
scCloud Documentation

Release 0.7.0

Bo Li, Siranush Sarkizova, Joshua Gould, Marcin Tabaka, Orr Ash

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scCloud is a tool for analyzing transcriptomes of millions of single cells. It is a command line tool, a python package and a base for Cloud-based analysis workflows.

VERSION 0.7.0 *OCTOBER 26, 2018*

Added support for CITE-Seq analysis.

VERSION 0.6.0 *OCTOBER 23, 2018*

Renamed scrtools to scCloud. Added demuxEM module for cell/nuclei-hashing.

VERSION 0.5.0 *AUGUST 21, 2018*

Fixed a problem related AnnData. Added support for BigQuery.

VERSION 0.4.0 *AUGUST 2, 2018*

Added mouse brain markers. Allow aggregate matrix to take 'Sample' as attribute.

VERSION 0.3.0 JUNE 26, 2018

scrttools supports fast preprocessing, batch-correction, dimension reduction, graph-based clustering, diffusion maps, force-directed layouts, and differential expression analysis, annotate clusters, and plottings.

5.1 Installation

Install **scCloud** locally via [Miniconda](#):

```
wget https://repo.continuum.io/miniconda/Miniconda3-latest-Linux-x86_64.sh .
bash Miniconda3-latest-Linux-x86_64.sh -p /users/foo/miniconda3
mv Miniconda3-latest-Linux-x86_64.sh /users/foo/miniconda3
conda create -n scCloud -y pip
source activate scCloud
conda install -y -c anaconda numpy
conda install -y -c anaconda cython
conda install -y -c conda-forge fftw
export CPATH=$CPATH:/users/foo/miniconda3/envs/scCloud/include
pip install pybind11
git clone https://github.com/nmslib/hnswlib hnswlib
cd hnswlib/python_bindings
python setup.py install
cd ../../
git clone https://github.com/broadinstitute/scRNA-Seq.git scRNA-Seq
cd scRNA-Seq/scCloud
pip install .
cd ../../
```

5.1.1 Use scCloud in UGER

First, you need to request a RedHat7 server:

```
qcrsh -q interactive -l h_vmem=4g -l os=RedHat7 -P regevlab
```

Then, if you have installed **scCloud**, you could activate the virtual environment:

```
source activate scCloud
```

Or, you can use an installed version by typing:

```
source /ahg/regevdata/users/libo/miniconda3/bin/activate scCloud
```

5.2 Use scCloud as a command line tool

scCloud can be used as a command line tool. Type:

```
scCloud -h
```

to see the help information:

```
Usage:
  scCloud <command> [<args>...]
  scCloud -h | --help
  scCloud -v | --version
```

scCloud has 10 sub-commands in 6 groups.

- Preprocessing:
 - aggregate_matrix** Aggregate cellranger-outputted channel-specific count matrices into a single count matrix. It also enables users to import metadata into the count matrix.
- Demultiplexing:
 - demuxEM** Demultiplex cells/nuclei based on DNA barcodes for cell-hashing and nuclei-hashing data.
- Analyzing:
 - cluster** Perform first-pass analysis using the count matrix generated from ‘aggregate_matrix’. This subcommand could perform low quality cell filtration, batch correction, variable gene selection, dimension reduction, diffusion map calculation, graph-based clustering, tSNE visualization. The final results will be written into h5ad-formatted file, which Seurat could load.
 - de_analysis** Detect markers for each cluster by performing differential expression analysis per cluster (within cluster vs. outside cluster). DE tests include Welch’s t-test, Fisher’s exact test, Mann-Whitney U test. It can also calculate AUROC values for each gene.
 - annotate_cluster** This subcommand is used to automatically annotate cell types for each cluster based on existing markers. Currently, it works for human/mouse immune/brain cells.
- Plotting:
 - plot** Make static plots, which includes plotting tSNEs by cluster labels and different groups.
 - iplot** Make interactive plots using plotly. The outputs are HTML pages. You can visualize diffusion maps with this sub-command.
- Subclustering:
 - view** View attribute (e.g. cluster labels) and their values. This subcommand is used to determine cells to run subcluster analysis.
 - subcluster** Perform sub-cluster analyses on a subset of cells from the analyzed data (i.e. ‘cluster’ output).
- Single cell portal:
 - scp_output** Generate output files for single cell portal.
- CITE-Seq:
 - merge_rna_adt** Merge RNA and ADT matrices into one 10x-formatted hdf5 file.

5.2.1 Quick guide

Suppose you have `example.csv` ready with the following contents:

```
Sample,Source,Platform,Donor,Reference,Location
sample_1,bone_marrow,NextSeq,1,GRCh38,/my_dir/sample_1/filtered_gene_bc_matrices_h5.h5
sample_2,bone_marrow,NextSeq,2,GRCh38,/my_dir/sample_2/filtered_gene_bc_matrices_h5.h5
sample_3,pbmc,NextSeq,1,GRCh38,/my_dir/sample_3/filtered_gene_bc_matrices_h5.h5
sample_4,pbmc,NextSeq,2,GRCh38,/my_dir/sample_4/filtered_gene_bc_matrices_h5.h5
```

You want to analyze all four samples but correct batch effects for bone marrow and pbmc samples separately. You can run the following commands:

```
scCloud aggregate_matrix --attributes Source,Platform,Donor example.csv example
scCloud cluster -p 20 --correct-batch-effect --batch-group-by Source -run-louvain --
↳run-tsne example_10x.h5 example
scCloud de_analysis --labels louvain_labels -p 20 --fisher example.h5ad example_de.
↳xlsx
scCloud annotate_cluster example.h5ad example.anno.txt
scCloud plot composition --cluster-labels louvain_labels --attribute Donor --style_
↳normalized --not-stacked example.h5ad example.composition.png
scCloud plot scatter --basis tsne --attributes louvain_labels,Donor example.h5ad_
↳example.scatter.png
scCloud iplot --attribute louvain_labels diffmap_pca example.h5ad example.diffmap.html
```

The above analysis will give you tSNE, louvain cluster labels and diffusion maps in `example.h5ad`. You can investigate donor-specific effects by looking at `example.composition.png`, `example.scatter.png` plotted tSNE colored by `louvain_labels` and Donor info side-by-side. You can explore the diffusion map in 3D by looking at `example.diffmap.html`. This html maps all diffusion components into 3D using PCA.

If you want to perform subcluster analysis by combining cluster 1 and 3, run the following command:

```
scCloud subcluster -p 20 --correct-batch-effect example.h5ad 1,3 example_sub
```

5.2.2 scCloud aggregate_matrix

The first step for single cell analysis is to generate one count matrix from cellranger's channel-specific count matrices. `scCloud aggregate_matrix` allows aggregating arbitrary matrices with the help of a CSV file.

Type:

```
scCloud aggregate_matrix -h
```

to see the usage information:

```
Usage:
    scCloud aggregate_matrix <csv_file> <output_name> [--restriction <restriction>
↳... --attributes <attributes> --google-cloud --select-only-singlets --minimum-
↳number-of-genes <ngene> --dropseq-genome <genome>]
    scCloud aggregate_matrix -h
```

- Arguments:

csv_file Input csv-formatted file containing information of each scRNA-Seq run. Each row must contain at least 2 columns — Sample, sample name and Location, location of the channel-specific count matrix in either 10x format (e.g. /sample/filtered_gene_bc_matrices_h5.h5) or dropseq format (e.g. /sample/sample.umi.dge.txt.gz). See below for an example csv:

```
Sample,Source,Platform,Donor,Reference,Location
sample_1,bone_marrow,NextSeq,1,GRCh38,/my_dir/sample_1/filtered_gene_
↳bc_matrices_h5.h5
sample_2,bone_marrow,NextSeq,2,GRCh38,/my_dir/sample_2/filtered_gene_
↳bc_matrices_h5.h5
sample_3,pbmc,NextSeq,1,GRCh38,/my_dir/sample_3/filtered_gene_bc_
↳matrices_h5.h5
sample_4,pbmc,NextSeq,2,GRCh38,/my_dir/sample_4/filtered_gene_bc_
↳matrices_h5.h5
```

output_name The output file name.

- Options:

--restriction <restriction>... Select channels that satisfy all restrictions. Each restriction takes the format of name:value,...,value or name:~value,...,value, where ~ refers to not. You can specify multiple restrictions by setting this option multiple times.

--attributes <attributes> Specify a comma-separated list of outputted attributes. These attributes should be column names in the csv file.

--google-cloud If files are stored in google cloud. Assuming google cloud sdk is installed.

--select-only-singlets If we have demultiplexed data, turning on this option will make scCloud only include barcodes that are predicted as singlets.

--minimum-number-of-genes <ngene> Only keep barcodes with at least <ngene> expressed genes.

--dropseq-genome <genome> If inputs are dropseq data, this option needs to turn on and provides the reference genome name.

-h, --help Print out help information.

- Outputs:

output_name_10x.h5 A 10x-formatted HDF5 file containing the count matrices and associated attributes.

- Examples:

```
scCloud aggregate_matrix --restriction Source:pbmc --restriction Donor:1 --
↳attributes Source,Platform,Donor example.csv example
```

5.2.3 scCloud demuxEM

If you have data generated by cell-hashing or nuclei-hashing, you can use `scCloud demuxEM` to demultiplex your data.

Type:

```
scCloud demuxEM -h
```

to see the usage information:

Usage:

```
scCloud demuxEM --hash-type <type> [options] <input_adt_csv_file> <input_raw_
→gene_bc_matrices_h5.h5> <output_name>
scCloud demuxEM -h
```

- Arguments:

input_adt_csv_file Input ADT (antibody tag) count matrix in CSV format.

input_raw_gene_bc_matrices_h5.h5 Input raw RNA expression matrix in 10x hdf5 format.

output_name Output name. All outputs will use it as the prefix.

- Options:

-hash-type <type> The hash type of the data. <type> can be 'cell-hashing' for cell-hashing and 'nuclei-hashing' for nuclei-hashing.

-p <number>, -threads <number> Number of threads. [default: 1]

-genome <genome> Reference genome name. If not provided, we will infer it from the expression matrix file.

-min-num-genes <number> We only demultiplex cells/nuclei with at least <number> expressed genes. [default: 100]

-max-background-probability <prob> Any cell/nucleus with no less than <prob> background probability will be marked as unknown. [default: 0.8]

-prior-on-samples <prior> The sparse prior put on samples.

-random-state <seed> The random seed used in the KMeans algorithm to separate empty ADT droplets from others. [default: 0]

-generate-diagnostic-plots Generate a series of diagnostic plots, including the background/signal between HTO counts, estimated background probabilities, HTO distributions of cells and non-cells etc.

-generate-gender-plot <genes> Generate violin plots using gender-specific genes (e.g. Xist). <gene> is a comma-separated list of gene names.

-h, -help Print out help information.

- Outputs:

output_name_demux_10x.h5 RNA expression matrix with demultiplexed sample identities in 10x's hdf5 format.

output_name_ADTs.h5ad Antibody tag matrix in h5ad format.

output_name_demux.h5ad Demultiplexed RNA count matrix in h5ad format.

output_name.ambient_hashtag.hist.png Optional output. A histogram plot depicting hashtag distributions of empty droplets and non-empty droplets.

output_name.background_probabilities.bar.png Optional output. A bar plot visualizing the estimated hashtag background probability distribution.

output_name.real_content.hist.png Optional output. A histogram plot depicting hashtag distributions of not-real-cells and real-cells as defined by total number of expressed genes in the RNA assay.

output_name.rna_demux.hist.png Optional output. A histogram plot depicting RNA UMI distribution for singlets, doublets and unknown cells.

output_name.gene_name.violin.png Optional outputs. Violin plots depicting gender-specific gene expression across samples. We can have multiple plots if a gene list is provided in ‘--generate-gender-plot’ option.

- Examples:

```
scCloud demuxEM -p 8 --hash-type cell-hashing --generate-diagnostic-plots example_
→adt.csv example_raw_gene_bc_matrices_h5.h5 example_output
```

5.2.4 scCloud cluster

Once we collected the count matrix `example_10x.h5`, we can perform single cell analysis using `scCloud cluster`.

Type:

```
scCloud cluster -h
```

to see the usage information:

```
Usage:
  scCloud cluster [options] <input_file> <output_name>
  scCloud cluster -h
```

- Arguments:

input_file Input file in 10x format. If first-pass analysis has been performed, but you want to run some additional analysis, you could also pass a h5ad-formatted file.

output_name Output file name. All outputs will use it as the prefix.

- Options:

-p <number>, --threads <number> Number of threads. [default: 1]

--processed Input file is processed and thus no PCA & diffmap will be run.

