

Single-cell RNA-seq data analysis using Chipster

January 2024

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CSC – Suomalainen tutkimuksen, koulutuksen, kulttuurin ja julkishallinnon ICT-osaamiskeskus

What will I learn?

- Analysis of single-cell RNA-seq (scRNA-seq) data
 - Find subpopulations (clusters) of cells and marker genes for them
 - Compare multiple samples (e.g. treatment vs control)
 - Identify cell types that are present in both samples
 - Obtain cell type markers that are conserved in both samples
 - Compare the samples to find cell-type specific responses to treatment
- Note: you can use the same analysis pipeline for single-nuclei RNA-seq (snRNA-seq)
- How to operate the Chipster software



Introduction to Chipster

Chipster

- User-friendly analysis software for high-throughput data
- Provides an easy access to over 500 analysis tools
- Command line tools
- R/Bioconductor packages
- Free, open source software
- What can I do with Chipster?
 - analyze high-throughput data
 - visualize data efficiently
 - share analysis sessions

Chipster website (<https://chipster.csc.fi/>)

A 3D rendering of a DNA double helix, oriented diagonally from the top left to the bottom right of the page.

Chipster
Open source platform for data analysis

Welcome to Chipster

Chipster is a user-friendly analysis software for high-throughput data such as Visium, single-cell and bulk RNA-seq. Chipster provides a web interface to over 500 analysis tools, and the actual analysis jobs run on the server side making use of CSC's computing environment.

If you would like to use Chipster hosted by CSC, you need a [user account](#). Please note that Chipster is also available for [local server installation](#) free of charge.

 [Launch Chipster](#)

Training:

- 29.-30.5.2023 [Single-cell RNA-seq data analysis](#)
- 25.10.2022 [Spatial transcriptomics \(Visium\) data analysis](#)
- 30.6.2021 [MOOC Single-cell RNA-seq data analysis using Chipster](#), instructions on [how to get started](#)

News and resources:

- ASV-based microbial community analysis using DADA2: [Tutorial videos](#)
- [Analysis of QuantSeq 3' UMI RNA-seq data enabled](#)
- [Chipster introduction video](#)
- [Instructions for moving data from Puhti to Chipster](#)
- [Video on how to convert tables to Chipster format and create phenodata file](#)
- [Lecture videos of advanced single-cell RNA-seq data analysis course](#)

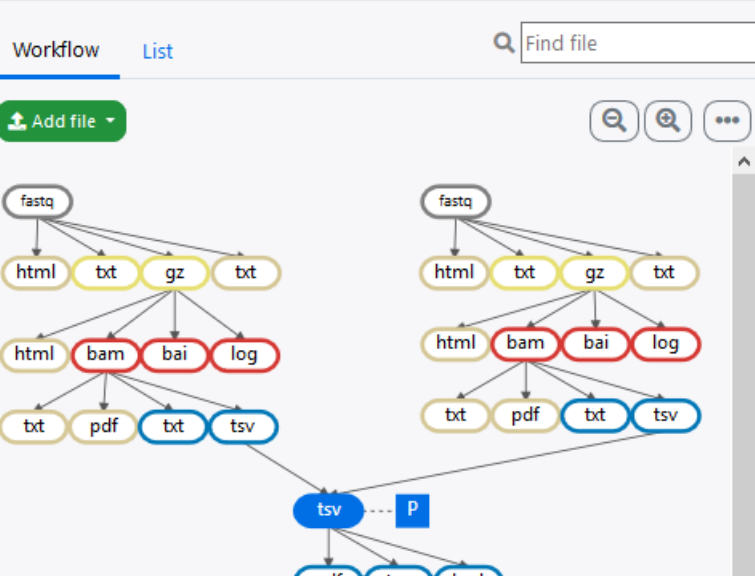
Chipster user interface (chipster.rahtiapp.fi)

Chipster Analyze Sessions Manual Contact ekorpela@csc.fi

Files

Workflow List Find file

Add file



Tools

NGS Microarray Misc Find tool

Category

- Quality control
- Preprocessing
- Utilities
- Matching sets of genomic regions
- Alignment
- Variants
- RNA-seq
- Small RNA-seq
- Single cell RNA-seq
- ChIP- and DNase-seq
- 16S rRNA sequencing
- CNA-seq

Tool

Read quality with FastQC Parameters Run

Read quality with MultiQC for many FASTQ files

Read quality statistics with FASTX

Read quality statistics with PRINSEQ

RNA-seq quality metrics with RseQC

RNA-seq strandedness inference and inner distance estimation using RseQC

Collect multiple metrics from BAM

PCA and heatmap of samples with DESeq2

Check FASTQ file for errors

Combine reports using MultiQC

Job 0

The tool runs FastQC on multiple FASTQ files, and then combines the reports using MultiQC. Input file is a single Tar package containing all the FASTQ files, which can be gzipped. This tool is based on the FastQC and MultiQC packages. [More info...](#)

File

ngs-data-table.tsv ...

Spreadsheet Text Expression Profile Scatter Plot Phenodata Details

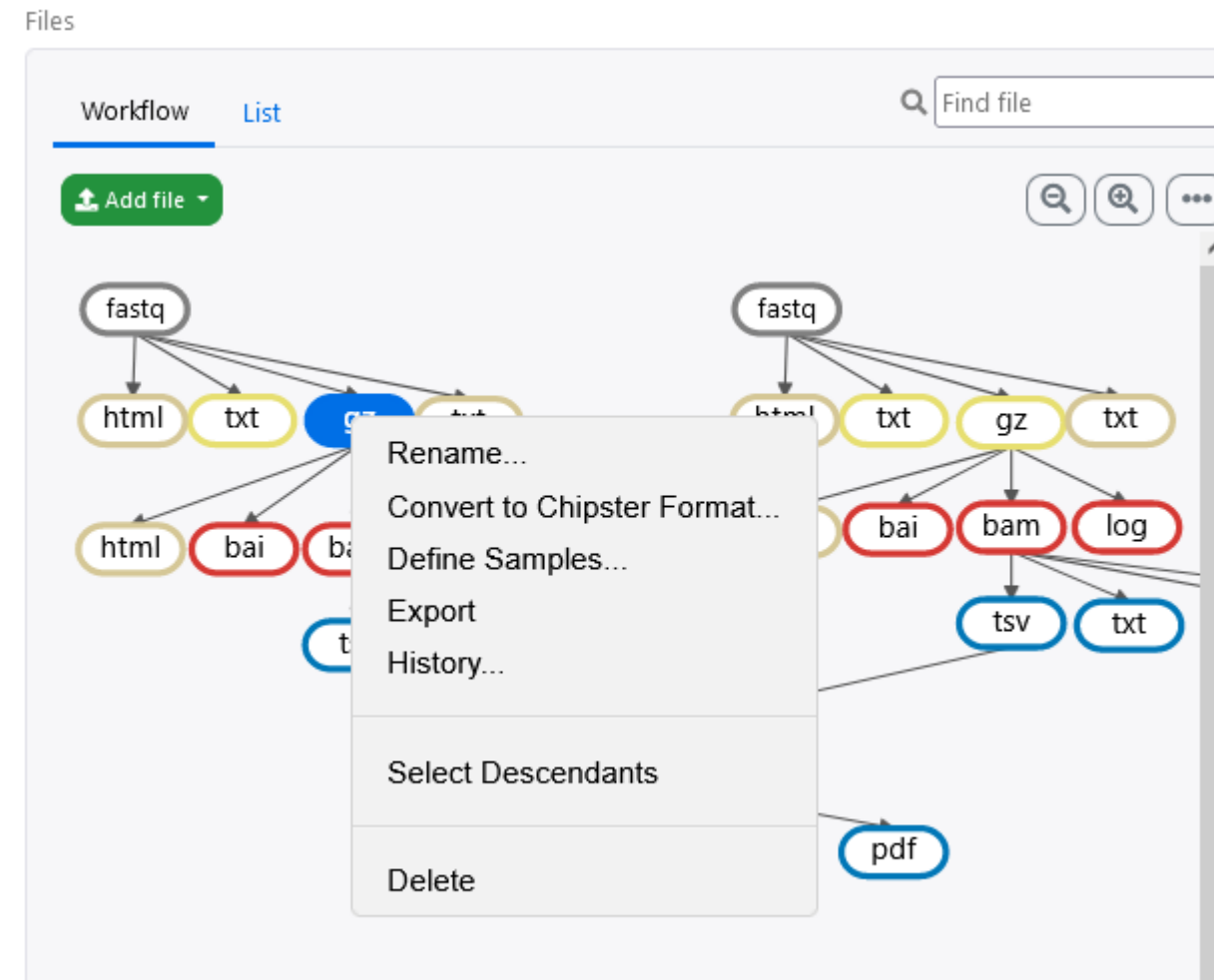
First 101 rows of 58396 [View in full screen](#) to see all the rows.

Full Screen

identifier	chr	start	end	length	sequence	chip.sample001.tsv	chip.sample002.tsv
ENSG000000000003	X	100627108	100639991	12883	NA	0	0
ENSG000000000005	X	100584801	100599885	15084	NA	0	0
ENSG00000000419	20	50934866	50958555	23689	NA	0	0

Workflow view

- Shows the relationships of the files
- You can move the boxes (files) around, and zoom in and out.
- Several files can be selected by
 - keeping the Ctrl/Cmd key down
 - drawing a box around them
- Right clicking a file allows you to
 - Download ("Export")
 - Delete
 - Rename
 - View history
 - Select descendants
 - Convert to Chipster format (for tables)
 - Define samples (for FASTQ files)



Options for importing data to Chipster



- Add file button
 - Upload files
 - Upload folder
 - Download from URL
- Sessions tab
 - Import session file
- Tools
 - Import from Illumina BaseSpace
 - Utilities / Retrieve data from Illumina BaseSpace
 - Access token needed
 - Import from SRA database
 - Utilities / Retrieve FASTQ or BAM files from SRA
 - Import from Ensembl database
 - Utilities / Retrieve data for a given organism in Ensembl
 - Import from URL
 - Utilities / Download file from URL directly to server

Analysis sessions

- Your analysis is saved automatically in the cloud
 - Session includes all the files, their relationships and metadata (what tool and parameters were used to produce each file).
 - Session is a single .zip file.
 - Note that cloud sessions are not stored forever! Remember to download the session when ready.
- You can share sessions with other Chipster users
 - You can give either read-only or read-write access
- If your analysis job takes a long time, you don't need to keep Chipster open:
 - Wait that the data transfer to the server has completed
 - Close Chipster
 - Open Chipster later and the results will be there

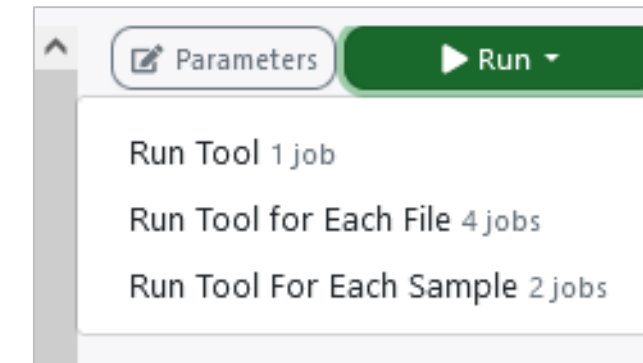
Running many analysis jobs at the same time



- You can have many analysis jobs running at the same time
 - No need to wait that one finishes before starting a new one

Run button gives several options:

- Run tool
 - Runs the selected analysis tool once
- Run tool for each file
 - Runs the selected analysis tool for each of the input files individually
- Run tool for each sample
 - If you have grouped paired end FASTQ files to samples using the Define samples –option, you can run the selected analysis tool for the input files in a sample specific manner.



Problems? Send us a support request

-request includes the error message and link to analysis session (optional)



Chipster Analyze Sessions Manual Contact ekorpela@csc.fi

Contact support

In case something doesn't work, please contact support.

[Contact support](#)

Contact information

If you have questions about using Chipster, you can send or view messages. For more information, see the [Chipster User's Guide](#).

chipster-users@lists.sourceforge.net
General list for Chipster users.
[Send message](#) | [View message](#)
chipster-tech@lists.sourceforge.net
Technical list for people installing and maintaining Chipster.
[Send message](#) | [View message](#)
chipster-announcements@lists.sourceforge.net
A very low traffic list for announcements about new versions etc. Only project administrators can post.
[View messages](#) | [Subscribe](#)

Contact support

Message

Please describe what happened

Attach session

Attach a copy of your last session **NGS_RNAseq_fromReadsToDifferentiallyExpressedGenes_ENCODE_2samples**
 Don't attach the session

Your email address

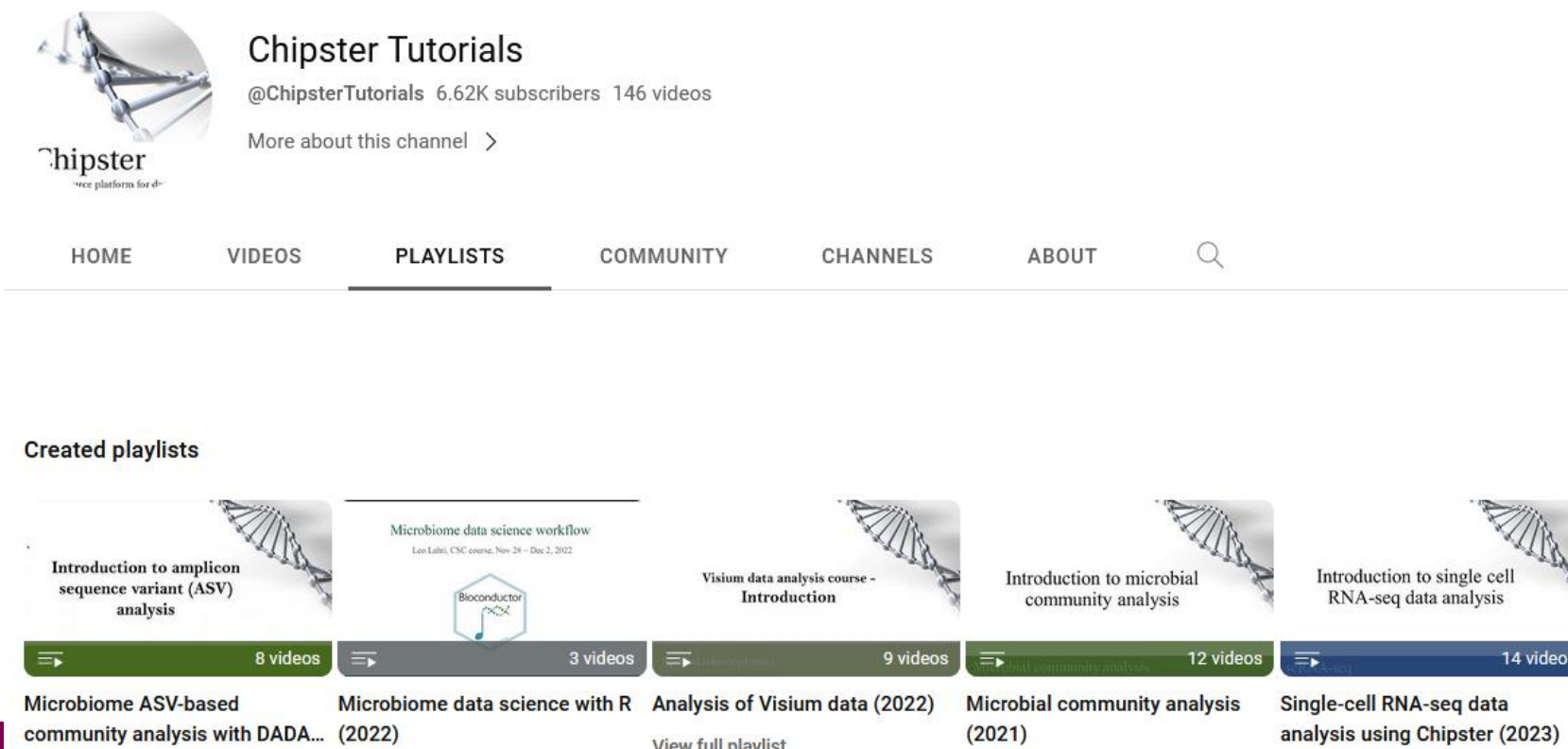
Eija.Korpelainen@csc.fi

Support personnel will use this address to contact you.
This email address was received from your login details. If it's not correct, please contact the organization that provided your login credentials to update it.

[Cancel](#) [Send](#)

More info

- chipster@csc.fi
- <http://chipster.csc.fi>
- Chipster tutorials in YouTube
- <https://chipster.csc.fi/manual/courses.html>



Chipster Tutorials

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Playlist	Video Count	View full playlist
Introduction to amplicon sequence variant (ASV) analysis	8 videos	View full playlist
Microbiome data science workflow	3 videos	View full playlist
Visium data analysis course - Introduction	9 videos	View full playlist
Introduction to microbial community analysis	12 videos	View full playlist
Introduction to single cell RNA-seq data analysis	14 videos	View full playlist

Microbiome ASV-based community analysis with DADA... (2022)

Microbiome data science with R

Analysis of Visium data (2022)

Microbial community analysis (2021)

Single-cell RNA-seq data analysis using Chipster (2023)

Introduction to single-cell RNA-seq data analysis

What will you learn

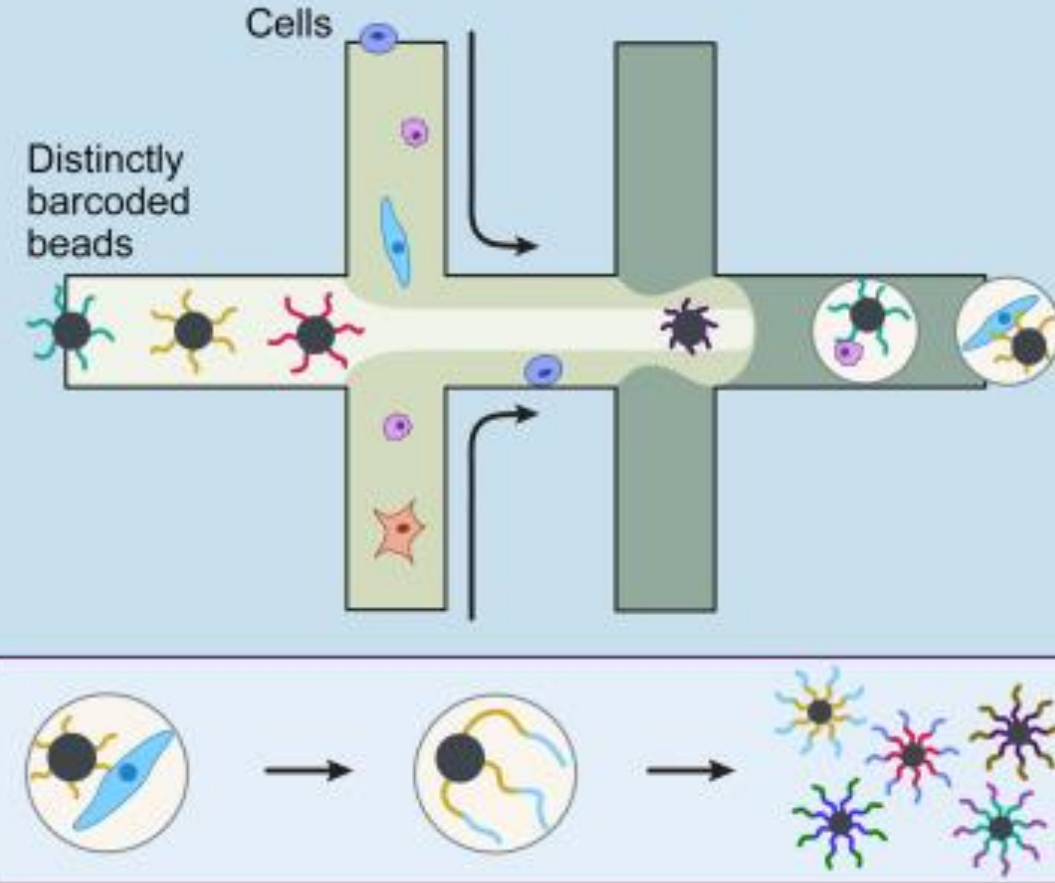
1. How does scRNA-seq work and what can go wrong
 - Empties, doublets and dropouts
 - What is a UMI and why do we use them
2. Why is scRNA-seq data challenging to analyze
3. What are the main analysis steps for clustering cells and finding cluster marker genes
4. What is Seurat

Single-cell RNA-seq

- Relatively new technology, data analysis methods are actively developed
- Gene expression profiling at single cell level has many applications
 - cell type detection, cellular differentiation processes, tumor heterogeneity and response to drugs, etc
- Many technologies for capturing single cell transcriptomes
 - Droplet-based (e.g. 10X Chromium, Drop-seq), plate-based and well-based
- Libraries are usually 3' tagged: only a short sequence at the 3' end of the mRNA is sequenced
- The size and scale of single-cell sequencing datasets is rapidly increasing

Bead: Cell barcode and unique molecular identifiers (UMIs)

Drop-seq single cell analysis



- Cell barcode: which cell the read comes from
- UMI: which mRNA molecule the read comes from (helps to detect PCR duplicates)

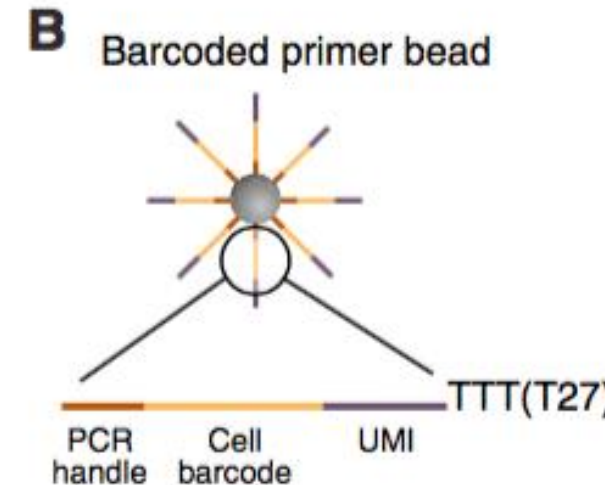
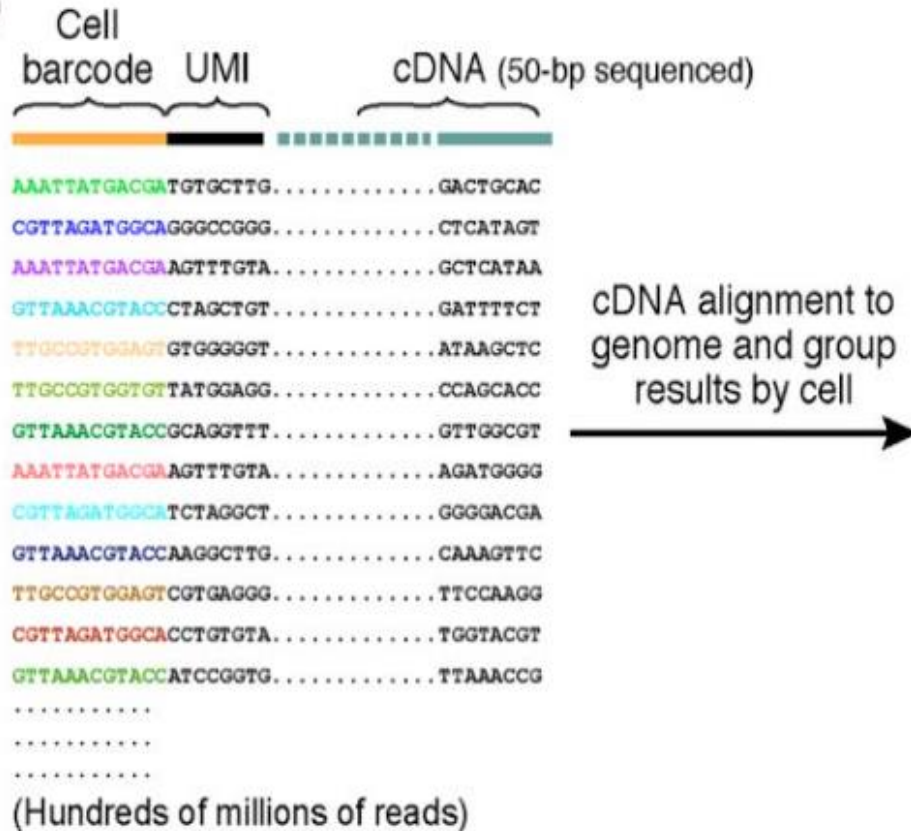


Figure by Macosko et al, *Cell*, 161:1202-1214, 2015

From reads to digital gene expression matrix (DGE)



Overview of DGE extraction



Cell 1	{	TTGCCGTGGAGT GTGGGGGT.....	ATAAGCTC] DDX51
		TTGCCGTGGTGTATGGAGG.....	CCAGCACC] NOP2
		TTGCCGTGGAGTCGTGAGGG.....	TTCCAAGG] ACTB
Cell 2	{	CGTTAGATGGCA GGGCCGGG.....	CTCATAGT] LBR
		CGTTAGATGGCA CCTGTGTA.....	TGGTACGT] ODF2
		CGTTAGATGGCA TCTAGGCT.....	GGGGACGA] HIF1A
Cell 3	{	AAATTATGACGA AGTTTGTA.....	GCTCATAA] ACTB
		AAATTATGACGA AGTTTGTA.....	AGATGGGG] RPS15
		AAATTATGACGA TGTGCTTG.....	GACTGCAC	
Cell 4	{	GTAAACGTACC CTAGCTGT.....	GATTTCT] GTPBP4
		GTAAACGTACC GCAGGTTT.....	GTGGCGT] GAPDH
		GTAAACGTACC AAGGCTG.....	CAAAGTTC] ARL1
		GTAAACGTACC ATCCGGTG.....	TTAAACCG	

Count unique UMIs for each gene in each cell

→

Create digital expression matrix

	Cell: 1	2	...	N
GENE 1	1	2		14
GENE 2	4	27		8
GENE 3	0	0		1
:	:	:		:
GENE M	6	2		0

Figure by Macosko et al, Cell, 161:1202-1214, 2015

What can go wrong?

1. Ideally there is one healthy cell in the droplet. However, sometimes
 - There is no cell in the droplet, just ambient RNA
 - Remove “empties” based on the small number of genes expressed
 - There are two (or more) cells in a droplet
 - Remove doublets (and multiplets) based on the large number of genes expressed
 - The cell in the droplet is broken/dead
 - Remove dead cells based on high percentage of mitochondrial transcripts
2. Sometimes barcodes have synthesis errors in them, e.g. one base is missing
 - Check the distribution of bases at each position and fix the barcode or remove the cell

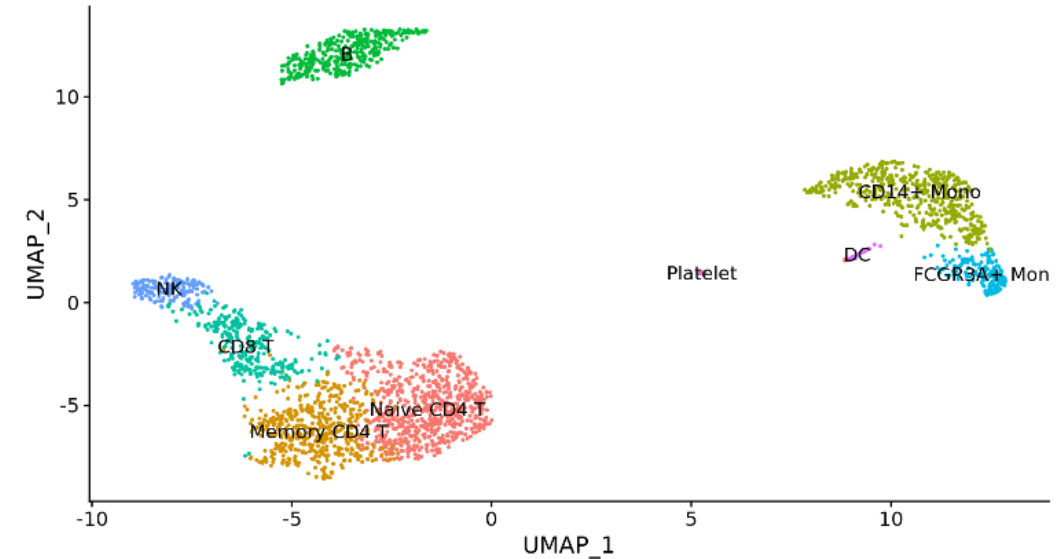
Single-cell RNA-seq data is challenging

- High number of dropouts
 - A gene is expressed but the expression is not detected due to technical limitations → the detected expression level for many genes is zero
 - Data is noisy. High level of variation between the cells due to
 - Capture efficiency (percentage of mRNAs captured)
 - Reverse transcription efficiency
 - Amplification bias (non-uniform amplification of transcripts)
 - Significant differences in sequencing depth (number of UMIs/cell)
 - Cell size and cell cycle stage
 - Complex distribution of expression values
 - Cell heterogeneity and the abundance of zeros give rise to multimodal distributions
- Analysis methods for bulk RNA-seq data don't work for single cell RNA-seq

Analysis steps for clustering cells and finding cluster marker genes

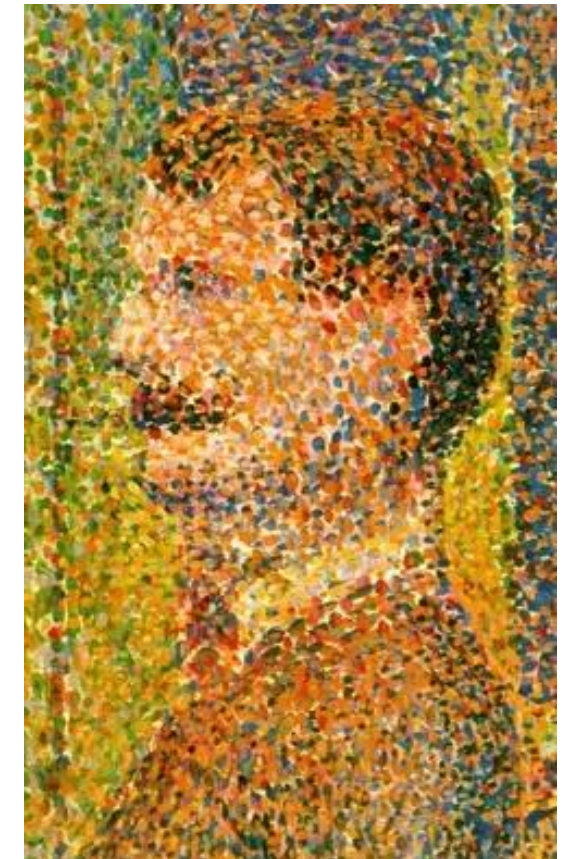


1. Check the quality of cells, filter genes
2. Filter out low quality cells
3. Normalize expression values
4. Identify highly variable genes
5. Scale data, regress out unwanted variation
6. Reduce dimensions using principal component analysis (PCA) on the variable genes
7. Determine significant principal components (PCs)
8. Use the PCs to cluster cells with graph based clustering
9. Visualize clusters with non-linear dimensional reduction (UMAP or tSNE) using the PCs
10. Detect and visualize marker genes for the clusters



Seurat

- One of the most popular R packages for scRNA-seq data analysis
- Provides tools for all the steps mentioned in the previous slide
 - Also tools for integrative analysis
- Stores data in Seurat object
 - Contains specific slots for different types of data like counts, PCA and clustering results, etc
- <http://satijalab.org/seurat>



Detail from La Parade (1889) by Georges Seurat

Analysis steps for clustering cells and finding marker genes



1. Create Seurat object, filter genes, check the quality of cells
2. Filter out low quality cells
3. Normalize expression values
4. Identify highly variable genes
5. Scale data, regress out unwanted variation
6. Reduce dimensions using principal component analysis (PCA) on the variable genes
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What will you learn

1. What kind of input files can be used
2. What is the structure of 10X Genomics matrix file
3. How to filter out genes
4. How to check the quality of cells and filter out bad ones

Start the analysis with one of these 3 types of input files in Chipster:

- 10X Genomics Market Exchange (MEX) format
 - Three files are needed: barcodes.tsv, features.tsv (genes.tsv) and matrix.mtx
 - the files need to be named exactly like this
 - You need to put the files in a tar package (use Chipster tool “Utilities / Make a Tar package”)
- 1. 10X Genomics CellRanger or CellBender HDF5 output format (.h5 file)
 - Hierarchical Data Format (HDF5 or H5) is a binary format that can compress and access data much more efficiently than text formats such as MEX, so it is especially useful for large datasets.
- 2. DGE (=digital gene expression) matrix (.tsv file)
 - DGE matrix made with DropSeq tools in Chipster, or import a ready-made matrix (.tsv file)
 - Genes as rows, cells as columns
- Check that the input file is correctly assigned!

