

**singleCellRNAseq**

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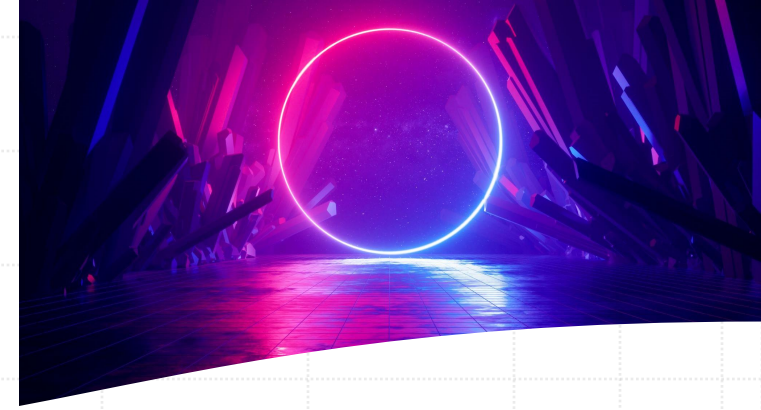
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# Introduction to Single-Cell RNA-seq

**Overview of scRNA-seq:** What it is, how it differs from bulk RNA-seq, key advantages.

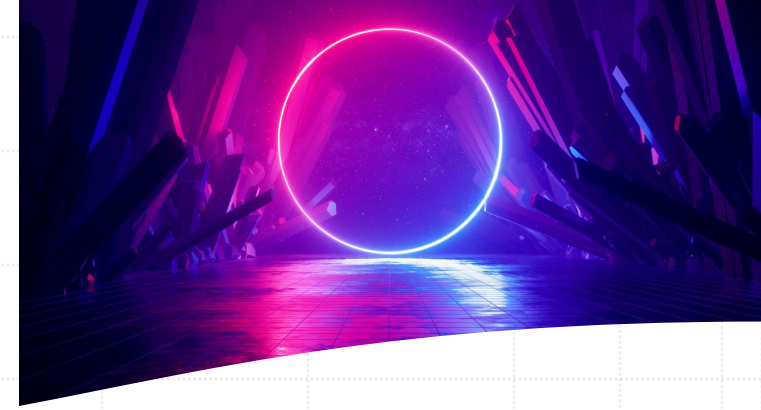
**Applications:** Understanding cellular heterogeneity, cell atlas projects, disease studies, developmental biology.



# Foundations of scRNA-seq

**Key Considerations:** Number of cells, sequencing depth, tissue preparation.

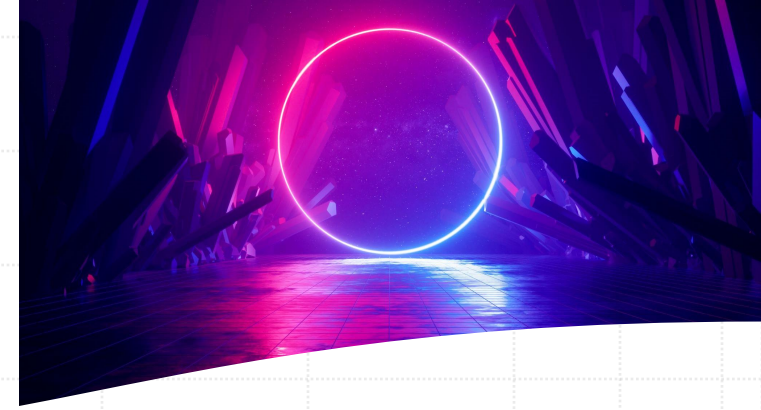
**Challenges:** Low RNA content per cell, technical noise, dropout events.



# Experimental Design

**General Workflow:** Cell isolation, library preparation, sequencing, data analysis.

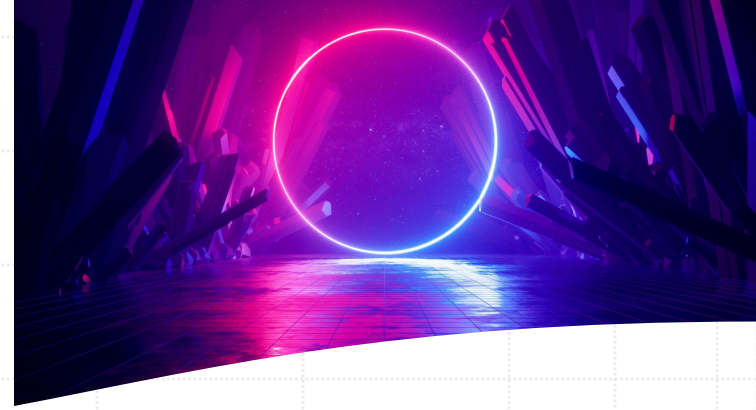
**Cell Capture Methods:** Microfluidics (e.g., 10X Genomics), FACS, Drop-seq.