--genome <genome> A string contains comma-separated genome names. `scCloud` will read all groups associated with genome names in the list from the hdf5 file. If genome is None, all groups will be considered.

--cite-seq Data are CITE-Seq data. `scCloud` will perform analyses on RNA count matrix first. Then it will attach the ADT matrix to the RNA matrix with all antibody names changing to ‘AD-’ + antibody_name. Lastly, it will embed the antibody expression using t-SNE (the basis used for plotting is ‘citeseq_tsne’).

--output-filtration-results <spreadsheet> Output filtration results into <spreadsheet>.

--output-seurat-compatible Output seurat-compatible h5ad file.

--output-loom Output loom-formatted file.

--correct-batch-effect Correct for batch effects.

--batch-group-by <expression> Batch correction assumes the differences in gene expression between channels are due to batch effects. However, in many cases, we know that channels can be partitioned into several groups and each group is biologically different from others. In this case, we will only perform batch correction for channels

within each group. This option defines the groups. If `<expression>` is `None`, we assume all channels are from one group. Otherwise, groups are defined according to `<expression>`. `<expression>` takes the form of either `'attr'`, or `'attr1+attr2+...+attrn'`, or `'attr=value11,...,value1n_1;value21,...,value2n_2;...;valuem1,...,valuemn_m'`. In the first form, `'attr'` should be an existing sample attribute, and groups are defined by `'attr'`. In the second form, `'attr1',..., 'attrn'` are `n` existing sample attributes and groups are defined by the Cartesian product of these `n` attributes. In the last form, there will be `m + 1` groups. A cell belongs to group `i` (`i > 0`) if and only if its sample attribute `'attr'` has a value among `valuei1,...,valuein_i`. A cell belongs to group 0 if it does not belong to any other groups.

- min-genes <number>** Only keep cells with at least `<number>` of genes. [default: 500]
- max-genes <number>** Only keep cells with less than `<number>` of genes. [default: 6000]
- mito-prefix <prefix>** Prefix for mitochondrial genes. [default: MT-]
- percent-mito <ratio>** Only keep cells with mitochondrial ratio less than `<ratio>`. [default: 0.1]
- gene-percent-cells <ratio>** Only use genes that are expressed in at `<ratio>` * 100 percent of cells to select variable genes. [default: 0.0005]
- counts-per-cell-after <number>** Total counts per cell after normalization. [default: 1e5]
- random-state <seed>** Random number generator seed. [default: 0]
- run-uncentered-pca** Run uncentered PCA.
- no-variable-gene-selection** Do not select variable genes.
- no-submat-to-dense** Do not convert variable-gene-selected submatrix to a dense matrix.
- nPC <number>** Number of PCs. [default: 50]
- nDC <number>** Number of diffusion components. [default: 50]
- diffmap-alpha <alpha>** Power parameter for diffusion-based pseudotime. [default: 0.5]
- diffmap-K <K>** Number of neighbors used for constructing affinity matrix. [default: 100]
- diffmap-full-speed** For the sake of reproducibility, we only run one thread for building kNN indices. Turn on this option will allow multiple threads to be used for index building. However, it will also reduce reproducibility due to the racing between multiple threads.
- calculate-pseudotime <roots>** Calculate diffusion-based pseudotimes based on `<roots>`. `<roots>` should be a comma-separated list of cell barcodes.
- run-louvain** Run louvain clustering algorithm.
- louvain-resolution <resolution>** Resolution parameter for the louvain clustering algorithm. [default: 1.3]
- louvain-affinity <affinity>** Affinity matrix to be used. Could be `'W_norm'`, `'W_diffmap'`, or `'W_diffmap_norm'`. [default: `W_norm`]
- run-kmeans** Run KMeans clustering algorithm on diffusion components.
- kmeans-n-clusters <number>** Target at `<number>` clusters for K means. [default: 20]
- run-hdbscan** Run hdbscan clustering algorithm on diffusion components.
- hdbscan-min-cluster-size <number>** Minimum cluster size for hdbscan. [default: 50]
- hdbscan-min-samples <number>** Minimum number of samples for hdbscan. [default: 50]
- run-approximated-louvain** Run approximated louvain clustering algorithm.
- approx-louvain-ninit <number>** Number of Kmeans tries. [default: 20]

-approx-louvain-nclusters <number> Number of clusters for Kmeans initialization. [default: 30]
-approx-louvain-resolution <resolution>. Resolution parameter for louvain. [default: 1.3]
-run-tsne Run multi-core t-SNE for visualization.
-tsne-perplexity <perplexity> t-SNE's perplexity parameter. [default: 30]
-run-fitsne Run FIt-SNE for visualization.
-run-umap Run umap for visualization.
-umap-on-diffmap Run umap on diffusion components.
-umap-K <K> K neighbors for umap. [default: 15]
-umap-min-dist <number> Umap parameter. [default: 0.1]
-umap-spread <spread> Umap parameter. [default: 1.0]
-run-fle Run force-directed layout embedding.
-fle-K <K> K neighbors for building graph for FLE. [default: 50]
-fle-n-steps <nstep> Number of iterations for FLE. [default: 10000]
-fle-affinity <affinity> Affinity matrix to be used. Could be 'W_diffmap', or 'W_diffmap_norm'. [default: W_diffmap]
-h, -help Print out help information.

- Outputs:

output_name.h5ad Output file in h5ad format. The clustering results are stored in the 'obs' field (e.g. 'louvain_labels' for louvain cluster labels). The PCA, t-SNE and diffusion map coordinates are stored in the 'obsm' field.

output_name.seurat.h5ad Optional output. Only exists if '-output-seurat-compatible' is set. This is the Seurat-readable h5ad file.

output_nameloom Optional output. Only exists if '-output-loom' is set. output_name.h5ad in loom format for visualization.

- Examples:

```
scCloud cluster -p 20 --correct-batch-effect --run-louvain --run-tsne example_10x.
↪h5 example
```

5.2.5 scCloud de_analysis

Once we have the clusters, we can detect markers using `scCloud de_analysis`.

Type:

```
scCloud de_analysis -h
```

to see the usage information:

```
Usage:
    scCloud de_analysis [--labels <attr> -p <threads> --alpha <alpha> --fisher --
↪mwu --roc] <input_h5ad_file> <output_spreadsheet>
    scCloud de_analysis -h
```

- Arguments:

input_h5ad_file Single cell data with clustering calculated. DE results would be written back.

output_spreadsheet Output spreadsheet with DE results.

