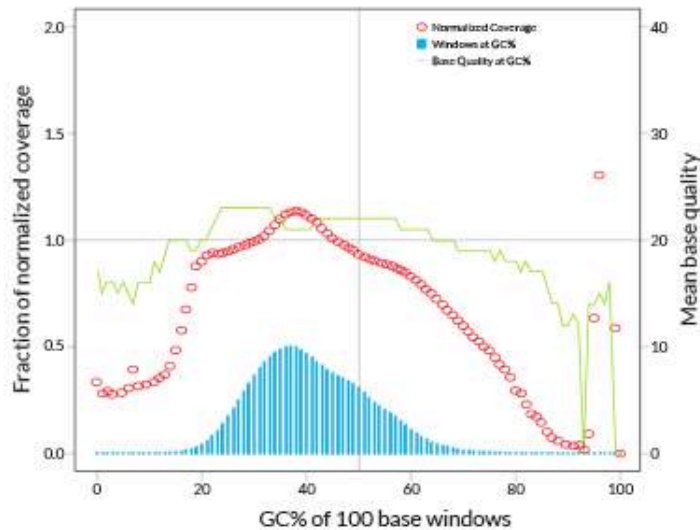
Sequencing artefacts: what are they?

PCR bias – important source of sequencing artefacts

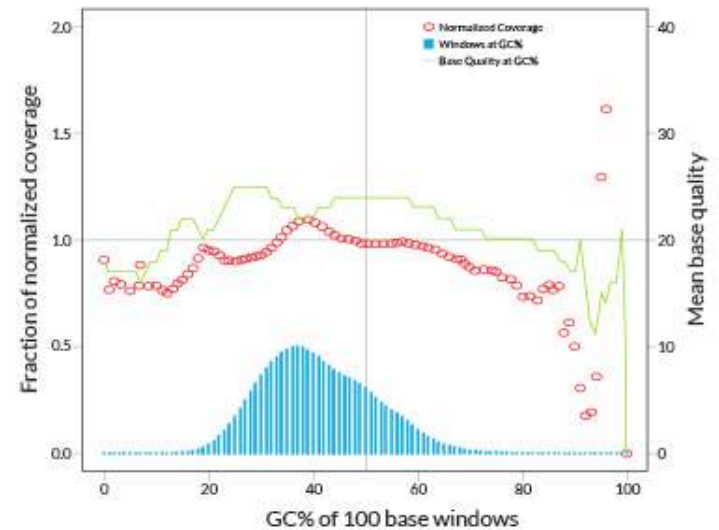


Clusters with shorter fragments grow faster -> quality signal from smaller clusters worsens

GC bias & genome coverage



Heavily amplified library



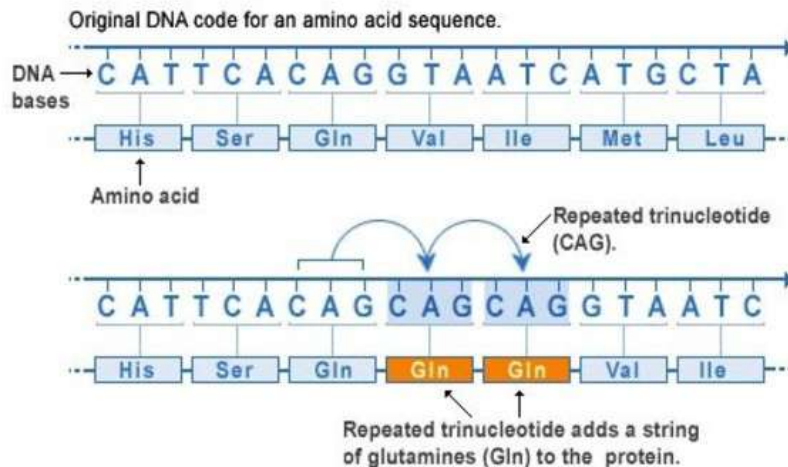
PCR-free library

Sequencing artefacts: what are they?

