

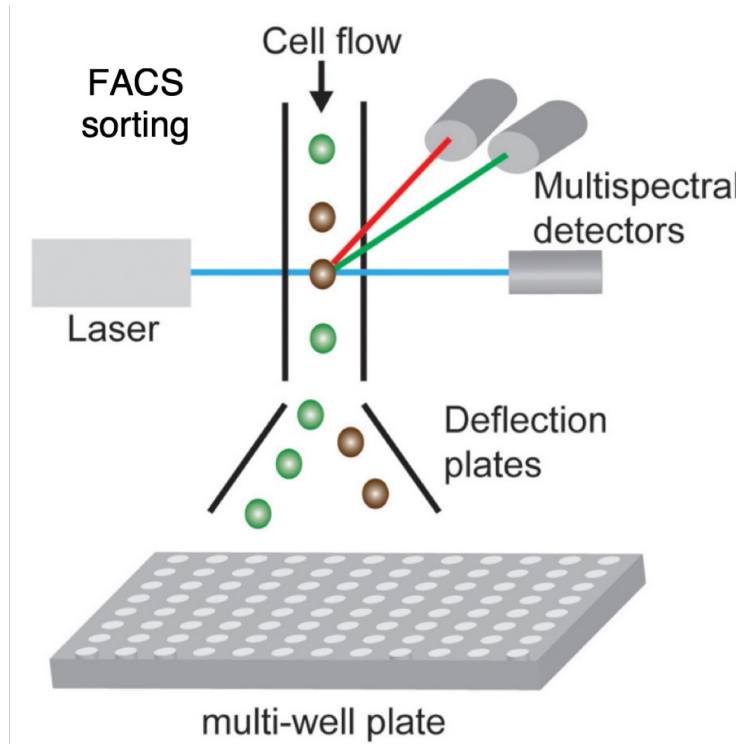
Introduction to Cell Ranger

Single Cell Transcriptomics in Python

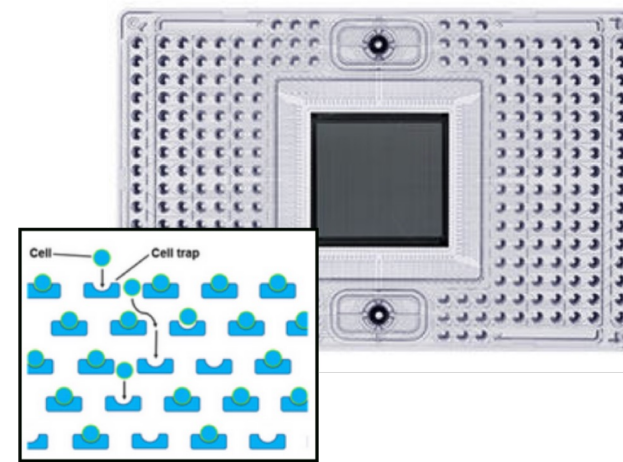
Alex Lederer

General overview of scRNA-seq technologies

