



# Atelier scRNA-seq

# Technology for scRNA-seq and data processing

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École de bioinformatique AVIESAN-IFB-INSERM 2024





# Organisation of this session

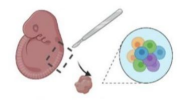
- From cells to nucleotide sequences (reads)
  - focus on the 10X genomics technology
  - $\circ$  how are the reads organised
- Preprocessing : from reads to raw count matrix
  - quality check (FASTQC)
  - mapping (STAR)
  - $\circ$  how is annotation used
  - barcode and UMI treatment

### Global overview of a scRNA-seq experiment

Tissue dissection + cell dissociation

Cell partitioning + mRNA capture

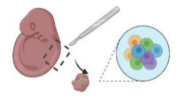
#### Library preparation + sequencing



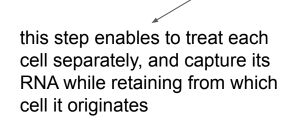


### Global overview of a scRNA-seq experiment

Tissue dissection + cell dissociation

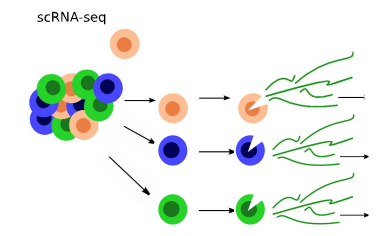


Cell partitioning + mRNA capture



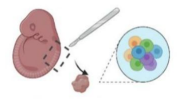
#### Library preparation + sequencing





### Global overview of a scRNA-seq experiment

Tissue dissection + cell dissociation

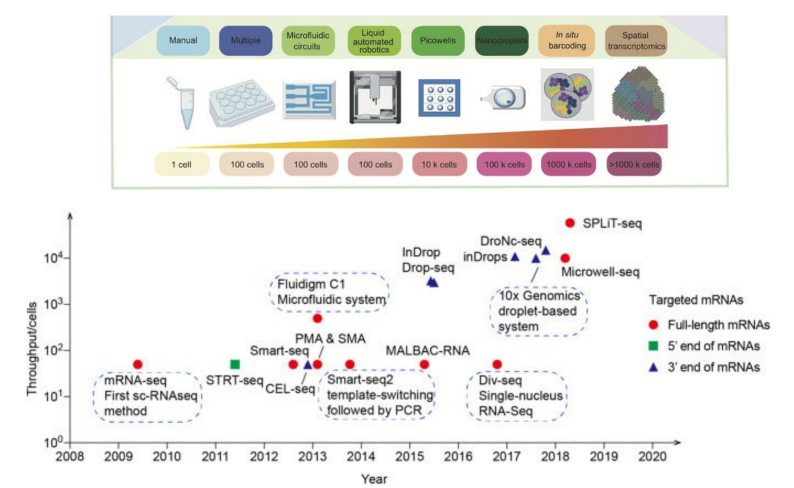


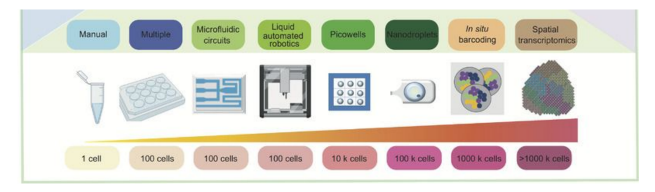
Cell partitioning + mRNA capture

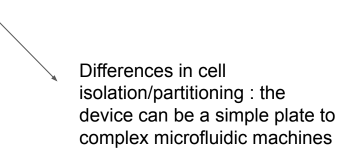
various technologies

developed over time for this specific step Library preparation + sequencing

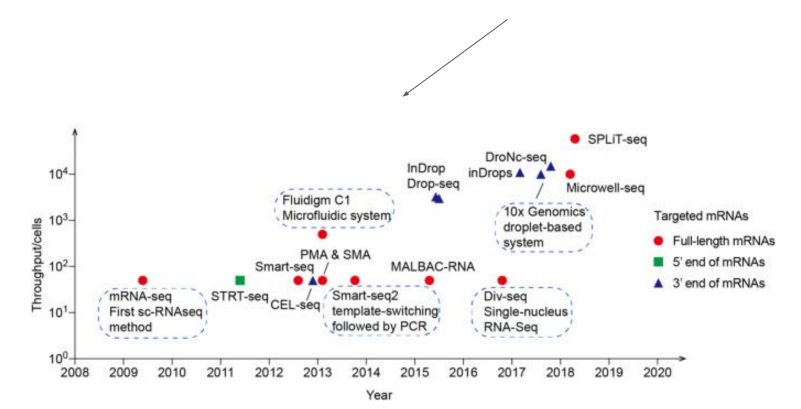








The number of cells that can be studied has grown from a handful to >10,000 in 10 years



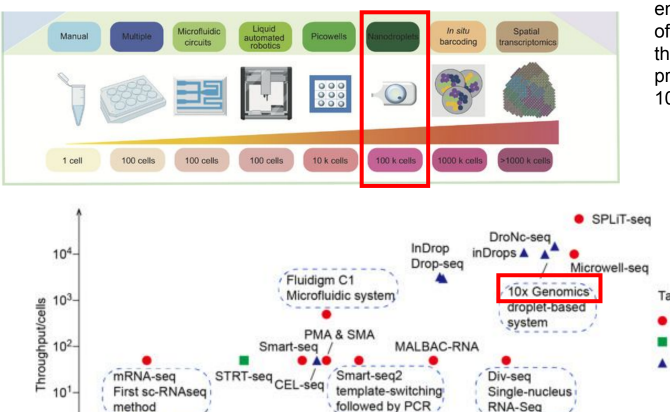
2009

2010

2011

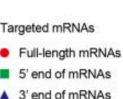
2012

2013



The technology that has enabled widespread usage of scRNA-seq approach is the droplet-based approach proposed by the company 10X Genomics.

