

# The EMOTIF database

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Received September 1, 2000; Revised and Accepted October 30, 2000

## ABSTRACT

**The EMOTIF database is a collection of more than 170 000 highly specific and sensitive protein sequence motifs representing conserved biochemical properties and biological functions. These protein motifs are derived from 7697 sequence alignments in the BLOCKS+ database (released on June 23, 2000) and all 8244 protein sequence alignments in the PRINTS database (version 27.0) using the EMOTIF-MAKER algorithm developed by Nevill-Manning *et al.* (Nevill-Manning, C.G., Wu, T.D. and Brutlag, D.L. (1998) *Proc. Natl Acad. Sci. USA*, 95, 5865–5871; Nevill-Manning, C.G., Sethi, K.S., Wu, T.D. and Brutlag, D.L. (1997) *ISMB-97*, 5, 202–209). Since the amino acids and the groups of amino acids in these sequence motifs represent critical positions conserved in evolution, search algorithms employing the EMOTIF patterns can identify and classify more widely divergent sequences than methods based on global sequence similarity. The EMOTIF protein pattern database is available at <http://motif.stanford.edu/emotif/>.**

## INTRODUCTION

Many newly discovered proteins do not share significant global sequence similarity to any known protein. These proteins, however, may contain modular domains that confer distinct structures and functions conserved throughout evolution. The conserved modules usually possess characteristic sequence elements, or sequence motifs, that only proteins containing the domains share. Many sequence analysis tools rely on discrete sequence motifs to classify newly discovered proteins into the proper families. The methods used to generate the discrete sequence motifs, therefore, play a vital role in determining the specificity and the sensitivity of the classification performed by these tools.

Most motif generation algorithms focus on finding a single 'best' motif for a given alignment of the homologous domain sequences from various species. These methods usually attempt to maximize sensitivity at the expense of specificity. Sequence analysis tools that use these specificity-compromised motifs are prone to classifying newly discovered proteins into the wrong families. For genomic-scale automated sequence classification to be effective, the expected number of misclassifications made by the discrete motifs must be minimized.

## DATABASE GENERATION AND FORMAT

The *e*MOTIF-MAKER program, previously known as EMOTIF, is a systematic method for determining regular expression patterns, or discrete sequence motifs, from aligned sets of protein sequences. Unlike other methods that allow only one motif to be reported for a given protein sequence alignment, *e*MOTIF-MAKER permits the methodical enumeration of multiple motifs for the same alignment such that motifs are generated from the complete alignment as well as subsets of that alignment. These multiple sequence alignment subsets usually represent subfamilies within the superfamily represented by the full alignment. Discrete sequence motifs constructed from these subsets can, therefore, classify novel sequences with higher specificity than the position-specific scoring matrices derived from entire multiple sequence alignments (1–3).

The *e*MOTIF-MAKER program was applied to 7697 non-PRINTS protein sequence alignments from the BLOCKS+ database released on June 10, 2000, and all 8244 protein sequence alignments from the PRINTS database, version 27.0 (1,2). The BLOCKS+ database and the PRINTS database both have an extensive collection of ungapped multiple sequence alignments that cover various protein domains known to be biologically relevant. A total of 170 294 regular expression patterns were generated from 15 863 ungapped multiple sequence alignments. Since these discrete motifs come from subsets of the alignments as well as full alignments, they can complement the BLOCKS+ and the PRINTS position-specific scoring matrices by assigning novel sequences to potential subfamilies within the known superfamilies (1–3). The details of the *e*MOTIF-MAKER algorithm have already been described by Nevill-Manning *et al.* (4,5). These highly specific and sensitive regular expression patterns, which will be referred to hereafter as *e*MOTIFS, can support both genome-scale sequence classification as well as single protein analysis.

Since the *e*MOTIF database is derived from ungapped multiple sequence alignments from the BLOCKS+ database and the PRINTS database, the *e*MOTIF database will be updated following each release of the BLOCKS+ database and the PRINTS database. The database of *e*MOTIFS exists as a single ASCII text file. Each line in the text file corresponds to a unique *e*MOTIF record. The tab-delimited fields in an *e*MOTIF record are, from left to right, the expected false positive frequency of the *e*MOTIF, the accession number of the *e*MOTIF's parent alignment, the descriptive name of that alignment, the regular expression pattern of the *e*MOTIF, and the sensitivity of the *e*MOTIF. The *e*MOTIFS have been sorted by their expected false positive frequencies, such that the *e*MOTIF with the

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smallest expected false positive frequency resides at the beginning of the file.