# Preprocessing of scRNA-seq Data

**Initial Steps:** FASTQ generation, quality control (QC), and alignment.

**Tools:** Cell Ranger (10X Genomics), Kallisto/BUStools, Salmon/Alevin, etc.





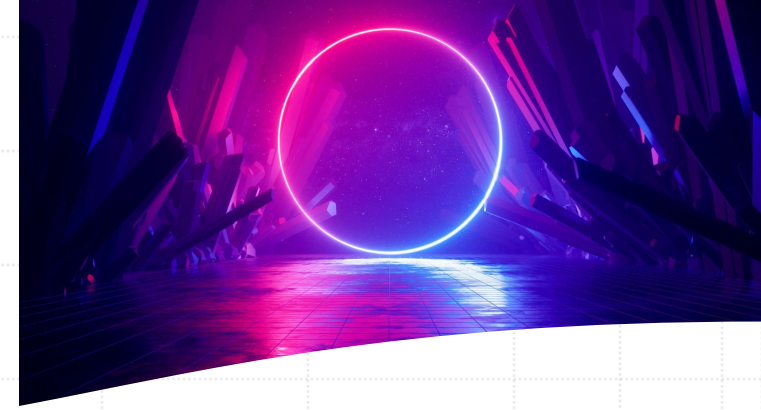
# Quantification of scRNA-seq Libraries

**UMI (Unique Molecular Identifiers):**

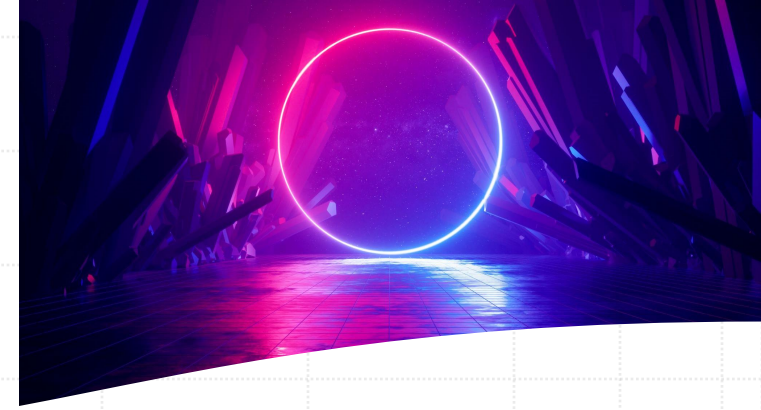
Role in counting transcripts.

**Gene-Cell Matrices:** Creation of the gene-cell count matrix.

**Dealing with Dropouts:** Technical artifacts and handling sparsity.



# Introduction to Seurat and Scanpy



**Seurat:** Overview of the R-based toolkit.

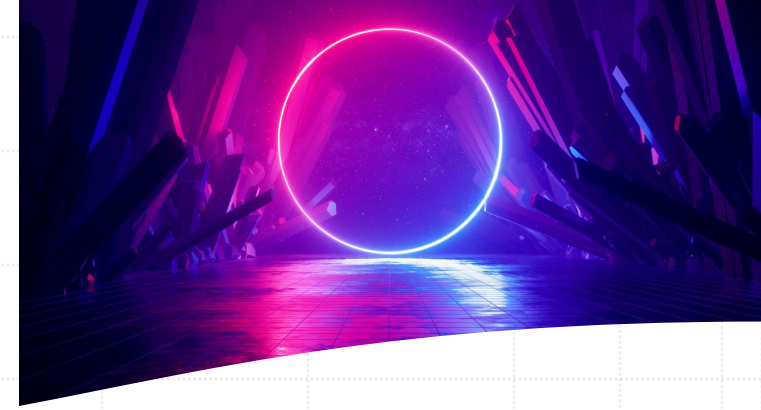
**Scanpy:** Python-based counterpart for scRNA-seq.

