

A DNA/protein sequence retrieval system: an Xtpanel application

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Abstract

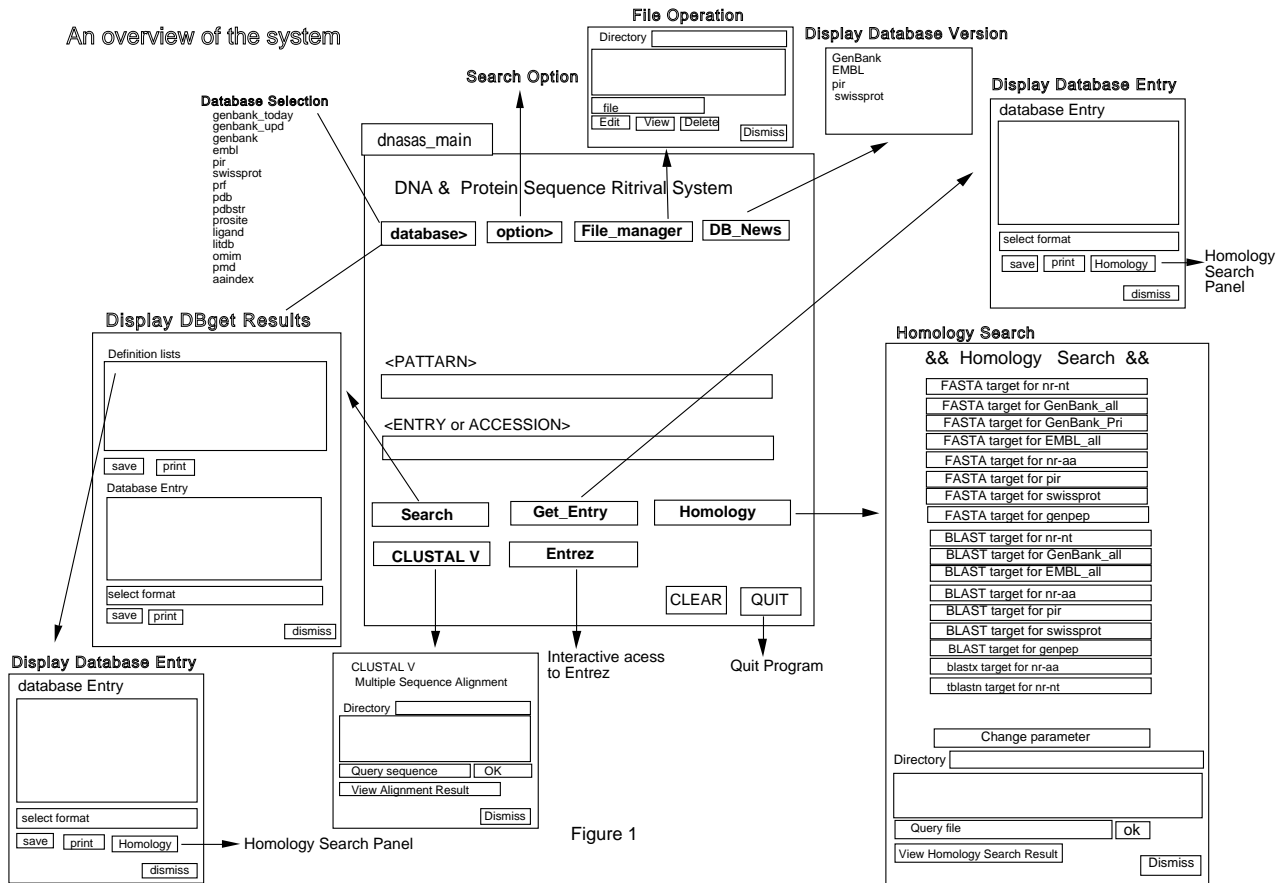
Computational methods are indispensable tools for researchers who determine a new nucleic acid or protein sequence. Such methods are used to obtain information to help identify and annotate the new sequence. Database searching is currently the most popular way to get this information. Using homology- search programs such as FASTA or BLAST, one can search a database for sequences similar to a given query sequence. In addition, the alignment of multiple nucleotide or amino acid sequences is important for drawing phylogenetic trees and for predicting the structure of proteins. The multiple sequence alignment of proteins is also used to discover motifs and biologically important patterns.

We have developed a retrieval system for nucleic acid and/or protein sequences. The system is an Xtpanel application that runs on Sun Workstations with the X-Window system. Xtpanel, which is written by Steve Cole and Dave Nichols of Stanford University, allows one to build interactive X-Window system interfaces without having to do conventional X programming.

It supports FASTA and BLAST for homology searches, bfind and bget for database entry retrievals, and CLUSTAL V for multiple sequence alignments.

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1 Function and graphical interface of the system

The system is easy to use because its functions are tightly coupled with its graphical user interface (see Figure 1 for an overview). The following functions are provided:

1. Keyword Searching
The user can get the contents of entries by specifying an entry name or an accession number. Alternatively, he can obtain the list of entry names whose contents involve a given pattern.
2. Homology Searching
The FASTA and BLAST programs are available.
3. Multiple Sequence Alignment
The CLUSTAL V program is available.
4. Entrez
The user can interactively access Entrez on the system.