What do the 10X files contain?

1. matrix.mtx

- Number of UMIs for a given gene in a given cell
- Sparse matrix (only non-zero entries are stored), in MEX format
 - Header: third line tells how many genes and cells you have
 - Each row: gene index, cell index, **number of UMIs**
- Make sure that you use the filtered feature barcode matrix (contains only those cell barcodes which are present in your data)

2. barcodes.tsv

- Cell barcodes present in your data

3. features.tsv (genes.tsv)

- Identifier, name and type (gene expression)

%%MatrixMarket matrix coordinate real			
%			
32738	2700	2286884	
32709	1	4	
32707	1	1	
32706	1	10	
32704	1	1	
32703	1	5	
32702	1	6	
32700	1	10	1
32699	1	25	2
			3
			...
			AAACATACAAACCAC-1
			AAACATTGAGCTAC-1
			AAACATTGATCAGC-1
			AAACCGTGTTCG-1
			AAACCGTGTATGCG-1
			AAACGCACTGGTAC-1
			AAACGCTGACCACT-1
			AAACGCTGGTTCTT-1
			AAACGCTGTAGCCA-1

Setting up a Seurat object, filtering genes



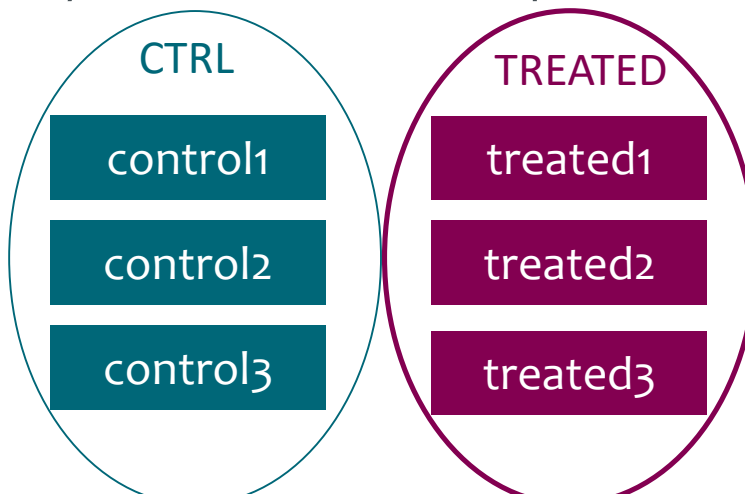
- Give a name for the project (used in some plots)
- Filtering genes
 - Keep genes which are expressed (= detected) in at least this number of cells

- Input files

- Assign correctly!

- Sample name & sample group

- Important if you have several samples



Seurat v4 -Setup and QC

Parameters

Project name for plotting

You can give your project a name. The name will appear on the plots. Do not use underscore _ in the names!

Keep genes which are expressed in at least this many cells

The genes need to be expressed in at least this many cells.

Sample name

Type the sample name or identifier here. For example control1, cancer3a. Do not use underscore _ in the names! Fill this field if you are combining samples later.

Sample group

Type the sample name or identifier here. For example CTRL, STIM, TREAT. Do not use underscore _ in the names! Fill this field if you are combining samples later.

Input files

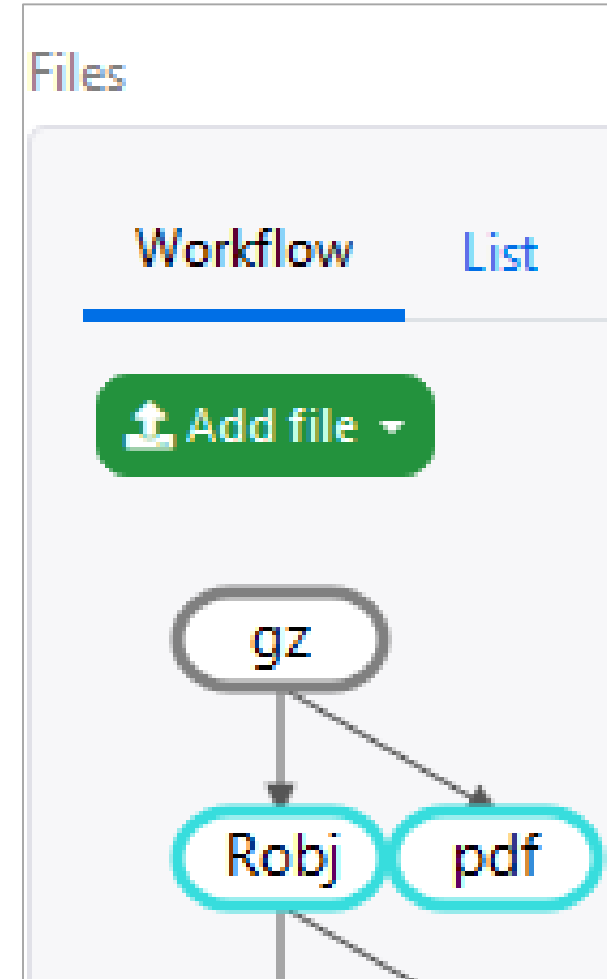
tar package of 10X output files

DGE table in tsv format

10X CellRanger hdf5 input file

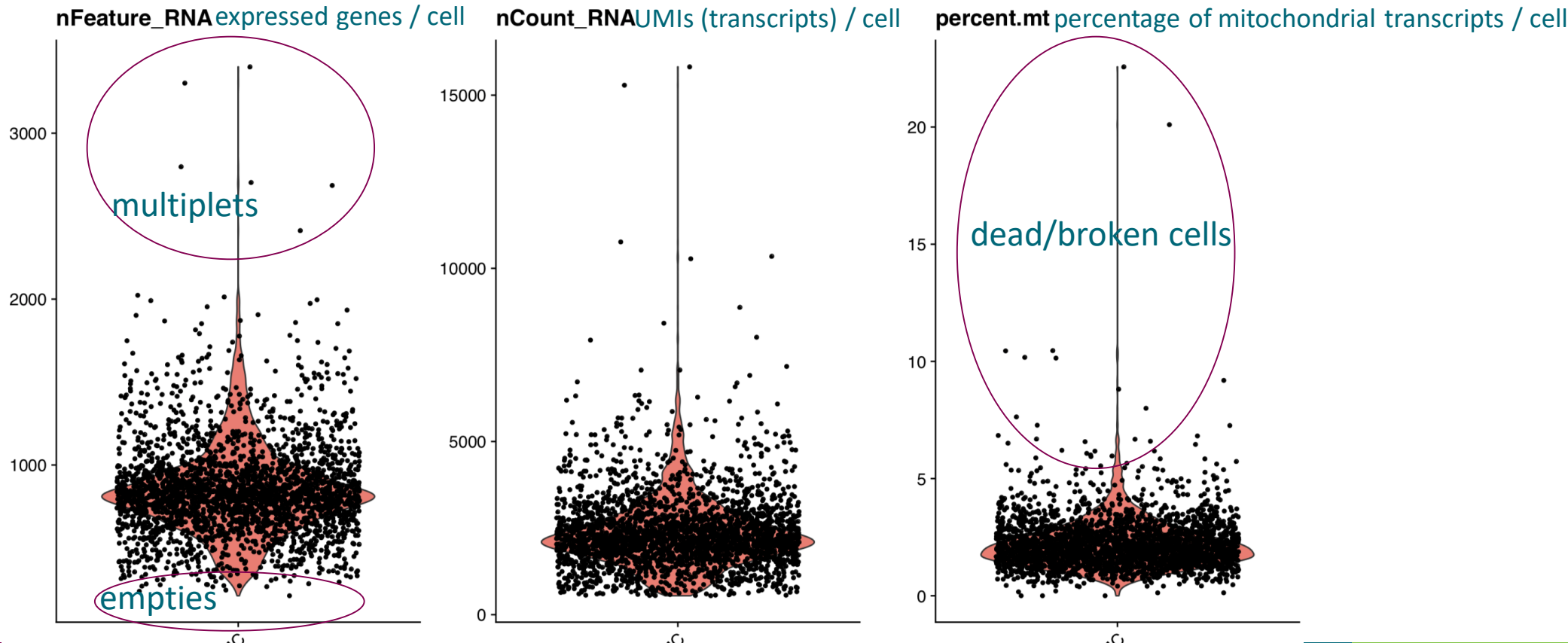
Output files

- Seurat object (Robj) that Seurat-based tools use to store data
 - Use this file as input for the next analysis tool
 - Contains specific slots for different types of data
 - View the contents using the tool Extract information from Seurat object
 - You cannot open the file in Chipster but you can import it to R
- Pdf file with quality control plots and cell number info
 - nFeature_RNA = number of expressed genes in a cell
 - nCount_RNA = number of transcripts in a cell
 - percent.mt = percentage of mitochondrial transcripts
 - percent.rb = percentage of ribosomal transcripts



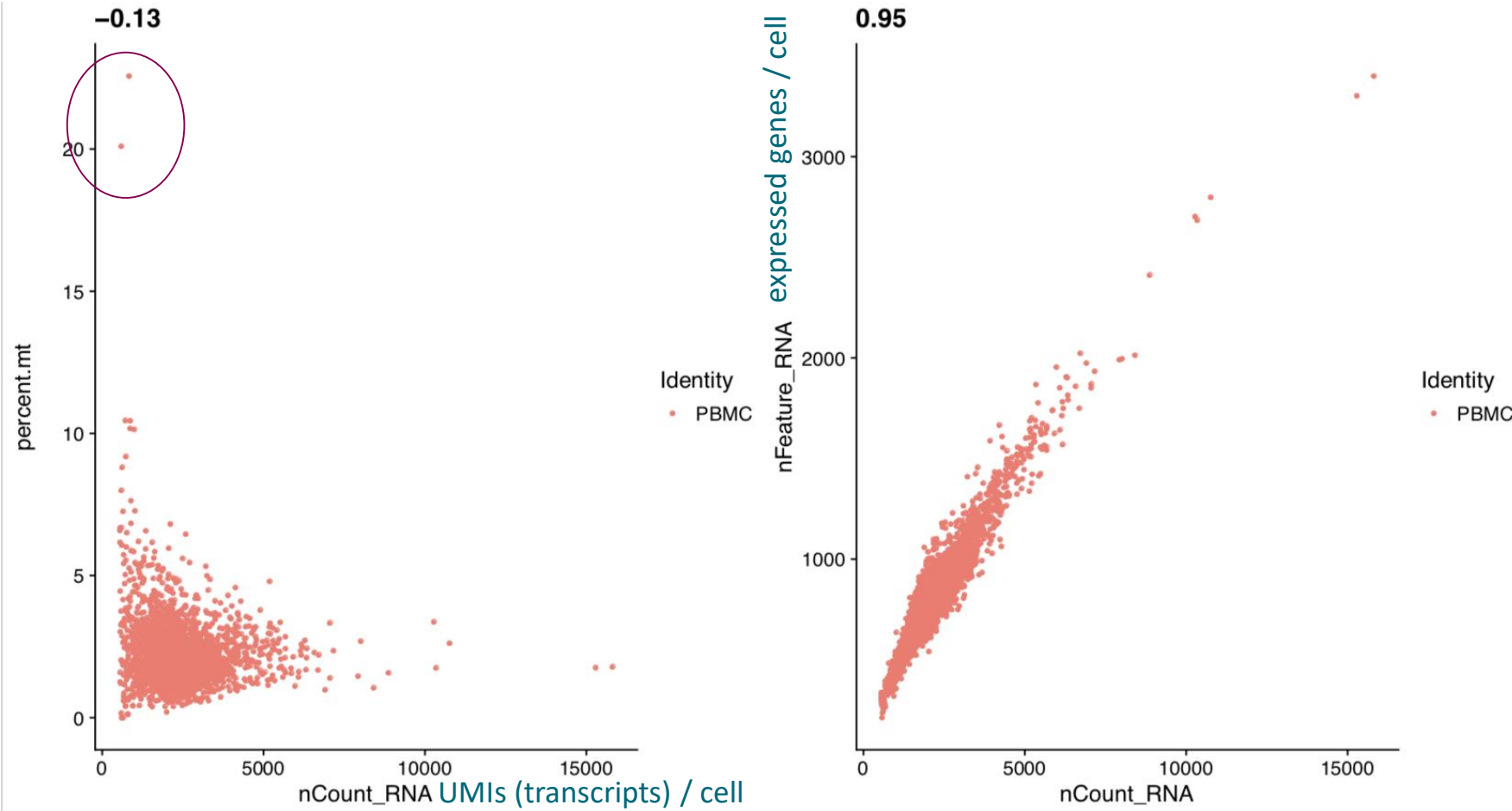
How to detect empties, multiplets and broken cells?

- Empty = no cell in droplet: low gene count (`nFeature_RNA < 200`)
- Doublet/multiplet = more than one cell in droplet: large gene count (`nFeature_RNA > 2500`)
- Broken/dead cell in droplet: lot of mitochondrial transcripts (`percent.mt > 5%`)



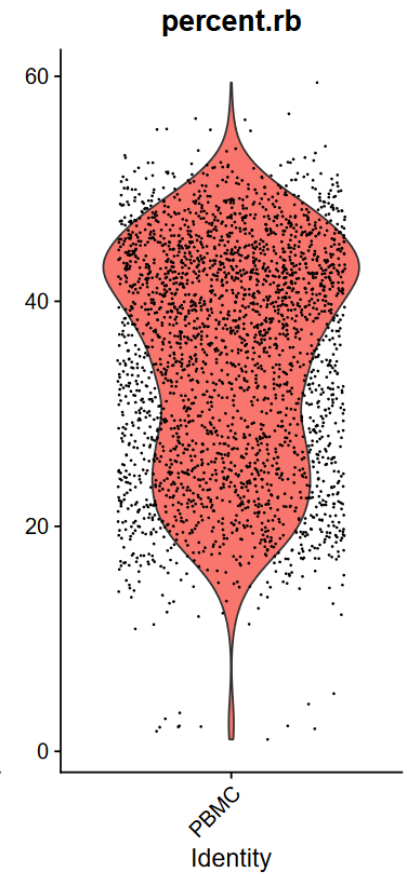
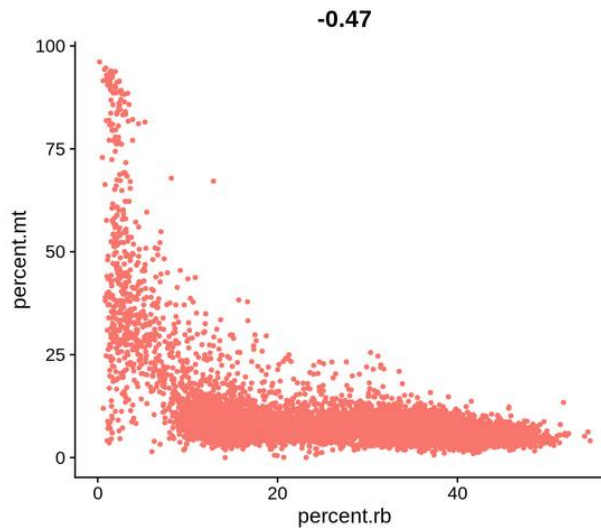
Scatter plots for quality control

- nCount_RNA vs percent.mt: are there cells with low number of transcripts and high mito%
- nCount_RNA vs nFeature_RNA: these should correlate.



Percentage of UMIs mapping to ribosomal genes (percent.rb)

- Ribosomal transcripts don't have polyA, so they should not be captured
- However, a large proportion of UMIs can be ribosomal
 - Percent.rb varies between cells for technical and biological reasons
- Percent.rb and percent.mito *anti-correlate*
 - You can filter out cells which have *lower* ribosomal percentage than x



Parameters for filtering out bad quality cells



Seurat v4 - Filter cells, normalize, regress and detect variable genes



Parameters

 Reset All

Filter out cells which have less than this many genes expressed Filter out empties. The cells to be kept must express at least this number of genes.	200
Filter out cells which have more than this many genes expressed Filter out multiplets. The cells to be kept must express less than this number of genes.	2500
Filter out cells which have higher mitochondrial transcript percentage Filter out dead cells. The cells to be kept must have lower percentage of mitochondrial transcripts than this if needed in your data.	5
Filter out cells which have lower ribosomal transcript percentage Filter out cells that have lower ribosomal transcript percentage.	0
Perform global scaling normalization For raw data, select yes.	yes
Scaling factor in the normalization Scale each cell to this total number of transcripts.	10000
Number of variable genes to return Number of features to select as top variable features, i.e. how many features returned.	2000
Regress out cell cycle differences Would you like to regress out cell cycle scores during data scaling? If yes, should all signal associated with cell cycle be removed, or only the difference between the G2M and S phase scores.	no

Analysis steps for clustering cells and finding marker genes



1. Create Seurat object, filter genes, check the quality of cells
2. Filter out low quality cells
3. **Normalize expression values**
4. Identify highly variable genes
5. Scale data, regress out unwanted variation
6. Reduce dimensions using principal component analysis (PCA) on the variable genes
7. Determine significant principal components (PCs)
8. Use the PCs to cluster cells with graph based clustering
9. Visualize clusters with non-linear dimensional reduction (tSNE or UMAP) using the PCs
10. Detect and visualize marker genes for the clusters

What will you learn

1. Why do we need to normalize gene expression values
2. What is a dropout
3. What does global scaling normalization do
4. When does it not work well

Normalizing scRNA-seq gene expression values

- We cluster cells based on differences in their gene expression profiles
- Variance of gene expression values should reflect biological variation across cells
 - We need to remove non-biological variation
- Single-cell gene expression values are noisy
 - Low mRNA content in a cell
 - Variable mRNA capture
 - Variable sequencing depth
- Normalization methods for bulk RNA-seq data don't work for single cell data
 - dropouts = genes whose expression is not detected → lot of zeros

Global scaling normalization

- Divide gene's UMI count in a cell by the total number of UMIs in that cell
- Multiply the ratio by a scale factor (10,000 by default)
 - This scales each cell to this total number of transcripts
- Transform the result by taking natural log

Parameters for normalization

Seurat v4 -Filter cells, normalize, regress and detect variable genes X

Parameters

 Reset All

Filter out cells which have less than this many genes expressed

Filter out empties. The cells to be kept must express at least this number of genes.



Filter out cells which have more than this many genes expressed

Filter out multiplets. The cells to be kept must express less than this number of genes.



Filter out cells which have higher mitochondrial transcript percentage

Filter out dead cells. The cells to be kept must have lower percentage of mitochondrial transcripts than this if needed in your data.



Filter out cells which have lower ribosomal transcript percentage

Filter out cells that have lower ribosomal transcript percentage.



Perform global scaling normalization

For raw data, select yes.



Scaling factor in the normalization

Scale each cell to this total number of transcripts.



Number of variable genes to return

Number of features to select as top variable features, i.e. how many features returned.



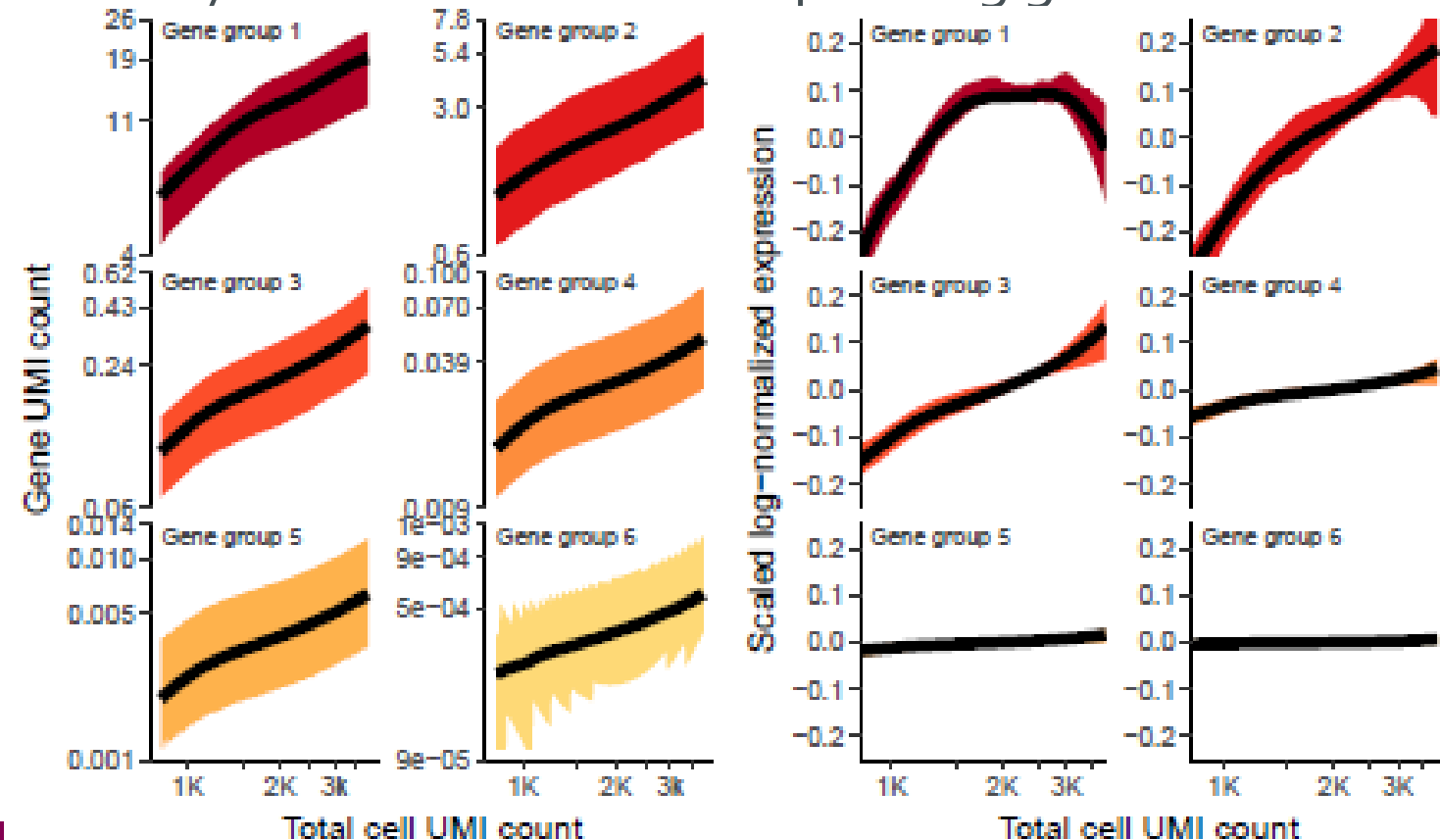
Regress out cell cycle differences

Would you like to regress out cell cycle scores during data scaling? If yes, should all signal associated with cell cycle be removed, or only the difference between the G2M and S phase scores.



Global scaling normalization: problem with high expressing genes

- Sequencing depth (number of UMIs per cell) varies significantly between cells
- Normalized expression values of a gene should be independent of sequencing depth
- The global scaling normalization works only for low to medium expressing genes
 - Expression values of high expressing genes correlate with sequencing depth
 - SCTransform can deal with this better
 - Hafemeister (2019): Normalization and variance stabilization of single-cell RNA-seq data using regularized negative binomial regression



SCTransform – alternative approach to normalization etc



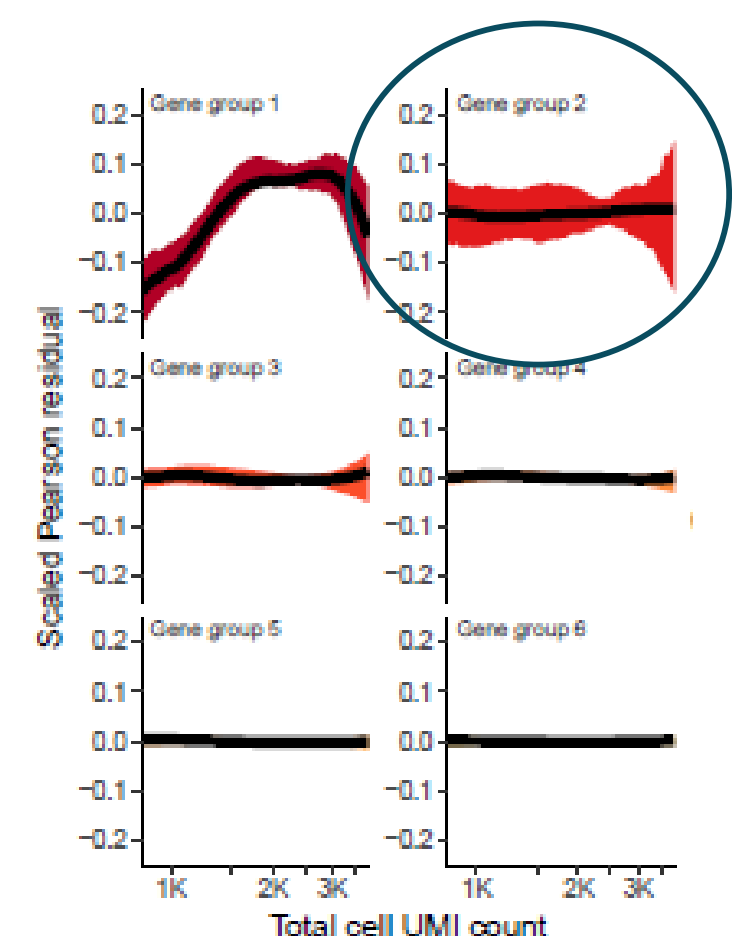
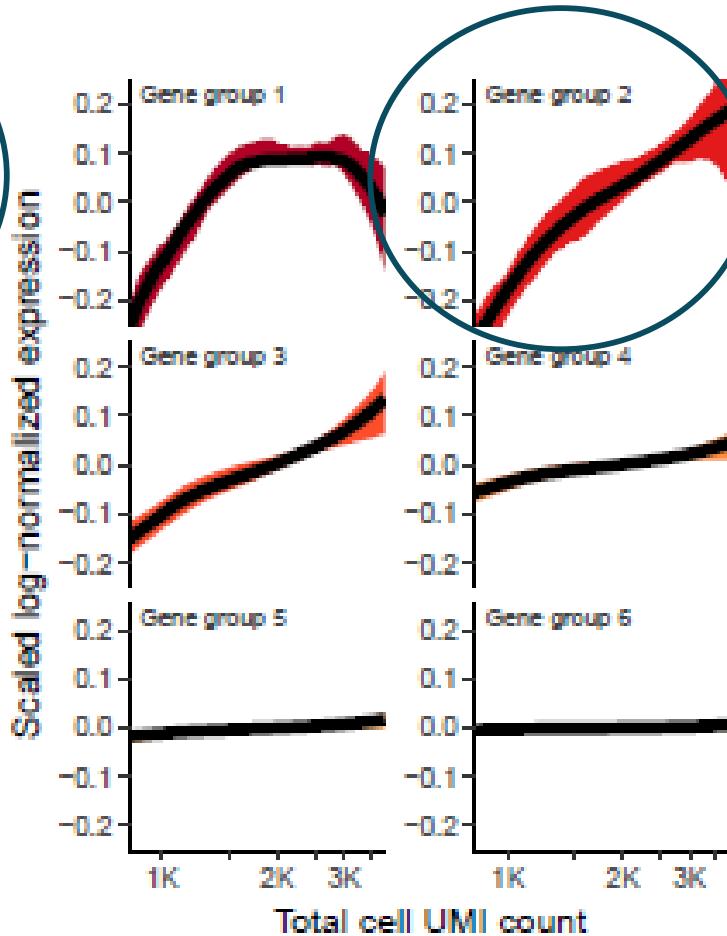
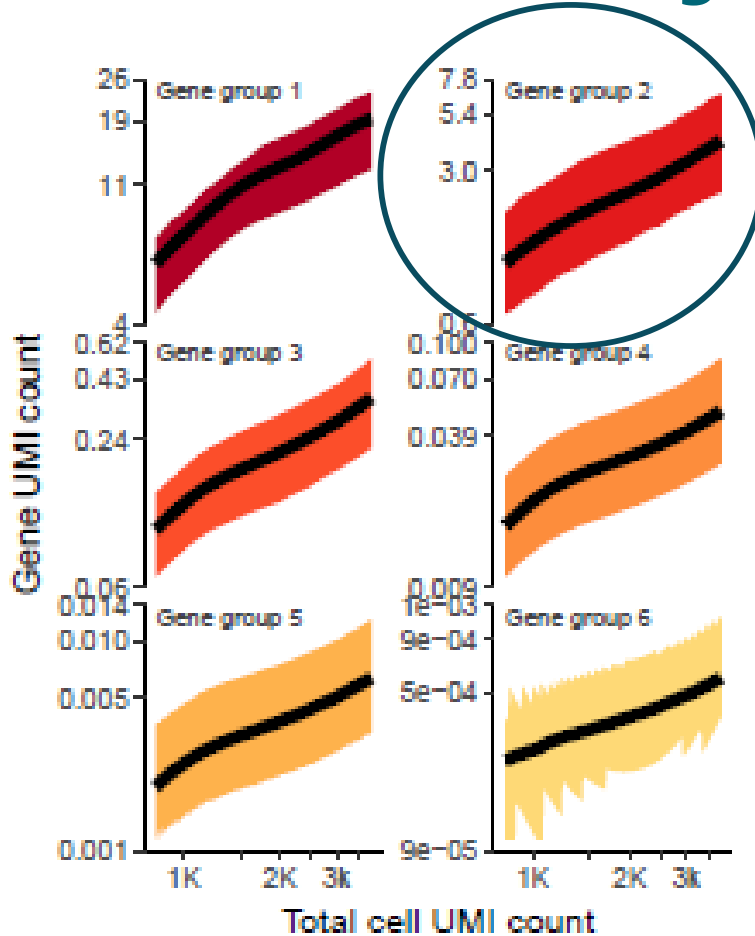
1. Create Seurat object, filter genes, check the quality of cells
 2. Filter out low quality cells
 3. ~~Normalize expression values~~
 4. ~~Identify highly variable genes~~
 5. ~~Scale data, regress out unwanted variation~~
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- 
- SCTransform**

SCTransform: modeling framework for normalization and variance stabilization



- Sequencing depth (number of UMIs per cell) varies significantly between cells
- Normalized expression values of a gene should be independent of sequencing depth
- The default log normalization works ok only for low to medium expressing genes
 - For high expressing genes the normalized expression values correlate with sequencing depth
 - High expressing genes show disproportionately high variance in cells with low sequencing depth
- SCTransform models gene expression as a function of sequencing depth using GLM
 - Constrains the model parameters through regularization, by pooling information across genes which are expressed at similar levels
 - Normalized expression values = Pearson residuals from regularized negative binomial regression
 - Pearson residual = response residual devided by the expected standard deviation (effectively VST)
 - Positive residual for a given gene in a given cell indicate that we observed more UMIs than expected given the gene's average expression in the population and the cellular sequencing depth

Normalization using Pearson residuals works best



Hafemeister (2019): Normalization and variance stabilization of single-cell RNA-seq data using regularized negative binomial regression

Parameters for SCTransform

Seurat v4 -SCTransform: Filter cells, normalize, regress and detect variable genes X

Parameters

Reset All

Filter out cells which have less than this many genes expressed
Filter out empties. The cells to be kept must express at least this number of genes.

 ^

Filter out cells which have more than this many genes expressed
Filter out multiplets. The cells to be kept must express less than this number of genes.

 ^

Filter out cells which have higher mitochondrial transcript percentage
Filter out dead cells. The cells to be kept must have lower percentage of mitochondrial transcripts than this.

 ^

Filter out cells which have lower ribosomal transcript percentage
Filter out cells that have lower ribosomal transcript percentage.

 ^

Number of variable genes to return
Number of features to select as top variable features, i.e. how many features returned. For SCTransform, the recommended default is 3000.

 ^

Regress out cell cycle differences
Would you like to regress out cell cycle scores during data scaling? If yes, should all signal associated with cell cycle be removed, or only the difference between the G2M and S phase scores.

