## **Importing 10x Genomics Matrix Files**

- Importing single cell data
  - Importing matrices into Partek Flow (this Market Exchange Format is popular for public repositories)
  - Importing matrices in h5 format (this Hierarchical Data Format is recommended for multiple samples)
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### Importing single cell data

Partek Flow supports the import of filtered gene-barcode matrices generated by 10x Genomics' Cell Ranger pipeline.

Below is a video summarizing the import of these files:

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# Importing matrices into Partek Flow (this Market Exchange Format is popular for public repositories)

To import the matrices into Partek Flow, create a new project and click **Add data** then select **Import scRNA count feature-barcode-mtx** under **Single cell > scRNA-Seq**.

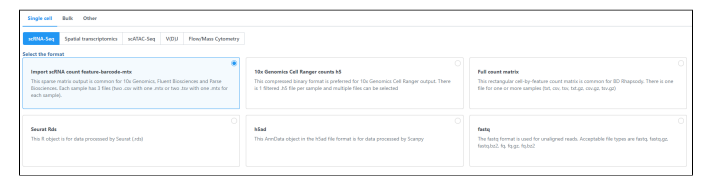


Figure 10. Importing single cell data

Samples can be added using the **Add sample** button. Each sample should be given a name and three files should be uploaded per sample using the **Brow** se button.

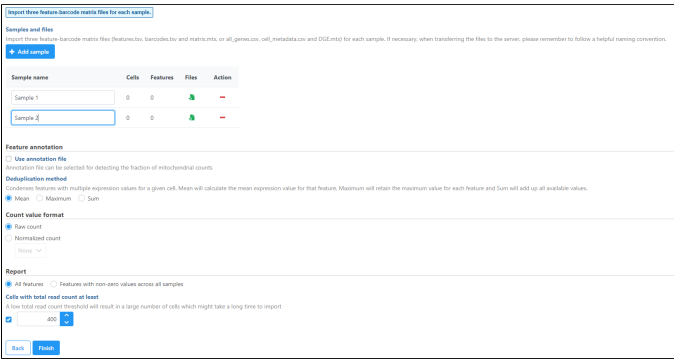


Figure 11. Import three feature-barcode matrix files for each sample

If you have not already, transfer the files to the server to be accessed when you click **Browse**. Follow the directions here to add files to the server. Make sure the files are **decompressed** before they are uploaded to the server.

By default, the Cell Ranger pipeline output will have a folder called filtered\_gene\_bc\_matrices (Figure 3). It is helpful to rename and organize the files prior to transfer using the File browser.

There are folders nested within the matrix folder, typically representing the reference genome it was aligned to. Navigate to the lowest subfolder, this should contain three files:

- barcodes.tsv
- genes.tsv
- matrix.mtx

Select all 3 files for import into Partek Flow

Name	Date modified	Туре	Size
analysis	3/9/2018 9:23 AM	File folder	
filtered_gene_bc_matrices	3/9/2018 9:23 AM	File folder	
raw_geນີe_bc_matrices	3/9/2018 9:23 AM	File folder	
cloupe.cloupe	3/9/2018 9:23 AM	CLOUPE File	36,529 KB
filtered_gene_bc_matrices_h5.h5	3/9/2018 9:23 AM	H5 File	9,736 KB
metrics_summary.csv	3/9/2018 9:23 AM	Microsoft Excel	1 KB
molecule_info.h5	3/9/2018 9:23 AM	H5 File	191,301 KB
possorted_genome_bam.bam	3/9/2018 9:33 AM	BAM File	32,421,469
possorted_genome_bam.bam.bai	3/9/2018 9:23 AM	BAI File	8,508 KB
raw_gene_bc_matrices_h5.h5	3/9/2018 9:23 AM	H5 File	17,052 KB
<ul> <li>web_summary.html</li> </ul>	3/9/2018 9:23 AM	Chrome HTML	3,710 KB

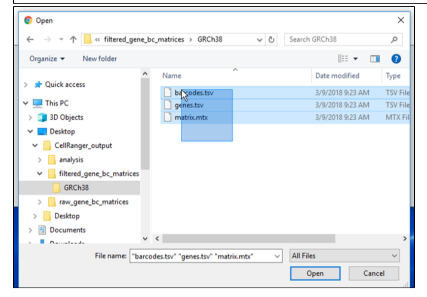


Figure 12. Filtered matrix folder from Cell Ranger pipeline

Specify the annotation file used when running the pipeline for additional information such as mitochondrial counts (Figure 4). Other information can also be specified, such as the count value format. All features can be reported or features with non-zero values across all samples can be reported and the read count threshold can be modified to make the import more efficient.



Figure 13. Configuring matrix metadata

Click Finish when you have completed configuration. This will queue the import task.

## Importing matrices in h5 format (this Hierarchical Data Format is recommended for multiple samples)

The Cell Ranger pipeline can also generate the same filtered gene barcode matrix in h5 format. This gives you the ability to select just one file per matrix and select multiple matrices to import in batch. To import an h5 matrix, select the **Import scRNA full count matrix or h5** option (Figure 1). **Browse** for the files and modify any configuration options. Remember the files need to be transferred to the server.

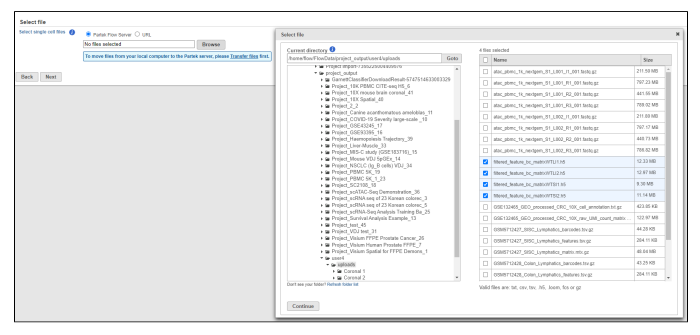


Figure 14. Importing matrix in h5 format

This feature is also useful for importing multiple samples in batch. Simply put all h5 files from your experiment on a single folder, navigate to the folder and select all the matrices you would like to import.