- Options:

-labels <attr> <attr> used as cluster labels. [default: louvain_labels]

-alpha <alpha> Control false discovery rate at <alpha>. [default: 0.05]

-fisher Calculate Fisher's exact test.

-mwu Calculate Mann-Whitney U test.

-roc Calculate area under curve in ROC curve.

-p <threads> Use <threads> threads. [default: 1]

-h, -help Print out help information.

- Outputs:

input_h5ad_file DE results would be written back to the 'var' fields.

output_spreadsheet An excel spreadsheet containing DE results. Each cluster has two tabs in the spreadsheet. One is for up-regulated genes and the other is for down-regulated genes.

- Examples:

```
scCloud de_analysis --labels louvain_labels -p 20 --fisher --mwu --roc example.
↪h5ad example_de.xlsx
```

5.2.6 scCloud annotate_cluster

Once we have the DE results, we could optionally identify putative cell types for each cluster using `scCloud annotate_cluster`. Currently, this subcommand works for human/mouse immune/brain cells. This command has two forms: the first form generates putative annotations and the second form write annotations into the h5ad object.

Type:

```
scCloud annotate_cluster -h
```

to see the usage information:

```
Usage:
  scCloud annotate_cluster [--json-file <file> --minimum-report-score <score> --
↪do-not-use-non-de-genes] <input_h5ad_file> <output_file>
  scCloud annotate_cluster --annotation <annotation_string> <input_h5ad_file>
  scCloud annotate_cluster -h
```

- Arguments:

input_h5ad_file Single cell data with DE analysis done by `scCloud de_analysis`.

output_file Output annotation file.

- Options:

- json-file <file>** JSON file for markers. Could also be human_immune/mouse_immune/mouse_brain/human_brain, which triggers scCloud to markers included in the package. [default: human_immune]
- minimum-report-score <score>** Minimum cell type score to report a potential cell type. [default: 0.5]
- do-not-use-non-de-genes** Do not count non DE genes as down-regulated.
- annotation <annotation_string>** Write cell type annotations in <annotation_string> into <input_h5ad_file>. <annotation_string> has this format: 'anno_attr:anno_1;anno_2;...;anno_n'. 'anno_attr' is the annotation attribute in the h5ad object and anno_i is the annotation for cluster i.
- h, -help** Print out help information.

- **Outputs:**

output_file This is a text file. For each cluster, all its putative cell types are listed in descending order of the cell type score. For each putative cell type, all markers support this cell type are listed. If one putative cell type has cell subtypes, all subtypes will be listed under this cell type.

- **Examples:**

```
scCloud annotate_cluster example.h5ad example.anno.txt
scCloud annotate_cluster --annotation "anno:T cells;B cells;NK cells;Monocytes"
↪example.h5ad
```

5.2.7 scCloud plot

We can make a variety of figures using `scCloud plot`.

Type:

```
scCloud plot -h
```

to see the usage information:

```
Usage:
    scCloud plot [options] [--restriction <restriction>...] <plot_type> <input_
↪h5ad_file> <output_file>
    scCloud plot -h
```

- **Arguments:**

plot_type Only 2D plots, chosen from 'composition', 'scatter', 'scatter_groups', 'scatter_genes', 'scatter_gene_groups', and 'heatmap'.

input_h5ad_file Single cell data with clustering done by Scanpy in h5ad file format.

output_file Output image file.

- **Options:**

-dpi <dpi> DPI value for the figure. [default: 500]

-cluster-labels <attr> Use <attr> as cluster labels. This option is used in 'composition', 'scatter_groups', and 'heatmap'.

-attribute <attr> Plot <attr> against cluster labels. This option is only used in 'composition'.

- basis <basis>** Basis for 2D plotting, chosen from 'tsne', 'fitsne', 'umap', 'pca', 'rpca', 'fle', or 'diffmap_pca'. This option is used in 'scatter', 'scatter_groups', 'scatter_genes', and 'scatter_gene_groups'. [default: tsne]
- attributes <attrs>** <attrs> is a comma-separated list of attributes to color the basis. This option is only used in 'scatter'.
- restriction <restriction>...** Set restriction if you only want to plot a subset of data. Multiple <restriction> strings are allowed. Each <restriction> takes the format of 'attr:value,value'. This option is used in 'composition' and 'scatter'.
- apply-to-each-figure** Indicate that the <restriction> strings are not applied to all attributes but for specific attributes. The string's 'attr' value should match the attribute you want to restrict.
- group <attr>** <attr> is used to make group plots. In group plots, the first one contains all components in the group and the following plots show each component separately. This option is used in 'scatter_groups' and 'scatter_gene_groups'. If <attr> is a semi-colon-separated string, parse the string as groups.
- genes <genes>** <genes> is a comma-separated list of gene names to visualize. This option is used in 'scatter_genes' and 'heatmap'.
- gene <gene>** Visualize <gene> in group plots. This option is only used in 'scatter_gene_groups'.
- style <style>** Composition plot styles. Can be either 'frequency', 'count', or 'normalized'. [default: frequency]
- not-stacked** Do not stack bars in composition plot.
- log-y** Plot y axis in log10 scale for composition plot.
- nrows <nrows>** Number of rows in the figure. If not set, scCloud will figure it out automatically.
- ncols <ncols>** Number of columns in the figure. If not set, scCloud will figure it out automatically.
- subplot-size <sizes>** Sub-plot size in inches, w x h, separated by comma. Note that margins are not counted in the sizes. For composition, default is (6, 4). For scatter plots, default is (4, 4).
- left <left>** Figure's left margin in fraction with respect to subplot width.
- bottom <bottom>** Figure's bottom margin in fraction with respect to subplot height.
- wspace <wspace>** Horizontal space between subplots in fraction with respect to subplot width.
- hspace <hspace>** Vertical space between subplots in fraction with respect to subplot height.
- alpha <alpha>** Point transparent parameter.
- legend-fontsize <fontsize>** Legend font size.
- use-raw** Use anndata stored raw expression matrix. Only used by 'scatter_genes' and 'scatter_gene_groups'.
- do-not-show-all** Do not show all components in group for scatter_groups.
- show-zscore** If show zscore in heatmap.
- heatmap-title <title>** Title for heatmap.
- h, -help** Print out help information.

