

User Guide



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I. What should your data look like?

1. For Differential expression, GSEA, ssGSEA, GO:

- Gene Expression File: .csv or .tsv or .xlsx or .xls file(.csv or .tsv could be more stable). Column name(header) should be sample name and first column should always be gene id. (Fig. 1a). Gene id could be Ensembl ID or EntrezID or Gene Symbol or Ensembl_Gene Symbol.
- Metadata: tab separated .txt file, header should be "sample" and "group" (Case sensitive) (Fig. 1b)

First column should always be gene id.

| | A | В | С | D | Sample name |
|---|--------------------------|---------|---------|---------|-------------|
| 1 | gene_id | sample1 | sample2 | sample3 | |
| 2 | ENSMUSG0000000001_Gnai3 | 5224 | 6084 | 6313 | |
| 3 | ENSMUSG00000000003_Pbsn | 0 | 0 | 0 | |
| 4 | ENSMUSG00000000028_Cdc45 | 1454 | 1766 | 1780 | |
| 5 | ENSMUSG00000000031_H19 | 0 | 0 | 0 | |
| 6 | ENSMUSG00000000037_Scml2 | 0 | 0 | 0 | |

b.

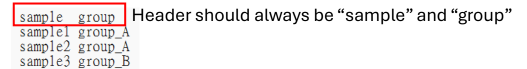


Figure 1. Differential expression, GSEA, ssGSEA, GO data example.

- (a) Expression matrix format (open in Excel).
- (b) Metadata format (open in Windows Notepad).



2. For SingleCell:

- Path to filtered_feature_bc_matrix: Fill in a directory, best for filtered_feature_bc_matrix from 10X platform, but .rds or .h5 or counts matrix and metadata(.csv or .tsv) are also acceptable.
 - For **filtered_feature_bc_matrix**, target folder should contain 3 files: **barcodes.tsv.gz**, **features.tsv.gz** and **matrix.mtx.gz**
 - For .rds or .h5, target folder should contain only 1 file, which will be your ***.rds or ***.h5
 - For counts matrix and metadata, target should contain only 1 counts file named ***_counts.csv (or .tsv) and may contain only 1 metadata file named ***_metadata.csv (or .tsv). Column name of count matrix should be cell barcode and first column should be gene id. (Fig. 2a). The first column of metadata should be the cell barcode, and the others are not specific. (Fig. 2b). If you need to apply an existing ident to counts data, directly change the column name to "assigned.ident". (Fig. 2b)

3. For Visium:

- Path to filtered_feature_bc_matrix : Fill in a directory, only filtered_feature_bc_matrix are accepted
- Path to spatial data: Fill in a directory, target folder should contain following files >>>
 - aligned_fiducials.jpg
 - detected_tissue_image.jpg
 - scalefactors_json.json
 - Thumbs.db
 - tissue_hires_image.png
 - tissue_lowres_image.png
 - tissue_positions_list.csv



| $\boldsymbol{\circ}$ | |
|----------------------|--|
| _ | |

| gene_id | AAACCCAGTCTTAGTG_V1 | AAACCCAGTTGTGGCC_V1 | AAACGAAAGGTGCATG_V1 |
|----------|---------------------|---------------------|---------------------|
| OR4F5 | 0 | 0 | 0 |
| OR4F29 | 0 | 0 | 0 |
| OR4F16 | 0 | 0 | 0 |
| SAMD11 | 0 | 0 | 2 |
| NOC2L | 1 | 0 | 0 |
| KLHL17 | 0 | 0 | 0 |
| PLEKHN1 | 0 | 0 | 0 |
| PERM1 | 0 | 0 | 0 |
| HES4 | 0 | 0 | 1 |
| ISG15 | 0 | 0 | 0 |
| AGRN | 1 | 0 | 1 |
| RNF223 | 0 | 0 | 0 |
| Clorf159 | 0 | 0 | 0 |
| TTLL10 | 0 | 0 | 0 |
| TNFRSF18 | 0 | 0 | 0 |
| | | | |

b.