The expected false positive frequency of an *e*MOTIF, as summarized by equation 1, is the probability that the *e*MOTIF will match a sequence of the same length by chance.

$$\text{expected false positive frequency (motif)} = \prod_{i=1}^{\text{motif length}} \left\{ \sum_{\text{amino acid} \in \text{group}} p(\text{amino acid}) \right\} \quad \mathbf{1}$$

where group represents the set of amino acids allowed at position *i* of the *e*MOTIF. This calculation assumes that the probability of finding a particular amino acid at a given position in a sequence, or *p*(amino acid), is equivalent to the distribution of that amino acid in the SWISS-PROT database and is independent of all other positions in the same sequence. By minimizing the expected false positive frequency of an *e*MOTIF, the *e*MOTIF-MAKER algorithm maximizes the specificity of that *e*MOTIF.

The sensitivity of an *e*MOTIF, defined as the percentage of sequences the *e*MOTIF can match from its parent multiple sequence alignment, measures the *e*MOTIF's ability to detect the presence of a particular domain whose conserved sequence elements have been captured by the *e*MOTIF. By maximizing an *e*MOTIF's coverage of its parent multiple sequence alignment, the *e*MOTIF-MAKER algorithm maximizes the sensitivity of that *e*MOTIF.

## SEARCH SOFTWARE

The *e*MOTIF-SEARCH program, previously known as IDENTIFY, provides a rapid and efficient search interface to the database of *e*MOTIFS. Given a protein sequence, *e*MOTIF-SEARCH will identify significant matches between subsequences within the query and the regular expression patterns from the *e*MOTIF database. Significance is determined by the number of false positive *e*MOTIF matches the user expects for a given protein sequence. As defined by equation 2, the expected number of false positive matches, or the expectation, for an *e*MOTIF-SEARCH query is the number of matches by chance the user is willing to accept, given the length of the query sequence and the content of the *e*MOTIF database.

$$\text{expectation} = [(\text{query length}) + 1] \sum_{i=1}^n f_i - \sum_{i=1}^n [(\text{motif length})_i \times f_i],$$

*f* = expected false positive frequency (motif) **2**

where *n* is the number of *e*MOTIFS the query sequence has encountered so far. The expectation for a query rises as the number of comparisons the *e*MOTIF-SEARCH program performs increases. The *e*MOTIFS have been sorted by their expected false positive frequencies, such that the more specific *e*MOTIFS, or the *e*MOTIFS with the smaller expected false positive frequencies, are compared against a query sequence before the less specific *e*MOTIFS, or the *e*MOTIFS with the larger expected false positive frequencies. As a result of this sorting, the *e*MOTIF-SEARCH program guarantees that matches with the more specific *e*MOTIFS are reported first.

The online version of *e*MOTIF-SEARCH provides an easy-to-use interface for single-protein analysis. After *e*MOTIF-SEARCH receives a cutoff expectation value and a query protein sequence over the network, it will return *e*MOTIF matches as long as the expectation for the query remains below the user-specified cutoff. A sample output of *e*MOTIF-SEARCH online is shown in Figure 1.

