

# Supplementary Materials: cellxgene VIP unleashes full power of interactive visualization, plotting and analysis of scRNA-seq data in the scale of millions of cells

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# 1 Getting started with cellxgene VIP

This is a cellxgene VIP tutorial book written in **Markdown**.

## 1.1 Why use cellxgene VIP?

To meet the growing demands from scientists to effectively extract deep insights from single cell RNA-seq datasets, we developed cellxgene VIP, a frontend interactive visualization plugin to cellxgene framework, which directly interacts with in-memory data to generate a comprehensive set of plots in high resolution, perform advanced analysis, and make data downloadable for further analysis. It makes large scale scRNA-seq data visualization and analysis more accessible and reproducible with the potential to become an ecosystem for the scientific community to contribute even more modules to the Swiss knife of scRNA-seq data exploration tool.

## 1.2 Getting Set up

### 1.2.1 Execute anaconda

```
bash ~/Downloads/Anaconda3-2020.02-Linux-x86_64.sh
```

If anaconda is not installed on server, you can install it following anaconda documentation (<https://docs.anaconda.com/anaconda/install/linux/>) ### Create and enable conda environment

```
# clone repo from cellxgene VIP github
git clone https://github.com/interactivereport/cellxgene_VIP.git
cd cellxgene_VIP

# conda environment
source <path to Anaconda3>/etc/profile.d/conda.sh (Default:
↪ /opt/anaconda3/etc/profile.d/conda.sh)
conda config --set channel_priority flexible
conda env create -n <env name, such as: VIP> -f VIP.yml (system-wide R) or
↪ VIP_conda_R.yml (local R under conda, no root privilege needed)
```

Activate conda environment

```
conda activate <env name, such as: VIP>
```

or

```
source activate <env name>
```

### 1.2.2 Cellxgene installation

Install cellxgene by running config.sh in “cellxgene\_VIP” directory

```
./config.sh
```

### 1.2.3 R dependencies

Install all required R packages on linux:

```
export LIBARROW_MINIMAL=false
# ensure that the right instance of R is used. e.g. system-wide: /bin/R or /usr/bin/R
↪ ; local R under conda: ~/.conda/envs/VIP_conda_R/bin/R
which R

R -q -e 'if(!require(devtools)) install.packages("devtools",repos =
↪ "http://cran.us.r-project.org")'
R -q -e 'if(!require(Cairo)) devtools::install_version("Cairo",version="1.5-12",repos
↪ = "http://cran.us.r-project.org")'
R -q -e 'if(!require(foreign))
↪ devtools::install_version("foreign",version="0.8-76",repos =
↪ "http://cran.us.r-project.org")'
R -q -e 'if(!require(ggpubr)) devtools::install_version("ggpubr",version="0.3.0",repos
↪ = "http://cran.us.r-project.org")'
```

```

R -q -e 'if(!require(ggtrastr))
  ↪ devtools::install_version("ggtrastr",version="0.1.9",repos =
  ↪ "http://cran.us.r-project.org")'
R -q -e 'if(!require(arrow)) devtools::install_version("arrow",version="2.0.0",repos =
  ↪ "http://cran.us.r-project.org")'
R -q -e 'if(!require(Seurat)) devtools::install_version("Seurat",version="3.2.3",repos
  ↪ = "http://cran.us.r-project.org")'
R -q -e 'if(!require(rmarkdown))
  ↪ devtools::install_version("rmarkdown",version="2.5",repos =
  ↪ "http://cran.us.r-project.org")'
R -q -e 'if(!require(tidyverse))
  ↪ devtools::install_version("tidyverse",version="1.3.0",repos =
  ↪ "http://cran.us.r-project.org")'
R -q -e 'if(!require(viridis))
  ↪ devtools::install_version("viridis",version="0.5.1",repos =
  ↪ "http://cran.us.r-project.org")'
R -q -e 'if(!require(BiocManager))
  ↪ devtools::install_version("BiocManager",version="1.30.10",repos =
  ↪ "http://cran.us.r-project.org")'
R -q -e 'if(!require(fgsea)) BiocManager::install("fgsea")'

# These should be already installed as dependencies of above packages
R -q -e 'if(!require(dbplyr)) devtools::install_version("dbplyr",version="1.0.2",repos
  ↪ = "http://cran.us.r-project.org")'
R -q -e 'if(!require(RColorBrewer))
  ↪ devtools::install_version("RColorBrewer",version="1.1-2",repos =
  ↪ "http://cran.us.r-project.org")'
R -q -e 'if(!require(glue)) devtools::install_version("glue",version="1.4.2",repos =
  ↪ "http://cran.us.r-project.org")'
R -q -e 'if(!require(gridExtra))
  ↪ devtools::install_version("gridExtra",version="2.3",repos =
  ↪ "http://cran.us.r-project.org")'
R -q -e 'if(!require(ggrepel))
  ↪ devtools::install_version("ggrepel",version="0.8.2",repos =
  ↪ "http://cran.us.r-project.org")'
R -q -e 'if(!require(MASS)) devtools::install_version("MASS",version="7.3-51.6",repos
  ↪ = "http://cran.us.r-project.org")'
R -q -e 'if(!require(data.table))
  ↪ devtools::install_version("data.table",version="1.13.0",repos =
  ↪ "http://cran.us.r-project.org")'

```

#### 1.2.4 Run cellxgene by h5ad file

You can also run cellxgene by specifying a h5ad file, which stores scRNA-seq data along with a host and a port. Use 'ps' to find used ports to spare. Please see <https://chanzuckerberg.github.io/cellxgene/posts/launch> for details

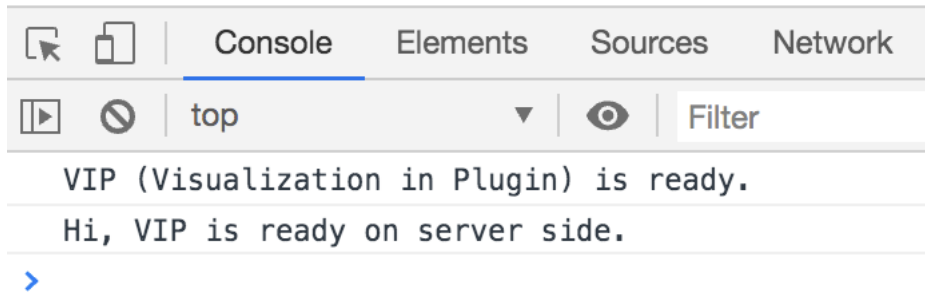
```

ps -ef | grep cellxgene
Rscript -e 'reticulate::py_config()'
# Run the following command if the output of the above command doesn't point to the
↪ Python in your env.
export RETICULATE_PYTHON=`which python`
cellxgene launch --host <xxx> --port <xxx> --disable-annotations --verbose <h5ad file>

```

### 1.2.5 Cellxgene on web browser

chrome is preferred, version 87.0.4280.88 or 87.0.4280.141 is used. Users can access **http(s)://host:port**. Following screenshot is what you should be able to see in console of chrome developer tools.



## 1.3 Authors

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## 2 Web source

### Cellxgene VIP

- **cellxgene VIP source code:** [https://github.com/interactivereport/cellxgene\\_VIP](https://github.com/interactivereport/cellxgene_VIP)
- **cellxgene VIP demo sites:** <https://cellxgenevip-ms.bxgenomics.com> [Schirmer / Rowitch MS, Human brain snRNAseq 46k cells, Nature 2019 Schirmer et al]

### Cellxgene

- **cellxgene:** <https://github.com/chanzuckerberg/cellxgene>
- **cellxgene tutorial:** <https://cellgeni.readthedocs.io/en/latest/visualisations.html>
- **cellxgene features:** <https://chanzuckerberg.github.io/cellxgene/posts/gallery>
- **cellxgene data preparation:** <https://chanzuckerberg.github.io/cellxgene/posts/prepare.html>

### Python packages

- **scanpy:** <https://github.com/theislab/scanpy>
- **scanpy plots:** <https://scanpy-tutorials.readthedocs.io/en/latest/visualizing-marker-genes.html>
- **diffxpy:** <https://github.com/theislab/diffxpy>
- **diffxpy tests:** <https://diffxpy.readthedocs.io/en/latest/tutorials.html#differential-testing>

## Others

- **Benchmarking of interactive data visualization of single-cell RNAseq data** — Batuhan Cakir : [https://www.youtube.com/watch?v=3nH2xi\\_Ni6I](https://www.youtube.com/watch?v=3nH2xi_Ni6I)
- **sceasy**: convertor of scRNA-Seq data formats <https://github.com/cellgeni/sceasy>
- **jupytertext**: <https://jupytertext.readthedocs.io> , Jupyter Notebooks as Markdown Documents, Julia, Python or R Scripts
- **nbconvert**: <https://nbconvert.readthedocs.io> , Convert a Jupyter .ipynb notebook document file into another static format including HTML, LaTeX, PDF, Markdown, and more

## 3 How to use Cellxgene VIP

### 3.1 Graphical user interface of cellxgene and VIP

The main window of cellxgene is divided into three regions, the left panel mainly displays categorical annotations, brief description of the data set and initial graphics setting, specifically embedding and coloring of cells. On the right panel, it hosts continuous variables, such as qc metrics shown in histogram with x, y corresponding to values of a measurement and numbers of cells, respectively. More importantly, cells shown as individual dots are presented in the center panel based on a selected embedding and colored by either categorical annotations or continuous variables, which is indicated by pressed rain drop icon.

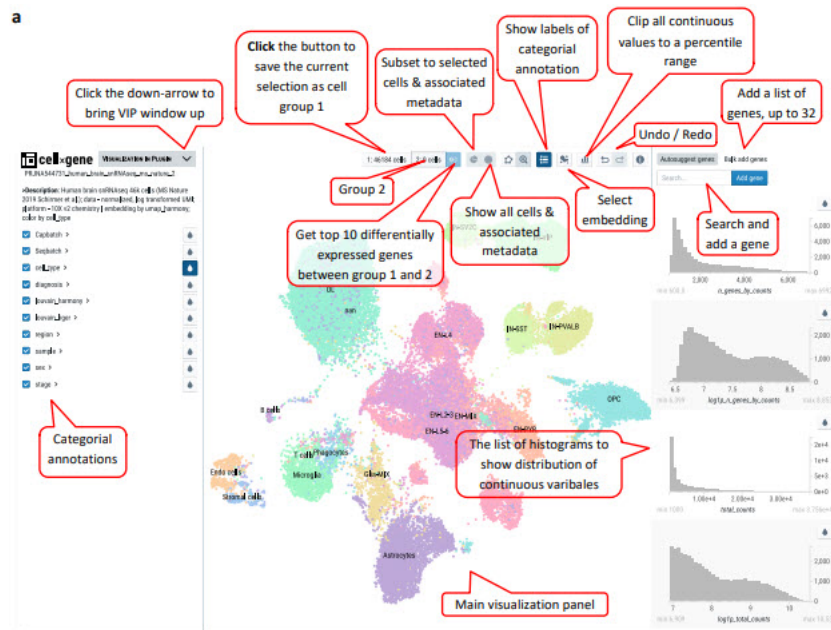


Figure 1: Cellxgene main window, functional icons and minimized VIP bar next to cellxgene logo

### 3.2 Cell selection by categorical annotations

It is an overlap operation when categories from multiple annotations are checked to make the final selection. E.g., if male from sex is also checked besides B cells, it means cells from B cells cluster of male samples are selected. Note: Click “1:” or “2:” button to save cell selection into group 1 or 2

### 3.3 Cell selection by brushing on distribution of continues variables

Note: Histograms of expression values of genes can be brushed as well to get cells expressing certain genes in the range.