 ^

SCTransform: things to take into account in analysis



- When the data is normalized with SCTransform, it is recommended to set
 - In normalization: Number of highly variable genes = 3000 (instead of 2000)
 - In PCA: Number of PCs to compute = 50 (instead of 20)
 - In clustering: Number of principal components to use = 30 (instead of 10), resolution = 0.8 (instead of 0.5)
- Why do we use a different number of highly variable genes and PCs after SCTransform?
 - SCTransform does a better job in normalization (variation in sequencing depth is not a confounding factor any more) → additional variable features are less likely to be driven by technical differences across cells, and instead may represent more subtle biological variability

Analysis steps for clustering cells and finding marker genes



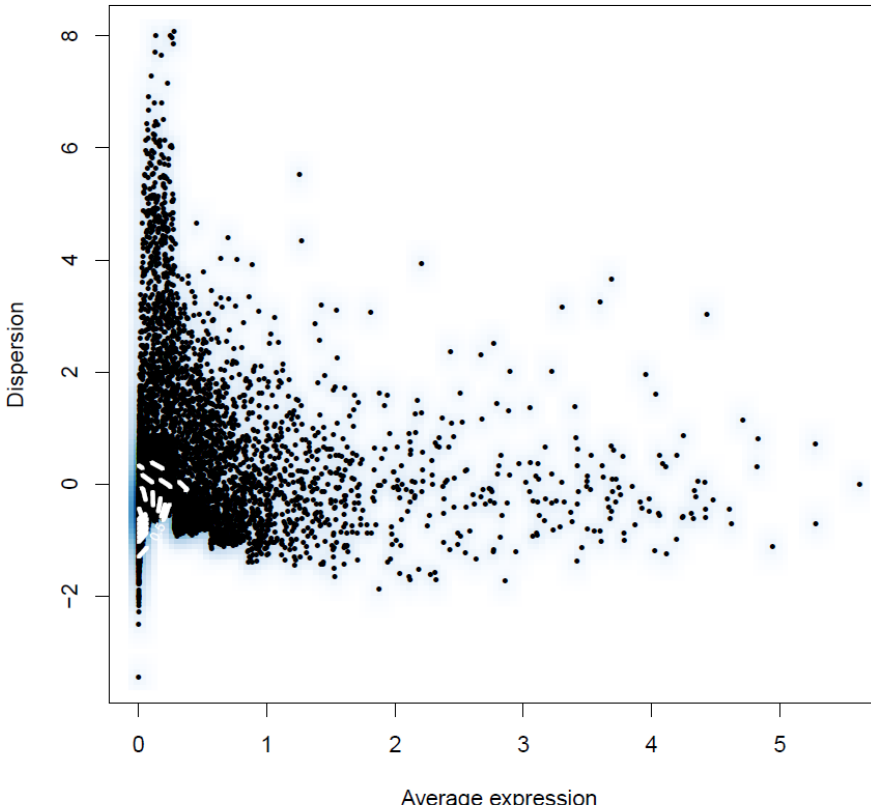
1. Create Seurat object, filter genes, check the quality of cells
2. Filter out low quality cells
3. Normalize expression values
4. **Identify highly variable genes**
5. Scale data, regress out unwanted variation
6. Reduce dimensions using principal component analysis (PCA) on the variable genes
7. Determine significant principal components (PCs)
8. Use the PCs to cluster cells with graph based clustering
9. Visualize clusters with non-linear dimensional reduction (tSNE or UMAP) using the PCs
10. Detect and visualize marker genes for the clusters

What will you learn

1. Why do we need to find highly variable genes
2. What kind of mean-variance relationship is there in scRNA-seq data
3. Why do we need to stabilize the variance of gene expression values

Selecting highly variable genes

- We want to cluster cells, so we need to find genes whose expression varies across the cells
 - Highly variable genes are used for PCA, and the PCs are used for clustering
 - We cannot select genes based on their variance, because scRNA-seq data has strong mean-variance relationship
 - low expressing genes have higher variance
- variance needs to be stabilized first

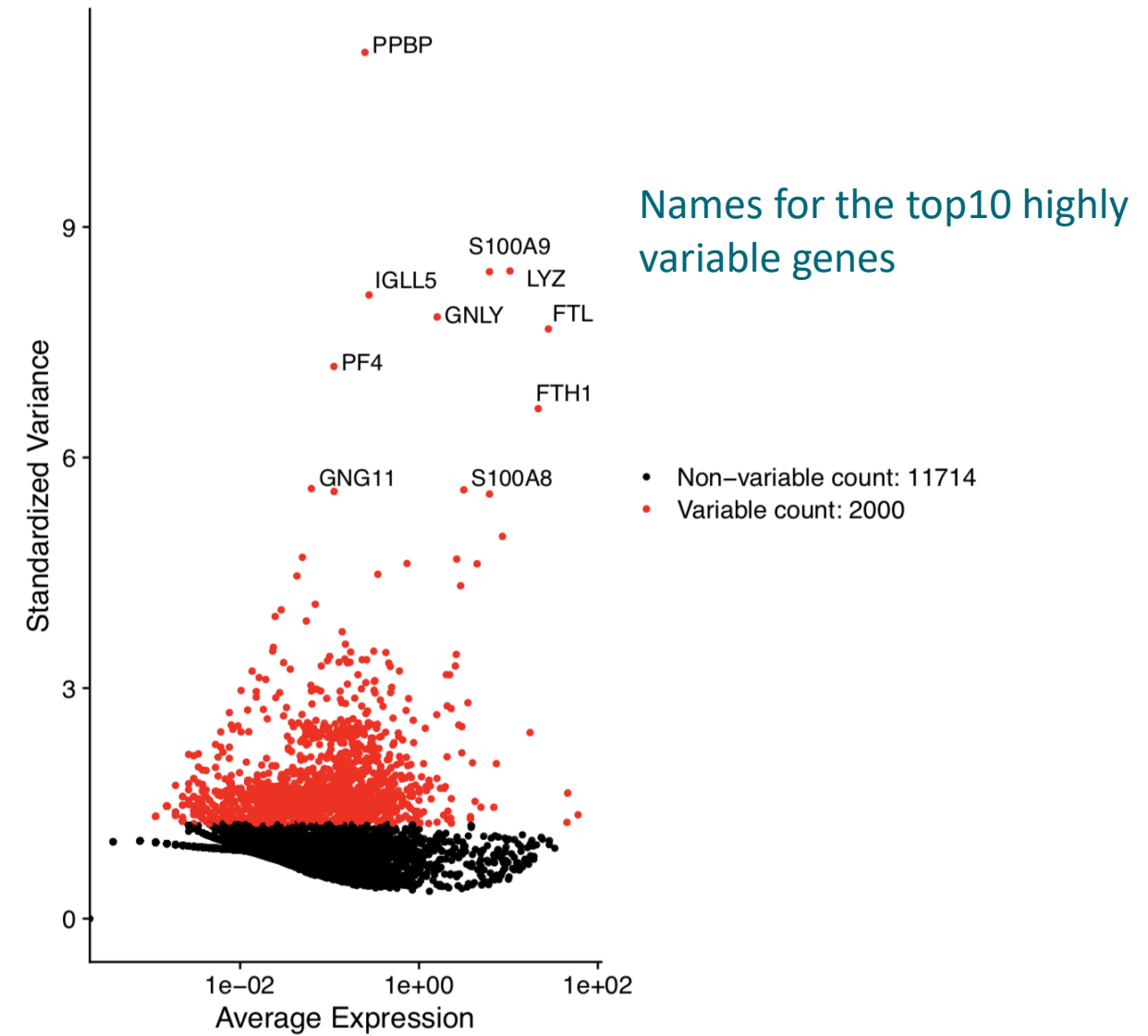
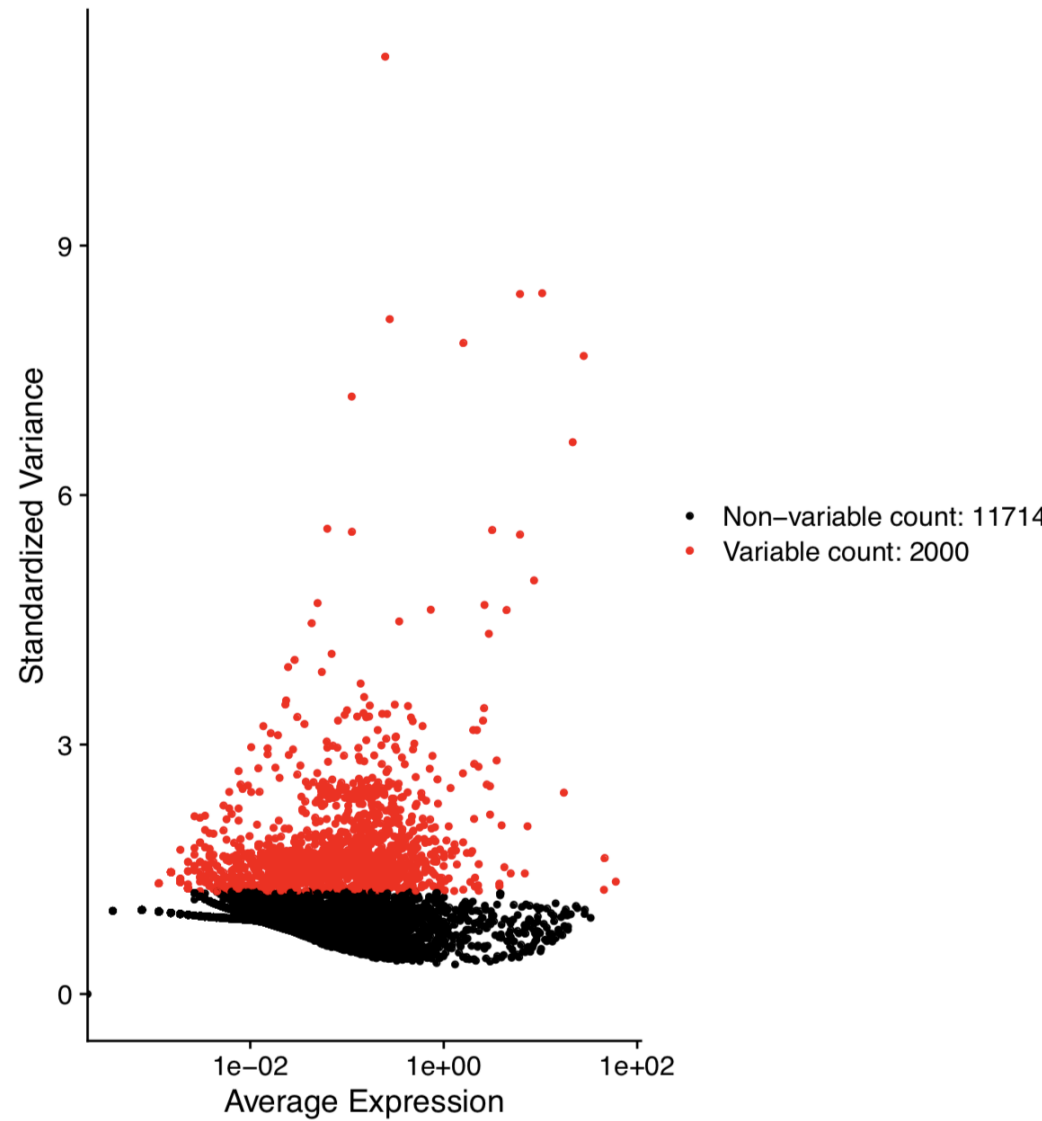


Variance stabilizing transformation (VST)



- Compute the mean and variance for each gene using the unnormalized UMI counts
 - Take \log_{10} of mean and variance
 - Fit a curve to predict the variance of each gene as a function of its mean expression
 - Standardized count = $(\text{expression}_{\text{geneXcellY}} - \text{mean expression}_{\text{geneX}}) / \text{predicted SD}_{\text{geneX}}$
 - reduce the impact of technical outliers: set the max of standardized counts to the square root of number of cells
 - For each gene, compute the variance of the standardized values across all cells
- Rank the genes based on their standardized variance and use the top 2000 genes for PCA and clustering

Detection of highly variable genes: plots



Parameter for detecting variable genes

Seurat v4 -Filter cells, normalize, regress and detect variable genes



Parameters

 Reset All

Filter out cells which have less than this many genes expressed

200



Filter out empties. The cells to be kept must express at least this number of genes.

Filter out cells which have more than this many genes expressed

2500



Filter out multiplets. The cells to be kept must express less than this number of genes.

Filter out cells which have higher mitochondrial transcript percentage

Filter out dead cells. The cells to be kept must have lower percentage of mitochondrial transcripts than this if needed in your data.

5



Filter out cells which have lower ribosomal transcript percentage

Filter out cells that have lower ribosomal transcript percentage.

0



Perform global scaling normalization

For raw data, select yes.

yes



Scaling factor in the normalization

Scale each cell to this total number of transcripts.

10000



Number of variable genes to return

Number of features to select as top variable features, i.e. how many features returned.

2000



Regress out cell cycle differences

Would you like to regress out cell cycle scores during data scaling? If yes, should all signal associated with cell cycle be removed, or only the difference between the G2M and S phase scores.

no



Analysis steps for clustering cells and finding marker genes



1. Create Seurat object, filter genes, check the quality of cells
2. Filter out low quality cells
3. Normalize expression values
4. Identify highly variable genes
5. **Scale data, regress out unwanted variation**
6. Reduce dimensions using principal component analysis (PCA) on the variable genes
7. Determine significant principal components (PCs)
8. Use the PCs to cluster cells with graph based clustering
9. Visualize clusters with non-linear dimensional reduction (tSNE or UMAP) using the PCs
10. Detect and visualize marker genes for the clusters

What will you learn

1. Why do we need to scale data prior to PCA
2. How is scaling done
3. How can we remove unwanted sources of variation

Scaling expression values prior to dimensional reduction



- Standardize expression values for each gene across all cells prior to PCA
 - This gives equal weight in downstream analyses, so that highly expressed genes do not dominate
- Z-score normalization in Seurat's ScaleData function
 - Shifts the expression of each gene, so that the mean expression across cells is 0
 - Scales the expression of each gene, so that the variance across cells is 1
- ScaleData has an option to regress out unwanted sources of variation
 - E.g. cells might cluster according to their cell cycle state rather than cell type

Regress out unwanted sources of variation



- Several sources of uninteresting variation
 - technical noise
 - batch effects
 - cell cycle stage, etc
- Removing this variation improves downstream analysis
- Seurat constructs linear models to predict gene expression based on user-defined variables
 - number of detected transcripts per cell, mitochondrial transcript percentage, batch,...
 - variables are regressed individually against each gene, and the resulting residuals are scaled and centered
 - scaled z-scored residuals of these models are used for dimensionality reduction and clustering
 - **In Chipster** the following effects are removed:
 - number of detected molecules per cell
 - mitochondrial transcript percentage
 - cell cycle stage (optional)

Parameter for regressing out unwanted sources of variation

Seurat v4 -Filter cells, normalize, regress and detect variable genes



Parameters

 Reset All

Filter out cells which have less than this many genes expressed

200

Filter out empties. The cells to be kept must express at least this number of genes.

Filter out cells which have more than this many genes expressed

2500

Filter out multiplets. The cells to be kept must express less than this number of genes.

Filter out cells which have higher mitochondrial transcript percentage

Filter out dead cells. The cells to be kept must have lower percentage of mitochondrial transcripts than this if needed in your data.

Filter out cells which have lower ribosomal transcript percentage

Filter out cells that have lower ribosomal transcript percentage.

Perform global scaling normalization

For raw data, select yes.

Scaling factor in the normalization

Scale each cell to this total number of transcripts.

Number of variable genes to return

Number of features to select as top variable features, i.e. how many features returned.

Regress out cell cycle differences

Would you like to regress out cell cycle scores during data scaling? If yes, should all signal associated with cell cycle be removed, or only the difference between the G2M and S phase scores.

5

0

yes

10000

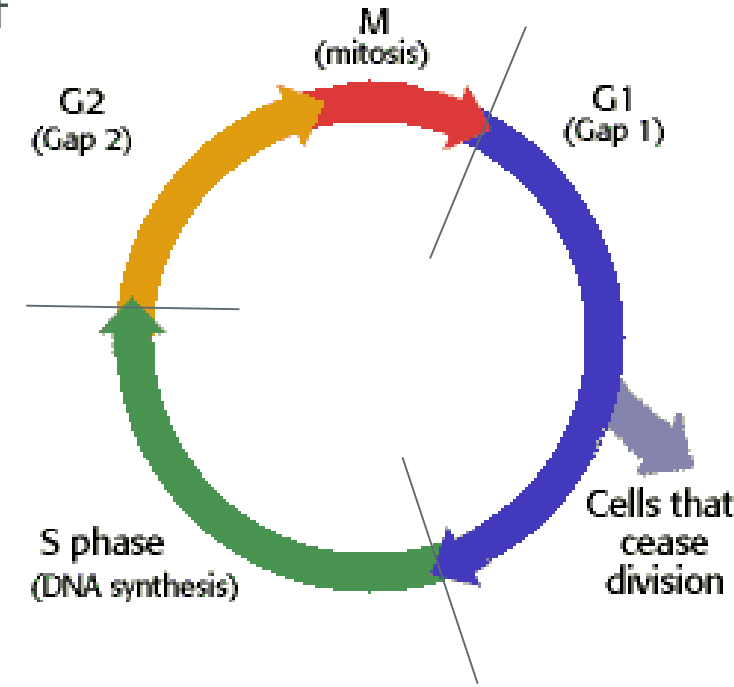
2000

no

Mitigating the effects of cell cycle heterogeneity



1. Compute cell cycle phase scores for each cell based on its expression of G₂/M and S phase marker genes
 - o These markers* are well conserved across tissues and species
 - o Cells which do not express markers are considered not cycling, G₁
2. Model each gene's relationship between expression and the cell cycle score
3. Two options to regress out the variation caused by different cell cycle stages
 1. Remove ALL signals associated with cell cycle stage
 2. Remove the DIFFERENCE between the G₂M and S phase scores.
 - o This preserves signals for non-cycling vs cycling cells, only the difference in cell cycle phase amongst the dividing cells are removed.
Recommended when studying differentiation processes



*list of cell cycle markers, from Tirosh et al, 2015

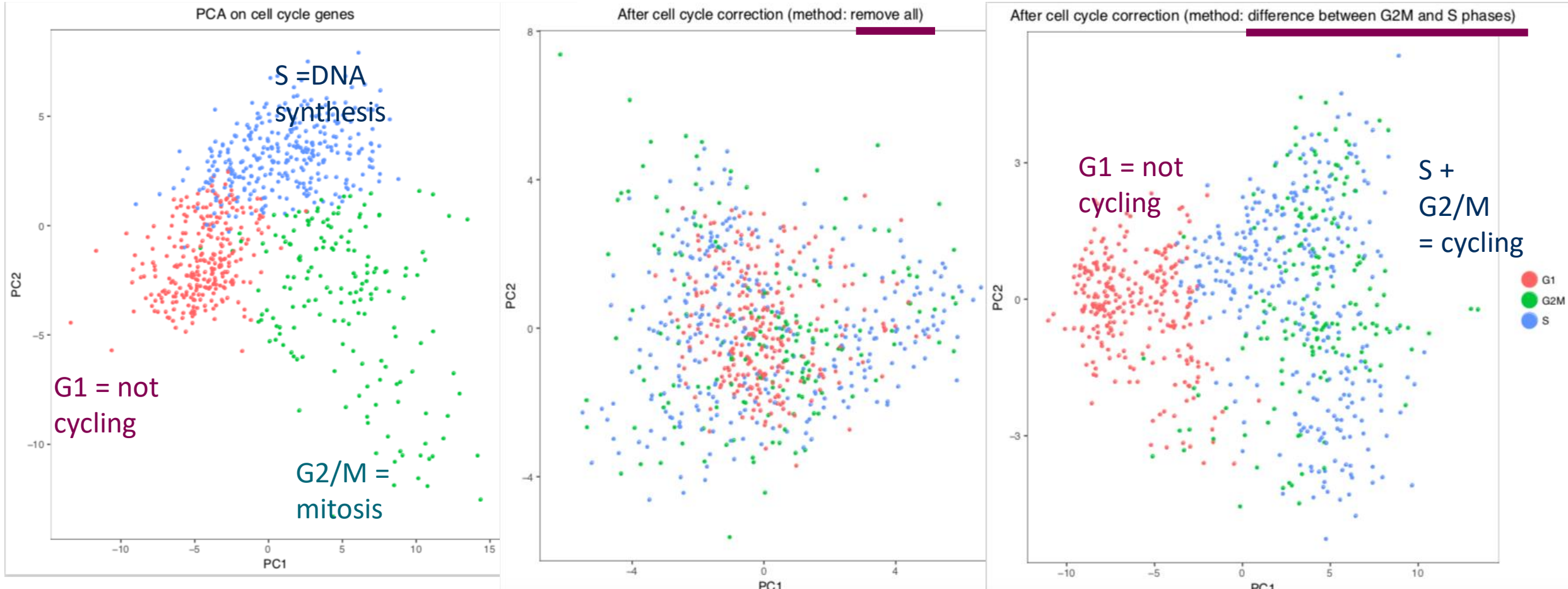
<https://www.ncbi.nlm.nih.gov/pmc/articles/PMC4944528/>

Regressing out the variation caused by different cell cycle stages



When? If we see clear distinction, and the cell differentiation process is not what we are interested in

- OR: perform the clustering without regression and see if we have clusters separated by cell cycle phase. If yes -> come back and perform the regression



PCA on cell cycle genes (dot = cell, colors = phases)

Analysis steps for clustering cells and finding marker genes



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10. Detect and visualize marker genes for the clusters

What will you learn

1. Why do we need to do dimensional reduction?
2. How dimensional reduction methods (PCA, tSNE, UMAP) work on intuitive level
3. Why we use both PCA and tSNE/UMAP?
4. How to select the principal components for the clustering step

Dimensionality reduction



- What for?
 1. Making clustering step easier (PCA)
 2. Visualization (tSNE, UMAP)
- Simplifies complexity so that the data becomes easier to work with
 - Cells are characterized by the expression values of all the genes → thousands of dimensions
 - We have thousands of genes and cells
- Removes redundancies in the data
 - The expression of many genes is correlated, we don't need so many dimensions to distinguish cell types
- Identifies the most relevant information in order to cluster cells
 - Overcomes the extensive technical noise in scRNA-seq data
- Can be linear (e.g. **PCA**) or non-linear (e.g. **tSNE, UMAP**)

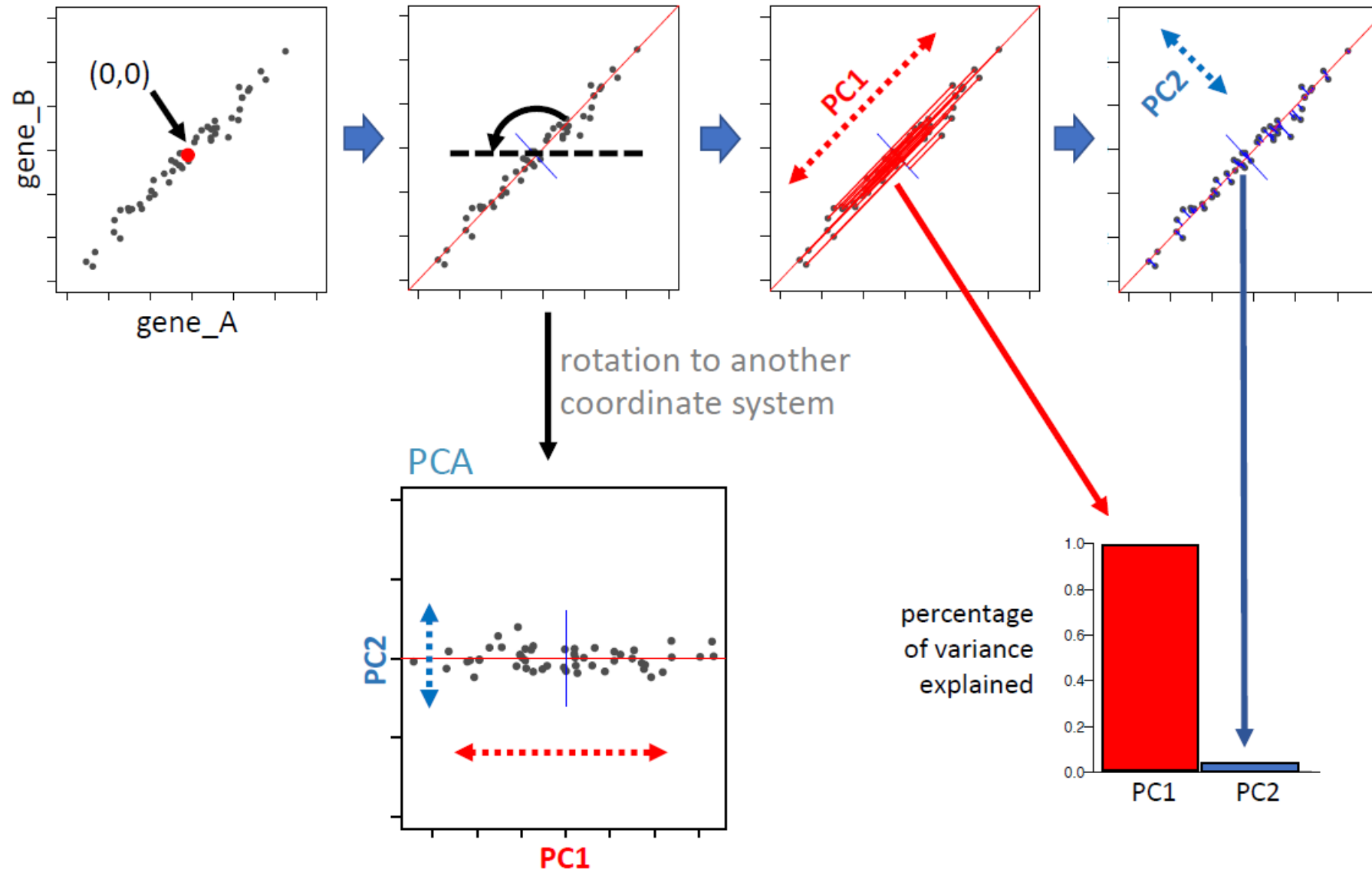
Principal Component Analysis (PCA)



- Finds principal components (PCs) of the data
 - Directions where the data is most spread out = where there is most variance
 - PC₁ explains most of the variance in the data, then PC₂, PC₃, ...
- We will select the most important PCs and use them for clustering cells
 - Instead of 20 000 genes we have now maybe 10 PCs
 - Essentially, each PC represents a robust 'metagene' that combines information across a correlated gene set
- Prior to PCA we scaled the data so that genes have equal weight in downstream analysis and highly expressed genes don't dominate
 - Shift the expression of every gene so that the mean expression across cells is 0 and the variance across cells is 1.

How PCA works

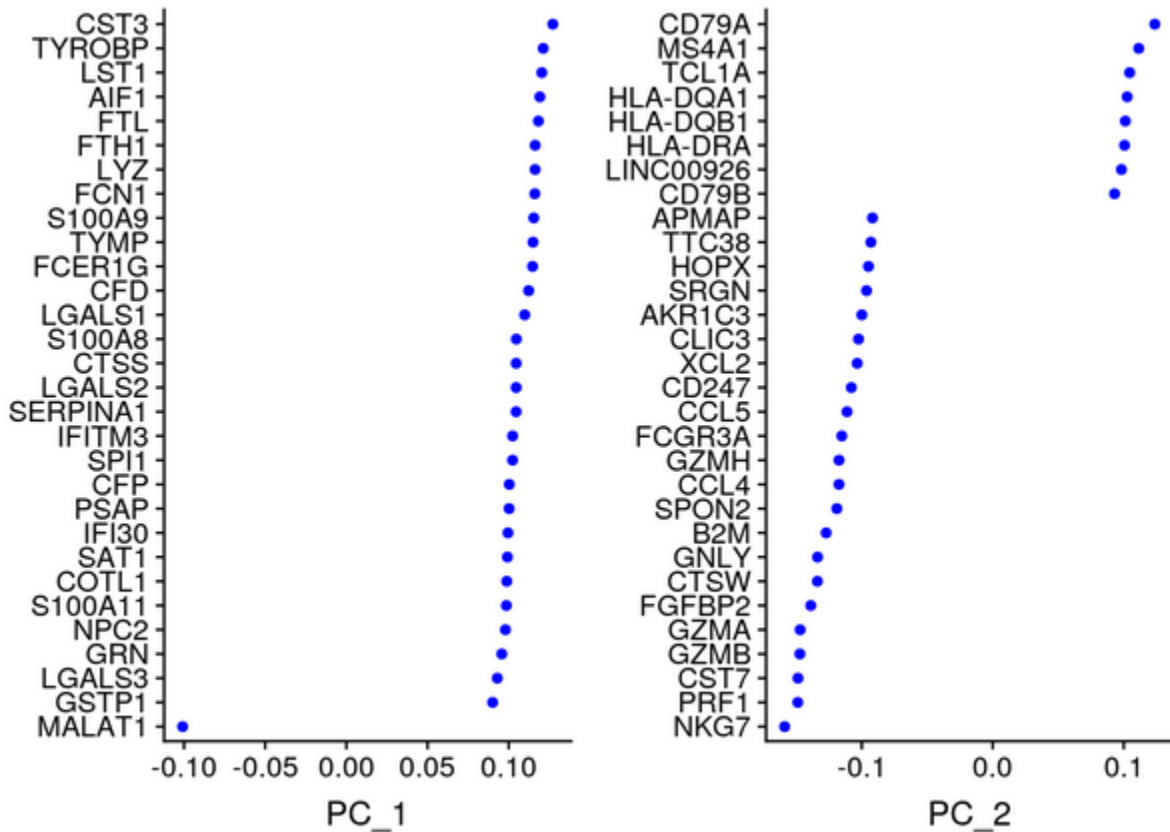
original data (Z-score)



- PC1 explains 98% of the variance
- => PC1 represents these two genes very well
- PC2 is nearly insignificant, and could be disregarded
- In real life, thousands of genes, and maybe tens of PCs

Visualizing PCA results: loadings

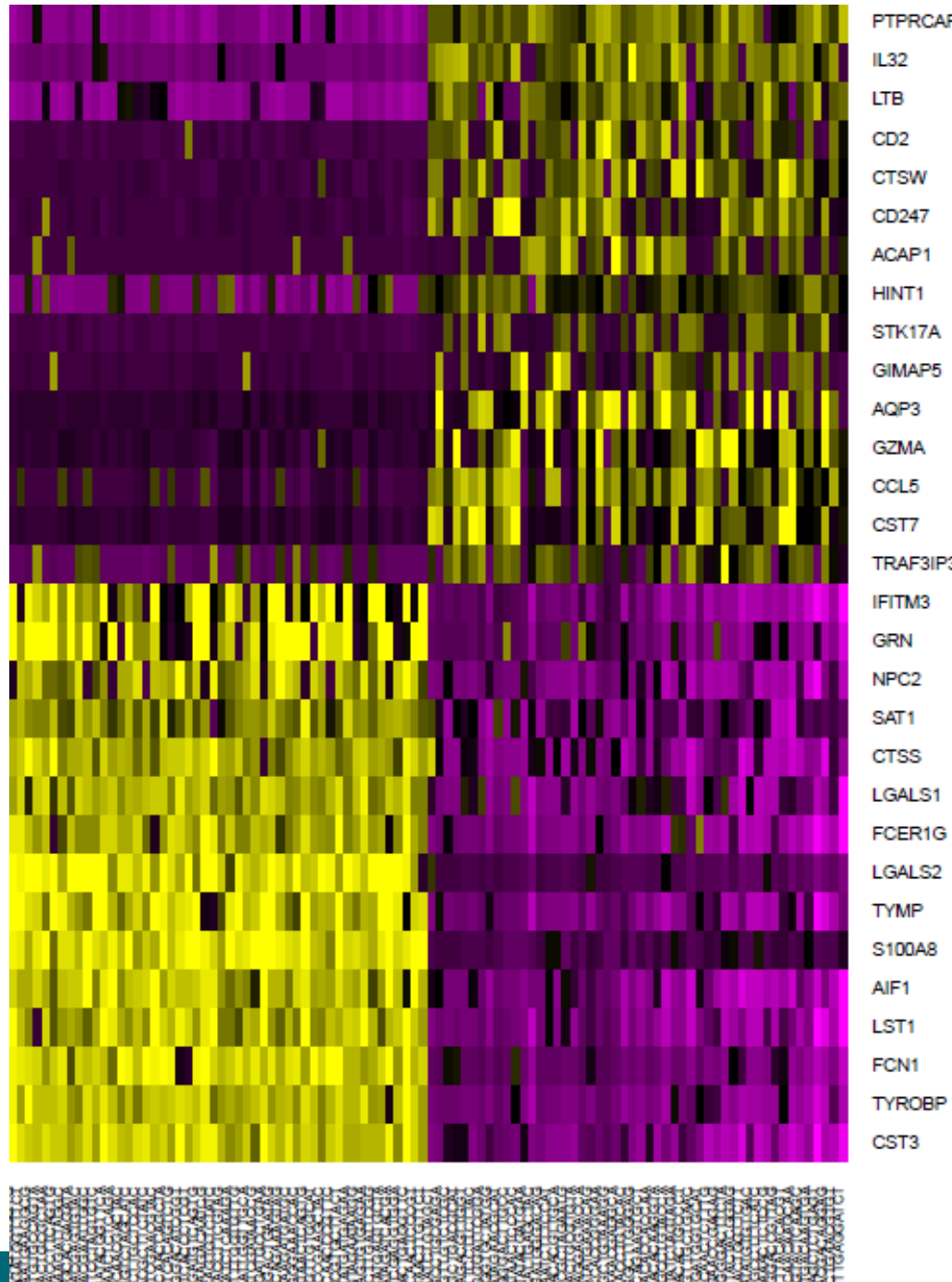
- Visualize top genes associated with principal components
 - = Which genes are important for PC1?)



