

#### 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.



#### 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



Role in counting transcripts.



**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.



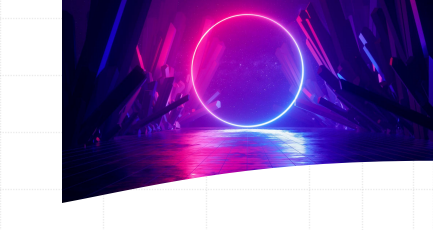
Filtering Low-Quality Cells: Threshold-based filtering in Seurat and Scanpy.

# Normalization and Scaling

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

Scaling Data: Z-score normalization.

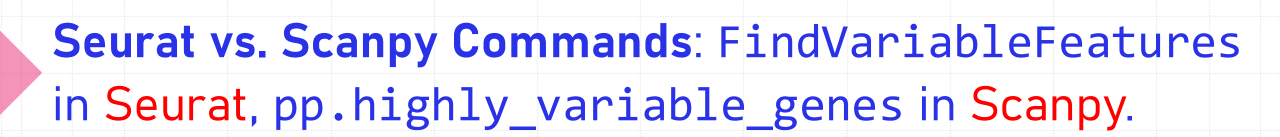
Normalization with Seurat and with Scanpy

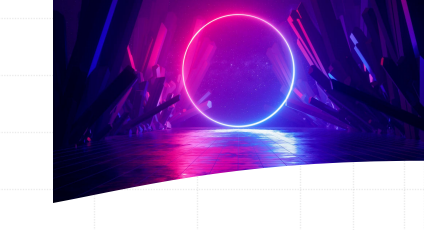


#### Feature Selection

Identifying Highly Variable Genes:

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





# **Dimensionality Reduction**

Principal Component Analysis (PCA):

First step for reducing data complexity.



Comparison of Seurat and Scanpy Methods for Dimensionality Reduction.



# **Clustering Cells**

Clustering Algorithms: Louvain and Leiden methods.





### Differential Expression Analysis

Identifying Marker Genes: Per-cluster analysis.





### **Annotation of Cell Types**

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



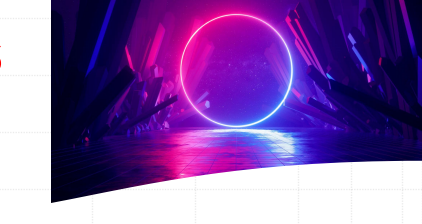


### Integration of Multiple Datasets

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

Seurat's Integration Workflow

Scanpy's Harmony Method:



#### Trajectory Analysis and Pseudotime



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.