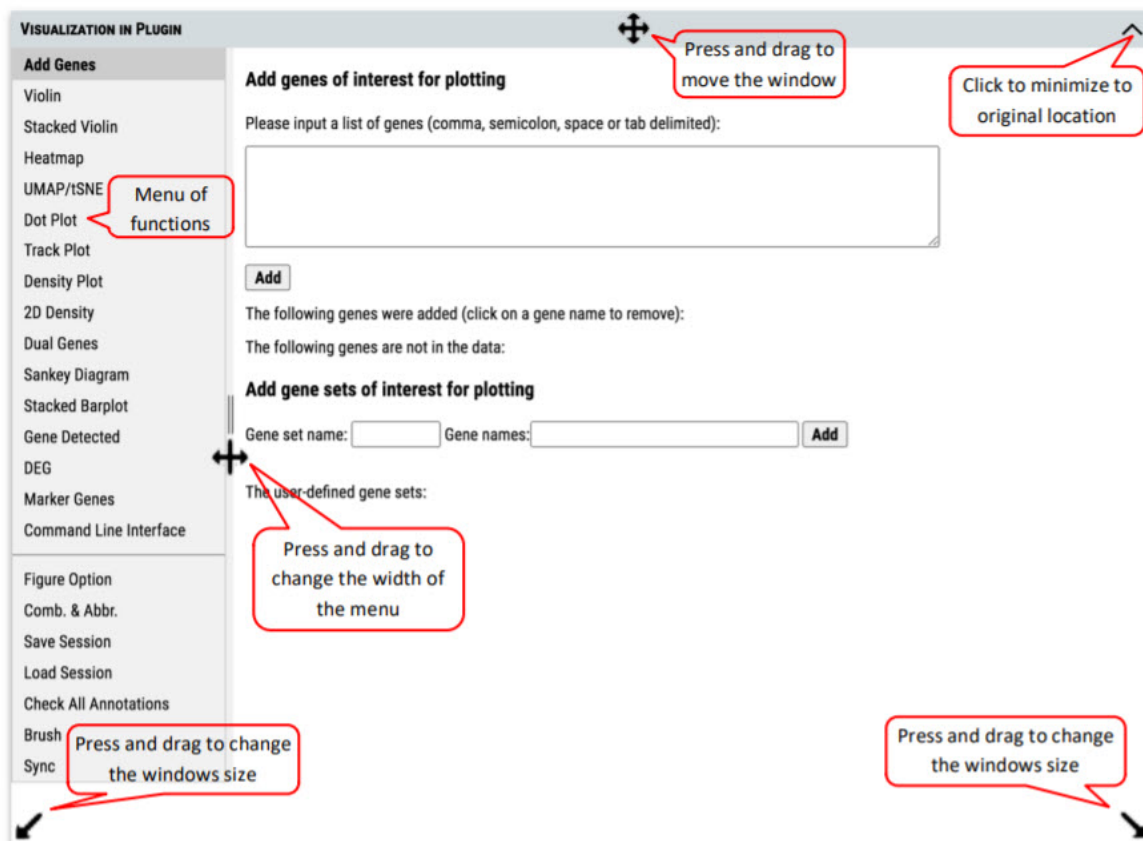


Figure 2: VIP (Visualization in Plugin) window and controls of user interface. The cursor will change to corresponding icon when mouse hovers over control anchors inside the window. In the case of missing title bar after operation, changing the size of outside browser window (not VIP window) will always bring the VIP window back to the original location near the cellxgene logo

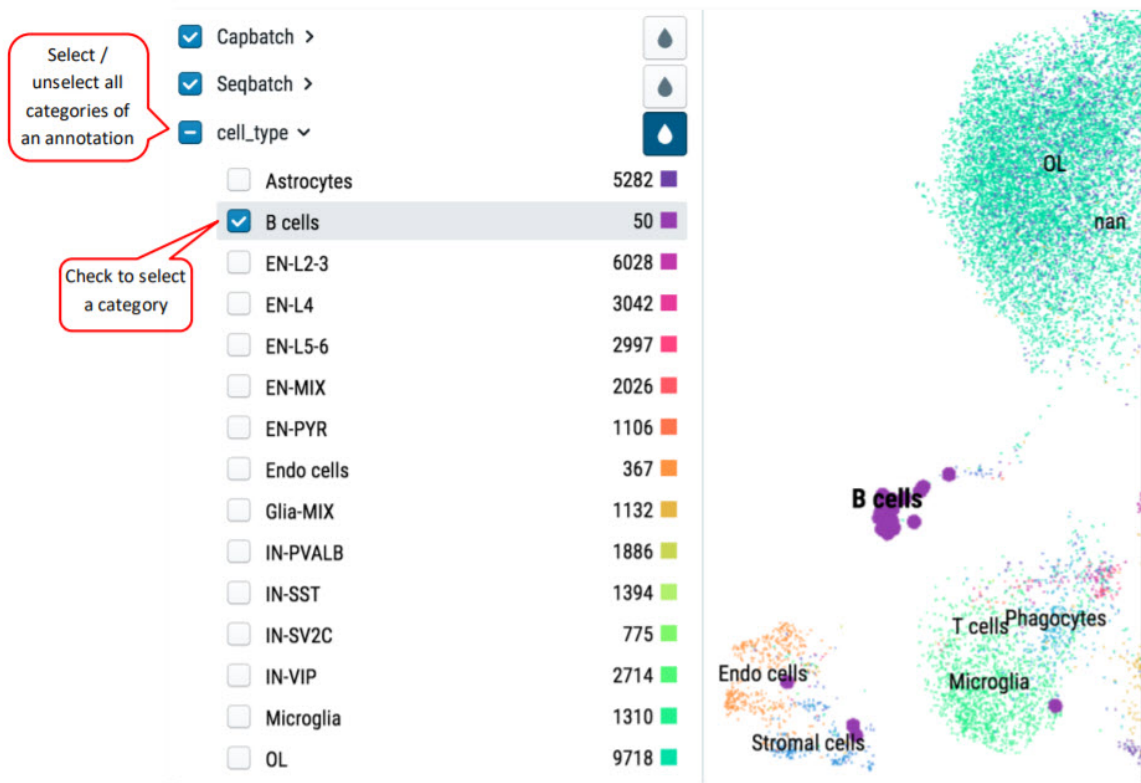


Figure 3: Cell selection by categorical annotation. Selected B cells are shown in bold dots and highlighted in purple color when hovering mouse over the cluster



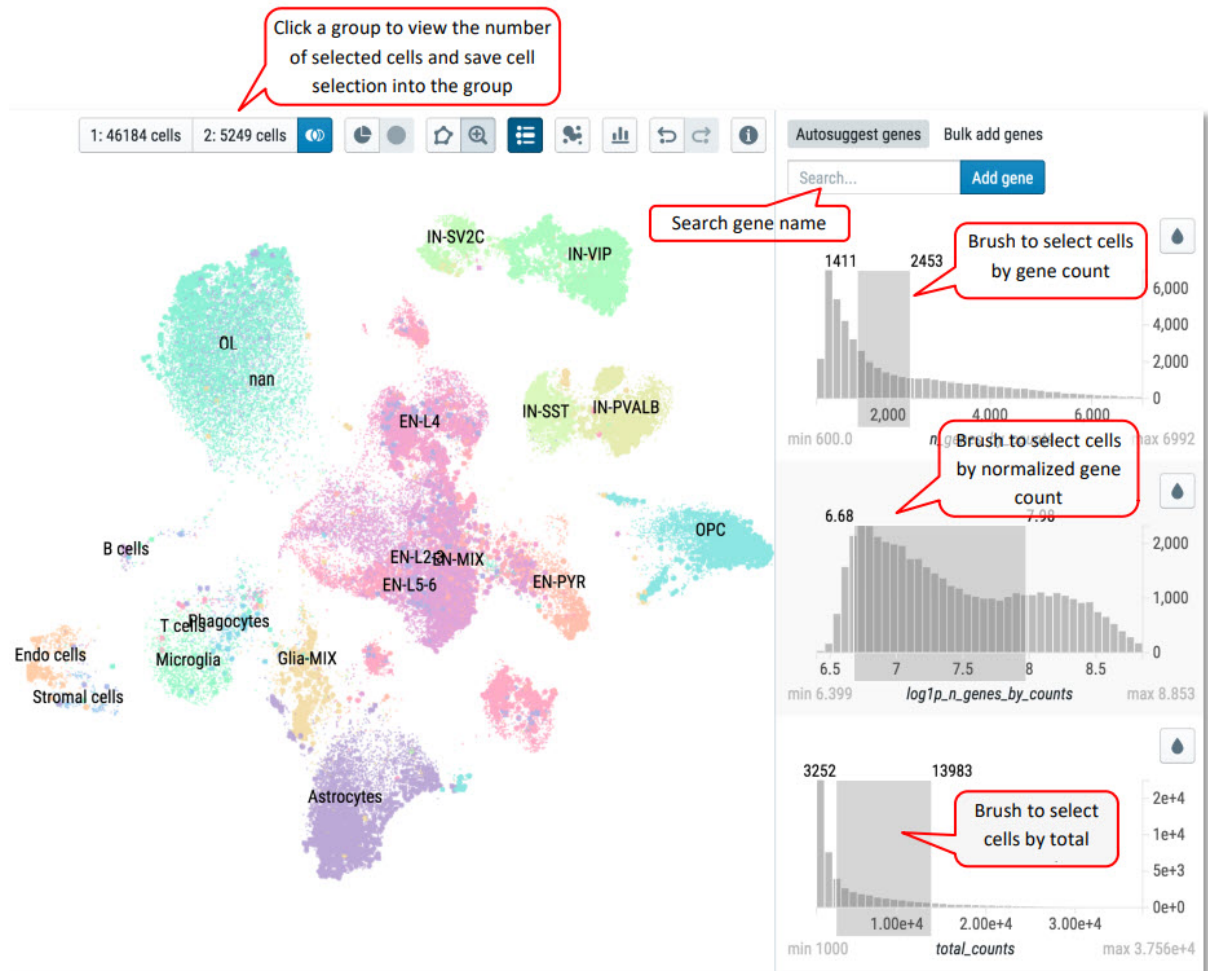


Figure 4: Cell selection by brushing the ranges of continuous variables. Low- and high-end values are shown at top corners of brushing boxes in dark gray

### 3.4 Free hand Lasso selection on dots representing cells

From the cell visualization panel, user can freely select a cluster of cells of interest by using ‘Lasso’ selection tool. The selected cluster of cells can also be added as a group for downstream analysis in cellxgene VIP.

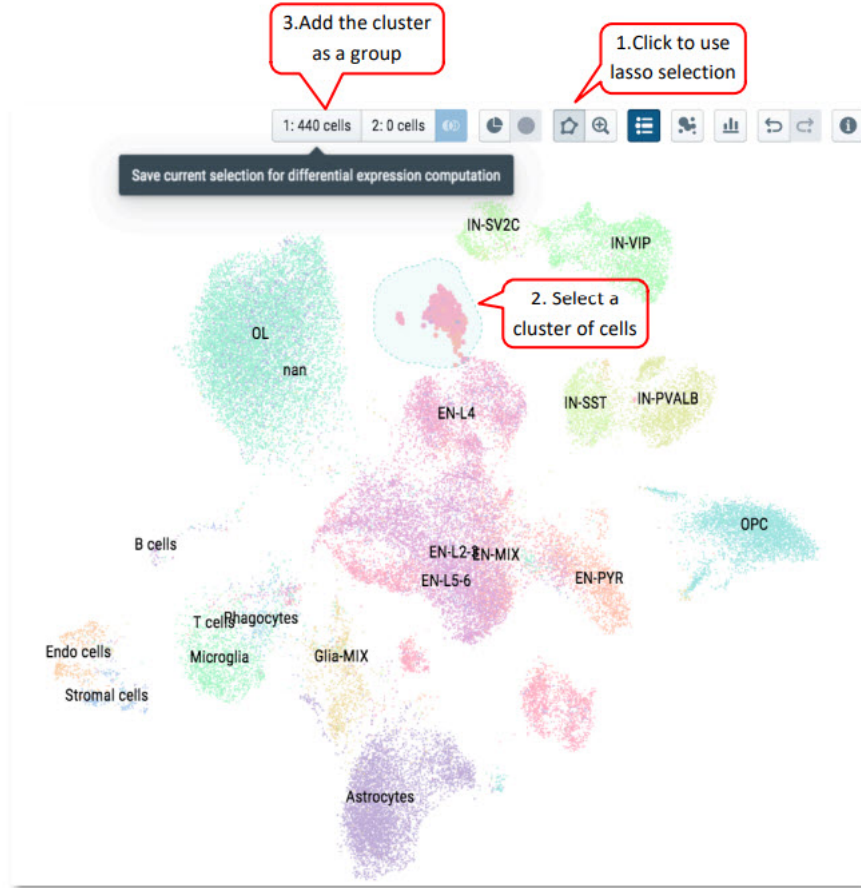


Figure 5: Select cells by using free hand Lasso selection tool and add these cells as a group for further analysis in cellxgene VIP

Note: Please try to draw as close as possible to the starting point in the end to make an enclosed shape to ensure successfully lasso selection.

### 3.5 VIP – Figure Option

User can set parameters for figure plotting that control plotting functions except CLI. ‘split\_show’ branch of Scanpy offers better representation of Stacked Violin and Dot Plot comparing to master branch.

Scaled data have zero mean and unit variance per gene. This was performed by calculating z-scores of the expression data using Scanpy’s scale function. (Scanpy pp.scale function: Scale data to unit variance and zero mean.)

We provide flexibility to allow 1) scale to unit variance or not; 2) Zero centered or not; 3) Capped at max value after scaling.

We recommend using scaled data for plotting/visualization while using non-scaled data for differential gene expression analysis.

Note: Dot plot is one exception in visualization category which uses non-scaled data for meaningful interpretation.

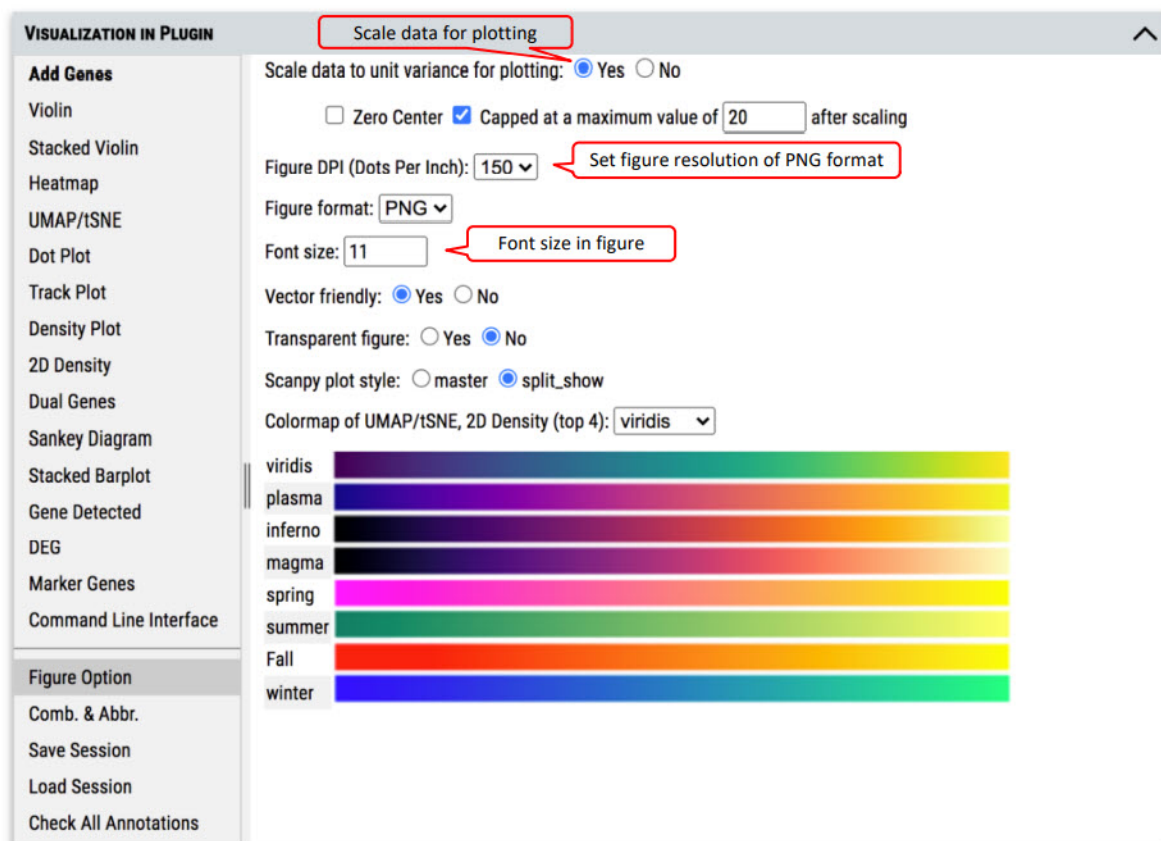


Figure 6: Setting parameters for figure plotting

### 3.6 VIP – Add Genes / Gene Sets

Cellxgene VIP allows user to add any genes or gene sets for extensive exploration and visualization. User can either type a list of gene in the textbox or create sets of genes to be grouped together in plots. Then the genes will be automatically listed for plotting in other functional modules after checking availability in the dataset.

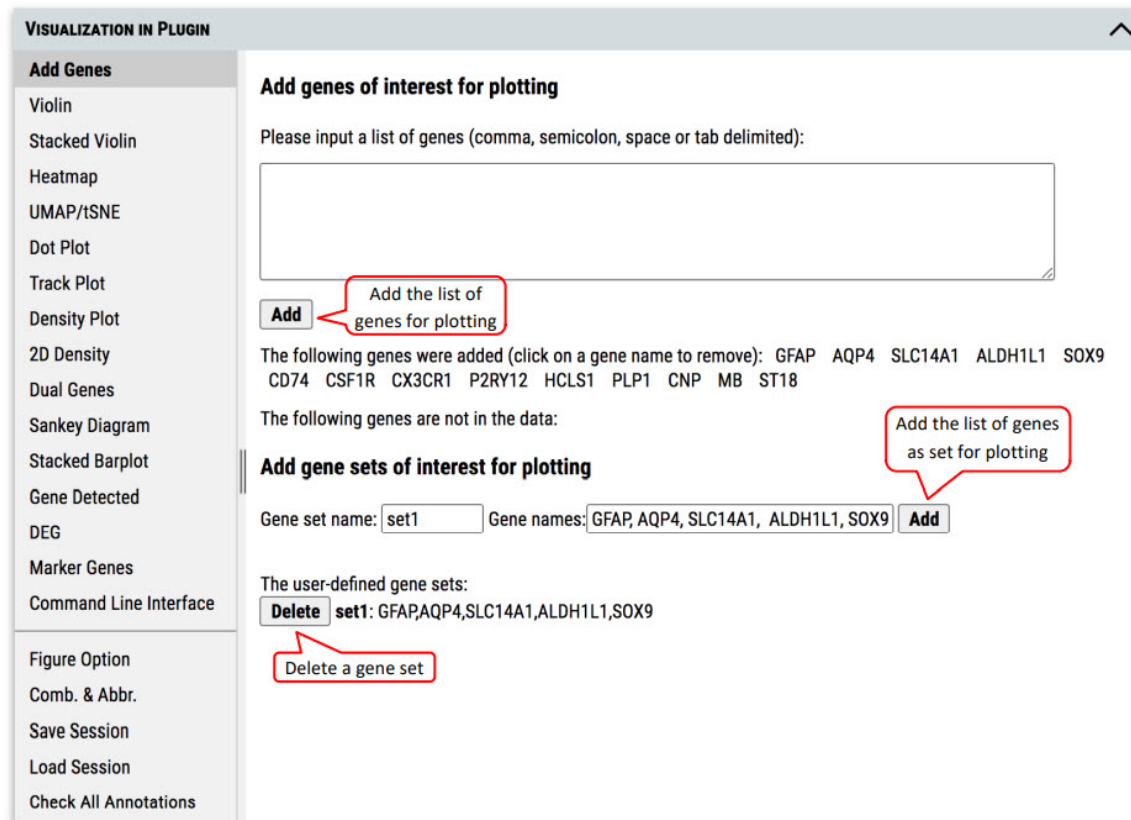


Figure 7: Add gene or gene sets for plotting

Note: The cursor will turn to cross icon while hovering over a gene name, then click to delete the gene.