PCR bias – important source of sequencing artefacts

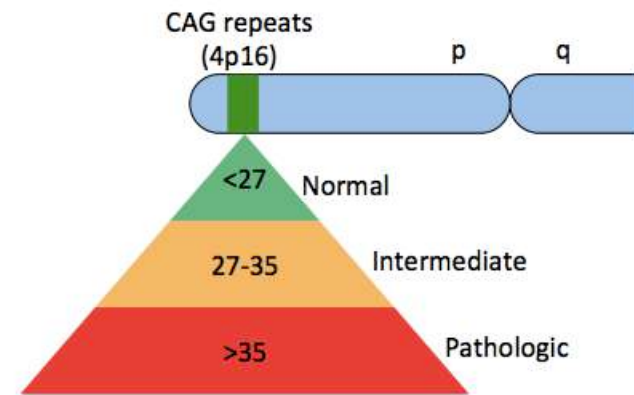
Polymerase slippage – low complexity regions

Repeat expansion mutation



U.S. National Library of Medicine

Huntington's Disease



Huntington's disease:

- Inherited disorder resulting in brain cell death
- Decline of motoric and cognitive functions
- Common onset: 30-50 years of age
- No cure
- Causative genetic variant: CAG-repeat expansion in *HTT* gene

Batch Effects

Batch effects and the effective design of single-cell gene expression studies

Po-Yuan Tung, John D. Blischak, Chiaowen Joyce Hsiao, David A. Knowles, Jonathan E. Burnett, Jonathan K. Pritchard & Yoav Gilad [✉](#)

Scientific Reports 7, Article number: 39921 (2017) | [Cite this article](#)

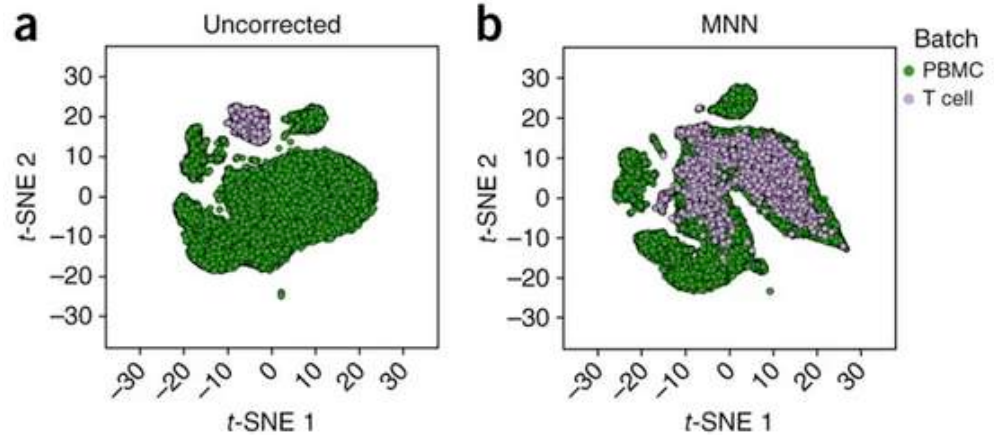


Single-cell & RNA sequencing

Batch effects in single-cell RNA-sequencing data are corrected by matching mutual nearest neighbors

Laleh Haghverdi, Aaron T L Lun, Michael D Morgan & John C Marioni [✉](#)

Nature Biotechnology 36, 421–427(2018) | [Cite this article](#)



Sequencing bias: SUMMARY

- Keep in mind that they are there
- Coverage varies across the genome
- One technology does not suit all the applications
- Beware of batch effects