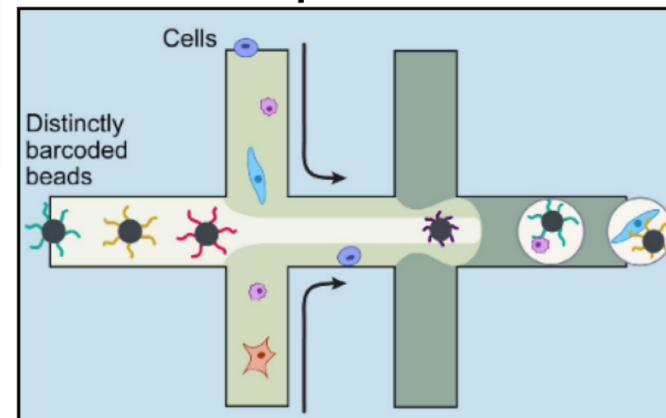
Microtitre Plates



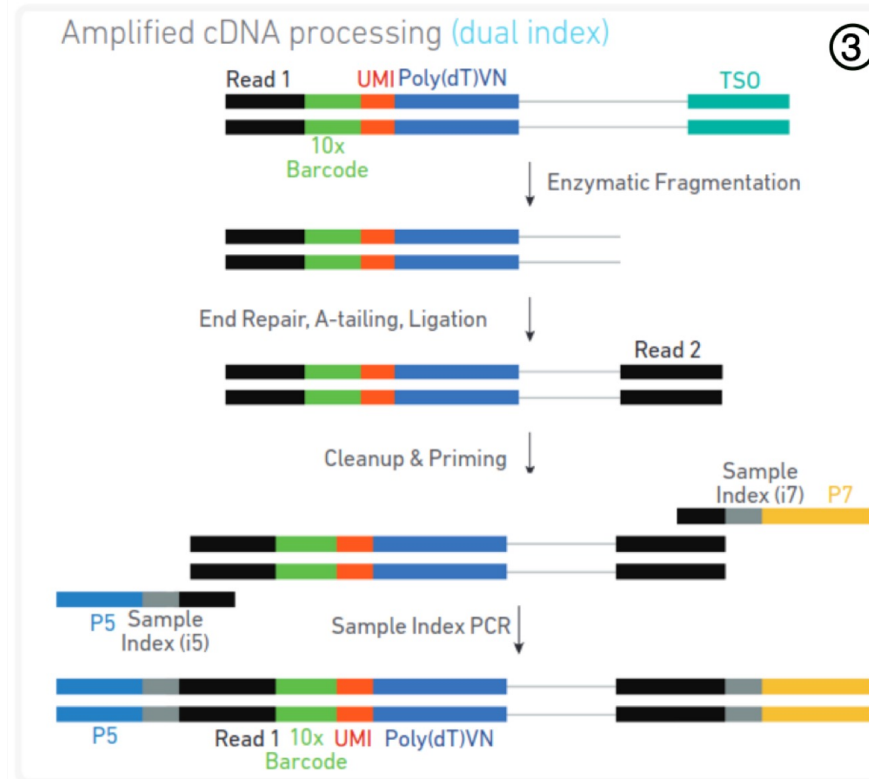
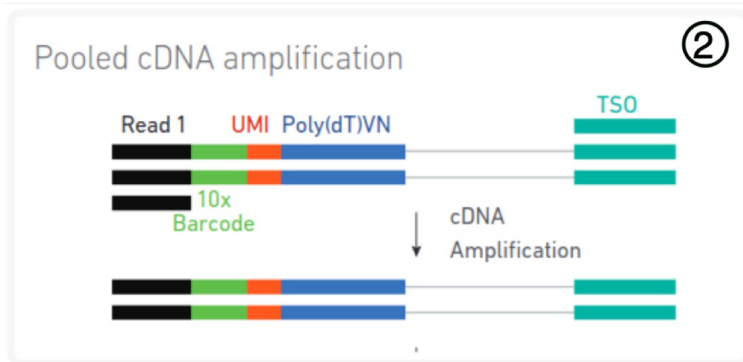
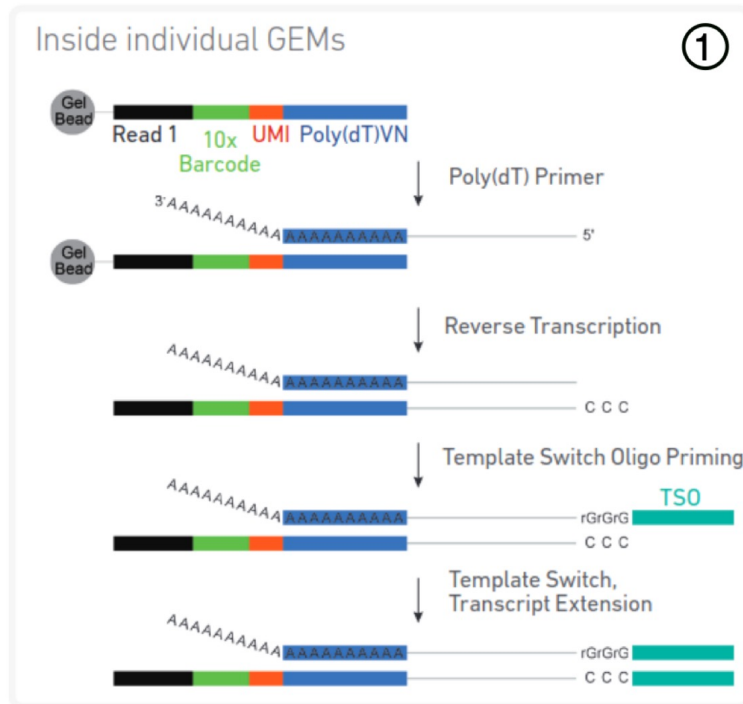
Microfluidic Arrays



