

### CALIFORNIA STATE SCIENCE FAIR **2014 PROJECT SUMMARY**

Name(s)

Min Jean Cho

**Project Number** 

# S1403

#### **Project Title**

## **Applying Bayes' Theorem to DNA Sequence for Identification of Pathogenic Bacteria**

Abstract

**Objectives/Goals** To develop an easy, simple method for identifying microorganisms based on their DNA sequences, Bayes' theorem was applied to DNA sequence analysis. It was hypothesized that the conditional probability of a DNA sequence from an unknown bacterial species being a member of a particular species could be the posterior probability, which could be estimated from prior probability and likelihood function using Bayes' theorem.

#### **Methods/Materials**

To test the hypothesis, 16S rRNA gene sequences of foodborne pathogens (eight bacterial species) were downloaded from NIH's GenBank (45 sequences from each bacterial species, 360 sequences in total) to construct a database. Bayes' theorem was used to estimate the posterior probability of a bacterial specie "Si" given an unknown sequence "Q",  $P(Si|Q) = P(Q|Si) \times P(Si) / P(Q)$ . To determine the likelihood, P(Q|Si), the DNA sequence "Q" was divided into words (k-size DNA sequence fragments), and P(Q|Si) was measured from the average probability of observing the word j from species Si, P(wj|Si). The prior probability, P(Si), and P(Q) were calculated from the database sequences.

#### Results

The size of word (k) affected values of P(Q|Si) and P(Q). The optimum size of word (k) was determined to be 39 nucleotides. All test sequences showed the highest P(Si|Q) values for the species to which they belong, which indicated that the developed method correctly identified the test sequences (accuracy = 100%).

#### **Conclusions/Discussion**

The hypothesized algorithm was proven to work in the experiments carried out with DNA sequences of bacterial species. Dividing the unknown DNA sequence Q into small-size words (wj) was especially important to determine P(Q|Si) and P(Q). An unknown sequence should be classified into the species with the highest P(Si|Q) value (rank-based identification), which indicated the most probable species among the species included in the database.

#### **Summary Statement**

Bayes; # theorem was applied to DNA sequence analysis in order to determine the conditional probability that a DNA sequence from an unknown species belongs to a particular species.

#### **Help Received**

My father assisted me with researching, and my mother read and edited my writing. I would also like to thank Ms. Julia Newman for her valuable advice and suggestions.