Take Home Final

Topic 1

Prior to 1970, protein and DNA sequence alignment was limited to visual comparison. This was a very tedious process; even proteins with global similarity can possess more regions of dissimilarity (eg. introns) than local similarity. In 1970, Needleman and Wunsch introduced the first computerized method for sequence alignment. Their method was based on the concept of dynamic programming. In dynamic programming, a similarity matrix for two sequences is constructed and computed such that each cell in the matrix is given a quantitative score that reflects the best previous alignment less any penalties assigned that result from making the alignment. The optimal alignment is therefore the highest scoring path through the entire matrix. In their original paper describing dynamic programming (Needleman & Wunsch (1971)), they assigned a value to each cell that was reflective of the number of identical matches that preceded that alignment.

The Needleman-Wunsch (NW) method of alignment is a global evaluation of homology. This method is used to compare two sequences in their entirety in an attempt to determine how similar they are overall. This method works well when the sequences are potentially equivalent (eg. human and mouse beta-globin). In 1981, Smith and Waterman derived a dynamic programming-based algorithm for local sequence alignment (Smith & Waterman (1981)). The Smith-Waterman (SW) local alignment algorithm was slightly different than the NW algorithm: they allowed for negative scores in their matrix such that optimal, high-scoring alignments (independent of size) could be identified amongst regions of dissimilarity. In other words, their algorithm allowed for short spans of similarity rather than global similarity. Today, short regions of similarity are critical for genome comparisons---sequence targeting signals, conserved domains, and protein motifs are all routinely identified using local alignment-based algorithms.

NW and SW algorithms are effective for comparing pairwise alignments and for evaluating global and local similarity, respectively. However, multiple alignments are critical for understanding and classifying genomic information. In multiple sequence alignments, clustering approaches are taken because this type of alignment is not amenable to dynamic programming. When a multiple alignment is constructed, the two most similar sequences are aligned pairwise and subsequent sequences are added on to this initial alignment. The ability to align multiple sequences has been critical to algorithm development today.

As sequence databases began to grow exponentially in the last decade, the need to rapidly search and extract information from these databases grew urgently as well. The first two algorithms designed specifically to address rapid database searches were FASTA (Pearson & Lipman (1988)) and BLAST (Altschul, et al. (1990)).

FASTA was the first program to become rapidly and widely utilized. FASTA first searches the database using several short sub-sequences (words) from the query
sequence to find identical matches; FASTA is a local alignment (SW-based) tool with multiple alignment capabilities. It is a more rapid search tool than SW, making it a more desirable tool for biological researchers. Unfortunately, however, sensitivity is sacrificed for speed in the case of FASTA as well as BLAST.

Following closely behind the advent of FASTA was BLAST. BLAST is an even more rapid alignment algorithm, and consequently, not as sensitive as other time consuming methods. BLAST employs a substitution matrix that is used to quantify the alignment such that each possible residue substitution is given a score reflective of the probability that the alignment could not have occurred by chance alone. The algorithm sought out not only high-scoring segment pairs (HSPs), but also HSPs with optimal neighboring ungapped alignments. In other words, the algorithm sought out the best ungapped local alignment that could not be improved with adjustments to the alignment (extensions or trimming). The search method is a two-step process. First, BLAST scans the database for words that possess a given score, thereby obtaining hits. Then second, BLAST verifies each hit by extending the alignment to neighboring pairs in search of HSPs.

One problem with BLAST is that it cannot generate gapped alignments. This problem has been addressed with Gapped BLAST (Altschul, et al. (1997)). Gapped BLAST is a modern counterpart of BLAST that surpasses its predecessor; it is more sensitive and just as fast. Gapped BLAST differs from the original BLAST in three ways. First, the algorithm establishes a cutoff such that sequences scoring above a given E-value threshold are omitted. Second, to increase the speed of the search, the criterion for extensions was modified from the original BLAST algorithm. Third, the ability to form gapped alignments was added. All of these improvements have allowed BLAST to remain one of the more commonly used search algorithms to date.

Another modern counterpart to BLAST is PSI-BLAST. PSI-BLAST is a method of searching for profiles and motifs that utilizes Gapped BLAST in its iteration scheme (Altschul, et al. (1997)). Compared with other BLAST derivatives, PSI-BLAST is slightly more sensitive. In PSI-BLAST, a profile group is created by aligning sequences for the database with the query sequence. Each time a new sequence is aligned to the profile group, the process of iteration is repeated, thereby generating a new multiple sequence alignment. The “new profile” is then reiterated and this process continues until all of the related sequences below a given E-value threshold are extracted from the database (Figure 1). With this algorithm, protein families are easily assembled; for this reason, this application is used when the query sequence contains a position-specific conserved domain. Consequently, PSI-BLAST is an essential tool for uncovering protein relationships.