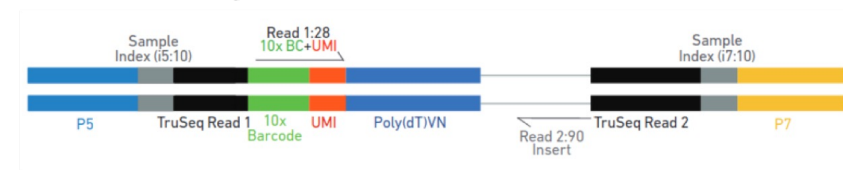
Microfluidic Droplets



10X Genomics Single-Cell RNA-sequencing protocol overview



Final library



Question

5k_pbmc_protein_v3 - 5k Peripheral blood mononuclear cells (PBMCs) from a healthy donor

Summary [Analysis](#)

5,247

Estimated Number of Cells

28,918

Mean Reads per Cell

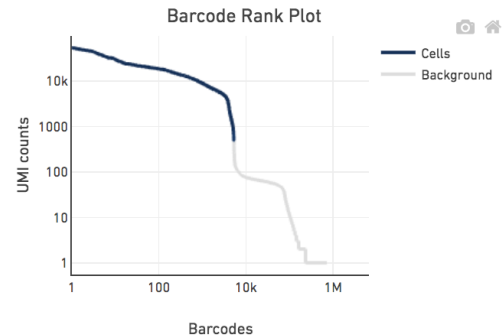
Sequencing [?]

Number of Reads	151,731,342
Valid Barcodes	97.5%
Valid UMIs	99.9%
Sequencing Saturation	52.4%
Q30 Bases in Barcode	95.8%
Q30 Bases in RNA Read	91.9%
Q30 Bases in Sample Index	89.8%
Q30 Bases in UMI	95.4%

Mapping [?]

Reads Mapped to Genome	94.3%
Reads Mapped Confidently to Genome	88.4%
Reads Mapped Confidently to Intergenic Regions	6.8%
Reads Mapped Confidently to Intronic Regions	25.0%
Reads Mapped Confidently to Exonic Regions	56.7%
Reads Mapped Confidently to Transcriptome	53.2%
Reads Mapped Antisense to Gene	1.3%

Cells [?]



Estimated Number of Cells	5,247
Fraction Reads in Cells	87.7%
Mean Reads per Cell	28,918
Median Genes per Cell	1,644
Total Genes Detected	20,822
Median UMI Counts per Cell	5,496

Sample

Sample ID	5k_pbmc_protein_v3
Sample Description	5k Peripheral blood mononuclear cells (PBMCs) from a healthy donor
Chemistry	Single Cell 3' v3
Transcriptome	GRCh38-3.0.0
Pipeline Version	3.1.0

How do I know if my sample is any good?

Sample Preparation

- Cell isolation and tissue dissociation
- **Potential issues:** *What if the tissue is not fully dissociated? Or agitated too much?*

Cell Encapsulation

- Gel Bead-In-Emulsions (GEMs) Formation: cells are encapsulated in droplets with gel beads coated with **barcoded cell-specific primers** and **molecule-specific UMIs**
- **Potential issues:** multiple cells per barcode, droplets without any cells, ambient RNA

