

PatternHunter: Faster And More Sensitive Homology Search

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Abstract

Genomics and proteomics studies routinely depend on homology searches based on the strategy of finding short seed matches which are then extended. The exploding genomic data growth presents a dilemma for DNA homology search techniques: increasing seed size decreases sensitivity whereas decreasing seed size slows down computation.

We present a new homology search algorithm “PatternHunter” that uses a novel seed model for increased sensitivity and new hit-processing techniques for significantly increased speed. At Blast levels of sensitivity, PatternHunter is able to find homologies between sequences as large as human chromosomes, in mere hours on a desktop.

PatternHunter is available at <http://www.bioinformaticssolutions.com>. It runs on all platforms that support Java.