# Getting started with the tutorial data set

- Creating a new project and importing the tutorial data set
  Filtering cells in single cell RNA-Seq data
- Filtering genes in single cell RNA-Seq data
- Normalizing single cell RNA-Seq data

### Creating a new project and importing the tutorial data set

The tutorial data set is available through Partek® Flow®.

• Click your avatar (Figure 1)

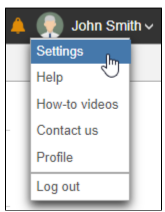


Figure 17. Location of the Settings link on the main page of Partek Flow

Click Settings

On the System information page, the Download tutorial data section includes pre-loaded data sets used by Partek Flow tutorials (Figure 2).

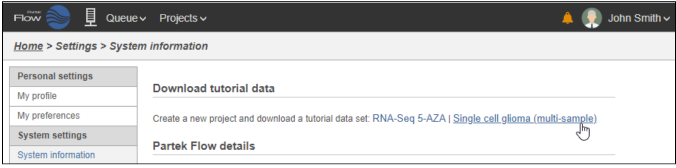


Figure 18. Tutorial data sets available through Partek Flow

#### • Click Single cell glioma (multi-sample)

The tutorial data set will be downloaded onto your Partek Flow server and a new project, Glioma (multi-sample), will be created. You will be directed to the Data tab of the new project. Because this is a tutorial project, there is no need to click on Import data, as the import is handled automatically (Figure 3).

Analyses	Data	Log	Project settings	Notebook	Data Viewer	Attachments				
)ata										
Import			You'll need so	You'll need some samples before you can run an analysis.						
Sample at	tributes				2	-				
Manage			To get started, click the "Import" menu option.							
Assign values from file										

Figure 19. The data tab during tutorial data import

You can wait a few minutes for the download to complete, or check the download progress by selecting **Queue** then **View queued tasks...** to view the *Que ue* (Figure 4).

1	Home > Queued tasks								
	There is 1 task in the queue.								
	Status	Task	Project	User	Submitted	End	Workers	Cancel	
	Download Tutorial Data 🛉 Glioma (multi-sample) Paul Fullerton 20 Mar 2018, 09:28 AM CDT 20 Mar 2018, 09:28 AM CDT 😢								
	<ul> <li>View recent activity</li> <li>T - Waiting for upstream tasks to complete R - Waiting for system resources A - Cannot run with current system configuration</li> <li>Time estimates are being continuously updated and will become more accurate.</li> </ul>								

Figure 20. Viewing the queue

Once the download completes, the sample table will appear in the Data tab, with one row per sample (Figure 5).

analyses Data Log	Proje	ct settings	Noteb	ook	Data Viewer	Attachments	
Data		Sample name		Attributes			
lana a d		Sample i	lame		Subtype	# Cells	
Import	1	MGH36	- 42	Olig	odendroglioma	788	
Sample attributes	2	MGH42		Astrocytoma		682	
Manage	3	MGH45	- 4	Astrocytoma		608	
Assign values	4	MGH53	- 42	Olig	odendroglioma	861	
	5	MGH54	- 4	Olig	odendroglioma	1225	
Assign values from file	6	MGH56	4	Astr	ocytoma	925	
Add system-wide attribute	7	MGH60	- 42	Olig	odendroglioma	430	
	8	MGH64	- 12	Astr	ocytoma	795	

Figure 21. Sample data table listing the name and the number of cells for each sample

The sample table is pre-populated with two sample attributes: # Cells and Subtype. Sample attributes can be added and edited manually by clicking *Managu* in the *Sample attributes* menu on the left. If a new attribute is added, click *Assign values* to assign samples to different groups. Alternatively, you can use the *Assign values from a file* option to assign sample attributes using a tab-delimited text file. For more information about sample attributes, see here.

For this tutorial, we do not need to edit or change any sample attributes.

# Filtering cells in single cell RNA-Seq data

With samples imported and annotated, we can begin analysis.

• Click Analyses to switch to the Analyses tab

For now, the *Analyses* tab has only a single node, *Single cell counts*. As you perform the analysis, additional nodes representing tasks and new data will be created, forming a visual representation of your analysis pipeline.

• Click on the Single cell counts node

A context-sensitive menu will appear on the right-hand side of the pipeline (Figure 9). This menu includes tasks that can be performed on the selected counts data node.