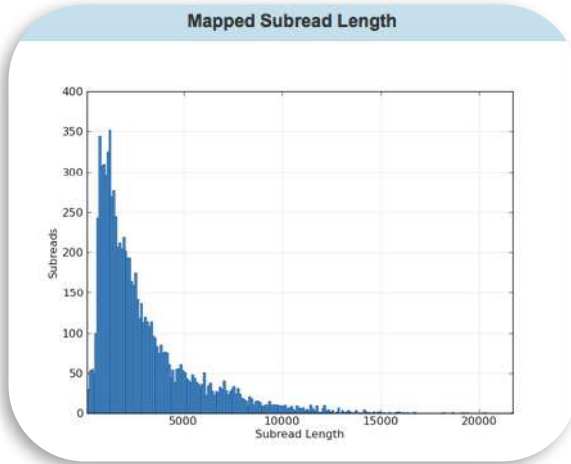
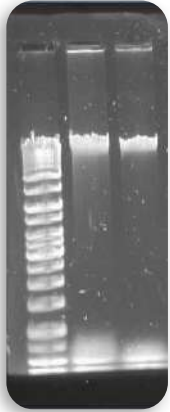
SAMPLE QUALITY REQUIREMENTS

Garbage in – garbage out:

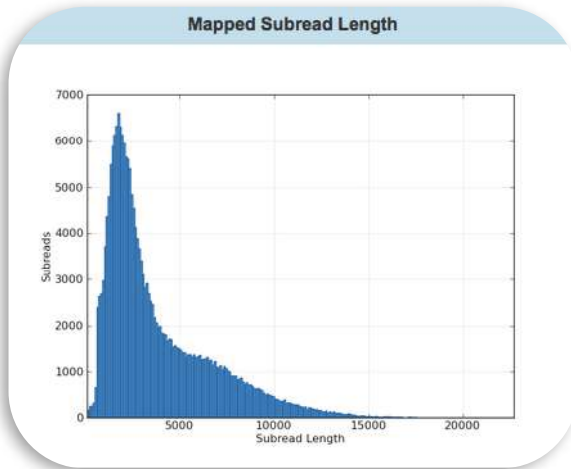
Sequencing success always depends
on the **sample quality**.

**NGS-quality DNA and
PCR-quality DNA
are two completely different things.**

2013: a wake-up call



Polished Contigs	223	Max Contig Length	36,298
N50 Contig Length	2,932	Sum of Contig Lengths	480,087



Polished Contigs	9	Max Contig Length	1,508,929
N50 Contig Length	1,353,702	Sum of Contig Lengths	7,813,244



For Long Reads one needs to have *long and pure DNA*

DNA quality and inhibition of sequencing

Short-read technologies: PCR inhibition

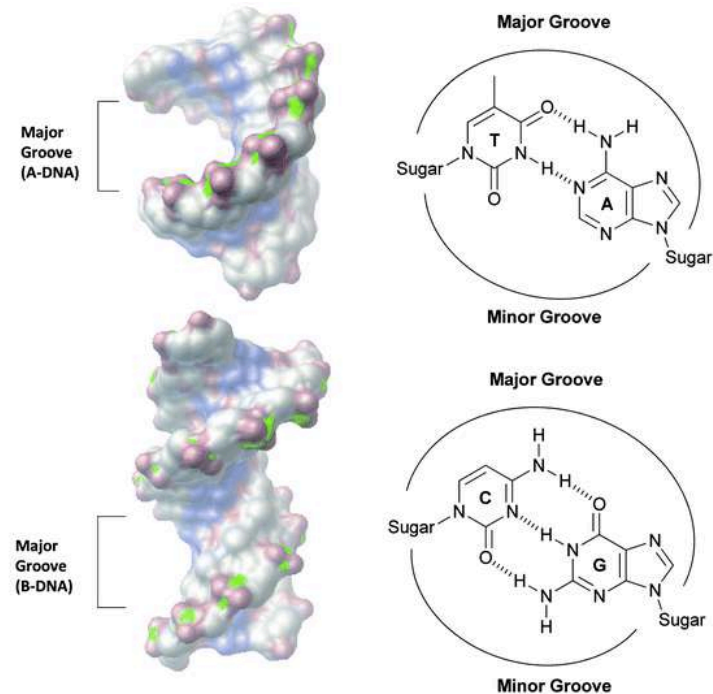
Long-read technologies are PCR-free, but one sequences native DNA “as is”.

DNA-binders:

- Proteins
- Polyphenols
- Secondary metabolites (e.g. toxins)
- Pigments
- Polysaccharides

Polymerase inhibitors:

- Salts
- Phenol
- Alcohols



[Hamilton & Arya, Nat. Prod. Rep., 2012, 29, 134-143](#)

Physical inhibiting factors – debris

What do absorption ratios tell us?