- Is the correlation direct (positive) or reverse (negative)?
 - Note however, that the signs are arbitrary
 - = if all the variables in a component are positively correlated with each other, all the loadings will be positive
 - if there are some negative correlations among the variables, some of the loadings will be negative

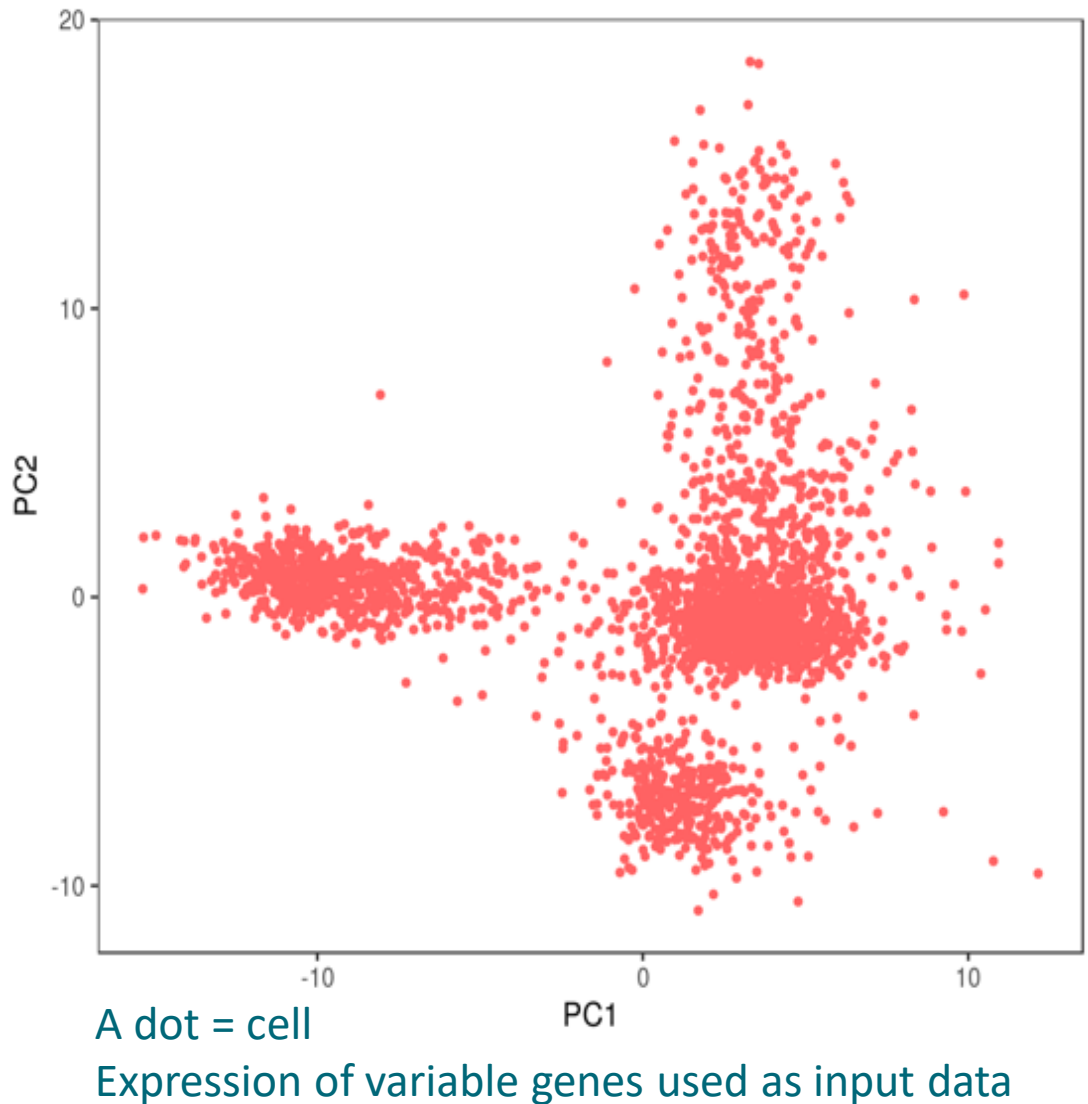
Visualizing PCA results: heatmap

- Which genes correspond to separating cells?
 - Check if there are cell cycle genes
 - Note: after clustering and DE gene analysis steps, we can also eyeball this from the data visualisations
- Both cells and genes are ordered according to their PCA scores. Plots the extreme cells on both ends of the spectrum



Visualizing PCA results: PCA plot

- Gene expression patterns will be captured by PCs → PCA *can* separate cell types
- Note however that PCA can also capture other things, like sequencing depth, cell size or cell heterogeneity/complexity!

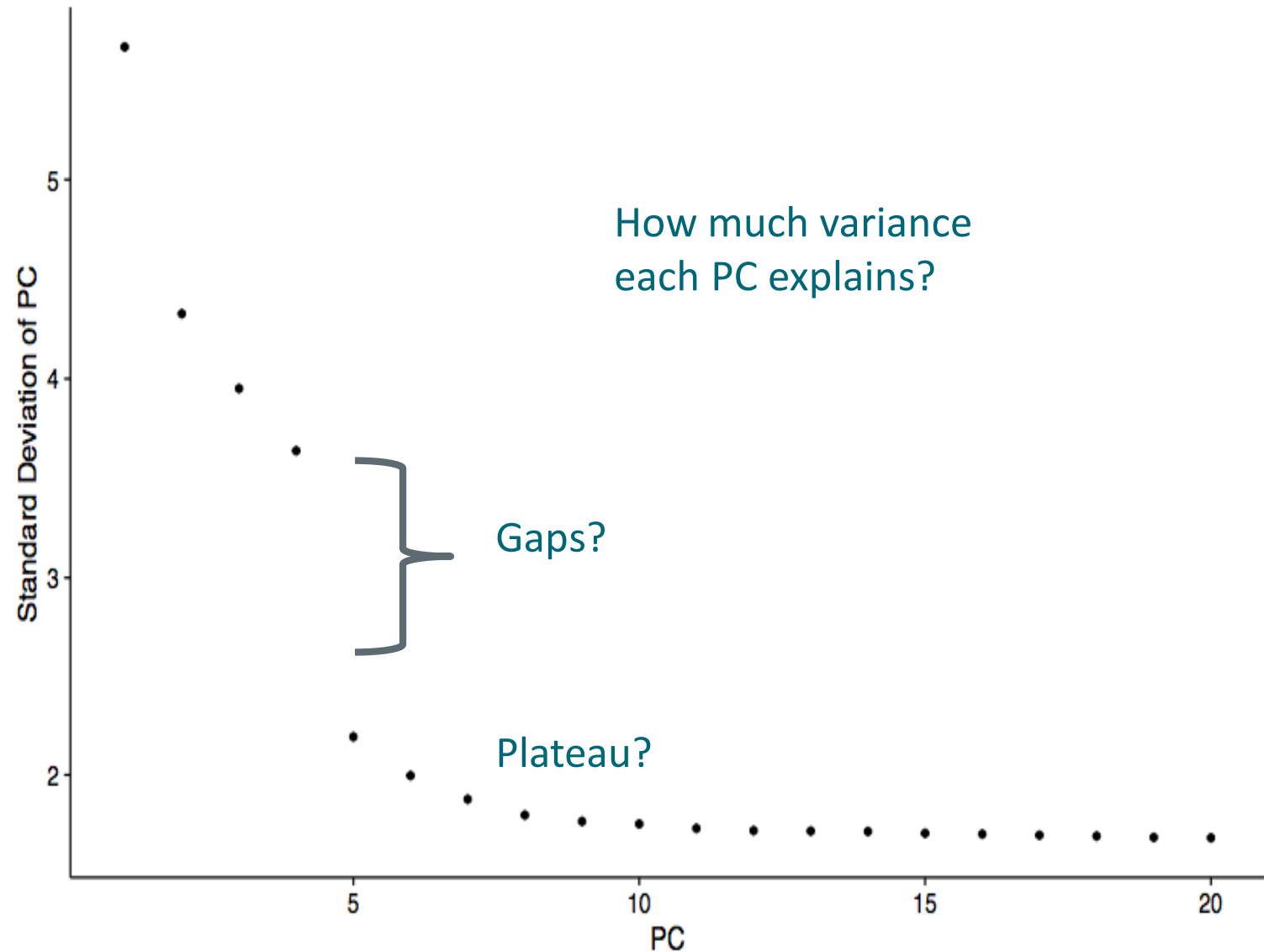


Determine the significant principal components

- It is important to select the significant PCs for clustering analysis
- However, estimating the true dimensionality of a dataset is challenging
- Seurat developers:
 - Try repeating downstream analyses with a different number of PCs (10, 15, or even 50!).
 - The results often do not differ dramatically.
 - Rather choose higher number.
 - For example, choosing 5 PCs does significantly and adversely affect results
- Chipster provides the following plots to guide you selecting the significant PCs:
 - Elbow plot
 - PC heatmaps

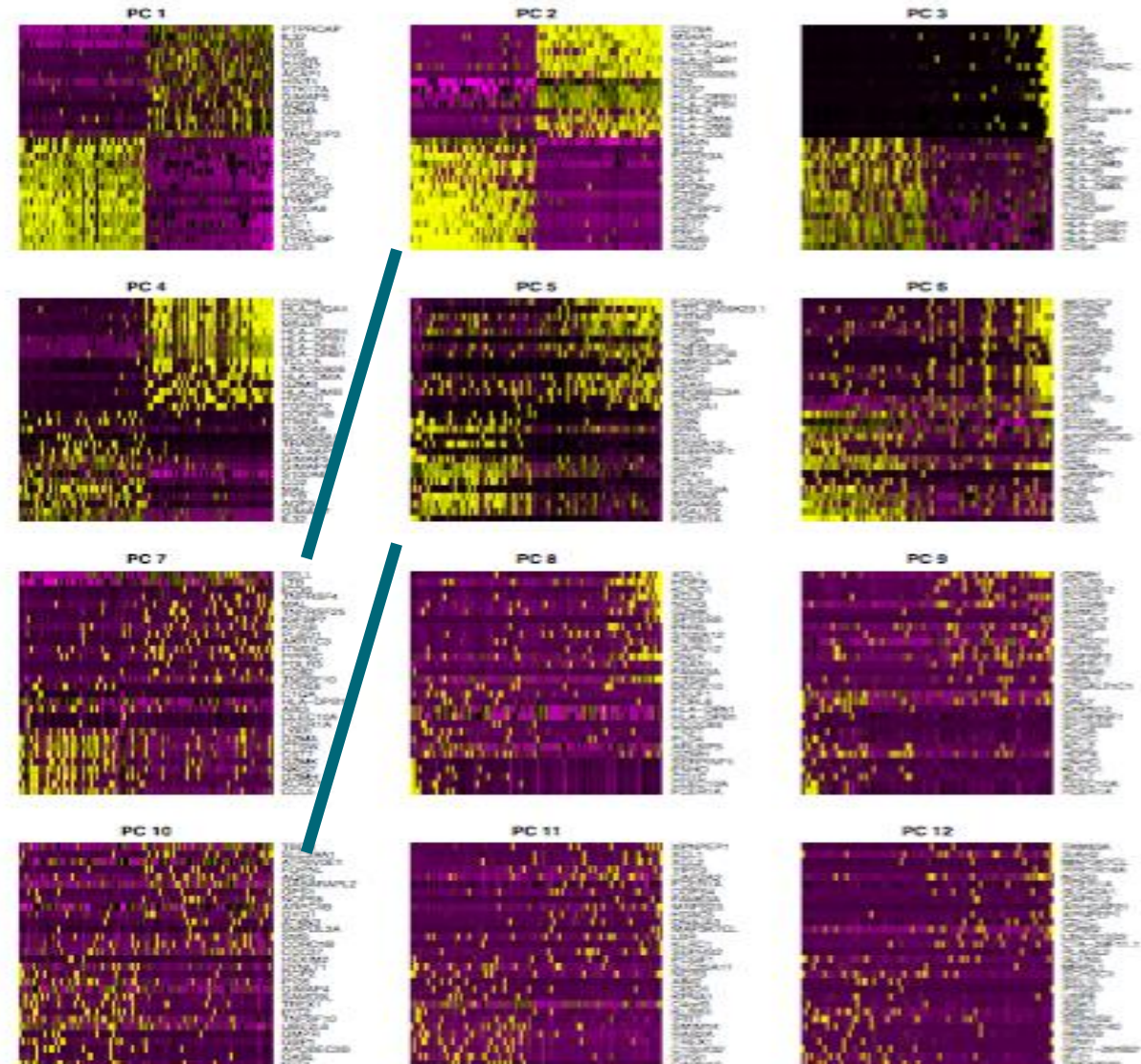
Elbow plot

- The elbow in the plot tends to reflect a transition from informative PCs to those that explain comparatively little variance.



Principal component heatmaps

- Check if there is still a difference between the extremes
- Exclude also PCs that are driven primarily by uninteresting genes (cell cycle, ribosomal or mitochondrial)



Other dimension reduction methods: used later for visualisation

- Graph-based, non-linear methods like tSNE and UMAP
- PCA, tSNE and UMAP available as options in most tools
- We use PCA for dimension reduction before clustering, and tSNE or UMAP for visualisation

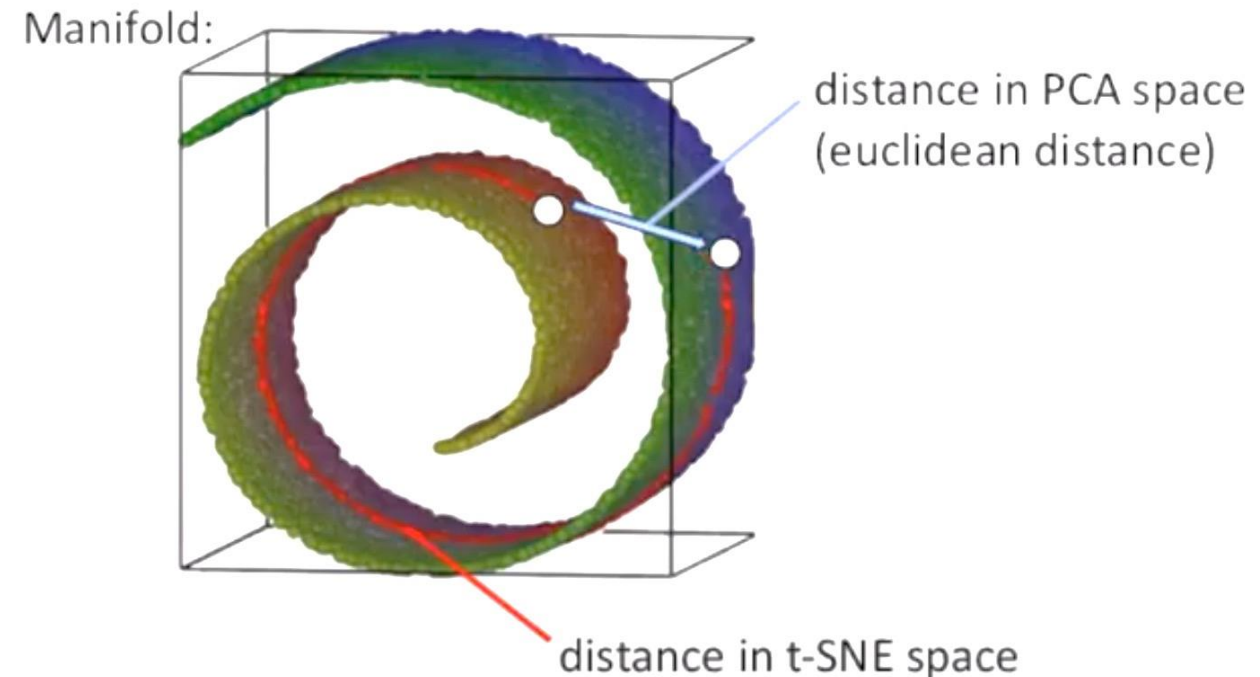
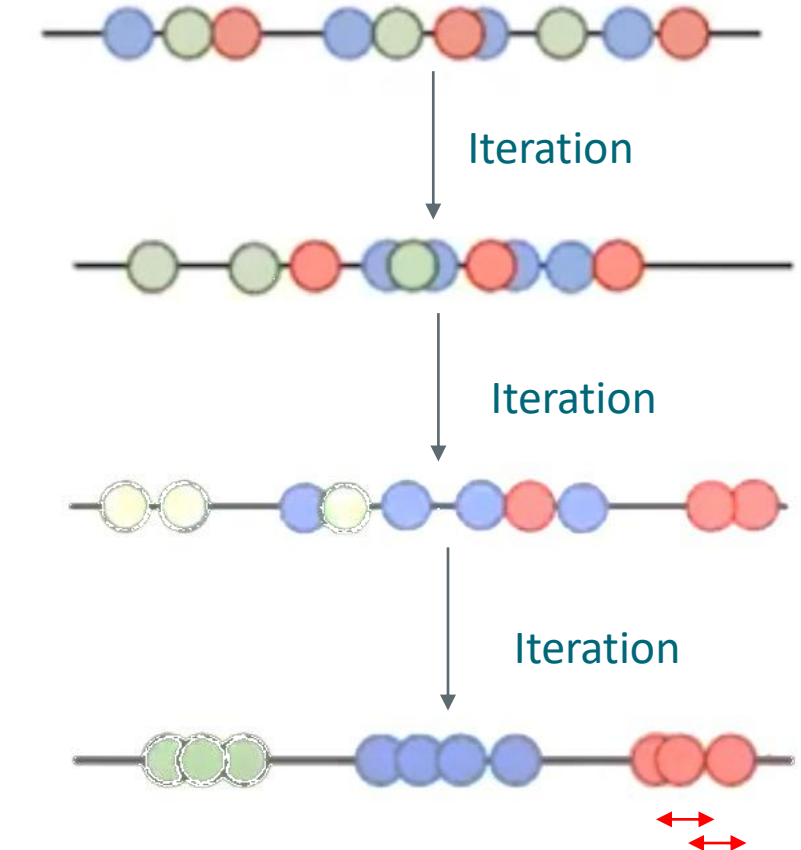
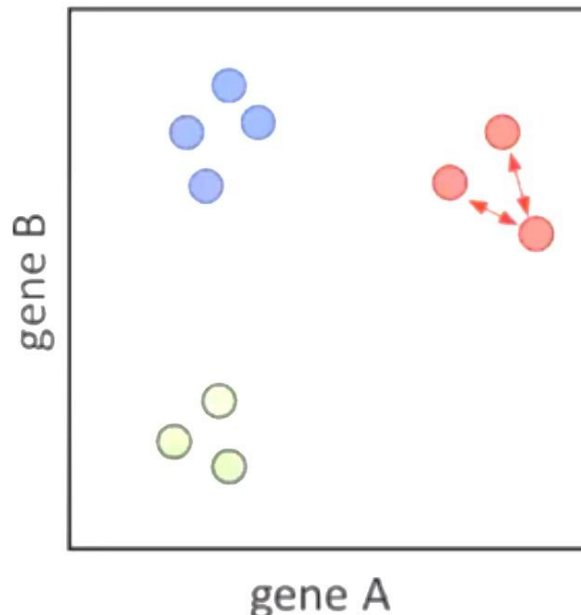


Image by Shigeo Takahashi et al, <http://web-ext.u-aizu.ac.jp/~shigeo/research/manifold/>

tSNE simplified

- Graph-based
- Non-linear
- Stochastic
- (Only) local distances preserved: distance between groups are not meaningful
- Gold standard
- Can be run on top of PCs
- Many parameters to optimize

Example: From 2D to 1D



Slide modified from Paulo Czarnewski's slides, image based on StatQuest

UMAP

- Non-linear graph-based dimension reduction method like tSNE
- Newer & efficient = fast
- Runs on top of PCs
- Based on topological structures in multidimensional space
- Unlike tSNE, you can compute the structure once (no randomization)
 - => faster
 - => you could add data points without starting over
- **Preserves the global structure** better than tSNE
- More info: video 6 at bit.ly/scRNA-seq
 - Dimensionality reduction explained by Paulo Czarnewski

Analysis steps for clustering cells and finding marker genes



1. Create Seurat object, filter genes, check the quality of cells
2. Filter out low quality cells
3. Normalize expression values
4. Identify highly variable genes
5. Scale data, regress out unwanted variation
6. Reduce dimensions using principal component analysis (PCA) on the variable genes
7. Determine significant principal components (PCs)
8. Use the PCs to cluster cells with graph based clustering
9. Visualize clusters with non-linear dimensional reduction (tSNE or UMAP) using the PCs
10. Detect and visualize marker genes for the clusters

What will you learn

1. Why is clustering a bit complex step?
2. What happens in the clustering step?
3. How to visualise the clusters

Clustering

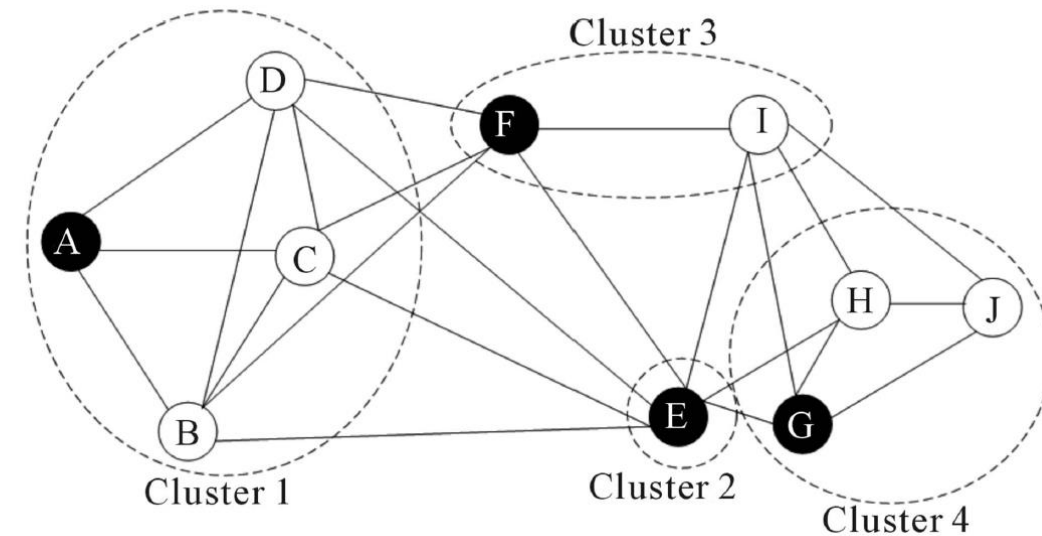
- Divides cells into distinct groups based on gene expression
- Our data is big and complex (lot of cells, genes and noise), so we use principal components instead of genes. We also need a clustering method that can cope with this.

→ Graph-based clustering

→ Shared nearest neighbor approach

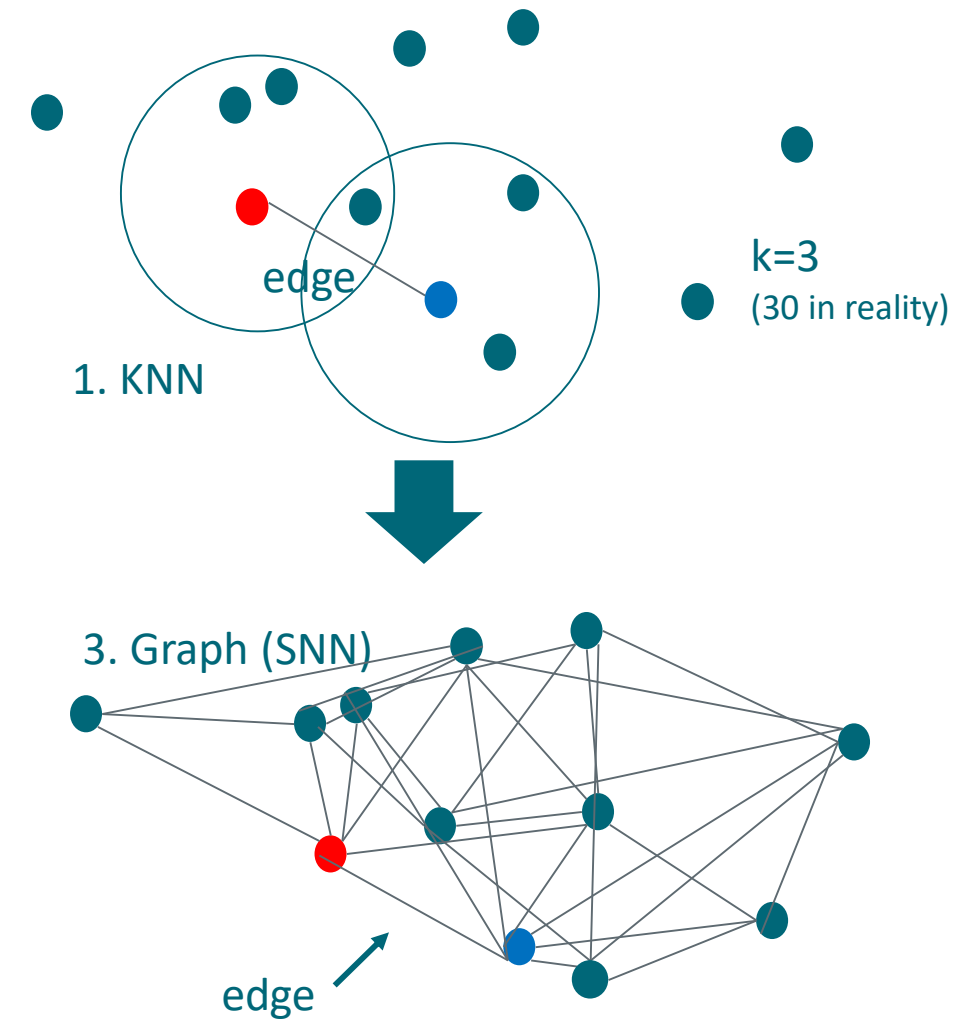
→ Graph cuts by Louvain method

Nodes → cells
Edges → similarity



Graph based clustering in Seurat

1. Identify k nearest neighbours of each cell
 - o Euclidean distance in PC space
2. Rank the neighbours based on distance
3. Build the graph: add an edge between cells if they have a shared nearest neighbour (SNN)
 - o Give edge weights based on ranking
4. Cut the graph to subgraphs (clusters) by optimizing modularity
 - o *Louvain algorithm* by default



Clustering parameters



- Number of principal components to use
 - Experiment with different values
 - If you are not sure, use a higher number
- Resolution for granularity
 - Increasing leads to more clusters
 - Values 0.4 - 1.2 can return good results for single cell datasets of around 3000 cells
 - Higher resolution is often optimal for larger datasets

Seurat v4 -Clustering

Parameters

Change this, if you used SCTransform

Normalisation method used previously

Which normalisation method was used in preprocessing, Global scaling normalization (default, NormalizeData function used) or SCTransform.

Number of principal components to use

How many principal components to use. User must define this based on the PCA-elbow and PCA plots from the setup tool. Seurat developers encourage to test with different parameters, and use preferably more than less PCs for downstream analysis.

Resolution for granularity

Resolution parameter that sets the granularity of the clustering. Increased values lead to greater number of clusters. Values between 0.6-1.2 return good results for single cell datasets of around 3K cells. For larger data sets, try higher resolution.

Perplexity, expected number of neighbors for tSNE plot

Perplexity, expected number of neighbors. Default 30. Set to lower number if you have very few cells. Used for the tSNE visualisation of the clusters.

Point size in tSNE and UMAP plots

Point size for the cluster plots.

Add labels on top of clusters in plots

Add cluster number on top of the cluster in UMAP and tSNE plots.

Give a list of average expression in each cluster

Returns an expression table for an 'average' single cell in each cluster.

Global scaling normalization

10

0.5

30

1

yes

no

Change this, if you have small data:

Visualization of clusters: tSNE or UMAP

- tSNE/UMAP plot is gray by default, we color it by clustering results from the previous step
 - Check how well the groupings found by tSNE/UMAP match with cluster colors
- Input data: same PCs as for the clustering
- 2 parameters:

Perplexity, expected number of neighbors for tSNE plot

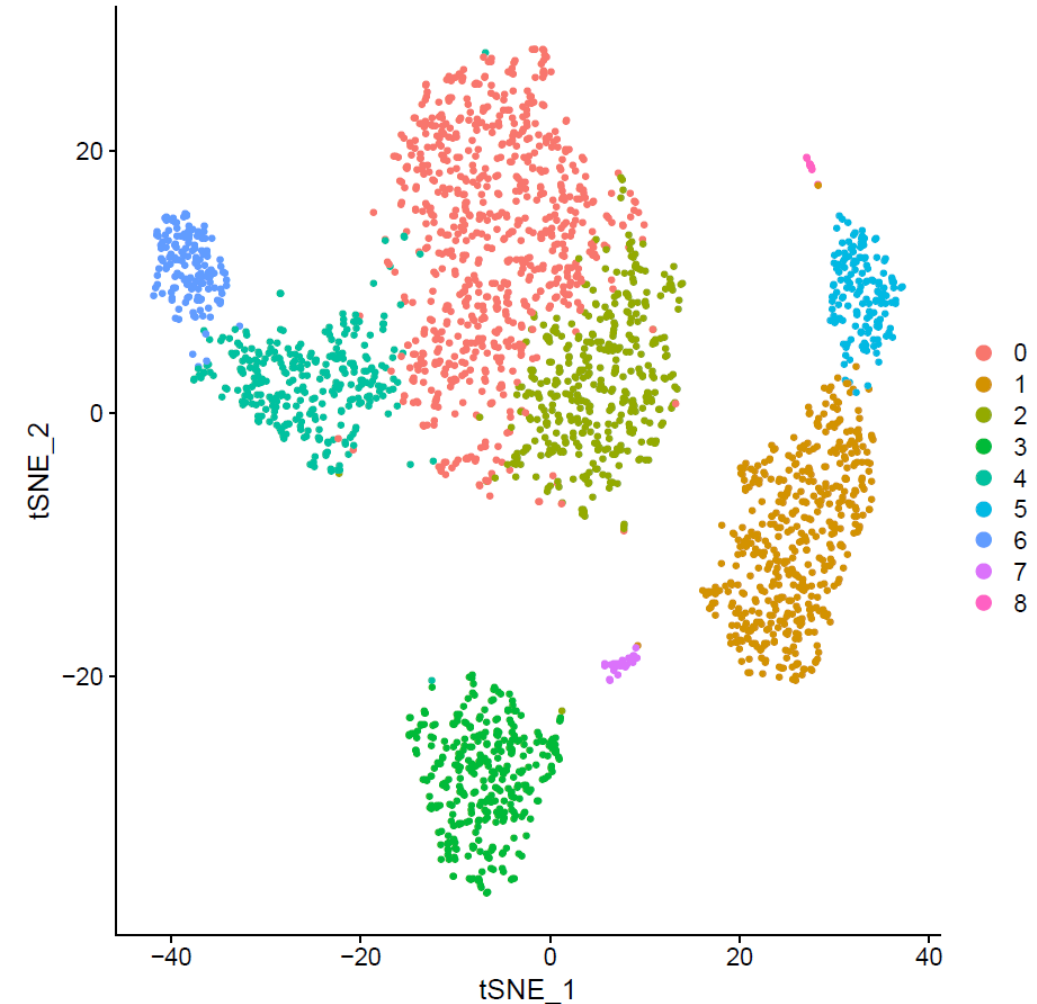
30

Perplexity, expected number of neighbors. Default 30. Set to lower number if you have very few cells. Used for the tSNE visualisation of the clusters.

