EDAT Documentation v.1

**Background**

**Objectives**

* The overall objective of the project was to create an app using R-Shiny in order to allow a person to explore gene expression data without extensive computational knowledge and rigor. The app will incorporate two dimensionality reduction algorithms, PCA and t-SNE, to help aid in data visualization. The user then may further visualize the data by selecting only specific genes, time points, and donors they wish to visualize.
* In order to accomplish this, some processing is needed to convert the raw fold change data into a usable format. Currently the program only accepts pre-processed gene expression data using the ration of change over the original expression level, but we hope to have the program process raw data in the future. Once the data has been processed, the user is able to subset the data using the UI which calls the subset() function in the program which extracts only the selected data. See the image in results for more information.
* In regards to the PCA performed by the program, we are using the prcomp() function which accepts the scale and center parameters which will soon be modifiable by the user. The variance of each dimension is also computer with the PCA and is displayed in a corresponding scree plot.
* For the t-SNE performed we are using the function Rtsne() which is part of the RTsne package. We will implement the function such that the user can adjust the dimension, initial\_dimension, perplexity, pca, and max\_iter parameters.

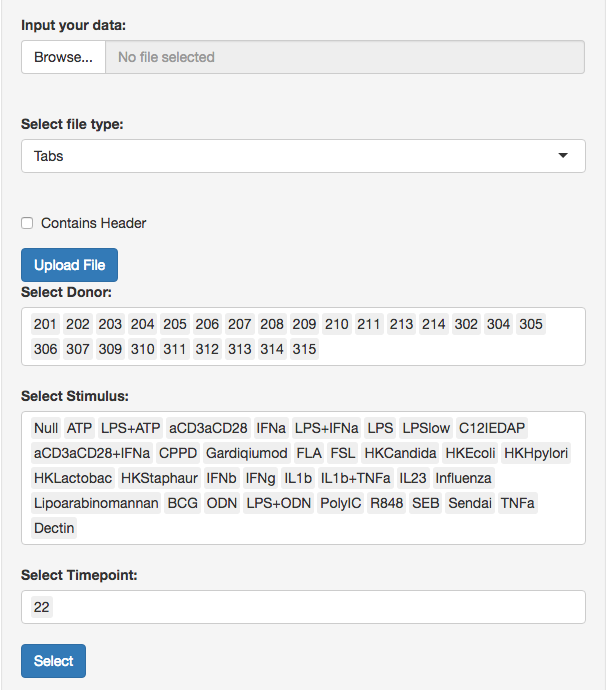
**Results**

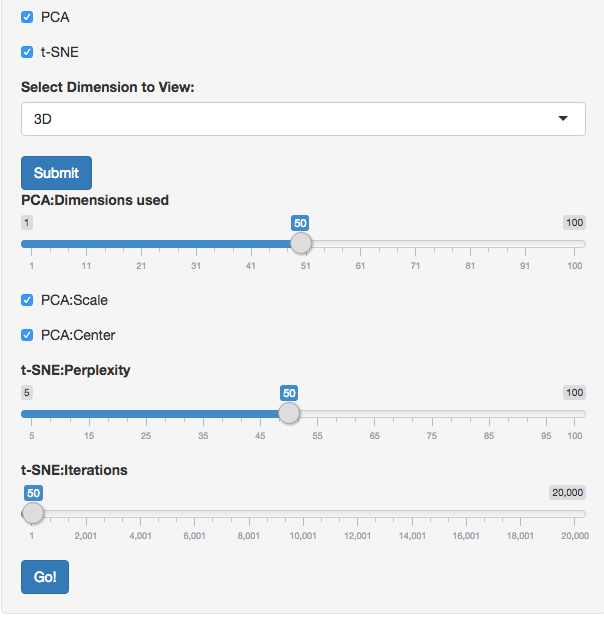
Pictured to the left is the sidebar for the application. It allows for the user to upload a file they wish to be parsed, select what the deliminator is and whether there is a header and begin upload. After the file is uploaded the selection menu is presented to the user.

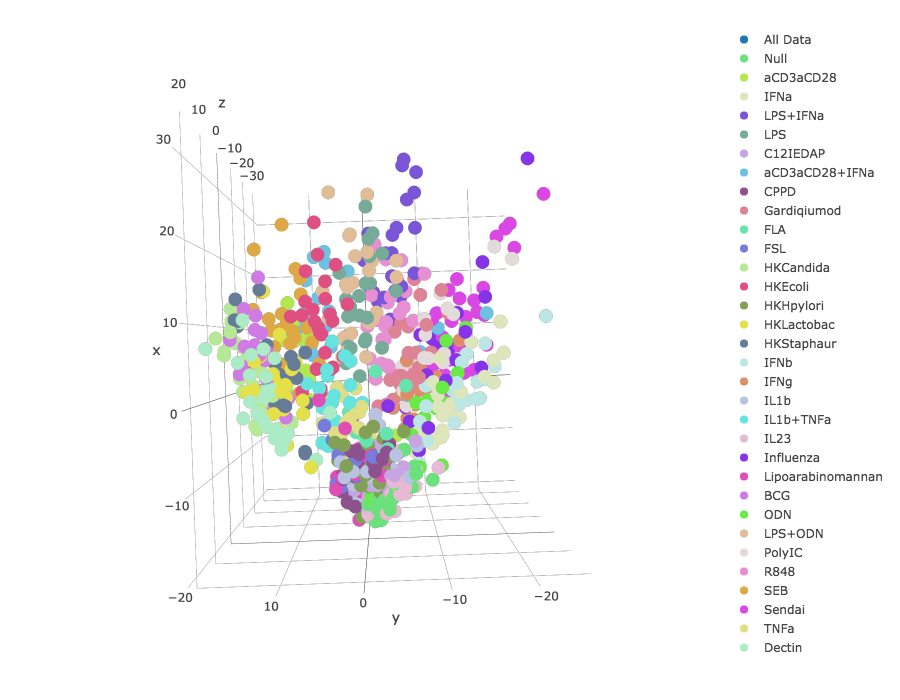
This menu is seen directly underneath the upload menu and allows for the user to select which donors, stimuli, and time points from the original file they want to include in the analysis.

