

Protein sequence search tool: A web-based interactive search engine

Protein sequence search tool (PSST) is a web-based interactive search tool developed to extract and analyse the protein/nucleic acid sequence and other related information from the database of protein and nucleic acid structures. A search could be performed using all the protein structures or in a selected subset of non-homologous proteins. The basic sequence information is updated every week and hence the results obtained using the search tool are up-to-date at any given time. The package PSST (Version 1.0) is available over the World Wide Web (www) at <http://pranag.physics.iisc.ernet.in/psst/>.

The number of available three-dimensional structures of proteins and nucleic acids is

increasing rapidly. The three-dimensional structures of nearly 11,800 proteins and nucleic acids are currently available in the public domain of the Protein Data Bank (PDB)¹. This database also carries the sequence information of all the known three-dimensional structures of proteins and nucleic acids. Prediction of the three-dimensional structure from the amino acid sequence is one of the central problems to be resolved in structural biology. While trying to understand this problem it is important to recognize structural characteristics of sequence patterns. In several DNA, RNA and protein structures similar sequence repeat units are often found, and it is essential to know the correlation with respect to

three-dimensional structures. Moreover, identifying the occurrence of functionally or structurally characteristic sequence motifs in various protein or nucleic acid structures is useful. Occurrence of a particular sequence pattern in a protein structure² might give insights on the possibility of the associated function and one could fine-tune the three-dimensional structure by means of protein engineering experiments to incorporate the function. Occurrence of sequence repeats in protein structures might suggest symmetries in them³. In order to analyse this large volume of database it is essential to have automated computational search tools such as SRS (Sequence Retrieval System)⁴. Towards this effort, several software packages