Pure DNA 260/280: 1.8 – 2.0

< 1.8:

Too little DNA compared to other components of the solution; presence of organic contaminants: proteins and phenol; glycogen - **absorb at 280 nm**.

> 2.0:

High share of RNA.

Pure DNA 260/230: 2.0 – 2.2

<2.0:

Salt contamination, humic acids, peptides, aromatic compounds, polyphenols, urea, guanidine, thiocyanates (latter three are common kit components) – **absorb at 230 nm**.

>2.2:

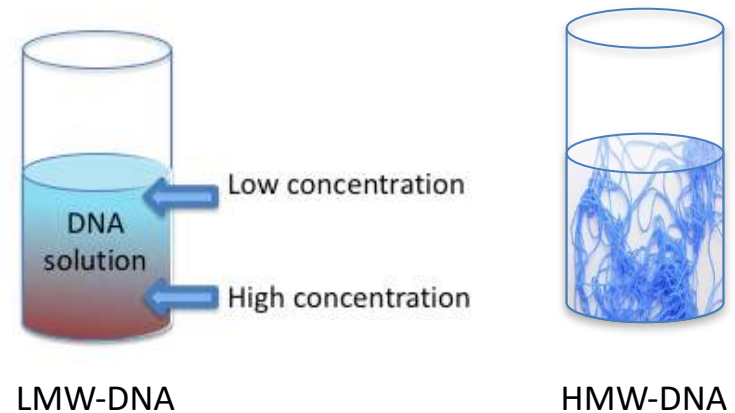
High share of RNA, very high share of phenol, **high turbidity**, dirty instrument, wrong blank.

*Photometrically active contaminants:
phenol, polyphenols, EDTA, thiocyanate, protein,
RNA, nucleotides (fragments below 5 bp)*

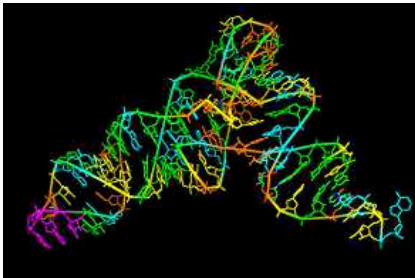
How to make a correct measurement

- Thaw DNA completely
- Mix gently (**never vortex!**)
- Put the sample on a thermoblock: 37°C, 15-30 min
- Mix gently
- **Dilute 1:100** (if HMW)
- Mix gently
- Make a measurement with an appropriate blank

- **NANODROP is Bad.** Point.
- Use Qubit, or PicoGreen.



What about RNA?



Transcriptome sequencing (RNA-seq)



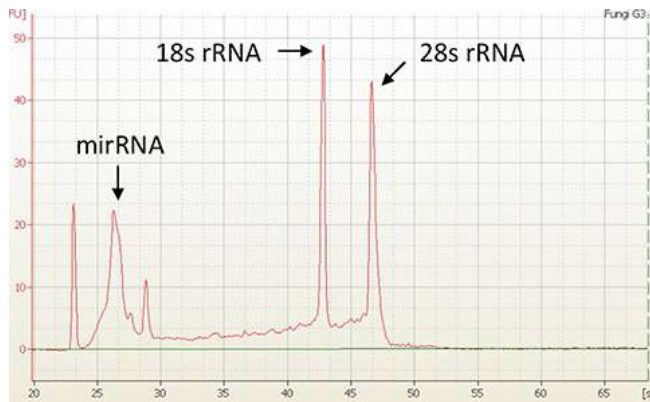
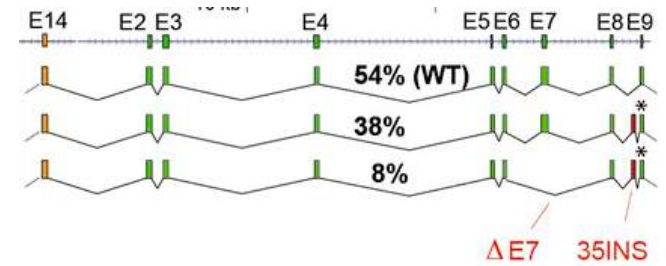
mRNA

