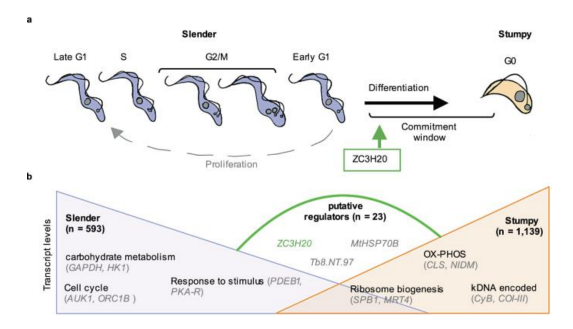
Single Cell RNA-Sequencing (scRNA-seq)

Note: this exercise uses TriTrypDB.org as an example database, but the same functionality is available on all VEuPathDB resources where this type of data is present.

Learning objectives:

- Find all genes with data from scRNA-seq experiments.
- Explore scRNA-seq data on specific gene pages.
- Explore scRNA-seq data using the cellxgene application.



Data used in this exercise is from Briggs, E.M., Rojas, F., McCulloch, R. *et al.* Single-cell transcriptomic analysis of bloodstream *Trypanosoma brucei* reconstructs cell cycle progression and developmental quorum sensing. *Nat Commun* **12**, 5268 (2021). https://doi.org/10.1038/s41467-021-25607-2

Slender markers:

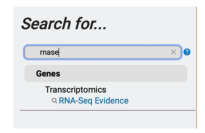
GAPDH: Tb927.6.4280 *PYK1*: Tb927.10.14140

Stumpy markers: *PAD1*: Tb927.7.5930 *PAD2*: Tb927.7.5940 *EP1*: Tb927.10.10260

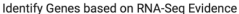
Development regulator:

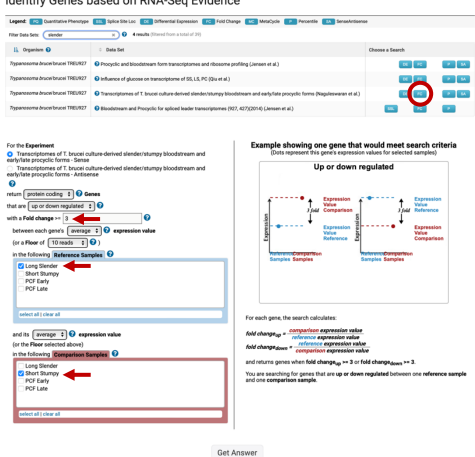
ZC3H20: Tb927.7.2660

- 1. Identify genes that are upregulated in the stumpy form compared to the slender form in different experiments.
 - a. Start by running a search based on RNA-Seq (hint: to find RNA-Seq searches start typing the word RNAseq in the search filter on the left of the home page). Click on RNA-Seq Evidence to view a list of all the available experiments and search types.

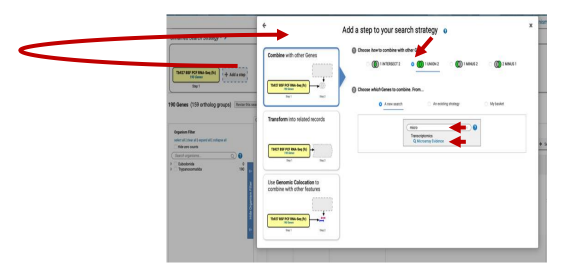


- b. Use the dataset filter to find all experiments that included slender parasites.
- c. Select the fold-change search associated with the data set: *Transcriptomes of T. brucei culture-derived slender/stumpy bloodstream and early/late procyclic forms* (*Naguleswaran et al*). Set up the search parameters to identify genes that are differentially regulated (up or down) by at least 3-fold between the slender and stumpy forms (see screenshot if you need help). Once you are happy with the parameters click on the "Get Answer" button.

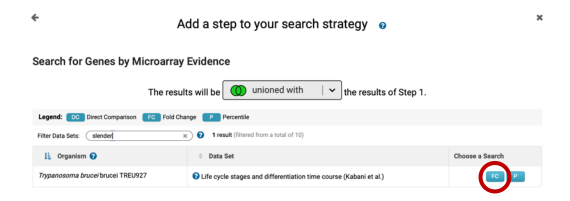


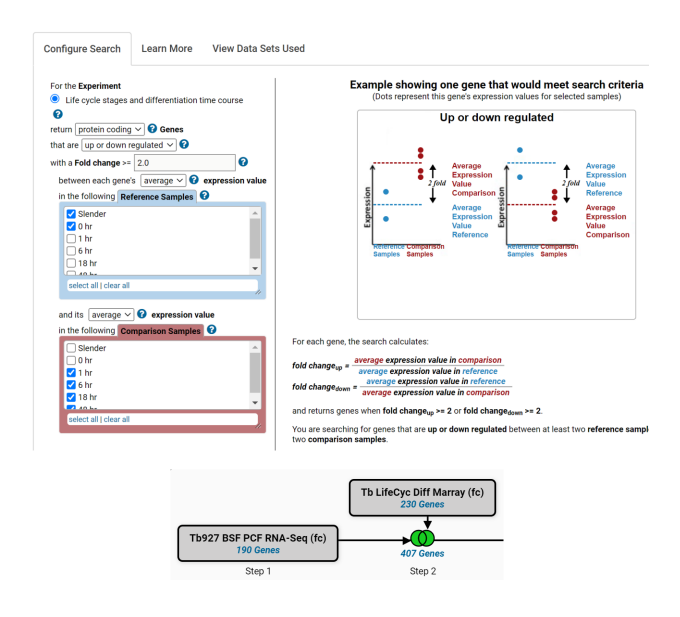


- d. Let's expand the number of genes that might be interesting by combining the results from the above search with results from another experiment that assayed the same stages. To do this follow these steps:
 - Click on the add step button in your search strategy panel.
 - Choose the union operator (why not intersect?).

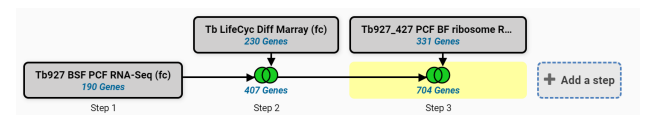


- e. Find the microarray searches by filtering the searches using the key word microarray, then click on the "microarray evidence" link.
- f. Filter the experiments and find one that includes slender stages. Select the fold change search associated with the Kabani et al. experiment. Configure the microarray search to find all genes that are differentially regulated by 2-fold between the slender and Ohr, and hours 1-48. When satisfied with your configuration click on "Run step".



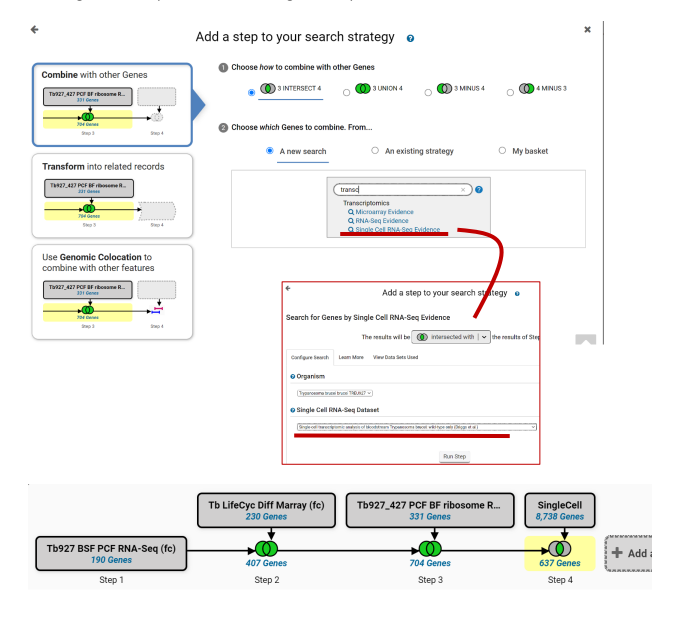


g. Using the same logic as above, add another step and find the RNA-Seq experiment from "Procyclic and bloodstream form transcriptomes and ribosome profiling (Jensen et al.)" and configure the fold change search to find all differentially expressed genes by 2-fold comparing the blood form (cBF mRNA) to the slender form (sIBF mRNA). (don't forget to use the union operator).



