

DNA Methylation Methods and Technologies

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Outline



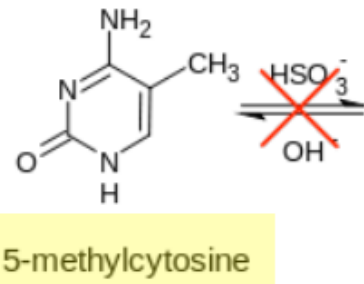
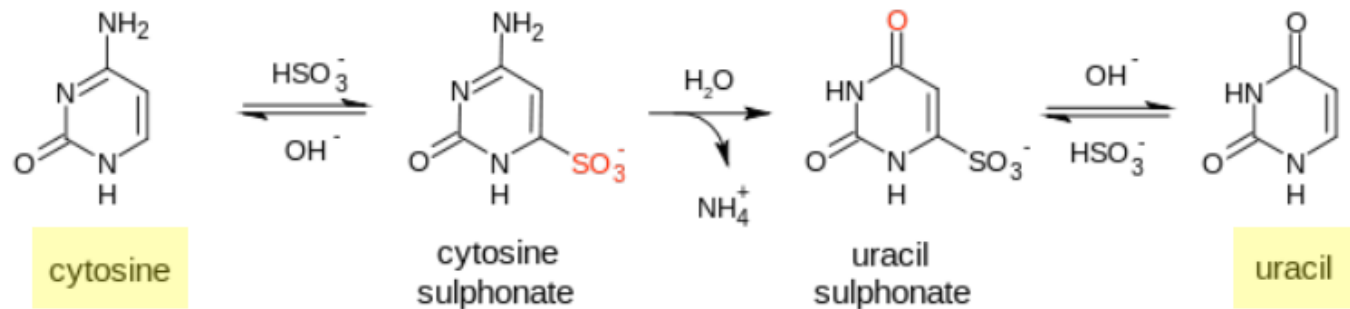
- i. Overview of methods for interrogation of DNA methylation
 - Overview of important concepts
 - Enrichment & targeted-based methods
 - Genome-wide methods
- ii. How to access epigenomics services for your research project at Sweden's National Genomics Infrastructure (NGI)

Short intro: Conversion



Bisulfite conversion has been the "Gold standard" for DNA methylation analysis.
Provides "single nucleotide resolution".

The Chemistry of Bisulfite Conversion of Cytosine to Uracil:

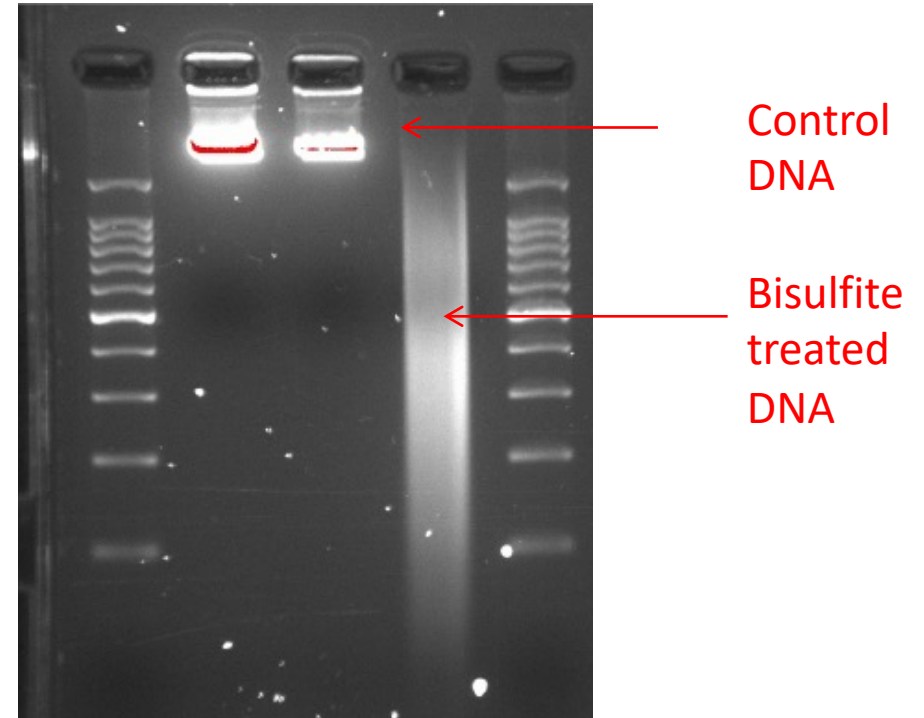


5-methyl-cytosine is resistant
to chemical conversion!!!!

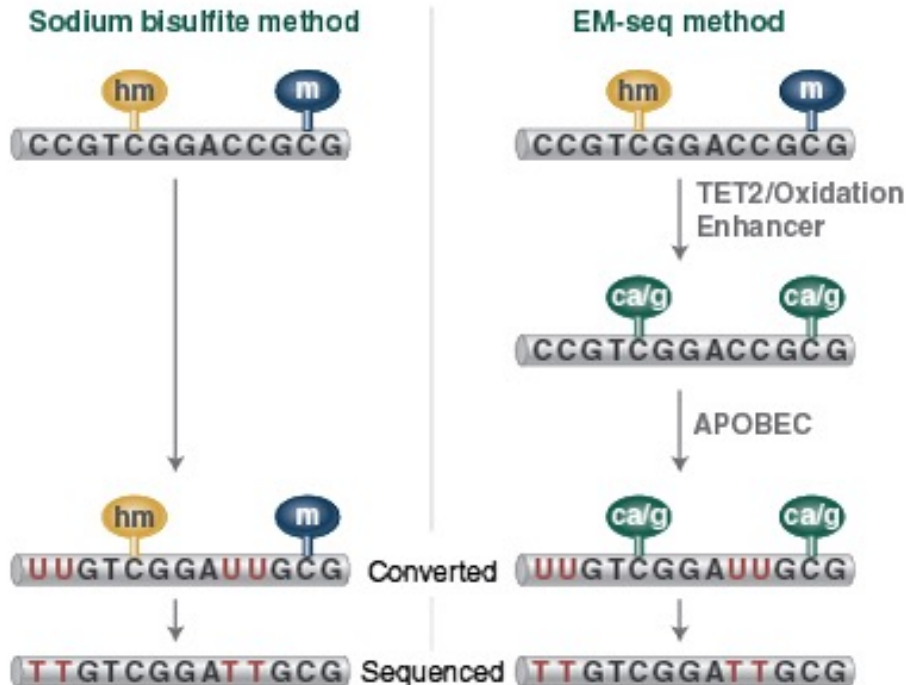
What you need to know about bisulfite conversion



- Very harsh chemical that degrades and fragments DNA



New innovation- Enzymatic conversion!



Gentle with little/no strand breakage!

WGBS is the gold standard for methylome analysis, but the chemical bisulfite reaction:

- I. Damages / degrades DNA
- II. Results in fragmentation / loss
- III. Can result in CG bias and uneven genome coverage

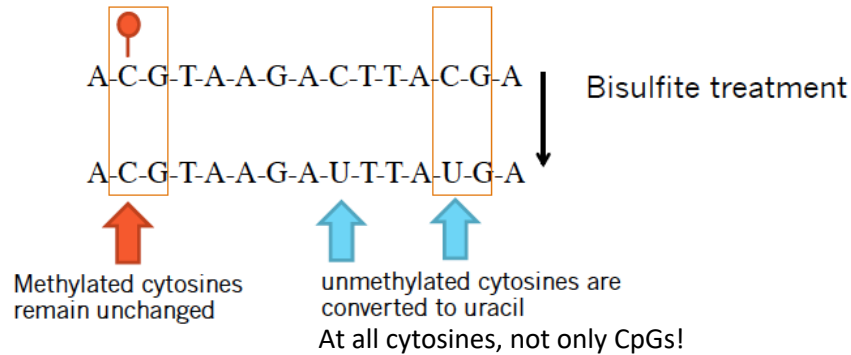
Enzymatic methylation sequencing (EM-seq)

TET2 enzymatically oxidizes 5mC and 5hmC through a cascade reaction into 5-carboxycytosine (5caC)

*5-methylcytosine (5mC) → 5-hydroxymethylcytosine (5hmC)
→ 5-formylcytosine (5fC) → 5-carboxycytosine (5caC)*

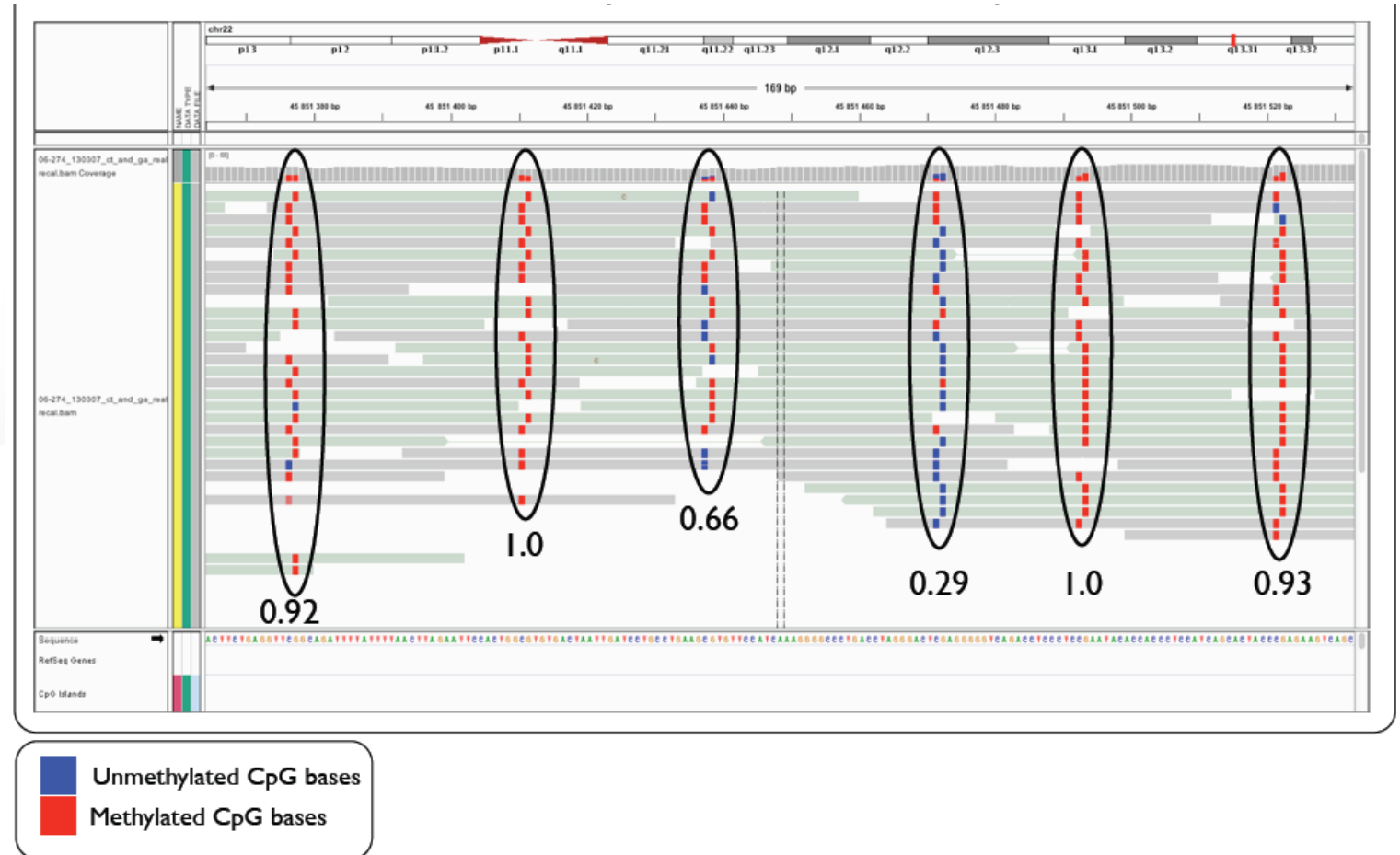
A second enzymatic step uses APOBEC to deaminate cytosine to uracil, but does not affect 5caC.