- **Dif.ex.**
- Annotation

Non-codingRNA **miRNA**

- Transcriptional regulation

Splice isoforms



Sample prep: RNA

mRNA degrades FAST

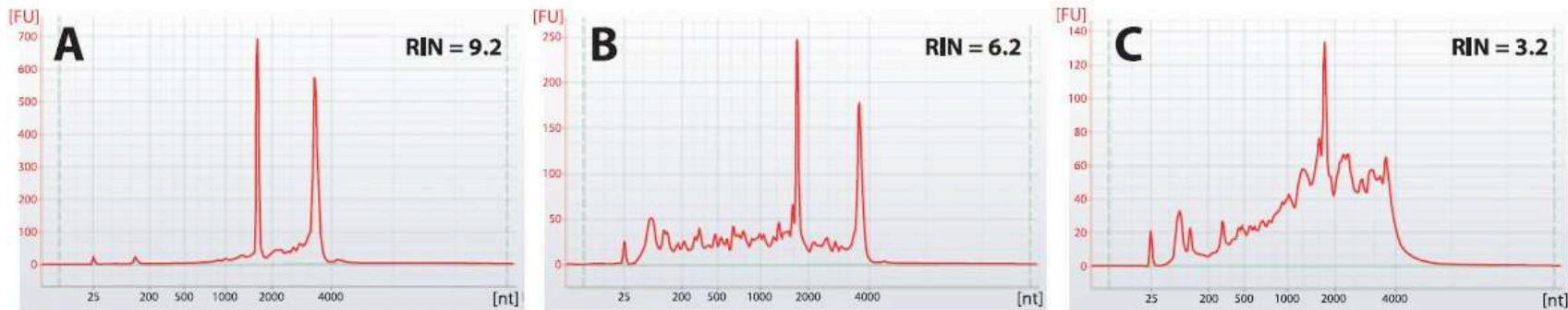
Freeze sample or place it in RNA-later within 30 sec (*if possible*)

Chose a correct kit for your particular application!

Always treat samples with DNase

Differential expression, miRNA – **RIN value over 8.0**

Aim for 4 biological replicates

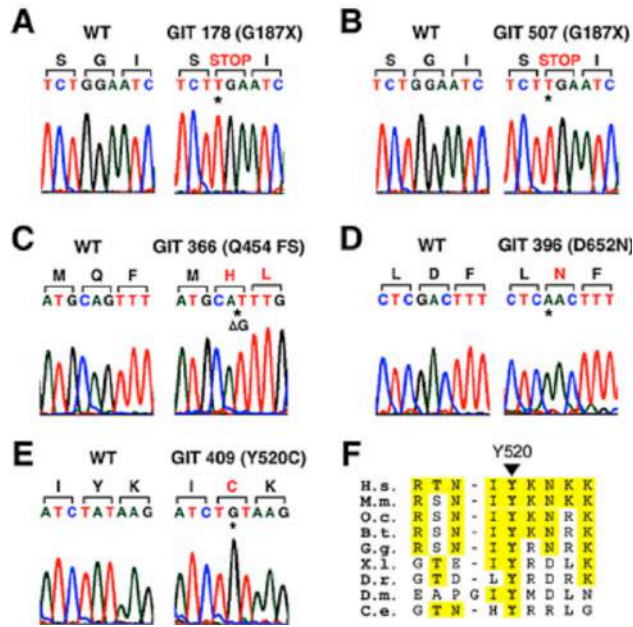


Sample prep: SUMMARY

- Sequencing success depends on the sample quality
- DNA quality is **essential** for PacBio and ONT sequencing
... as well as PCR-free Illumina libraries & linked reads!
- Basic understanding of biochemistry is needed
- **NGS-grade sample \neq PCR-grade sample**
- **Be cautious with data interpretation**

1 minute of phylosophy

Genome is not a linear string of bases!!



