

A fast Peptide Match service for UniProt Knowledgebase

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ABSTRACT

Summary: We have developed a new web application for peptide matching using Apache Lucene-based search engine. The Peptide Match service is designed to quickly retrieve all occurrences of a given query peptide from UniProt Knowledgebase (UniProtKB) with isoforms. The matched proteins are shown in summary tables with rich annotations, including matched sequence region(s) and links to corresponding proteins in a number of proteomic/peptide spectral databases. The results are grouped by taxonomy and can be browsed by organism, taxonomic group or taxonomy tree. The service supports queries where isobaric leucine and isoleucine are treated equivalent, and an option for searching UniRef100 representative sequences, as well as dynamic queries to major proteomic databases. In addition to the web interface, we also provide RESTful web services. The underlying data are updated every 4 weeks in accordance with the UniProt releases.

Availability: <http://proteininformationresource.org/peptide.shtml>

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Supplementary information: Supplementary data are available at *Bioinformatics* online.

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1 INTRODUCTION

Locating occurrences of a specific peptide in a protein sequence database is important for protein identification in proteomics studies as well as for sequence-based protein retrieval. The Protein Information Resource (Wu *et al.*, 2003) has hosted a peptide match service for >10 years. It is one of the most popular tools on the Protein Information Resource website.

Although exact peptide matching is algorithmically simple, quickly locating a peptide in a large protein sequence database has become challenging, as the number of protein sequences exponentially increases. So far, to our knowledge, no other peptide match services provide all of the key characteristics in our new service, including search speed, support for any query peptide of ≥ 3 amino acids long, coverage of all UniProtKB sequences including isoforms (UniProt Consortium, 2013) and frequent update. Other major peptide match methods are UniPept (Mesuere *et al.*, 2012), which indexes *in silico* digested tryptic peptides, and Pep2Pro (Askenazi *et al.*, 2010), which uses indexes based on 6-mers. A detailed comparison of these and our method

is provided in the Supplementary Material (Supplementary Table S1). Another related work is SANS (Koskinen and Holm, 2012), which uses suffix arrays to effectively identify protein sequence similarities in the range of 50–100% identity.

As brute-force peptide matching is slow, we need effective ways to index protein sequences, analogous to indexing in relational databases. The recent advance in search engine technology, such as the Apache Lucene™ (McCandless *et al.*, 2010) information retrieval software library, provides a scalable and high-performance index structure as well as powerful and efficient search algorithms to build a fast search engine for our Peptide Match service. Here we present our new peptide matching application, with an enhanced web user interface.

2 METHODS

The source protein sequence database is UniProtKB (including isoform sequences), currently containing >40 million protein sequences. The index engine is built based on Apache Lucene. We index the original protein sequences as well as leucine and isoleucine equivalent sequences. We include accession number, protein name, organism, taxonomic group and lineage from UniProtKB records to annotate the matched proteins. We also provide protein cross-references to proteomic/peptide spectral databases if the matched proteins are present in those databases. We store the corresponding UniProtKB accession numbers that are mapped to NIST Peptide Libraries (Stein and Rudnick, 2009), PeptideAtlas (Desiere *et al.*, 2006), PRIDE (Vizcaino *et al.*, 2013) and Immune Epitope Database (IEDB) (Vita *et al.*, 2009) using our ID mapping (Huang *et al.*, 2011) services. These informational fields are stored, and key fields are indexed to support field-specific search and fast retrieval.

The protein sequence is analysed by Lucene NGramTokenFilter, which generates a series of 3-gram terms from the original sequence, and therefore, allows searching for a tri-peptide. These 3-gram terms filter search space analogous to other approaches: tryptic peptides in UniPept, 6-mers in Pep2Pro and suffixes in SANS. Internally, Lucene index maintains an inverted table for a term and its corresponding protein accession numbers, and stores the term frequency and position occurrence. The Lucene's Term Query method can only find the exact match with the same length term from the indexes. For example, the Term Query for 3-gram terms would match 'MKE' but not 'MKEV'. We, therefore, use Lucene's Phrase Query to search for overlapping consecutive 3-gram terms derived from the query peptide. We also use Lucene's Boolean Query to combine the Phrase Query of sequence field and Term Query of other informational fields if the peptide search is restricted by a specific set of organisms, taxonomic groups or lineage.

