# User documentation of the 'Superviolin' Python package

The superviolin package can be used to create Violin SuperPlots. As well as a tutorial on how to use the package, this documentation provides an introduction to Python installation and package management to enable non-programmers to easily setup a Python workspace to use this software.

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### 1. Python installation for non-programmers

If you haven't used Python before, you will be installing a Python distribution called Anaconda, which is widely used by scientific Python developers. A distribution is a bundle that contains the components necessary to run Python, as well as some domain-specific packages. For example, the Anaconda distribution is intended for use as a data science solution and contains packages for machine learning, scientific data analysis, and much more. It ships with over 200+ packages, including the package dependencies for Superviolin.

However, as this guide is aimed at non-programmers, we will be installing the Miniconda distribution, which has the same functionality as Anaconda, but saves on hard drive space and installation time by only installing the basic packages required for it to function. If there are other packages you want to use in the future, you can install them manually later on.

#### To install Miniconda:

- 1. Download the Python 3.x Miniconda installer for your OS at <a href="https://docs.conda.io/en/latest/miniconda.html">https://docs.conda.io/en/latest/miniconda.html</a>.
  - Superviolin requires the installed Python version to be at least 3.6.2
- 2. Run the installer and follow the on-screen instructions, using the default settings you are prompted with.
- Congratulations! You now have a functional Python environment on your computer!

You can now proceed to installing Superviolin. Python has a so-called package manager which is used to install, update, and remove packages. It ensures that all package

dependencies are maintained and will warn you if something would be broken by a new package installation.

Superviolin uses basic functions from Numpy, Scipy, and Pandas which aren't likely to be changed dramatically, so you don't need to worry about the version number. As long as you installed Python 3.6.2 or later, Superviolin should install just fine.

Installing Python packages requires you to enter text commands in the terminal (Mac, Linux) or in Anaconda Prompt (Windows). Spacing and upper/lower case must be maintained to ensure the commands run as expected. Most common programming errors occur due to these mistakes and they affect both experienced and beginners alike, though experienced programmers tend to get these errors less often.

To install Superviolin, open up the terminal or Anaconda Prompt and enter:

The package manager will list the dependencies required and install them for you. Once you see a message stating superviolin was successfully installed, you can proceed to the next section and try out the package with your data.

# 2. Superviolin command overview

The superviolin CLI has 3 commands:

#### demo

This command creates a Violin SuperPlot using dummy data that ships with the package. Run superviolin demo from your command prompt after installation to be sure the package is working. It should generate a figure similar to **Supplemental Figure S2** above. If you get an error, please contact Martin Kenny at <a href="mailto:mkenny5@tcd.ie">mkenny5@tcd.ie</a> to report your issue so that it can be corrected.

#### init

The superviolin init command generates an 'args.txt' file in the current directory, which will be used to generate a Violin SuperPlot based on the Excel or csv file of your data, located in the same folder.

### plot

The superviolin plot command renders the Violin SuperPlot as a figure. This layout can be edited prior to saving. A paired or unpaired t-test is run if there are 2 conditions to be compared. A one-way ANOVA followed by Tukey's tests with Bonferroni corrections are run when there are 3 or more conditions. Statistical test results are output on the Anaconda command line interface, as well as saved in a 'posthoc\_statistics.txt' file in the same folder as the data. There is no strict limit for the number of conditions that can be plotted/compared, apart from the size of the plot window which limits the navigation of the plot.

NOTE: To run any of these commands, enter it into your terminal or Anaconda Prompt.

### 3. Generating a Violin SuperPlot

Superviolin comes with a command-line interface (CLI) so you can generate SuperPlots using the terminal or Anaconda Prompt. You can also import the "Superviolin" class into your Python scripts and extend its functionality to create your own specific brand of Violin SuperPlot, but first, let's understand how it works. This section also contains a simple how-to for non-programmers.