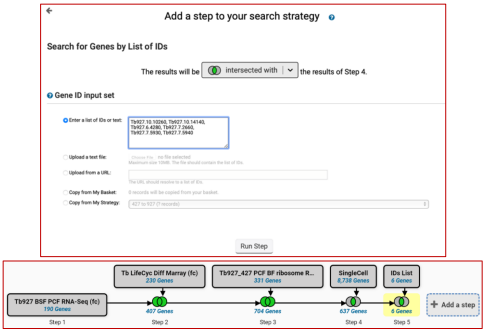
2. Identify genes with expression in single cell RNA-Seq experiments.

a. How many of the genes from #1 above were also identified in the single cell RNA-Seq experiment: "Single-cell transcriptomic analysis of bloodstream Trypanosoma brucei: wild-type only (Briggs et al.)". Using the same logic as above for adding searches, find the single cell RNA-seq experiments and use the intersect operation to find genes from your previous search that have results in the scRNA-Seq experiment. Why are some genes not represented in the single cell experiment?



b. Does this list of genes include any of the markers described in the paper? You can add another step and search using a list of IDs. Copy and paste the following IDs into the

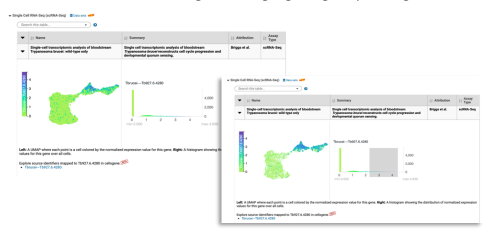
search window: Tb927.10.10260, Tb927.10.14140, Tb927.6.4280, Tb927.7.2660, Tb927.7.5930, Tb927.7.5940



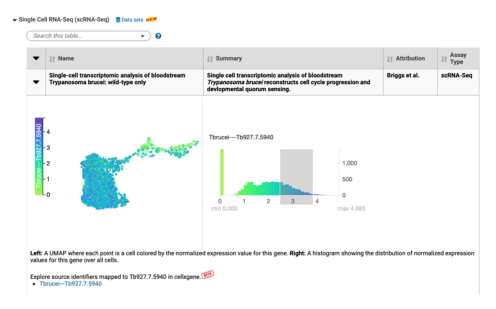
c. Visit the gene page for glyceraldehyde 3-phosphate dehydrogenase (*GAPDH*: Tb927.6.4280) and go to the single cell RNA-Seq section of the page. You can quickly do this by filtering the categories on the left side of the gene page.



d. Expand the first experiment showing wild-type only cells. What does the UMAP plot show? Where are the cells with the highest expression of this gene? You can click and drag in the histogram panel on the right to highlight cells in the left panel. Choose the area between 3 and 4 on the histogram to highlight high expressing cells on the graph.



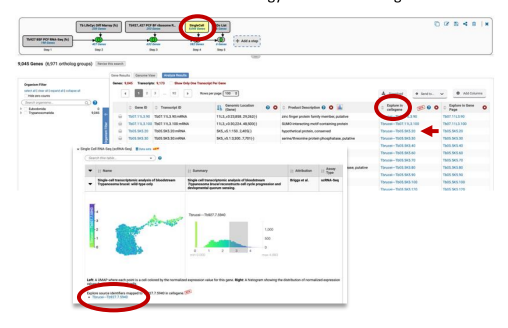
e. Try the same thing with "Protein Associated with Differentiation" (*PAD2*: Tb927.7.5940). Do cells expressing elevated levels of *PAD2* and *GAPDH* coincide on the UMAP or are they in different regions of the plot? Since *GAPDH* is a slender marker and *PAD2* is a stumpy marker, what can you conclude about the cells that coincide with those markers?