Base-pair resolution and quantitative measurement of methylation levels

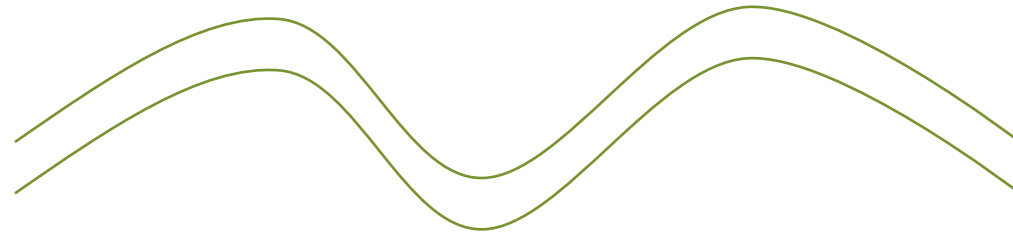


C = methylated

U->T = unmethylated



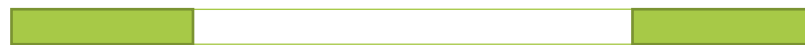
Short intro: "NGS" libraries



Double stranded genomic DNA

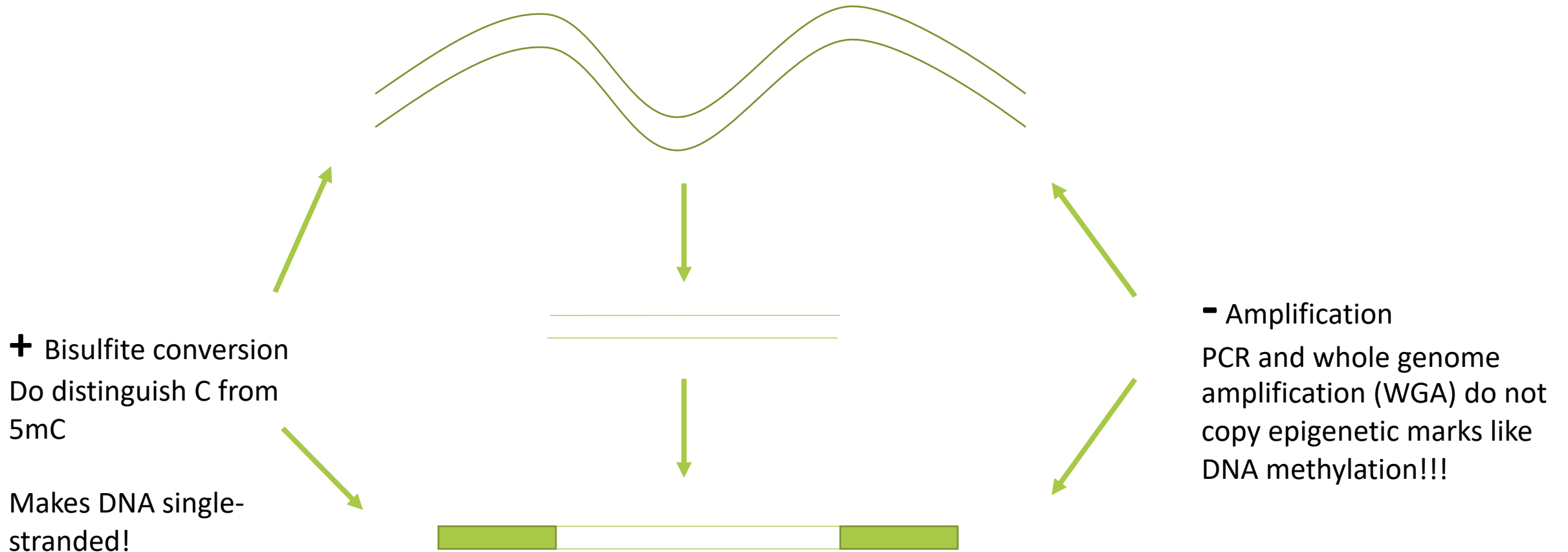


Shearing to make DNA fragments shorter (with bisulfite treatment optional)



Ligate platform-specific sequencing adapters

Short intro: "NGS" libraries



Enrichment & targeted-based methods



Different approaches to reduce the genome to regions of interest (typically those with many CpG sites)

- Cost saving (less sequencing required)
- Less computationally intensive (less data generated)
- High throughput (some approaches)

Enrichment-based methods



Capture of methylated DNA fragments using methyl-binding protein or a anti-methyl-cytosine antibody

- MeDIP-seq (Methylated DNA immunoprecipitation):
 - ✓ Genome-wide coverage
 - ✓ ~150bp resolution.
 - ✓ Anti-body against 5-Hydroxy-methyl-cytosine
 - ✓ Relatively cost-efficient
- MBD-seq (Methylated DNA binding domain):
 - ✓ Genome-wide coverage
 - ✓ ~150bp resolution.
 - ✓ Only capture CpG methylation not CHH
 - ✓ Relatively cost-efficient

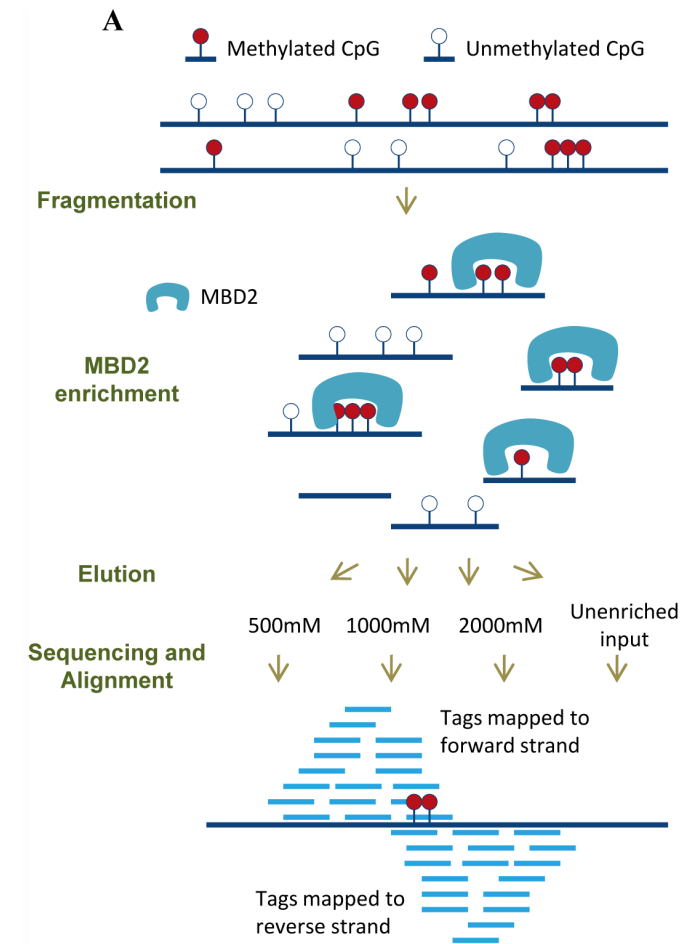
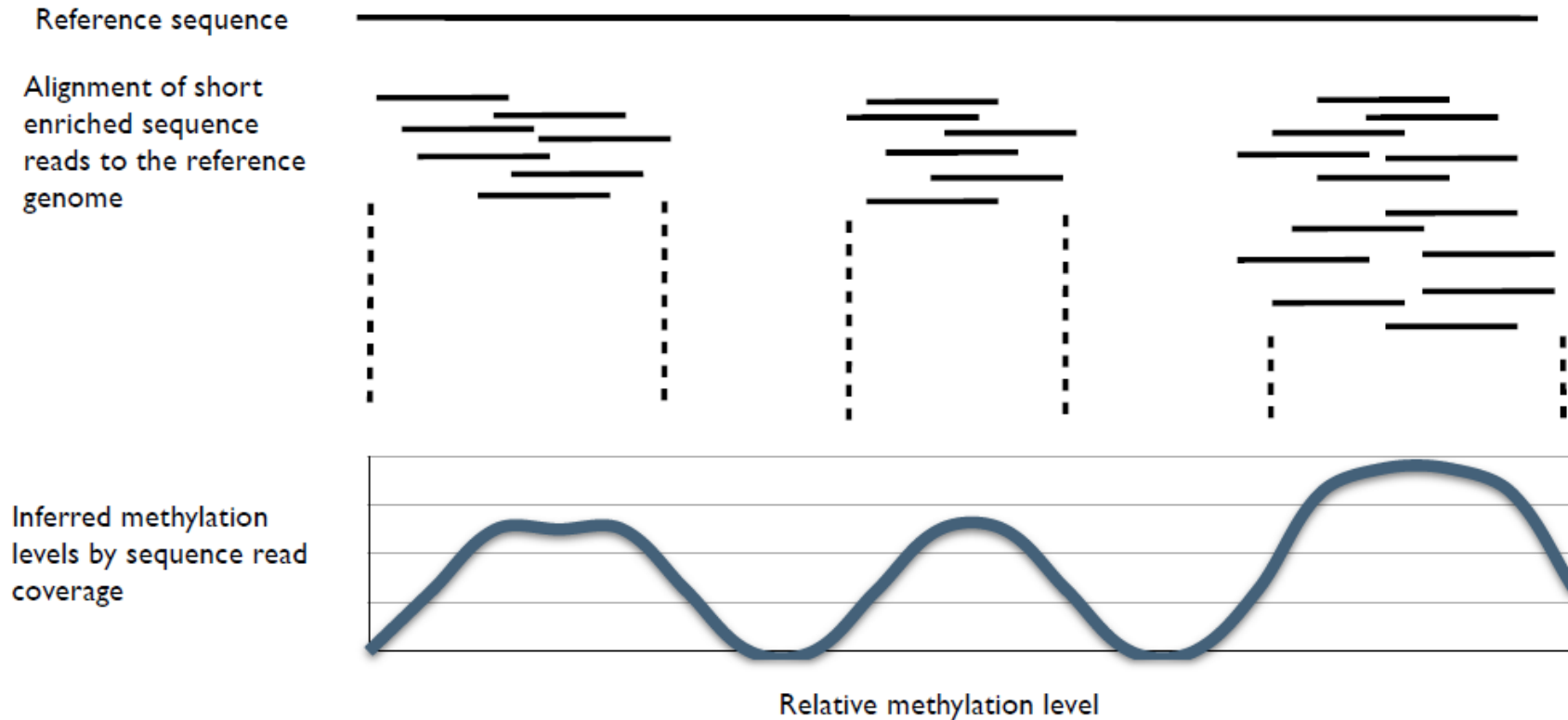


Figure from:
Lan, et al. (2011) High Resolution Detection and Analysis of CpG
Dinucleotides Methylation Using MBD-Seq Technology.
<https://doi.org/10.1371/journal.pone.0022226>

Enrichment-based methods



The depth of sequence reads is taken as an indirect measurement of Methylation levels

Pros:

- Works for different species

Cons:

- Not base-pair resolution
- Indirect measurement of DNA methylation can be more difficult to interpret
- Lab-intensive and not easily automated

Target-Capture



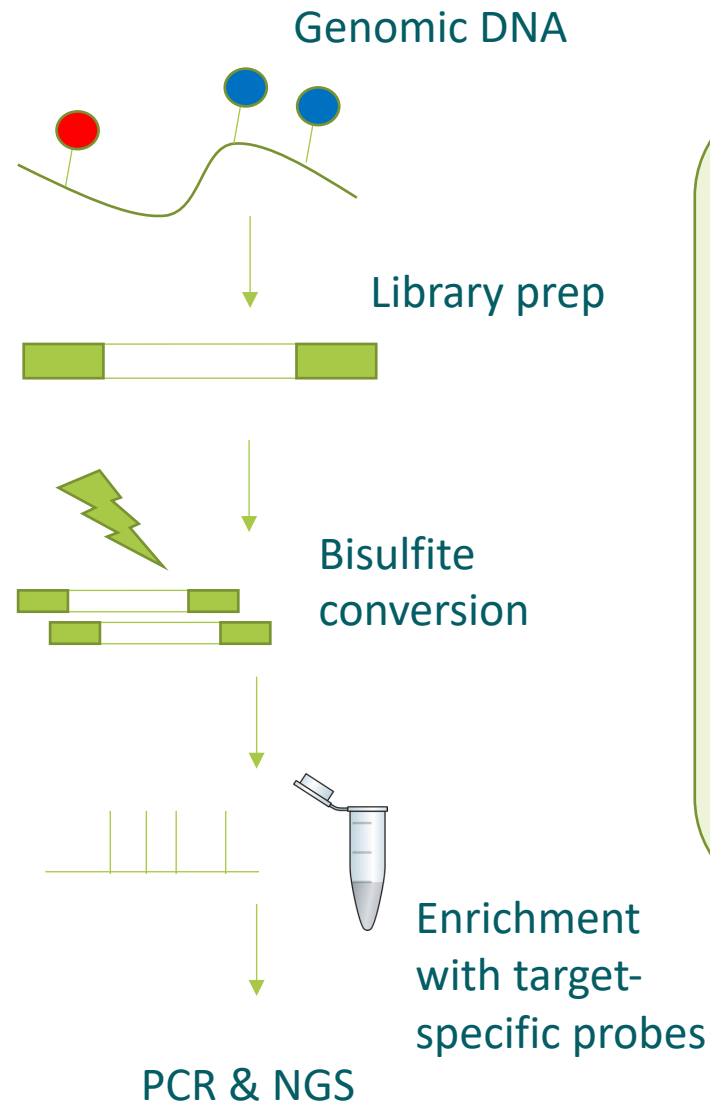
Target-capture of pre-defined genomic regions, NGS library preparation, uses bisulfite conversion.