**Comparison:** Strengths of each tool.

# Quality Control and Filtering

**QC Metrics:** Mitochondrial content,  
number of detected genes per cell,  
total RNA per cell.

► **Filtering Low-Quality Cells:** Threshold-based filtering  
in Seurat and Scanpy.



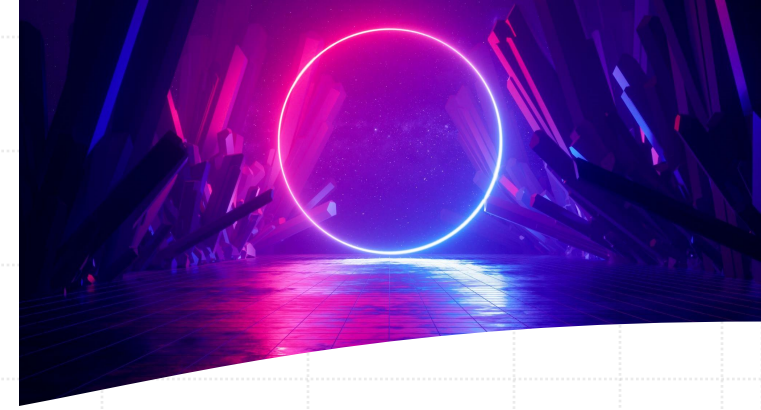


# Normalization and Scaling

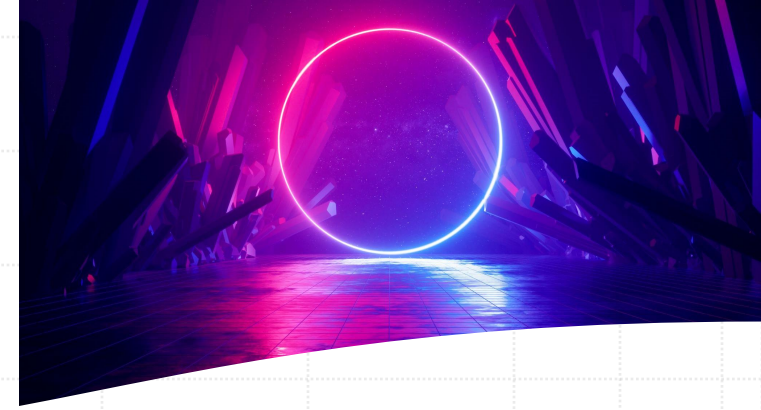
**Log-Normalization:** Adjusting for differences in sequencing depth.

**Scaling Data:** Z-score normalization.

Normalization with Seurat and with Scanpy



# Feature Selection



## Identifying Highly Variable Genes:

Why it's important, how it's done.

Seurat vs. Scanpy Commands: FindVariableFeatures  
in Seurat, `pp.highly_variable_genes` in Scanpy.

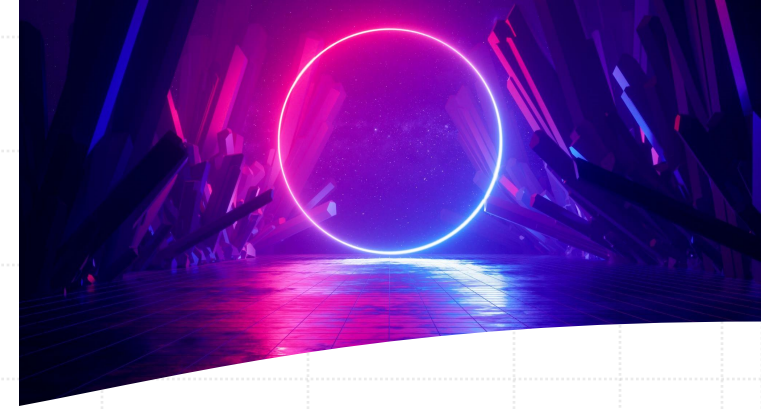
# Dimensionality Reduction

**Principal Component Analysis (PCA):**

First step for reducing data complexity.