3. Explore scRNA-Seq data in the cellxgene application.

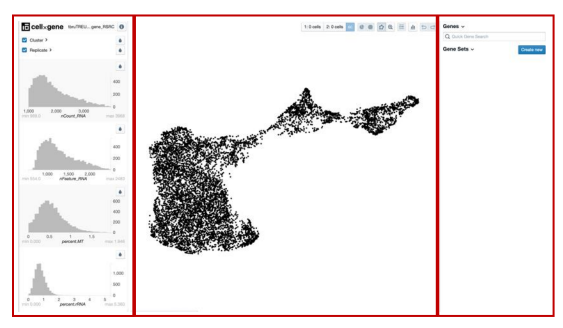
Cellxgene (cell-by-gene) is an open-source data visualization and exploration tool designed to help interrogate high dimensional data. We use cellxgene in VEuPathDB as a supplement to allow investigators to explore scRNA-Seq data.

a. Start with the Briggs et al. wild-type experiment. There are two ways to open cellbygene. From the gene page there is a link below the graphs for each experiment. You can also add a column to the strategy result for the Single Cell search.

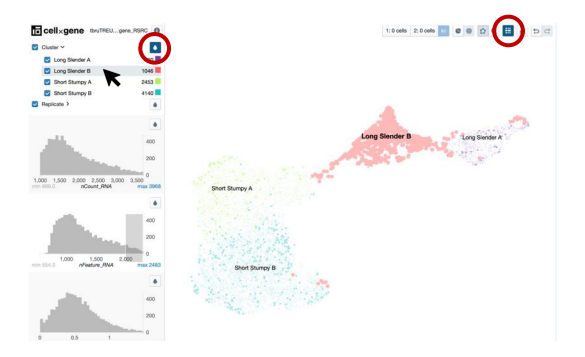


- b. Your initial view will be a UMAP plot of all cells from this experiment. This may be black and white, or may be colored to show expression of a specific gene depending on how you got there.
- c. The left-hand panel includes *metadata* while the right-hand panel includes *gene feature data* where data for any gene measured in the dataset can be explored. The central area is the *cell visualization and exploration* panel.

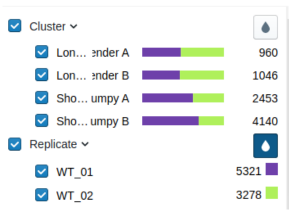




- d. Note that the metadata section includes numerical metadata represented as interactive histograms and categorical metadata such as the cluster assignments or replicates. The exact data shown here will vary by experiment.
- e. The droplet icon can be used to color the cells in the central panel with metadata from the left panel or gene expression data from the right panel. Try this:
 - Expand the "Cluster" metadata category to see the cluster names. Note that these have been annotated by the author of the dataset
 - Use the droplet icon to color the cells by cluster. Do the annotations fit with what you saw when you looked at GAPDH and PAD2 on the gene pages earlier?
 - Hover over the cluster names to bring them into focus in the UMAP.
 - Label the UMAP with the cluster names by clicking on the labels button in the central panel menu.

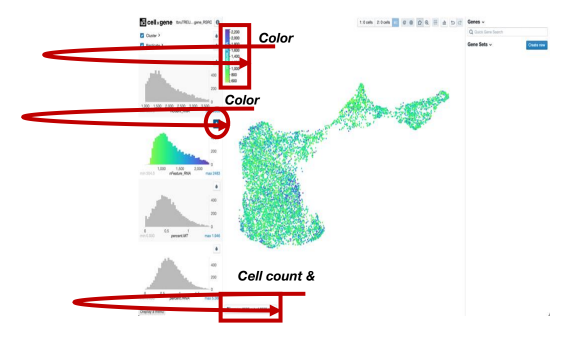


Turn the labeling off again. Expand the "Replicate" metadata category. Use the droplet icon to color based on replicate.
 Mouseover the replicates to see how they are distributed in the UMAP. Notice the bars that appear for the cluster categories showing the proportion of cells from each replicate in each cluster. Do these look like good replicates?