> 10X GENOMICS®



2015

2016

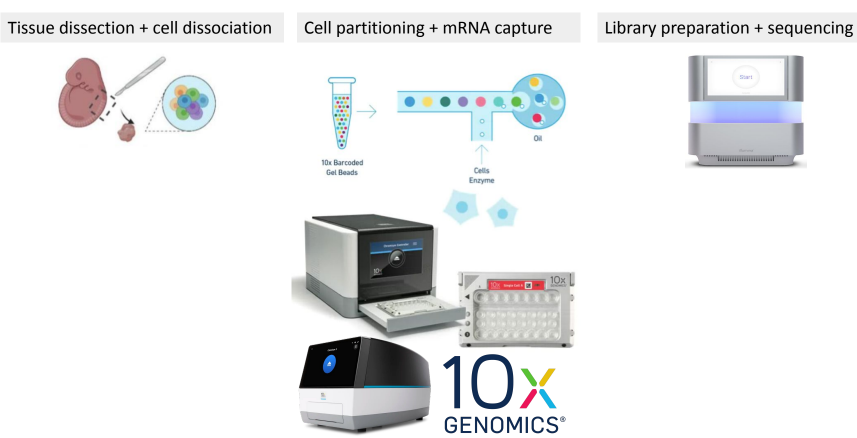
2017

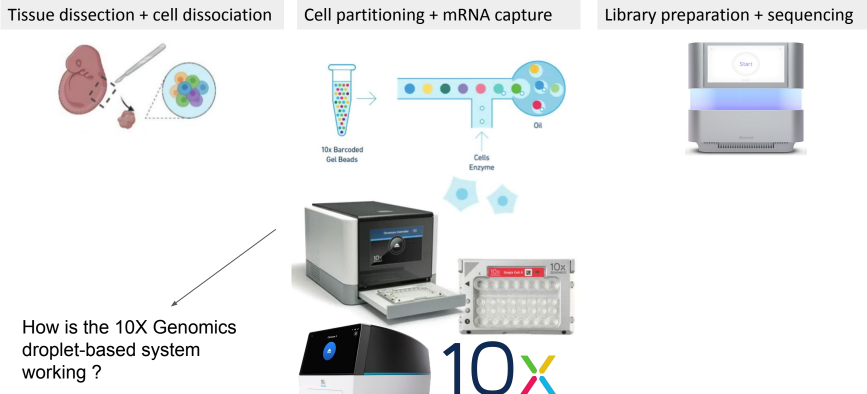
2018

2019

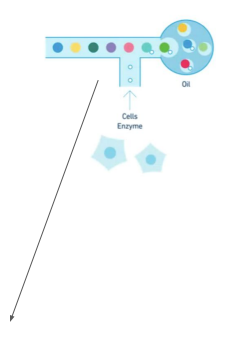
2020

2014

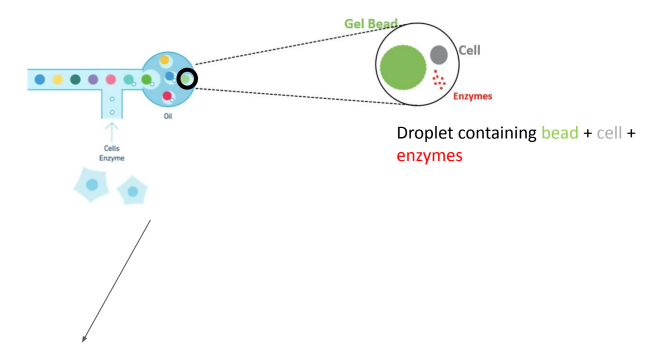




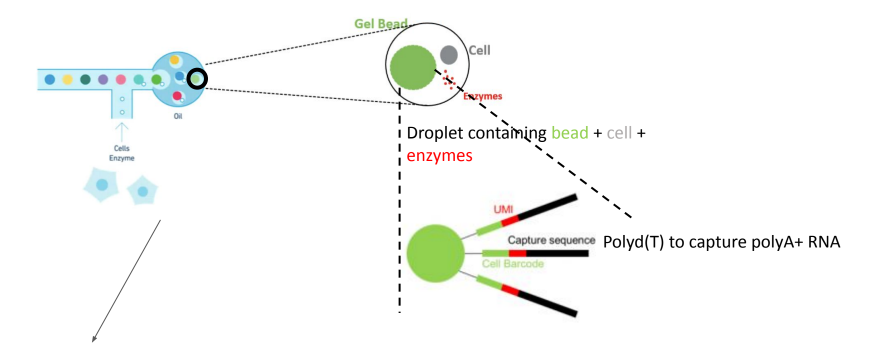
**GENOMICS®** 



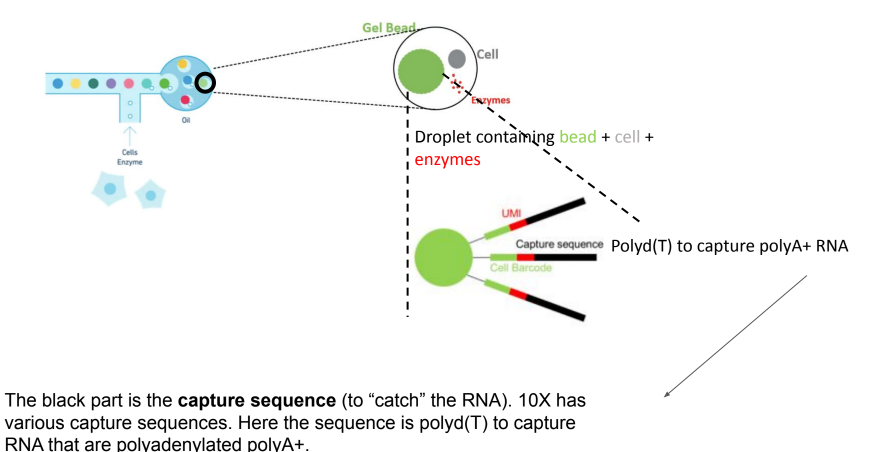
Cells and gel beads arrive in the device from 2 separate channels

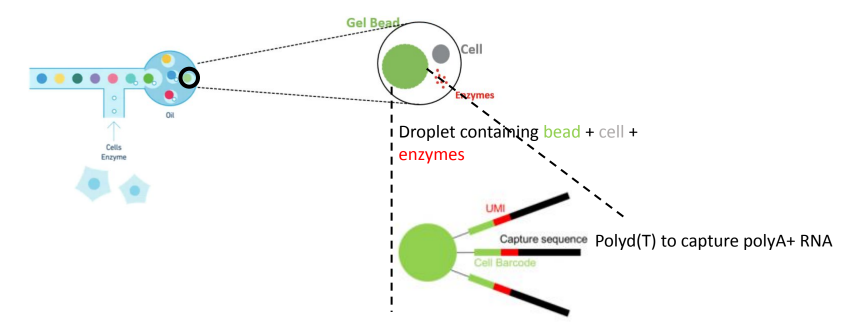


A single cell and a single gel bead (+ enzymes) are then encapsulated in a droplet



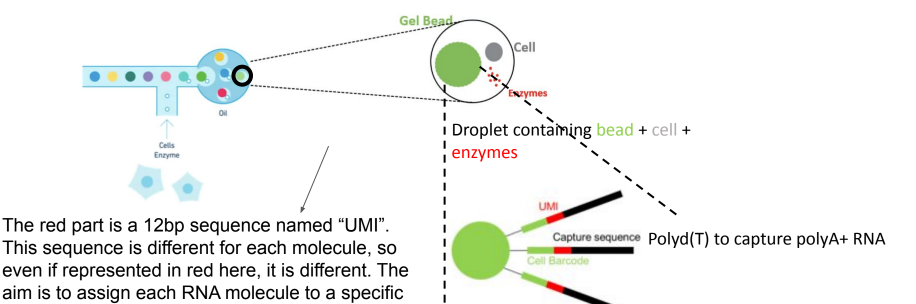
The gel bead is special : it is covered with molecules made of 3 parts





**Cell Barcode (16bp)** = sequence specific to each bead (so each cell)

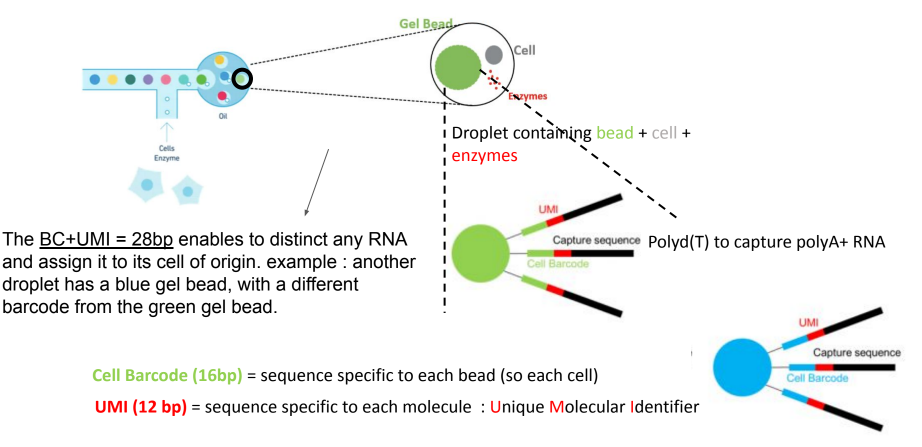
The green part is a 16bp sequence named "barcode". This same sequence is all over the bead. These barcodes are created by 10X and the list is available.

