



Step-by-step guidelines to Run the AutoNSGACytoNet framework:

1. Collect data from <https://xenabrowser.net/datapages/>: Collect GDC TCGA gene expression RNAseq TPM data for BRCA, LAML, LUAD, LUSC, COAD, SKCM, GBM and LIHC Pancancer Cohorts.
Already downloaded data can be found in the [dataset](#)
2. The [Dataset](#) folder contains directory-wise carcinogenic Data. Here, each directory is named after each cancer type, and each directory contains a .tsv file format for each cancer type. These initial tsv files need to be Transposed so that we can use the genes as features and merge together to result in a final Dataset. Using the [merging](#) script, the final dataset can be achieved.

PS: Step 2 has one directory-related dependency. We need to set the directory/path of [Dataset](#) manually to run the code.

3. Step 3 is the preprocessing step. Data in the [file](#) can be used or generated using the [merging](#) script. The notebook [Preprocessing Data](#) can be used to create preprocessed CSV files. It also has director/path dependency. Need to set the path manually. The outcome is already stored as [Preprocessed 8 cancer genes.csv](#) in Google Drive.
4. Step 4 is the autoencoder-based feature selection step. Here, the input is as [Preprocessed 8 cancer genes.csv](#) file. After running the notebook [autoencoders-with-cv.ipynb](#) we will get out csv files [top 0.5 percent features cv.csv](#), [top 0.25 percent features cv.csv](#), [top 1.0 percent features cv.csv](#). Again, this notebook has a directory/path dependency. We need to change accordingly.
5. The next step is the NSGA-2 Step. This step is implemented in [nsga2-with-rf-cv2.ipynb](#) notebook. It will take these files ([Preprocessed 8 cancer genes.csv](#), [top 0.5 percent features cv.csv](#), [top 0.25 percent features cv.csv](#), [top 1.0 percent features cv.csv](#)) as input. We need to set the path manually here as well.
6. After running the notebook [nsga2-with-rf-cv2.ipynb](#), we will get three output files [NSGA2 77 compression 1.csv](#), [NSGA2 308 compression 3.csv](#), [NSGA2 154 compression 2.csv](#) with gene subsets.
7. At step 7, we will need the gene subset from [NSGA2 308 compression 3.csv](#) file. We will upload the gene set in [STRING database](#) to generate Protein-Protein-Interaction PPI network. The network needs to be downloaded in .tsv format. Which is also available in Google Drive as [string_interactions.tsv](#)
8. At step 8 we need to upload the [string_interactions.tsv](#) network in Cytoscape software with cytohubba dependency. It is an open-source tool. At this step, we will select 13 hub genes. These genes will be used for evaluation purposes.
9. This is the final evaluation step. Here we need to run the [classification.ipynb](#) notebook. This notebook requires [Preprocessed 8 cancer genes.csv](#) Dataset. The gene subset acquired from Cytoscape is already hardcoded in the notebook. After running the notebook, we will get desired evaluation metrics.

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