

CALIFORNIA STATE SCIENCE FAIR 2011 PROJECT SUMMARY

Name(s) Christian Liu	Project Number S1421
Project Title Effect of Gap Penalty on Global Alignm	ent Accuracy
Objectives/Goals Abstract Objectives/Goals The Needleman-Wunsch global alignment algorithm uses a mismatch, and gap penalty to compare two large sequences of the gap penalty on sequence alignments when the match Methods/Materials Sequence-modifying, global alignment, and alignment score compare alignments. The sequence-modifying program cor 50, 200, 350, 500, and 600 characters in an original 626-ch were then globally aligned to the original DNA sequence u Finally the scoring program judged the accuracy of the align Results At a gap penalty of 0, there were gaps throughout the align gap penalty of -5, the algorithm avoided gaps for all alignmet Conclusions/Discussion At very small gap penalties, scores are low because the algorithm quickly stops aligning as penalties values still large enough to discourage unnecessary gaps.	a scoring system consisting of a match, s of data. This research aims to examine effects a score is 1 and the mismatch score is 0. ring programs were written in Java and used to ntrolled how similar sequences were by altering paracter long sequence. These altered sequences using 0, -1, -2, -3, and -5 as gap penalties. gnment. ments and alignment scores were low, but at a nents. The ideal gap penalty was -2. orithm doesn't penalize for using many gaps. grow, so the most effective penalties are small
Summary Statement Five gap penalties were used to align sequences against an on the accuracy of the global alignment algorithm.	original and model the effect of gap penalties
Help Received My advisor Dr. James Li reviewed the programs I wrote ar	nd helped me correct errors.