Configure all the relevant sample metadata, including sample name and the annotation that was used to generate the matrices, and click **Finish** when completed. Note that all matrices *must have been generated using the same reference genome and annotation* to be imported into the same project.

### Importing spatial data

#### Importing Xenium Output Bundle

Raw output data generated by the 10x Genomics' Xenium Onboard Analysis pipeline consists of decoded transcript counts and morphology images. The raw output and other standard output files derived from them are compiled into a zipped file called Xenium Output Bundle.

To import the Xenium Output Bundle into Partek Flow, create a new project and click **Add data**, then select **Import 10x Genomics Xenium** under **Single cell > Spatial**, click **Next.** 

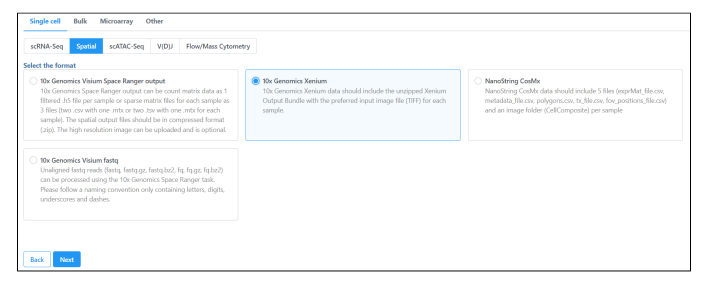


Figure 15. Importing Xenium spatial data

Samples can be added using the **Add sample** button. Each sample should be given a name and a folder containing the required 6 files: cell\_feature\_matrix.h5, cells.csv.gz, cell\_boundaries.csv.gz, nucleus\_boundaries.csv.gz, transcripts.csv.gz, morphology\_focus.ome.tif should be uploaded per sample using the **Browse** button. The required 6 files should be all included in the Xenium Output Bundle folder.

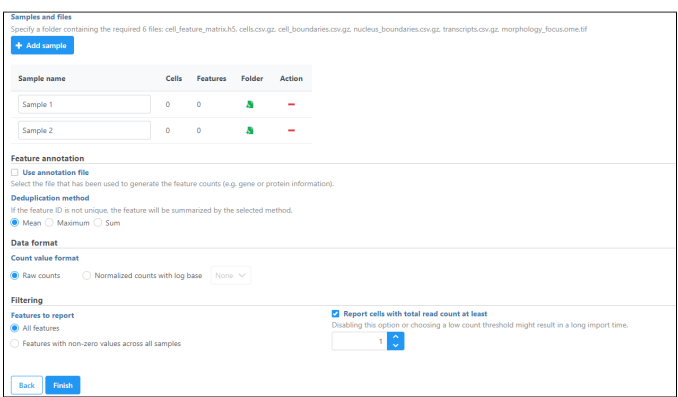


Figure 16. Import Xenium Output Bundle folder for each sample

If you have not already, transfer the files to the server to be accessed when you click **Browse**. Follow the directions here to add files to the server. You will need to **decompress** the Xenium Output Bundle zip file before they are uploaded to the server. After decompression, you can **drag and drop** the entire folder into the Transfer files dialog, all individual files in the folder will be listed in the Transfer files dialog after drag & drop, with no folder structure. The folder structure will be restored after upload is completed.

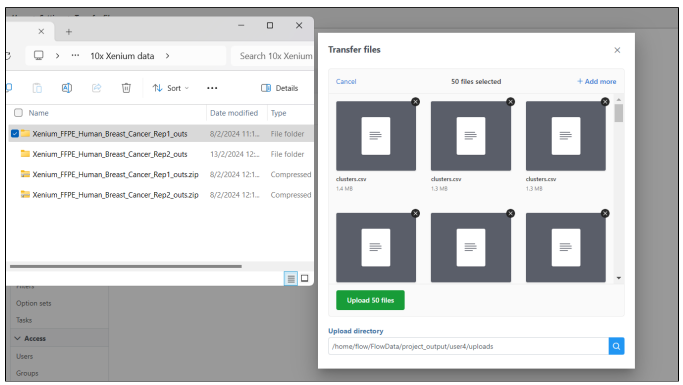


Figure 17. Drag & drop unzipped Xenium Output Bundle folder into Transfer files dialog

Once you have uploaded the folder into the server, you can continue to select the folder for each sample from **Browse**. Once the folder is selected, the *Celli* and *Features* values will auto-populate. You can choose an annotation file that matches what was used to generate the feature count. Then, click **Finish** to start importing the data into your project.

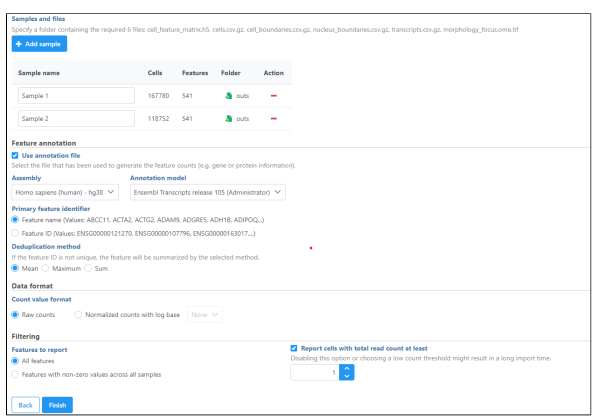


Figure 18. Add Xenium Output Bundle and select annotation

#### Additional Assistance

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