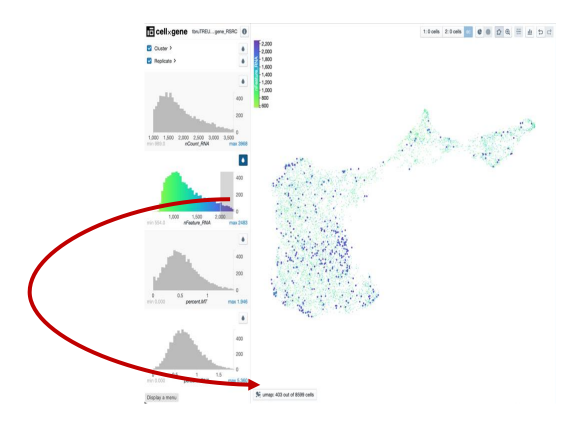


- f. The droplet icon can also be used to color cells based on continuous metadata. Generally, the continuous metadata available is provided for QC purposes. Try this: click on the droplet icon for the nFeature_RNA (number of genes detected in each cell). How many cells are displayed?
- g. In single cell data, it is common to capture a variable number of genes from each cell. How many cells were captured in which 2000 or more genes were observed? To find this, click and drag the histogram area in the left panel to highlight the area representing

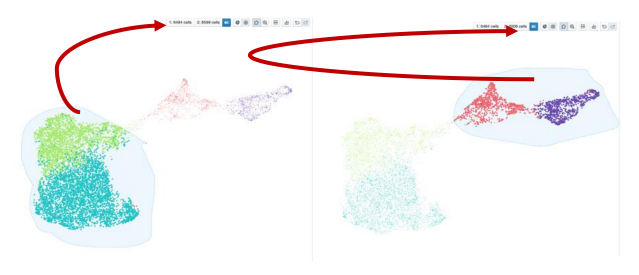
2000 and above. Note: don't worry about being exact here, you are just trying to get an idea of what the data looks like.



h. Do you think you have a higher percentage of stumpy cells with more genes assayed than the slender forms? Can you get the number of cells in each of the stages that met the >/= 2000 genes representation (See step f)? To do this, click and draw around the

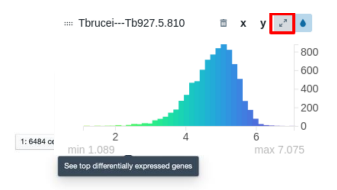


- stumpy cells in the central panel or use the checkboxes next to the cluster labels to deselect the slender cells.
- i. Do the same thing for the slender population. Do you see a difference in the number of cells? Don't forget to take account of the overall number of cells in each population. You can see this in the left panel.
- j. Now let us identify genes that differentiate between the stumpy and slender populations. Follow these steps to do this:
 - Select the stumpy population (both A and B). You can do this by clicking and drawing round them, or by using the check boxes in the left pane.
 - Click on population 1 in the menu bar to save the selection for differential expression.
 - Repeat the same process to select the slender population and save it as

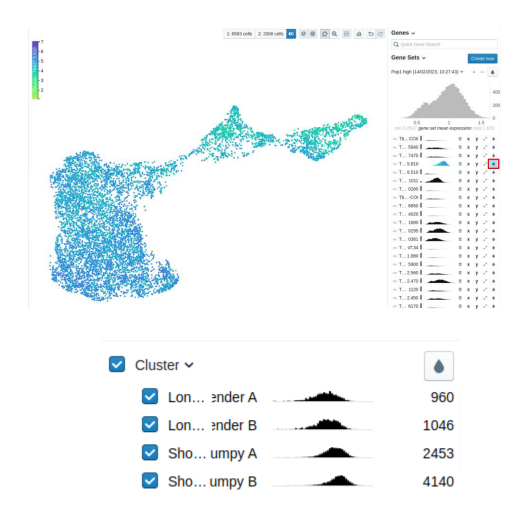


population 2.