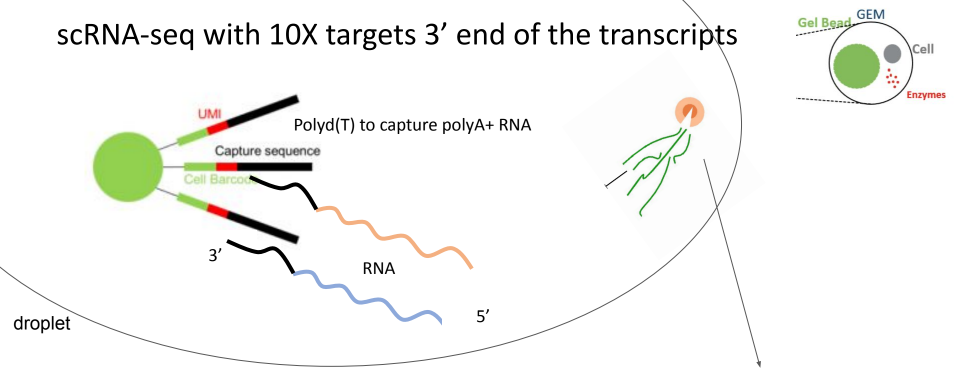


**Cell Barcode (16bp)** = sequence specific to each bead (so each cell)

UMI.

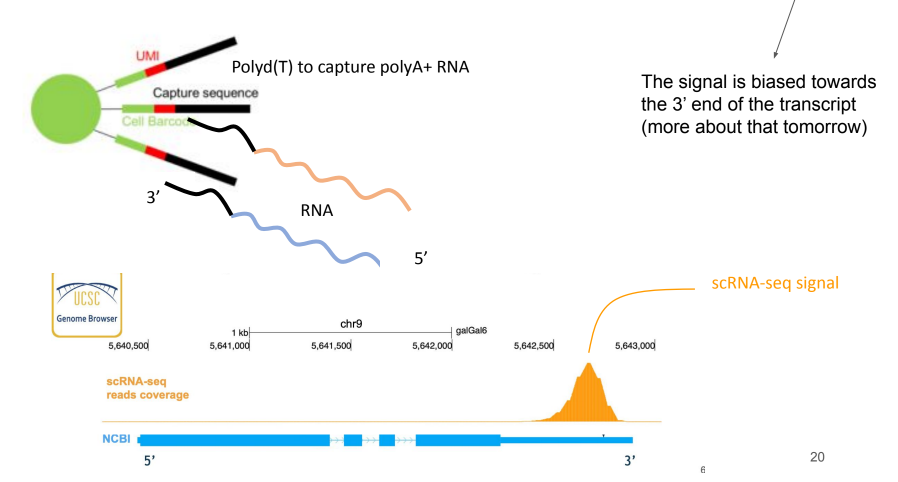
UMI (12 bp) = sequence specific to each molecule : Unique Molecular Identifier





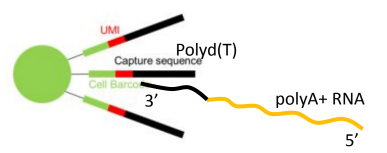
The cell is then lysed within the droplet. RNA is released (but contained in the droplet). polyA+ RNA are captured from the 3'end on the polyd(T) sequence

### scRNA-seq with 10X targets 3' end of the transcripts



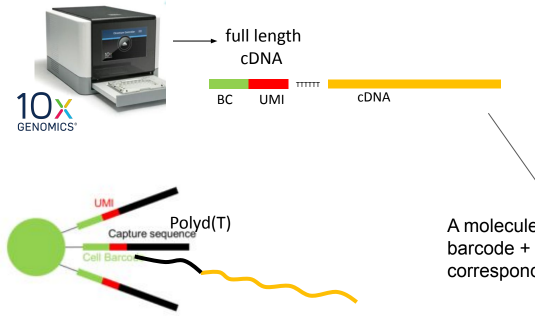


At the end of this step, what actually comes out of the 10X Genomics device ?



**Cell Barcode** = sequence specific to each cell

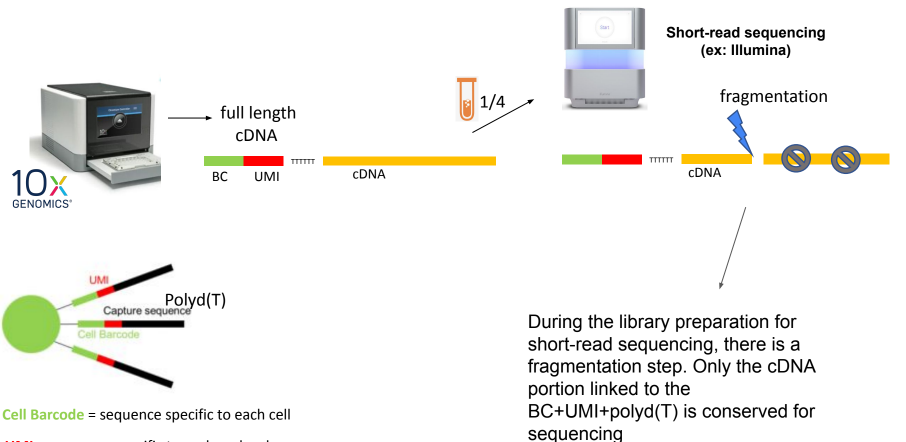
**UMI** = sequence specific to each molecule : Unique Molecular Identifier



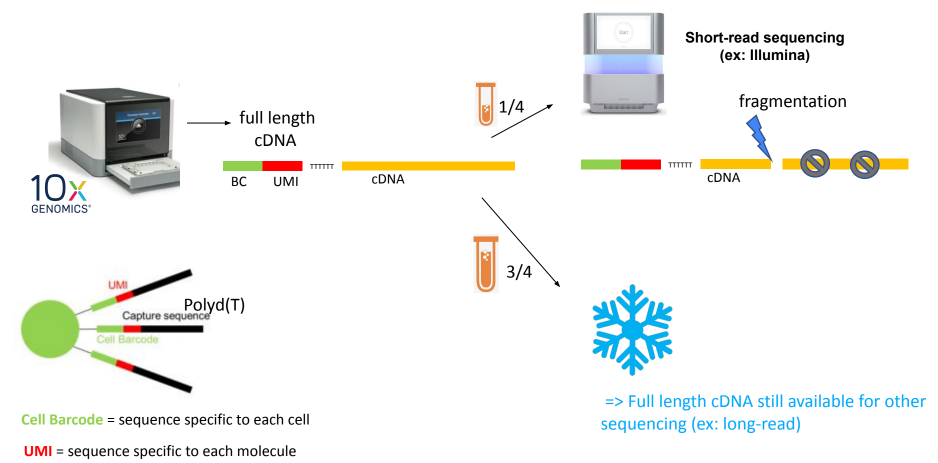
A molecule construction made of the barcode + UMI + polyd(T) + cDNA corresponding to the captured RNA

