

Algorithms in Bioinformatics

Stabilization matrix method - Mini project

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ABSTRACT

Summary We look at two Stabilization matrix methods [1] used for predicting binding affinity of immunogenic peptides to major histocompatibility complex [2] (MHC) molecules. The testing data available was from 35 different MHC molecules from HLA-A and HLA-B, with number of peptides ranging from 59 to 3089.

INTRODUCTION

Blabla bla

$$E = \frac{1}{2} \sum_{i} (O_i - t_i)^2 + \lambda \sum_{l} w_i^2$$
 (1)

BlaBlaBla

MATERIAL AND METHODS

We were given semi-completed code for the implementation of the Stabilization matrix method. The minimization procedures were gradient descent in one case and monte carlo in the other. The testing data available was from 35 different MHC molecules from HLA-A and HLA-B, with number of peptides ranging from 59 to 3089. In addition to finishing the code we added some optimizations like skipping unnecessary memory allocations at performance critical locations. We ran the matrix creation on each 35 MHC molecules to get the broadest perspective. The data for each MHC molecule was splited into five parts, equally (or nearly equally) sized. Each part was then used for the validation of the model trained on the rest (4/5) of the data, resulting in 5 iterations to be used for (5-fold) cross-validation. As noted in [1] the use of the λ value suppresses the effect of noise in the measured data. In the case of zero value λ , the optimal entries for the weight vector w minimize the difference between predicted and measured values. Minimizing with a non-zero value for λ results in a shift of the optimal entries in w towards values closer to zero. It should be noted here that the λ value used as a parameter in the code is the per_target normalized λ value, not the global one.

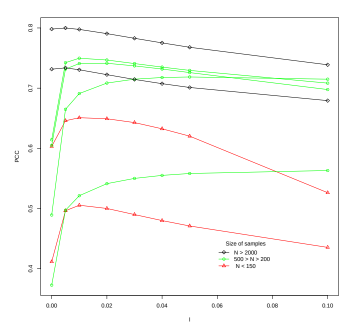


Fig. 1. Pearson's correlation coefficient of different sample using increasing values of λ . The sample were grouped by the size of the dataset. We can see how dataset of similar size show a similar trend of the Pearson correlation λ depending.

RESULTS

Figure 1 shows the trend for Pearson's correlation coefficient (PCC, prediction accuracy) using different λ values for various data sizes. As a general rule, as the data size gets larger the less important the effect of a non-zero λ becomes. For large data sets a λ value of zero seems to give the best results. A large dataset should average out the noise. Similar graph showing results from the Monte Carlo version says the same story (Figure 2). For the small datasets an optimal λ value is less than 0.02. Next we run the smm algorithms on a data from four different kinds of MHC molecules showing the error as a function of iterations. Here the default λ value of 0.05 is used (Figure 3).

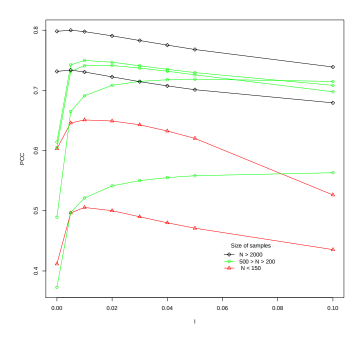


Fig. 2. Pearson's correlation coefficient of different sample using increasing values of λ . The sample were grouped by the size of the dataset. We can see how dataset of similar size show a similar trend of the Pearson correlation λ depending.

REFERENCES

- [1]B. Peters and A. Sette. Generating quantitative models describing the sequence specificity of biological processes with the stabilized matrix method. *BMC Bioinformatics*, 6(132), 2005.
- [2]Wikipedia. Major histocompatibility complex, 2010. [Online; accessed 18-October-2010].