Pros:

- Focused set of targets regions: can achieve high coverage on target
- “Cost-effective”
- Captures millions of CpG sites (3-5M)

Cons:

- Typically only for Human, other species possible on some platforms
- Standard conversion cannot distinguish between 5mC and 5hmC



Seq-Cap enrichment
(Roche) –
5M CpGs

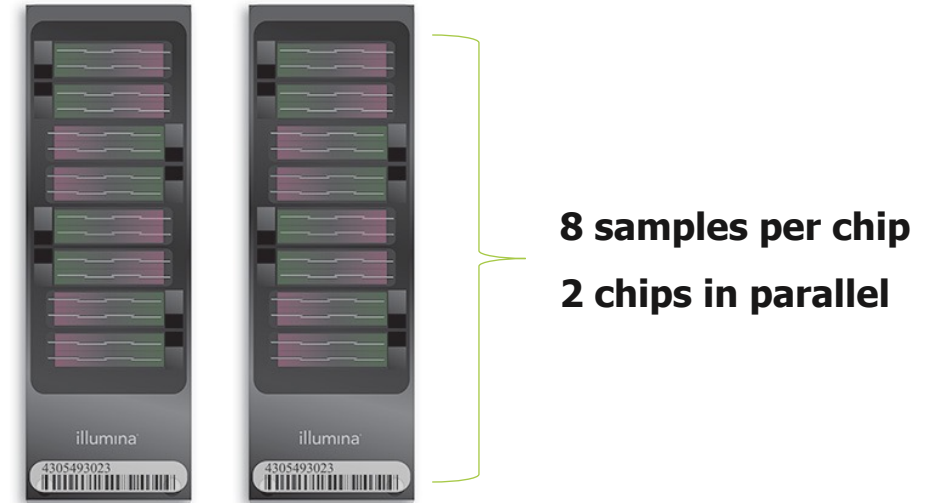
SureSelect^{XT} Methyl-Seq
Target Enrichment Kit
(Agilent Technologies) –
3.7M CpGs

**Twist Custom Methylation
Panel – 3.2M CpGs**

DNA methylation arrays



- Bisulfite converted DNA
- >800,000 CpG sites
- 96% CpG islands
- 99% Refseq genes
- CpG sites outside of CpG islands
- Non-CpG methylated sites identified in human stem cells
- Differentially methylated sites found in cancer and several tissue types
- FANTOM 4 promoters
- DNase hypersensitive sites
- miRNA promoters



$$\text{Beta value } (\beta) = \frac{M}{M + U + 100}$$

DNA methylation arrays

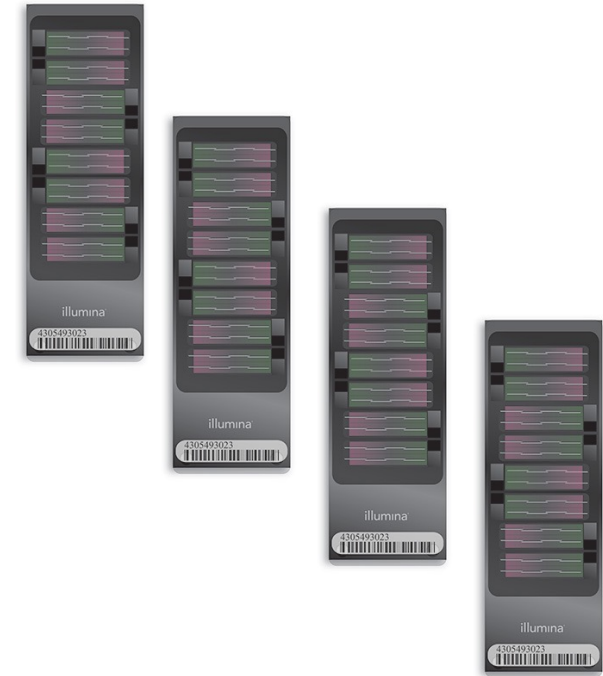


Pros:

- The most popular method on the market
- Base-pair resolution
- Compatible with FFPE DNA
- Compatible with 5hmC detection
- Many **R packages** available for data analysis and publically available datasets

Cons

- Human only* (Mouse Methylation BeadChip available with 285k CpG sites or flexible iSelect Methyl Custom BeadChip, but \$\$)
- 850k out of 29M CpG sites



Reduced Representation Bisulfite Sequencing (RRBS)



- “reduces” the genome to informative regions with high CG content
- Based on restriction digestion with an enzyme that cuts at CCGG (MspI)

Pros:

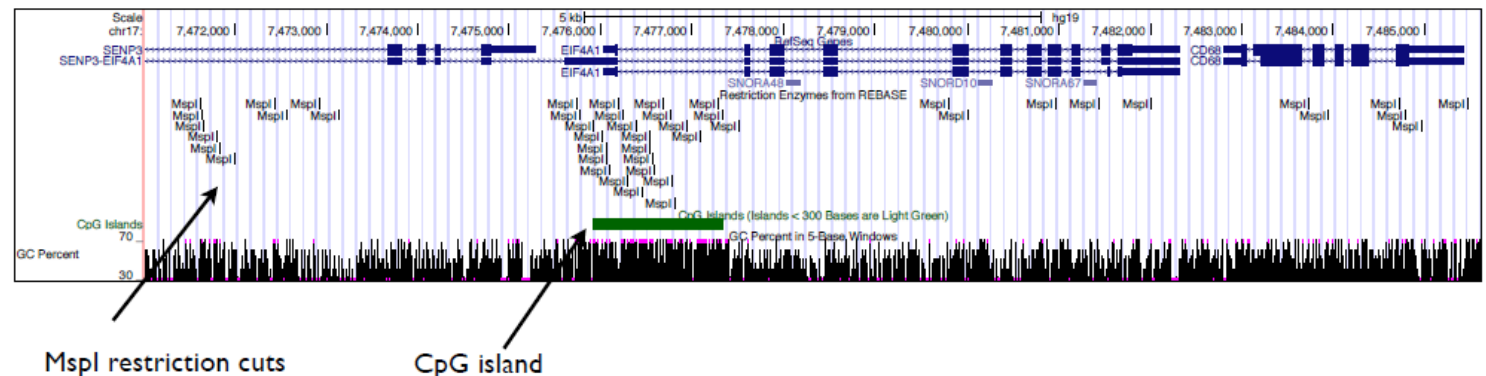
- Compatible with most species
- Low cost
- Base-pair resolution (bisulfite)
- reads are heavily concentrated to CpG islands
- High throughput

Cons

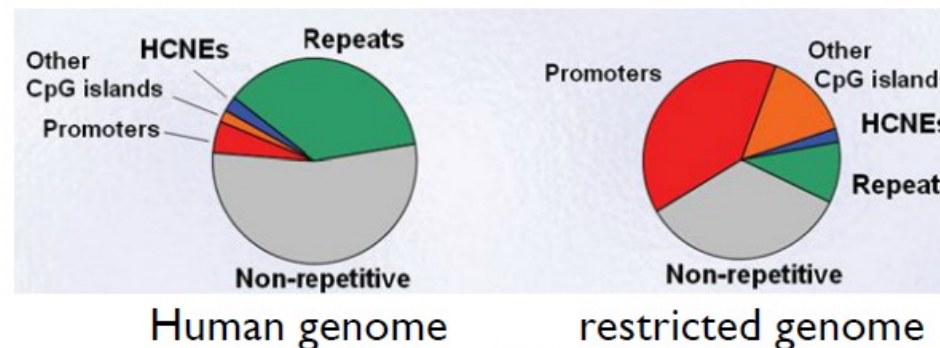
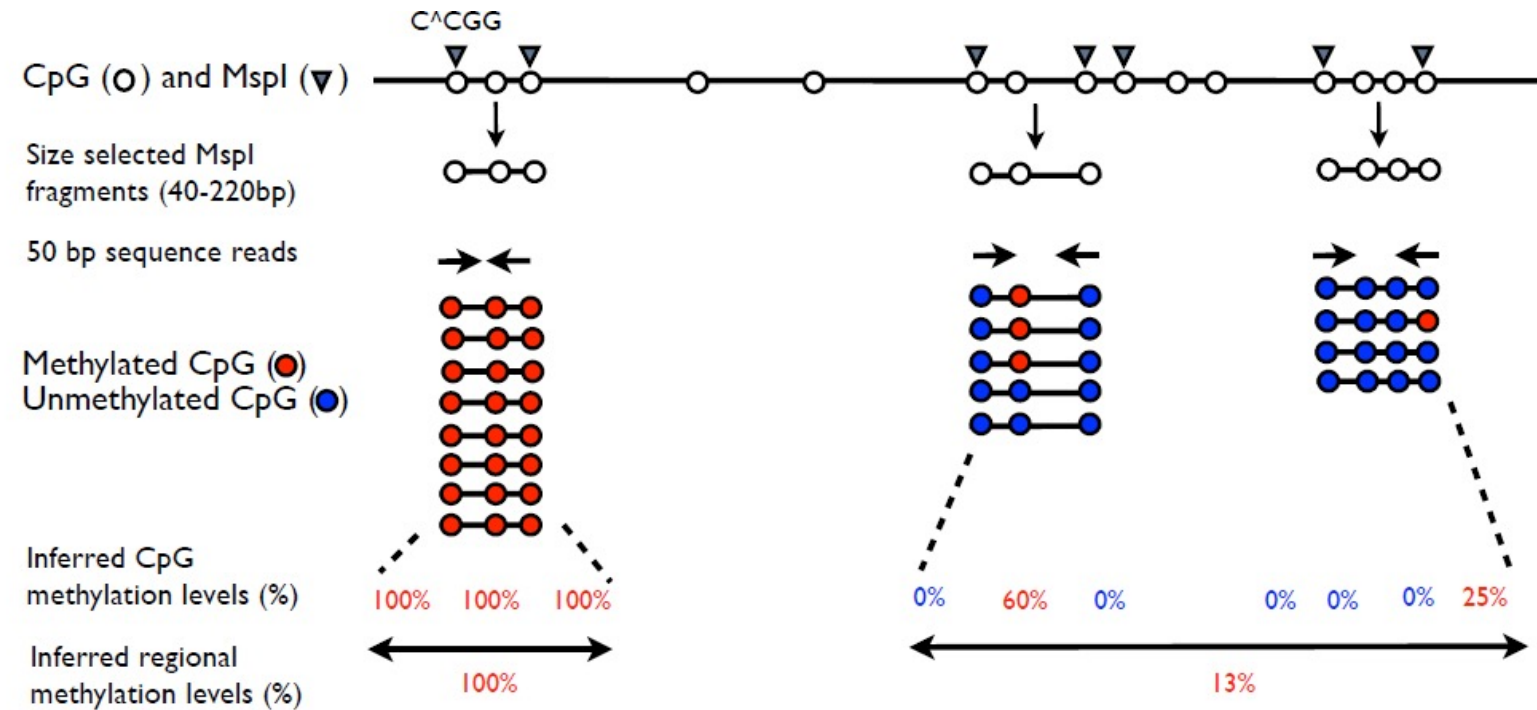
- Does not capture all promoters or CpG islands
- Results can vary depending on input DNA quality / contaminants in the sample

- MspI (C^ACGG)
 - TaqI (T^ACGA)
- Methylation insensitive restriction enzymes

- Base-pair resolution



RRBS



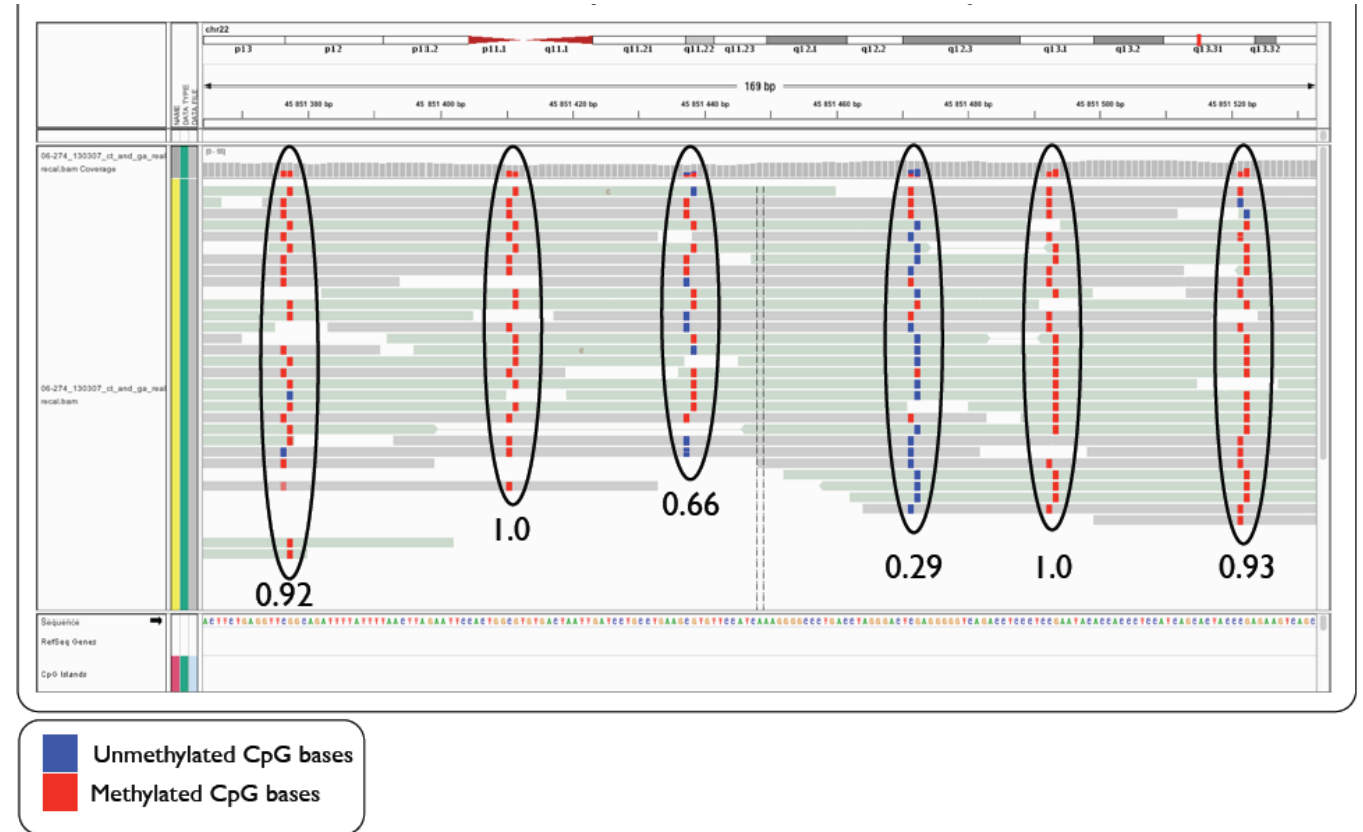
Genomic structures
enrichment after digestion
and size selection

HCNE:
highly conserved non-coding element

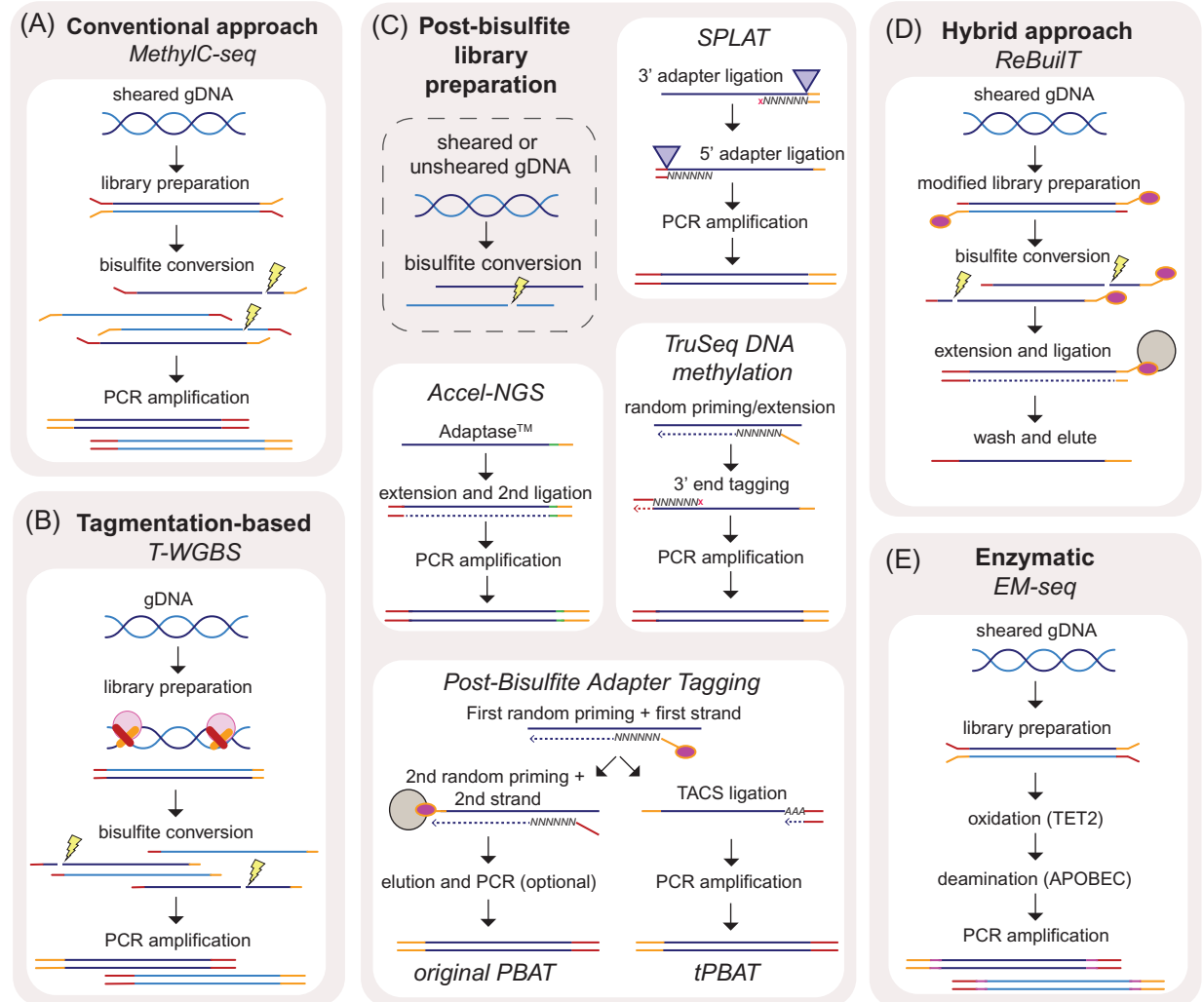
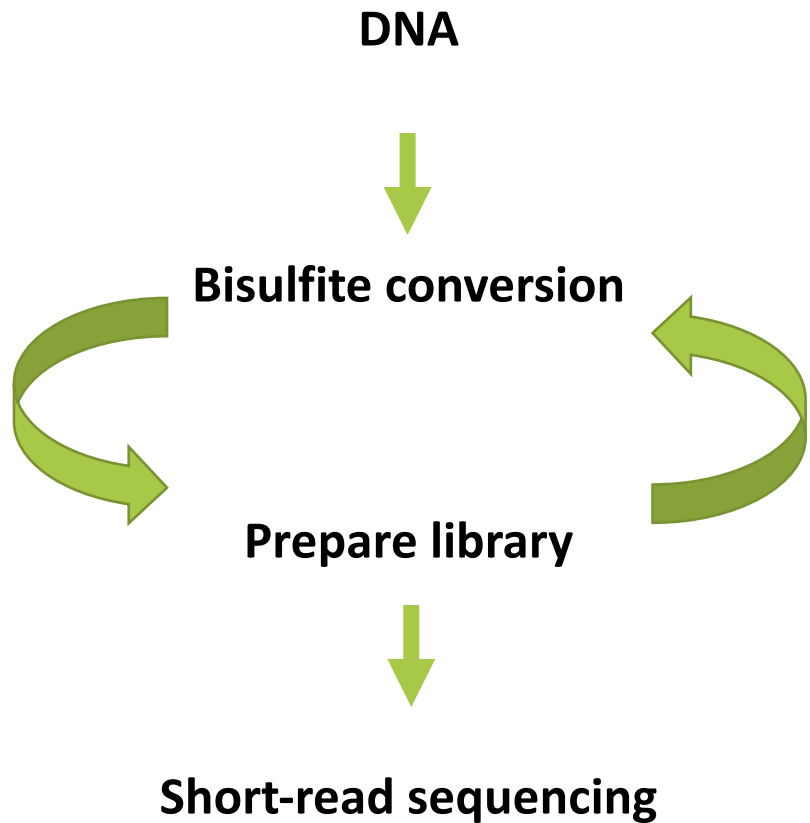
Whole Genome Methylome Sequencing



- Many acronyms ; WGBS, MethylC-seq, BS-seq
- “Unbiased” – no selection or enrichment
- Genome-wide coverage of all cytosines
- Base-pair resolution
- Uses bisulfite conversion or enzymatic conversion to distinguish methylated from unmethylated cytosines



Many different approaches ...



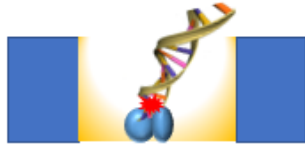
Whole Genome Methylome Sequencing



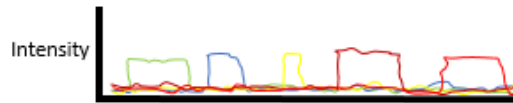
Direct read out of DNA modifications by single molecule, long read technologies (PacBio, Oxford Nanopore)

PacBio SMRT seq

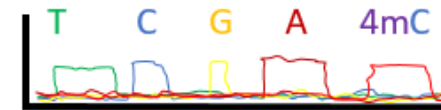
DNA passes thru
polymerase in an
illuminated volume



Raw output is fluorescent signal
of the nucleotide incorporation,
specific to each nucleotide

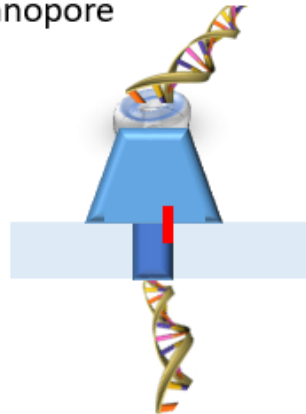


A,C,T,G have known pulse
durations, which are used to
infer methylated nucleotides

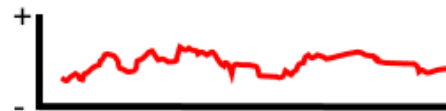


Oxford Nanopore

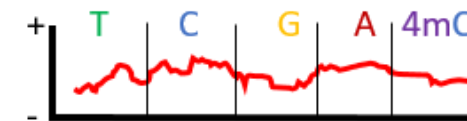
DNA passes thru
nanopore



Raw output is electrical signal
caused by nucleotide blocking
ion flow in nanopore



Each nucleotide has a specific
electric "signature"



In theory can detect all sorts of DNA modification-Challenge is to train models to correctly detect specific modifications

Cons; need a lot of native DNA for sequencing + compute

Pros: Phased information! Allele specific methylation. Imprinting

Short vs long-read sequencing, what's the difference?



Short-read

Illumina

Pros:

- Low cost
- High throughput
- Detect 5mC & 5hmC *depending on library prep applied
- Species agnostic

Cons:

- Requires conversion of (un)modified bases DNA with chemicals or enzymes
- 5mC cannot be distinguished from 5hmC (and other types of marks) without specific conversion approaches

Long-read

PacBio/ONT

Pros:

- Base modification can be read directly from sequencing
- Maintain phasing information
- Detect 4mC, 5mC, 5hmC, 5fC, 5caC, 6mA, etc
- Species agnostic

Cons:

- Cost (high coverage needed) – limiting for large genomes
- Difficult to detect signals
- Low throughput

Reproducibility & quality



Foxx J, Nordlund J, et al. The SEQC2 epigenomics quality control (EpiQC) study. Genome Biol 2021:
<https://doi.org/10.1186/s13059-021-02529-2>

EPIC arrays

- duplicate/triplicate at 3 labs

WGBS

- TruSeq DNA methylation (Illumina)
- Accel-NGS methyseq (Swift)
- SPLAT (Raine et al, NAR 2017)

OXBS

- TrueMethyl oxBS-seq (NuGEN)

Enzymatic deamination

- EM-seq (NEB)

ONT: direct methylation calling

7 cell lines

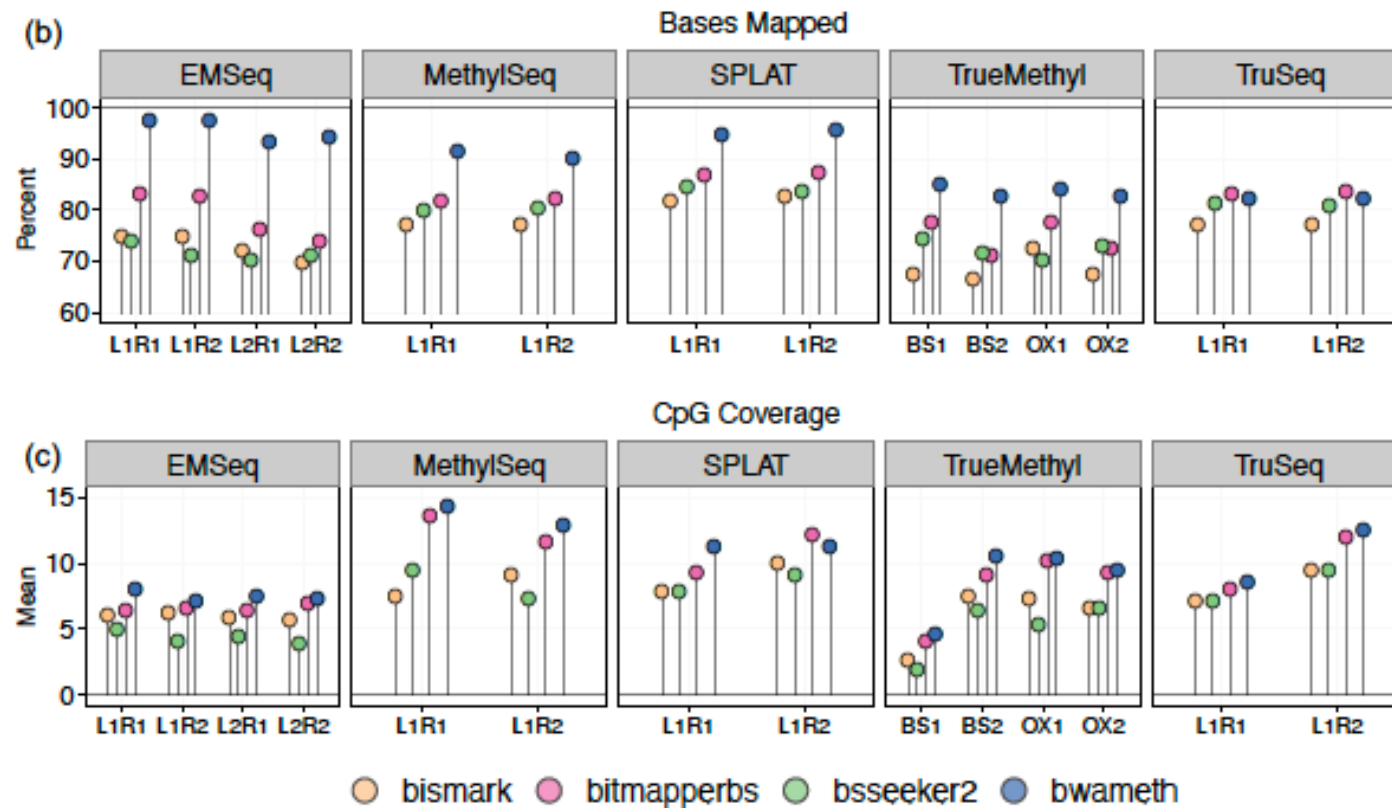
Alignment and methylation calling:

- BISMARK
- BitMapperBS
- BSSeeker2
- Bwa-meth
- Gem-bs

Microarray normalization

- 26 between-array and within-array normalization methods

Reproducibility & quality



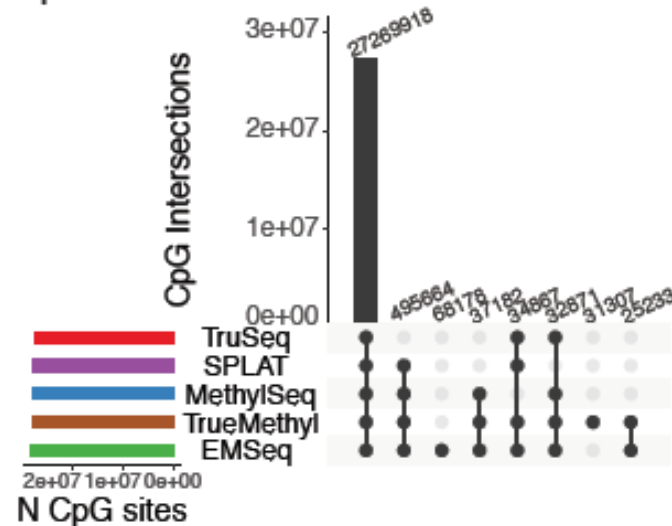
Overall, no major quantitative difference between pipelines but bwa-meth was easiest to implement and retained most data.

Noticeable inter- and intra-library differences

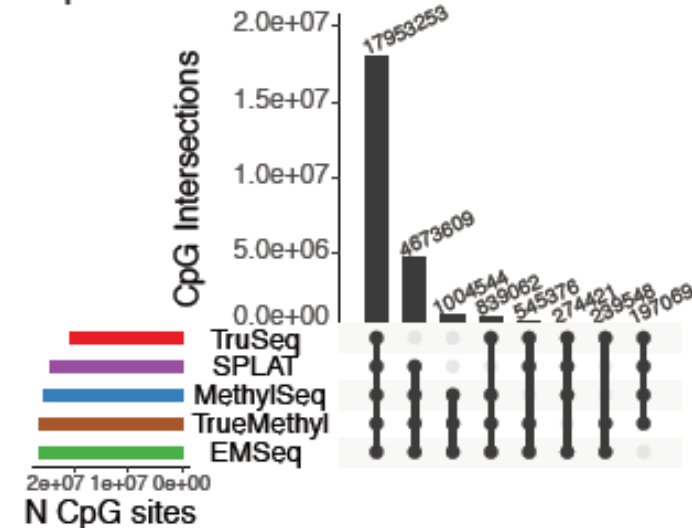
Reproducibility & quality



Average 20x GC coverage:
CpGs $\geq 1x$



CpGs $\geq 10x$



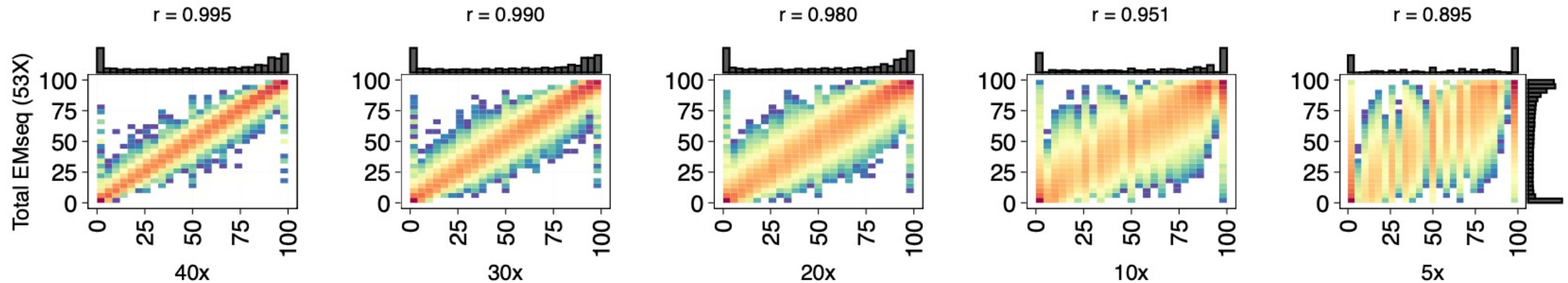
Overall, no major quantitative difference between methylation (beta-values) called after libraries were normalized for nr reads mapped (see next slide).

But they did differ in number of CpG sites detected!

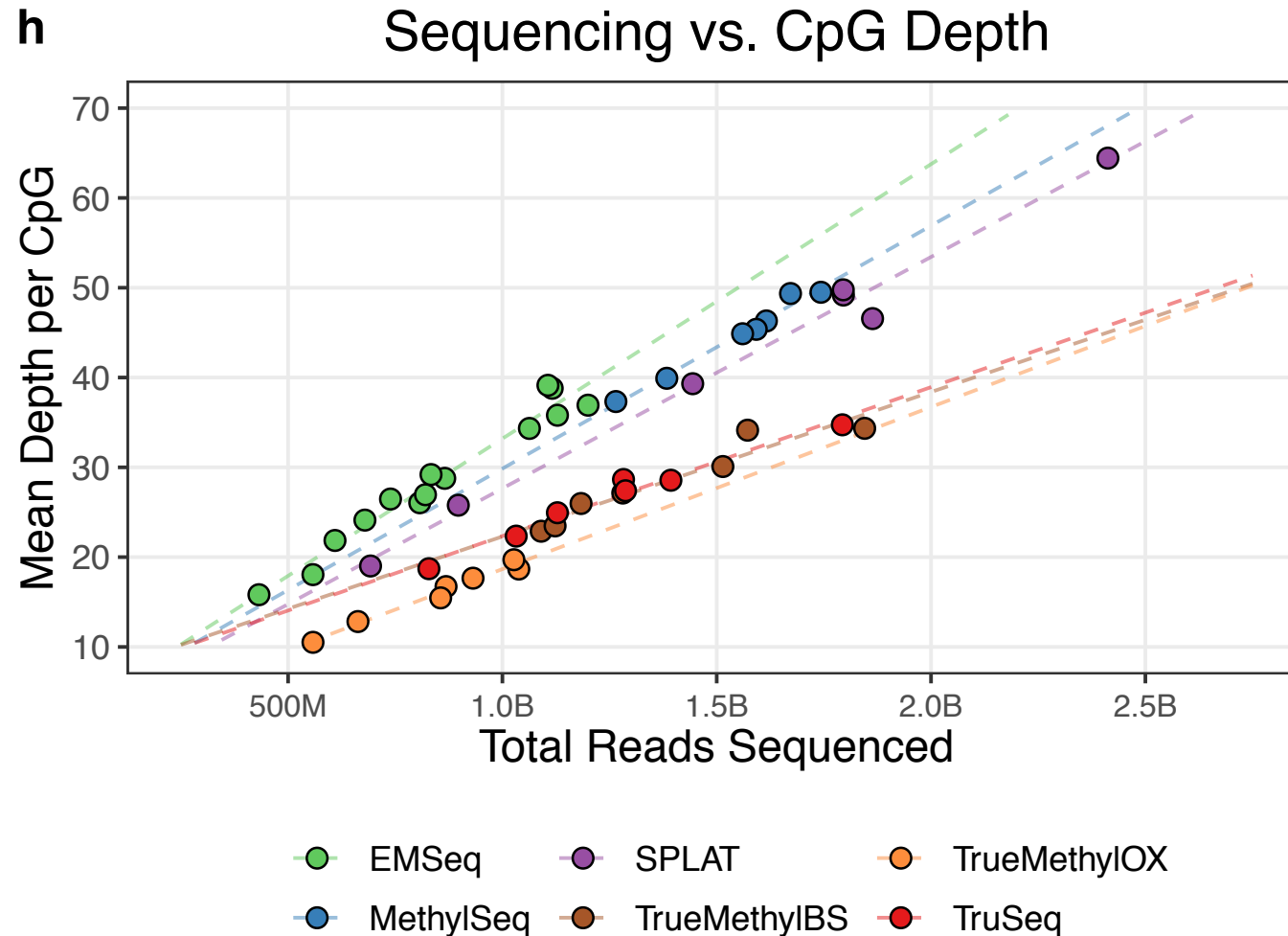
Reproducibility & quality



Correlation in DNA methylation estimation decreases as coverage decreases →



Reproducibility & quality

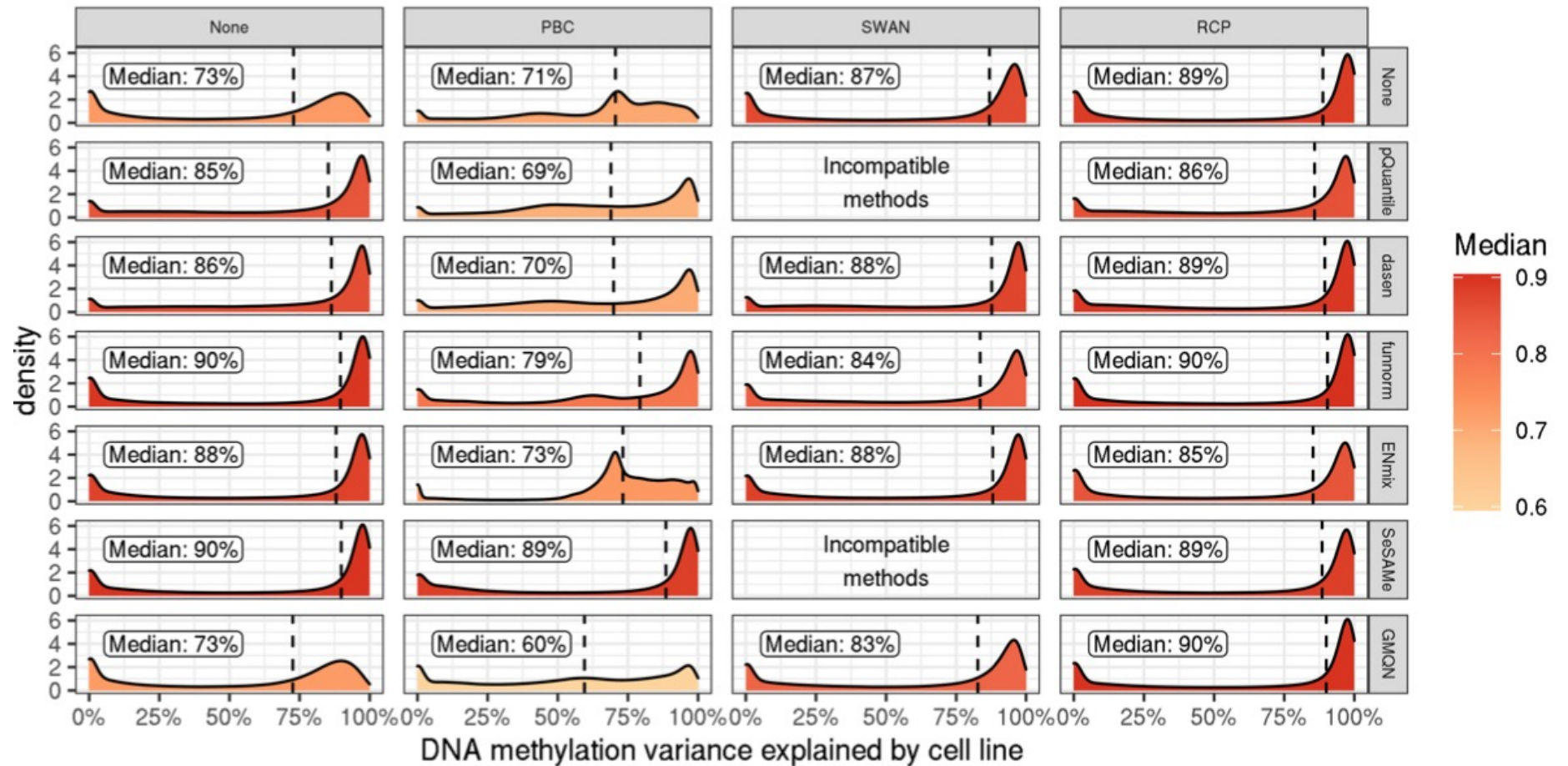


Reproducibility & quality

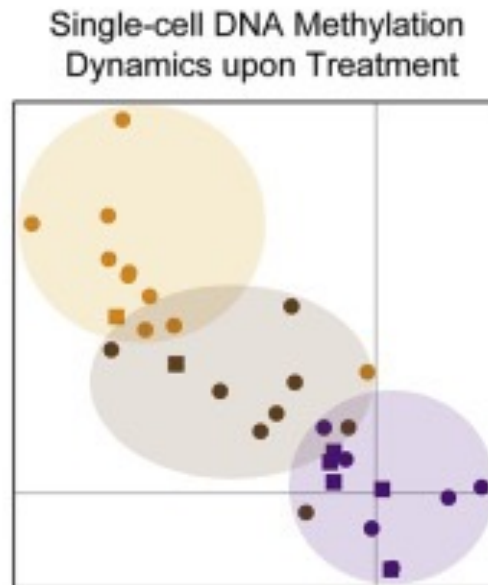
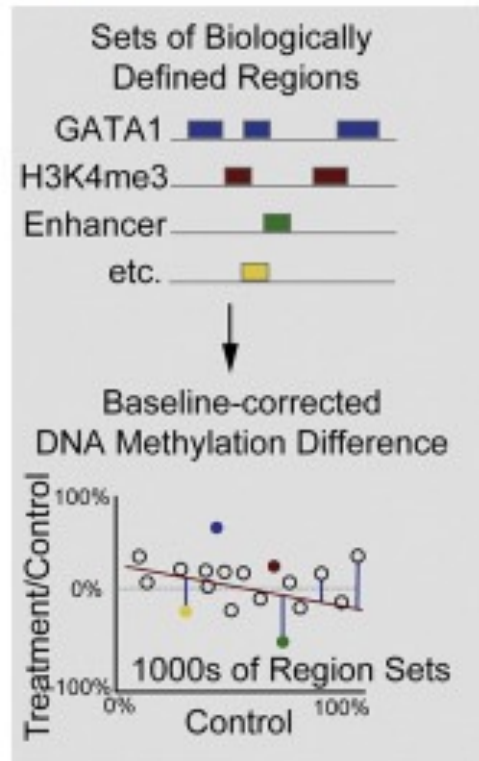
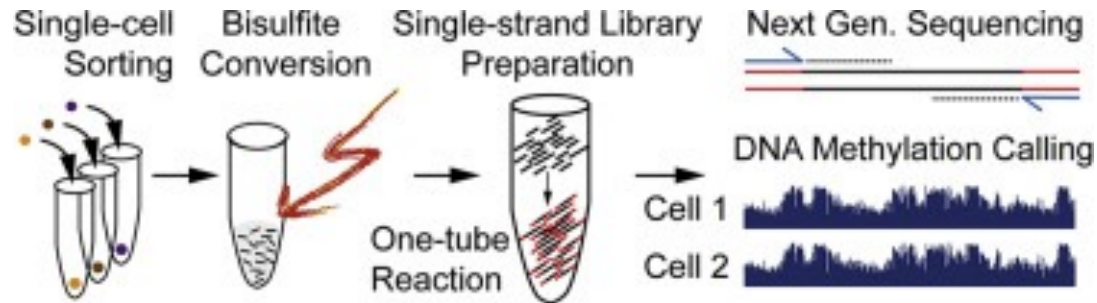


funnorm + RCP
worked best on
these samples

(a) Concordance between microarray replicates across the epigenome, by normalization pipeline



Single-cell WGBS



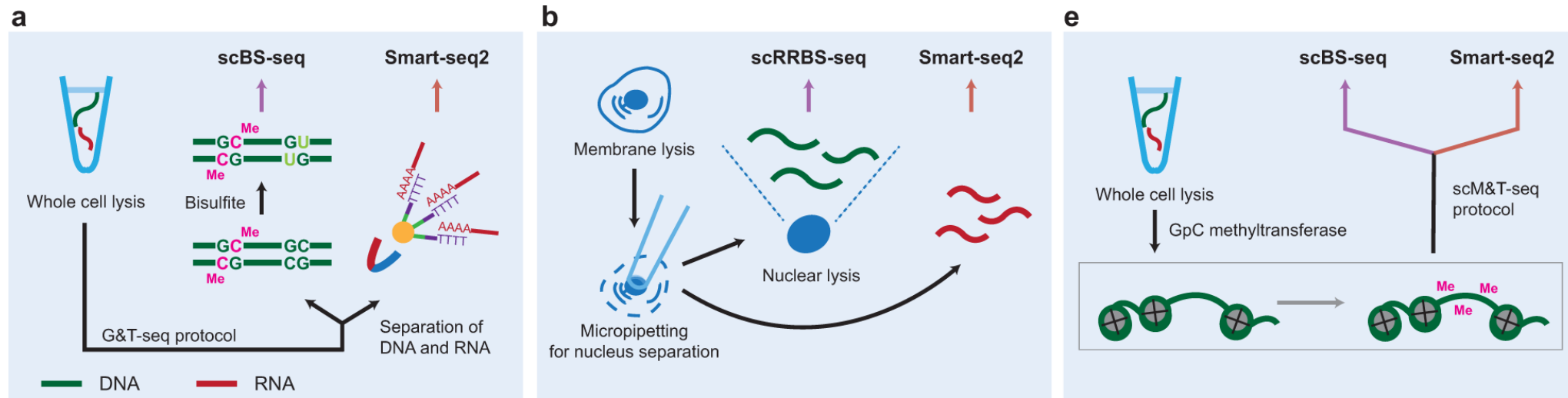
Single cell WGBS

- ✓ Single stranded library prep
- ✓ FACS sorting required (384 plates)
- ✓ Plate- based low throughput (although automation enable throughput of >1000 cells/exp)
- ✓ Expensive
- ✓ Sparse information-At most 50% CpG sites coverage, usually a lot less

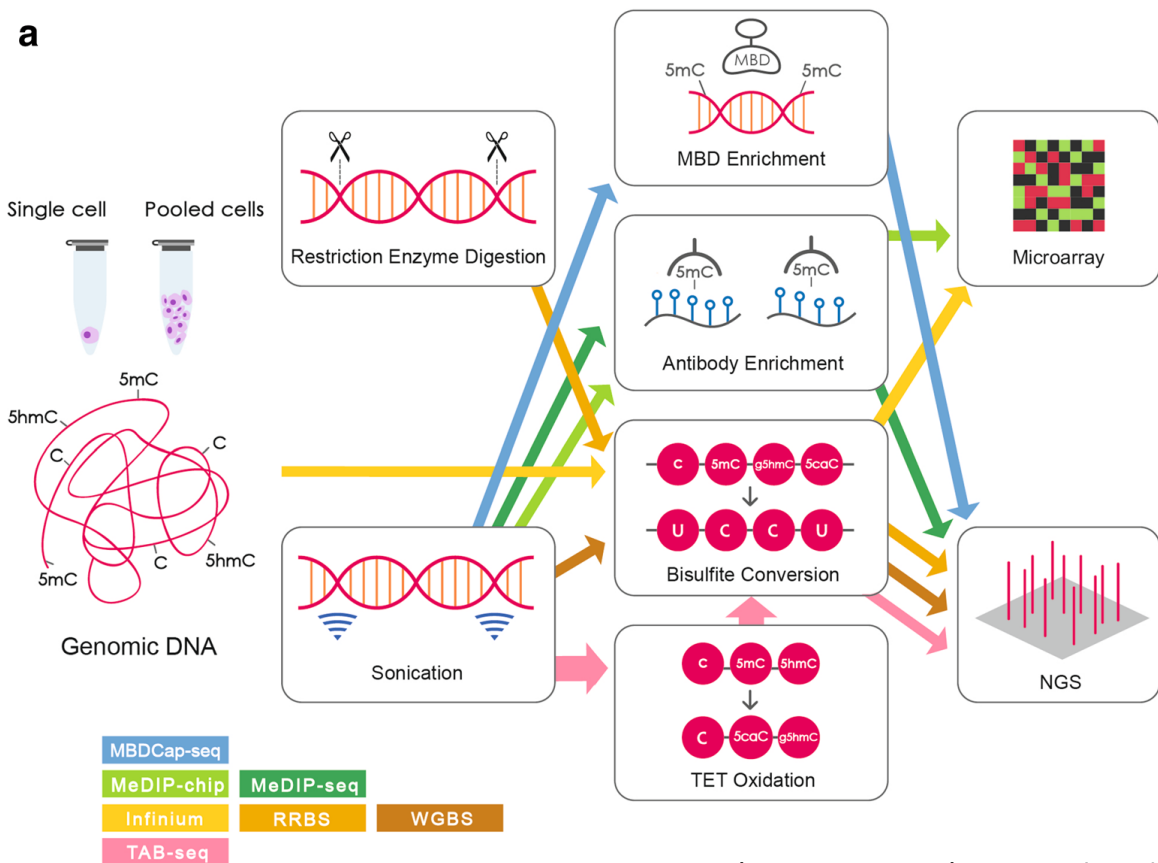
From "bulk" to single cells



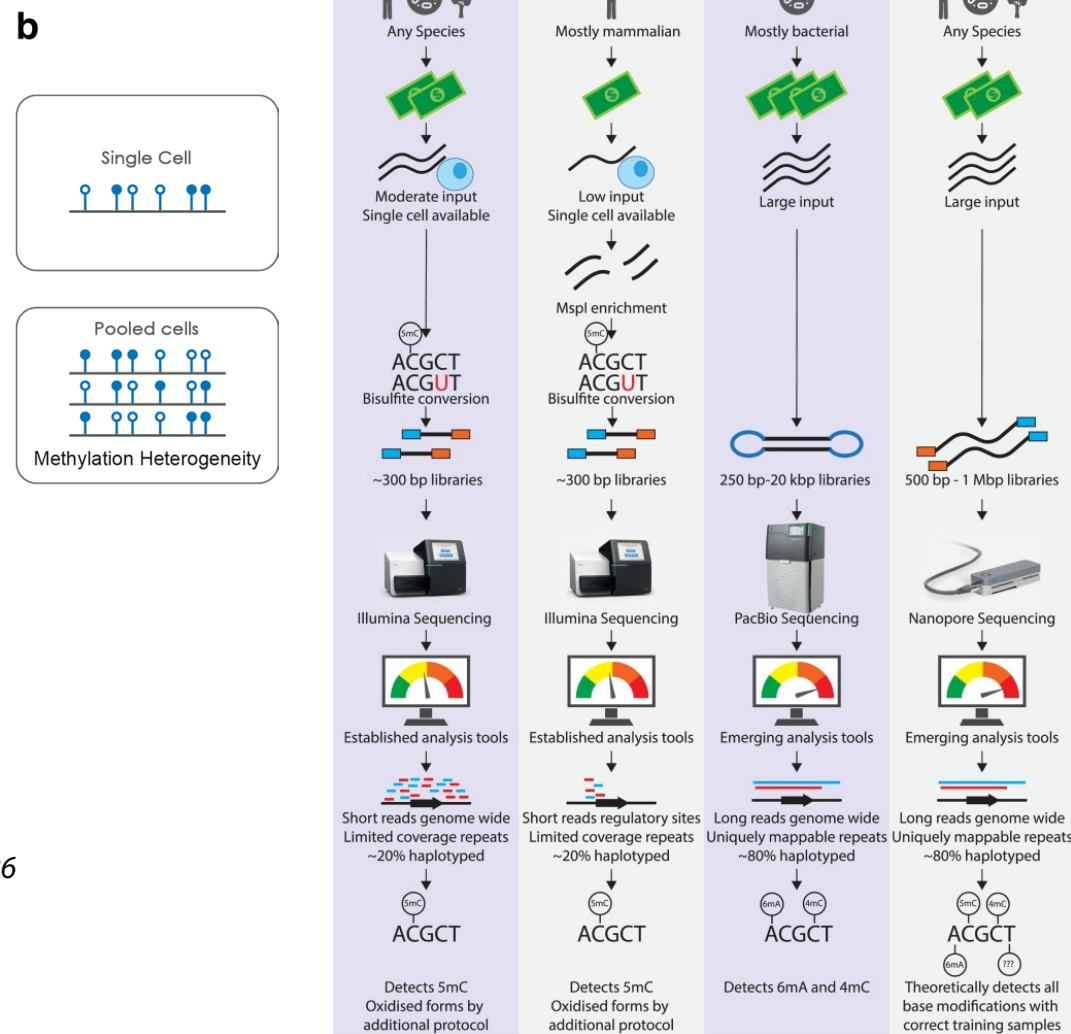
Numerous protocols exist for scWGSB, RRBS, etc – and even integrate transcriptomics in and DNA methylation measurements from the same cell!



In summary, there are many approaches for studying DNA methylation



Yong et al. *Epigenetics & Chromatin* (2016) 9:26
DOI 10.1186/s13072-016-0075-3

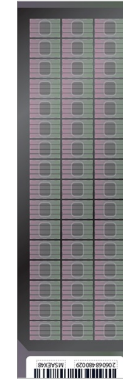


New methods coming



illumina®

Infinium Methylation Screening Array (48 samples/array)
Ultra high throughput, reduced cost
~270k CpG sites
50ng DNA input



illumina®

Illumina 5-base library prep

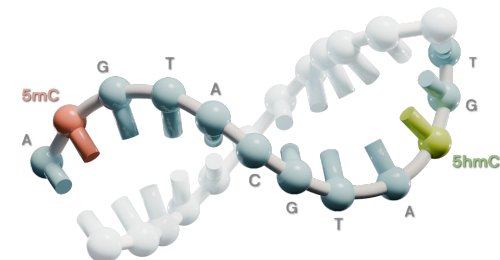


TET-assisted pyridine borane sequencing (TAPS)

**Convert methylated cytosine instead of unmethylated cytosine

bioRxiv modal

Whole genome analysis of 6-bases in 1* experiment



So which method should I choose?



nature
biotechnology

ANALYSIS

Comparison of sequencing-based methods to profile DNA methylation and identification of monoallelic epigenetic modifications

R Alan Harris^{1,*}, Ting Wang², Cristian Coarfa¹, Raman P Nagarajan³, Chibo Hong³, Sara L Downey³, Brett E Johnson³, Shaun D Fouse³, Allen Delaney⁴, Yongjun Zhao⁴, Adam Olshen³, Tracy Ballinger⁵, Xin Zhou², Kevin J Forsberg², Junchen Gu², Lorigail Echipare⁶, Henriette O'Geen⁶, Ryan Lister⁷, Mattia Pelizzola⁷, Yuanxin Xi⁸, Charles B Epstein⁹, Bradley E Bernstein⁹⁻¹¹, R David Hawkins¹², Bing Ren^{12,13}, Wen-Yu Chung^{14,15}, Hongcang Gu⁹, Christoph Bock^{9,16-18}, Andreas Gnirke⁹, Michael Q Zhang^{14,15}, David Haussler³, Joseph R Ecker⁷, Wei Li⁸, Peggy J Farnham⁶, Robert A Waterland¹⁻¹⁹, Alexander Meissner^{9,16,17}, Marco A Marra⁴, Martin Hirst⁴, Aleksandar Milosavljevic¹ & Joseph F Costello³

Fox et al. *Genome Biology* (2021) 22:332
<https://doi.org/10.1186/s13059-021-02529-2>

Genome Biology

RESEARCH

Open Access

The SEQC2 epigenomics quality control (EpiQC) study



Jonathan Fox^{1,2†}, Jessica Nordlund^{3,4†}, Claudia Lalancette^{5†}, Ting Gong^{6†}, Michelle Lacey^{7†}, Samantha Lent^{8†}, Bradley W. Langhorst⁹, V. K. Chaitanya Ponnaluri⁹, Louise Williams⁹, Karthik Ramaswamy Padmanabhan⁵, Raymond Cavalcante⁵, Anders Lundmark^{3,4}, Daniel Butler¹, Christopher Mozsary¹, Justin Gurvitch¹, John M. Grealis¹⁰, Masako Suzuki¹⁰, Mark Menor⁶, Masaki Nasu⁶, Alicia Alonso^{1,11}, Caroline Sheridan^{1,11}, Andreas Scherer^{4,12}, Stephen Bruinsma¹³, Gosia Golda¹⁴, Agata Muszynska¹⁵, Pawel P. Łabaj¹⁵, Matthew A. Campbell⁹, Frank Wos¹⁶, Amanda Raine^{3,4}, Ulrika Liljedahl^{3,4}, Tomas Axelsson^{3,4}, Charles Wang¹⁷, Zhong Chen¹⁷, Zhaowei Yang^{17,18}, Jing Li^{17,18}, Xiaopeng Yang¹⁹, Hongwei Wang²⁰, Ari Melnick¹, Shang Guo²¹, Alexander Blume²², Vedran Franke²², Inmaculada Ibanez de Caceres^{4,23}, Carlos Rodriguez-Antolin^{4,23}, Rocio Rosas^{4,23}, Justin Wade Davis⁹, Jennifer Ishii¹⁶, Dalila B. Megherbi²⁴, Wenming Xiao²⁵, Will Liao¹⁶, Joshua Xu²⁶, Huixiao Hong²⁶, Baitang Ning²⁶, Weida Tong²⁶, Altuna Akalin²², Yunliang Wang^{21*}, Youping Deng^{6*} and Christopher E. Mason^{1,2,7,28*}

Essays in Biochemistry (2019) 63:639–648
<https://doi.org/10.1042/EBIC20190027>



Review Article

Latest techniques to study DNA methylation

Quentin Gouli^{1,2} and Andrew Keniry^{1,2}

¹Epigenetics and Development Division, Walter and Eliza Hall Institute of Medical Research, Parkville, Australia; ²Department of Medical Biology, University of Melbourne, Parkville, Australia

Lee et al. *Experimental & Molecular Medicine* (2020) 52:1428–1442
<https://doi.org/10.1038/s12276-020-0420-2>

Experimental & Molecular Medicine

REVIEW ARTICLE

Open Access

Single-cell multiomics: technologies and data analysis methods

Jeongwoo Lee¹, Do Young Hyeon¹ and Daehee Hwang¹

nature
genetics

PERSPECTIVE

<https://doi.org/10.1038/s41588-018-0290-x>

Single-cell and single-molecule epigenomics to uncover genome regulation at unprecedented resolution

Efrat Shema^{1,2,4}, Bradley E. Bernstein^{1,2} and Jason D. Buenrostro^{1,2,3*}

Chromatin (2016) 9:26
0075-3

Epigenetics & Chromatin

REVIEW

Open Access

Profiling genome-wide DNA methylation

Wai-Shin Yong^{1†}, Fei-Man Hsu^{2†} and Pao-Yang Chen^{1*}



Epigenomics services offered by the National Genomics Infrastructure (NGI)



NGI is a facility within the
**SciLifeLab Genomics
Platform** located at two nodes:

NGI-Uppsala

- SNP&SEQ Technology Platform (UU)
- Uppsala Genome Centre (UU)

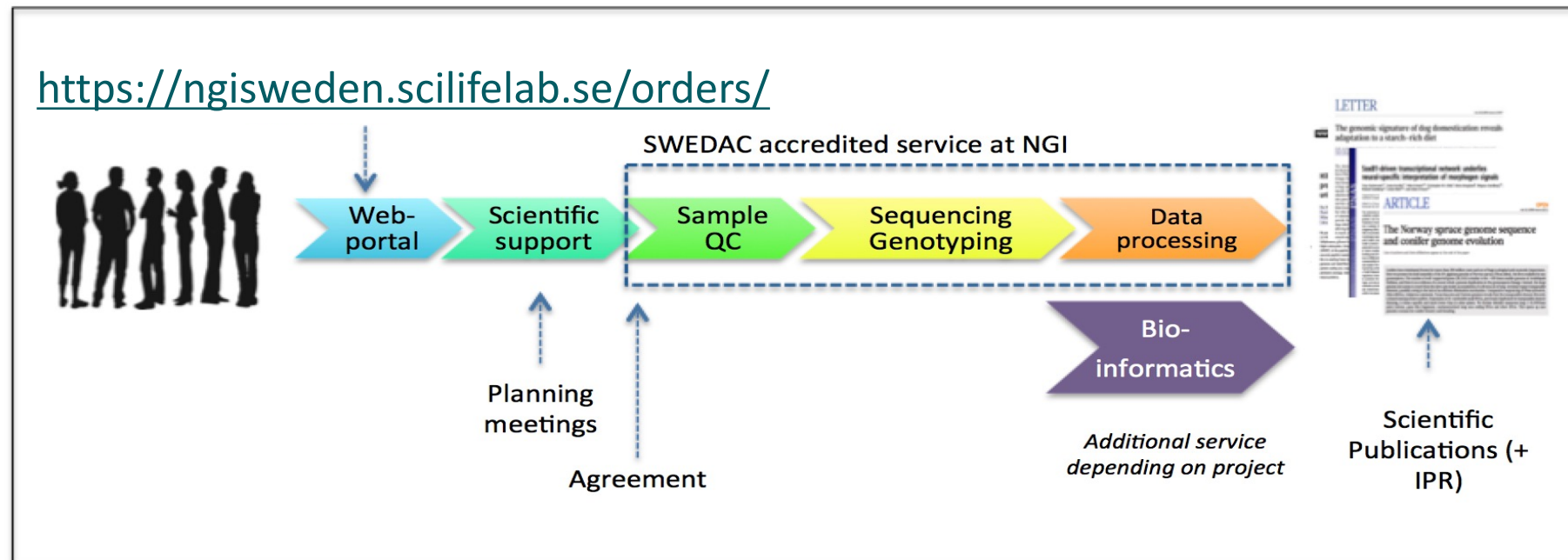
NGI-Stockholm

- SciLifeLab Solna (KTH, KI, SU)

NGI's project portal



- All projects submitted through a **common order system**
- Projects are dynamically allocated between Stockholm/Uppsala depending on type of application, queue situation, or request by researcher

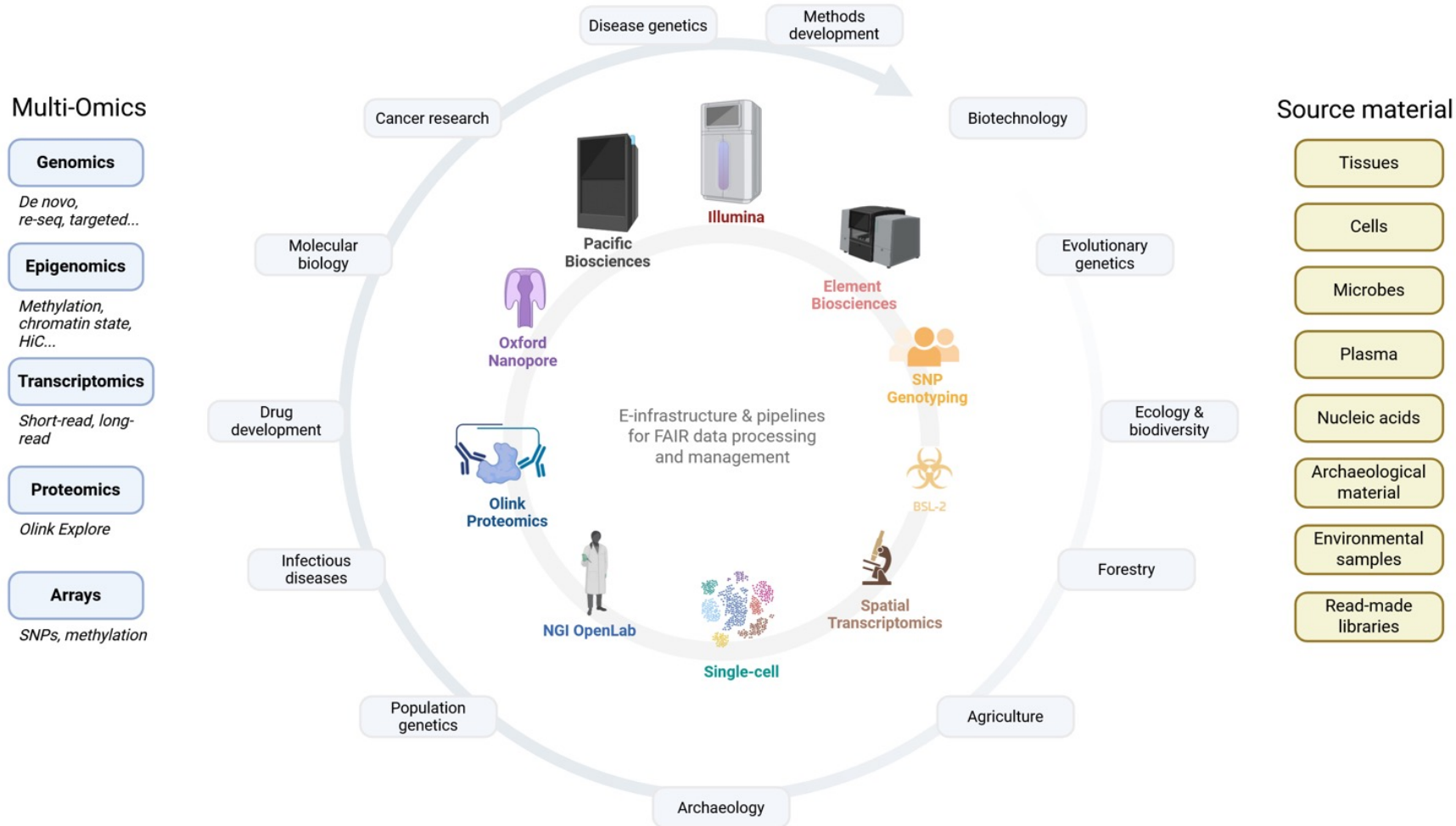


From single SNPs to whole genomes



Genomics Platform:

Expertise, infrastructure and services in genomic technologies and applications

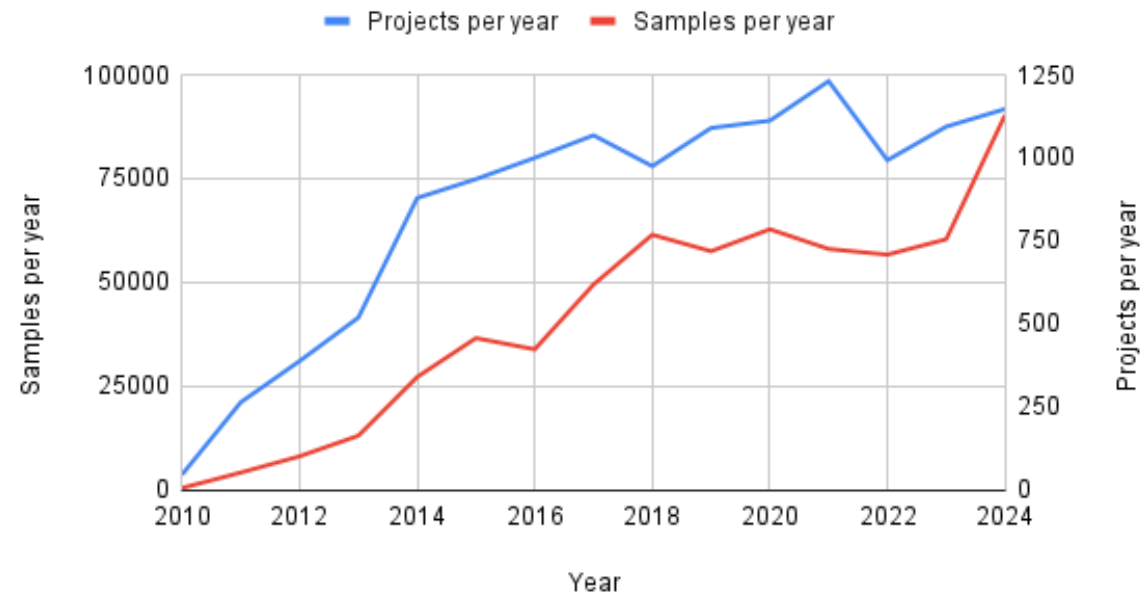


Our mission:

- Enabling research that would not otherwise be possible
- Added value by pre- and post-project consulting
- Flexibility for most sample types and quality
- High data quality & reproducibility – **ISO17025**
- Up to date equipment
- Cheap
- Flexible
- ...



