

## Recent trends in Remote homology detection: an Indian Medley

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### Abstract:

The development of remote homology detection methods is a challenging area in Bioinformatics. Sequence analysis-based approaches that address this problem have employed the use of profiles, templates and Hidden Markov Models (HMMs). These methods often face limitations due to poor sequence similarities and non-uniform sequence dispersion in protein sequence space. Search procedures are often asymmetrical due to over or under-representation of some protein families and outliers often remain undetected. Intermediate sequences that share high similarities with more than one protein can help overcome such problems. Methods such as MulPSSM and Cascade PSI-BLAST that employ intermediate sequences achieve better coverage of members in searches. Others employ peptide modules or conserved patterns of motifs or residues and are effective in overcoming dependencies on high sequence similarity to establish homology by using conserved patterns in searches. We review some of these recent methods developed in India in the recent past.

**Keywords:** Sequence analysis; Remote homology detection; PSI-BLAST; Protein Evolution

### Background:

The last few decades have seen spectacular developments in the growth of protein sequence and structure data and in tools to analyse them. Large-scale experimental determination of the biochemical roles of proteins is a phenomenal exercise. Bioinformatics shows exciting promise in the development of methods that allow quick resolution of newly sequenced proteins to their closest experimentally verified relatives. Sequence search procedures such as PSI-BLAST [1], can detect protein relationships effectively when sequence similarities are high. However, when sequence similarities become poor (20-30%), such detection becomes a non-trivial task.

Proteins dissimilar in sequence can adopt similar structure, perform similar functions and also be homologous. This is exemplified in structural databases, which classify proteins with high similarity in sequence, structure and function into families and group families of similar structure and function into superfamilies. For superfamily members, which show poor sequence similarity, it is difficult to determine evolutionary relationship in the absence of structure. Such proteins, related despite low sequence similarity, lie in the 'twilight zone' and are termed 'remote homologues'.

Structure-based approaches can detect such relationships since protein structures are less perturbed by sequence changes. Some approaches consider overall structural similarity in defining relatedness while others such as Bhaduri *et al.*, [2] have shown that conserved spatial interactions in a protein superfamily are excellent constraints in the identification of more members in the superfamily.

### Description:

#### Sequence analysis based approaches for remote homology detection:

Methods like PSI-BLAST build profiles iteratively during searches in sequence databases. [1] The use of 'intermediate protein sequences' that share sequence features of more than one protein is quite effective in detecting distant protein similarities. [3] As seen in Figure 1, such sequences populate sequence space and relate proteins, traditionally difficult to relate, due to poor sequence similarities. 'Intermediate sequences' that share high similarities with more than one protein, if detected and employed in profile generation can improve effectiveness of sequence analysis-based approaches. Sandhya *et al.*, [4] discuss an application of such sequences in PSI-BLAST searches in fold-specific databases to detect relationships not evident through simple sequence searches.

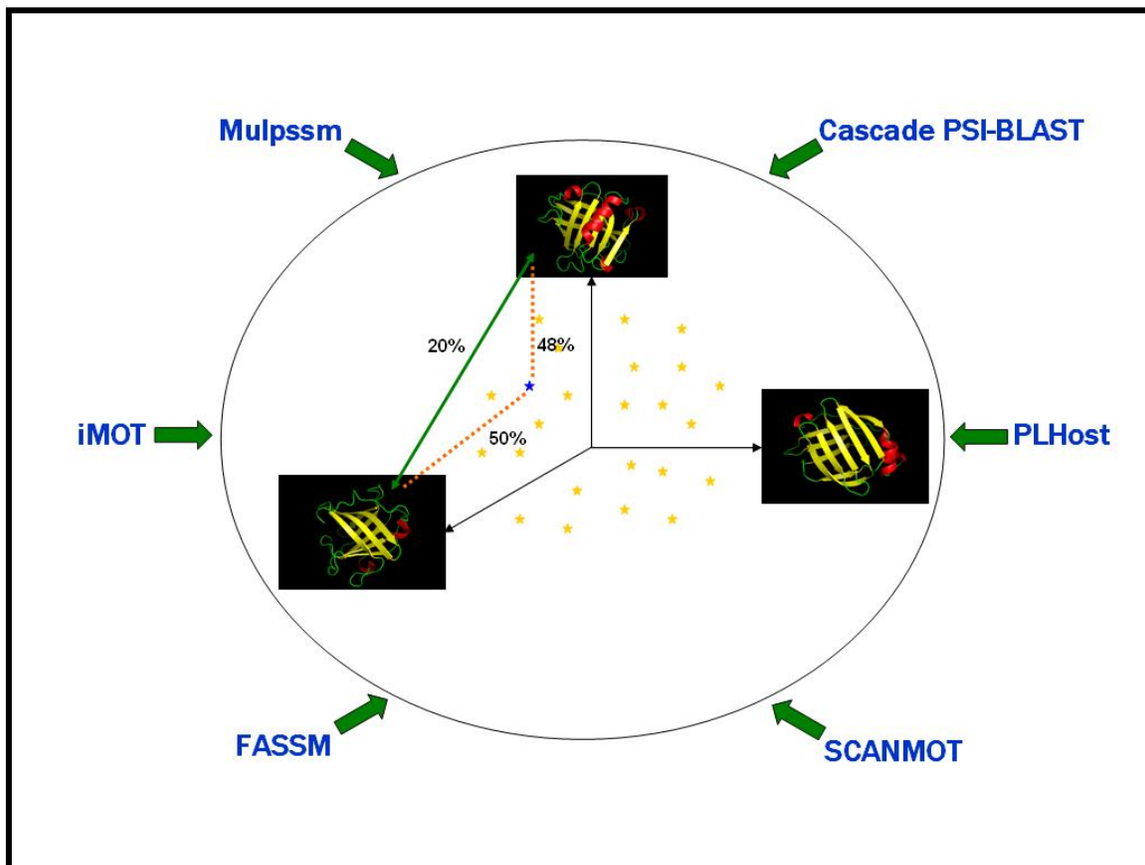
Several exciting and interesting efforts in remote homology search methods have been made in the Indian sub-continent in the last few years. These methods (Figure 1) show promise and effectively detect such deep relationships in proteins. In this review, we highlight salient features of these approaches.