| cell | orig.ident | nCount_RNA | nFeature_RNA | percent.mt | cluster_id_res0.3 |
|---------------------|------------|------------|--------------|-------------|-------------------|
| AAACCCACACCTTCGT_V1 | V1 | 1550 | 814 | 17.5483871 | 1 |
| AAACCCAGTCTTAGTG_V1 | V1 | 2000 | 943 | 0.45 | 1 |
| AAACCCAGTTGTGGCC_V1 | V1 | 14502 | 4084 | 4.261481175 | 0 |
| AAACGAAAGGTGCATG_V1 | V1 | 5287 | 1973 | 7.603555892 | 0 |
| AAACGAAGTTGCATAC_V1 | V1 | 1364 | 749 | 1.173020528 | 1 |
| AAACGCTTCCTCACTG_V1 | V1 | 1796 | 1038 | 12.08240535 | 2 |
| AAAGGGCAGTATGACA_V1 | V1 | 8489 | 3019 | 8.76428319 | 0 |
| AAAGGGCCAGCAGTGA_V1 | V1 | 1017 | 592 | 3.539823009 | 1 |
| AAAGGTACATTGGGAG_V1 | V1 | 10843 | 3161 | 2.914322604 | 0 |
| AAAGGTAGTAGAGTTA_V1 | V1 | 13434 | 3274 | 1.793955635 | 1 |



Change target column name to "assigned.ident"

| cell | assigned.ident | nCount_RNA | nFeature_RNA | percent.mt | cluster_id_res0.3 |
|---------------------|----------------|------------|--------------|-------------|-------------------|
| AAACCCACACCTTCGT_V1 | V1 | 1550 | 814 | 17.5483871 | 1 |
| AAACCCAGTCTTAGTG_V1 | V1 | 2000 | 943 | 0.45 | 1 |
| AAACCCAGTTGTGGCC_V1 | V1 | 14502 | 4084 | 4.261481175 | 0 |
| AAACGAAAGGTGCATG_V1 | V1 | 5287 | 1973 | 7.603555892 | 0 |
| AAACGAAGTTGCATAC_V1 | V1 | 1364 | 749 | 1.173020528 | 1 |
| AAACGCTTCCTCACTG_V1 | V1 | 1796 | 1038 | 12.08240535 | 2 |
| AAAGGCAGTATGACA_V1 | V1 | 8489 | 3019 | 8.76428319 | 0 |
| AAAGGCCAGCAGTGA_V1 | V1 | 1017 | 592 | 3.539823009 | 1 |
| AAAGGTACATTGGGAG_V1 | V1 | 10843 | 3161 | 2.914322604 | 0 |
| AAAGGTAGTAGAGTTA_V1 | V1 | 13434 | 3274 | 1.793955635 | 1 |

Figure 2. SingleCell counts matrix and metadata example.

- (a) Counts matrix format of SingleCell (open in Excel).
- (b) Metadata format of SingleCell and method of applying an existing ident to counts data (open in Excel).



4. For Multi-sample SingleCell:

 Path to Data List: tab separated .txt file, header should be "sample_name" and "data_path" (Case sensitive) (Fig. 3). "data_path" should be a directory to target folder.

Figure 3. Multi-sample SingleCell data list example (open in Windows Notepad).

5. Others:

- Target Gene File: .csv or .tsv or .xlsx or .xls or tab separated .txt file
 with only one column named "id" (Case sensitive). Only gene
 symbols are accepted (Fig. 4a). Expression barplot will be drown for
 each gene and correlation will be performed on every combination of
 target genes.
- Path to Target Gene Set: .csv or .tsv or .xlsx or .xls or tab separated .txt file, header should be your gene set name or gene name(a single gene can also be accepted, mainly used in SingleCell and Visium). One column will be treated as one gene set and only gene symbols are accepted (Fig. 4b).

