Project 2 – Signature creation and iLINCS API

This assignment builds upon the R/shiny class and expands the API example.

- 1. For the assignment use the dataset TCGA_breast_cancer_ERstatus_allGenes.tsv. You may choose to "debug" your code with a smaller dataset TCGA_breast_cancer_ERpositive_vs_ERnegative_PAM50.tsv or TCGA_breast_cancer_LumA_vs_Basal_PAM50.tsv.
- 2. Your assignment is to develop code using R Shiny to present user interface that allows a user to upload a tsv file, presents an option to split the samples into two groups based on the available metadata and create a signature. The signature is then submitted to the iLincs API which retrieves concordant signatures.
- 3. Template is provided in the Project 2 github project (https://github.uc.edu/uc-datascience/Project2.g it).
- 4. The template is missing the calculation of the differential expression please use t-test to calculate t-statistic / p-value.
- 5. Expand the template to allow users to filter the input file to L1000 genes only (See the include L1000.txt file).
- 6. Further expand the template to allow users to submit only top 100 differentially expressed genes.
- 7. Compare results with iLincs
- 8. Extra credit for a heatmap or other visualizations.

The assignment is due on – March XX, 2020 midnight.

The submission should be zip compressed file named "project2-[your UC username].zip" (e.g. "project2-lastnfi.zip") which includes any supporting R files. The zip file should be uploaded canopy. The assignment entry in Canopy will be created shortly.