### 3.7 VIP – Violin Plot

To plot expression of gene among categories of an annotation, e.g., cell type, sex, or batch etc. Step 1. User needs to select the group(s) of cells for plotting. These groups can be created by using selection tools illustrated in tutorial section 2, 3 and/or 4. Initially, all of cells are gathered in 'Group 1' by default. Step 2. Select a gene from the gene list which could be added as shown in section 6. An expression level cutoff can be set to further filter out cells with low level expression of such gene. Step 3. Select the annotation to group cells for plotting. Step 4. Execute plotting, get plotting data (i.e., gene expression), manipulate image (e.g., zoom in/out) or save the image.



Note: Figure resolution and format can be set in “Figure Option” tab as shown in tutorial section 5.

### 3.8 VIP – Stacked Violin

Beyond plotting expression values of a gene, stacked violin allows plotting of multiple genes together.

Note: If collapsing of gene sets is set to ‘Yes’, average gene expression of genes in a set is used for plotting.

### 3.9 VIP – Heatmap

To show or compare the expression level (i.e., expression value or expression Z-score) of multiple genes among the selected group of cells.

### 3.10 VIP – UMAP/tSNE

To plot the embedding of cells in the selected group(s). One of pre-computed and loaded embeddings can be selected.

User can color cells in the embedding plots by multiple annotations (e.g., cell\_type, diagnosis).

Besides coloring cells by annotations, user can color cells based on gene expression level of selected genes in the embedding plots.

### 3.11 VIP – Dot Plot

To show the fraction of cells (annotated by dot size) expressing a gene in each group and the averaged expression level of the gene (annotated by color intensity) in the group.

Note: The number of cells represented by side bar chart are always numbers of cells distributed in each category of certain annotation without filtering. It will give accurate estimate of number of cells in each bubble in the plot. The use of the plot is only meaningful when the counts matrix contains



Figure 8: Stacked Violin plot of multiple genes and/or gene set



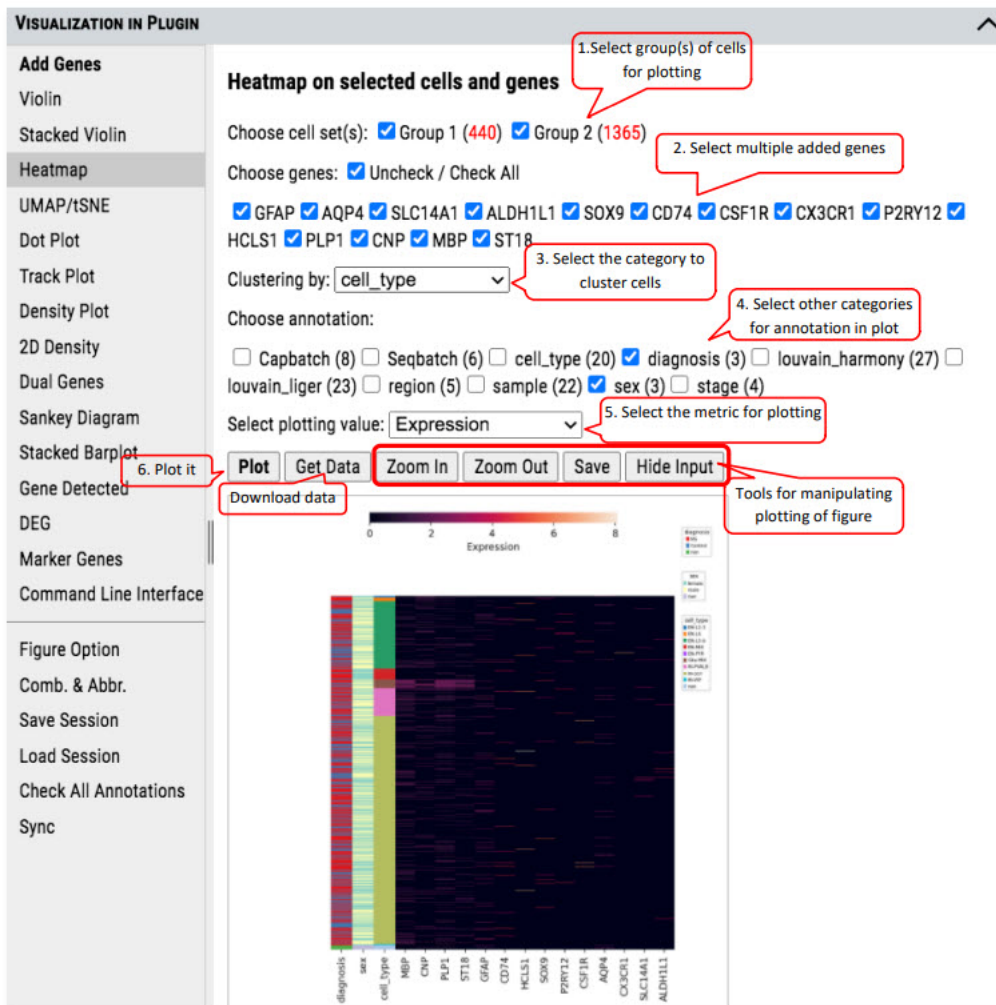


Figure 9: Heatmap of gene expression in cells grouped by annotations.

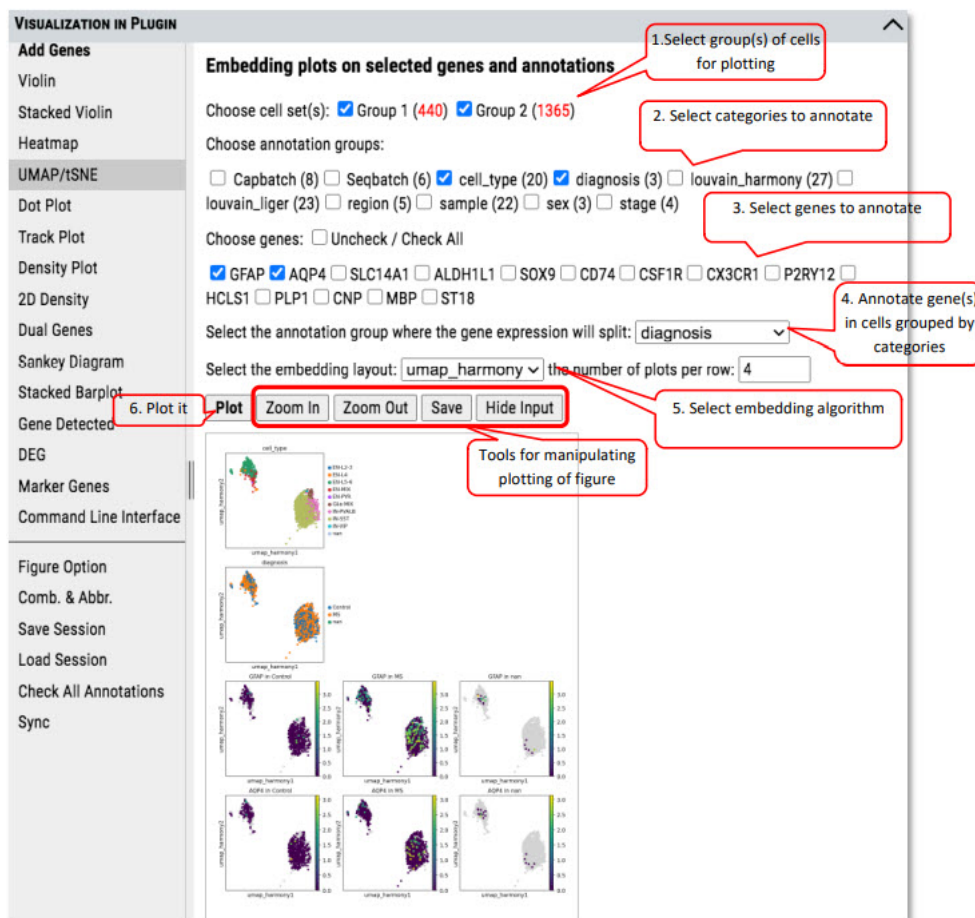


Figure 10: Embedding plotting of expression level of genes or gene set in the cells split by categories of an annotation



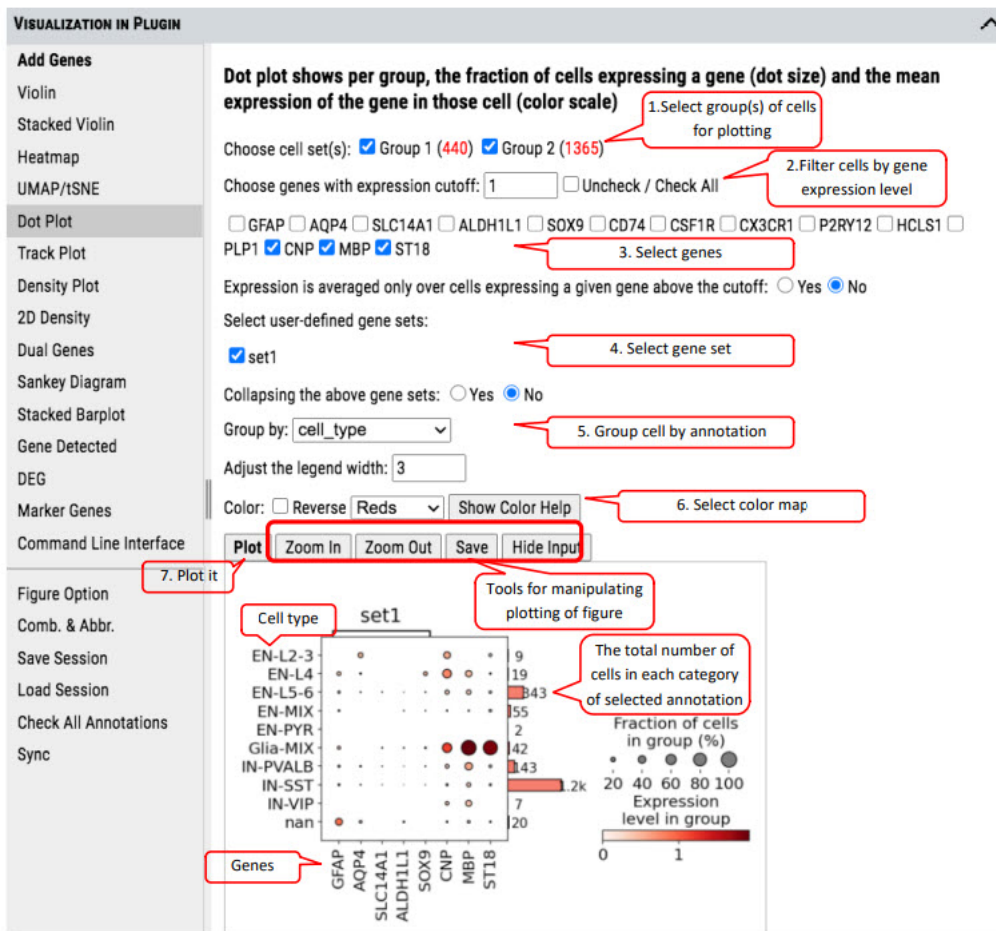


Figure 11: Dot plotting of the fraction of cells expressing genes above a cutoff in each categorie of the selected annotation

zeros representing no gene counts. Its visualization does not work for scaled or corrected matrices in which zero counts had been replaced by other values, see <https://scanpy-tutorials.readthedocs.io/en/multiomics/visualizing-marker-genes.html#Dot-plots>.