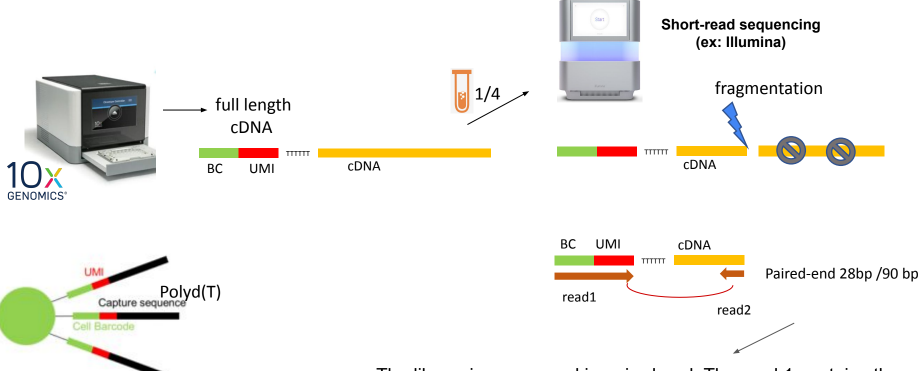
**Cell Barcode** = sequence specific to each cell

**UMI** = sequence specific to each molecule



**UMI** = sequence specific to each molecule



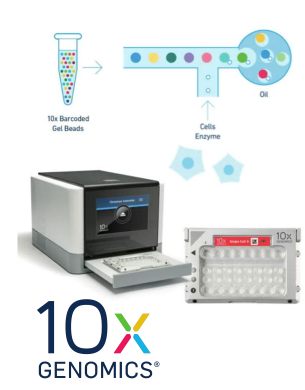


Cell Barcode = sequence specific to each cell UMI = sequence specific to each molecule The library is sequenced in paired-end. The read 1 contains the BC+UMI (28bp). The read2 contains a 90bp portion of cDNA. Only read2 corresponds to genomic/biological DNA. Read1 stems from synthetic molecules, not the transcriptome.

### 10X scRNA-seq in a nutshell

#### Tissue dissection + cell dissociation

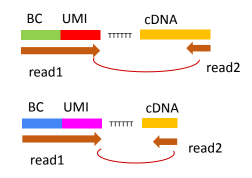
#### Cell partitioning + mRNA capture



#### Library preparation + sequencing



Paired-end 28bp /90 bp



### Biases/limitations of 10X Genomics technology

- Only the 3' end is sequenced (with short-read protocol + 3' kit)
- Max 10,000 cells (but millions on the newer device Chromium X)
- Cell size < 30um otherwise clog microfluidic channels
- 30% polyA+ transcripts captured per cell
- A droplet may contain 2 cells (= doublet)
- Some cell sub-population may be completely depleted/unfound

### Considerations on experiments

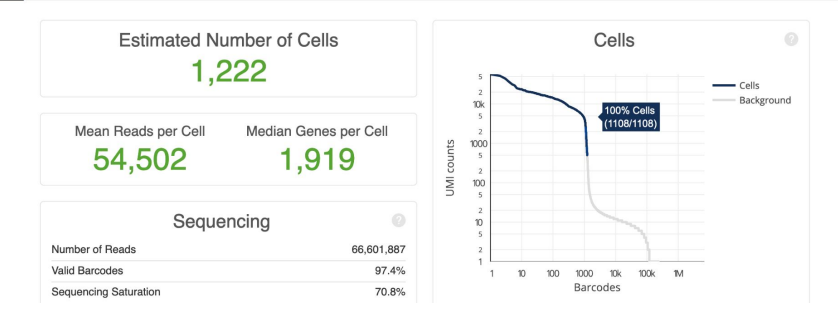
- Fresh cells : time between dissociation and 10X experiment should be <30min, otherwise cells start to die and result in RNAs wrongly assigned to cells (RNA "soup") and many expressed genes linked to cell death
- Frozen cells : does not work on all cells
- **FFPE** : only in human + mouse, restricted to certain tissues
- **Dissociation + Fixation** with ACME protocol (acetic acid + methanol + glycerol): requires optimisation but successful on exotic species (GenomiqueENS)
- Charge a bit more cells (25,000)
- Many tests have been done on **PBMCs** (immune cells) that are natively dissociated. Results do not necessarily reproduce on cells dissociated from tissues
- Q&A section of 10X website is very informative : https://kb.10xgenomics.com/hc/en-us/categories/360000149952-Single-Cell-Gene-E xpression

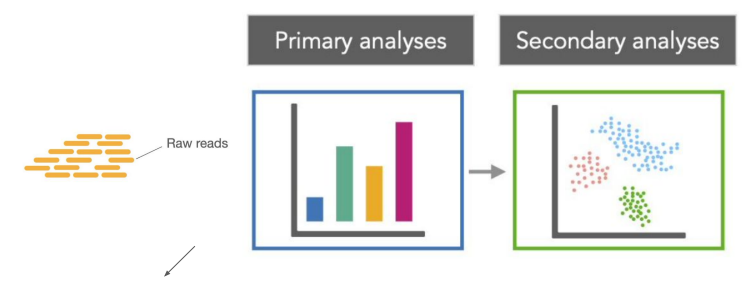


Which result file(s) did you obtain from the sequencing core facility ?

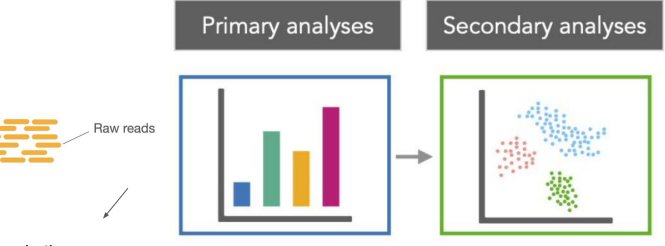
Cell Ranger · pbmc\_1k\_v3 · Peripheral blood mononuclear cells (PBMCs) from a healthy donor

SUMMARY ANALYSIS



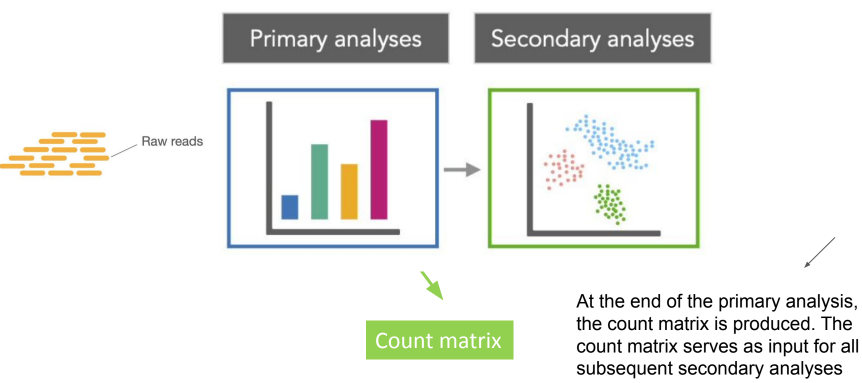


Raw data are the sequence reads. Then the bioinformatics analysis are in 2 phases : Primary (= preprocessing) and secondary

