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

Prof. Manpreet S KatariTA. Jaime Cascante VegaEmail. jc12343@nyu.edu

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. A bootstrap sample of a dataset \mathscr{D} of size N, $\mathscr{D} = [x_1, x_2, x_3, \dots, x_N]$ correspond to sample with replacement with equal probabilities each x_i and forming a new dataset $\mathscr{B} = [b_1, b_2, \dots, b_N]$.

In the N = 2, $\mathscr{D} = [x_1, x_2]$ and the possible bootstrap samples are $\mathscr{B}_1 = [x_1, x_1]$, $\mathscr{B}_2 = [x_1, x_2]$, $\mathscr{B}_3 = [x_2, x_2]$, $\mathscr{B}_4 = [x_2, x_1]$.

How many possible bootstrap samples are in a dataset of size N?

- 2. Let $X_1, X_2, ..., X_n$ be independent and identically distributed (iid) random variables with mean μ and variance σ^2 . Consider a bootstrap sample of those variables of size n denoted by $Y_1, Y_2, ..., Y_n$.
 - 2.1 Calculate $\mathbb{E}[Y_i]$ and $\text{Var}[Y_i]$.
 - 2.2 Denote \bar{Y} the mean of the bootstrap sample as show below. Calculate the conditional expected value of the mean and conditional variance of the bootstrap sample given the original dataset, i.e. calculate $\mathbb{E}[\bar{Y}|X_1,X_2,\ldots,X_n]$ and $\text{Var}[\bar{Y}|X_1,X_2,\ldots,X_n]$.

$$\bar{Y} = \frac{1}{n} \sum_{i=1}^{n} Y_i$$

- 3. How does the first two moments, mean and variance, of the prediction accuracy change with the numbers of bootstraps?
- 4. Use just the 5 genes significant in the ANOVA (volcano plot shown in the Figure below) 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.
- 5. Repeat 3. again but just with the genes significant in the ANOVA.

- 6. Compare 3. and 5., 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?
- 7. Write the leave-one-out cross validation code with partitions of 10 each (some will have more because there are 53 data-points).
 - 7.1 How does the expected estimated prediction accuracy compare to that one with bootstrapping? Use figures to present your results and discuss.
 - 7.2 How does the variance of the prediction accuracy compare to the one estimated with bootstrapping?