## APPLICATION

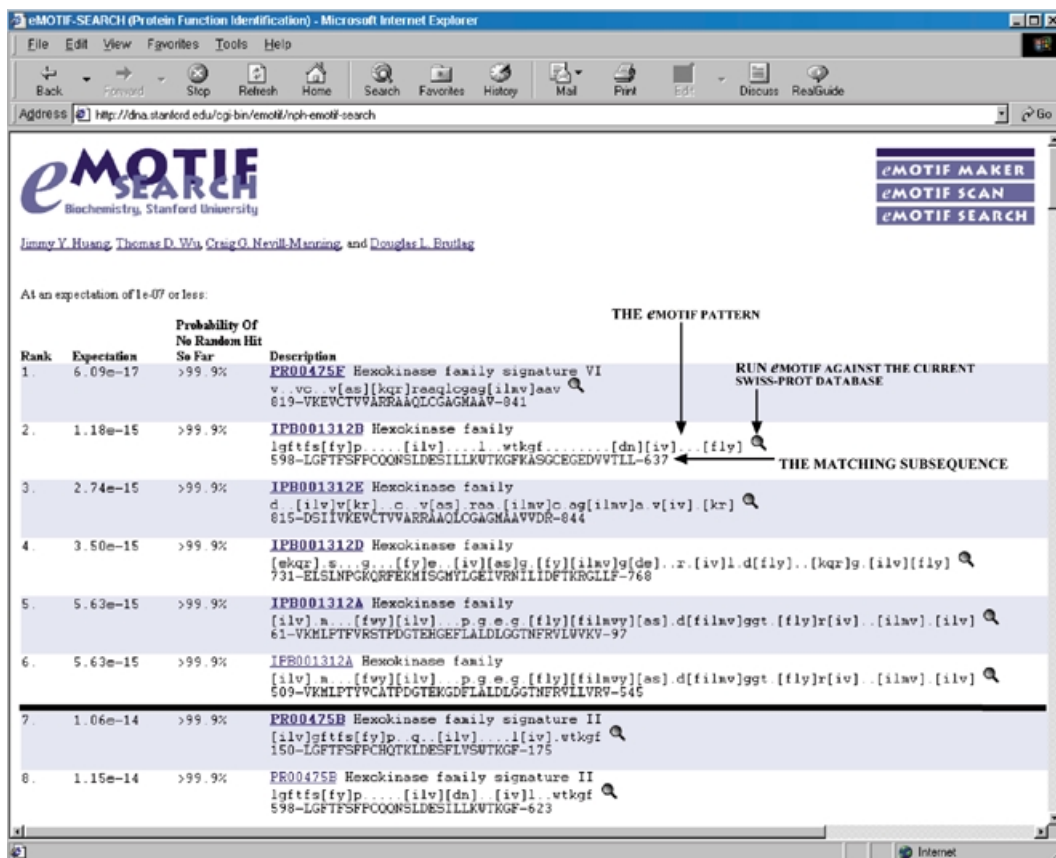
To facilitate efficient automated genomic-scale protein classification, *e*MOTIF-SEARCH and all the necessary preprocessing programs have been repackaged into the *e*MOTIF Batch Analysis Suite, or *e*BAS. The core of this software bundle is a Makefile script that has streamlined and automated the steps involved in running the database of *e*MOTIFS against multiple ORFs, genomes, and proteomes. *e*BAS accepts FASTA-formatted ORF sequences, FASTA-formatted genomic DNA sequences, and FASTA-formatted protein sequences as input. *e*BAS assumes the ORFs are reported in the right frame; therefore, *e*BAS will only translate one frame of the ORF sequences. Genomic DNA sequences, on the other hand, may or may not code for proteins; *e*BAS, therefore, will translate all six frames of the genomic DNA sequences. The input protein and translated sequences are then piped to *e*MOTIF-SEARCH for automated classification. *e*BAS has been applied to coding sequences derived from the *Arabidopsis thaliana* genomic sequences; the coding sequence derivation was performed by C. J. Palm at the Stanford DNA Sequencing and Technology Center using the Genscan program (6). The results from *e*BAS may be accessed at <http://baggage.stanford.edu/group/arabprotein/>.

## CONCLUSION

The extensive collection of ungapped multiple sequence alignments in the BLOCKS+ database and the PRINTS database covers various protein domains known to be biologically relevant. From these multiple sequence alignments discrete patterns like *e*MOTIFS or position-specific scoring matrices like those available through the BLOCKS+ database and the PRINTS database may both be generated (1–3). *e*MOTIFS have the advantage over the position-specific scoring matrices derived from the entire multiple sequence alignments in that the *e*MOTIFS are able to classify novel proteins into protein subfamilies. In addition *e*MOTIFS focus on just those highly conserved residues; this focus greatly improves the signal-to-noise ratios of the *e*MOTIFS and enhances the specificity of the automated sequence classification. The *e*MOTIF database, therefore, can complement the BLOCKS+ and the PRINTS position-specific scoring matrices in identifying novel protein sequences.

## AVAILABILITY

The *e*MOTIF database is available at <http://motif.stanford.edu/emotif/> for interactive use. Local installation of the *e*MOTIF database and the associated software is also available upon request. Not-for-profit institutions may contact Douglas L. Brutlag ([brutlag@stanford.edu](mailto:brutlag@stanford.edu)) for a royalty-free license; for-profit institutions should contact Richard Scholes ([rich.scholes@stanford.edu](mailto:rich.scholes@stanford.edu)).



**Figure 1.** The protein sequence of type II human hexokinase (P52789) was submitted to eMOTIF-SEARCH online. The cutoff expectation for the query was set at  $1e^{-7}$ . The matches were ranked in order, so that matches with the more specific eMOTIFS were reported first. Along with each hit, eMOTIF-SEARCH calculates the expectation for the query based on the fraction of the eMOTIF database it has already run against the input sequence. In addition eMOTIF-SEARCH determines the probability that the query sequence has not matched a single eMOTIF from the ones processed so far by chance. The reporting of these two values allows the user to rerun the search with a lower value of the cutoff expectation for a given query. The black line between the sixth and the seventh eMOTIF match is a sample readjustment performed on this particular sample query. The cutoff expectation was lowered from  $1e^{-7}$  to  $1e^{-15}$ . Only the first six hits were considered significant by the new, more stringent cutoff.

## ACKNOWLEDGEMENTS

This work was supported by a grant from the National Genome Research Institute R01 HG02235-07. J.Y.H. was supported by the Cell and Molecular Biology training grant 2T32GM07276-26 from the National Institutes of Health.

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