An important step in analyzing single cell RNA-Seq data is to filter out low-quality cells. A few examples of low-quality cells are doublets, cells damaged during cell isolation, or cells with too few counts to be analyzed.

Expand the QA/QC section of the task menu

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• Click on Single cell QA/QC (Figure 6)

Analyses	Data	Log	Project settings	Notebook	Data Viewer	Attachments	🔊 List ger	nerator
							× Single cell counts	
							▼ QA/QC	
Single cell cou	nts						Feature distribution	
							Single cell QA/QC	
							Cell barcode QA/QC	
							► Pre-analysis tools	
							► Filtering	
							Normalization and sca	aling
							► Differential analysis	
							► Exploratory analysis	
							► Pipelines	
							Download data (393 MB)	ц <b>г</b>
							Create project from data not	de
							Publish node to data reposit	tory

Figure 22. Selecting the Single cell QA/QC task from the task menu

A task node, *Single cell QA/QC*, is produced. Initially, the node will be semi-transparent to indicate that it has been queued, but not completed. A progress bar will appear on the *Single cell QA/QC* task node to indicate that the task is running.

- Click the Single cell QA/QC node once it finishes running
- Click Task report on the task menu (Figure 7)

Analyses	Data	Log	Project settings	Notebook	Data Viewer	Attachments	S List generator
							× Single cell QA/QC
	)	Single coll					▼ Task results
Single cell co	unts	Single cell QA/QC					Task report
							Task details
							Task actions

Figure 23. Selecting the task report for any task node opens a report with any tables or charts the task produced

The *Single cell QA/QC* report opens in a new data viewer session. There are interactive violin plots showing the most commonly used quality metrics for each cell from all samples combined (Figure 8). For this data set, there are two relevant plots: the total count per cell and the number of detected genes per cell. Each point on the plots is a cell and the violins illustrate the distribution of values for the y-axis metric. Typically, there is a third plot showing the percentage of mitochondrial counts per cell, but mitochondrial transcripts were not included in the data set by the study authors, so this plot is not informative for this data set.

• Remove the % mitochondrial counts and the extra text box in the bottom right by clicking **Remove plot** in the top right corner of each plot (Figure 8).

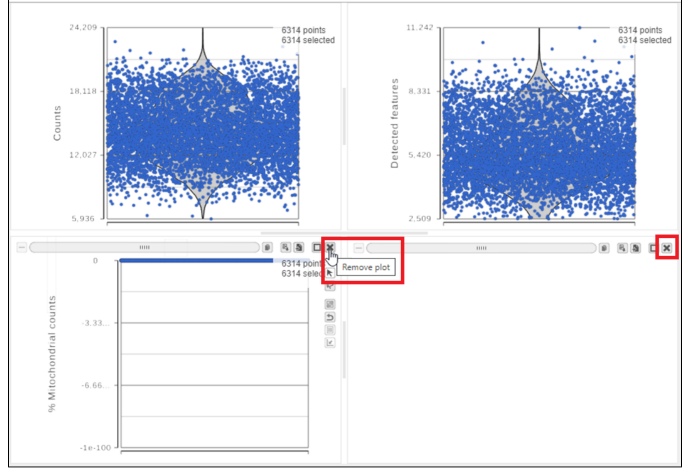


Figure 24. Each cell is shown as a point on the plot. Remove the % mitochondrial counts and empty text box using the X icons

The plots are highly customizable and can be used to explore the quality of cells in different samples.

- Click on Single cell counts in the Get Data icon on the left (Figure 9)
- Click and drag the Sample name attribute onto the Counts plot and drop it onto the X-axis
- Repeat this for the *Detected genes* plot

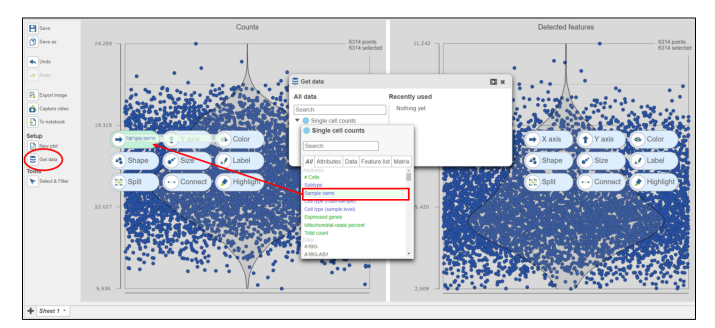


Figure 25. Click and drag the Sample name attribute onto the X-axis for each plot

