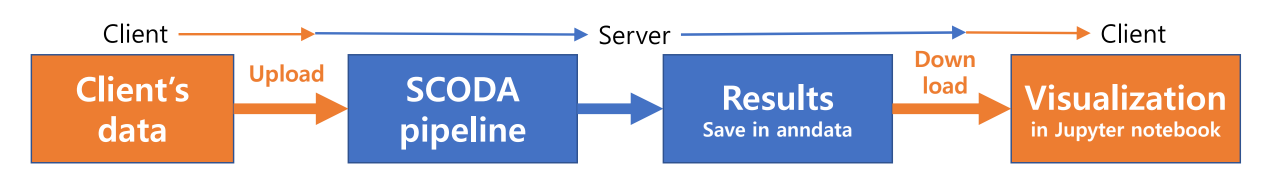


# SCODA pipeline overview (Single-Cell Omics Data Analysis pipeline)



### **Supported format**

- 1. Compressed 10x mtx files sets + (optional) meta data (csv file)
- Compressed CSV
  (cell-by-gene matrix)
  + (optional) meta
  data (csv file)
- 3. Compressed h5ad file

#### **Functions**

- 1. Celltype annotation using HiCAT (v0.6.7)
- 2. DEG & GSA/GSEA per minor celltype
- 3. Inference of cell-cell interaction using CellPhoneDB (v2.1.7)
- 4. (optional) Tumor cell identification using InferCNV (v0.4.2) + icnv addon
- 5. More will be added soon

### Results saved in anndata format (v0.8.0)

- 1. Celltype annotation in obs field
- 2. DEG & GSA/GSEA results for each celltype in uns field
- 3. CellPhoneDB results in uns field
- 4. (optional) Tumor cell identification results in obs field

### **Exploring the results**

- 1. Use SCANpy & SCODA-viz package (open source)
- 2. Example Jupyter notebook provided
- 3. With a little bit of programming, users can get deeper insight into the data by comparing cells in different condition

## SCODA-viz: visualization library