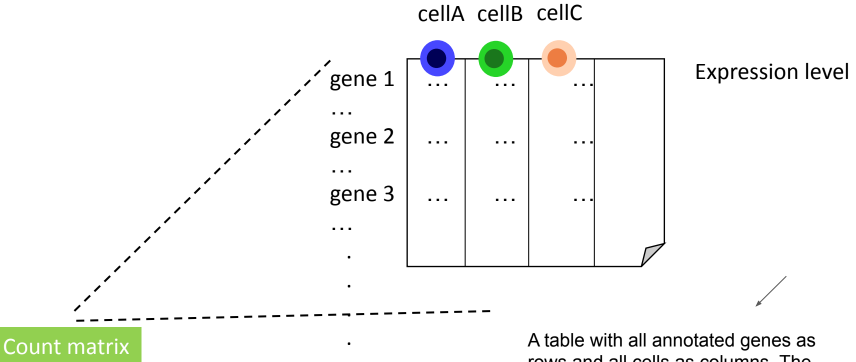


Cell Ranger is the program developed by 10X Genomics to perform the primary analysis (and a bit of secondary)



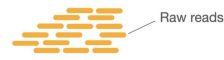


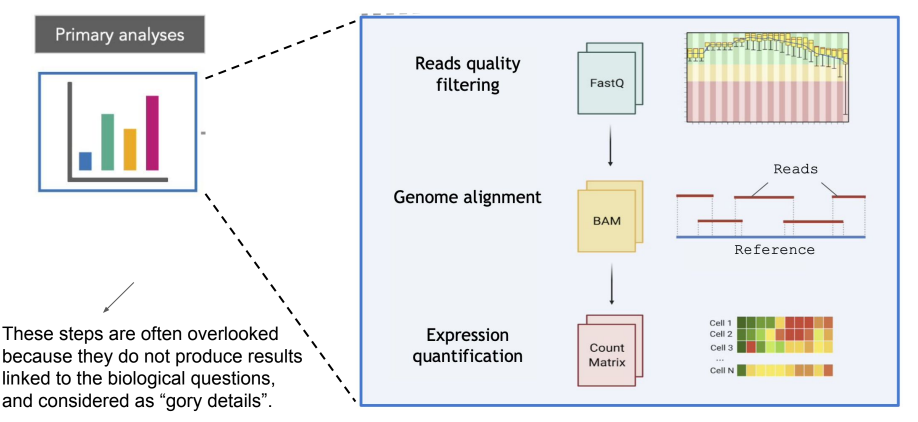
### What is a count matrix ?



A table with all annotated genes as rows and all cells as columns. The content of the table are expression levels (measured as read counts)

### The processing steps that are often overlooked





### Primary analyses : Reads





- Results starts by a BCL file (raw base calling from the sequencer). This file needs to be treated to produce the FASTQ files containing the reads
- This steps is done by the program bcl2fastq from Illumina (step "mkfastq" in CellRanger)

this is a detail, we just indicate it here in case you read further about Cell Ranger and step upon the notion of BCL

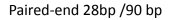
# Primary analyses : Reads quality checking

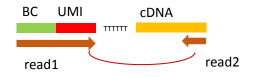




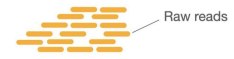
Primary analyses

- 2 FASTQ files :
  - one contains all the read1
  - one contains all the read2







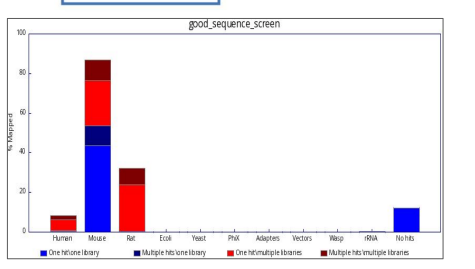


- Dataset name : pbmc\_1k\_v3 => 1000 human peripheral blood mononuclear cells (PBMCs) in human, freely available from 10X genomics website
- 2 files :
  - pbmc\_1k\_v3\_S1\_L001\_R1\_001.fastq.gz
  - pbmc\_1k\_v3\_S1\_L001\_R2\_001.fastq.gz

10X provides other public datasets, for each application, chemistry...

# Primary analyses : Reads quality checking





- As for any other NGS experiment, check the quality of the reads with FASTQC.
- FastqScreen enables to check for contaminations with other organisms
- These steps are usually done by the sequencing core facility, ask for these results if not provided





Thu 6 Oct 2022

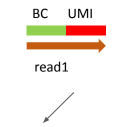
#### **Report**

#### Summary

Basic Statistics
Per base sequence quality
Per tile sequence quality
Per sequence quality scores
Per base sequence content
Per sequence GC content
Per base N content

#### Basic Statistics

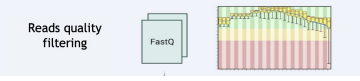
Measure	Value	
Filename	2022-006sc_S1_L001_R1_001.fastq.gz	
File type	Conventional base calls	
Encoding	Sanger / Illumina 1.9	
Total Sequences	494792037	
Sequences flagged as poor quality	0	
Sequence length	28	
%GC	49	



2022-006sc\_S1\_L001\_R1\_001.fastq.gz

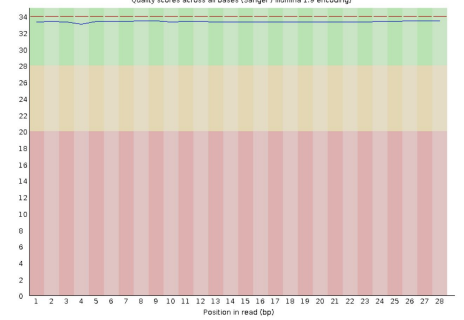
Read1 : 28bp 494M reads





#### Per base sequence quality

Quality scores across all bases (Sanger / Illumina 1.9 encoding)



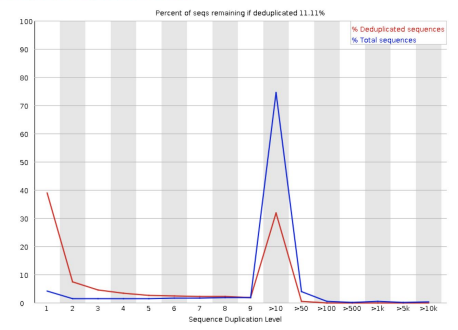


quality is excellent





#### **Osequence Duplication Levels**



normal to have duplication level because some BC+UMI have amplification biaises





# Basic Statistics

Measure	Value	
Filename	2022-006sc_S1_L001_R2_001.fastq.gz	
File type	Conventional base calls	
Encoding	Sanger / Illumina 1.9	
Total Sequences	494792037	
Sequences flagged as poor quality	0	
Sequence length	90	
%GC	46	



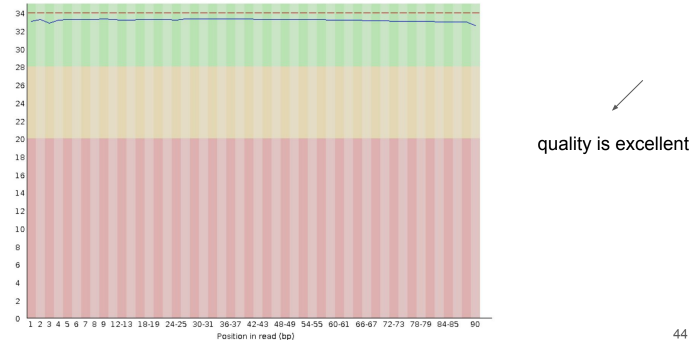
Read2 : 90bp 494M reads



**Reads quality** filtering FastQ

#### Per base sequence quality

Quality scores across all bases (Sanger / Illumina 1.9 encoding)



