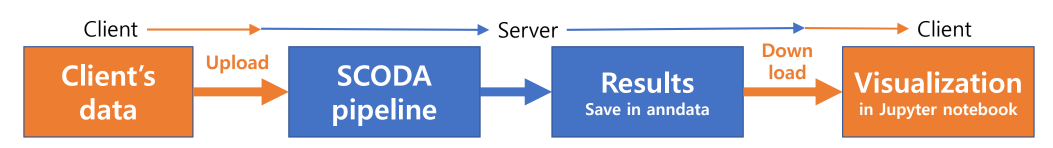
What is SCODA?

☐ Automated one-stop computing service for Single-Cell (transcript)Omics Data Analysis (single-cell RNA-seq) ☐ Especially useful for • In-vivo Tissue/tumor micro environment (TME) study. Immune cell profiling in, for example, autoimmune disease & cancer ☐ Simply uploading your count data, you get standardized functional analysis results in ten-minutes or so (depending on your data size) ☐ Results include High precision cell-type annotation (major-type, minor-type and subset) Condition specific DEG and GSA results per-cell-type Possible cell-cell interactions between different cell-types per-condition CNV estimates and tumor-cell identification results



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

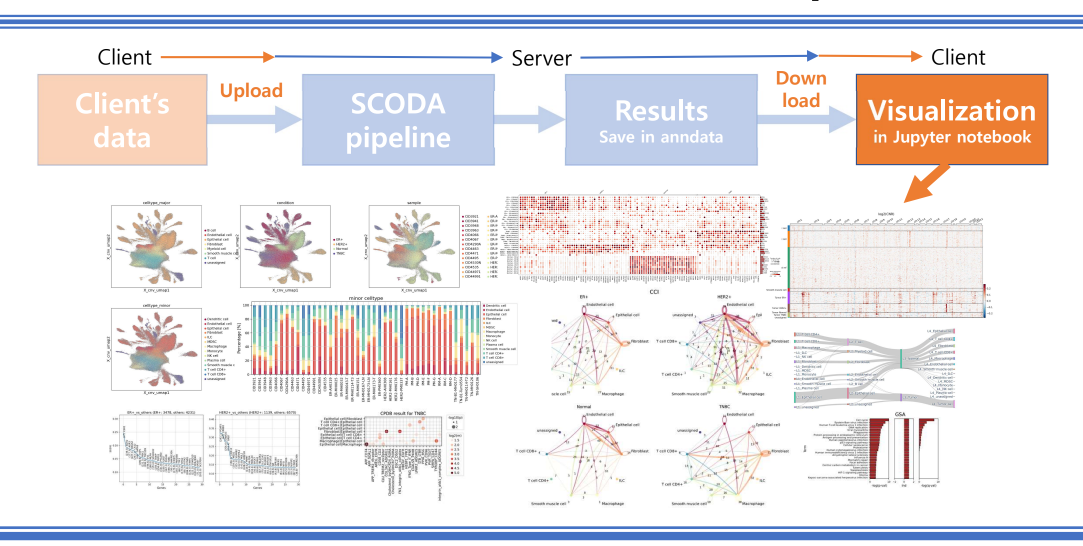
Results saved in anndata format (v0.8.0)