The cells are now separated into different samples along the x-axis (Figure 10)

- · Hold Control and left-click to select both plots
- Open the Style icon on the left under Configure
- Under *Color*, use the slider to reduce the **Opacity**
- Open the Axis icon on the left
- Adjust the X-rotation on the plots to 90

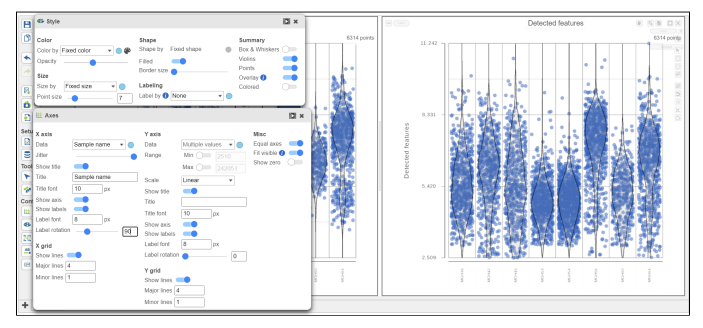


Figure 26. Counts and detected genes plots can be customized to compare cells from different samples

Note how both plots were modified at the same time.

Cells can be selected by setting thresholds using the Select & Filter tool. Here, we will select cells based on the total count

- Open Select & Filter under Tools on the left
- Under *Criteria*, Click **Pin histogram** to see the distribution of counts
- Set the Counts thresholds to 8000 and 20500

Selected cells will be in blue and deselected cells will be dimmed (Figure 11).

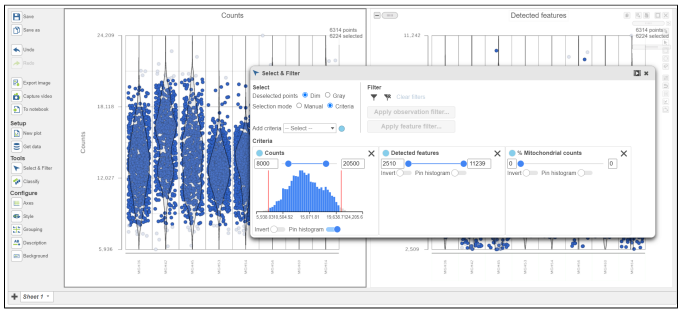


Figure 27. Previewing a filter using the Single cell QA/QC violin plots

Because this data set was already filtered by the study authors to include only high-quality cells, this count filter is sufficient.

<ul> <li>Click Under <i>Filter</i> to include the selected cells</li> <li>Click Apply observation filter</li> <li>Click the Single cell counts data node in the pipeline preview (Figure 12)</li> <li>Click Select</li> </ul>								
Select data to filter								
Single cell courts Single cell Single cell courts Single cell								
Select Cancel								

Figure 28. After the Apply filter button is selected, you will be presented with a preview of your pipeline. You need to select the appropriate data node to apply the filtering to

A new task, *Filter counts*, is added to the *Analyses* tab. This task produces a new *Filter counts* data node (Figure 13).

- Click on the Glioma (multi-sample) project name at the top to go back to the Analyses tab
- Your browser may warn you that any unsaved changes to the data viewer session will be lost. Ignore this message and proceed to the Analyses ta

Analyses Da	ta Log	Project settings	Notebook	Data Viewer	Attachments	🔊 List generator
Single cell counts	Filter counts	Filtered counts				

Figure 29. Applying a cell quality filter

Most tasks can be queued up on data nodes that have not yet been generated, so you can wait for filtering step to complete, or proceed to the next section.

#### Filtering genes in single cell RNA-Seq data

A common task in bulk and single-cell RNA-Seq analysis is to filter the data to include only informative genes. Because there is no gold standard for what makes a gene informative or not, ideal gene filtering criteria depends on your experimental design and research question. Thus, Partek Flow has a wide variety of flexible filtering options.