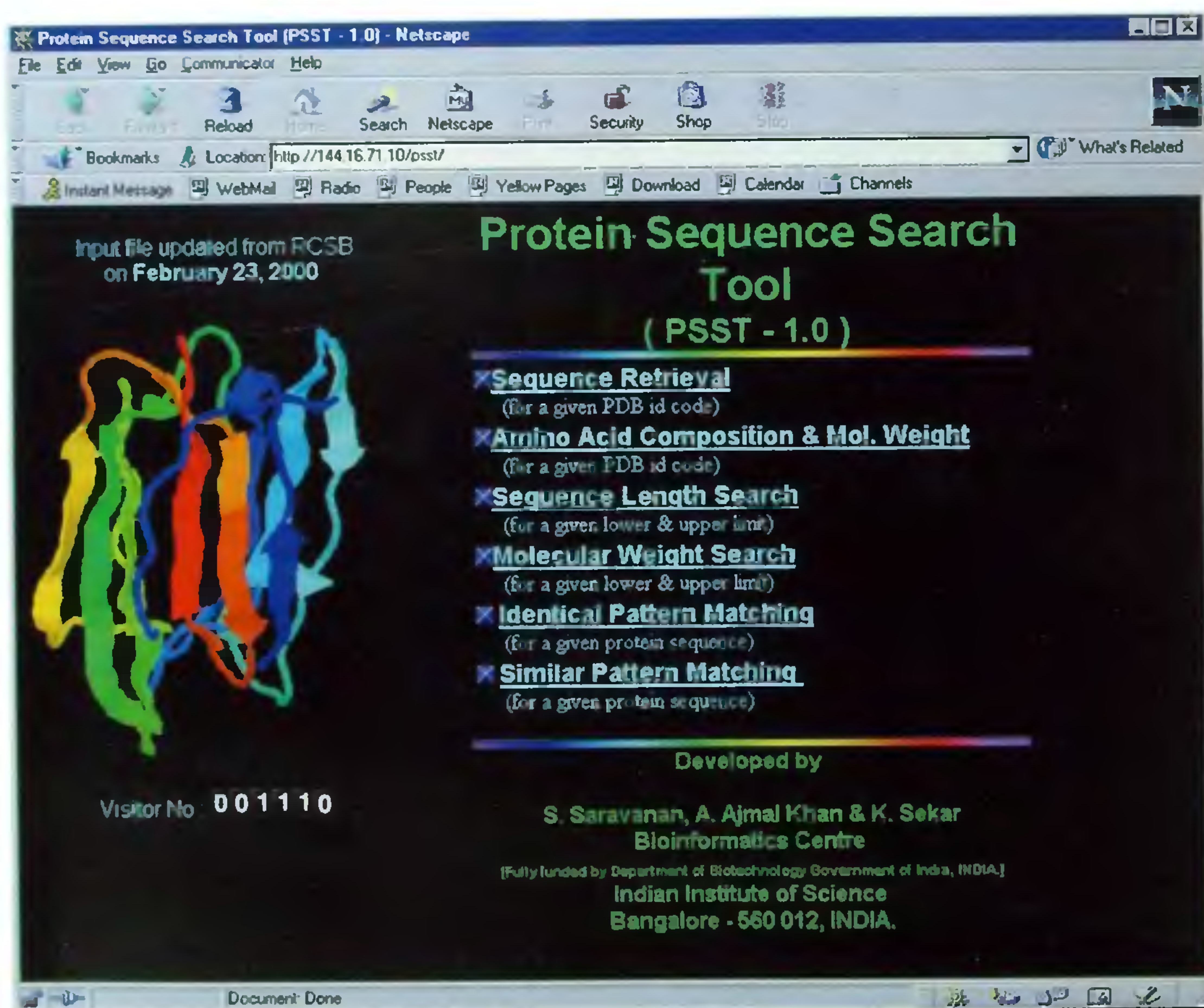


Figure 1. Web page of PSST along with the available search facilities. Clicking on the appropriate search facility activates the search engine. The molecule on the left side is a lectin from the seeds of jackfruit. The crystal structure¹⁰ of this lectin has four subunits, but only one of them is shown. This picture has been created using the graphics software RASMOL¹¹.

have been developed for identifying the similarity in sequence patterns and their compatibilities with the available databases like SWISS-PROT⁵ and PIR (Protein Identification Resource)⁶. To the best of our knowledge, a tool to facilitate search for a well-defined molecular weight and also to perform a search for sequence patterns in a set of non-homologous protein sequences is not yet available. PSST addresses these issues too. The front page of the search tool is shown in Figure 1. PSST offers two different input options for the user to address her/his query. The user has the option of choosing all the protein structures or only using the non-homologous protein structures derived by Hobohm and Sander⁷. In this set of non-homologous protein structures no two proteins have sequence identity of over 30%. At present there are about 1,100 protein chains in the non-homologous category.

The PSST has the following types of search facility: (a) Sequence retrieval in

PIR format, (b) Amino acid composition, (c) sequence length search, (d) molecular weight search, (e) identical pattern matching and (f) similar pattern matching. The basic input for (a) and (b) is in a four-letter PDB-id code that usually appears in research papers dealing with the three dimensional macromolecular crystal structures. The search tool displays the corresponding results in an easy and simple format. For example, the amino acid composition search output displays the name of the amino acid (full name, three-letter code and single-letter code), the number of the particular residue and its contribution towards molecular weight. It also displays the percentage content of all the amino acid residues in the input query protein sequence. At the end of the output it displays the total molecular weight of the protein. The search engine requires the lower and upper limits to pick the protein structures, whose lengths satisfy the two-number input (lower and upper bound) criterion for the sequence

length search. Similarly, the molecular weight search requires the lower and upper limit values of the molecular weight in Daltons. A sample output of the user-desired lower and upper limit criterion is shown in Figure 2. The amino acid sequence for the proteins and nucleotide sequences for DNA/RNA structures is required as the search input for carrying out identical/similarity pattern matching. Moreover, PSST has an option of the user-desired number of mismatches. For example, in the similarity search the default value for the number of mismatches is 3 and it can be modified while submitting the web form at the user end. The identical and similar patterns blink with a different colour on the screen. The results obtained from the searches correspond to the most recent information available in the PDB¹. As mentioned earlier, PSST is available on the www and the users can easily submit their queries. In the trial runs, the results appear in about 1–2 min depending on the network speed.