- 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

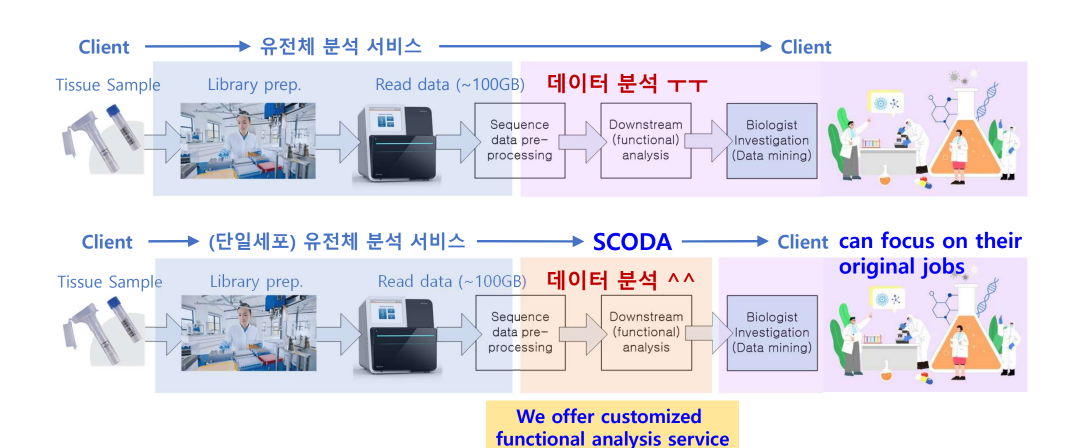
Exploring the results

- 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 lib. (free open source)



Why SCODA?



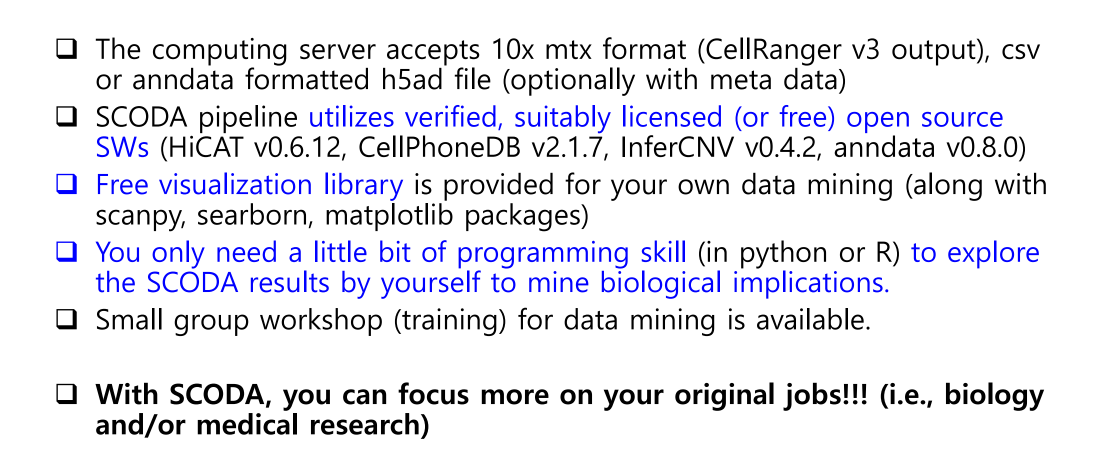
How to use SCODA?

- ☐ With your raw datasets (e.g., CellRanger v3 count data)
- 1. Go to SCODA home https://mlbi-lab.net to upload and get results.
- 2. Get Jupyter notebook to explore the results: https://github.com/combio-dku/scoda_explorer
- 3. Mine the SCODA results in Jupyter notebook on your own computer (or in Google Colab)
- 4. Contact us if you have larger datasets
- ☐ Contact: mlbi.dku@gmail.com for any inquiries or online workshop for using SCODA-viz

Why SCODA?

Trend in bio/medical research: convergence of data science & bio/medicine Requires complicated communications between biologist and informatician.
SCODA is an automated one-stop computing service using verified, standardized functional analysis tools for single-cell omics data.
With SCODA, you don't need to spend your time to do something that are not your field of expertise, e.g., of for complicated computing environment setup of develop pipelines to preprocess your datasets
Can accelerate the research with single-cell RNA-seq data. The only thing you need is a little-bit of programming skill, e.g., handling data frame, for your own data mining.
It provides you with a much higher degree of freedom for your research.

What is SCODA?



Try SCODA now

Are you ready to unravel the mysteries of single-cell gene expression? And gain deeper insights into cellular heterogeneity?
Look no further!
Our single-cell RNA-seq data analysis pipeline was designed to empower your research with state-of-the-art computational capabilities.
The visualization and data mining is fully customizable with a little bit of programming skill.
You can tailor the analysis pipeline to your specific needs.
With optional analysis parameter settings, you have control to adapt workflows for your specific requirement.
Don't hesitate to contact us (mlbi.dku@gmail.com). We're here to help you.