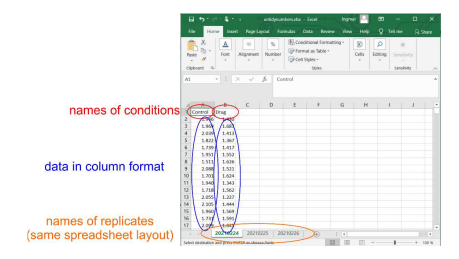
1. Prepare your data as a \*.csv or Excel file and place it into an empty folder. The data should be arranged in either (A) the tidy data format where you have a column for each variable/property and a row for each observation, or (B) 'untidy' data in an Excel workbook where each sheet is an experimental replicate containing columns for each condition. The tidy data format is preferred because it is more flexible. For tidy data, the Condition column specifies if an observation belongs to a different condition (control vs drug, different drug concentrations, wild type vs mutant, etc.), the Value column(s) specify the value(s) of the measured variable(s) (spreading area, circularity, intensity, etc.), and the Replicate column specifies which replicate the observation is from (donor ID, date, well, etc.).

Condition	Area	Ellipticity	Replicate
Control	25.2	1.434	Donor 1
Control	17.1	1.017	Donor 1
Drug	26.6	1.418	Donor 1

Drug	12.4	1.592	Donor 1
Control	28.9	1.481	Donor 2
Control	35.3	1.294	Donor 2
Control	39.5	1.374	Donor 2
Drug	9.8	1.767	Donor 2
Drug	18.9	1.660	Donor 2
Drug	32.2	1.376	Donor 2
Control	34.1	1.374	Donor 3
Drug	25.5	1.810	Donor 3

Example of a data file in tidy format (either csv or Excel file).

<u>For 'untidy' data</u>, column names have to be exactly the same across replicates to allow the software to combine the data for plotting; the data in each column doesn't have to be paired data, i.e. can have different number of entries. Column names should not contain special characters or sub/super-script.



Example of a data file in an 'untidy' format (several Excel spreadsheets).

- 2. Open the terminal or Anaconda Prompt. Navigate to the folder containing the data file by typing & copy-pasting cd drive:\full\path\to\your\data\folder into the command line (replace the 'drive:\...' by your actual path).
- 3. Run superviolin init to generate the args.txt file in this folder.
- 4. Edit the args.txt file with the parameters specific to your data file. Optional arguments are detailed in section 6. The required parameters are:
  - i. **filename**, the name of your data file. Must be in CSV or Excel format.

ii. **data\_format**, whether the data follows the format of tidy data (table above) or 'untidy' data (figure above). NOTE: when using the untidy format, the package creates a tidy format dataframe from the Excel sheets, and the columns of this dataframe are named from the following 3 arguments.

For tidy data, additional inputs are required:

- iii. **condition**, the column in your dataset which specifies the experimental conditions.
- iv. **value**, the variable to be plotted from your dataset.
- v. **replicate**, the name of the replicate column in your dataset.
- 5. Save the args.txt file.
- 6. Run superviolin plot to generate your Violin SuperPlot which will open in the "Figure 1" window. Please note that the Figure window needs to be closed to be able to access the command line / Anaconda prompt again.

Simply changing the filename in the args.txt file will allow you to apply those settings to other CSV or Excel files with the same structure, provided the new filename is in the same directory as the args.txt file.

### 4. Editing your Violin SuperPlot

Most of the editing of the Violin SuperPlot is done directly in the args.txt file. The basic Violin SuperPlot generated by the program can then be further adapted to edit the amount of whitespace around your plot. **Supplemental Figure S2** shows the Violin SuperPlot generated by the superviolin demo command. The whitespace around the plot is already optimized within the code, but the size of these spaces can be adjusted by clicking on the slider button to open a menu of sliders corresponding to the whitespace to the top, bottom, left, and right of the plot.