- When done with your selections and saving populations, click on the differential expression icon.
- Click on population 1 in the right-hand gene feature panel to reveal the top stumpy genes. Click on the expand icon to view a gene more clearly.



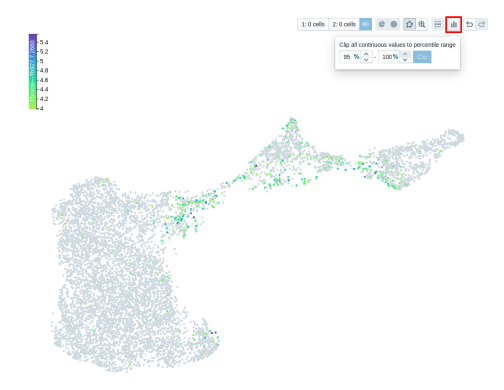
- The histogram in the right panel shows the expression of this gene over all the cells. You can color the UMAP by clicking on the droplet icon next to each gene. The expression of this gene in each cluster can be viewed as histograms in the left panel.



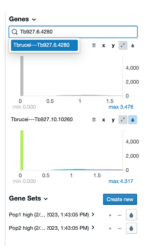
- Copy one of the gene IDs and explore it in TriTrypDB. Can you come up with a rational reason why your selected gene might be important in stumpy development? Note that copying gene IDs from cellxgene is frustrating. If you click on the expand icon for the individual gene, it becomes easier to copy the gene ID.
- Repeat this for the slender forms.
- k. How do the gene sets you identified in your differential expression compare to the marker genes used in the paper? You search for specific genes by pasting the gene ID in

the quick gene search window in the right-hand panel. If the gene is found, you can select it to explore it further. Here is the list of marker genes: Tb927.10.10260, Tb927.10.14140, Tb927.6.4280, Tb927.7.2660, Tb927.7.5930, Tb927.7.5940

- I. The authors identified one gene as a putative regulator of slender to stump transition. This is a zinc-finger protein which has been described as having a role post-transcriptional regulation. Let's look at the expression of this protein.
 - The gene id is Tb927.7.2660. Find this gene using the quick search, and color the UMAP with expression values for this gene.
 - Which cells are expressing this gene at the highest levels?
 Is it easy to see a pattern just by coloring for this gene?
 - We can explore this further in two different ways. First, try clicking and dragging on the expression histogram for this gene to highlight cells where the expression value is > 4.5. You have done this already using the nFeature RNA histogram
 - The second method is to use the clipping tool. Select the clipping tool in the top menu. Leave the upper value at 100%. Change the lower value to 95% and click "Clip". You are now coloring only the cells in the 95th percentile of expression for this gene.

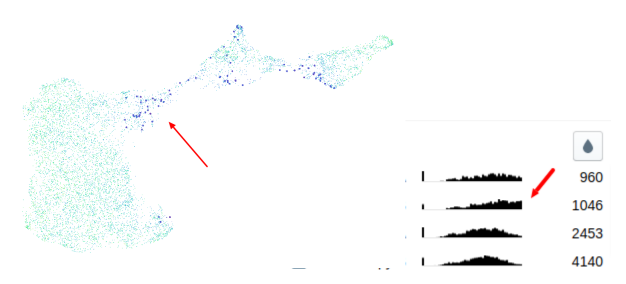


- What happens to the UMAP and the histogram? Is it easier to find the cells with the highest expression levels for this gene now?



