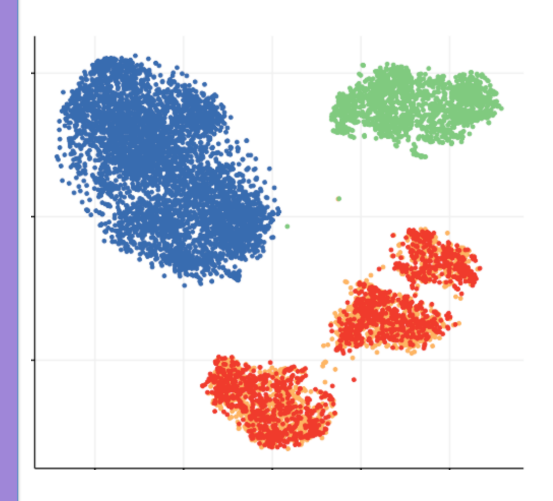
Introduction to single cell RNA sequencing

CRUK Bioinformatics Summer School 2018

> Mike Morgan Comp Bio Postdoc Marioni Group





Why study single cells?

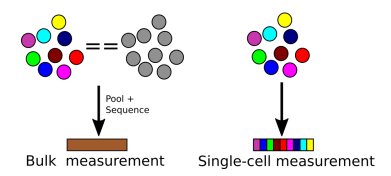
Unravel tissue heterogeneity:

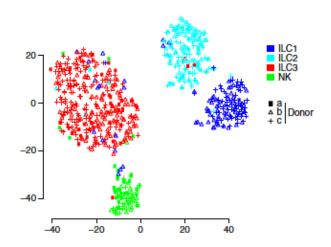
- Novel and rare cell types
- Unknown cellular states

Transcriptional dynamics

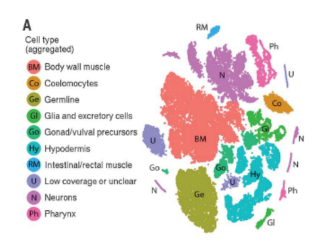
Can also measure single-cell:

- Chromatin accessibility
- Mutation & CNV (scDNAseq)
- Methylation

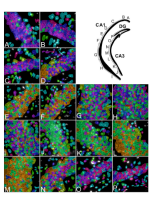




Innate-lymphoid cells
Bjorklund *et al.*, Nature Immunology (2016)



Whole *C. elegans* larva Cao *et al.*, Science (2017)



Mouse hippocampus Shah *et al.*, Neuron (2017)



How can we study single cells?

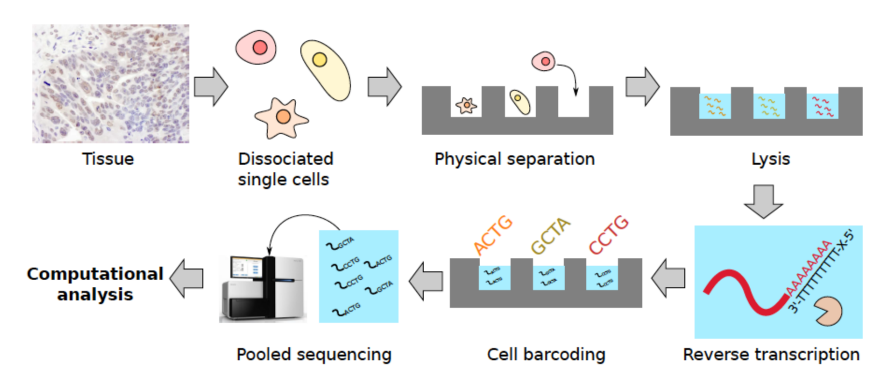
Technology	Measurements (P)	Cells (N)	Throughput	Pro	Con
Flow cytometry	1-15	1k-100k	big N, small P	Technically easy	Limited targets
Mass cytometry	20-50	1k-100k	big N, medium P	>P than flow	Limited targets
RNA FISH	1	~100	small N, small P	Spatial resolution	Technically hard, low throughput
Multiplex FISH	~100	100's	medium N, medium P	Spatial resolution	Technically and analytically hard
SS2 scRNA-seq	~20,000	100-1000	medium N, big P	High throughput	Sparse, low input material
Droplet scRNA-seq	~20,000	100-1M	big N, big P	High throughput	Very sparse, low input material

NB – every method has it's pros and cons. There is no all-encompassing single cell methodology.