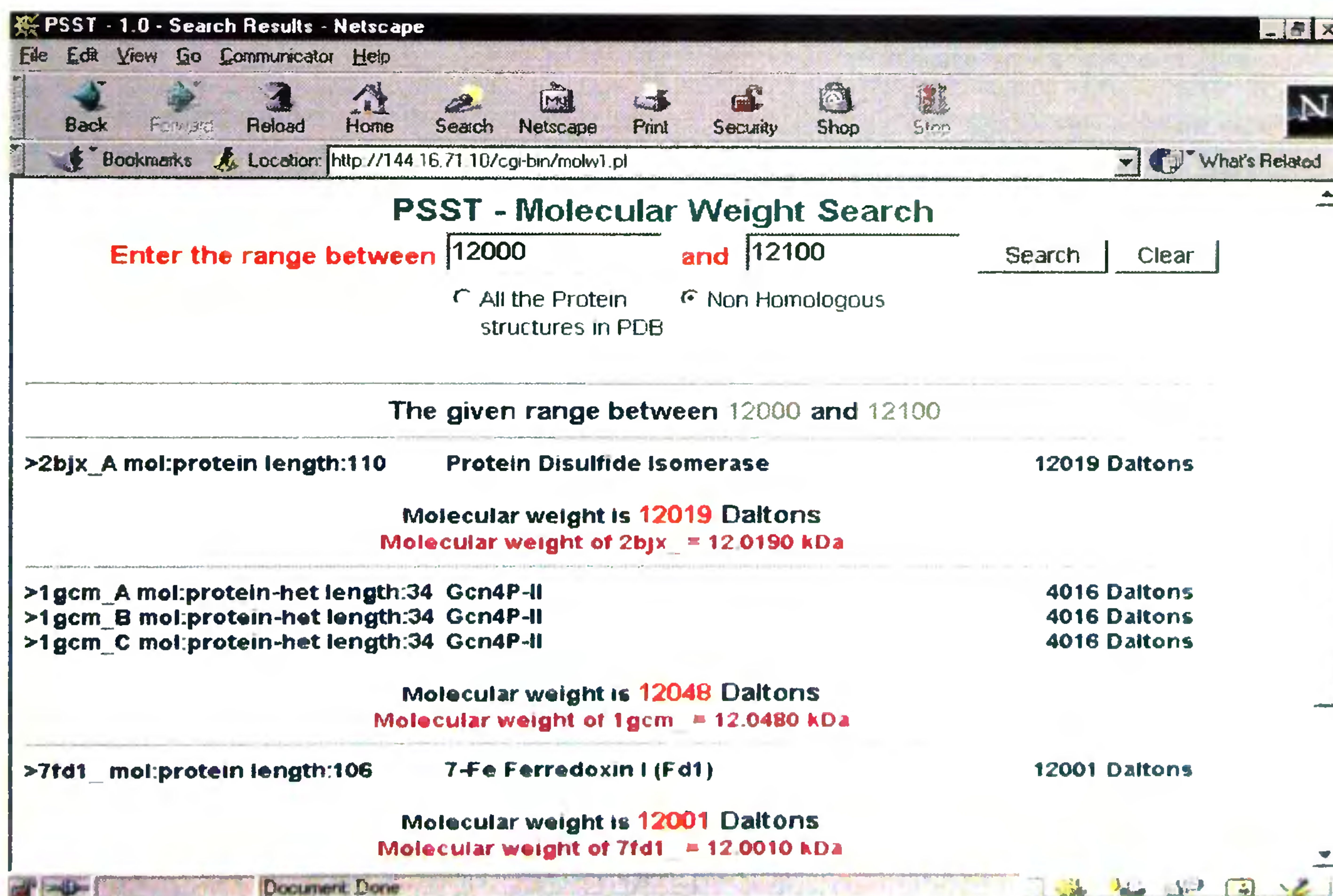


Figure 2. Typical output of molecular weight search using the lower and upper limits of the user-desired query from the non-homologous protein sequence database.