44



In CellRanger report



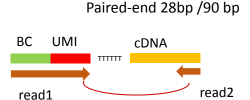
Sequencing	
Number of Reads	66,601,887
Valid Barcodes	97.4%
Sequencing Saturation	70.8%
Q30 Bases in Barcode	94.1%
Q30 Bases in RNA Read	90.2%
Q30 Bases in Sample Index	91.1%
Q30 Bases in UMI	92.7%

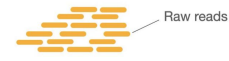
check the "sequencing" section of the report. The Q30 means "very high quality of bases"

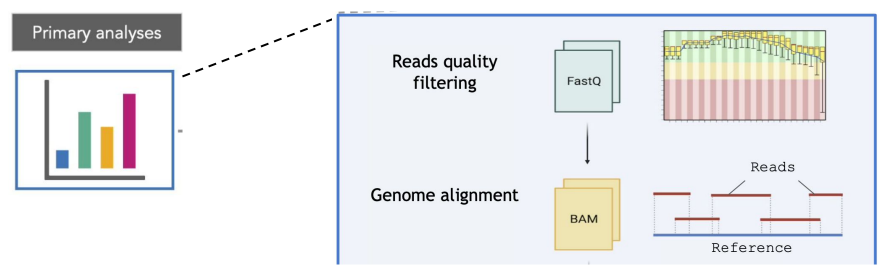
## Primary analyses : Reads quality checking



- Make sure read1 is of high quality because it contains the BC + UMI, later used to trace back the cell from which originates the RNA
- Ns and highly repeated sequences would impair read assignment.
- Any wrong base => lost read and barcode

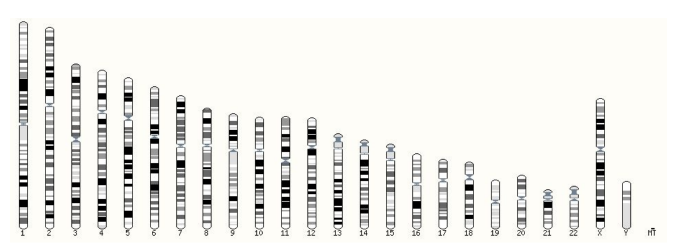


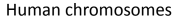




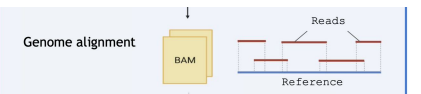
- Read1 and Read2 are then treated separately
- Read2 corresponds to genomic sequence => mapping step (=infer the position on the genome from which the read originates)



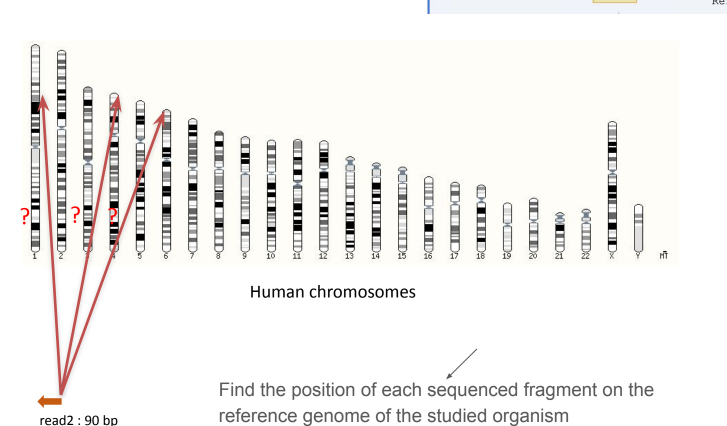


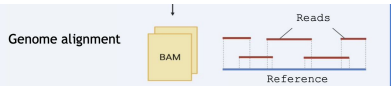


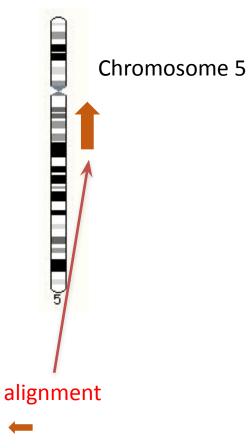


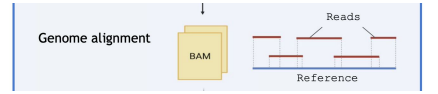


read2 : 90 bp



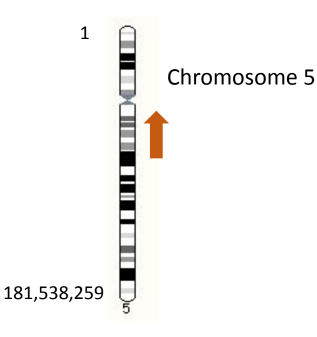


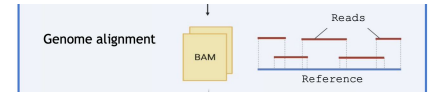




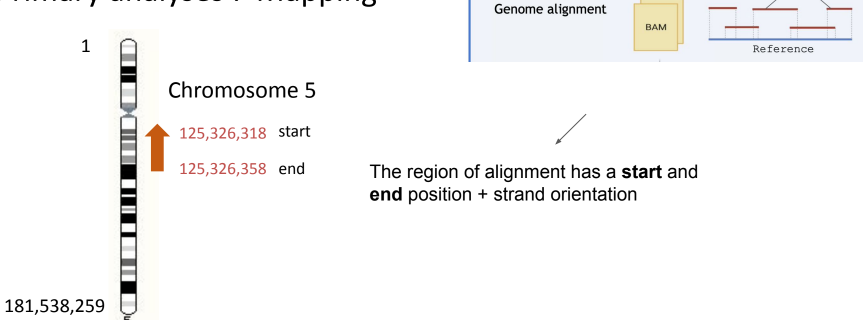
The best alignment is found for this read over the whole genome. Here it is on chromosome 5



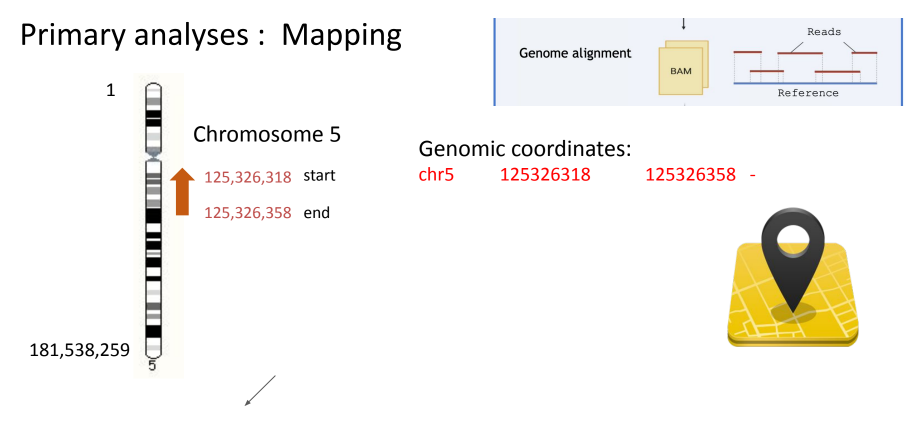




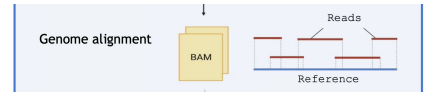
To precise the localisation of the best alignment, a coordinate system is used. First, each position of the chromosome has a particular value, corresponding to its distance from the beginning of the chromosome



Reads



The genomic coordinates is like "GPS coordinates" to locate regions on a genome. The format is : chromosome start end strand



- The mapping step enables to obtain the genomic coordinates of all reads2 for which an alignment has been found.
- The output file is in **BAM** format
- Not all reads can be aligned (contaminations, differences between the sample and reference genome, ...)
- Programs that perform this mapping step are often called "mappers"

read2

cDNA

55

sequences or genomes

requires more work because

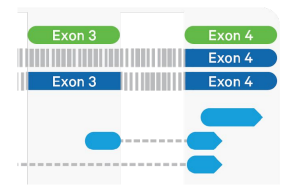
the genome index must be built (computer-intensive)

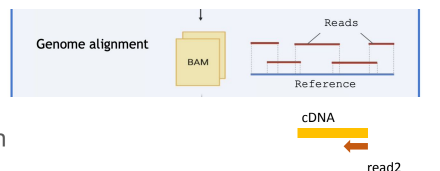
- CellRanger internally uses **STAR** as the program to align the reads on the reference genome
- The reference genome must be provided in the form of an index
- Ready-to-use genomes index:
  - human (hg19, GRCh38)
  - o mouse (mm10)
  - both (xenografts)
- For other organisms :
  - Use the genome in FASTA format
  - convert it with *cellranger mkgtf* and *cellranger mkref*.
- If you use some specific sequences (transgenes), don't forget to provide the sequence and rebuild the index ! (otherwise, no reads will be mapped to this region)

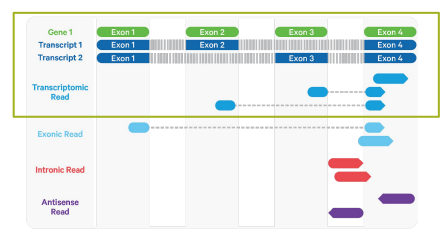




 STAR deals with RNA splicing, a read can be artificially "cut" to map to distant regions from which it originates (=different exons)

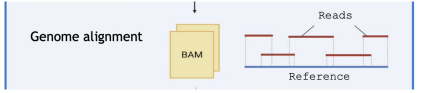






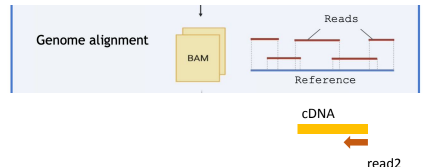


In CellRanger report



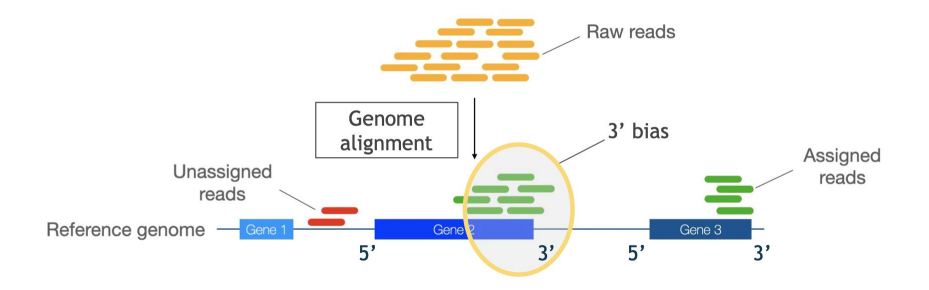
Mapping		
Reads Mapped to Genome	95.4%	
Reads Mapped Confidently to Genome	92.4%	
Reads Mapped Confidently to Intergenic Regions	4.8%	
Reads Mapped Confidently to Intronic Regions	31.1%	
Reads Mapped Confidently to Exonic Regions	56.5%	
Reads Mapped Confidently to Transcriptome	53.7%	
Reads Mapped Antisense to Gene	1.0%	

It is normal to have <100% reads aligned to the genome, because the reference genome is not exactly the genome of the studied sample. % will decrease with huge rearrangements (cancer or cell lines) or many SNPs (wild animals)

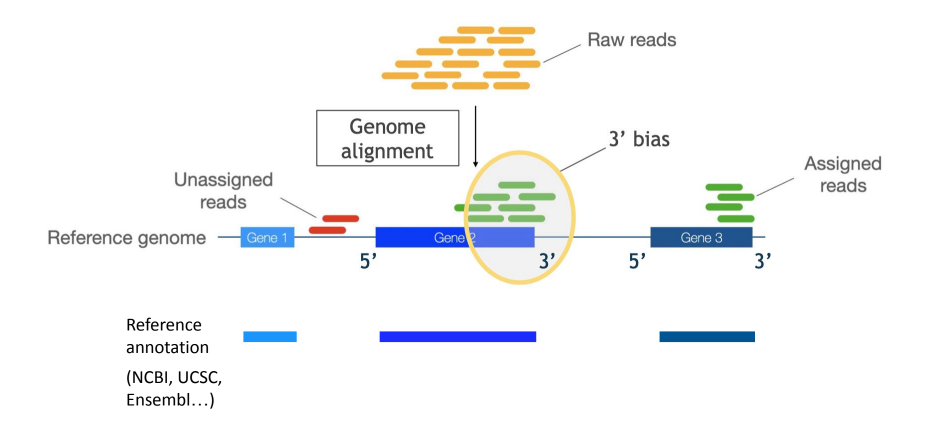


- Then, the **genome annotation** is used to assign the reads to genes
- Annotation is provided by genome portals (NCBI, Ensembl, UCSC) or consortiums of researchers working on a same organism
- genome annotation is generally provided as a file in the format GFF or GTF

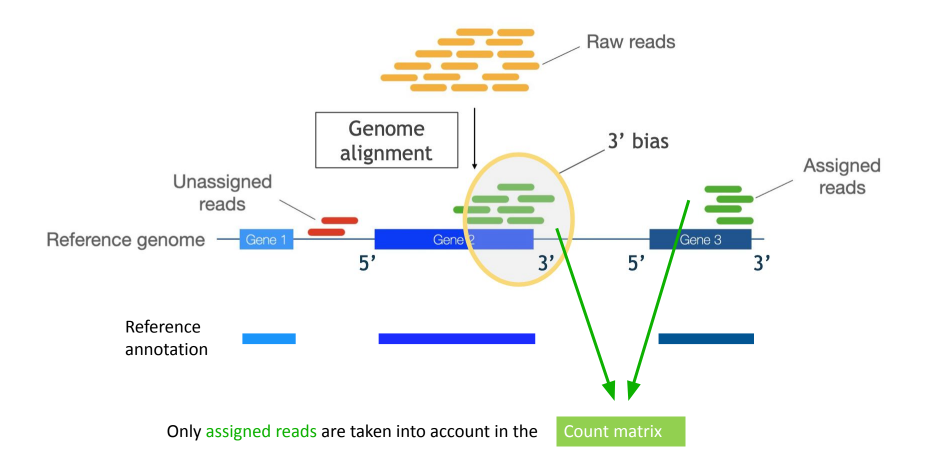
#### How is the genome reference annotation used ?