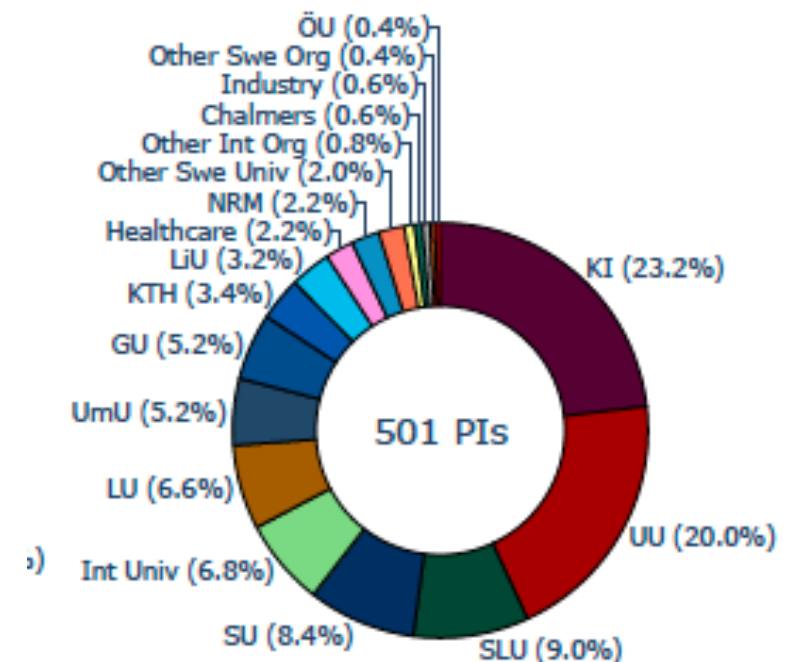
Sequencing: Samples and Projects per Year



Statistics for 2024:

- 1147 projects / 90,000 samples (70k sequencing, 20k genotyping)
- **985 Terabases** (10^{12}) of sequence data

Users 2024



As of Jan 1, 2025 NGI has delivered a total of 8.3 Petabases (10^{15}) of sequencing data

Support



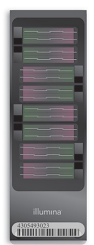
Pre support

- **Project design** via discussions with expert project coordinators
- **Advise** in sample collection and/or preparation
- **DNA extraction services available** for specific applications
- **Sample quality (QC)** for all incoming samples and user-made libraries

Post support

- Control over produced data: making sure data meet our **high standards** in terms of quality and yield.
- Open source Bioinformatic pipelines for a wide range of applications: *NF-core*
- Data delivered via **SciLifeLab Data Delivery System (DDS)**

Epigenetic methods available at NGI



EPIC Arrays:

500 ng DNA

Minimum sample size 15 samples: lower cost per sample for large projects

Short-read

Whole genome methylome sequencing with SPLAT (WGBS) or EM-Seq

Twist targeted methylation

~500 ng DNA

Long-read

whole genome sequencing (+base modifications)

PacBio Sequell II / Oxford Nanopore PromethION

Cost depends on genome size and epigenetic marks analyzed

Single-cell:

scATAC-seq (10x Genomics)

scWGBS with SPLAT

RRBS:

500 ng DNA
~**2000** SEK/sample

**limited availability*

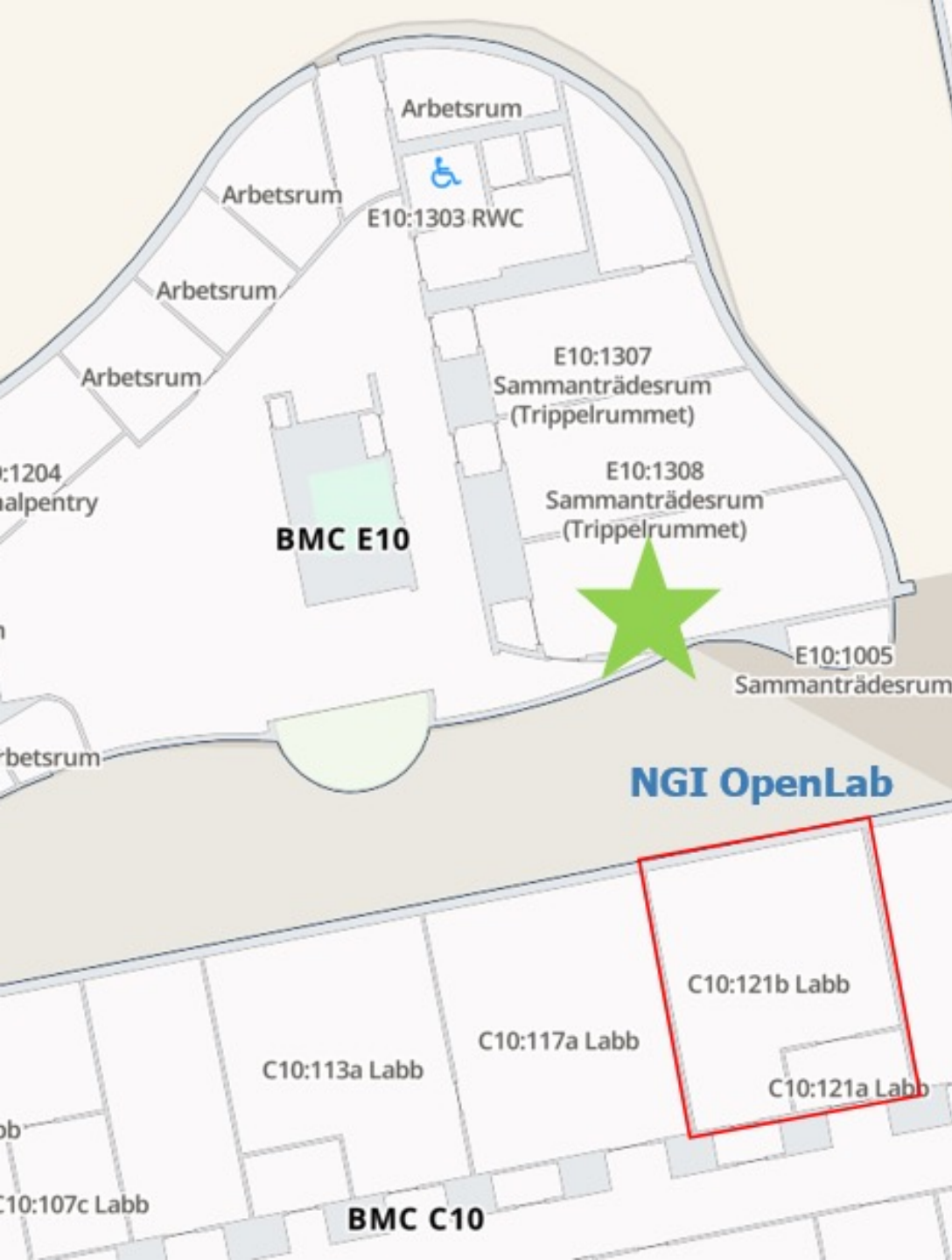
ATAC-seq

>50.000 cells
~**2000** SEK/sample

**limited availability*

HiC

method for mapping genome-wide DNA contacts

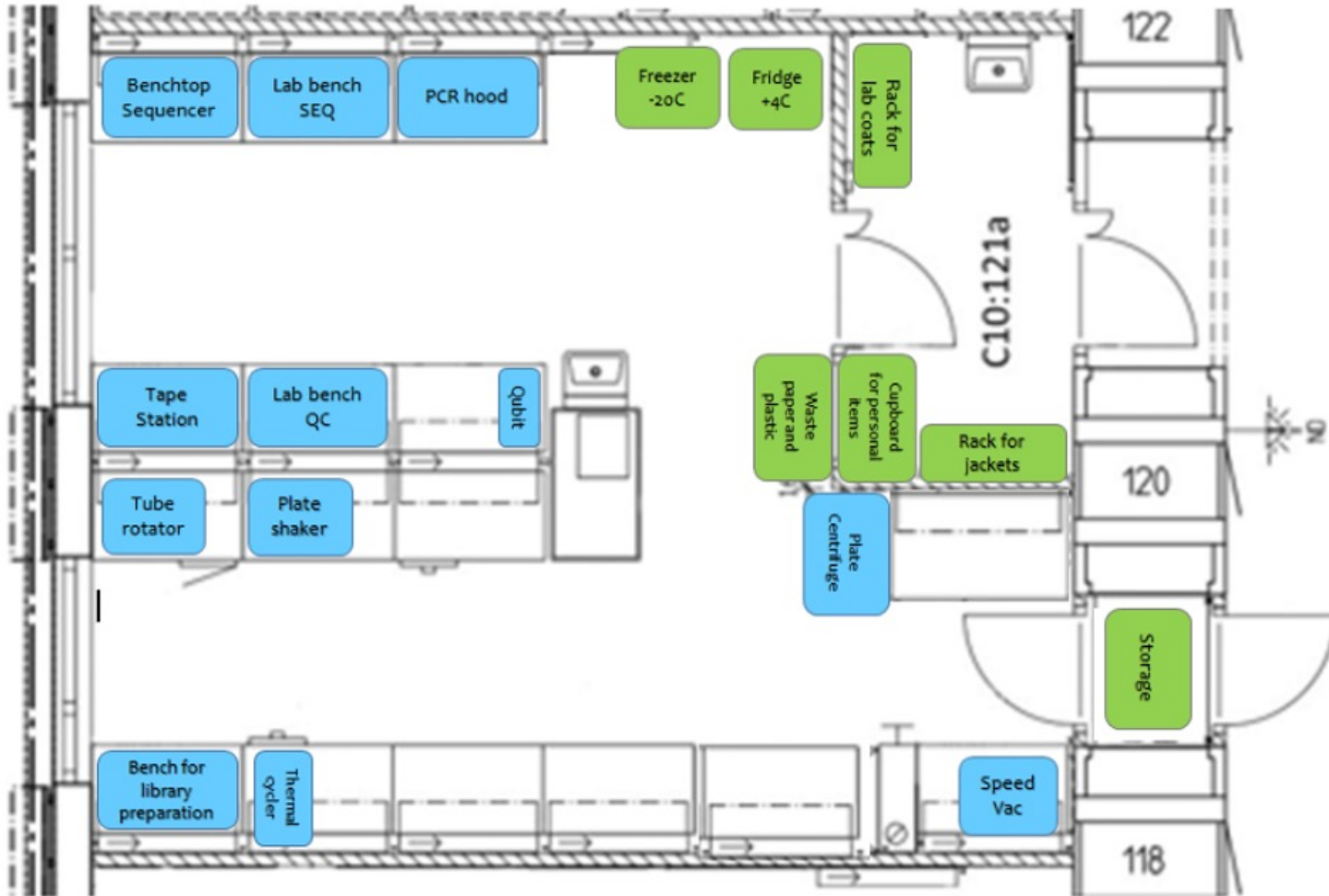


NGI OpenLab

- Located at BMC C10:1
- Open for users from Swedish academia, access available to others if capacity allows (full cost model)
- Operates on trust, mutual respect and a shared commitment for taking care of the equipment and common labs pace

<https://ngisweden.scilifelab.se/applications/ngi-openlab/>

NGI OpenLab



Workstations

- Sample quality control
- Sequencing Library preparation
- Sequencing

Coming soon- Single-cell



Follow us!



LinkedIn: **SciLifeLab National Genomics Infrastructure - NGI**



Information about sequencing applications that NGI supports:

<https://ngisweden.scilifelab.se>

Don't hesitate to reach out to NGI's project coordinators:

support@ngisweden.se

-or me-

Jessica.nordlund@medsci.uu.se



Swedish
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