### 3.12 VIP – Track Plot

To show the expression of gene(s) of individual cells as vertical lines grouped by the selected annotation on x-axis. Instead of a color scale, the gene expression is represented by height.



Figure 12: Track plotting of expression of genes or gene set in each category of the selected annotation. Gene expression levels are represented by the heights of vertical lines

### 3.13 VIP – Density Plot

To show the density of gene(s) expression in the cells annotated by category in the selected group(s) of cells. A density plot is a representation of the distribution of a numeric variable. It uses a kernel density estimate to show the probability density function of the variable (see more). It is a smoothed version of the histogram and is used in the same concept.

The bandwidth defines how close to a value point the distance between two points must be to influence the estimation of the density at the point. A small bandwidth only considers the closest values, so the estimation is close to the data. A large bandwidth considers more points and gives a smoother estimation.

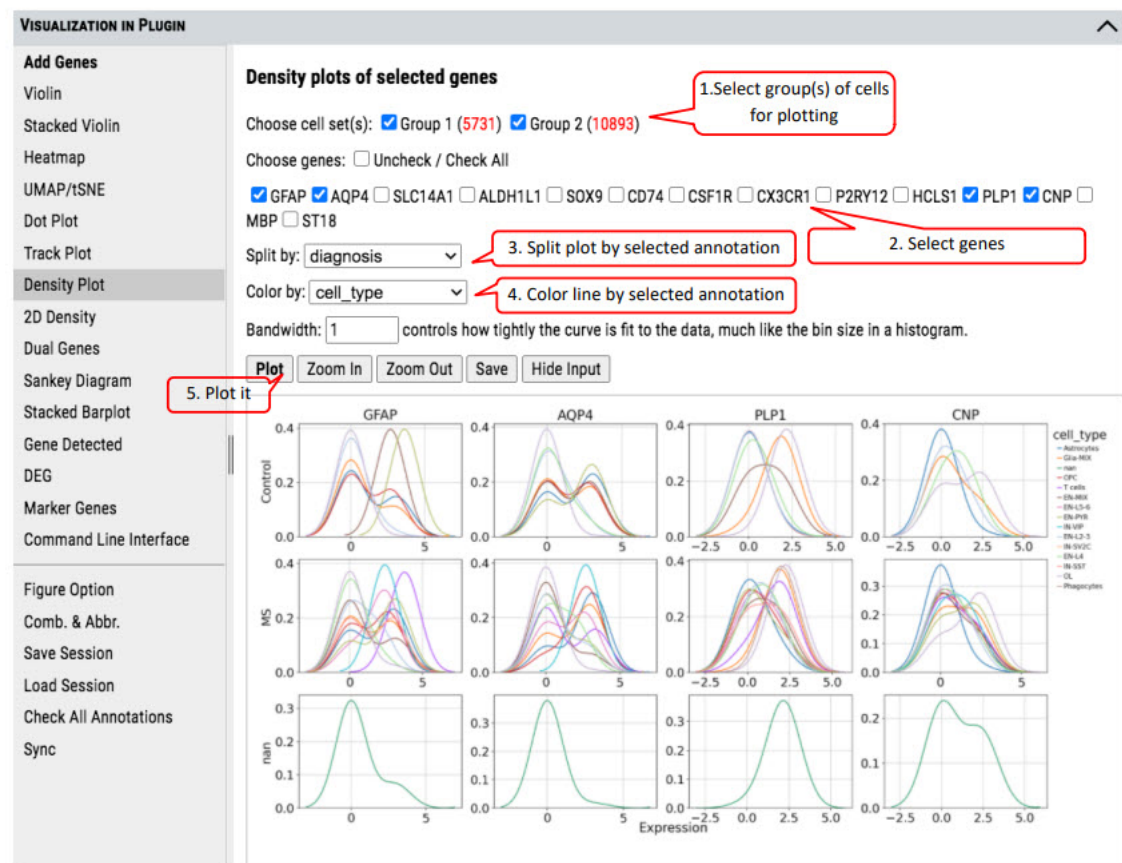


Figure 13: Density plotting of the expression of genes in each group split by one annotation while colored by another.

### 3.14 VIP – 2D Density Plot

Besides plotting of expression density of single gene, 2D density plot allows to explore the joint expression density of two genes in the cells expressing both genes above a cutoff.

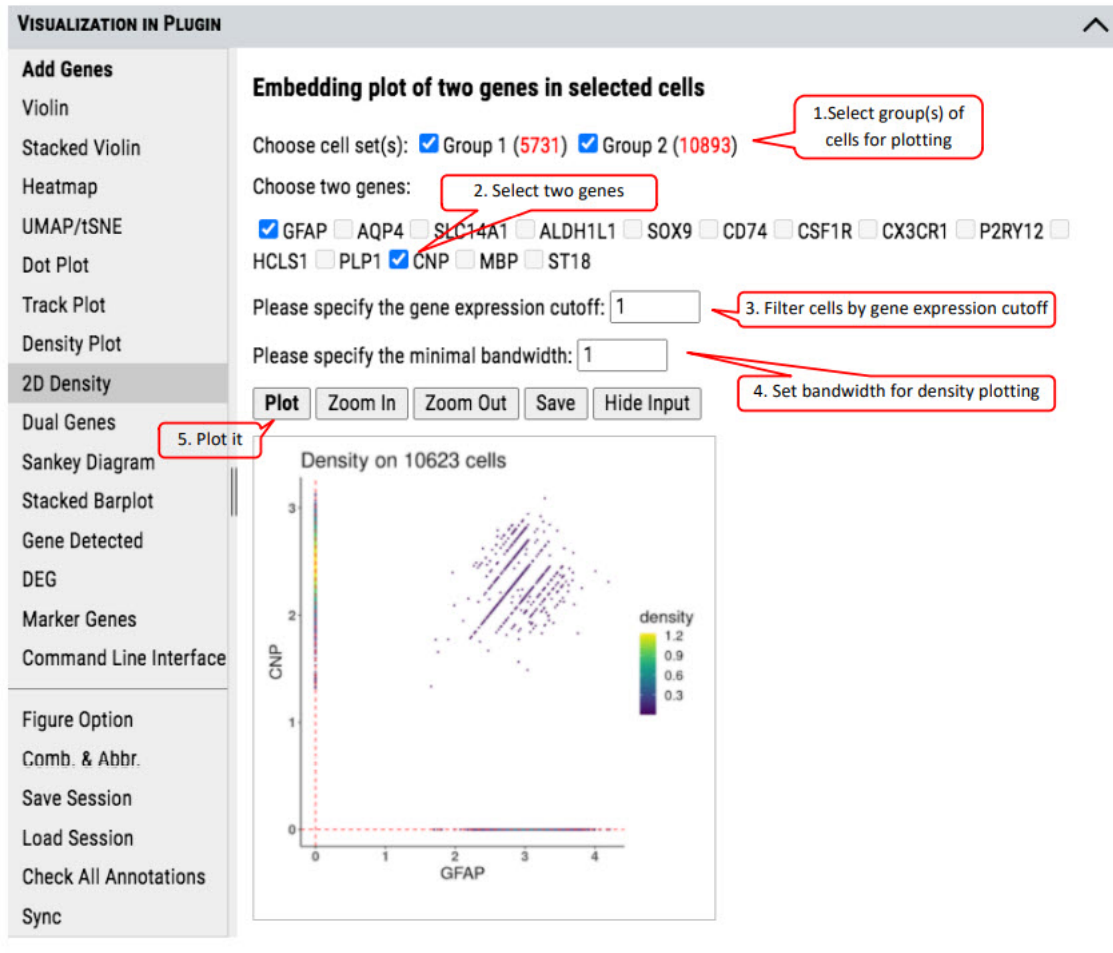


Figure 14: 2D Density plotting of expression of two genes in the selected cells

### 3.15 VIP – Dual Genes

To view the relationship of expression levels of two genes in selected cells. It is based on the embedding plot of cells while coloring cells according to the expression levels of gene(s) in each cell.

### 3.16 VIP – Sankey Diagram

Sankey diagram shows the flow of gene expression and annotations linked by cells. Gene expression is divided equally into bins so user can view distribution relationship between gene expression and annotations.

The diagram is also shown in an interactive way that user can change the layout by selecting several items (e.g., thin or thick on color bar, small or large space) from the panel. Also, user can drag these small boxes on the plot to get preferred layout and save it as high resolution SVG figure.

In addition, when you hover over mouse on a box, you can get detailed information about the source and target of flow



Figure 15: Embedding plotting of the expression of two genes in the selected cell group(s)



Figure 16: Sankey diagram provides quick and easy way to explore the inter-dependent relationship of variables.

### 3.17 VIP – Stacked Barplot

To show the distribution of cells among categories of an annotation and/or ranges of expression of a gene. Only two factors from annotations or genes can be chosen. The plot allows user to explore the distribution of cells in different views interactively.



Figure 17: The distribution of cells in the selected group(s) regarding categories of an annotation and expression ranges of a gene

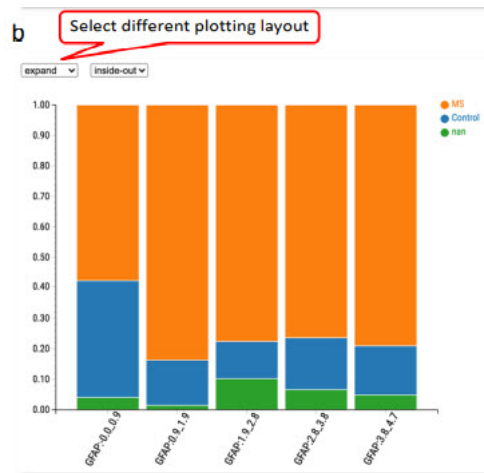


Figure 18: Expand view to show percentage instead of numbers of cells

### 3.18 VIP – Gene Detected

To show the number of genes expressed above the specified expression cut-off in the selected group(s) of cells.



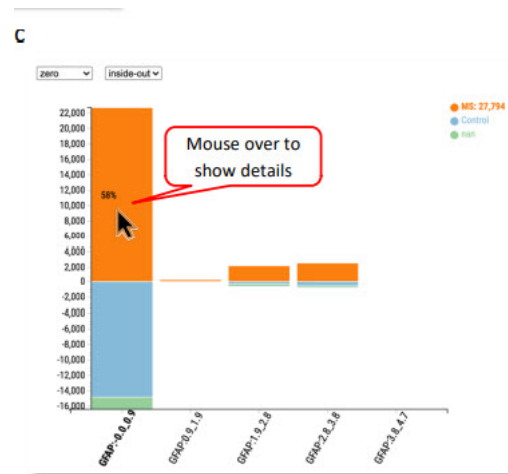


Figure 19: Mouse over a bar to show details. Y-axis is re-located for better view and the number of cells represented by the bar is also shown in legend.

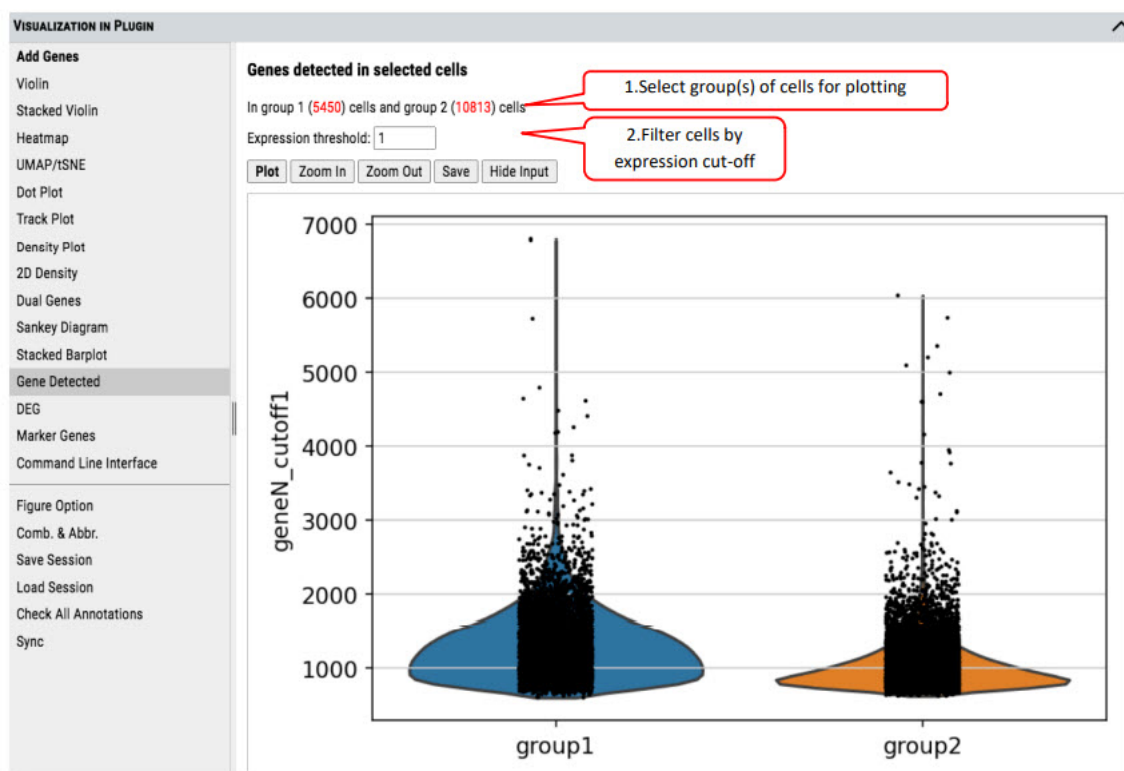


Figure 20: The number of genes with expression over the cut-off in the cells from the selected group(s) of cells

### 3.19 VIP – DEG (Differential Expressed Genes)

Besides plotting functions, cellxgene VIP also provides differential analysis between two selected groups of cells to identify differential expressed genes.

Three differential analysis statistical test methods are provided including Welch's t-test, Wilcoxon rank test and Wald's test. The statistical test results are presented in a table format including log2 Fold change, p-value and padj value (i.e, FDR value). Please note, we provide users with simple test methods for quick exploration within the interactive framework. However, there would be covariates need to be considered in a proper statistical test. Please consult your stats experts for appropriate test by using the right test method and right model.

Volcano plotting is also provided to show the log2FC vs. -log10(FDR) relationship for all genes. User can select the gene(s) from the pre-selected gene list to be highlighted with text in the volcano plot.

**a**

**1. Select group(s) of cells for plotting**

**2. Top DEGs for listing in table**

**3. Select statistical method**

**4. Select gene(s) to be highlighted in Volcano plot**

**DEG statistical output table**

Gene	Log Fold Change	p-value	padj
KCNN3	-4.38	0.0000E+00	0.0000E+00
ARHGAP31	-4.98	0.0000E+00	0.0000E+00
PSEN1	1.70	0.0000E+00	0.0000E+00
PAPLN	-7.40	0.0000E+00	0.0000E+00
HIP1	1.90	0.0000E+00	0.0000E+00
TTYH2	1.74	0.0000E+00	0.0000E+00
EGFR	-5.61	0.0000E+00	0.0000E+00
COBL	1.72	0.0000E+00	0.0000E+00
NRXN3	1.96	0.0000E+00	0.0000E+00
STON2	-4.18	0.0000E+00	0.0000E+00

Figure 21: DEG analysis between the selected group(s)

Note: The data used by DEG is unscaled (please refer to description of the dataset to find out what preprocessing was done on the data). Scaling control in the Figure Option does not apply to DEG. The three methods: 'Welch's t-test' uses t-test (assuming underlining data with normal distributions) this uses cellxgene t-test implementation, 'Wilcoxon rank test' uses Wilcoxon rank-sum test (does not assume known distributions, non-parametric test) and 'Wald's test' uses Wald Chi-Squared test which is based on maximum likelihood. 'Wilcoxon rank test' and 'Wald's test' use diffxpy's implementation.

### 3.20 VIP – Marker Genes

This functional module allows user to identify marker genes in the selected group(s) (more than 2, if 2 groups, please use DEG) of cells by annotation categories.

Four methods are provided for detecting marker genes including logreg, t-test, Wilcoxon, and t-testoverest\_var. For each identified marker gene, the gene name, scores (the z-score underlying the computation of a p-value for each gene for each group) and assigned group are listed in the output table.



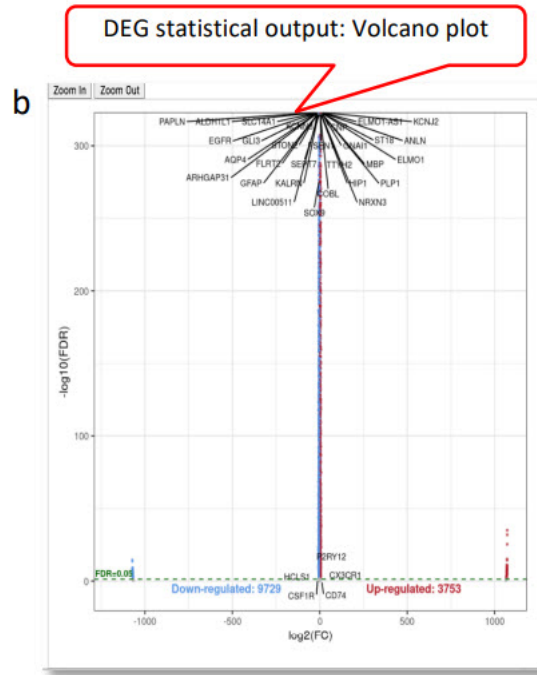


Figure 22: Volcano plotting of identified DEGs.

In each annotation category, top ranked marker genes (this example shows top 2) will be plotted by score in comparison to the rest of the categories.

Note: The four methods implementations by calling `scanpy.tl.rank_genes_groups` function: ‘logreg’ uses logistic regression, ‘t-test’ uses t-test, ‘wilcoxon’ uses Wilcoxon rank-sum, and ‘t-test\_overestim\_var’ overestimates variance of each group.

### 3.21 VIP – Command Line Interface

Although cellxgene VIP provides a rich set of visualization modules as shown above, command line interface is also built to allow unlimited visualization and analytical capabilities by power user who know how to program in Python / R languages.

Note: In CLI the AnnData (adata) object is available by default, and it is processed as ‘Description’ of the dataset states (i.e.: normalized and log transformed, but no scaled etc.). Settings in ‘Figure Option’ tab won’t apply to CLI.

### 3.22 VIP – Comb. & Abbr.

User can combine multiple annotations to create combinatorial annotation to group cells in various of plotting, e.g., stacked violin and dot plot. Firstly, user ‘uncheck all annotations’ and, secondly go to the annotation panel in main window to select annotation categories to be combined, e.g., diagnosis (Control and MS) combined with sex (female and male). After clicking on ‘Create’ button, all possible combinatorial names will be listed and ‘Custom\_combine’ will be automatically available as an option in ‘Group by’ drop down menu of many plotting functions.

User can also rename each annotation by creating abbreviations to shorten axis labels in figures.

### 3.23 VIP – Other Functions

There are other convenient functions available to user, such as ‘Save’ or ‘Load’ session, ‘Check All Annotations’ and ‘Brush’.

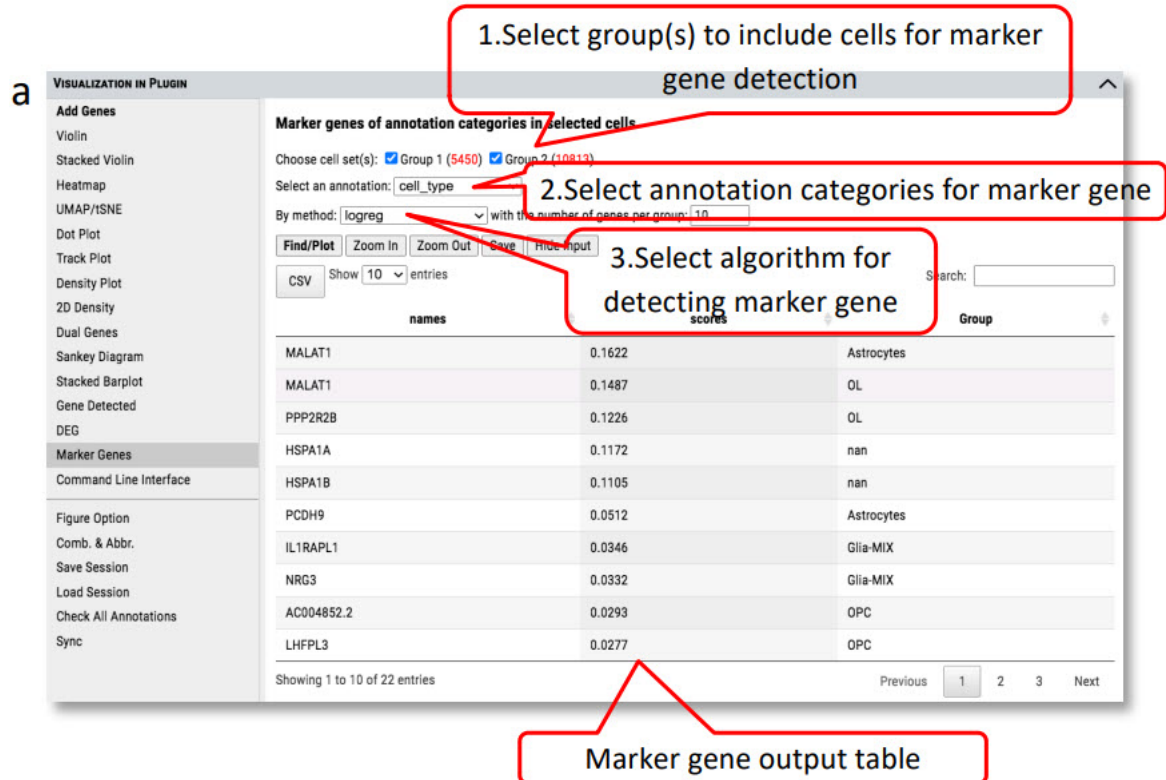


Figure 23: Marker genes detection in the selected group(s) of cells regarding to the selected annotation categories

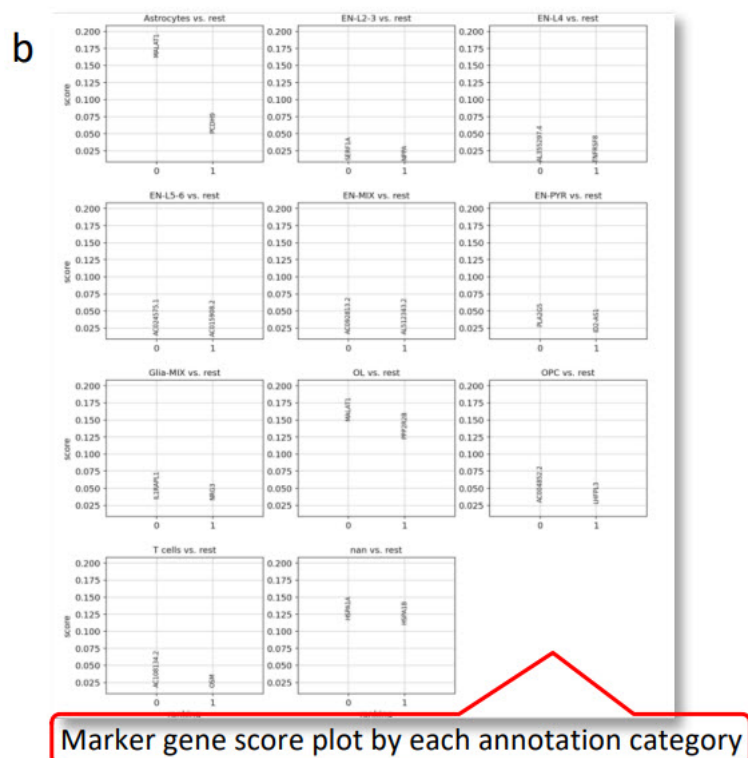


Figure 24: Plotting of identified marker genes by each category

how to program in Python / R languages.

Don't select any gene unless you operate on these genes only. In that case, it will speed up the execution of code in CLI by creating a much smaller AnnData

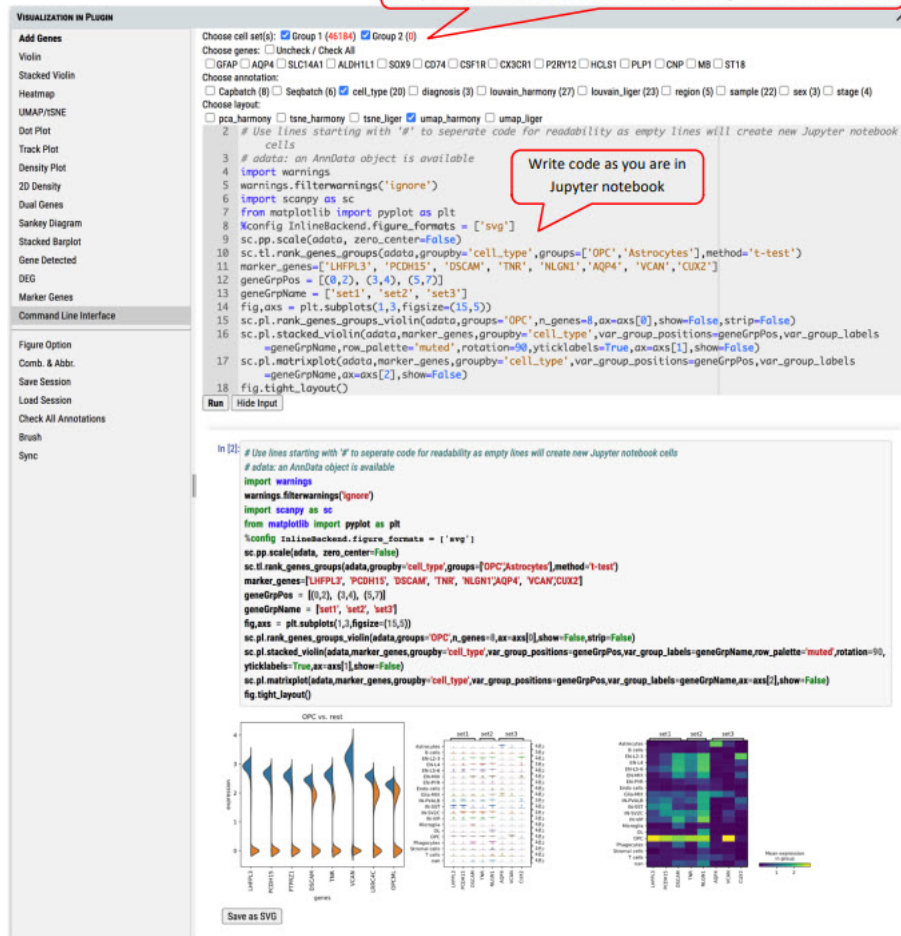


Figure 25: Command line interface for user to program for advanced plotting and statistical analysis.

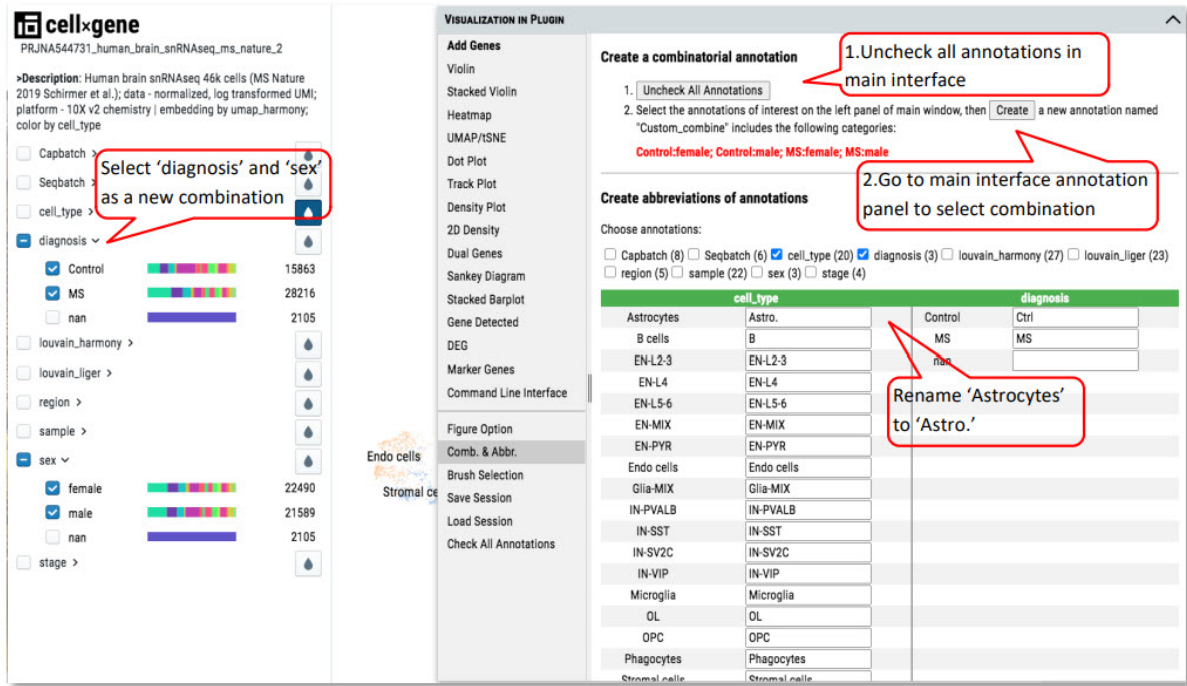


Figure 26: Comb. & Abbr. function allows user to create new annotation by combining multiple annotations and abbreviations to shorten axis labels in figures especially when custom combinatorial names are used

'Save' or 'Load' session are used to save the current cell selections and parameter settings to text file or load previously saved session file in the tool for visualization.

'Check All Annotations' is used to check all categorical selection boxes of annotations on the left panel.

'Brush' is to display exactly these selected ranges from histograms of variables on the right panel in a nice table that is not available in original cellxgene.

## 4 Methods

### Client-side Integration by a jsPanel Window (VIP)

Following section in config.sh file.

```
<script src="static/jquery.min.js"></script>
<link href='static/jspanel/dist/jspanel.css' rel='stylesheet'>
<script src='static/jspanel/dist/jspanel.js'></script>
<script src='static/jspanel/dist/extensions/modal/jspanel.modal.js'></script>
<script src='static/jspanel/dist/extensions/tooltip/jspanel.tooltip.js'></script>
<script src='static/jspanel/dist/extensions/hint/jspanel.hint.js'></script>
<script src='static/jspanel/dist/extensions/layout/jspanel.layout.js'></script>
<script
  ↪ src='static/jspanel/dist/extensions/contextmenu/jspanel.contextmenu.js'></script>
<script src='static/jspanel/dist/extensions/dock/jspanel.dock.js'></script>
```

```
<script>
  // execute JavaScript code in panel content
  var setInnerHTML = function(elm, html) {
```