Point size in tSNE and UMAP plots

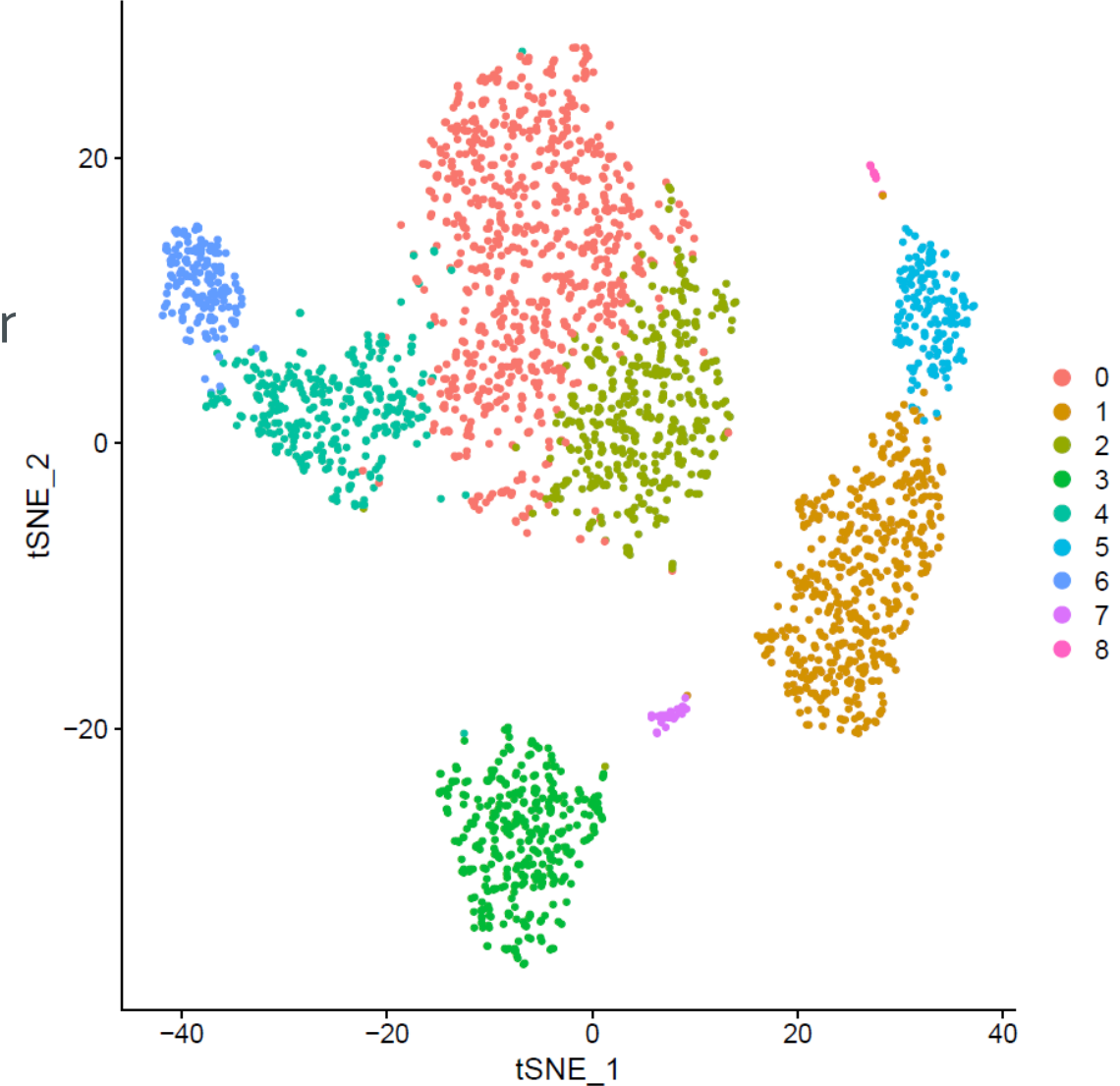
1

Point size for the cluster plots.



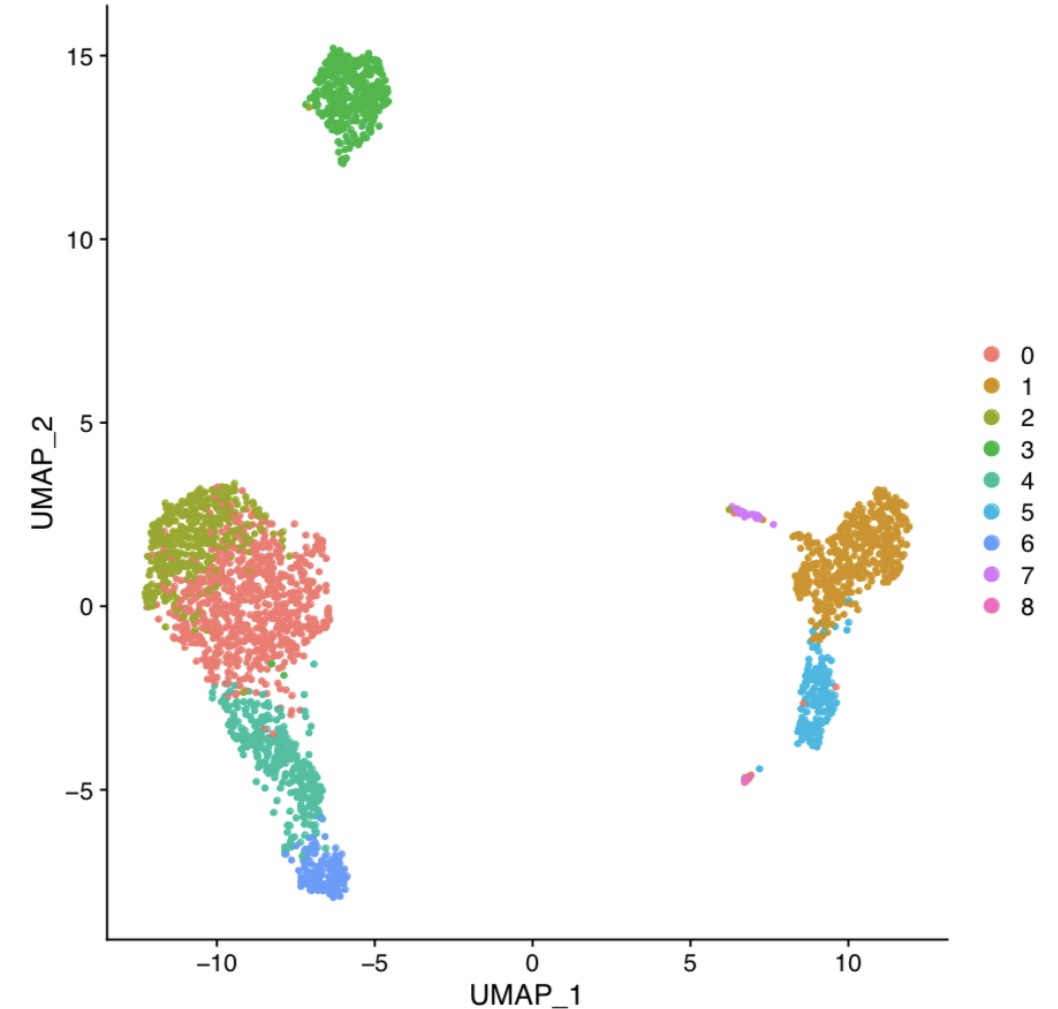
tSNE plot for cluster visualization

- t-distributed Stochastic Neighbor Embedding
- Graph-based non-linear dimensional reduction
 - Different transformations to different regions
- Specialized in local embedding
 - Distance between clusters is not meaningful
 - <https://distill.pub/2016/misread-tsne/>
- Perplexity = number of neighbors to consider
 - Default 30, lower for small datasets



UMAP plot for cluster visualization

- UMAP = Uniform Manifold Approximation and Projection
- Non-linear graph-based dimension reduction method like tSNE
 - Preserves more of the global structure than tSNE



Analysis steps for clustering cells and finding marker genes



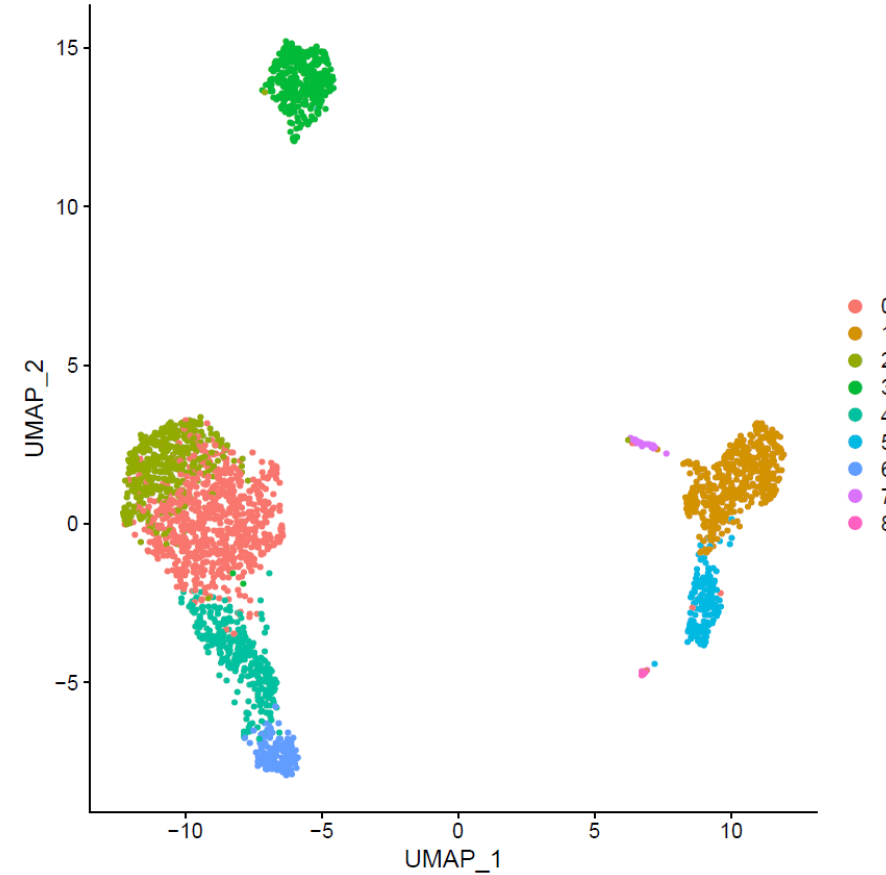
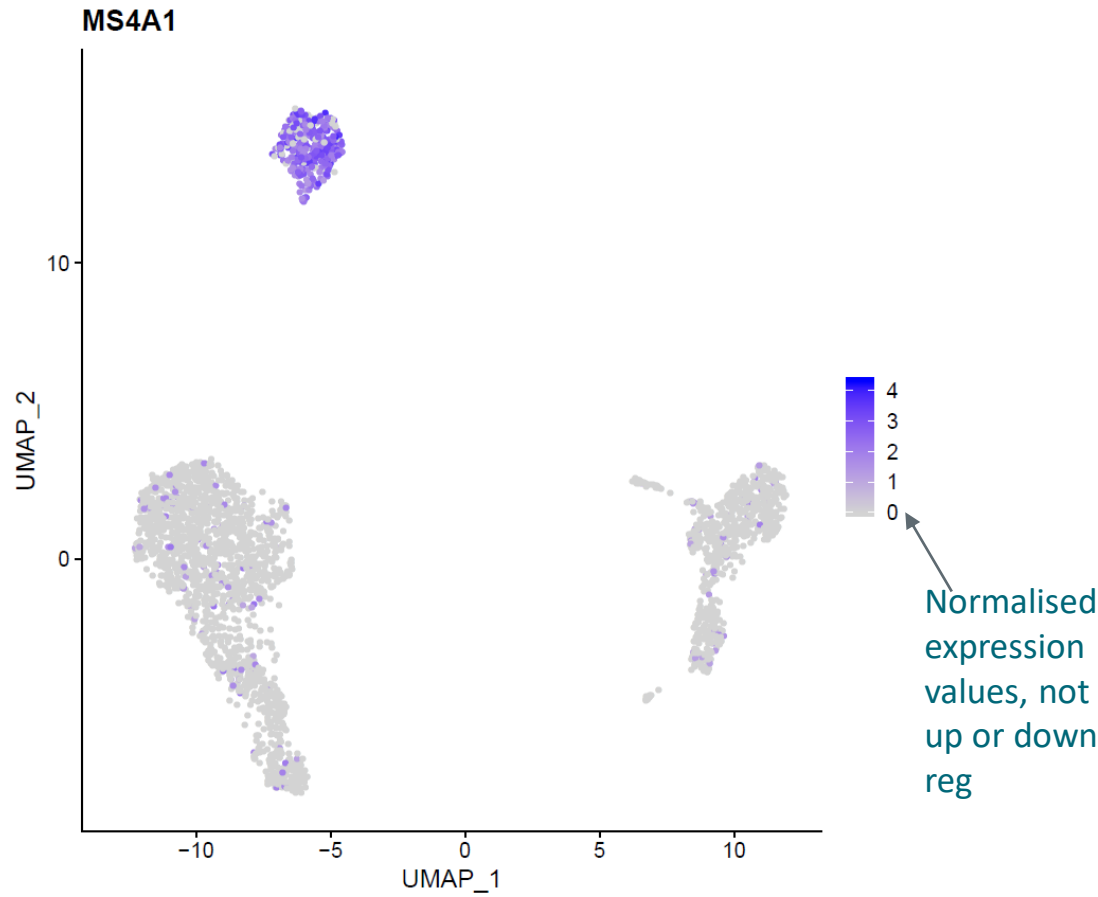
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- 10. Detect and visualize marker genes for the clusters**

What will you learn

1. What is a marker gene
2. What aspects of scRNA-seq data complicate differential expression analysis
3. Why do we want to filter out genes prior to statistical testing

Marker gene for a cluster

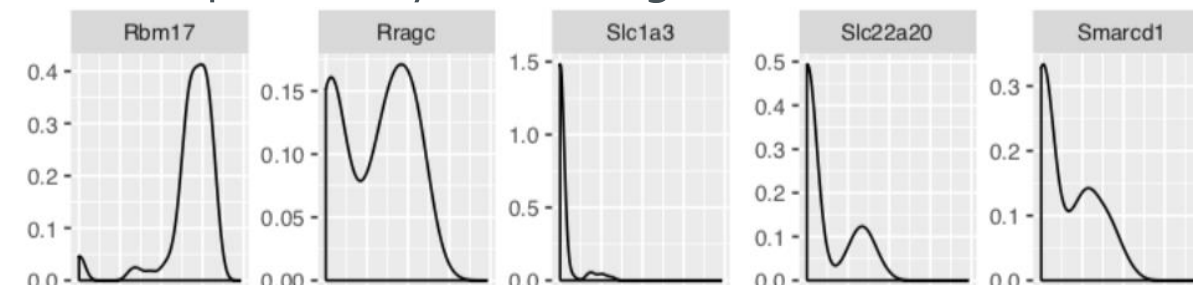
- Differentially expressed between the cluster and all the other cells



Differential expression analysis of scRNA-seq data



- Challenging because the data is noisy
 - low amount of mRNA → low counts, high dropout rate, amplification biases
 - uneven sequencing depth
- Non-parametric tests, e.g. Wilcoxon rank sum test (Mann-Whitney U test)
 - Can fail in the presence of many tied values, such as the case for dropouts (zeros) in scRNA-seq
- Methods specific for scRNA-seq, e.g. MAST
 - Take advantage of the large number of samples (cells) for each group
 - MAST accounts for stochastic dropouts and bimodal expression distribution
- Methods for bulk RNA-seq, e.g. DESeq2
 - Based on negative binomial distribution, works ok for UMI data.
 - Note: you should *not* filter genes, because DESeq2 models dispersion by borrowing information from other genes with similar expression level
 - Very slow! Use only for comparing 2 clusters



Wilcoxon rank-sum test

Gene A

Cluster1	Rank1	Cluster2	Rank2
21	7	5	1.5
5	1.5	13	6
29	8	6	3
10	5	8	4
	21.5		14.5



values	Rank
5	1.5
5	1.5
6	3
8	4
10	5
13	6
21	7
29	8

$$U\text{-stat} = \frac{\text{Rank sum} - n(n+1)}{2}$$

$$UA = 21.5 - 4(4+1)/2 = 21.5 - 10 = 11.5$$

$$UB = 17.5 - 4(4+1)/2 = 14.5 - 10 = 4.5$$

U-stat = 4.5 (use the smallest from above)

U-critical = 0 (for alpha=0.05)

U-stat > U-critical (no significant difference)

P-value = 0.342857

Filtering out genes prior to statistical testing – why?



- We test thousands of genes, so it is possible that we get good p-values just by chance (false positives)
 - Multiple testing correction of p-values is needed
 - The amount of correction depends on the number of tests (= genes)
 - Bonferroni correction: adjusted p-value = raw p-value * number of genes tested
 - If we test less genes, the correction is less harsh → better p-values
- Filtering also speeds up testing

Detect cluster marker genes

Find all markers = every cluster is compared to all the other cells

OR

Compare the cluster of interest to all others or to another cluster

- Limit testing to genes which
 - are expressed in at least this fraction of cells in either of the two groups (default 10%)
 - show at least this **log₂** fold change between the two groups (default 0.25)

Parameters

 Reset All

FALSE

Find all markers

Give as an output a large table with markers for all the clusters. Each cluster is compared to all the other clusters. This parameter overwrites the two cluster number parameters below. You will want to filter this table with the tool in Utilities category.

1

Cluster of interest

The number of the cluster of interest.

1

Cluster to compare with

Number(s) of the cluster(s) to compare to. By default the cluster of interest is compared to cells in all other clusters. You can also compare to another cluster or a group of clusters, just separate the cluster numbers with a comma.

all others

Limit testing to genes which are expressed in at least this fraction of cells

Test only genes which are detected in at least this fraction of cells in either of the two populations. Meant to speed up testing by leaving out genes that are very infrequently expressed.

0.1

Limit testing to genes which show at least this fold difference

Test only genes which show on average at least this log₂ fold difference, between the two groups of cells. Increasing the threshold speeds up testing, but can miss weaker signals.

0.25

Which test to use for detecting marker genes

Seurat currently implements Wilcoxon rank sum test, bimod (likelihood-ratio test for single cell gene expression), roc (standard AUC classifier), Students t-test, Tobit-test, MAST (GLM-framework that treats cellular detection rate as a covariate), poisson, negbinom and DESeq2. The latter three should be used on UMI datasets only, and assume an underlying poisson or negative-binomial distribution. Note that DESeq2 is very slow and should be used only for comparisons between two clusters.

wilcox

Report only positive marker genes

When this parameter is set to true, only genes with positive log₂ fold change are listed in the result file.

TRUE

Cluster marker gene result table



- p_val = p-value
- p_val_adj = p-value adjusted using the Bonferroni method
- avg_logFC = \log_2 fold change between the groups
- cluster = cluster number
- pct1 = percentage of cells where the gene is detected in the first group

markers.tsv ***

Spreadsheet Text Details

Showing the first 100 of 477 rows. View in [full screen](#) to see all rows.

	p_val	avg_log2FC	pct.1	pct.2	p_val_adj	cluster	gene
LTB	6.219516e-123	1.348424	0.958	0.600	8.529445e-119	0	LTB
IL32	3.455676e-113	1.186582	0.893	0.413	4.739115e-109	0	IL32
LDHB	1.155309e-111	1.059929	0.913	0.578	1.584391e-107	0	LDHB
CD3D	1.235473e-109	1.113704	0.872	0.376	1.694328e-105	0	CD3D
IL7R	2.138245e-94	1.278283	0.699	0.281	2.932389e-90	0	IL7R
CD2	1.398161e-60	1.141928	0.551	0.223	1.917438e-56	0	CD2
S100A9	0.000000e+00	5.563093	0.996	0.216	0.000000e+00	1	S100A9
S100A8	0.000000e+00	5.482122	0.973	0.122	0.000000e+00	1	S100A8
LGALS2	0.000000e+00	3.804741	0.908	0.060	0.000000e+00	1	LGALS2
FCN1	0.000000e+00	3.390813	0.952	0.151	0.000000e+00	1	FCN1

How to filter the gene list?

- You can filter the result table for example based on the adjusted p-value column using the tool **Utilities / Filter table by column value** using the following parameters:

Parameters

Column to filter by

p_val_adj

Data column to filter by

Does the first column lack a title

Specifies whether the first column has a title or not.

yes

Cut-off value

0.05

Cut-off for filtering

Filtering criteria

smaller-than

Smaller or larger than the cutoff is filtered. Use the "within" or "outside" options to filter symmetrically around two cut-offs, useful for example when searching for up- and down-regulated genes.

How to retrieve marker genes for a particular cluster?

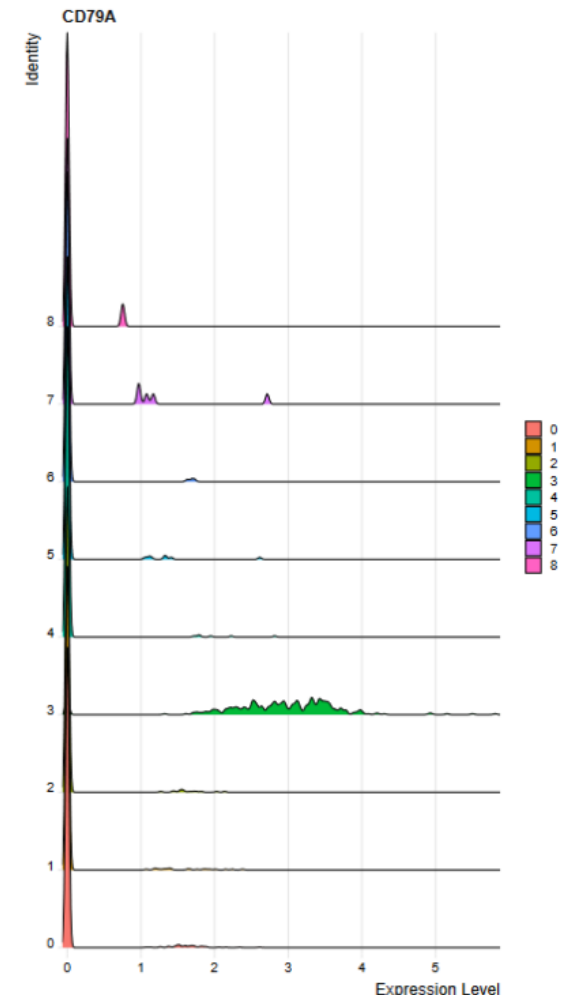
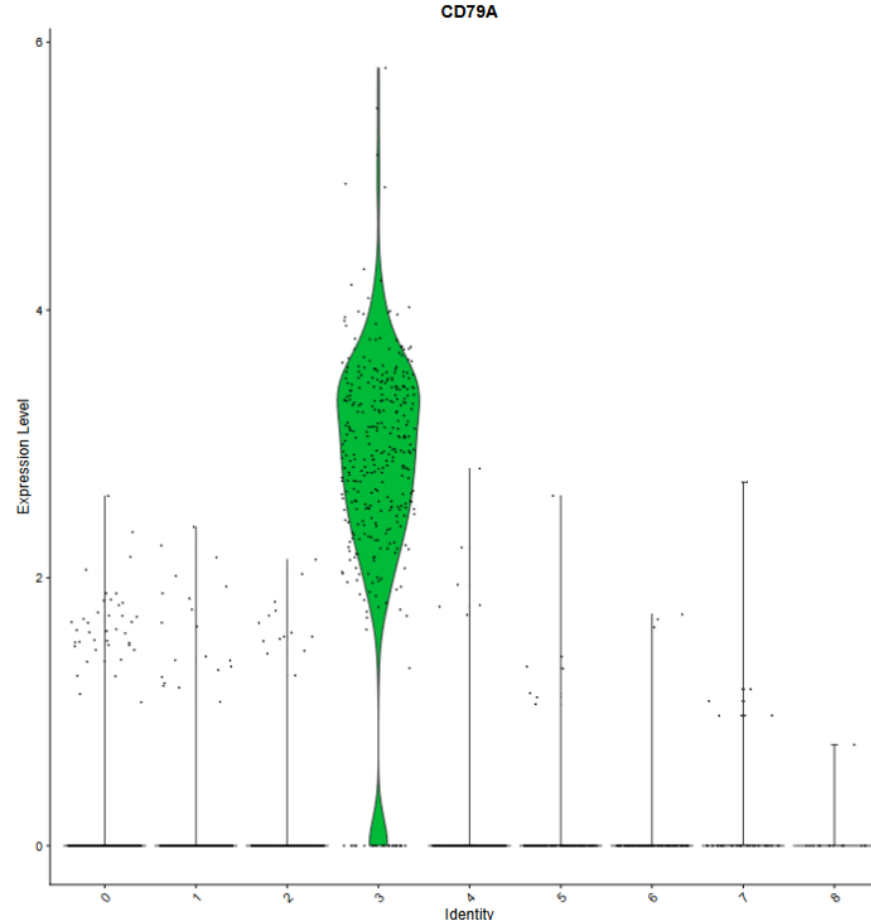
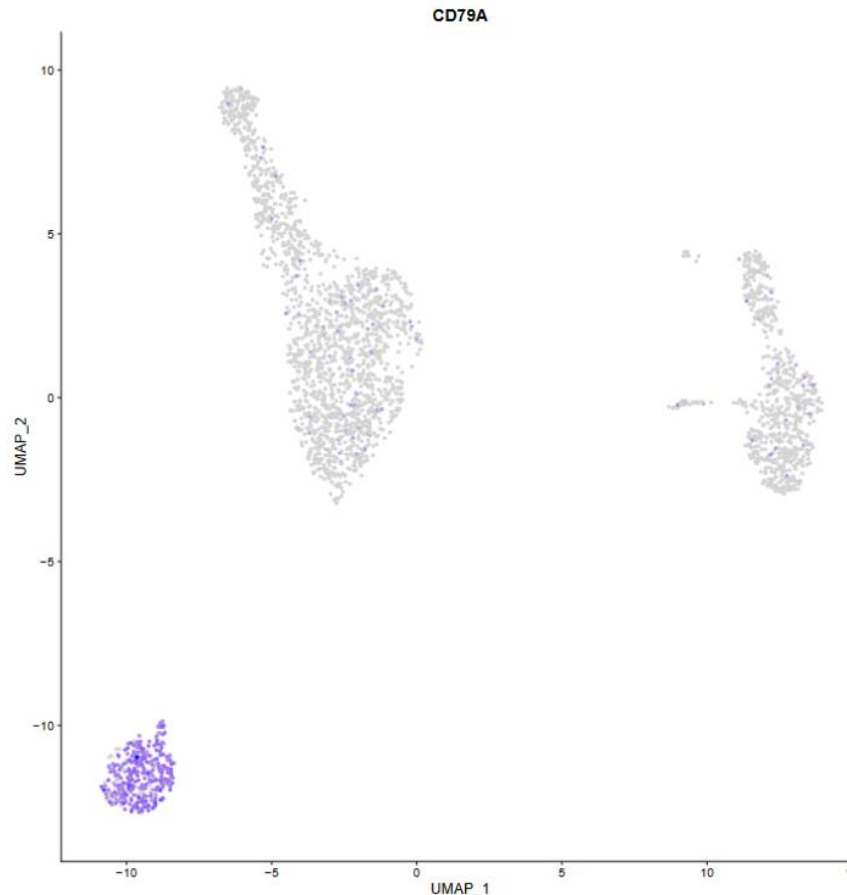


- If you had set Find all markers = TRUE, the result table contains marker genes for all the clusters
- You can filter the result table based on the cluster column using the tool **Utilities / Filter table by column value** using the following parameters

Parameters	
Column to filter by	<input type="text" value="cluster"/>  
Data column to filter by	
Does the first column lack a title	<input type="text" value="yes"/>  
Specifies whether the first column has a title or not.	
Cut-off value	<input type="text" value="3"/>  
Cut-off for filtering	
Filtering criteria	<input type="text" value="equal-to"/>  
Smaller or larger than the cutoff is filtered. Use the "within" or "outside" options to filter symmetrically around two cut-offs, useful for example when searching for up- and down-regulated genes.	

Visualize cluster marker genes

- UMAP, tSNE or PCA plot colored with marker gene expression
- Violin plot
- Ridge plot



Tool “Visualize genes”



Seurat v4 -Visualize genes X

Parameters

Gene name(s) Reset All
Name(s) of the biomarker gene to plot. If you list multiple gene names, use comma (,) as separator.

Point size in cluster plot ^
Point size for tSNE and UMAP plots.

Add labels on top of clusters in plot ▼
Add cluster number on top of the cluster in UMAP plot.

Visualisation with tSNE, UMAP or PCA ▼
Which dimensionality reduction plot to use.

Plotting order of cells based on expression ▼
Plot cells in the order of expression. Can be useful to turn this on if cells expressing given feature are getting buried.

For each gene, list the average expression and percentage of cells expressing it in each cluster ▼
Returns two tables: average expression and percentage of cells expressing the user defined genes in each cluster.

Input files

Seurat object ▼

Optional text file of the gene name(s) ▼
The gene name(s) you wish to plot can also be given in the form of a text file, separated by comma. In case the text file is provided, the gene parameter is ignored.

Result tables



- Gene's average expression level in each cluster
- Percentage of cells expressing the gene in each cluster

percentage_of_cells_expressing.tsv ...									
Spreadsheet Text Open in New Tab Details									
Showing all 3 rows.									
	0	1	2	3	4	5	6	7	8
MS4A1	4.56	5.64	5.29	86.01	4.61	8.18	5.56	3.23	7.14
LYZ	50.98	100	49.37	42.57	41.78	98.74	43.06	96.77	50
PF4	0.13	1.67	0.5	1.46	0.99	3.77	0	6.45	100

average_expressions.tsv ...									
Spreadsheet Text Open in New Tab Details									
Showing all 3 rows.									
	0	1	2	3	4	5	6	7	8
MS4A1	0.192	0.217	0.221	11.749	0.265	0.256	0.255	0.063	0.469
LYZ	3.211	183.343	2.874	3.223	2.688	30.414	2.892	127.022	11.558
PF4	0.006	0.099	0.022	0.059	0.111	0.152	0	0.216	158.976

Extract information from Seurat R-object

- Seurat R-object consists of specific data slots which contain more slots
- Chipster tool “Extract information from Seurat object” allows you to check
 - what the scRNA-seq data set includes
 - How many cells and genes
 - What genes
 - What are the highly variable genes
 - whether the data has already been normalised and how (SCTransform or global scaling)
 - which Seurat functions were used
- If you get a Seurat object from somebody and import it to Chipster, you can see what has been done

Result tables

- Text file including the different slots in the object such as the counts and assays
- Meta data table containing additional information associated with the cells or features of the object

slots.txt

File size 1.0 kB.

```
[1] "Assays in the seurat object: "
$RNA
Assay data with 13714 features for 2700 cells
First 10 features:
AL627309.1, AP006222.2, RP11-206L10.2, RP11-206L10.9, L
KLHL17, PLEKHN1, RP11-5407.17, HES4

[1] "Active assay in the object: "
[1] "RNA"
[1] "Active cluster identity in the cluster: "
AACATACAACCAC-1 AAACATTGAGCTAC-1 AAACATTGATCAGC-1 AAACC
PBMC PBMC PBMC
AAACCGTGTATGCG-1 AAACGCACTGGTAC-1
PBMC PBMC

Levels: PBMC
[1] "List of graph objects in the seurat object: "
list()
[1] "List of neighbor objects in the seurat object: "
list()
[1] "List of dimensional reductions for this object: "
list()
[1] "List of spatial image objects in this object: "
list()
[1] "Name of the project: "
[1] "PBMC"
[1] "A list of miscellaneous information in the Seurat object: "
list()
[1] "Version of Seurat this object was built under: "
[1] '4.1.1'
[1] "A list of logged commands run on this Seurat object: "
list()
[1] "A list of miscellaneous data generated by other tools: "
list()
```

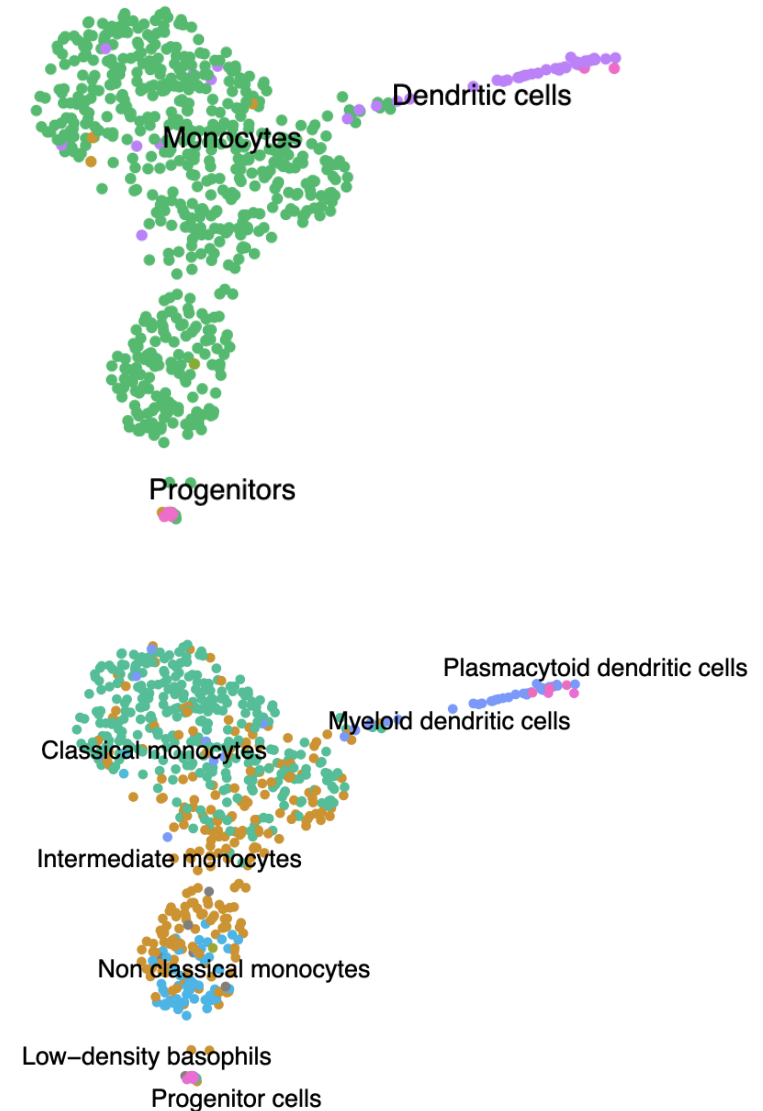
meta_data.tsv

Showing all 2700 rows.