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- Click the Filter counts node produced by the Filter counts task
- Click Filtering in the task menu
  Click Filter features (Figure 14)

Analyses	Data	Log	Project settings	Notebook	Data Viewer	Attachments	💿 List generator
							× Filtered counts
		Υ_					Task results
Single cell cou	nts	ilter counts	Filtered counts				► QA/QC
	$\Box$	<u>th</u>					Pre-analysis tools
	:	Single cell QA/QC					▼ Filtering
							Filter features
							Filter groups
							Split by attribute
							Downsample cells
							Normalization and scaling
							Differential analysis
							Exploratory analysis
							Pipelines
							Download data (386 MB)
							Create project from data node
							Publish node to data repository

Figure 30. Invoking Filter features

There are four categories of filter available - noise reduction, statistics based, feature metadata, and feature list (Figure 15).

For a detailed explanation of options,	please see the <u>documentation</u> for this task.
Noise reduction filter	
Exclude features where	maximum v <= v 1.0 *
○ Statistics based filter	
Filter features by	Counts      Percentiles
	Keep the top 100.0 🖕 features with highest variance v
○ Feature metadata filter 0	
Filter features by	include V Chromosome V in V Terms V OR
	AND
○ Feature list filter	
Filter criteria	Include O Exclude
Case sensitive	$\sqrt{2}$
Feature identifier 🏼 🧃	Gene symbol (Values: A1BG, A1BG-AS1, A1CF, A2M, A2M-AS1, A2ML1)
	O gene_id (Values: ENSG00000121410, ENSG00000268895, ENSG00000148584,)
	gene_name (Values: A1BG, A1BG-AS1, A1CF, A2M, A2M-AS1, A2ML1)
	O transcript_id (Values: ENST00000600966, ENST00000595302, ENST00000414883,)
Back Finish	

Figure 31. Viewing the filtering options

The noise reduction filter allows you to exclude genes considered background noise based on a variety of criteria. The statistics based filter is useful for focusing on a certain number or percentile of genes based on a variety of metrics, such as variance. The feature list filter allows you to filter your data set to include or exclude particular genes.

We will use a noise reduction filter to exclude genes that are not expressed by any cell in the data set but were included in the matrix file.

- Click the **Noise reduction filter** checkbox
- Set the Noise reduction filter to Exclude features where value <= 0 in 99% of cells using the drop-down menus and text boxes (Figure 16)
- Click **Finish** to apply the filter

For a detailed explanation of options, p	lease see the <u>documentation</u> for this task.
Noise reduction filter	
Exclude features where	value $\checkmark$ $0$ $\checkmark$ in at least $99$ $\checkmark$ $\%$ of the cells
○ Statistics based filter 0	
Filter features by	Counts O Percentiles       Keep the top     100.0 + features with highest     variance
Feature metadata filter ()	
Filter features by	include V Chromosome V in V Terms V OR
○ Feature list filter	AND
Filter criteria	Include O Exclude
Case sensitive	$\checkmark$
Feature identifier 🏼 👔	<ul> <li>Gene symbol (Values: A1BG, A1BG-AS1, A1CF, A2M, A2M-AS1, A2ML1)</li> <li>gene_id (Values: ENSG00000121410, ENSG00000268895, ENSG00000148584,)</li> <li>gene_name (Values: A1BG, A1BG-AS1, A1CF, A2M, A2M-AS1, A2ML1)</li> <li>transcript_id (Values: ENST00000600966, ENST00000595302, ENST00000414883,)</li> </ul>
Back Finish	

Figure 32. Configuring a noise reduction filter to exclude genes not expressed in the data set

This produces a *Filtered counts* data node. This will be the starting point for the next stage of analysis - identifying cell types in the data using the interactive t-SNE plot.

## Normalizing single cell RNA-Seq data

We are omitting normalization in this tutorial because the data has already been normalized.

The tutorial data set is taken from a published study and has already been normalized using TPM (Transcripts per million), which normalizes for the length of feature and total reads, and transformed as log2(TPM/10+1). This normalization and transformation scheme can be performed in Partek Flow, along with other commonly used RNA-Seq data normalization methods.

For more information on normalizing data in Partek Flow, please see the Normalization section of the user manual.

« Single Cell RNA-Seq Analysis (Multiple Samples) Classify cells from multiple samples using t-SNE »

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# Additional Assistance

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