PSST - Identical Pattern Matching

Enter the Pattern :

☒ All the Protein structures in PDB ☐ Non Homologous

☒ Protein ☐ Nucleic Acid

The Search String is CCXXHXXC

Matching position : 43 - 50
 >1a2a_A mol:protein length:122 Phospholipase A2
 NLLQFNKMIK EETGKNAIPF YAFYGCYCGG GGNGKPKDGT DRCCFVHDCC YGRLVNCNTK SDIY
 GYITCGKGTN CEEQICECDR VAAECFRRNL DTYNNGYMFY RDSKCTETSE EC

Matching position : 43 - 50
 >1a2a_B mol:protein length:122 Phospholipase A2
 NLLQFNKMIK EETGKNAIPF YAFYGCYCGG GGNGKPKDGT DRCCFVHDCC YGRLVNCNTK SDIY
 GYITCGKGTN CEEQICECDR VAAECFRRNL DTYNNGYMFY RDSKCTETSE EC

Matching position : 43 - 50
 >1a2a_C mol:protein length:122 Phospholipase A2
 NLLQFNKMIK EETGKNAIPF YAFYGCYCGG GGNGKPKDGT DRCCFVHDCC YGRLVNCNTK SDIY
 GYITCGKGTN CEEQICECDR VAAECFRRNL DTYNNGYMFY RDSKCTETSE EC

Matching position : 43 - 50
 >1a2a_D mol:protein length:122 Phospholipase A2
 NLLQFNKMIK EETGKNAIPF YAFYGCYCGG GGNGKPKDGT DRCCFVHDCC YGRLVNCNTK SDIY
 GYITCGKGTN CEEQICECDR VAAECFRRNL DTYNNGYMFY RDSKCTETSE EC

Matching position : 43 - 50
 >1a2a_E mol:protein length:122 Phospholipase A2
 NLLQFNKMIK EETGKNAIPF YAFYGCYCGG GGNGKPKDGT DRCCFVHDCC YGRLVNCNTK SDIY
 GYITCGKGTN CEEQICECDR VAAECFRRNL DTYNNGYMFY RDSKCTETSE EC

Matching position : 43 - 50
 >1a2a_F mol:protein length:122 Phospholipase A2
 NLLQFNKMIK EETGKNAIPF YAFYGCYCGG GGNGKPKDGT DRCCFVHDCC YGRLVNCNTK SDIY
 GYITCGKGTN CEEQICECDR VAAECFRRNL DTYNNGYMFY RDSKCTETSE EC

Matching position : 43 - 50
 >1a2a_G mol:protein length:122 Phospholipase A2
 NLLQFNKMIK EETGKNAIPF YAFYGCYCGG GGNGKPKDGT DRCCFVHDCC YGRLVNCNTK SDIY
 GYITCGKGTN CEEQICECDR VAAECFRRNL DTYNNGYMFY RDSKCTETSE EC

Matching position : 43 - 50
 >1a2a_H mol:protein length:122 Phospholipase A2
 NLLQFNKMIK EETGKNAIPF YAFYGCYCGG GGNGKPKDGT DRCCFVHDCC YGRLVNCNTK SDIY
 GYITCGKGTN CEEQICECDR VAAECFRRNL DTYNNGYMFY RDSKCTETSE EC

Figure 3. Sample output frame of the consensus pattern search (only first output frame is shown here). The input search string is CCXXHXXC.

A sample output of the result of a typical search for a sequence pattern in all the proteins is shown in Figure 3. The pattern searched is taken from PROSITE⁸ corresponding to a key consensus pattern involved in the function of phospholipase A₂. The pattern is CCXXHXXC (where C is cysteine, X is any amino acid and H is histidine) and histidine is the active site residue. As can be seen from Figure 3 almost all the hits correspond to various structures of phospholipase A₂. This simple search helped in the recognition of all the homologous structures in a family without performing global alignment of complete amino acid sequences.

A simple variant of the input sequence pattern could result in the recognition of a non-phospholipase A₂. Depending upon the three-dimensional structure of this hit one might introduce, by site-directed mutagenesis, the residue necessary for the function of phospholipase A₂ in an otherwise non-phospholipase A₂.

The sequence and structure databases are being updated every week from PDB and fully incorporated with the PSST search tool. The search engine is written in Perl, version 5 (ref. 9) and it can be executed on our NT-4.0 Bioinformatics server (a 300 MHz Pentium II processor, 128 Mbytes of main memory). The front

end input data part of this tool is written in HTML and JavaScript and allows user-friendly web forms.

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