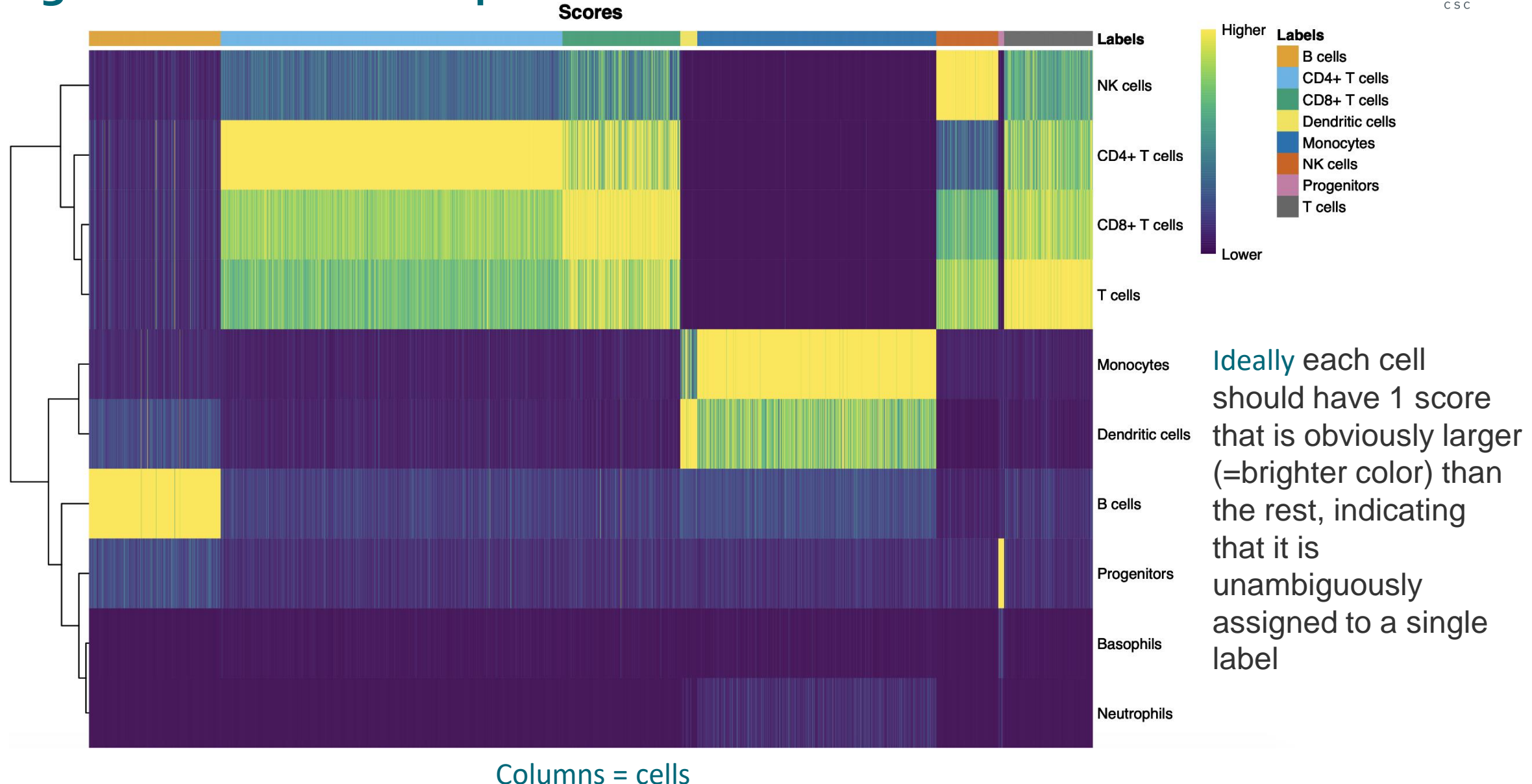
	orig.ident	nCount_RNA	nFeature_RNA	percent.mt
AAACATACAACCAC-1	PBMC	2419	779	3.0177759
AAACATTGAGCTAC-1	PBMC	4903	1352	3.7935958
AAACATTGATCAGC-1	PBMC	3147	1129	0.8897363
AAACCGTGTCTCCG-1	PBMC	2639	960	1.7430845
AAACCGTGTATGCG-1	PBMC	980	521	1.2244898
AAACGCACTGGTAC-1	PBMC	2163	781	1.6643551
AAACGCTGACCACT-1	PBMC	2175	782	3.8160920
AAACGCTGGTTCTT-1	PBMC	2260	790	3.0973451
AAACGCTGTAGCCA-1	PBMC	1275	532	1.1764706
AAACGCTGTTCTG-1	PBMC	1103	550	2.9011786
AAACTTGAAAACG-1	PBMC	3914	1112	2.6315789
AAACTTGATCCAGA-1	PBMC	2388	747	1.0887772
AAAGAGACGAGATA-1	PBMC	2410	864	1.0788382
AAAGAGACGCGAGA-1	PBMC	3033	1058	1.4177382
AAAGAGACGGACTT-1	PBMC	1151	457	2.3457863
AAAGAGACGGCATT-1	PBMC	792	335	2.3989899
AAAGATCTGGCAA-1	PBMC	1347	551	5.9391240
AAAGCAGAAGCCAT-1	PBMC	1158	567	5.0949914
AAAGCAGATATCGG-1	PBMC	4584	1422	1.3961606
AAAGCCTGTATGCG-1	PBMC	2928	1013	1.7076503
AAAGGCTGTCTAG-1	PBMC	4973	1445	1.5282526
AAAGTTTGATCACG-1	PBMC	1268	444	3.4700315
AAAGTTTGGGTGA-1	PBMC	3281	1015	2.5906736
AAAGTTTGTAGAGA-1	PBMC	1102	417	1.5426497
AAAGTTTGTACCGT-1	PBMC	2683	877	2.4972046
AAATCAACAATGCC-1	PBMC	2319	787	1.1642950
AAATCAACACCAGT-1	PBMC	1412	508	1.9830028
AAATCAACCAGGAG-1	PBMC	2800	823	2.2500000
AAATCAACCCATT-1	PBMC	5676	1541	2.4312896
AAATCAACGGAAGC-1	PBMC	3473	996	1.7564066

SingleR annotations to clusters

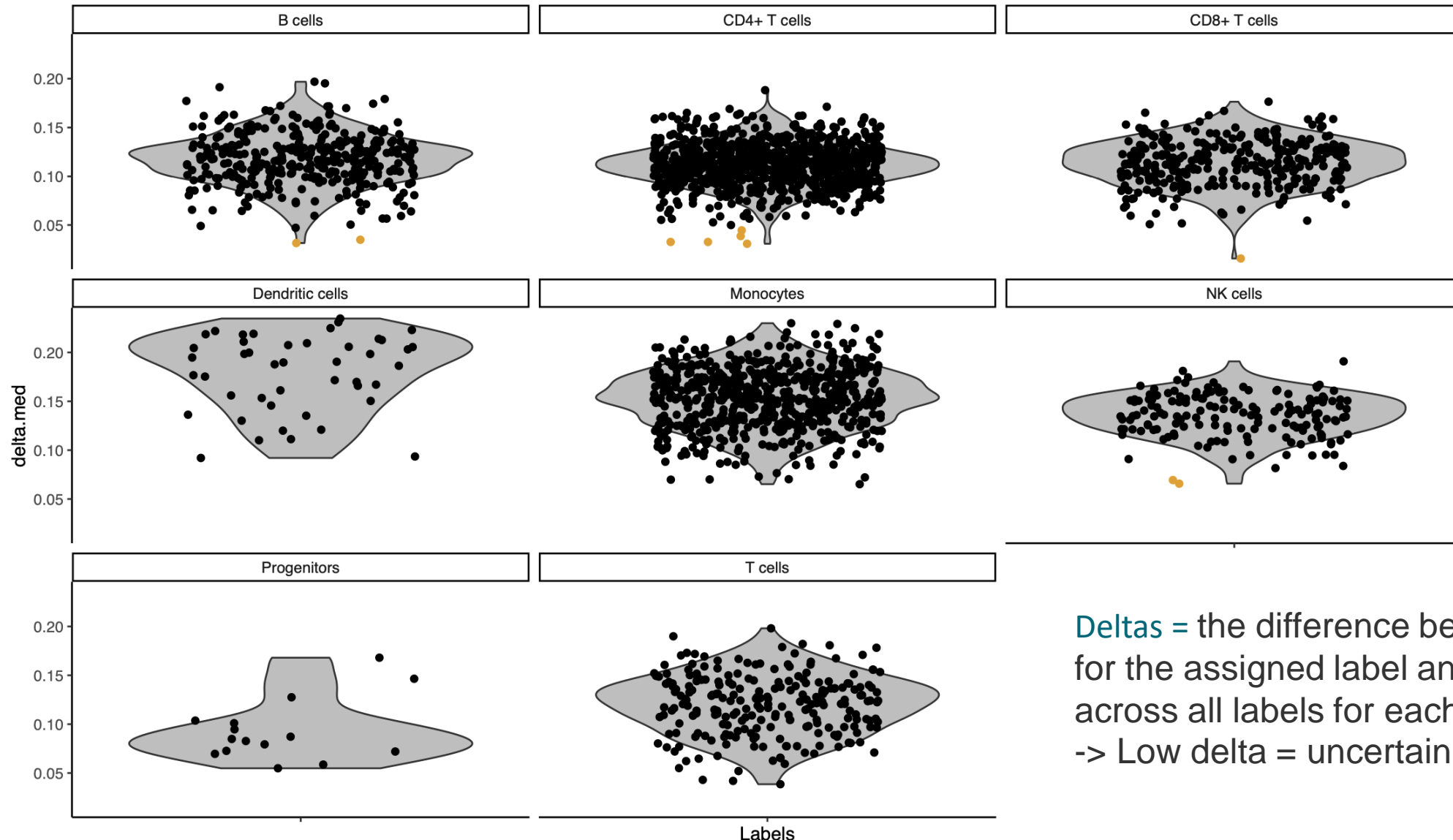
- **SingleR** is an automatic annotation method for scRNA-seq data
- Labels cells from the query dataset based on similarity to the reference dataset with known labels
- The **CellDex reference package** provides access to several reference datasets (mostly derived from bulk RNA-seq or microarray data) through dedicated retrieval functions -> sometimes, connection issues
- User can select the CellDex package to be used as reference
- Main level & fine level annotations



SingleR annotation: QC plots



SingleR annotation: QC plots



Pruned = potentially poor-quality or ambiguous assignments are removed based on the deltas

Pruned

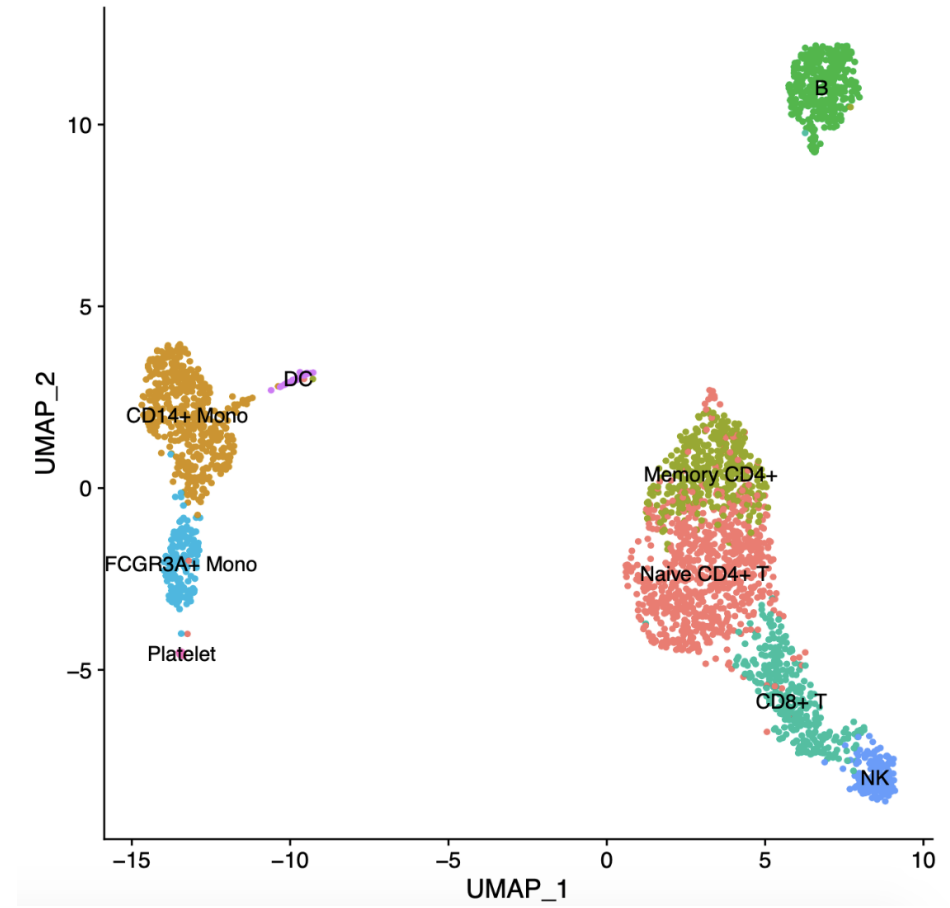
- FALSE
- TRUE

Deltas = the difference between the score for the assigned label and the median across all labels for each cell
-> Low delta = uncertain assignment

Rename clusters

- Based on previous knowledge and/or the SingleR results
- Import a table like this:

Cluster ID	Cluster name
0	Naive CD4+ T
1	CD14+ Mono
2	Memory CD4+
3	B
4	CD8+ T
5	FCGR3A+ Mono
6	NK
7	DC
8	Platelet



Analysis steps for clustering cells and finding marker genes



1. OPTIONAL: Remove background with CellBender
2. Create Seurat object, filter genes, check the quality of cells
3. Filter out low quality cells
4. Normalize expression values
5. Identify highly variable genes
6. Scale data, regress out unwanted variation
7. Reduce dimensions using principal component analysis (PCA) on the variable genes
8. Determine significant principal components (PCs)
9. Use the PCs to cluster cells with graph based clustering
10. Visualize clusters with non-linear dimensional reduction (tSNE or UMAP) using the PCs
11. Detect and visualize marker genes for the clusters

What you will learn

- What is background contamination
- How to remove it with CellBender in Chipster
 - You can include this optional step before the "Setup and QC" tool

Outline

- Problem of background contamination
- Feature-barcode matrices from Cell Ranger and CellBender
- Example: effects of CellBender background removal in downstream analyses
- Brief introduction into CellBender method
- How to use CellBender in Chipster

Problem of background contamination

- Good-quality cells may contain background contamination counts from **cell-free ambient RNA** or **barcode swapping** events
 - Observed UMI counts are a sum of **true biological counts** and **background contamination counts**
 - One study found that background contamination is **highly variable across replicates and cells** (3-35% of UMI counts per cell) [1]
- Potential source of **batch effects** and **spurious differential gene expression results** [2]
- Background contamination has a small effect on clustering [1]

[1] Janssen, P., Kliesmete, Z., Vieth, B. et al. The effect of background noise and its removal on the analysis of single-cell expression data. *Genome Biol* 24, 140 (2023).
<https://doi.org/10.1186/s13059-023-02978-x>

[2] Stephen J Fleming, Mark D Chaffin, Alessandro Arduini, Amer-Denis Akkad, Eric Banks, John C Marioni, Anthony A Phillipakis, Patrick T Ellinor, and Mehrtash Babadi. Unsupervised removal of systematic background noise from droplet-based single-cell experiments using CellBender. *Nature Methods*, 2023.
<https://doi.org/10.1038/s41592-023-01943-7>

CellBender uses raw (unfiltered) feature-barcode matrix as input

All barcodes →
 Estimated non-empty cells by Cell Ranger →

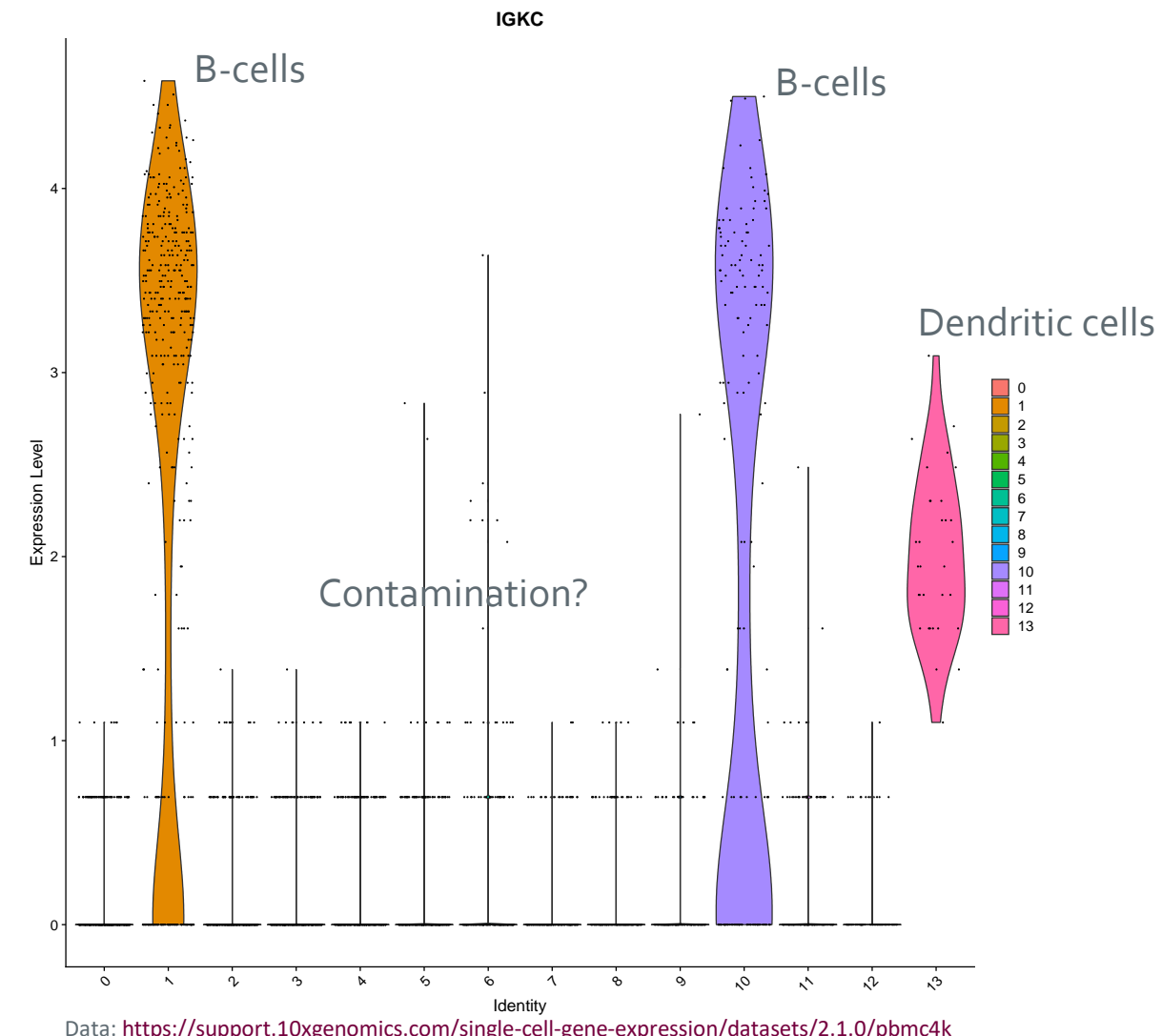
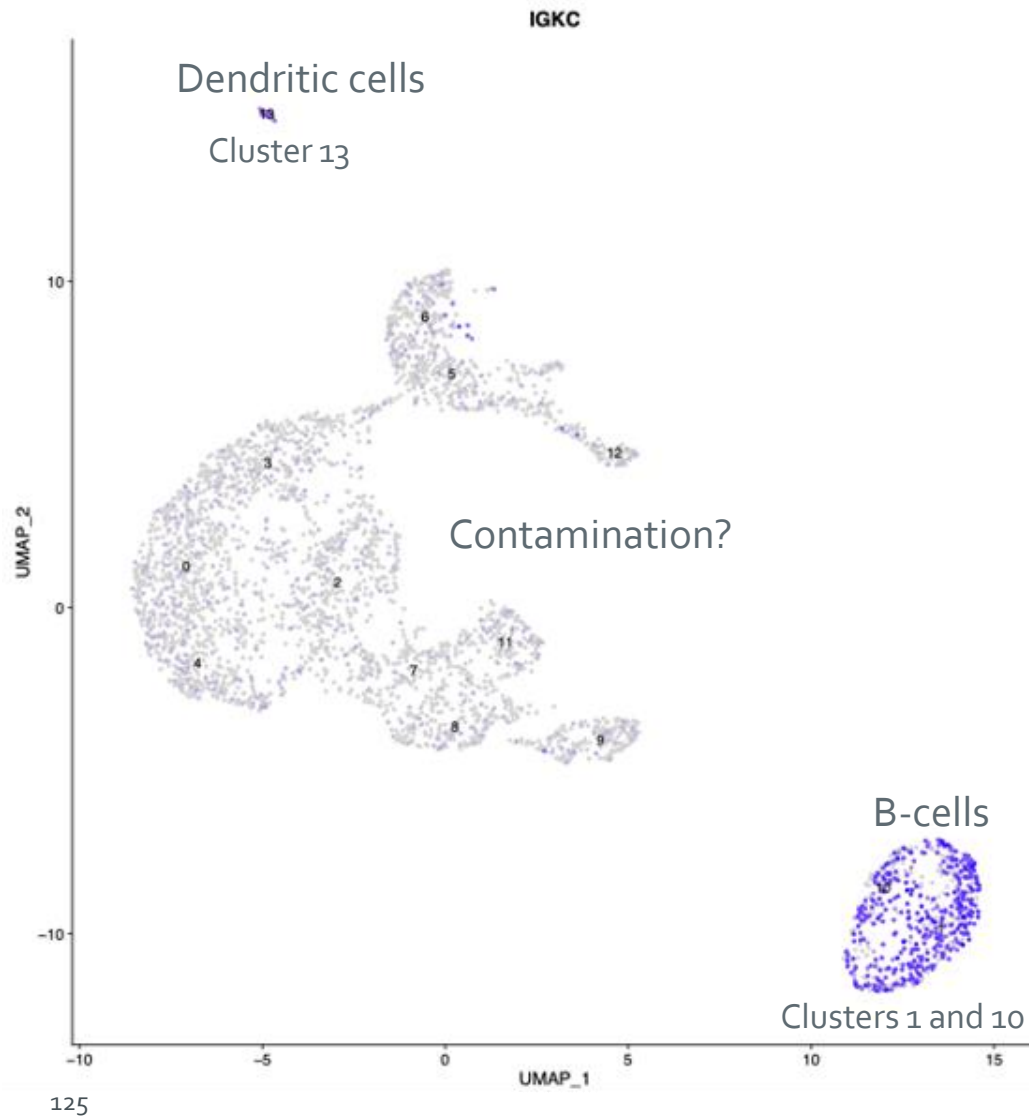
Type	Description
Unfiltered feature-barcode matrix	<p>Contains every barcode from the fixed list of known-good barcode sequences that has at least one read. This includes background and cell-associated barcodes.</p> <p><code>count: outs/raw_feature_bc_matrix/</code> <code>multi: outs/multi/count/raw_feature_bc_matrix/</code></p>
Filtered feature-barcode matrix	<p>Contains only detected cell-associated barcodes. For Targeted Gene Expression samples, non-targeted genes are removed from the filtered matrix.</p> <p><code>count: outs/filtered_feature_bc_matrix/</code> <code>multi: outs/per_sample_outs/count/sample_filtered_feature_bc_matrix/</code></p>

[3] <https://support.10xgenomics.com/single-cell-gene-expression/software/pipelines/latest/output/matrices>

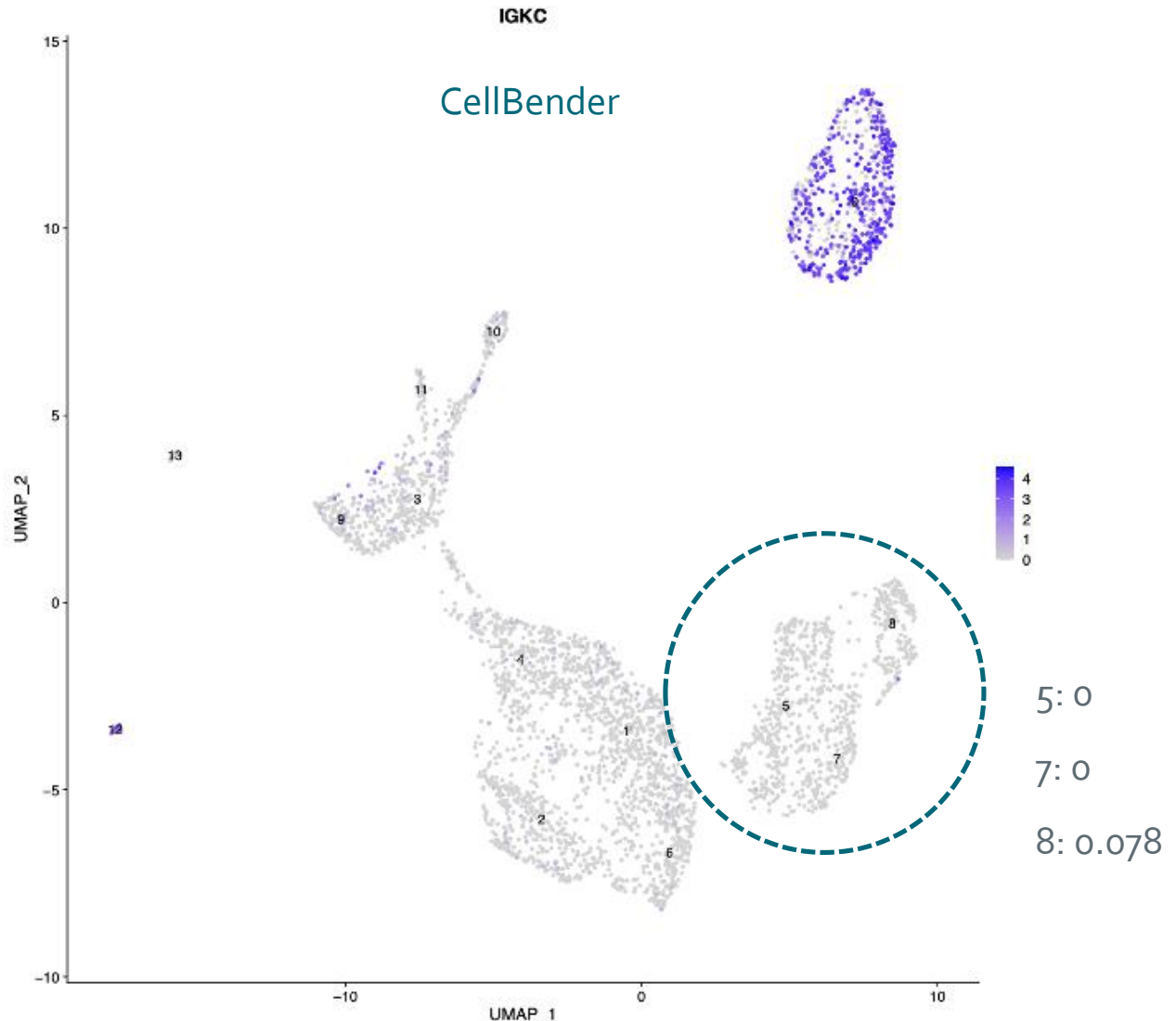
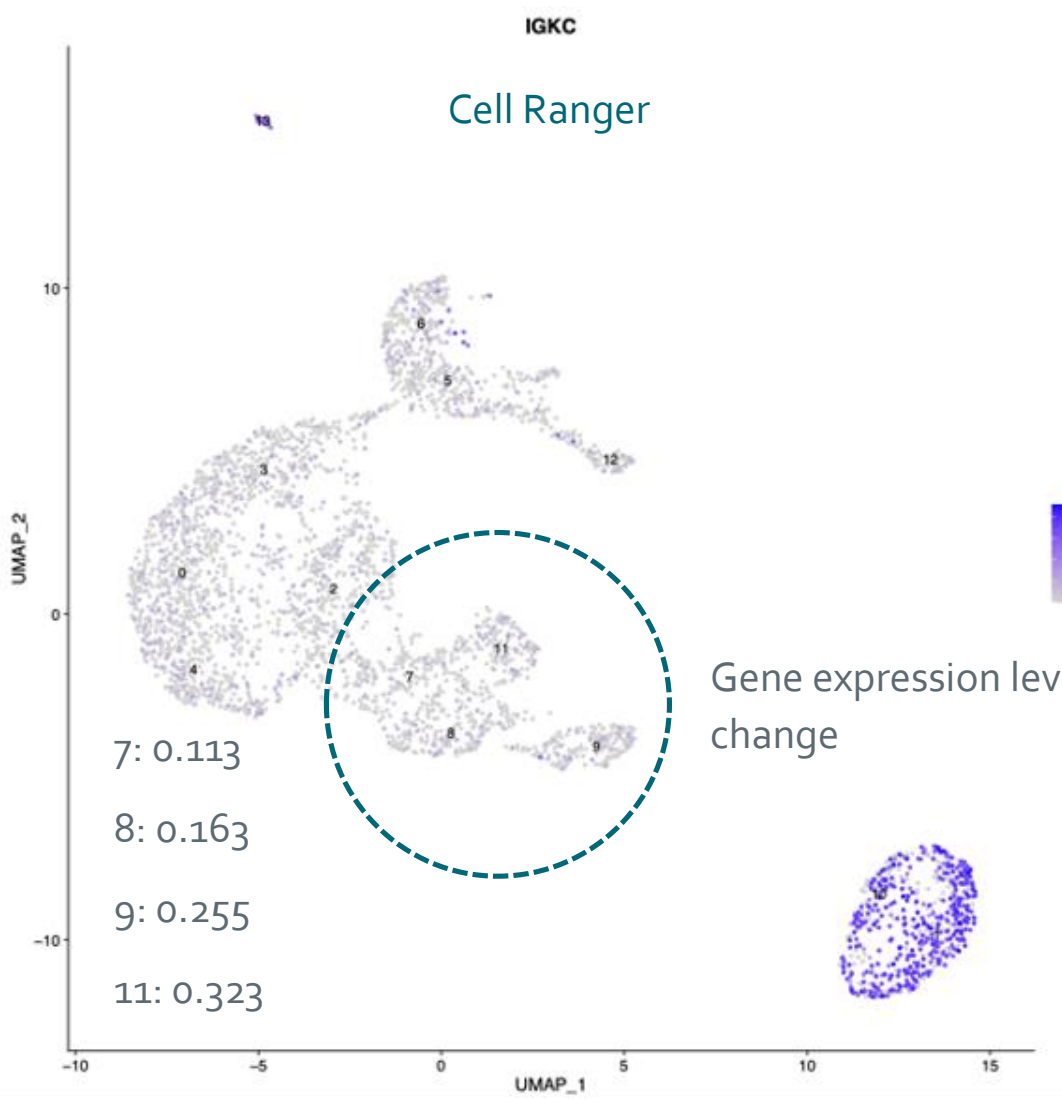
CellBender estimates new filtered feature-barcode matrix

- Background contamination counts have been removed
- Different approach to estimate cells compared to Cell Ranger
 - Contains slightly different set of cells

Example: immunoglobulin IGKC gene expression counts in all cell clusters?



