**mmsearch: a motif arrangement language and search program**

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**ABSTRACT**

**Summary:** This paper presents a language for describing arrangements of motifs in biological sequences, and a program that uses the language to find the arrangements in motif match databases. The program does not by itself search for the constituent motifs, and is thus independent of how they are detected, which allows it to use motif match data of various origins.

**Availability:** The program can be tested online at http://hits.isb-sib.ch and the distribution is available from ftp://ftp.isrec.isb-sib.ch/pub/software/unix/mmsearch-1.0.tar.gz

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**Supplementary information:** The full documentation about mmsearch is available from http://hits.isb-sib.ch/~tjunier/mmsearch/doc.

**INTRODUCTION**

In biological sequences, the joint occurrence of several conserved motifs is often more informative than the presence of a single one, and the arrangement of the motifs along the sequence can carry even more information as to the sequence’s function. One way of modelling arrangements is to build a descriptor for the whole region. This is analogous to the concatenation of individual profiles for the constituent motifs, with the intervening regions modelled by states with higher entropy and insertion probability. This approach is used by Meta-MEME (Grundy et al., 1997), and a similar one by FingerPRINTSscan (Scordis et al., 1999).

The mmsearch program differs from these in two ways. First, it does not look for the motifs themselves, but only for arrangements, which the user specifies at runtime. The position of motifs along sequences (or match data) is supplied to mmsearch in a tab-separated format such as GFF (Durbin, 1997). The advantage of this approach is that it does not depend on a particular motif search algorithm, and that motif match data of diverse origin (including, for example, database annotations) can be freely intermixed. Secondly, and more importantly, the technique can handle motif overlaps and inclusions, which are hard or impossible to handle with HMMs, profiles or similar techniques (see Section Examples).

**EXAMPLES**

**Simple arrangements**

Most cytokine receptors share two conserved regions: a series of Cys residues near the N-terminus, and a W-S-x-W-S pattern near the membrane. Furthermore, all such receptors feature a signal peptide, and most are membrane-bound (Figure 1a). The two conserved regions are modeled by PROSITE (Hofmann et al., 1999) patterns PS00241 and PS00340. These have 20 and 53 false positives in SWISS-PROT release 38, respectively. (see PROSITE documentation entry PDOC00214.) These figures can be lowered by running pattern search programs and predictors for signal peptides and transmembrane domains, and by conserving only the proteins which match the following arrangement:

\[
\text{SIGNAL} = \text{PS00241} = \text{PS00340} = \text{TRANSMEM}. \tag{1}
\]

The patterns also miss a number of true matches (false negatives). This can be solved by rewriting the patterns in a less restrictive manner, the arrangement condition keeping out most false positives.

**Feature ends**

The thioredoxin domain is shared by a variety of proteins, some of which have an active site consisting of a disulfide bridge (Figure 1b). If we wish to select only those proteins that contain a thioredoxin domain with an active site, we could use an expression of the form

\[
<\text{THIORED} = \text{SS} = \text{THIORED}> \tag{2}
\]

where ‘<’ and ‘>’ mean the start and end of a motif, respectively; and SS stands for a protein region flanked by cysteine residues forming a disulfide bridge. These are handy for dealing with overlaps and inclusions.

**Equivalence**

There may be more than one predictor for the same motif, for example a PROSITE profile and a Pfam HMM (Bateman et al., 2000) for the same domain. The different predictors rarely agree exactly: they may start or stop a...
Fig. 1. Examples of domain arrangements that can be located with a metamotif search.

few residues apart, or either one may be absent altogether. Figure 1c presents six different arrangements which in fact represent the same biological fact, i.e. the presence of a protein kinase domain. This is handled by the 'equivalence' operator, in which the possible motifs are separated by pipes ('|') and included in square brackets ('[ ]'):

\[ \text{PFAM_PK|PROSITE_PK} \]  (3)

which reads 'a PROSITE PK, or a Pfam PK, or both; and in that case, arrangement is irrelevant'.

Other features

mmsearch’s syntax include controlling the distance between motifs, repeating part of a metamotif, alternating between possible arrangements, and anchoring the metamotif at the beginning or end of the sequence. This is explained in detail, along with a complete grammar, in the documentation (see Section Abstract).

THE SEARCH PROGRAM

The program generates a state automaton by parsing the user-supplied expression. The match data are converted into a string containing motif names and positions. This string is fed to the automaton, but unlike in pure pattern matching, some state transitions are governed not only by the next input symbol, but by the values of numbers or names parsed from the input string.

The mmsearch program was written as a stand-alone, command-line application using the Python language. A Web interface to mmsearch is available on the Hits Web site (URL given in the Section Abstract) where it can be used with PROSITE and Pfam motifs in the SwissProt, TrEMBL, trGEN and trEST databases (Pagni et al., 2001).

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REFERENCES