#### Cascade PSI-BLAST: Hops through intermediates detect remote homologues:

Sandhya *et al.*, [5] have recently reported the development of a method called Cascade PSI-BLAST that propagates PSI-BLAST searches in a non-directed manner to detect distant similarities between proteins. In this method, that extensively employs intermediate sequences, a PSI-BLAST search termed "first generation" is initiated for a query in a database. Hits detected are allowed to propagate independent searches in the same database to detect more

new hits ("second generation" search etc.). Typically, the authors recommend up to three generations of search for better coverage and detection of relationships. An assessment of the approach on the detection of existing relationships from the PALI database [6] shows that the coverage in detecting relationships in protein families is

improved by 15% and in protein superfamilies by ~35% over traditional use of PSI-BLAST. This method is being made available for use in public domain through a web server (<http://crick.mbu.iisc.ernet.in/~CASCADE>) (manuscript communicated).



**Figure 1:** A superfamily of proteins whose members share poor sequence similarity (<20%). 'Intermediate sequences' (in yellow) populate protein space and owing to their high similarities with more than one protein (40-50%) can effectively detect such remote homologues. Methods developed recently in India address the problem of remote homology detection effectively with patterns/ intermediate sequences.

**MulPSSM: A database of multiple family profiles corresponding to a constant alignment:**

Searches against sequence databases using PSSMs or profiles are effective in identifying distant relationships compared to searches involving pair wise sequence alignments. [1] The effectiveness of any profile-based search depends on the quality and diversity of sequences in the multiple sequence alignment used in profile generation.

Anand *et al.*, [7] demonstrate that generating multiple family profiles enables the reliable detection of distant relationships in protein family and superfamily. In order to assess the effectiveness of multiple family profiles, they have generated a database of multiple and single-family

profiles and profile-HMMs for all structural families from integrated sequence-structure database from PALI. [6] Searches against these three databases were assessed and compared in terms of Sensitivity, Specificity and Error rates as functions of E-values. All three parameters support that generating multiple family profiles improves detection of more superfamily relationships. The CPU time for searches in a multiple family profile database is also economical. This database is available at <http://crick.mbu.iisc.ernet.in/~mulpssm>. [8]

**FASSM: Enhanced Function Association in whole genome analysis using Sequence and Structural Motifs:**

Motif-based approaches use patterns or features unique to a class of proteins to detect remote similarities. Gaurav *et al.*, [9] have developed a neural network-based method called 'FASSM' that employs the positional conservation of sequences and ordering and positions of family specific inter-motif lengths, in a scoring scheme to guide family associations. Their assessments of the approach in detecting over 30 superfamily level relationships indicate an improved performance over other methods. An exciting application of this method is its ability to detect relationships in proteins despite discontinuous domains and circular permutations.

**SCANMOT: A simultaneous scan of multiple sequence motifs to search for similar sequences:**

Motif-based search methods that are available in the public domain often scan for a single pattern at a time. Chakrabarti *et al.*, [10] have developed a multiple motif-based search engine 'SCANMOT' that combines multiple-pattern searching and a search for hits with statistically significant sequence similarity to detect relationships between proteins sharing a common set of motifs. This method is available through a web-based server interface (<http://www.ncbs.res.in/~faculty/mini/scanmot/scanmot.html>).

**PLHOST: Peptide Library based Homology search tool:**

Brahmachari and Dash [11] have developed a tool for whole genome comparisons that can compare several thousand proteins of a genome with proteins of other genomes. The principle belying their method is that proteins use a limited set of peptide modules in various genomes. They show in comparisons of octa-peptide libraries of individual organisms that some invariant peptides are present in related proteins from multiple organisms. Such invariant peptides of variable lengths distributed over a large number of functionally vital proteins can serve as signatures in functional annotation of hypothetical ORFs. They suggest that such a peptide-library based approach can overcome problems associated with low sequence similarity and in identifying polypeptide stretches unique to an organism.

**Conclusion:**

In this review, we have highlighted the recently developed remote homology detection methods from India. These methods apply either intermediate sequences in profile-based searches or motif-based principles to expand protein families and help identify novel evolutionary relationships between proteins. The methods described in this review offer at least a partial solution to the problem. Continued progress in the development of sequence and structure alignment methods will increase the sensitivity of remote homology detection.

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