## BIOL-GA 2031. Statistics and Machine Learning in Genomics Homework 02

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In the jupyter notebook LOGISTIC\_REGRESSION\_PY.IPYNB I wrote some Python code for loading the data and performing a logistic regression + Bootstrapping on some Dengue disease data. You could use that code I wrote or write one yourself to answer the following numerals. The equivalent R code I wrote can be found in LOGISTIC\_REGRESSION\_R.IPYNB, I'm not super versed in R so that code might have a lot of bits to improve.

- 1. How does the first two moments, mean and variance, of the expected prediction accuracy change with the numbers of bootstraps?
- 2. Use just the 5 genes significant in the ANOVA (volcano plot) and do the bootstrapping+logistic regression again. How does the estimate of the expected prediction accuracy change? Use Figures to present and discuss your results.
- 3. Repeat 1. again but just with the genes significant in the ANOVA.
- 4. Compare 1. and 3., and discuss. Does the previous knowledge about the significantly expressed genes during disease contribute to the estimate of the expected prediction accuracy? Did you require a lower number of bootstrap samples?
- 5. Write the leave-one-out cross validation code with partitions of 10 each (some will have more because there are 53 data-points).
  - 5.1 How does the expected estimated prediction accuracy compare to that one with bootstrapping? Use figures to present your results and discuss.
  - 5.2 How does the variance of the prediction accuracy compare to the one estimated with bootstrapping?