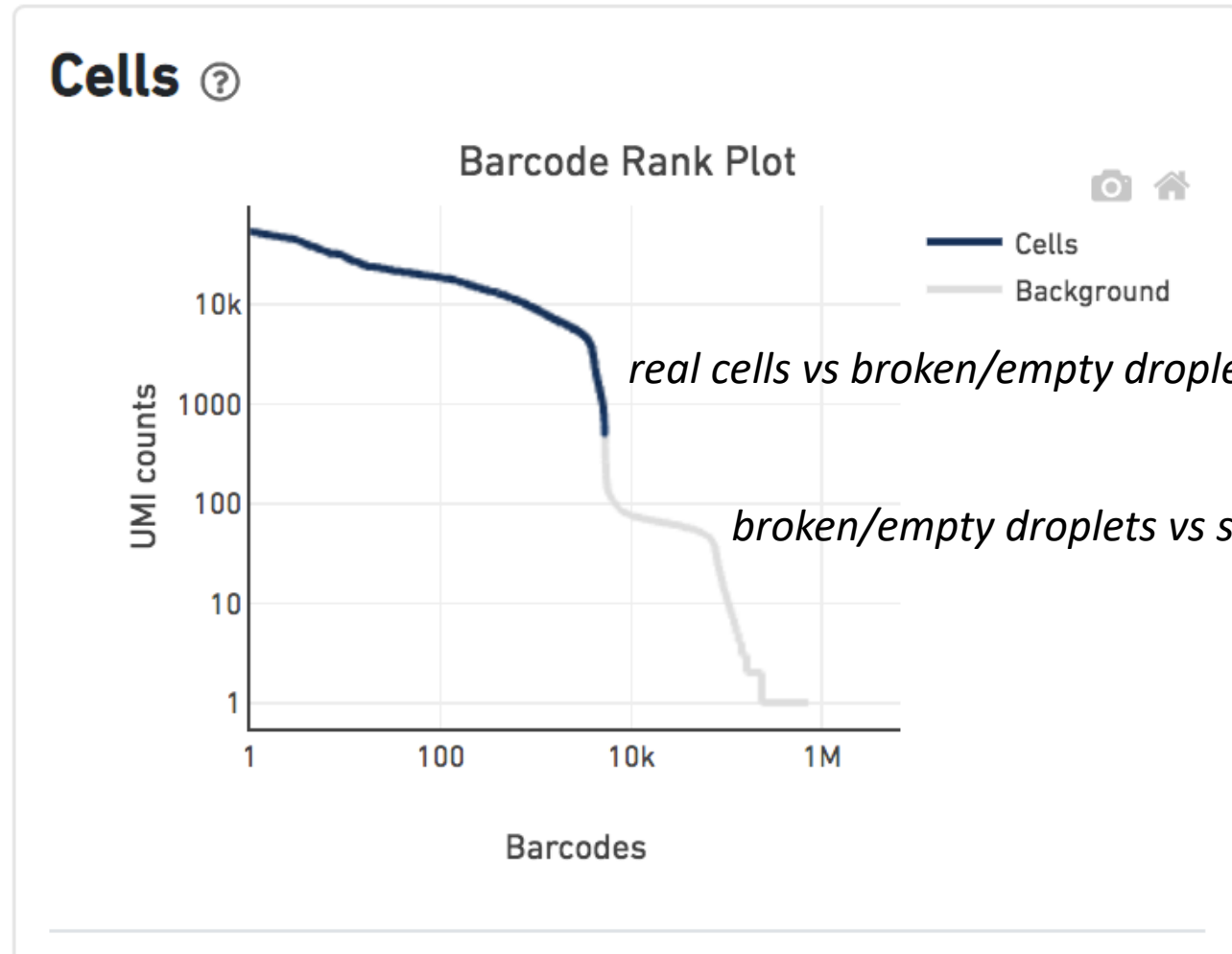
Reverse Transcription

- Converts mRNA into cDNA within each droplet.
- **Potential issues:** degradation of unstable RNA molecules

Post-Emulsion Breakage Processing

- Emulsion Breakage and cDNA Cleanup: The emulsions are broken, and the pooled cDNA is cleaned up, usually using magnetic beads. The cleaned-up cDNA is PCR-amplified to generate sufficient material for library preparation (PCA amplification) and sequencing.
- **Potential issues:** incorrect size selection with magnetic beads; amplification of primers or other small DNA fragments

The barcode rank plot



Question