Examples:

```

scCloud plot composition --cluster-labels louvain_labels --attribute Donor --style_
↳normalized --not-stacked example.h5ad example.composition.png
scCloud plot scatter --basis tsne --attributes louvain_labels,Donor example.h5ad_
↳example.scatter.png
scCloud plot scatter_groups --cluster-labels louvain_labels --group Donor example.
↳h5ad example.scatter_groups.png
scCloud plot scatter_genes --genes CD8A,CD4,CD3G,MS4A1,NCAM1,CD14,ITGAX,IL3RA,CD38,
↳CD34,PPBP example.h5ad example.genes.png
scCloud plot scatter_gene_groups --gene CD8A --group Donor example.h5ad example.gene_
↳groups.png
scCloud plot heatmap --cluster-labels louvain_labels --genes CD8A,CD4,CD3G,MS4A1,
↳NCAM1,CD14,ITGAX,IL3RA,CD38,CD34,PPBP --heatmap-title 'markers' example.h5ad_
↳example.heatmap.png

```

5.2.8 scCloud iplot

We can also make interactive plots in html format using `scCloud iplot`. These interactive plots are very helpful if you want to explore the diffusion maps.

Type:

```
scCloud iplot -h
```

to see the usage information:

```

Usage:
    scCloud iplot --attribute <attr> [options] <basis> <input_h5ad_file> <output_
↳html_file>
    scCloud iplot -h

```

- Arguments:

basis Basis can be either ‘tsne’, ‘fitsne’, ‘umap’, ‘diffmap’, ‘pca’, ‘rpca’ or ‘diffmap_pca’.

input_h5ad_file Single cell data with clustering done in h5ad file format.

output_html_file Output interactive plot in html format.

- Options:

-attribute <attr> Use attribute <attr> as labels in the plot.

-is-real <attr> is real valued.

-is-gene <attr> is a gene name.

-log10 If take log10 of real values.

-h, -help Print out help information.

- Examples:

```

scCloud iplot --attribute louvain_labels tsne example.h5ad example.tsne.html
scCloud iplot --attribute louvain_labels diffmap_pca example.h5ad example.diffmap.
↳html

```

5.2.9 scCloud view

We may want to further perform sub-cluster analysis on a subset of cells. This sub-command helps us to define the subset.

Type:

```
scCloud view -h
```

to see the usage information:

```
Usage:
    scCloud view [--show-attributes --show-gene-attributes --show-values-for-
    ↪attributes <attributes>] <input_h5ad_file>
    scCloud view -h
```

- Arguments:
 - input_h5ad_file** Analyzed single cell data in h5ad format.
- Options:
 - show-attributes** Show the available sample attributes in the input dataset.
 - show-gene-attributes** Show the available gene attributes in the input dataset.
 - show-values-for-attributes <attributes>** Show the available values for specified attributes in the input dataset. <attributes> should be a comma-separated list of attributes.
 - h, --help** Print out help information.
- Examples:

```
scCloud view --show-attributes example.h5ad
scCloud view --show-gene-attributes example.h5ad
scCloud view --show-values-for-attributeslouvain_labels,Donor example.h5ad
```

5.2.10 scCloud subcluster

If there is a subset of cells that we want to further cluster, we can run `scCloud subcluster`. This sub-command will outputs a new h5ad file that you can run `de_analysis`, `plot` and `iplot` on.

Type:

```
scCloud subcluster -h
```

to see the usage information:

```
Usage:
    scCloud subcluster [options] --subset-selection <subset-selection>... <input_
    ↪file> <output_name>
    scCloud subcluster -h
```

- Arguments:
 - input_file** Single cell data with clustering done in h5ad format.
 - output_name** Output file name. All outputs will use it as the prefix.

- Options:

- subset-selection <subset-selection>...** Specify which cells will be included in the subcluster analysis. Each <subset_selection> string takes the format of 'attr:value,...,value', which means select cells with attr in the values. If multiple <subset_selection> strings are specified, the subset of cells selected is the intersection of these strings.
- p <number>, -threads <number>** Number of threads. [default: 1]
- correct-batch-effect** Correct for batch effects.
- output-seurat-compatible** Output seurat-compatible h5ad file.
- output-loom** Output loom-formatted file.
- random-state <seed>** Random number generator seed. [default: 0]
- run-uncentered-pca** Run uncentered PCA.
- no-variable-gene-selection** Do not select variable genes.
- no-submat-to-dense** Do not convert variable-gene-selected submatrix to a dense matrix.
- nPC <number>** Number of PCs. [default: 50]
- nDC <number>** Number of diffusion components. [default: 50]
- diffmap-alpha <alpha>** Power parameter for diffusion-based pseudotime. [default: 0.5]
- diffmap-K <K>** Number of neighbors used for constructing affinity matrix. [default: 100]
- diffmap-full-speed** For the sake of reproducibility, we only run one thread for building kNN indices. Turn on this option will allow multiple threads to be used for index building. However, it will also reduce reproducibility due to the racing between multiple threads.
- calculate-pseudotime <roots>** Calculate diffusion-based pseudotimes based on <roots>. <roots> should be a comma-separated list of cell barcodes.
- run-louvain** Run louvain clustering algorithm.
- louvain-resolution <resolution>** Resolution parameter for the louvain clustering algorithm. [default: 1.3]
- louvain-affinity <affinity>** Affinity matrix to be used. Could be 'W_norm', 'W_diffmap', or 'W_diffmap_norm'. [default: W_norm]
- run-kmeans** Run KMeans clustering algorithm on diffusion components.
- kmeans-n-clusters <number>** Target at <number> clusters for K means. [default: 20]
- run-hdbscan** Run hdbscan clustering algorithm on diffusion components.
- hdbscan-min-cluster-size <number>** Minimum cluster size for hdbscan. [default: 50]
- hdbscan-min-samples <number>** Minimum number of samples for hdbscan. [default: 50]
- run-approximated-louvain** Run approximated louvain clustering algorithm.
- approx-louvain-ninit <number>** Number of Kmeans tries. [default: 20]
- approx-louvain-nclusters <number>** Number of clusters for Kmeans initialization. [default: 30]
- approx-louvain-resolution <resolution>**. Resolution parameter for louvain. [default: 1.3]
- run-tsne** Run multi-core t-SNE for visualization.
- tsne-perplexity <perplexity>** t-SNE's perplexity parameter. [default: 30]
- run-fitsne** Run FIt-SNE for visualization.

