Import single cell data

- Import single cell data for different assay types and formats
- · Import single cell data from count matrix text file(s) using Full count matrix as the data format

Import single cell data for different assay types and formats

Select **Single cell**, choose the **assay** type (scRNA-Seq, Spatial transcriptomics, scATAC-Seq, V(D)J, Flow/Mass cytometry), and select the data **format** (Figure 1). Use the **Next** button to proceed with import.

Single cell Bulk Other		
scRNA-Seq Spatial transcriptomics scATAC-Seq V(D)J Flow/Mass Cytometry		
Select the format		
 Import scRNA count feature-barcode-mtx This sparse matrix output is common for 10x Genomics. Fluent Biosciences and Parse Biosciences. Each sample has 3 files (two.csv with one.mtx or two.tsv with one.mtx for each sample). 	10x Genomics Cell Ranger counts h5 This compressed binary format is preferred for 10x Genomics Cell Ranger output. There is 1 filtered .h5 file per sample and multiple files can be selected	 Full count matrix This rectangular cell-by-feature count matrix is common for BD Rhapsody. There is one file for one or more samples (bit, csv, tsv, bt.gz, csv.gz, tsv.gz)
Seurat Rds This R object is for data processed by Seurat (rds)	h5ad This AnnData object in the h5ad file format is for data processed by Scanpy	fastq fastq fastq format is used for unaligned reads. Acceptable file types are fastq. fastq.gz, fastq.bz2, fq. fq.gz, fq.bz2

Figure 3. Choose import single cell data option

Import single cell data from count matrix text file(s) using Full count matrix as the data format

Partek Flow supports single cell data analysis in count matrix text format using the Full count matrix data format (Figure 2). Each matrix text file is assumed to represent on sample, each value in the matrix represents expression value of a feature (e.g. a gene, or a transcript) in a cell. The expression value can be raw count, or normalized count. The requirement of the format of each text file should be the same as count matrix data.

Specify text file location, only one text file (in other words one sample) can be imported at once, preview of the file will be displayed, configuration of the file format is the same as Import count matrix data. In addition, you need to specify the details about this file.



Figure 4. Choose Full count matrix as the data format

Click Finish, the sample will be imported, on the data tab, number of cells in the sample will be displayed.

To import multiple samples, repeat the above steps by clicking **Import data** on the *Metadata* tab or within the task menu (toolbox) on the *Analyses* tab. Make the same previous selections using the cascading menu.

Additional Assistance

If you need additional assistance, please visit our support page to submit a help ticket or find phone numbers for regional support.



Copyright © 2018 by Partek Incorporated. All Rights Reserved. Reproduction of this material without express written consent from Partek Incorporated is strictly prohibited.