Sequence alignment methods have progressed greatly in the last 30 years, each new improvement building on previous algorithms and meeting the growing demands of bioinformatics. As databases continue to grow, future advances should focus on creating algorithms that increase the speed with which alignments are carried out, without sacrificing sensitivity in the process.
FIGURE 1
PSI-BLAST is a modern counterpart of BLAST that incorporates many sequence alignment algorithms into its iteration scheme.
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Topic 2

In light of the recent sequencing of the unicellular biflagellar green alga, T. samaroo, I propose a plan for functionally identifying and characterizing flagellar proteins in this organism. Given that my interests lie in studying intraflagellar transport (IFT), I propose the following steps for the initial characterization of proteins localized to this organelle.

AIM 1: To create an expression profile using microarrays to compare mRNA expression levels in T. samaroo cells possessing flagella and those lacking flagella

T. samaroo is a biflagellar unicellular organism. Its cellular body has polarity, such that its apical and basolateral surfaces are distinctly different. The primary difference between the two membrane surfaces is the presence of two flagella on its apical membrane. In culture, the flagella can be detached by exposing cells to low pH medium and, subsequently, the flagella can be purified from the culture. Given this fact, I propose to prepare a DNA chip consisting of cDNAs encoding the 5,000 ORFs of T. samaroo. I will then isolate and fluorescently label mRNA from cells possessing flagella (wild-type) and those in which the flagella have been removed. The labeled mRNAs will then be hybridized to the DNA chip and subsequently scanned. The data will then be analyzed and the expression levels normalized and scaled. From this experiment I hope to identify ORFs that are significantly overexpressed in the cells possessing flagella. These ORFs could encode proteins that are localized to the flagella and/or are involved in IFT.

AIM 2: To identify T. samaroo intraflagellar transport homologs in C. reinhardtii using PSI-BLAST and BLAST

Given that T. samaroo is a green algae that is closely related to C. reinhardtii, there is high probability that the two organisms share regions of highly conserved sequence. Comparison with this organism whose genome has also been sequenced will be useful in identifying homologs. I propose to first utilize a motif-finding program to identify motifs in flagellar ORFs that are highly conserved in IFT proteins: 1) dynein-binding domain, 2) kinesin-binding domain, 3) dynein ATPase domain, and 4) microtubule-binding domain. Once I have identified proteins containing any one of these motifs, I will use multiple alignment algorithms (BLAST, PSI-BLAST, and Smith-Waterman) to align these sequences with known C. reinhardtii proteins. Through this
analysis I will identify proteins in *T. samaroo* that are homologous to proteins in *C. reinhardtii*. Furthermore, this analysis will allow me to further narrow the pool of flagellar proteins down to IFT-related proteins. This information will allow me to further characterize the IFT mechanics of *T. samaroo* by comparing these proteins to known components of intraflagellar transport.

**AIM 3: To tandem-affinity purify (TAP) *T. samaroo* IFT proteins for large-scale isolation and identification of their protein complex formation using MALDI-TOF; to create a higher-order organizational map for those proteins involved in IFT**

Given that from Aims 1 and 2, potential IFT proteins have been identified, a higher-order map of protein networks may serve as a tool for further identifying protein function. For this experiment, I propose to PCR-generate gene-specific TAP cassettes. These cassettes will then be inserted into their specific genes through homologous recombination. For each successful transformation, a TAP-tag fusion protein will be affinity purified in a two-step process and subject to SDS-PAGE. Upon separation of the proteins, each band will be digested with trypsin and subject to MALDI-TOF for sequence identification (Gavin, et al. (2002)). Given that each TAP-fusion protein should associate with native complexes in vivo, each affinity purification will not only pull-down the fusion protein, but other complexed proteins as well. By using each IFT protein as a separate entry-point into a complex, a higher-order map of protein networks could be constructed. In comparing this *T. samaroo* protein network map with one of *C. reinhardtii*, more homologs can be identified, and therefore more function assigned to novel proteins.

**AIM 4: To use Bayesian networks to predict additional protein-protein interactions**

Bayesian networks have been previously shown to be strong predictors of protein interactions (Jansen, et al. (2003)). Given the known sequence information for *C. reinhardtii*, this model will be useful in expanding and confirming a flagellar protein network map. A Bayesian network prediction model may be further useful in establishing protein function through interaction and flagellar localization to proteins that may have been overlooked thus far.
FIGURE 2
Scheme for the functional characterization of *T. samaroo* IFT proteins
REFERENCES