It depends on your biological question!



A typical scRNA-seq experiment



- Dissociation can be easy (blood) or hard (collagenous tissue)
- Separation and RT differ by protocol

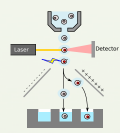
Physical separation defines main scRNA-seq protocols

Microfluidic device



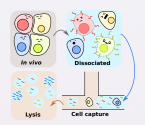
- 96 or 800 well format
- Physically check presence of cells
- High capture efficiency
- Doublet issues
- Expensive
- Full-length cDNA (SMART-seq{2})
- Spike-in control RNA
- High gene coverage

Plate-based



- 96 or 384 well format
- Sort specific population(s) of cells
- High capture efficiency
- Experimental design considerations
- Full-length cDNA (SMART-seq(2) or endtagging; UMIs)
- Spike-in control RNA
- High gene coverage

Droplet-based

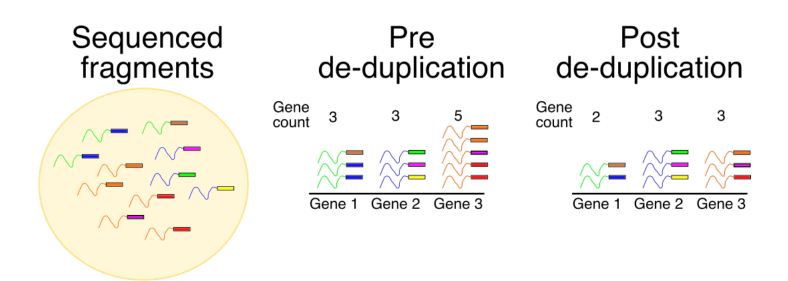


- 100-1000's of cells
- Doublet issues
- Variable capture efficiency
- Low per-cell cost
- 3' end tag; UMIs
- No spike-in control RNA
- High cell coverage

What are UMIs?

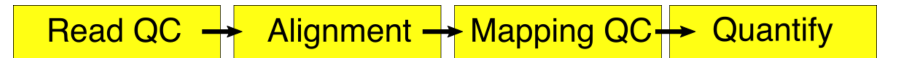
<u>Unique molecular identifiers give (almost) exact molecule counts in sequencing experiments.</u>

They reduce the amplification noise by allowing (almost) complete de-duplication of sequenced fragments.





A typical SMART-seq workflow

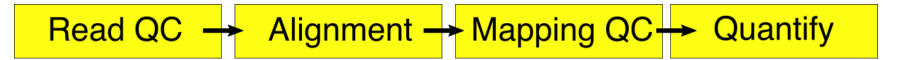


The same tools used for bulk RNA-seq, e.g. FastQC, Star, PicardTools (Deduplication is essential)

Typically 1 library per cell, potentially many 100's of FASTQ files Need to be able to handle many files in parallel – e.g. high performance computing cluster.

Pipelining tools exist (beyond the scope of this tutorial – see resources).

A typical SMART-seq workflow

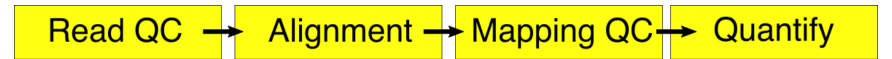


The same tools used for bulk RNA-seq, e.g. FastQC, Star, PicardTools (Deduplication is essential)



Single-cell specific tools (generally performed in R; Practical 1)

A typical SMART-seq workflow



The same tools used for bulk RNA-seq, e.g. FastQC, Star, PicardTools (Deduplication is essential)



Single-cell specific tools (generally performed in R; Practical 1)



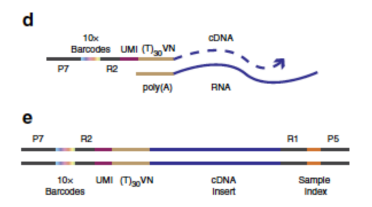


A typical droplet workflow

Droplet-based methods create a new problem, and solution:

- Many 100's-1000's cells == 1000's small FASTQ files
- Prohibitively expensive to sequence 20,000 cells to 1M reads

Solution: multiplex cells using barcodes

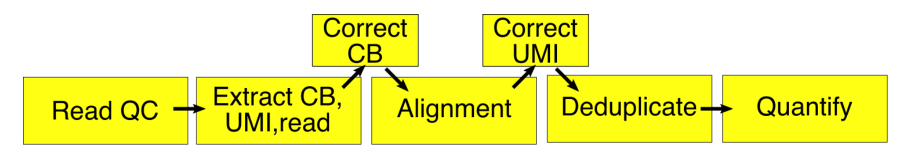


A single 10X Genomics Chromium library generates 3 FASTQ files: R1, R2, Index

10X Genomics Chromium v1 chemistry design Zheng *et al.*, Nature Comms (2017)



A typical droplet workflow



Generally run in a single pipeline, e.g. Cellranger (10X specific), DropSeq (Macosko *et al.*) or custom (not recommended if just starting).

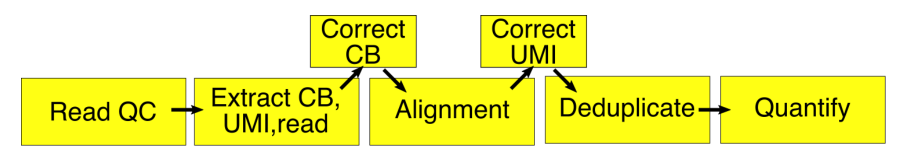
Sequencing errors in cell barcodes and UMIs are a source of technical noise – must be dealt with

Recent development: Rob Patro & co have a new end-to-end (i.e. FASTQ to counts matrix) lightweight pipeline:

https://salmon.readthedocs.io/en/latest/alevin.html



A typical droplet workflow



Generally run in a single pipeline, e.g. Cellranger (10X specific), DropSeq (Macosko *et al.*) or custom (not recommended if just starting).



Single-cell specific tools (generally performed in R; Practical 1)





Dealing with single cells

Regardless of technology, our goal is to derive/extract real biology from technically noisy data.



Single cell analysis workflow

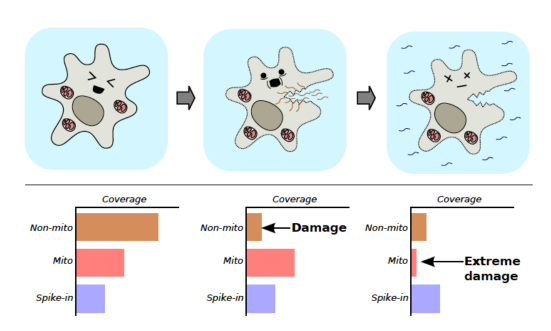
Starting with a counts matrix:

- Quality control
- Normalization
- Batch correction [if required]
- Dimensionality reduction and visualisation (part 2)
- Clustering (part 2)
- Differential expression testing (same as bulk RNA seq... mostly)

Cell	Α	В	C	D
Gene X	10	20	30	40
Gene Y	15	30	45	60
Gene Z	20	40	60	80

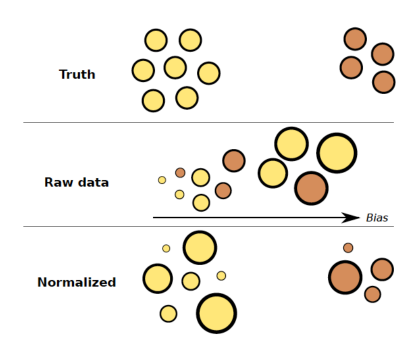
Quality control on cells

- Low sequencing depth
- Low numbers of expressed genes (i.e. any non-zero count)
- High spike-in (if present) or mitochondial content



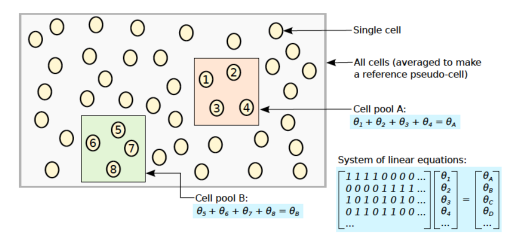
Normalization

- The aim is bring all cells onto the same distribution to remove biases between them
- We want to preserve biological variability, not introduce new technical variation
- Primary source of bias is sequencing depth – scale down counts accordingly
- Need a method that is robust to sparsity and composition bias
 - TMM & DESeq size factors are not!