Mutations in coding regions only



Transcriptional & post-transcriptional regulation



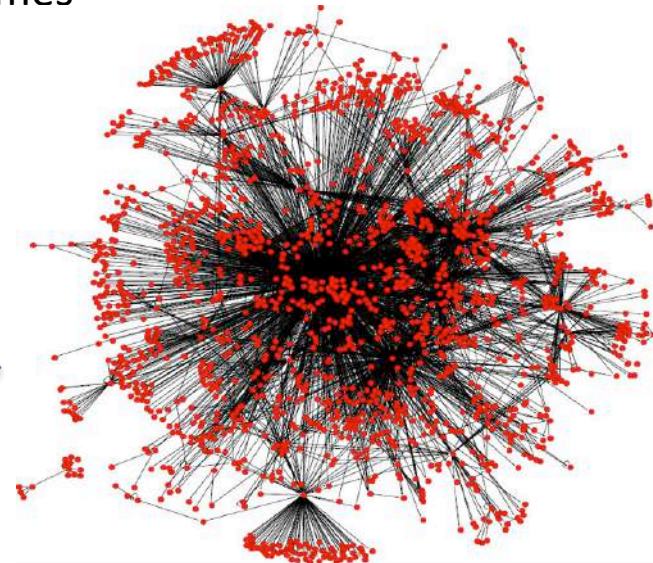
Epigenetics



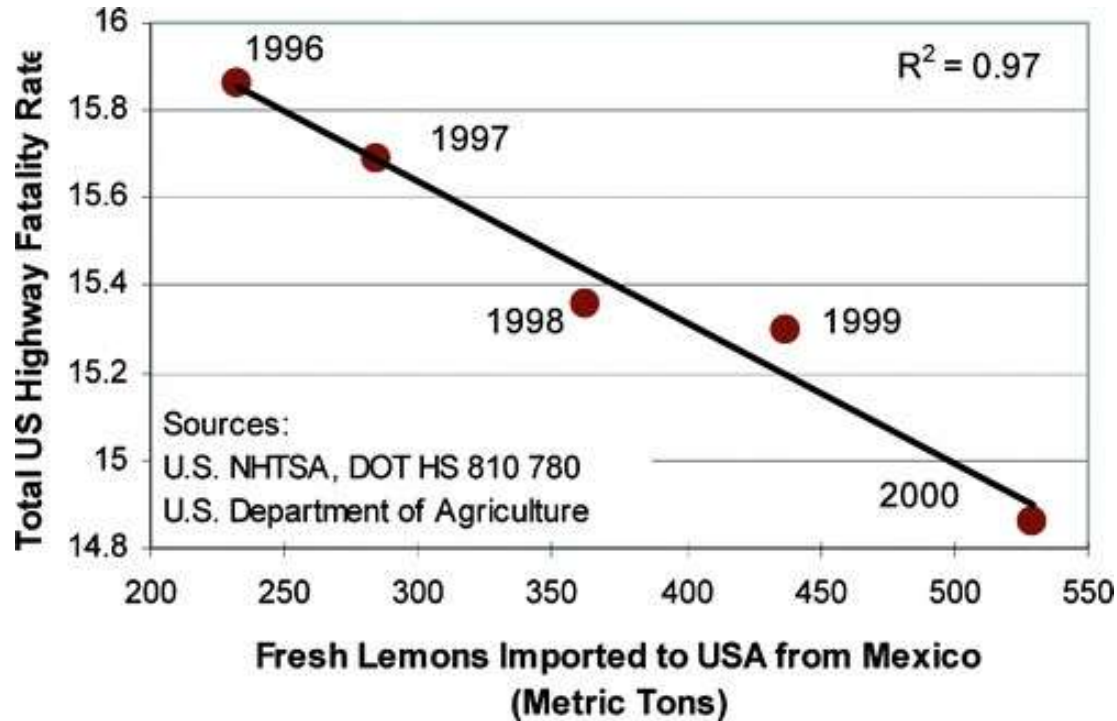
Proximity in chromosomes

>gi|30018278|ref|NC_004722.1| *Bacillus cereus* ATCC 14579, complete genome

```
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ACAATTTGTGTATAACATGTGGACAGTTTTAATCACATGTGGGTAAGTAGTTGTCCACATTTGCTTTTTT
TGTCGAAAACCCCTTCTCAATACAAACGACGTTTTAGGTTTTAAAATCGTTTTTCGTATAAAATATACATT
TAATTTATTAGGTTGTACATTTGTTGCACAACCTTTATTCTTTTACCAACTTAGTAAAGGAGGGACACCT
TTGGAAAATATCTCTGATTTATGGAATAGTGCCTTAAAAGAATTAGAAAAAAGGTAAGCAAGCCTAGTT
ATGAGACATGGTTAAAATCCACAACGGCTCATAACTTGAAGAAAAGACGTATTAACGATTACAGCTCCGAA
TGAATTTGCTCGTGACTGGCTAGAACTCTCAATTACTCCGAACATAATTCAGAAAACACTATACGATTTAACA
GGGGCAAAATTAGCAATTGCTTTTATTATCCCCAAGTCAAGCTGAAGAGGACATTGATCTGCCTCCAG
TTAAGCCGAATCCAGCACAAGATGATTGAGCTCATTACCACAGAGCATGTTAAATCCAAAATATACATT
CGATACATTTGTTATTGGCTCTGGTAACCGTTTTGCCCATGCAGCTTCATTAGCTGTAGCTGAGGCGCCA
GCTAAAGCGTATAATCCACTCTTTATTACGGGGGAGTTGGACTTGGAAAAGACACATTTAATGCAGGCAA
TTGGTCATTATGTAATTGAACATAATCCAAATGCAAAAGTTGTATATTTATCATCAGAAAAATTTACAAA
TGAATTTATTAACCTCTATTGCTGATAATAAAGCTGTTGATTTTCGTAATAAATATCGTAACGTAGATGTT
TTATTGATAGATGATATTCAATTTCTTGCTGGAAAAGAACAGACTCAAGAAGAGTTTTTCCATACATTTA
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```

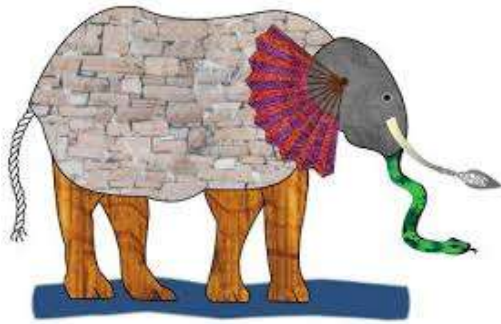


Never forget: Correlation vs Causation



Reduction in export of fresh lemons from Mexico causes significant reduction of highway traffic fatality rates in the US!