# 5. Saving your Violin SuperPlot

Once satisfied with the plot, you have the option to save it either as a rendered image or a vector graphics image. Clicking the save icon will open a dialog to choose the location for saving your figure and allow you to select the appropriate output format from the dropdown menu.

Rendered images: preferred are lossless formats (TIFF, PNG). The plot is saved at the resolution specified during creation of the plot (see section 6 below for optional arguments in the 'args.txt' file).

For maximal resolution and further editing in other programs, it is recommended to save the figure in SVG vector graphics format.

## 6. Optional arguments in args.txt used to customize the Violin SuperPlot

For example usages of optional arguments, see Supplemental Figure S3.

- i) Xlabel and Ylabel indicate the labels you wish to use for the x and y axes. These arguments can be left unchanged if you don't wish to have a label on either axis. Special characters and super/sub-script formatting can be specified using Python's string formatting: <a href="https://pythonforundergradengineers.com/unicode-characters-in-python.html">https://pythonforundergradengineers.com/unicode-characters-in-python.html</a>
  - For example, the y-axis label "spreading area ( $\mu$ m<sup>2</sup>)" in Supplemental Figure S2 is input into the args.txt file as "spreading area ( $\mu$ m<sup>2</sup>)", where  $\mu$  and the  $\mu$ 72\$ correspond to the Greek letter  $\mu$  and the  $\mu$ 7.
- ii) **Order**, the order of the experimental conditions to be displayed on the x axis. The values in this argument MUST be separated by a single comma and single space otherwise it will throw an error. There is no default order.
  - In the above example: 'Control, Drug'
- iii) **Middle\_vals**, the central measure of each replicate. Displayed as circles over each replicate stripe. Default is mean. Accepted values are "mean", "median", or "robust". The "robust" mean ignores the upper and lower 2.5% of the data before taking the mean of the inner 95%.
- iv) **Centre\_val**, the central tendency (mean or median) calculated from replicate means (or medians) and shown between the error bars on the plot. Default is mean.
- v) **Error\_bars**, whether to use the standard error of the mean (SEM), the standard deviation (SD), or 95% confidence interval (CI) for plotting error bars. Default is SEM.

- vi) **Bw**, the bandwidth of the kernel density estimator, a decimal smoothing factor for stripe and violin outlines. The default value is "None", which means an optimal value will be calculated using Scott's Rule and the software will show this quantity in the Anaconda Prompt or Terminal window.
- vii) **Paired\_data**, whether to run a paired samples t-test or a non-paired t-test for statistical comparison of replicates. Values are either "yes" or "no". Only applicable if exactly two conditions are provided/present.
- viii) **Stats\_on\_plot**, choose whether to display t-test statistics on the plot. Only applicable for 2 experimental conditions. Either "yes" or "no".
- ix) **Ylimits**, the minimum and maximum values to display on the y-axis. Should be in the form "low, high" e.g. "0.5, 1.5". If this argument is to be ignored, leave the value as "None".
- x) **Total\_width**, the overall normalised width of each violin in decimal form. Should be between 0 and 1. Default is 0.8.
- xi) **Linewidth**, the width of lines (in points) used to plot summary statistics and the outline of each violin. Default is 1.
- xii) **Show\_legend**, whether to show a legend or not. The legend links the names of each replicate to the corresponding colour-coded stripe in each Violin SuperPlot.
- xiii) **Cmap**, the colourmap specifying the colours used for the replicates. Set2 is the default colourmap, but users can specify a list of colours or a colourmap of their choosing from <a href="https://matplotlib.org/stable/gallery/color/named\_colors.html">https://matplotlib.org/stable/tutorials/colors/colormaps.html</a>
- xiv) **Dpi**, the dots per inch used for saving the plots. Plots are rendered at 300dpi for display purposes, but can be saved at much higher resolution, as Python cleverly manages an object for the figure which can be rendered and saved at different dpi values. The default value for saving is 600dpi, but can accommodate other values e.g. 300 or 1200.