To allow users to determine whether the given query peptide has previously been observed in mass spectrometry (MS)-based proteomics

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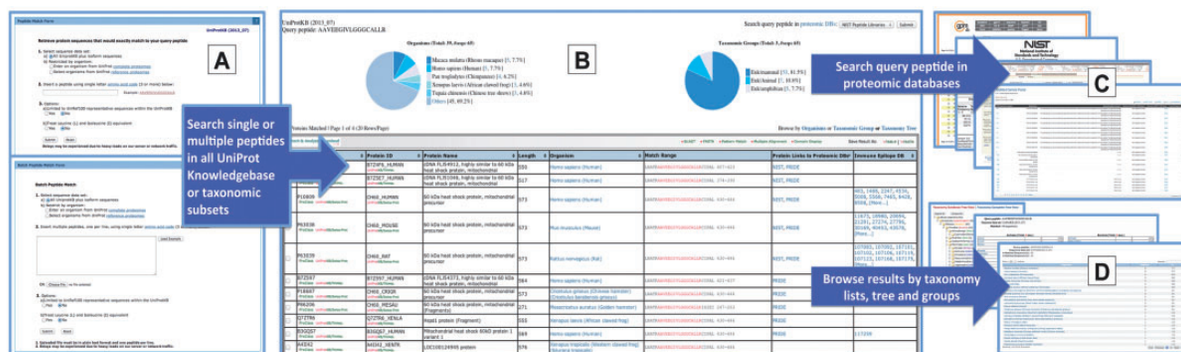


Fig. 1. The search interfaces and functionalities of the Peptide Match server. (A) search interfaces; (B) matched result summary; (C) dynamic proteomic databases search results; (D) taxonomic views

experiments, we provide a dynamic search of the query peptide against four major proteomic/peptide spectral databases: gpmDB (Craig *et al.*, 2004), NIST Peptide Libraries, PeptideAtlas and PRIDE from our web application using the search engines of those individual databases. The current implementation of Peptide Match service uses Apache Lucene 3.5 and Apache Solr™ 3.5 for indexing and searching. The web application is developed in Java Server Pages/Servlets running on Apache Tomcat 6.

3 RESULTS

The search engine is fast and accurate. A typical peptide match from our web interface takes only a second, including time spent rendering the results. On the search time itself, our test results showed an average of 0.1 s from a random sample of ~11 million peptides of 10-mers derived from the UniProtKB Human complete proteome. For UniProtKB release 2013_07 containing ~40 million protein sequences with annotations (24 GB), the index size is 98 GB and the indexing time is 5 h on a Linux server with Intel® Xeon® 2.00 Ghz CPUs and 128 GB RAM.

The Peptide Match web interface provides multiple functionalities, both for input query and output navigation (Fig. 1). A user can input a query peptide and specify whether to search the entire UniProtKB or limit the search to sequences from one or more selected organisms (Fig. 1A). For the latter, a user can either enter an organism name/taxonomic ID of a UniProt complete proteome or select a set of organisms from the UniProt reference proteomes. Additional options are provided to (i) restrict the matches in UniRef100 (Suzek *et al.*, 2007) representative sequences to reduce redundancy and (ii) equate isobaric leucine and isoleucine to support MS-based proteomics. The match results are summarized by organism and taxonomic group as pie charts and listed as tables. The matched proteins are shown in paginated tables (Fig. 1B), along with annotations, matched region(s) in the protein sequence and hypertext links to corresponding databases. The match results can be sorted by different fields and browsed by organisms, taxonomic group or taxonomy tree (Fig. 1D). As an example (Fig. 1), peptide ‘AAVEEGIVLGGGCALLR’ matches 65 proteins in UniProtKB, among which 53 are from mammals, and 10, 5, 4 have links to corresponding proteins in PRIDE, NIST Peptide Libraries and IEDB, respectively. For direct proteomic evidence of the query peptide, the dynamic search using search engines of the proteomic/peptide spectral databases further displayed the peptide with links to the source MS data (Fig. 1C).

For a given peptide, the matching protein entries from UniProtKB database can be directly accessed using a simple

URL (e.g. <http://proteininformationresource.org/peptide/SVQYDDVPEYK>), and used for other servers as in PeptideAtlas. We also provide a web interface for batch submission of peptides to our service as well as RESTful web services. The underlying data of our new Peptide Match server are updated every 4 weeks in accordance with the UniProt releases. The Peptide Match services will be forthcoming from the UniProt website (www.uniprot.org).

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Conflict of Interest: none declared.

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