Blind men & an elephant



Letter

Genome-wide association study identifies five new schizophrenia loci

The Schizophrenia Psychiatric Genom Article

Nature Genetics **43**, 969–976 (2011)

doi:10.1038/ng.940

[Download Citation](#)

Genome-wide association analysis identifies 30 new susceptibility loci for schizophrenia

Zhiqiang Li, Jianhua Chen [...] Yongyong Shi

Nature Genetics **49**, 1576–1583 (2017)

doi:10.1038/ng.3973

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Schizophrenia and the dynamic genome

Patrick F. Sullivan

177 9:22

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Current opinion in psychiatry

Author Manuscript

HHS Public Access

Genome-wide association studies (GWAS) of schizophrenia: does bigger lead to better results?

Sarah E. Bergen, PhD and Tracey L. Petryshen, PhD

Summary

Copy number variation (CNV) is a widely replicated risk factor for psychiatric disorders, including schizophrenia, although the mechanisms by which CNVs confer risk are unclear. Recent studies have provided robust evidence of CNVs associated with schizophrenia, and have highlighted a potential role for schizophrenia risk-associated

NGS and its challenges: SUMMARY

- Technologies develop VERY FAST.
- Beware of sequencing bias.
- Sequencing result depends on sample quality.
- Consult experts when it comes to experimental design and technology choice.
- ***Do not forget the elephant.....***

THANK YOU!