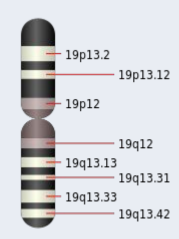
With current gene mapping techniques, can I predict the location of a gene based on its composition?

Beth Fawcett

Background

- DNA
- Chromosomes
- Genes
 - Locus
- Proteins
 - Strand +/-
 - Amino acids





Business Case

- Drug Development
- Individualized Medicine
- Advancement in understanding mechanism behind chromosomal based disease processes
- Health applications

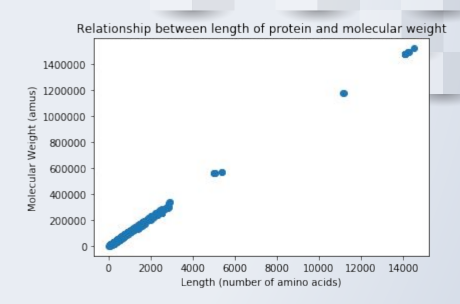
Methodology

- Supervised Learning
 - **Techniques**
- Loci with > 50 occurrences
- □ Chromosome 19 has 48
 - instances of loci with > 50
 - occurrences
- Locus Study
- Strand Study

- Obtain Data
- Scrub Data
- Explore Data
- Modeling
- Interpret

Findings

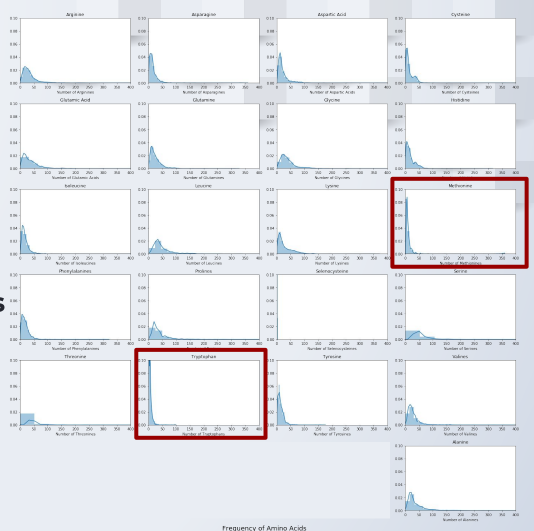
A commonly known fact is that as length of amino acids in protein increases, the molecular weight increases.



Findings

Tryptophan, a heavy protein, was frequently found.

Methenamine, was also found frequently, but it is the start of all proteins.



Findings

Cystine and
 Tryptophan's
 weights are high for
 the distribution.

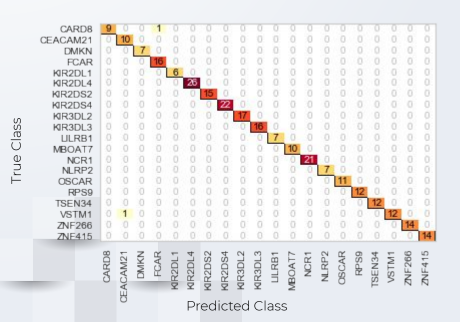


Findings - Random Forest

Locus Study

Test Accuracy: 99%

Strand Study
Test Accuracy: 73%





Future Work

- Grid Search with PCA
- Other types of classifiers: SVM, XGBoost
- Additional chromosomes
- SMOTE (Synthetic Minority Over-sampling Technique) to assist with expanding to all loci in chromosome
- Neural Network
- Build a chromosome from predicted values

Thank you!

For more information:

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