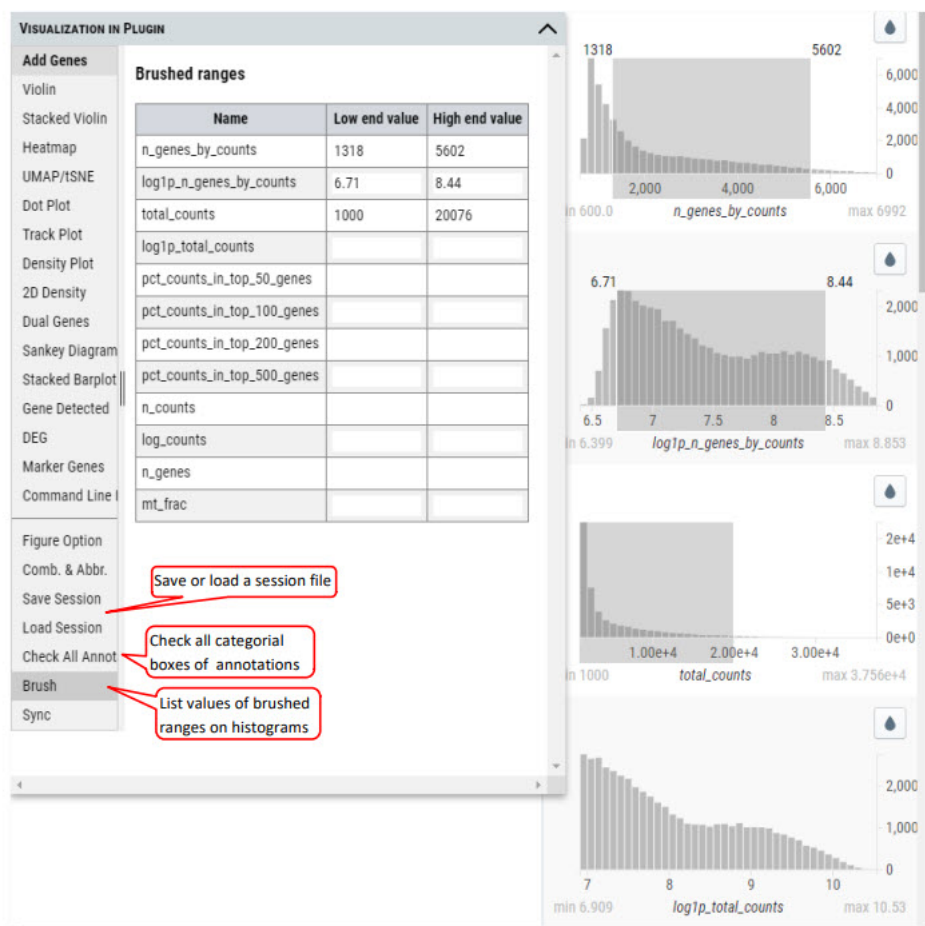


Figure 27: Other functions allow user to ‘Save’ or ‘Load’ session information, ‘Check All Annotations’ and show values of brushed ranges on histograms

```

elm.innerHTML = html;
Array.from(elm.querySelectorAll('script')).forEach( oldScript => {
  const newScript = document.createElement('script');
  Array.from(oldScript.attributes)
    .forEach( attr => newScript.setAttribute(attr.name, attr.value) );
  newScript.appendChild(document.createTextNode(oldScript.innerHTML));
  oldScript.parentNode.replaceChild(newScript, oldScript);
});
}
var plotPanel = jsPanel.create({
  panelSize: '190 0',
  position: 'left-top 160 6',
  dragit: { containment: [-10, -2000, -4000, -2000] }, // set dragging range of VIP
  ↪ window
  boxShadow: 1,
  border: "solid #D4DBDE thin",
  contentOverflow: 'scroll scroll', // adding scrolling bars
  headerControls:{
    close: 'remove',
    minimize: 'remove',
    maximize: 'remove'
  },
  headerTitle: function () {return '<strong>Visualization in Plugin</strong>'},
  contentAjax: {
    url: 'static/interface.html',
    done: function (panel) {
      setInnerHTML(panel.content, this.responseText);
    }
  },
  onwindowresize: function(event, panel) {
    var jptop = parseInt(this.currentData.top);
    var jpleft = parseInt(this.currentData.left);

    if (jptop<-10 || window.innerHeight-jptop<10 || window.innerWidth-jpleft<10 ||
    jpleft+parseInt(this.currentData.width)<10) {
      this.reposition("left-top 160 6");
    }
  },
  onunsmallified: function (panel, status) {
    this.reposition('center-top -370 180');
    this.resize({ width: 740, height: function() { return Math.min(480,
    ↪ window.innerHeight*0.6);} });
  },
  onsmallified: function (panel, status) {
    this.reposition('left-top 160 6');
    this.style.width = '190px';
  }
}).smallify();
plotPanel.headerbar.style.background = "#D4DBDE";

```

```

</script>
EOF
insertL=$(sed -e 's/[&\\\/]/\\&/g; s/|/\\|/g; s/$/\\$/;' -e '$s/\\$// ' <<<"$insertL")
sed -i "s|<div id=\"root\"></div>|$insertL\n&|" "cellxgene/client/index_template.html"

```

All functional VIP HTML and JavaScript code will be in “interface.html” that is independent of cellxgene code bases.

## Server-side Integration

Following section in config.sh file.

```
echo '
from server.app.biogenInterface import route
@webbp.route("/biogen", methods=["POST"])
def biogen():
    return route(request.data,current_app.app_config)' >> cellxgene/server/app/app.py
.
.
.

strPath="$(python -c 'import site; print(site.getsitepackages())')"
strPath=${strPath//["'"/]}
strPath=${strPath//["'"/]}
strweb="${strPath}/server/common/web/static/."
echo $strweb
cp interface.html $strweb
cp jquery.min.js $strweb
cp color_map.png $strweb

cp -R DataTables $strweb
cp -R jspanel $strweb

cp cellxgene/server/test/decode_fbs.py $strPath/server/app/.
cp VIPInterface.py $strPath/server/app/.
```

## Communication between VIP and cellxgene web GUI

Cellxgene client utilizes React Redux that is the official React binding for Redux. It lets your React components read data from a Redux store, and dispatch actions to the store to update data.

So, this line of code is appended to the end of client/src/reducers/index.js of cellxgene source code to expose the store to the browser.

```
window.store = store;
```

By doing this, Redux store holding client data and user selections are visible to VIP to access variables and dispatch actions to control cellxgene user interface. For example,

- Unselect / select a feature. GUI is refreshed automatically after dispatching.

```
window.store.dispatch({type: "categorical metadata filter deselect", metadataField:
↪ "louvain", categoryIndex: 5})
window.store.dispatch({type: "categorical metadata filter select", metadataField:
↪ "louvain", categoryIndex: 5})
```

- Get state of just finished action and synchronize gene input and cell selections from main window to VIP if corresponding action was performed.

```
const unsubscribe = window.store.subscribe(() => {
  if (window.store.getState()["@@undoable/filterState"].prevAction) {
    actionType = window.store.getState()["@@undoable/filterState"].prevAction.type;
    if (actionType.includes("user defined gene success") ||
```



```

        actionType.includes("store current cell selection as differential set")) {
            sync();
        }
    }
});

```

## Diffxpy Integration

This is the sample pseudocode, please see VIPInterface.py for actual implementation.

```

import scanpy as sc
import pandas as pd
import diffxpy.api as app
# set 1 of cells as cell1; set 2 of cells as cell2

with app.get_data_adaptor() as data_adaptor:
    X1 = data_adaptor.data.X[cell1]
    X2 = data_adaptor.data.X[cell2]

adata = sc.AnnData(pd.concat([X1,X2]),pd.DataFrame(['grp1' for i in
    ↪ range(X1.shape[0])] + ['grp2' for i in range(X2.shape[0])],columns=['comGrp']))
deg = de.test.two_sample(adata,'comGrp').summary()
#deg is a dataframe contains the folloing columns ['gene','log2fc','pval','qual']

```

### 4.1 Create h5ad file from Seurat object

First, export the following from Seurat object in R: **expression matrix (assume normalized), meta-data and coordinates (pca, tsne, umap)** as separate txt files.

Next in Python, create an AnnData object from 10x (scanpy.read\_h5ad function) as a starting point. Then replace the expression matrix, meta data and coordinates as following, a h5ad file would be generated.

```

import sys
import scanpy as sc
import pandas as pd
import numpy as np
import seaborn as sns
from numpy import ndarray, unique
from scipy.sparse.csc import csc_matrix

adata= sc.read_h5ad("previous generated .h5ad")

# read clustering res
xpca = pd.read_csv("./data/harmony_clustered.h5ad.pca_coordinates.txt", sep='\t',
    ↪ encoding='utf-8')
xtsne = pd.read_csv("./data/harmony_clustered.h5ad.tsne_coordinates.txt", sep='\t',
    ↪ encoding='utf-8')
xumap = pd.read_csv("./data/harmony_clustered.h5ad.umap_coordinates.txt", sep='\t',
    ↪ encoding='utf-8')
xobs = pd.read_csv("./data/harmony_clustered.h5ad.meta_data.txt", sep='\t',
    ↪ encoding='utf-8')

```



```

xpca.set_index('index', inplace=True)
xtsne.set_index('index', inplace=True)
xumap.set_index('index', inplace=True)
xobs.set_index('index', inplace=True)

adata.obsm['X_pca'] = np.array(xpca.loc[adataRaw.obs.index])

adata.obsm['X_tsne'] = np.array(xtsne.loc[adataRaw.obs.index])
adata.obsm['X_umap'] = np.array(xumap.loc[adataRaw.obs.index])
adata.obs = xobs.loc[adataRaw.obs.index] # this is a pandas dataframe

# read in expression matrix as numpy.ndarray as following:
exp_mat = np.loadtxt(fname="expression matrix .txt")
adata.X = exp_mat

# convert dense matrix into sparse matrix to save storage space and memory usage
adata.X = csc_matrix(adata.X)_matrix

# add short description and initial graph settings. "|" and "by" are delimiters for
↪ VIP to parse the initial settings. Please follow the same rule for your own h5ad
↪ files.
adata.obs['>Description'] = ['Human brain snRNAseq 46k cells (MS Nature 2019 Schirmer
↪ et al.); data normalized, log transformed and scaled UMI; platform - 10X v2
↪ chemistry | embedding by umap; color by cell_type']*adata.n_obs

# Then last step to save h5ad:
adata.write_h5ad("final output.h5ad")

```

## 5 Helpful Tips

### 5.1 Handle nulls in categorical annotation

Such nan's in categorical annotation would cause trouble in VIP because it cannot be converted to string. Here is how to handle it, let's call the annotation X\_annotation:

```

# Cast to str from categorical
adata.obs = adata.obs.astype({'X_annotation': 'str'})

# replace all of nan by 'nan'
adata.obs["X_annotation "][adata.obs["X_annotation "].isnull()] = 'nan'

```

### 5.2 Display full traceback stack for debugging in VIP

It follows the global setting. Please set “—verbose” to launch cellxgene server.

### 5.3 Pitfall of using special characters

In the mode which allows user to create manual annotation in cellxgene, user should try to avoid using hyphen (“-”) in name label. It would cause client-side issue. Please try to use underscores.

## 5.4 Potential use for bulk or pseudo bulk sample dataset

Once the data matrix is replaced by sample x gene matrix, cellxgene VIP framework can handle regular bulk / pseudobulk RNAseq datasets. Simply replace “cells” by “samples”. All plotting functions would still work.