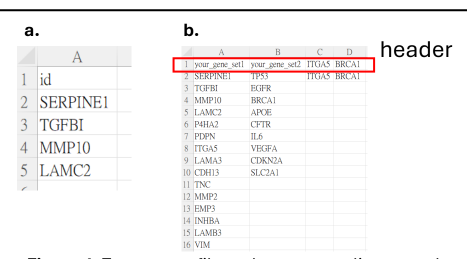


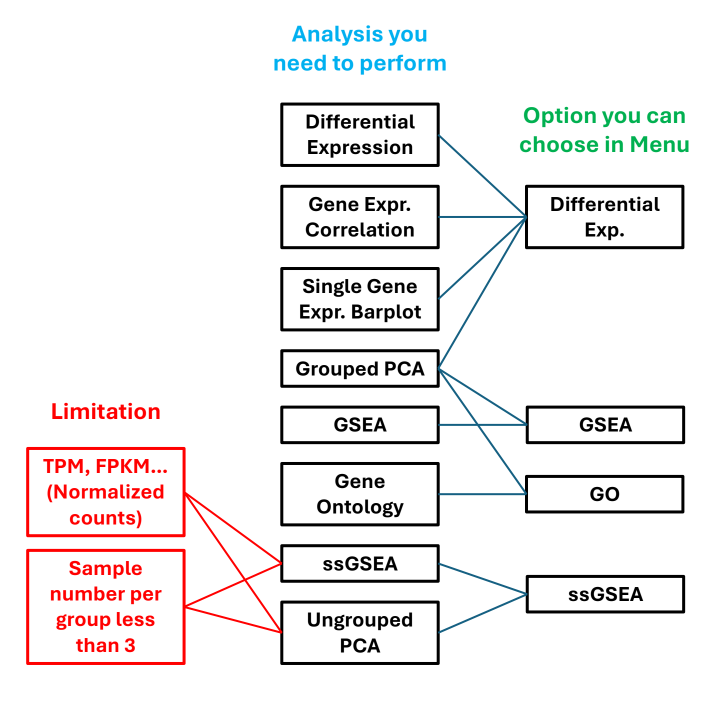
Figure 4. Target gene file and target gene list example.

- (a) Target gene file (open in Excel).
- (b) Target gene list (open in Excel).



II. Workflow

1. Basic Analysis:

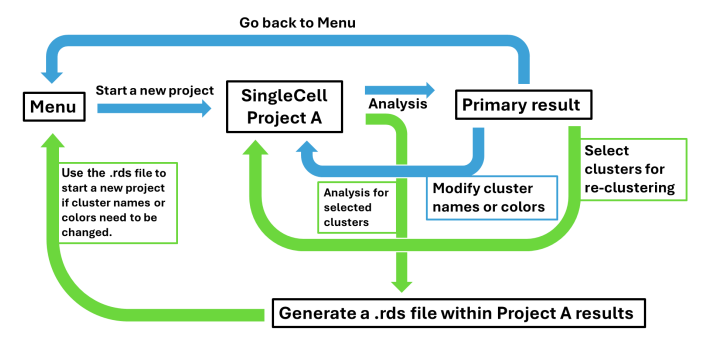


NOTE:

If your data are in TPM, FPKM, CPM format (or other normalized data), or the sample number per group is less than 3 (DESeq2 not accepted), you can only perform ssGSEA in this app.



2. SingleCell and Visium:



NOTE:

After the initial run, you will only be able to view the Primary Results (blue arrow). From there, you have three options:

- (1) Accept Results and Return to Menu.
- (2) Modify Colors or Cluster Names (for all clusters).
- (3) Select Clusters for Re-clustering.

Important: Options 2 and 3 cannot be performed simultaneously. If you choose to modify colors or cluster names (Option 2), all clusters must be selected. If you select specific clusters for re-clustering (Option 3), you cannot change their colors or names.

If you need to modify the color or cluster name for selected clusters after reclustering, return to the Menu and start a new project using the .rds file generated from previous project, which will be located in the folder named "Recluster for Selected Clusters".