Example: CellBender removes IGKC background



CellBender

- CellBender is a **probabilistic model** of noisy scRNA-seq data
- Takes into account key steps of data-generation process
- Observed UMI counts are a sum of **true (biological) cell UMI counts** and **background contamination counts**



[2] Stephen J Fleming, Mark D Chaffin, Alessandro Arduini, Amer-Denis Akkad, Eric Banks, John C Marioni, Anthony A Phillipakis, Patrick T Ellinor, and Mehrtash Babadi. Unsupervised removal of systematic background noise from droplet-based single-cell experiments using CellBender. *Nature Methods*, 2023.

<https://doi.org/10.1038/s41592-023-01943-7>

[4] <https://cellbender.readthedocs.io/en/latest/>

Modeling background contamination

- Background counts as Poisson distribution
- Two types of background contamination:
 - physically encapsulated exogenous molecules (cell-free ambient RNA)
 - barcode misassignment (barcode swapping)

$$c_{ng}^{\text{noise}} \sim \text{Poisson} \left[\underbrace{(1 - \rho_n) \epsilon_n d_n^{\text{drop}} \chi_g^a}_{\text{ambient noise rate}} + \underbrace{\rho_n \epsilon_n (y_n d_n^{\text{cell}} + d_n^{\text{drop}}) \bar{\chi}_g}_{\text{barcode swapping}} \right]$$

Modeling true cell UMI counts

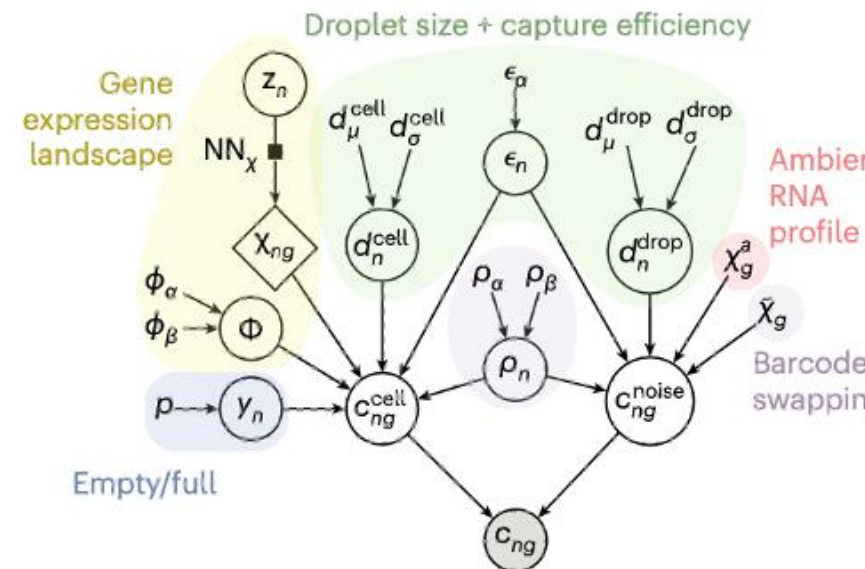
- True cell UMI counts as negative binomial distribution

Prior on true gene expression determined by neural network

$$c_{ng}^{\text{cell}} | \mathbf{z}_n \sim \text{NegBinom} \left[(1 - \rho_n) \epsilon_n y_n d_n^{\text{cell}} \overbrace{\chi_{ng}[\mathbf{z}_n]}^{\text{Prior on true gene expression}}[\mathbf{z}_n], \Phi \right]$$

Full model

- Variational autoencoder inside a structured probabilistic model of noisy scRNA-seq data
- New feature-barcode matrix obtained by subtracting the likely noise-counts from observed counts



$$\begin{aligned}
 z_n &\sim N(0, 1) \\
 X_{ng} &= \text{NN}_X(z_n) \\
 d_n^{\text{drop}} &\sim \text{lognormal}(d_\mu^{\text{drop}}, d_\sigma^{\text{drop}}) \\
 d_n^{\text{cell}} &\sim \text{lognormal}(d_\mu^{\text{cell}}, d_\sigma^{\text{cell}}) \\
 y_n &\sim \text{Bernoulli}(p) \\
 \rho_n &\sim \text{Beta}(\rho_\alpha, \rho_\beta) \\
 \epsilon_n &\sim \text{Gamma}(\epsilon_\alpha, \epsilon_\alpha) \\
 \Phi &\sim \text{Gamma}(\phi_\alpha, \phi_\beta) \\
 c_{ng}^{\text{cell}} &\sim \text{NegBinom} [\mu = (1 - \rho_n) \epsilon_n d_n^{\text{cell}} X_{ng} \Phi] \\
 c_{ng}^{\text{noise}} &\sim \text{Poisson} [(1 - \rho_n) \epsilon_n d_n^{\text{drop}} X_g^a \\
 &\quad + \rho_n \epsilon_n (y_n d_n^{\text{cell}} + d_n^{\text{drop}}) \bar{X}_g] \\
 c_{ng} &= c_{ng}^{\text{cell}} + c_{ng}^{\text{noise}}
 \end{aligned}$$

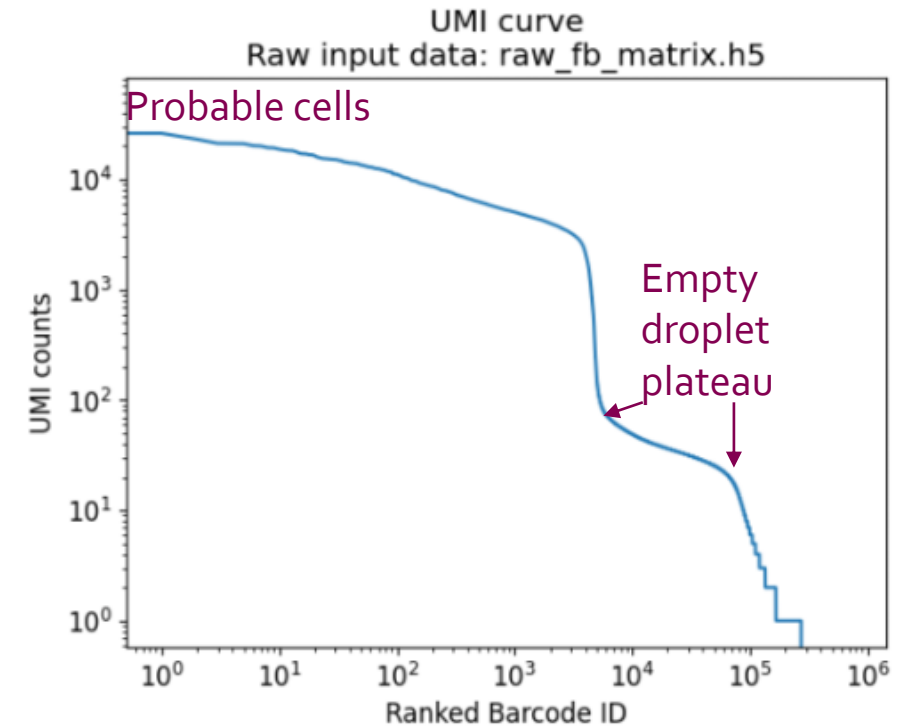
[2] Stephen J Fleming, Mark D Chaffin, Alessandro Arduini, Amer-Denis Akkad, Eric Banks, John C Marioni, Anthony A Phillipakis, Patrick T Ellinor, and Mehrtash Babadi. Unsupervised removal of systematic background noise from droplet-based single-cell experiments using CellBender. *Nature Methods*, 2023. <https://doi.org/10.1038/s41592-023-01943-7>

How to use CellBender in Chipster

- Input: Run tool as first step before Setup and QC!
 - Raw 10x feature-barcode matrix in **hdf5** (.h5) file format
- Parameters:
 - Expected number of non-empty cells [auto]
 - Total number of included droplets [auto]
 - Number of epochs [150] Run CellBender using defaults first!
 - Learning rate [0.0001]
 - Nominal false positive rate [0.01]
- Output:
 - Feature-barcode matrix filtered with CellBender
 - Report (.html) Check report and log files for possible warnings!
 - Log (.log)

UMI curve plot

- Barcodes ranked by their UMI counts
- **Expected number of cells (exp_num):** all droplets that reasonably surely contain cells
- **Total number of included droplets (tot_num):** a number that goes a few thousand barcodes into the empty droplet plateau

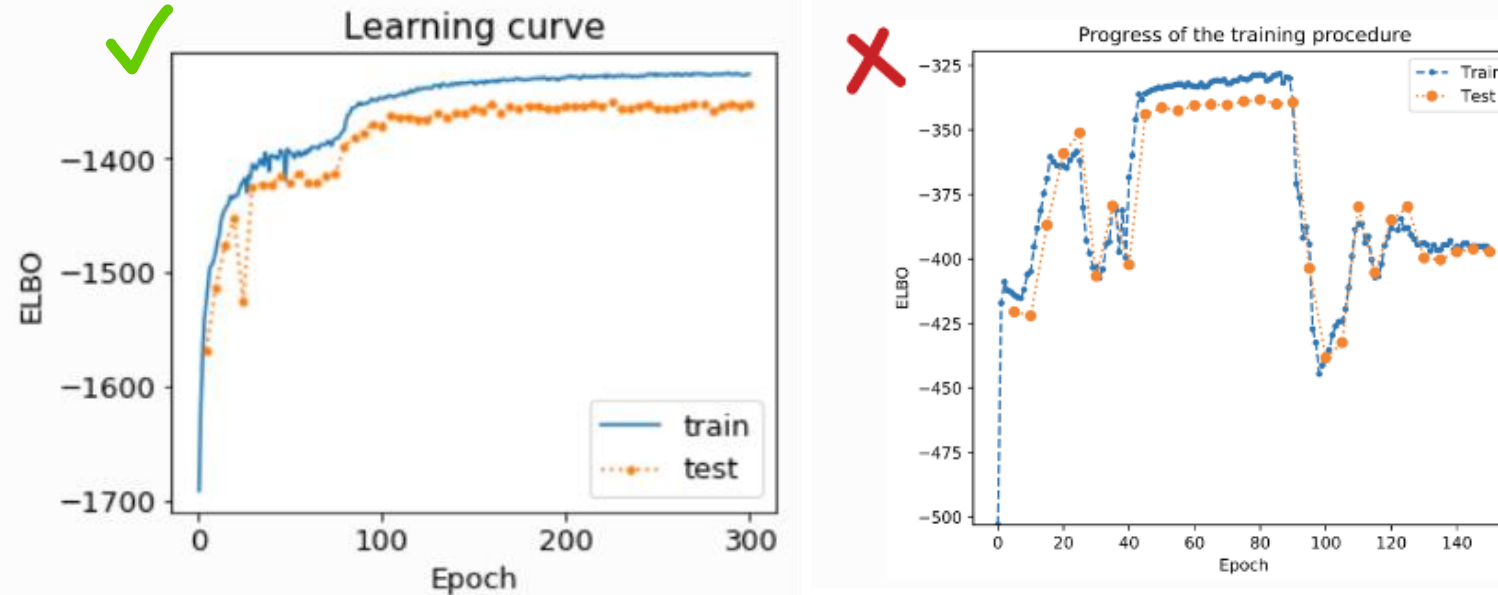


Log file: exp_num = 2383

tot_num = 2383 + 9531 = 11914

```
ound: Excluding barcodes with counts below 18
ound: Using 2383 probable cell barcodes, plus an additional 9531 barcodes, and
ound: Largest surely-empty droplet has 44 UMI counts.
ound: Attempting to unpack tarball "ckpt.tar.gz" to /tmp/tmpv_gf0ff6
```

Learning curve



- ELBO should increase as training epochs increase and converge at some high plateau
- Look for large downward dips in ELBO
- **Learning rate:** if large dips, try to reduce the learning rate by a factor of two
- **Number of epochs:** increase number if learning curve has not converged but generally do not go beyond 300!

[5] <https://cellbender.readthedocs.io/en/latest/troubleshooting/index.html>

Nominal false positive rate

- Involves trade-off between removing noise and retaining signal
- Imposes an upper bound on the amount of falsely removed signal counts
- Default (0.01) fairly conservative and appropriate for most analyses
- Value 1 represents removal of nearly every UMI count (signal and noise)

Check the report and log files for any warnings!

WARNING: The expression of the highly-expressed gene [REDACTED] decreases quite markedly after CellBender. Check to ensure this makes sense!

WARNING: Only 1500 barcodes in the input file. Ensure this is a raw (unfiltered) file with all barcodes, including the empty droplets.

Summary

- Background contamination can be a potential source of batch effects and spurious differential gene expression
- Consider removing background contamination, **when high level of systematic background contamination** is present in your data
- Use CellBender as a first step in your analysis
- Use **raw 10x feature-barcode matrix in hdf5** format as input
- Run CellBender tool using **default parameter values** and look at the report and log files to see if changes are needed
- Use the new CellBender filtered feature-barcode matrix as as input to the tool "Setup and QC" and continue with downstream analyses

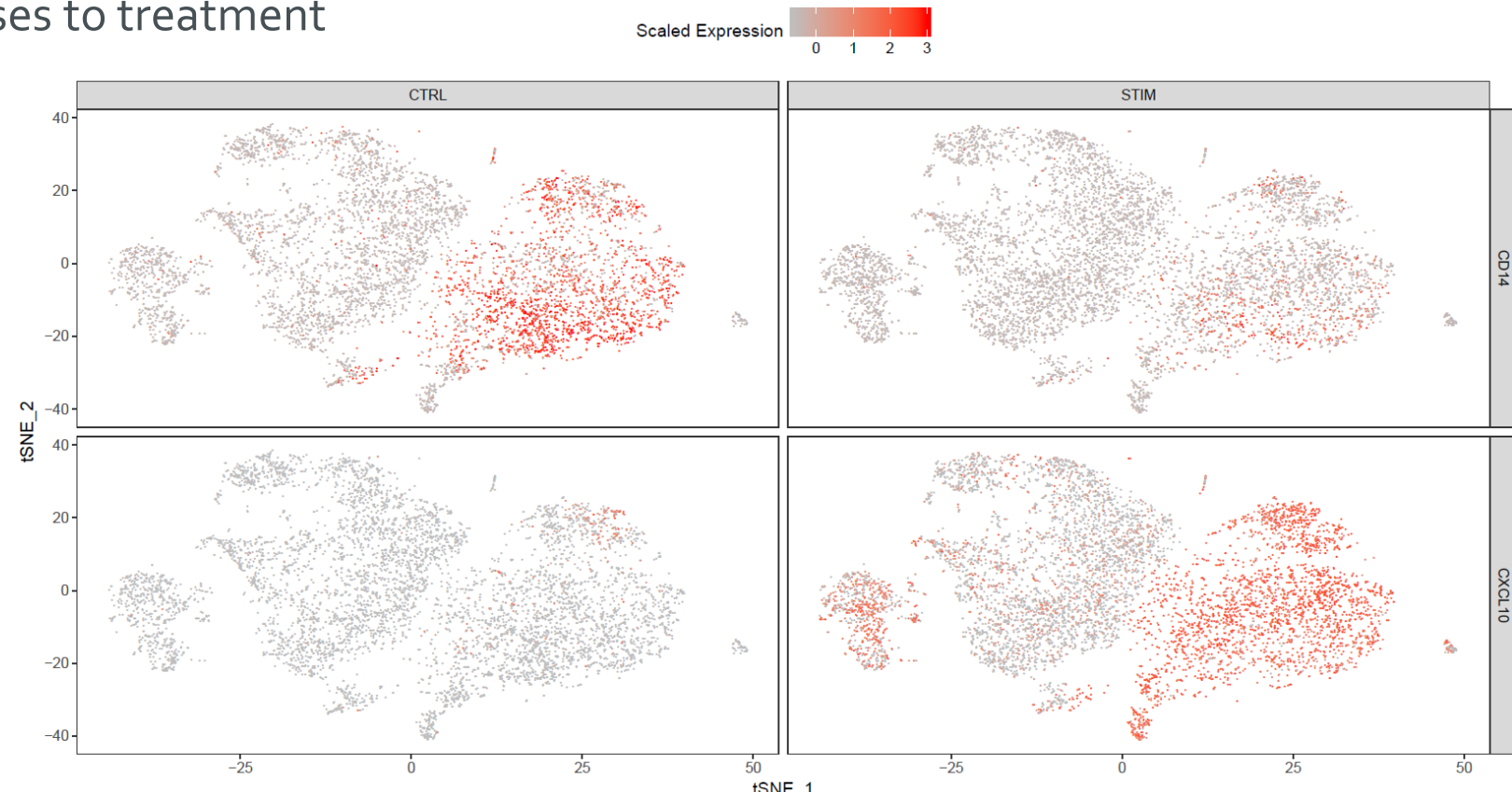
Integrated analysis of multiple samples

What will you learn

1. What we need to consider when comparing samples
2. How to integrate samples
3. How to find conserved cluster marker genes
4. How to find differentially expressed genes between samples, within clusters
5. How to visualize interesting genes

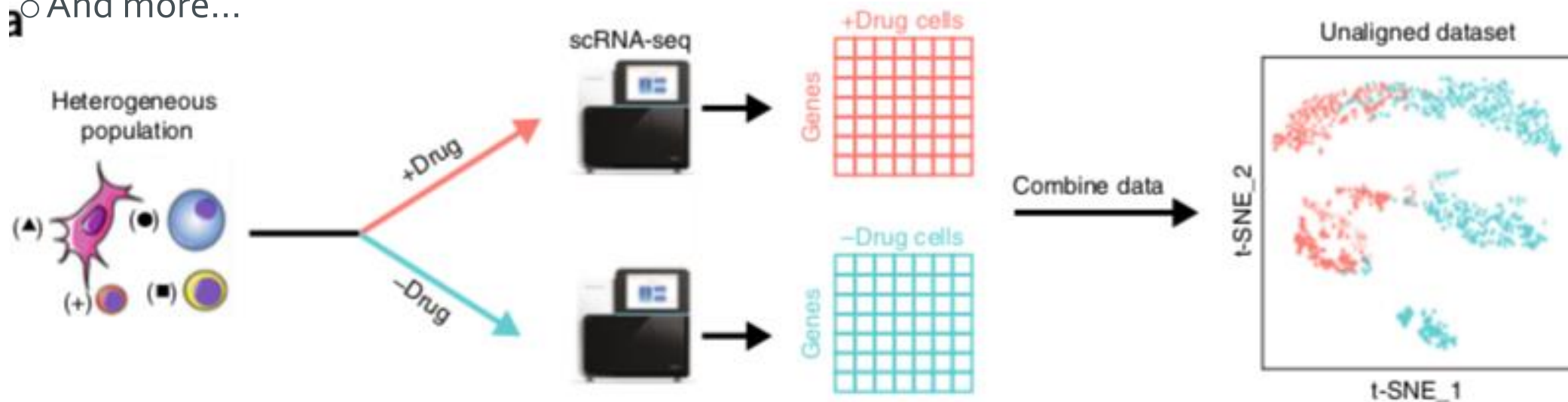
Goals of integrated analysis

- When comparing two samples, e.g. control and treatment, we want to
 - Identify cell types that are present in both samples
 - Obtain cell type markers that are conserved in both control and treated cells
 - Find cell-type specific responses to treatment



When comparing samples we need to correct for batch effects

- We need to find corresponding cells in the samples
 - Technical and biological variability can cause batch effects which make this difficult
- Several batch effect correction methods for single cell RNA-seq data available, e.g.
 - Seurat v2: Canonical correlation analysis (CCA) + dynamic time warping
 - Seurat v3-v4: CCA + anchors
 - Mutual nearest neighbors (MNN)
 - And more...

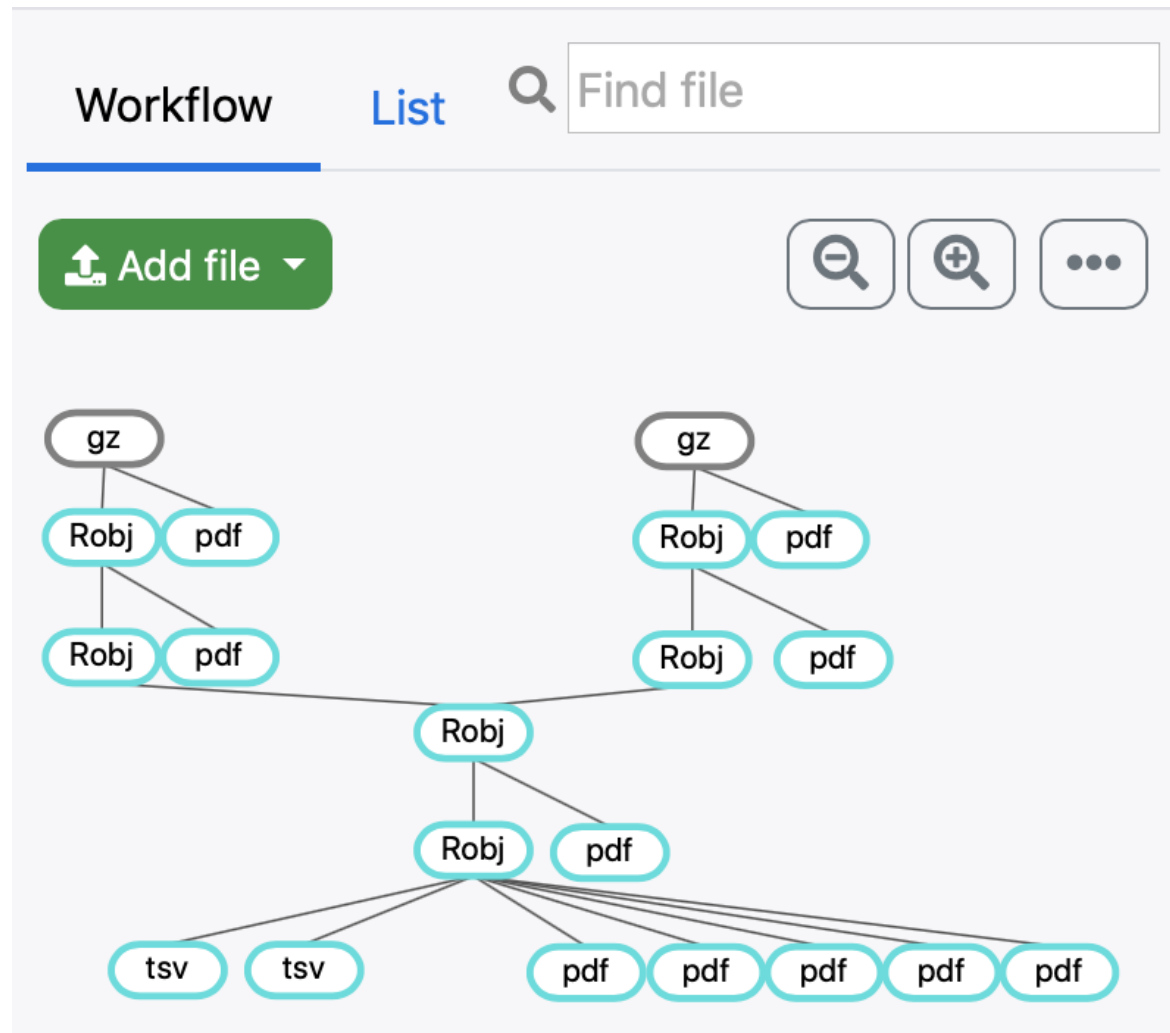


Analysis steps for integrated analysis

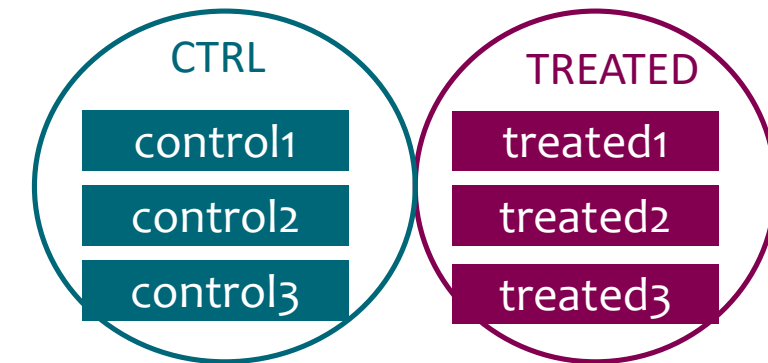


1. Create Seurat objects, filter genes, check the quality of cells
2. Normalize expression values
3. Identify highly variable genes
4. Integrate samples and perform CCA, align samples
5. Scale data, perform PCA
6. Cluster cells, visualize clusters with tSNE or UMAP
7. Find conserved biomarkers for clusters
8. Find differentially expressed genes between samples, within clusters
9. Visualize interesting genes

Integrated analysis: Setup, QC, filtering



- Perform the Seurat object setup, QC and filtering steps separately for the samples
 - Same as before, just remember to name:
 - the samples: e.g. control_1, control_2...
 - and sample groups: e.g. CTRL and TREATED



Parameters

Project name for plotting

You can give your project a name. The name will appear on the plots.

Do not use underscore _ in the names!

My_project_name

control1

CTRL

Sample name

Type the sample name or identifier here. For example control1, cancer3a. Do not use underscore _ in the names! Fill this field if you are combining samples later.

Sample group

Type the sample name or identifier here. For example CTRL, STIM, TREAT. Do not use underscore _ in the names! Fill this field if you are combining samples later.

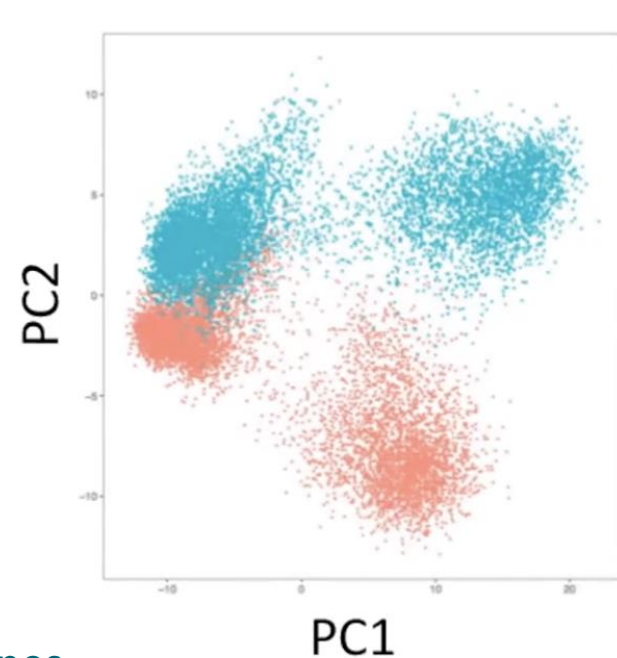
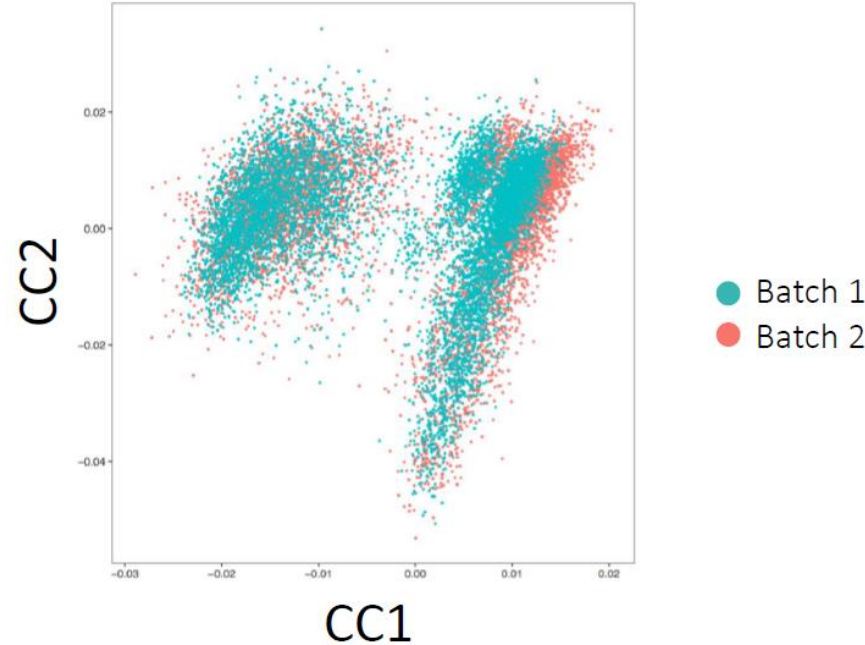
Analysis steps for integrated analysis



1. Create Seurat objects, filter genes, check the quality of cells
2. Normalize expression values
3. Identify highly variable genes
4. Integrate samples and perform CCA, align samples
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Canonical correlation analysis (CCA)

- Dimension reduction, like PCA
- Captures common sources of variation between two datasets
 - Aim: place datasets in a shared, low-dimensional space
- Produces canonical correlation vectors, CCs
 - Effectively capture correlated gene modules that are present in both datasets
 - Represent genes that define a shared biological space
- Why not PCA?
 - It identifies the sources of variation, even if present only in 1 sample (e.g. technical variation)
 - We want to integrate, so we want to find the *similarities*

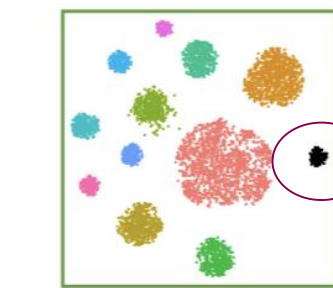
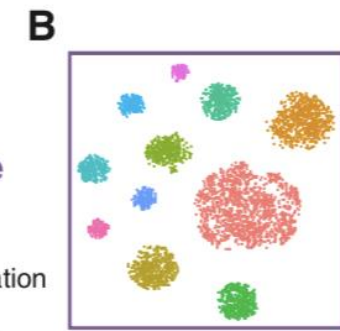
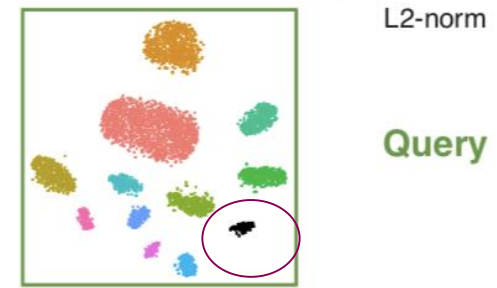
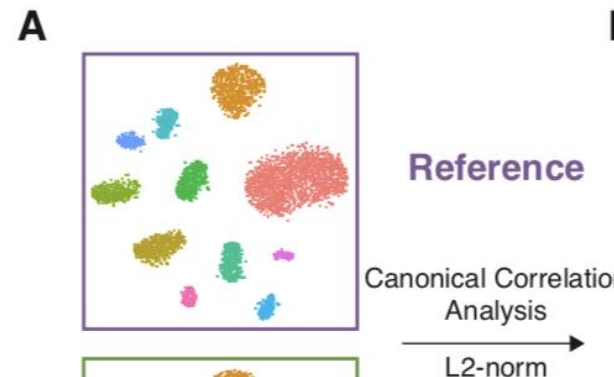


Input:
Highly variable genes

Aligning two samples (Seurat v3/v4)

See the Seurat paper:
[https://www.cell.com/cell/fulltext/S0092-8674\(19\)30559-8](https://www.cell.com/cell/fulltext/S0092-8674(19)30559-8) 

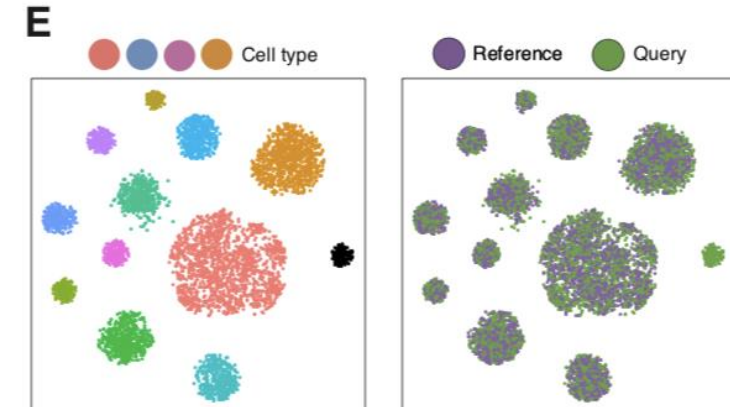
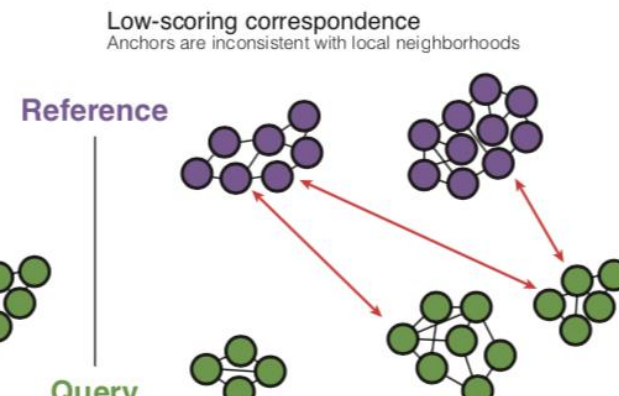
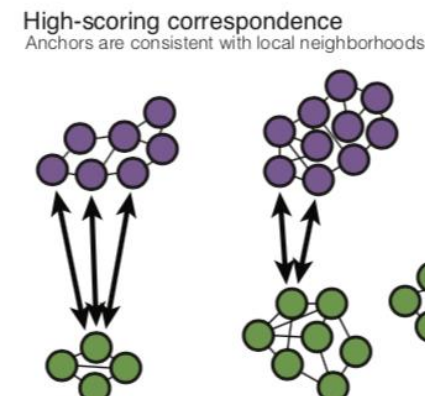
1. Canonical correlation analysis
+ L2-normalisation of CCVs for scaling
→ shared space



2. Identify pairs of mutual nearest neighbors (MNN) → "anchors"

3. Filter & score anchors **D**
(based on neighborhood, in PC space)

4. Anchors + scores → correction vectors



Combine multiple samples tool

1. Identify “anchors” for data integration
 - o Parameter: how many CCs to use in the neighbor search [20]
2. Integrate datasets together
 - o Parameter: how many PCs to use in the anchor weighting procedure [20]

Parameters

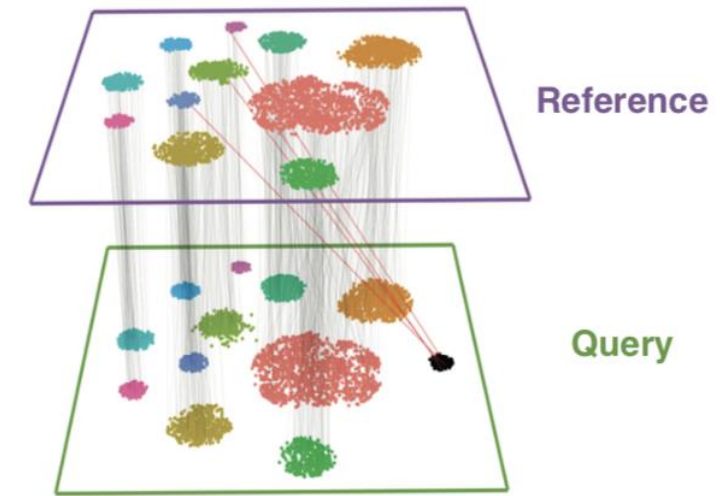
Number of CCs to use in the neighbor search

Which dimensions to use from the CCA to specify the neighbor search space. The neighbors are used to determine the anchors for the alignment.