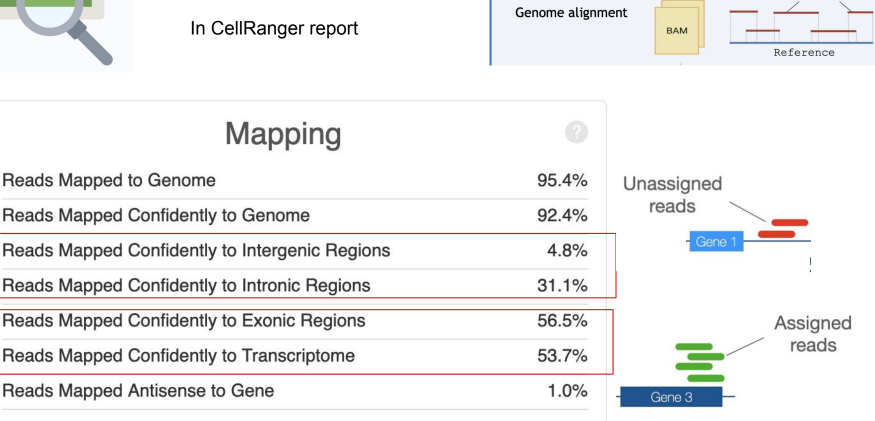
#### How is the genome reference annotation used ?



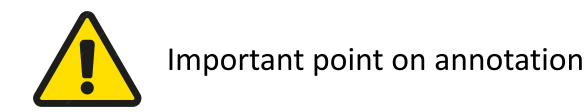
#### How is the genome reference annotation used ?







Reads



- Annotation is a crucial parameter (largely *underestimated*), as reads outside the annotated exons will not be taken into account !
- CellRanger will warn you on the report with the Alert below. In such cases, you <u>need</u> to visualise your signal in a genome browser (more on this tomorrow) and suspect the annotation may be problematic

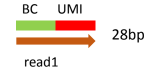
#### Alerts

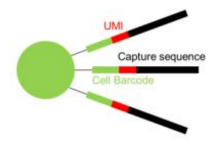
The analysis detected  $\triangle$  1 warning.

	Alert	Value	Detail
▲	Low Fraction Reads	51.5%	Ideal > 30%. This can indicate use of the wrong reference transcriptome, a reference transcriptome with
	Confidently Mapped To		overlapping genes, poor library quality, poor sequencing quality, or reads shorter than the recommended
	Transcriptome		minimum. Application performance may be affected.

# Primary analyses : barcode and UMI

- Read1 is made of BC + UMI
- Barcode and UMI are treated separately





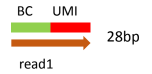
Reminder : barcode enables to trace back the read to the cell of origin ; UMI enables to distinguish each individual molecule

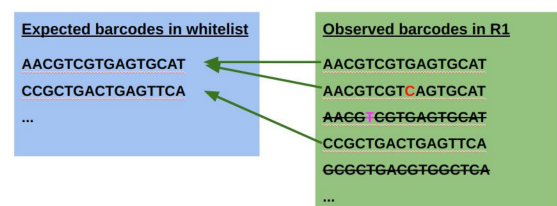
**Cell Barcode (16bp)** = sequence specific to each bead (so each cell)

UMI (12 bp) = sequence specific to each molecule : Unique Molecular Identifier

# Primary analyses : barcode

- Barcode is extracted (16bp)
- 10X provides a whitelist containing all possible barcodes used on the gel beads (~3 million barcodes for the v3 chemistry)
- All barcodes are compared to this whitelist
- **Correction**: barcodes with 1 difference (1 mismatch) from the whitelist are corrected.
- **Filtering**: keep only BC in the whitelist.







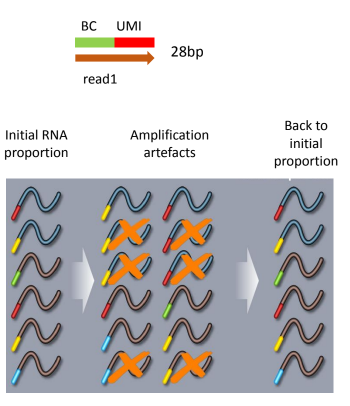
In CellRanger report

Sequencing	
Number of Reads	66,601,887
Valid Barcodes	97.4%
Sequencing Saturation	70.8%
Q30 Bases in Barcode	94.1%
Q30 Bases in RNA Read	90.2%
Q30 Bases in Sample Index	91.1%
Q30 Bases in UMI	92.7%

% of valid barcodes is indicated in the report

# Primary analyses : UMI

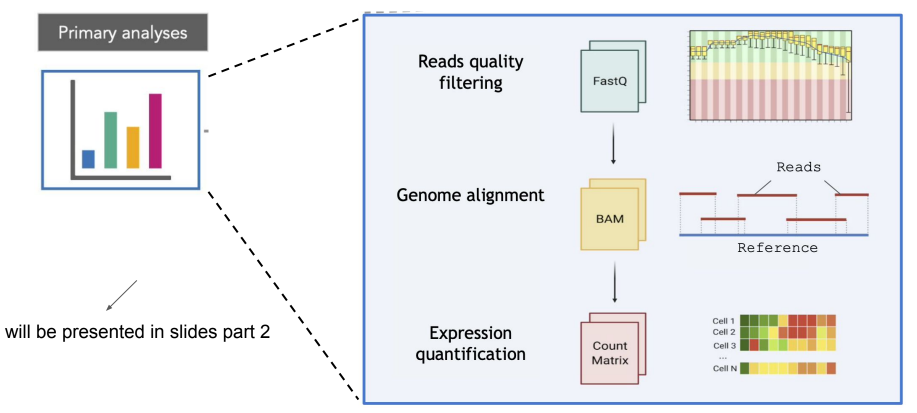
- UMI is extracted (12bp)
- UMI are randomised sequences, there is no whitelist
- **Correction**: UMI with 1 difference (1 mismatch) from a higher-count UMI are corrected to the higher count UMI if they share a cell barcode.
- Filtering: remove incorrect UMIs:
  - homopolymers (e.g. AAAAAAAAAA)
  - Contains 1 or several N
  - contains any base with BASEQ < 10



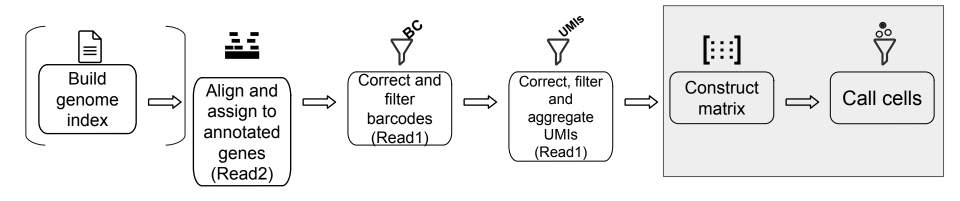
UMI aims at correcting amplification artefacts (more details tomorrow)

#### Last step : generation of the count matrix





### Overview of the workflow for primary analysis



# Take-home messages

- **Primary analysis is important !** If this step has issues, the resulting count matrix will have issues that will be propagated to all downstream analyses
- These steps are often overlooked
- **Cell Ranger** : program provided by 10X Genomics that perform primary analysis (and a bit more). Cell Ranger is reliable but it is necessary to understand what it does and its limits
- You will hear that "the raw data is the count matrix" => this is wrong, remember the raw data are the reads
- Only **read2 is mapped** to the genome ; read1 is synthetic Barcode+UMI
- Alternative ways to perform primary analysis exist

## Acknowledgements

 Some illustrations were created by Nathalie Lehmann and Rémi Montagne