- run-umap** Run umap for visualization.
- umap-on-diffmap** Run umap on diffusion components.
- umap-K <K>** K neighbors for umap. [default: 15]
- umap-min-dist <number>** Umap parameter. [default: 0.1]
- umap-spread <spread>** Umap parameter. [default: 1.0]
- run-fle** Run force-directed layout embedding.
- fle-K <K>** K neighbors for building graph for FLE. [default: 50]
- fle-n-steps <nstep>** Number of iterations for FLE. [default: 10000]
- fle-affinity <affinity>** Affinity matrix to be used. Could be 'W_diffmap', or 'W_diffmap_norm'. [default: W_diffmap]
- h, -help** Print out help information.

- Outputs:

output_name.h5ad Output file in h5ad format. The clustering results are stored in the 'obs' field (e.g. 'louvain_labels' for louvain cluster labels). The PCA, t-SNE and diffusion map coordinates are stored in the 'obsm' field.

output_name.seurat.h5ad Optional output. Only exists if '-output-seurat-compatible' is set. This is the Seurat-readable h5ad file.

output_name.loom Optional output. Only exists if '-output-loom' is set. output_name.h5ad in loom format for visualization.

- Examples:

```
scCloud subcluster --subset_selection louvain_labels:1,3 --subset_selection_
↳Donor:1 -p 20 --correct-batch-effect example.h5ad example_sub
```

5.2.11 scCloud scp_output

If we want to visualize analysis results on single cell portal (SCP), we can generate required files for SCP using this subcommand.

Type:

```
scCloud scp_output -h
```

to see the usage information:

```
Usage:
  scCloud scp_output <input_h5ad_file> <output_name>
  scCloud scp_output -h
```

- Arguments:

input_h5ad_file Analyzed single cell data in h5ad format.

output_name Name prefix for all outputted files.

- Options:

-h, -help Print out help information.

- Outputs:

output_name.scp.metadata.txt, output_name.scp.barcodes.tsv, output_name.scp.genes.tsv, output_name.scp.matrix.mtx
Files that single cell portal needs.

- Examples:

```
scCloud scp_output example.h5ad example
```

5.2.12 scCloud merge_rna_adt

If we have CITE-Seq data, we can merge RNA count matrix and ADT (antibody tag) count matrix into one file using this subcommand.

Type:

```
scCloud merge_rna_adt -h
```

to see the usage information:

```
Usage:
  scCloud merge_rna_adt <input_raw_gene_bc_matrices_h5.h5> <input_adt_csv_file>
  ↪<antibody_control_csv> <output_10x.h5>
  scCloud merge_rna_adt -h
```

- Arguments:

input_raw_gene_bc_matrices_h5.h5 Input raw RNA expression matrix in 10x hdf5 format.

input_adt_csv_file Input ADT (antibody tag) count matrix in CSV format.

antibody_control_csv A CSV file containing the IgG control information for each antibody.

output_10x.h5 Merged output file in 10x hdf5 format.

- Options:

-h, --help Print out help information.

- Outputs:

output_10x.h5 Output file in 10x hdf5 format. This file contains two groups — one is for RNAs and the other is for ADTs.

- Examples:

```
scCloud merge_rna_adt example_raw_h5.h5 example_adt.csv antibody_control.csv ↪
  ↪example_merged_raw_10x.h5
```

5.3 API

scrtools can also be used as a python package. Import scrtools by:

```
import scrtools
```

5.3.1 Tools:

Aggregate channel-specific count matrices

<code>tools.aggregate_10x_matrices(csv_file, ...)</code>	Aggregate channel-specific 10x count matrices into one big count matrix.
--	--

`tools.aggregate_10x_matrices`

`tools.aggregate_10x_matrices(csv_file, restrictions, attributes, output_file, google_cloud=False, select_singlets=False, ngene=None, is_dropseq=None)`

Aggregate channel-specific 10x count matrices into one big count matrix.

This function takes as input a `csv_file`, which contains at least 2 columns — Sample, sample name; Location, folder that contains the count matrices (e.g. `filtered_gene_bc_matrices_h5.h5`). It outputs a 10x-formatted HDF5 file containing on merged matrix per genome.

Parameters

- **csv_file** (*str*) – The CSV file containing information about each 10x channel.
- **restrictions** (*list[str]*) – A list of restrictions used to select channels, each restriction takes the format of `name:value,...,value` or `name:~value,...,value`, where `~` refers to not.
- **attributes** (*list[str]*) – A list of attributes need to be incorporated into the output count matrix.
- **output_file** (*str*) – The output count matrix file name.
- **google_cloud** (*bool*, optional (default: `False`)) – If the channel-specific count matrices are stored in a google bucket.
- **select_singlets** (*bool*, optional (default: `False`)) – If we have demultiplexed data, turning on this option will make scCloud only include barcodes that are predicted as singlets.
- **ngene** (*int*, optional (default: `None`)) – The minimum number of expressed genes to keep one barcode.
- **is_dropseq** (*str*, optional (default: `None`)) – If `None`, assume data are generated from 10 assays; otherwise, assume data are generated using Drop-Seq with reference genome `<is_dropseq>`.

Returns

Return type `None`

Examples

```
>>> tools.aggregate_matrix('example.csv', ['Source:pbmc', 'Donor:1'], ['Source',
↪ 'Platform', 'Donor'], 'example_10x.h5')
```

Preprocess

<code>tools.read_input(input_file[, genome, ...])</code>	Load data into memory.
<code>tools.update_var_names(data)</code>	
<code>tools.filter_data(data[, mito_prefix, ...])</code>	

Continued on next page

Table 2 – continued from previous page

<code>tools.log_norm(data, norm_count)</code>	Normalization and then take log
<code>tools.run_pca(data[, standardize, ...])</code>	
<code>tools.run_rpca(data[, scale, max_value, ...])</code>	smooth outliers, then no center/scale data
<code>tools.get_anndata_for_subclustering(data, ...)</code>	

tools.read_input

`tools.read_input(input_file, genome=None, return_a_dict=False, demux_ngene=None, mode='r+')`
Load data into memory.

This function is used to load input data into memory. Inputs can be in .h5, .h5ad, .dge.txt.gz or .csv format.

Parameters

- **input_file** (*str*) – Input file name.
- **genome** (*str*, .h5, optional (default: None)) – A string contains comma-separated genome names. scCloud will read all groups associated with genome names in the list from the hdf5 file. If genome is None, all groups will be considered.
- **return_a_dict** (*boolean*, .h5, optional (default: False)) – If input file contains multiple genome groups, if concatenate them into one h5ad object or return a dictionary of genome-h5ad pairs. If this option is on, return a dict.
- **demux_ngene** (*int*, .h5, optional (default: None)) – Minimum number of genes to keep a barcode for demultiplexing.
- **mode** (*str*, .h5ad, optional (default: r+)) – If input is in h5ad format, the backed mode for loading the data. mode could be 'a', 'r', 'r+'. 'a' refers to load all into memory.