**t-SNE and UMAP:** Techniques for visualizing high-dimensional data.

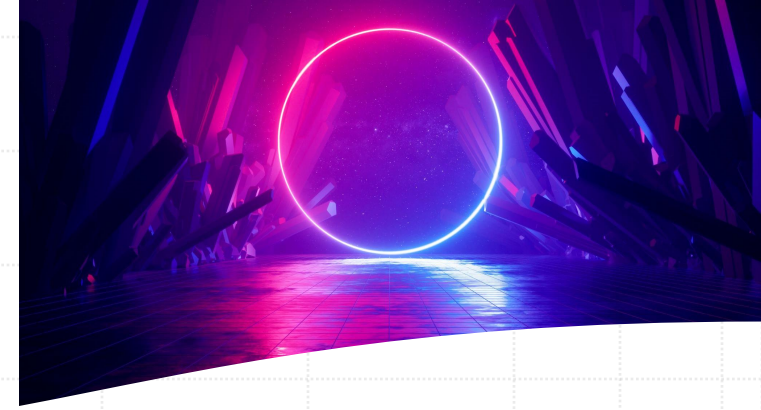
Comparison of Seurat and Scanpy Methods for Dimensionality Reduction.



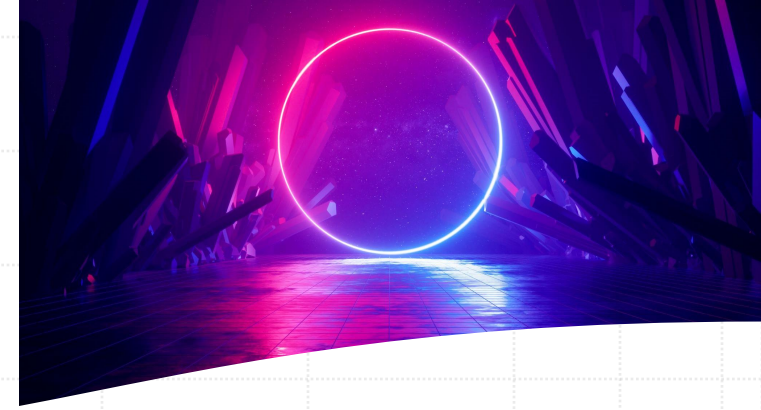
# Clustering Cells

**Clustering Algorithms:** Louvain and Leiden methods.

**Choosing Resolution:** Impact of resolution on cluster granularity.



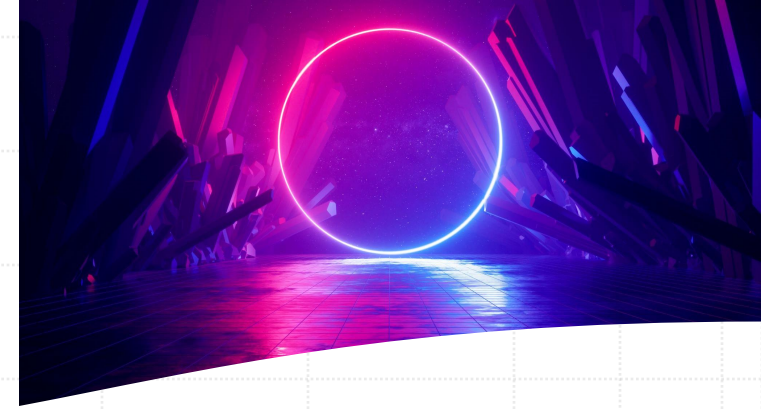
# Differential Expression Analysis



**Identifying Marker Genes:** Per-cluster analysis.

**Comparing DE Methods:** Wilcoxon Rank Sum, t-test, etc.

# Annotation of Cell Types

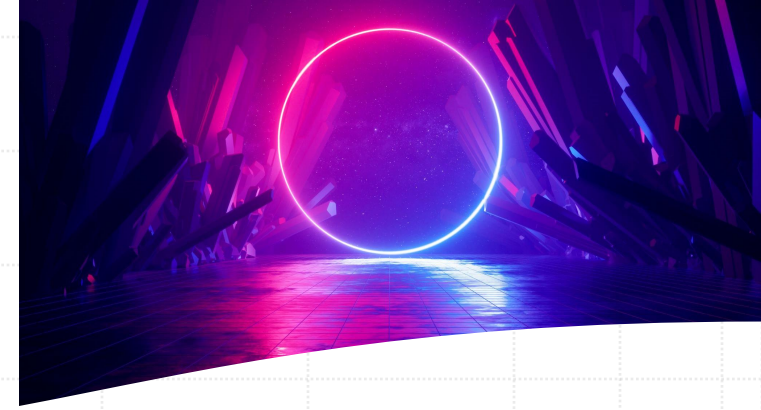


**Using Reference Datasets:** Cell type identification using tools like SingleR, Azimuth.

▶ **Manual Annotation:** Using marker gene databases and literature.