▼

Number of PCs to use in the anchor weighting

Number of PCs to use in the anchor weighting procedure. The anchors and their weights are used to compute the correction vectors, which allow the datasets to be integrated.

▼


Same question as before:
What is the dimensionality
of the data?

Dimensionality –how many CCs / PCs to choose for downstream analysis?

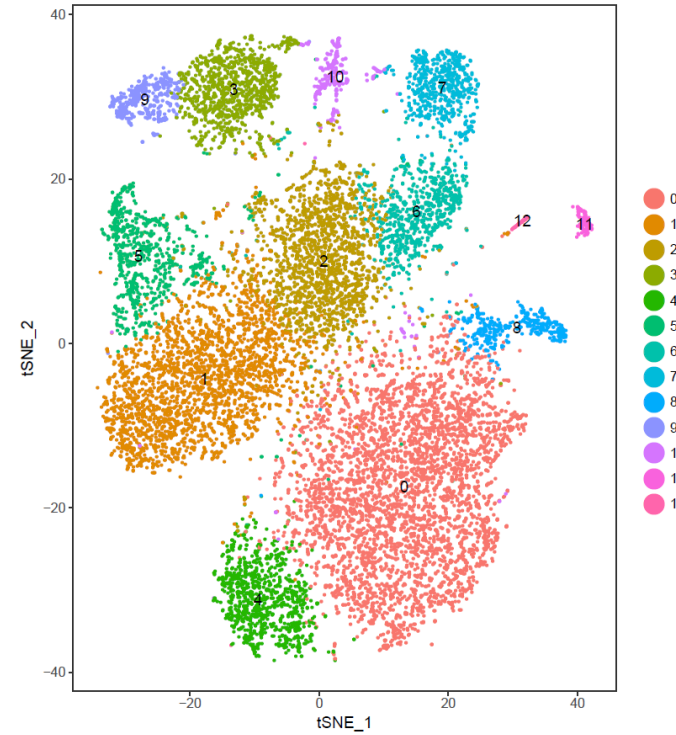
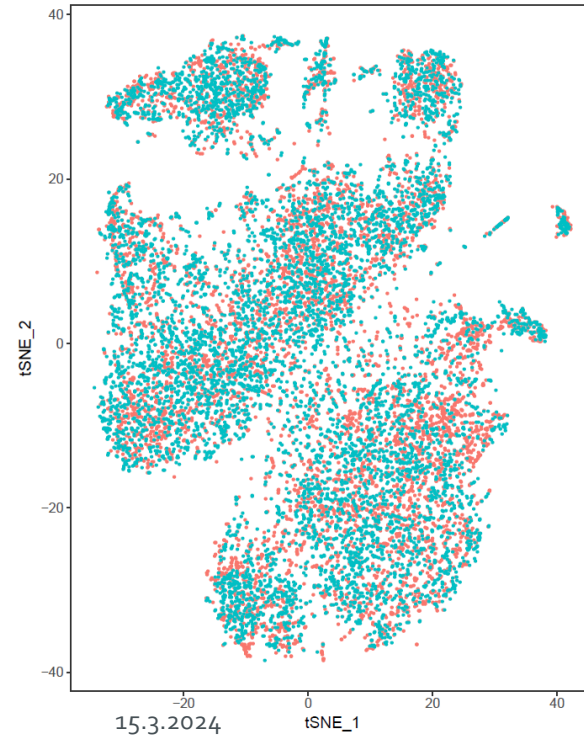


- In the article* by Seurat developers, they “neglect to finely tune this parameter for each dataset, but still observe robust performance over diverse use cases”.
 - For all neuronal, bipolar, and pancreatic analyses: dimensionality of 30.
 - For scATAC-seq analyses: 20.
 - For analyses of human bone marrow: 50
 - The integration of mouse cell atlases: 100
- Higher numbers: for significantly larger dataset and increased heterogeneity

* <https://www.biorxiv.org/content/biorxiv/early/2018/11/02/460147.full.pdf>

Integrated analysis of two samples –tools (v4)

1. Cluster cells
 - As before
2. Visualize clustering
 - tSNE or UMAP, as a parameter



Total number of cells: 13997
Number of cells in each cluster:

	CTRL	STIM
0	2172	2126
1	973	1579
2	870	848
3	512	547
4	400	553
5	351	478
6	295	328
7	297	318
8	296	222
9	185	200
10	86	121
11	51	80
12	37	20
13	23	29

Larger datasets 1: Using reference samples in integration

- Why?
 - Memory and time savings (too long jobs are killed, and memory can run out)
 - By default, anchors are identified between all pairs of samples (i.e. for 10 samples, there are 45 comparisons).
- The "**Samples to use as references**" parameter allows users to list sample names to be used as integration references.
 - Users can type the reference sample names (separated with comma)
 - Make sure you type the sample name correctly, exactly like you typed it in the Setup tool!
 - For example, 10 samples, 1 reference → 9 comparisons
- Select representative samples as references!
 - For example, if you have samples from male and female patients, pick one reference from both

Large datasets 2: Anchor identification method (CCA → rPCA)



- CCA = default
 - Might lead to overcorrection, especially when large proportion of cells are non-overlapping
 - Recommended when:
 - When cell types are conserved, but there's still big difference between the samples/experiments → experimental condition/disease causes very strong expression shift
 - Cross-modality mapping
 - Cross-species mapping
- rPCA = reciprocal PCA
 - **Faster**, more conservative: cells in different biological states are less likely to “align”
 - Each dataset is projected into the others PCA space and the anchors are constrained by the same mutual neighbourhood requirement
 - Recommended when:
 - A substantial fraction of cells in one dataset have no matching type in the other
 - Datasets originate from the same platform (i.e. multiple lanes of 10x Genomics)
 - There are a large number of datasets or cells to integrate

https://satijalab.org/seurat/articles/integration_rpc.html

https://satijalab.org/seurat/articles/integration_large_datasets.html

Analysis steps for integrated analysis



1. Create Seurat objects, filter genes, check the quality of cells
2. Normalize expression values
3. Identify highly variable genes
4. Integrate samples and perform CCA, align samples
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Find conserved cluster marker genes in multiple samples



- Conserved marker gene = marker for a given cluster in *all* samples
 - Give a cluster as a parameter
 - Compares gene expression in cluster X vs all other cells
 - This is done in each sample, and then the p-values are combined using Wilkinson's method
- Uses Wilcoxon rank sum test

Seurat v4 -Find conserved cluster markers and DE genes in multiple samples



Parameters



Normalisation method used previously

Which normalisation method was used in preprocessing, Global scaling normalization (default, NormalizeData function used) or SCTransform.

Global scaling normalization

Name of the cluster

Name of the cluster of which you want to identify the differentially expressed. By default, the clusters are named with numbers starting from 0.

3

Return only positive marker genes

Tool only returns positive markers as default. Change the parameter here if you want to also include the negative markers.

TRUE

Conserved markers: Fold change in log2 scale

Genes with an average fold change smaller than this are not included in the analysis.

0.25

Conserved markers: Adjusted p-value cutoff

Cutoff for the adjusted p-value of the conserved cluster marker genes: by default, adjusted p-values bigger than 0.05 in any sample are filtered out.

0.05

Conserved markers: Limit testing to genes which are expressed in at least this fraction of cells

Test only genes which are detected in at least this fraction of cells in the cluster in question or in all the other cells. Meant to speed up testing by leaving out genes that are very infrequently expressed.

0.1

Conserved markers: Minimum number of cells in one of the groups

How many cells at least there needs to be in each sample in the cluster in question.

3

Differentially expressed genes: Fold change in log2 scale

Genes with an average fold change smaller than this are not included in the analysis.

0.25

Differentially expressed genes: Adjusted p-value cutoff

Cutoff for the adjusted p-value of the DE genes: by default, adjusted p-values bigger than 0.05 are filtered out.

0.05

Differentially expressed genes: Limit testing to genes which are expressed in at least this fraction of cells

Test only genes which are detected in at least this fraction of cells in either of two samples being compared in the cluster of question. Meant to speed up testing by leaving out genes that are very infrequently expressed.

0.1

Showing all 399 rows.

	PBMC_control_p_val	PBMC_control_avg_log2FC	PBMC_control_pct.1	PBMC_control_pct.2	PBMC_control_p_val_adj	PBMC_stim_p_val	PBMC_stim_avg_log2FC	PBMC_stim_pct.1	PBMC_stim_pct.2
MS4A1	0	3.00964815667874	0.583	0.015	0	0	2.62960740854217	0.466	0
CD79B	0	2.21759625961027	0.384	0.017	0	4.64055698459073e-127	1.09663557662942	0.139	0
CD79A	0	3.82731299763922	0.806	0.029	0	0	3.33403518707591	0.673	0
CD74	6.8422885025142e-223	2.27808118754189	0.998	0.661	9.6154680325832e-219	1.72427249916692e-275	2.06125382725302	0.994	0
BANK1	1.66963534002066e-196	1.31368036905531	0.194	0.004	2.34633854333103e-192	1.23235576431409e-265	1.58515860222653	0.239	0
TNFRSF13B	1.54363717399291e-212	1.57262716485063	0.19	0.003	2.16927332061224e-208	1.24170154864956e-207	1.35511769802397	0.174	0
BLNK	3.6266469581841e-168	1.40999622559305	0.207	0.009	5.09652697033611e-164	6.72781546727165e-208	1.64446692862033	0.28	0
IRF8	8.25570610845141e-66	1.29665063307448	0.24	0.047	1.16017437942068e-61	1.67971662266102e-199	2.08503639380049	0.545	0
KIAA0226L	2.08720921799168e-193	1.63628212605763	0.242	0.011	2.93315511404371e-189	1.70445839800893e-184	1.2049627683166	0.174	0
TCL1A	3.36767246176029e-174	1.38193266241839	0.181	0.005	4.73259011051174e-170	1.53474080152144e-112	1.10856360324414	0.121	0
FCRLA	1.79270329956639e-162	1.13552720513697	0.148	0.002	2.51928594688064e-158	8.30439587710791e-130	0.836853125240639	0.108	0
C7orf50	1.14014217765531e-70	1.35985597431791	0.282	0.061	1.602241802259e-66	1.28353768100081e-158	1.66938921961221	0.319	0
CD83	1.63516776125378e-157	2.25944038088898	0.581	0.133	2.29790125488993e-153	1.46029122999176e-82	1.58463347318392	0.463	0

PBMC_stim_pct.2	PBMC_stim_p_val_adj	max_pval	minimump_p_val
0.012	0	0	0
0.007	6.52137473044535e-123	4.64055698459073e-127	0
0.02	0	0	0
0.662	2.42312014307927e-271	6.8422885025142e-223	3.4485449983335e-275
0.007	1.7318295555906e-261	1.66963534002066e-196	2.46471152862819e-265
0.004	1.74496318631723e-203	1.24170154864956e-207	3.08727434798585e-212
0.022	9.45459907615685e-204	3.6266469581841e-168	1.34556309345435e-207
0.124	2.36050576982553e-195	8.25570610845141e-66	3.359433245322e-199

Find cell-type specific differentially expressed genes between samples

- We are now looking for differential expression between *samples in one cluster*
- Uses Wilcoxon rank sum test
- Parameters for filtering the table:
 - Adjusted p-value cutoff for differentially expressed genes (default = 0.05)
 - Fold change threshold for differentially expressed genes in log2 scale (default = 0.25)
- If there are >2 samples, a table for each sample is given as output
 - named: de-list_sample1VsAllOthers.tsv, de-list_sample2VsAllOthers.tsv...

de-list_PBMC_control_vs_PBMC_stim.tsv ...

Spreadsheet Text Open in New Tab Details

Showing the first 100 of 778 rows. View in full screen to see all rows.

	p_val	avg_log2FC	pct.1	pct.2	p_val_adj	aver_expr_ident1	aver_expr_ident2
ISG15	1.7296497545789e-178	-4.653740532325	0.231	0.998	2.43067680010973e-174	2.5918	89.4134
IFIT3	7.75442106128566e-172	-4.51511176245437	0.046	0.959	1.08972879174247e-167	0.4133	31.3164
ISG20	8.79249690050336e-169	-2.99108560051549	0.651	1	1.23560958942774e-164	9.8323	85.1244
IFI6	6.95069852044584e-167	-4.19106395111127	0.076	0.953	9.76781663078254e-163	0.6552	29.2335
IFIT1	1.0058042691094e-150	-4.11242648599536	0.028	0.886	1.41345673937944e-146	0.2613	20.8171

Find cell-type specific differentially expressed genes between samples



- We are now looking for differential expression between *samples* in one cluster
 - Actually, these tests are looking at the cells in that sample & in that cluster
 - Each cell is thus treated as an independent replicate, and the inherent correlations between cells originating from the same sample are ignored → Large number of false positives*
 - Use caution when interpreting!
- "Solution": **pseudobulk** analysis
 - Sum together gene counts of all the cells from the sample & cluster → one expression profile, samples treated as individual observations. Then compare to the cell-wise results. [In Seurat v5](#)
- 3 tools:
 - Seurat v4 -Find conserved cluster markers and DE genes in multiple samples
 - Seurat v4 -Find DE genes between chosen samples
 - Seurat v4 -Find DE genes between sample groups

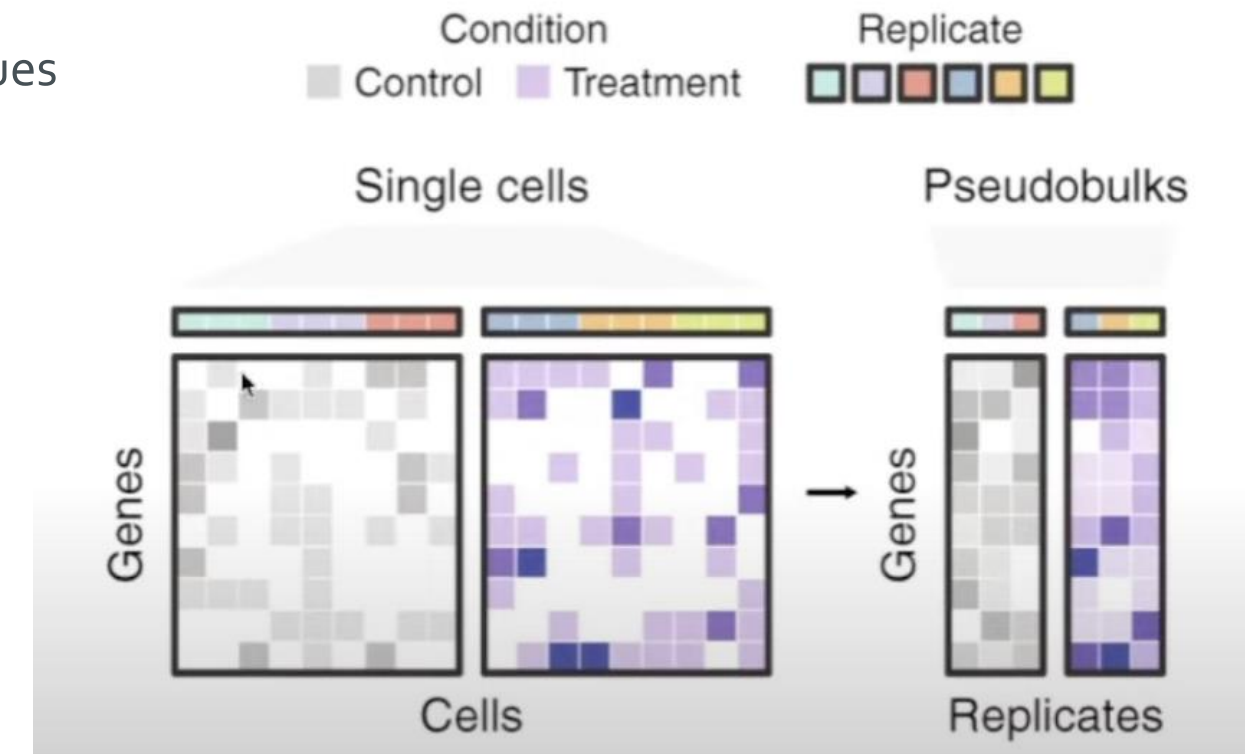
All sample-wise comparisons at once

Choose the samples you want to compare

Choose to compare sample groups (from the setup tool)

What pseudo bulk analysis?

- Why is it better than scRNASeq or bulk RNASeq?
 - Each cell treated as sample → inflated p-values → false positives
 - Tendency to identify high expressors as DE genes
 - Ignoring the variation across population / unmodelled correlation between samples
 - Get the benefit of single-cell resolution (= recognition of cell types) AND the statistical rigor of the existing bulk-RNASeq methods



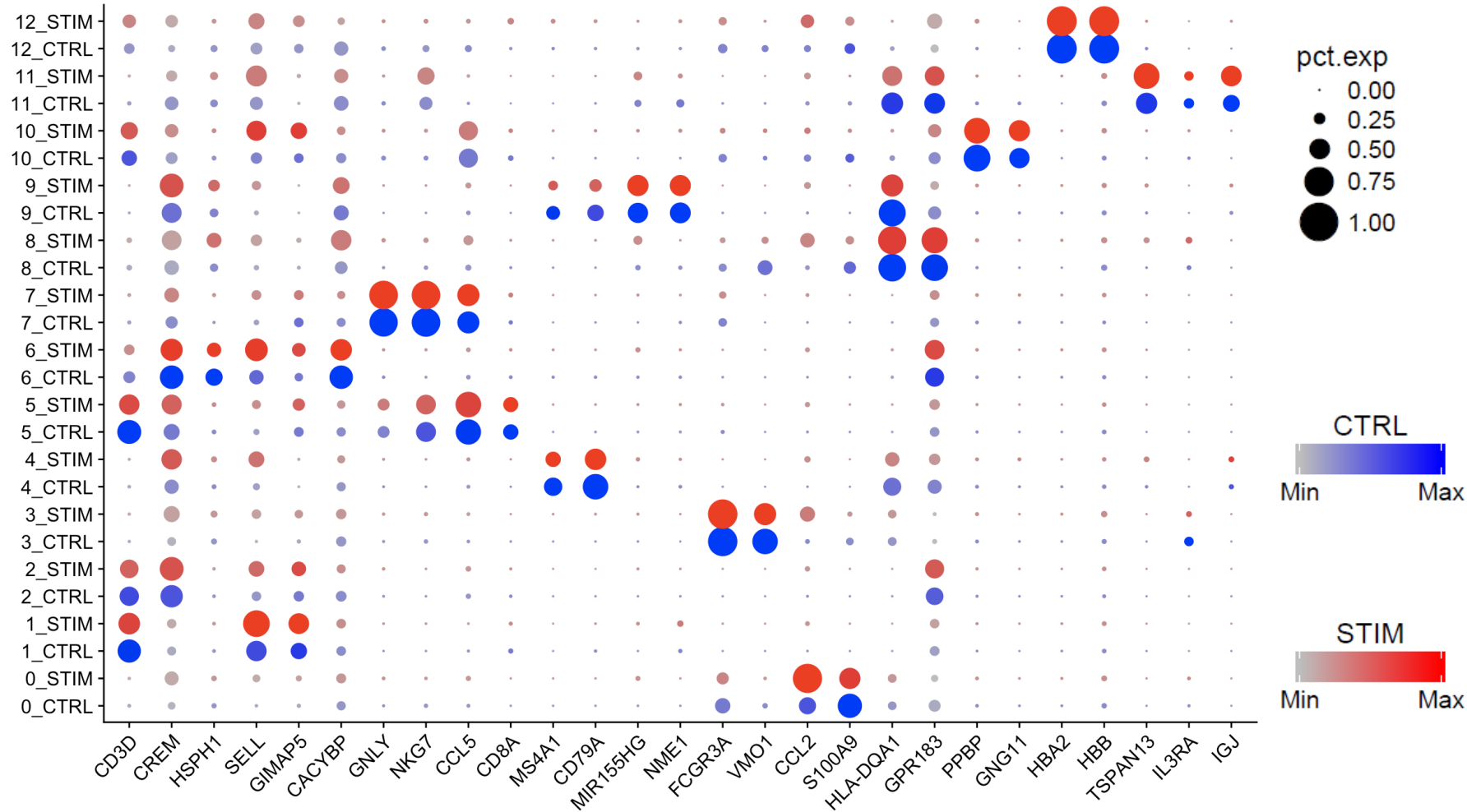
Analysis steps for integrated analysis



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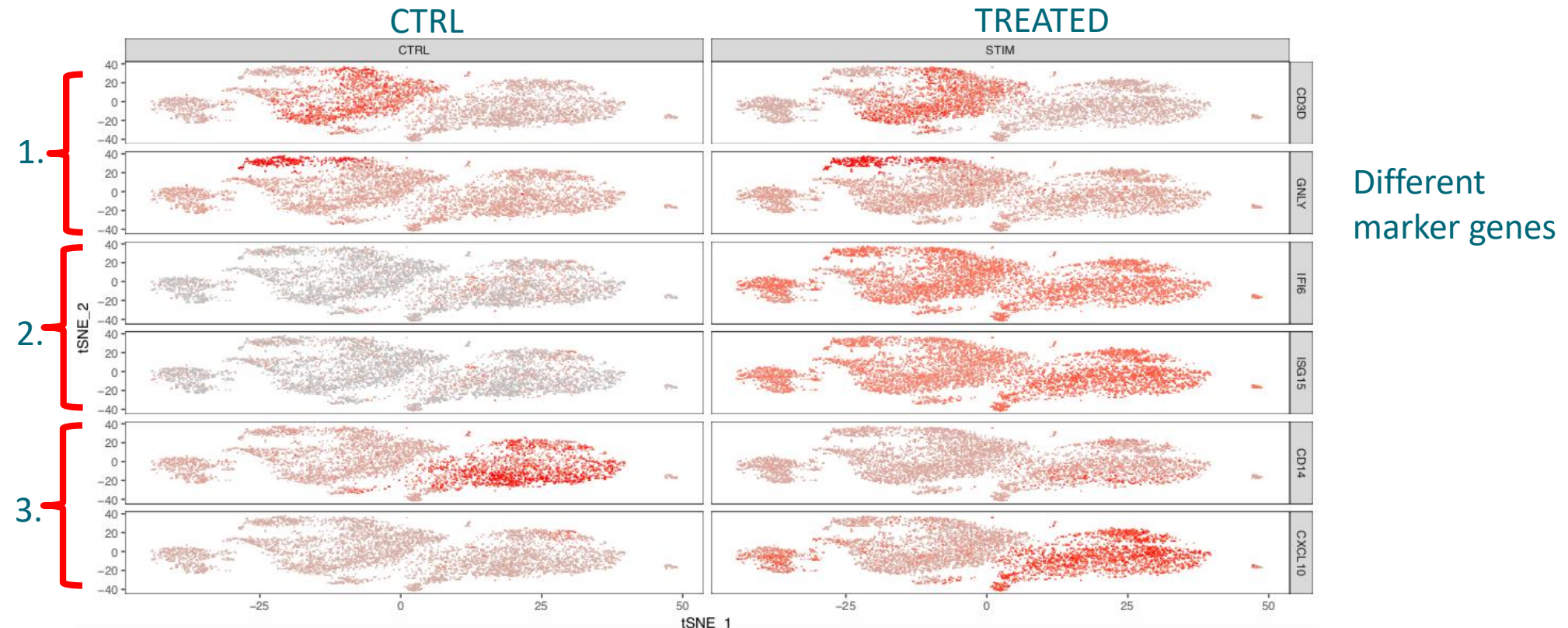
Visualize interesting genes in split dot plot

- Size = the percentage of cells in a cluster expressing a given gene
- Brightness = the average expression level in the expressing cells in a cluster



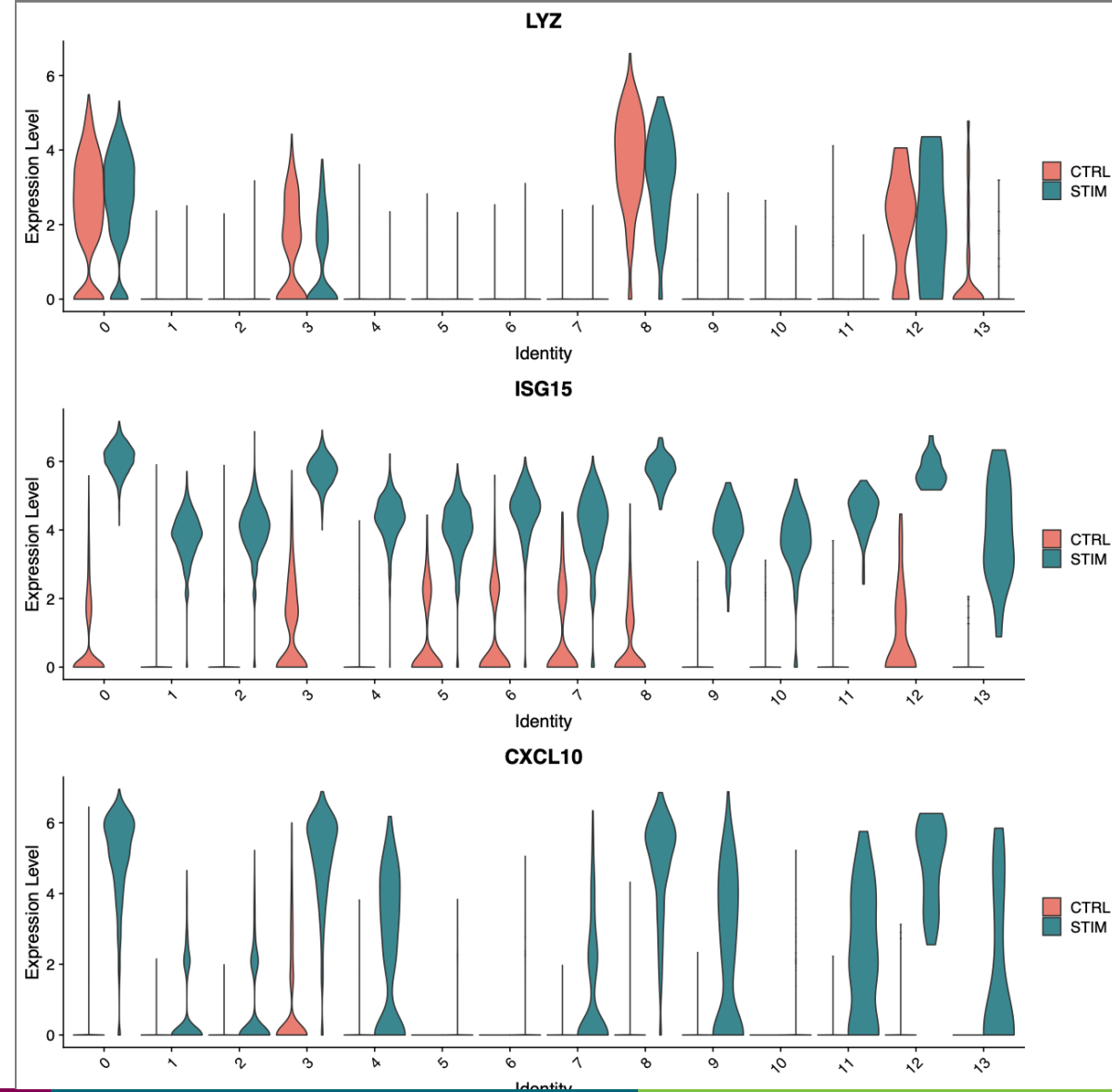
Visualize interesting genes in tSNE/UMAP plots

1. No change between the samples: conserved cell type markers
2. Change in all clusters: cell type independent marker for the treatment
3. Change in one/some clusters: cell type dependent behavior to the treatment



Visualize interesting genes in violin plots

1. No change between the samples: conserved cell type markers
2. Change in all clusters: cell type independent marker for the treatment
3. Change in one/some clusters: cell type dependent behavior to the treatment



Seurat v5: differences to v4

- “Assays”: instead of separate R-objects for each sample, one joined R-object with an “assay” for each sample. “Layers” contain the different data types (such as raw data, normalised data...)
- Pseudobulk analysis for differential expression
- SCTransform v2, improves:
 - speed and memory consumption
 - the stability of parameter estimates
 - variable feature identification in subsequent steps
- *presto* package makes differential expression analysis faster

Seurat v4 vs v5: Analysis steps for integrated analysis