- Looking at the expression levels, why do you think the authors chose this transcriptional regulator for further study?
- **4.** The dataset you have just explored is one of two from the same paper. In the first dataset, the authors explored the transition from the replicative slender form of *T. brucei* to the non-replicative, transmissible stumpy form. During this work, they identified a putative regulator, a zinc finger protein ZC3H20: Tb927.7.2660. We looked at the expression of this gene at the end of part 3.

Note that this gene is most highly expressed in the slender B population, and the cells that are transitioning between slender to stumpy. This can be observed in the histograms in the left pane, or by using the gene expression histogram in the right pane to select the cells with the highest expression levels.



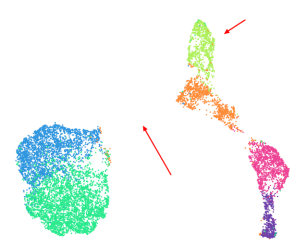
In a subsequent experiment, the group knocked this gene out. The sequenced data from the knockout was integrated with the wild-type cells you've already looked at. This data can be viewed in TriTrypDB using the methods you learnt about above, or it can be directly accessed in cellsgene here:

https://tritrypdb.org/cellxgene/view/tbruTREU927 briggs ZC3H20 KO cellxgene RSRC.h5ad/

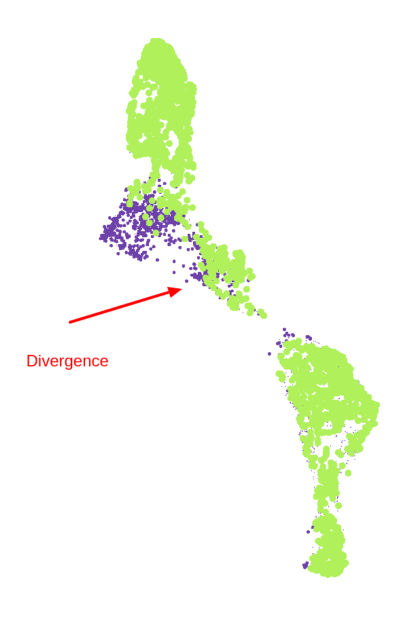
This exercise has fewer instructions - explore this dataset using the tools you learnt about above.

- a. Look at the UMAP for the integrated experiment. How does this differ compared to the wild-type cells?
 - Are there any cell populations you didn't see before?
 - Is there still a clear transition from slender to stumpy?

What do you think is the effect of knocking out this gene?



- b. Color the UMAP for the expression of the gene that was knocked out (Tb927.7.2660). Use the tools you've already learnt about to explore this.
 - Does this look as you expect?
 - Are there any cell populations where this gene is not expressed?
 - Do you think the knockout was successful?
- c. Expand the "Line" metadata category in the left menu. Color the UMAP based on this. Mouseover to highlight the different populations.
 - How are the knockout cells different from the wild-type cells?
 - Expand the "Cluster" metadata category to see the proportions of cells.
- d. It appears that knock-out cells cannot differentiate from slender to stumpy. Instead, they form a novel cluster labeled "Long Slender B.2". Let's see what genes are highly expressed in this cluster:
 - Uncheck all the clusters except "Long Slender B.2". Add this to Population 1
 - Inverse this operation. Add the other clusters to Population 2.
 - Do the differential expression. Can you find any genes in the Pop 1 high list (upregulated in B.2) that look interesting to explore further?
- e. Color the UMAP by the "Line" metadata category again
 - Mouseover the categories and watch what happens in the Long Slender B.1 population
 - It looks like the cells diverge in their differentiation programme here



- The best way to explore this is through trajectory inference. Unfortunately, cellxgene does not offer this. However, we can begin to explore this with differential expression
- Select only the Long Slender B.1 population. Use the check box, then draw around it to remove outlier cells.
- Use the subset button to remove the other cells.
- Now add the ZC3H20_KO cells in this subset to population 1, add the WT cells in this subset to population 2, and do a differential expression.
- What genes in this gene set do you think might be worth exploring further?