# Integration of Multiple Datasets

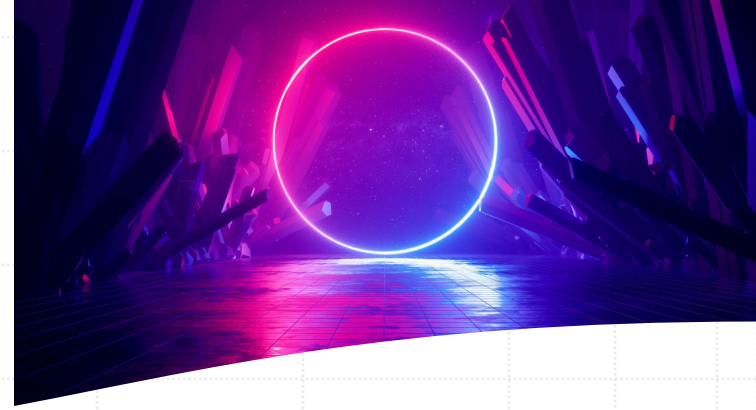


**Why Integration Matters:** Correcting for batch effects, combining multiple datasets.

Seurat's Integration Workflow

Scanpy's Harmony Method:

# Trajectory Analysis and Pseudotime



## Overview of Trajectory Analysis:

Understanding lineage progression and differentiation.

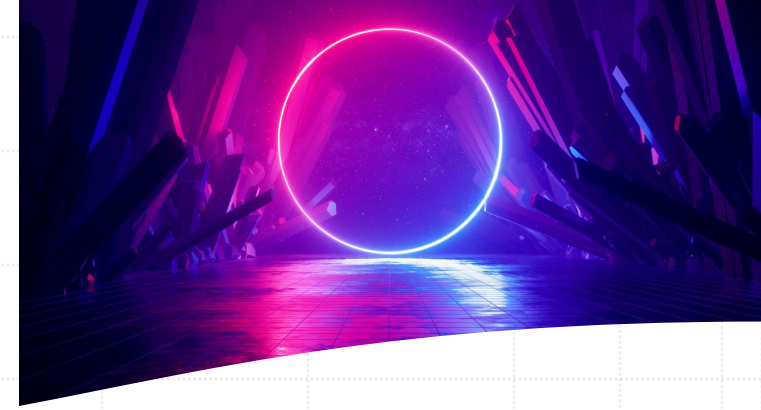
**Tools:** Monocle3, Slingshot.

**Seurat/Scanpy Integration:** Compatibility and workflows.

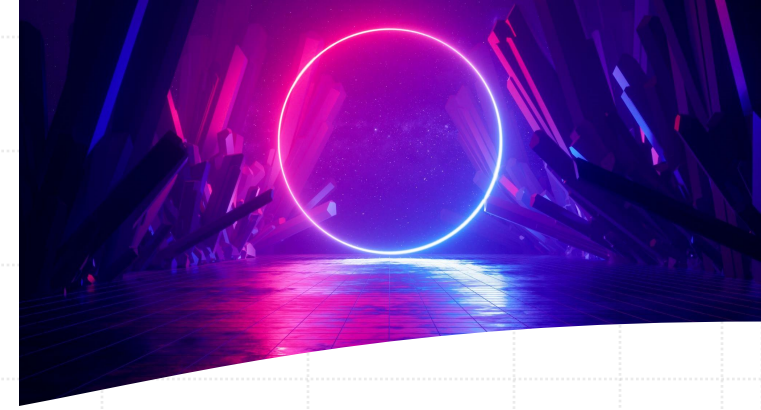
# Advanced Topics in scRNA-seq

**RNA Velocity:** Predicting future states of cells using spliced and unspliced mRNA.

**Seurat/Scanpy Integration:** scVelo in Scanpy.



# Conclusion and Future Directions



**Emerging Trends:** Multi-omics integration (CITE-seq, scATAC-seq), spatial transcriptomics.

**Challenges and Opportunities:** Scalability, data interpretation, computational demands.



A stylized illustration of a human head profile in silhouette, facing left. The interior of the head is filled with a dense network of colorful neurons. The neurons have cell bodies in shades of purple, blue, and orange, with long, branching dendrites and axons extending throughout the brain area. The background of the head is a dark, starry space. The overall style is artistic and scientific.

# Join us for this workshop...

- It will be a lot of fun!!!