Returns An *anndata* object or a dictionary of *anndata* objects containing the count matrices.

Return type *anndata* object or a dictionary of *anndata* objects

Examples

```
>>> adata = tools.read_input('example_10x.h5', genome = 'mm10')
>>> adata = tools.read_input('example.h5ad', mode = 'r+')
>>> adata = tools.read_input('example_ADT.csv')
```

tools.update_var_names

`tools.update_var_names(data)`

tools.filter_data

`tools.filter_data(data, mito_prefix='MT-', filt_xlsx=None, min_genes=500, max_genes=6000, percent_mito=0.1, percent_cells=0.0005)`

tools.log_norm

`tools.log_norm(data, norm_count)`
 Normalization and then take log

tools.run_pca

`tools.run_pca(data, standardize=True, max_value=10, nPC=50, random_state=0)`

tools.run_rpca

`tools.run_rpca(data, scale=False, max_value=10.0, nPC=50, random_state=0)`
 smooth outliers, then no center/scale data

tools.get_anndata_for_subclustering

`tools.get_anndata_for_subclustering(data, subset_selections)`

Batch correction

```
tools.set_group_attribute(data, attribute_string)
tools.estimate_adjustment_matrices(data)
tools.filter_genes_dispersion(data, ...[, ...])
tools.collect_variable_gene_matrix(data, ...)
tools.correct_batch_effects(data)
```

tools.set_group_attribute

`tools.set_group_attribute(data, attribute_string)`

tools.estimate_adjustment_matrices

`tools.estimate_adjustment_matrices(data)`

tools.filter_genes_dispersion

`tools.filter_genes_dispersion(data, consider_batch, min_disp=0.5, max_disp=None, min_mean=0.0125, max_mean=7)`

tools.collect_variable_gene_matrix

`tools.collect_variable_gene_matrix(data, gene_subset)`

tools.correct_batch_effects

`tools.correct_batch_effects` (*data*)

Diffusion map

```
tools.run_diffmap(data, rep_key[, n_jobs, ...])
```

```
tools.run_pseudotime_calculation(data,
```

```
roots)
```

tools.run_diffmap

`tools.run_diffmap` (*data*, *rep_key*, *n_jobs=1*, *n_components=100*, *alpha=0.5*, *K=100*, *random_state=0*, *knn_method='hns'*, *eigen_solver='randomized'*, *M=20*, *efC=200*, *efS=200*, *full_speed=False*)

tools.run_pseudotime_calculation

`tools.run_pseudotime_calculation` (*data*, *roots*)

Cluster algorithms

```
tools.run_louvain(data[, affinity, ...])
```

```
tools.run_hdbscan(data, rep_key[, n_jobs, ...])
```

```
tools.run_kmeans(data, rep_key, n_clusters)
```

```
tools.run_approximated_louvain(data,
```

```
rep_key)
```

tools.run_louvain

`tools.run_louvain` (*data*, *affinity='W_norm'*, *resolution=1.3*, *random_state=0*)

tools.run_hdbscan

`tools.run_hdbscan` (*data*, *rep_key*, *n_jobs=1*, *min_cluster_size=50*, *min_samples=25*)

tools.run_kmeans

`tools.run_kmeans` (*data*, *rep_key*, *n_clusters*, *n_init=10*, *n_jobs=1*, *random_state=0*)

tools.run_approximated_louvain

`tools.run_approximated_louvain` (*data*, *rep_key*, *n_jobs=1*, *resolution=1.3*, *random_state=0*, *n_clusters=30*, *n_init=20*)

Visualization algorithms

```
tools.run_tsne(data, rep_key, n_jobs[, ...])
```

```
tools.run_fitsne(data, rep_key, n_jobs[, ...])
```

Continued on next page

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```
tools.run_umap(data, rep_key[, ...])
```

```
tools.run_force_directed_layout(data,
```

```
...[, ...])
```

tools.run_tsne

```
tools.run_tsne (data, rep_key, n_jobs, n_components=2, perplexity=30, early_exaggeration=12, learning_rate=1000, random_state=0, out_basis='tsne')
```

tools.run_fitsne

```
tools.run_fitsne (data, rep_key, n_jobs, n_components=2, perplexity=30, early_exaggeration=12, random_state=0)
```

tools.run_umap

```
tools.run_umap (data, rep_key, n_components=2, n_neighbors=15, min_dist=0.1, spread=1.0, random_state=0)
```

tools.run_force_directed_layout

```
tools.run_force_directed_layout (data, file_name, n_jobs, affinity='W_diffmap', K=50, layout='fa', n_steps=10000, memory=20)
```

Differential expression analysis

```
tools.run_de_analysis(input_file, ...)
```

tools.run_de_analysis

```
tools.run_de_analysis (input_file, output_excel_file, labels, n_jobs, alpha, run_fisher, run_mwu, run_roc)
```

5.3.2 Annotate clusters:

```
annotate_cluster.
```

```
annotate_clusters(data, ...)
```

annotate_cluster.annotate_clusters

```
annotate_cluster.annotate_clusters (data, json_file, thre, fout=<_io.TextIOWrapper  
name='<stdout>' mode='w' encoding='UTF-8'>, ignoreNA=False)
```

5.3.3 Plotting:**Static plots**

<code>plotting.plot_composition(data, cluster, attr)</code>	Generate a composition plot, which shows the percentage of cells from each condition for every cluster.
<code>plotting.plot_scatter(data, basis, attrs[, ...])</code>	
<code>plotting.plot_scatter_groups(data, basis, ...)</code>	
<code>plotting.plot_scatter_genes(data, basis, genes)</code>	
<code>plotting.plot_scatter_gene_groups(data, ...)</code>	
<code>plotting.plot_heatmap(data, cluster, genes)</code>	

plotting.plot_composition

`plotting.plot_composition` (*data*, *cluster*, *attr*, *style*='frequency', *stacked*=True, *logy*=False, *subplot_size*=(6, 4), *left*=0.15, *bottom*=None, *wspace*=0.3, *hspace*=None, *restrictions*=[])

Generate a composition plot, which shows the percentage of cells from each condition for every cluster.

This function is used to generate composition plots, which are bar plots showing the cell compositions (from different conditions) for each cluster. This type of plots is useful to fast assess library quality and batch effects.

Parameters

- **data** (*anndata* object) – Single cell expression data as an *anndata* object.
- **cluster** (*str*) – A string represents cluster labels, e.g. `louvain_labels`.
- **attr** (*str*) – A sample attribute representing the condition, e.g. `Donor`.
- **style** (*str*, optional (default: *frequency*)) – Composition plot style. Can be either *frequency*, *count*, or 'normalized'. Within each cluster, the *frequency* style show the ratio of cells from each condition over all cells in the cluster, the *count* style just shows the number of cells from each condition, the *normalized* style shows the percentage of cells from the condition in this cluster over the total number of cells from the condition for each condition.
- **stacked** (*bool*, optional (default: *True*)) – If stack the bars from each condition.
- **logy** (*bool*, optional (default: *False*)) – If show the y-axis in log10 scale
- **subplot_size** (*tuple*, optional (default: (6, 4))) – The plot size (width, height) in inches.
- **left** (*float*, optional (default: 0.15)) – This parameter sets the figure's left margin as a fraction of subplot's width (`left * subplot_size[0]`).
- **bottom** (*float*, optional (default: 0.15)) – This parameter sets the figure's bottom margin as a fraction of subplot's height (`bottom * subplot_size[1]`),
- **wspace** (*float*, optional (default: 0.2)) – This parameter sets the width between subplots and also the figure's right margin as a fraction of subplot's width (`wspace * subplot_size[0]`).
- **hspace** (*float*, optional (default: 0.15)) – This parameter sets the height between subplots and also the figure's top margin as a fraction of subplot's height (`hspace * subplot_size[1]`).
- **restrictions** (*list[str]*, optional (default: *[]*)) – This parameter is used to select a subset of data to plot.

Returns A *matplotlib.figure.Figure* object containing the composition plot.

Return type *Figure* object

Examples

```
>>> fig = plotting.plot_composition(data, 'louvain_labels', 'Donor', style =
↳ 'normalized', stacked = False)
```

plotting.plot_scatter

```
plotting.plot_scatter(data, basis, attrs, restrictions=[], nrows=None, ncols=None, subplot_size=(4,
4), left=None, bottom=None, wspace=None, hspace=None, alpha=None, leg-
end_fontsize=None, apply_to_all=True)
```

plotting.plot_scatter_groups

```
plotting.plot_scatter_groups(data, basis, cluster, group, restrictions=[], nrows=None,
ncols=None, subplot_size=(4, 4), left=None, bottom=None,
wspace=None, hspace=None, alpha=None, legend_fontsize=None,
showall=True)
```

plotting.plot_scatter_genes

```
plotting.plot_scatter_genes(data, basis, genes, nrows=None, ncols=None, subplot_size=(4, 4),
left=None, bottom=None, wspace=0.3, hspace=None, alpha=None,
use_raw=False)
```

plotting.plot_scatter_gene_groups

```
plotting.plot_scatter_gene_groups(data, basis, gene, group, nrows=None, ncols=None, sub-
plot_size=(4, 4), left=None, bottom=None, wspace=0.3,
hspace=None, alpha=None, use_raw=False)
```

plotting.plot_heatmap

```
plotting.plot_heatmap(data, cluster, genes, use_raw=False, showzscore=False, title="", **kwargs)
```

Interactive plots

<code>plotting.scatter(df, output_file)</code>
<code>plotting.scatter_real(df, output_file[, log10])</code>
<code>plotting.scatter3d(df, output_file)</code>
<code>plotting.scatter3d_real(df, output_file[, log10])</code>

plotting.scatter

```
plotting.scatter(df, output_file)
```

plotting.scatter_real

```
plotting.scatter_real(df, output_file, log10=False)
```

plotting.scatter3d

```
plotting.scatter3d(df, output_file)
```

plotting.scatter3d_real

```
plotting.scatter3d_real(df, output_file, log10=False)
```

5.3.4 Miscellaneous:

<code>misc.search_genes(data, gene_list[, measure])</code>	Extract and display gene expressions for each cluster from an <i>anndata</i> object.
<code>misc.search_de_genes(data, gene_list[, ...])</code>	Extract and display differential expression analysis results of markers for each cluster from an <i>anndata</i> object.

misc.search_genes

```
misc.search_genes(data, gene_list, measure='percentage')
```

Extract and display gene expressions for each cluster from an *anndata* object.

This function helps to see marker expressions in clusters via the interactive python environment.

Parameters

- **data** (*anndata* object) – An *anndata* object containing the expression matrix and differential expression results.
- **gene_list** (*list[str]*) – A list of gene symbols.
- **measure** (*str*) – Can be either *percentage* or *mean_log_expression*. *percentage* shows the percentage of cells expressed the genes and *mean_log_expression* shows the mean log expression.

Returns A data frame containing marker expressions in each cluster.

Return type *pandas.DataFrame*

Examples

```
>>> results = misc.search_genes(data, ['CD3E', 'CD4', 'CD8'], measure =  
↪ 'percentage')
```

misc.search_de_genes

```
misc.search_de_genes(data, gene_list, test='fisher', thre=1.5)
```

Extract and display differential expression analysis results of markers for each cluster from an *anndata* object.

This function helps to see if markers are up or down regulated in each cluster via the interactive python environment. ++ indicates up-regulated and fold change \geq threshold, + indicates up-regulated but fold change $<$ threshold, - indicates down-regulated and fold change $\leq 1 / \text{threshold}$, - indicates down-regulated but fold change $> 1 / \text{threshold}$, '?' indicates not differentially expressed.

Parameters

- **data** (*anndata* object) – An *anndata* object containing the expression matrix and differential expression results.
- **gene_list** (*list[str]*) – A list of gene symbols.
- **test** (*str*, optional (default: *fisher*)) – Differential expression test to look at, could be either *t*, *fisher* or *mwu*.
- **thre** (*float*, optional (default: *1.5*)) – Fold change threshold to determine if the marker is a strong DE (++) or weak DE (+ or -).

Returns A data frame containing marker differential expression results for each cluster.

Return type *pandas.DataFrame*

Examples

```
>>> results = misc.search_de_genes(data, ['CD3E', 'CD4', 'CD8'], test = 'fisher',
↳ thre = 2.0)
```

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