Normalization by deconvolution

- Estimate cell-specific size factors.
- Handles sparsity and is robust to DE.



Lun et al., Genome Biology (2016)

- 1. Cluster cells together
- 2. Pool cells to increase counts, reduce 0's
- 3. Robust estimate of each pool size factor
- 4. Wash & repeat for multiple pools
- 5. Solve the linear system of equations to obtain *per-cell* size factors

Confounders and batch correction

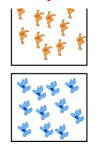
- A segue into proper experimental design
- Some batch effects cannot be avoided
- Some can, make sure you know which is which

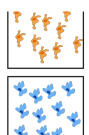
Confounded design

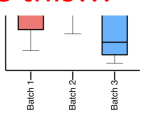
Biological

Processing

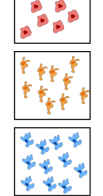
Please don't design your experiment like this!!!

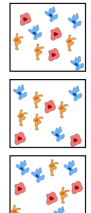


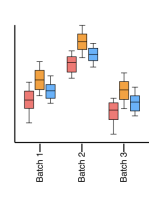




Not confounded design Biological Processing groups groups







What if I still have batch effects?

Good experimental design doesn't remove batch effects, it prevents them from biasing your results (hopefully)

If you still have batch effects then they can be dealt with (if necessary) <- important for clustering and visualization



Simple batch correction

If you have a single cell type and multiple conditions:

Use a linear model to regress gene expression on batch

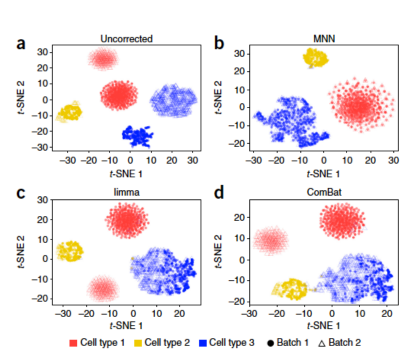


More complex batch correction

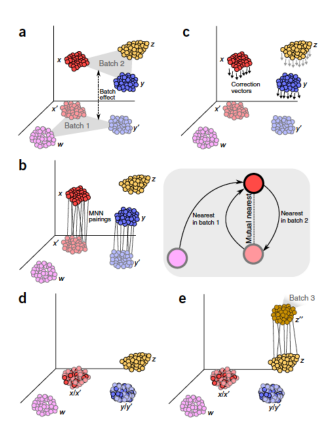
Linear models (and bulk batch correction methods) can't handle composition differences between batches.

Need a method that handles multiple batches, i.e. > 2, and corrects expression values properly

Match cells between batches that share the same biological subspace, remove the orthogonal components (mnnCorrect).









Pseudotime



Resources

Single Cell Resources:

Single cell course (Hemberg Lab; Wellcome Sanger Institute):

http://hemberg-lab.github.io/scRNA.seq.course/index.html

Aaron Lun's single cell workflow (very detailed):

https://www.bioconductor.org/packages/release/workflows/html/simpleSingleCell.html

Cellranger pipeline:

https://support.10xgenomics.com/single-cell-gene-expression/software/pipelines/latest/what-is-cell-ranger

Resources

Workflow Resources:

Snakemake (Python):

http://snakemake.readthedocs.io/en/stable/#

Nextflow (Java/agnostic):

https://www.nextflow.io

Ruffus (Python):

http://www.ruffus.org.uk

make (bash):

https://www.tutorialspoint.com/unix commands/make.htm

Recommended reading

Study design

Hicks et al., bioRxiv (2015):

https://www.biorxiv.org/content/biorxiv/early/2015/08/25/025528.full.pdf

Batch correction:

Haghverdi et al, Nature Biotech (2018):

https://www.nature.com/articles/nbt.4091

Butler et al., Nature Biotech (2018):

https://www.nature.com/articles/nbt.4096