After pressing select, the user is then prompt to choose the dimension and algorithm they wish to use. They are given a choice between Principal Component Analysis (PCA), t-Distributed Stochastic Neighbor Embedding (t-SNE) or both.

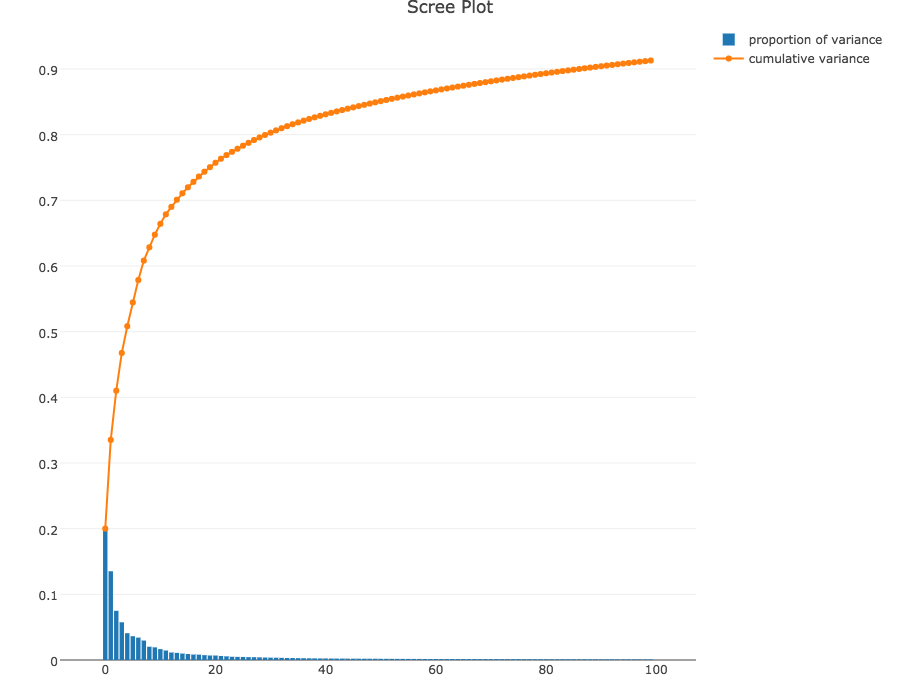
Once those two settings are submitted, the user is able to tune parameters relevant to the algorithm they chose and then produce the plot by pressing Go.

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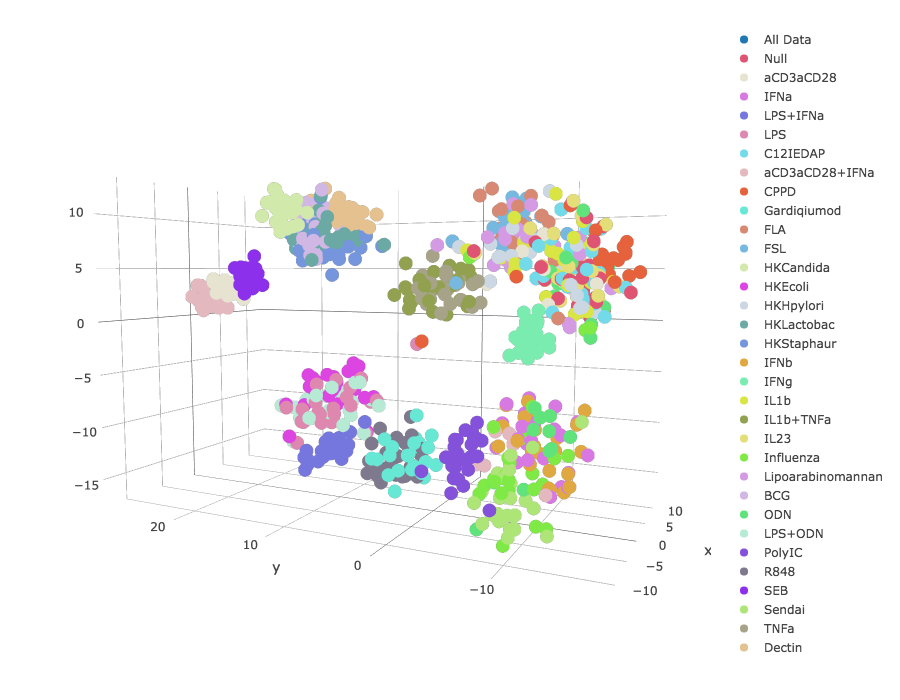
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[EXPLANATION OF IMAGE]



For each run of PCA a corresponding scree plot is displayed showing the variance captured by each dimension and the cumulative variance captured over a span of dimensions. This plot allows the user to see how much information is lost with the reduction of the data into the 2 or 3 dimensions seen in the PCA plot.



[EXPLANATION]

[DIFFICULTIES AND IMPROVEMENTS]

* Ui
* Labels hover
* Optimization
* Parameter selection
* Display progress

**Conclusion**

[NEW IDEAS AND WORK REMAINING]

**Usage**

With Processed Data:

With Raw Data: