pLM-BLAST – distant homology detection based on direct comparison of sequence representations from protein language models

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Abstract

Summary: Homology detection by sequence comparison is a typical first step in the study of protein function and evolution. Here, we describe a new homology detection tool, pLM-BLAST, that uses a modified Smith-Waterman algorithm for unsupervised comparison of single-sequence representations obtained from a protein language model (such as ProtT5) trained on millions of sequences. In our benchmarks, pLM-BLAST has shown the ability to detect homology between highly divergent proteins, demonstrating its applicability to tasks such as protein classification, domain annotation, and function prediction. Availability and Implementation: pLM-BLAST is available as a web server in the MPI Bioinformatics Toolkit (https://toolkit.tuebingen.mpg.de/tools/plmblast), where it can be used to search precomputed databases. It is also available as a standalone tool to build custom databases and run batch searches (https://github.com/labstructbioinf/pLM-BLAST).

Introduction

Homology, i.e., descent from a common ancestor, and its inference are fundamental to comparative, evolutionary, and molecular biology. In the case of proteins, statistically significant local or global sequence similarity is accepted as the primary marker for inferring homology. When the similarity between the protein sequences being compared is high (>=30%), homology can be readily detected using methods based on sequence-to-sequence and sequence-to-profile comparisons such as BLAST or PSI-BLAST (Altschul et al., 1997; Eddy, 2011). However, when similarity is low (<30%), methods based on profile HMMs, such as HMMER and HHsearch, are currently our best tools to infer homology (Steineggger et al., 2019).

In the case of highly distant evolutionary relationships, sequences may have diverged to the point where we can no longer detect their relatedness. Because structures diverge much more slowly than sequences, their similarity is often used to infer homology in such cases. In fact, owing to the recent revolution in structure prediction (Lin et al., 2022; Jumper et al., 2021; Li et al., 2022), predicted structures are available for a large proportion of known proteins, and structural similarity is increasingly being used to infer homology. However, similar structures, particularly at the domain and sub-domain levels, may have evolved convergently due to the limited number of structural solutions available to a folded polypeptide chain, and thus structural similarity is often not conclusive evidence of common
ancestry. To address this issue, several deep learning-based methods have sought to detect weak sequence signals between highly divergent proteins in recent years (Zheng et al., 2019; Gao and Skolnick, 2021; Li et al., 2017). The most promising of these methods, such as knnProtT5 (Schütze et al., 2022), TM-Vec (Hamamsy et al., 2022), and DEDAL (Llinares-López et al., 2022), rely on single-sequence representations obtained from protein language models (pLMs) (Bepler and Berger, 2021). pLMs are neural networks trained on a large number of natural protein sequences for tasks such as guessing masked residues based on contextual information (Elnaggar et al., 2021; Lin et al., 2022). Once trained, pLMs can be used to rapidly compute the aforementioned representations (also referred to as embeddings), i.e., numerical descriptors of protein sequences that place them in the context of the total knowledge collected by the network. Because of their high information content, sequence embeddings have been successfully applied to many other tasks, including prediction of tertiary structures (Lin et al., 2022), transmembrane segments (Bernhofer and Rost, 2022), and signal peptides (Teufel et al., 2022).

In this report, we describe pLM-BLAST, a new tool for detecting local homology between protein sequences that combines pLM representations with a local similarity detection algorithm inspired by BLAST (Altschul et al., 1997). In contrast to TM-vec and DEDAL, pLM-BLAST is based on an unsupervised approach that does not require training of a specialized deep-learning model or defining positive labels based on structurally similar or homologous protein pairs. To this end, we implemented a modified Smith-Waterman algorithm in which the substitution matrix is generated directly from the raw pLM representations of the two sequences to be compared. In this work, we used the pLM ProtT5 (Elnaggar et al., 2021), but this approach can be used in conjunction with other pLMs, such as ESM-2 (Lin et al., 2022), to generate local alignments. In a benchmark using domain pairs from the ECOD protein classification database (Cheng et al., 2014), pLM-BLAST demonstrated its suitability for fast pairwise comparisons and database searches to detect distant homology.

**Methods**

pLM-BLAST extends the concept of BLAST by replacing invariant substitution matrices, such as BLOSUM62 (Henikoff and Henikoff, 1992), with per-residue similarities between ProtT5
embeddings. Consequently, the similarity between a given pair of residues is entirely context-dependent. Such sequence context information has been shown to significantly improve the sensitivity of sequence search methods and enable the detection of nontrivial conserved patterns (Biegert and Söding, 2009; Remmert et al., 2011).

**Substitution matrix**

The embedding of a sequence \(seq_i\) is given by matrix \(E_i \in \{e_{ij}\}\) of size \(n \times m\), where \(n\) and \(m\) denote the number of residues and the size of the embedding, respectively. To obtain standardized and comparable results, the values in each row are scaled by dividing them by the Euclidean norm of the row

\[
e_{ij}^+ = \frac{1}{\sqrt{\sum_{j=m}^{n} e_{ij}^2}} e_{ij}
\]

where \(e_{ij}\) denotes the \(E_i\) matrix element in the \(i\)-th row and \(j\)-th column, and \(\dagger\) is the normalization operator. In pLM-BLAST, a substitution matrix \(S_{ik}\) for two sequences \(seq_i\) and \(seq_k\) is calculated as

\[
S_{ik} = E_i^{\dagger T} E_k^+ 
\]

The element \(s_{ij}\) of the matrix \(S_{ik}\) is in the interval \((-1,1)\) and denotes the embedding similarity of the \(i\)-th residue of \(seq_L\) and the \(j\)-th residue of \(seq_K\). In other words, the above equations are equivalent to calculating the cosine similarity of each possible pair of residues from \(seq_L\) and \(seq_K\) and placing them in a two-dimensional array.

**Scoring matrix**

A scheme adapted from the Smith-Waterman algorithm is used to create a scoring matrix \(H_{LK} \ni \{h_{ij}\}\) of size \((|L|+1) \times (|K|+1)\), where the first row and column are filled with zeros and the rest of the matrix is filled with the following procedure

\[
h_{ij} = \max \left\{ h_{i-1,j-1} + s_{ij}, \max_p \{h_{i-p,j}\}, \max_s \{h_{i,j-s}\} \right\}
\]
Unlike the original Smith-Waterman algorithm, pLM-BLAST does not use gap penalties because the $s_{ij}$ values for dissimilar regions are negative due to the use of the cosine similarity property. Gap penalties were implemented, but in our tests they did not provide any significant benefits, so they are turned off by default.

**Traceback procedure**

To extract matching regions for a given pair of sequences, a traceback procedure is used. For this purpose, $n$ possible paths $P$ are constructed, starting from the right and bottom edges of the matrix $H$. The path $p_n$ is a set of coordinates from the matrix $H_{LK}$, where $n \in N$ and $N$ equals $|L| + |K|$. 

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**Input:** $H$

**Output:** $P$

**For** $n \in N$ **do**

- Initialize path $p_n$
- Set pair $(i', j')$ to $n$ border coordinates

**While** $i \geq 1$ and $j \geq 1$ **do**

  - $(i', j') = \text{argmax}_{i,j} \{H_{i-1,j-1}, H_{i,j-1}, H_{i-1,j}\}$
  - **If** $H_{i',j'} \leq 0$ **then**
    - Break
  - **Else**
    - Add $(i', j')$ to $p_n$
    - Decrement by one $i, j$

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**Local alignment**

Each path is scanned for the presence of high-scoring sub-paths (local alignments) using the substitution matrix. A moving average is applied to each path $p_n$ and all continuous subpaths where the average is greater than a specified threshold are treated as local alignments (the threshold is set by default to 1 standard deviation of the substitution matrix). Each local alignment is assigned a score calculated as the mean of substitution matrix values at coordinates defined by its subpath. In the case of database searches, an optional procedure may be applied in which local alignments to a single database entry are merged. Our tests showed that the standard deviation of the substitution matrix tends to assume a value of around 0.05, whereas the path score for meaningful alignments is an
order of magnitude higher. However, increasing (>1) or decreasing (<1) the standard deviation cut-off will make the algorithm more strict or permissive, respectively.

Benchmark

The sequences of all ECOD domains (version 20220912), excluding nested domains, with a length of 50-600 amino acids were obtained with localpdb ECOD plugin (Ludwiczak et al., 2022), and clustered using MMSeqs2 at a sequence identity cut-off of 30% (Steinegger and Söding, 2017). The longest sequences were selected as clusters’ representatives and those belonging to H-groups with less than five members were removed. The final benchmark set was created by randomly selecting five domains from each Homology(H)-group, an ECOD level that groups together domains with common ancestry. This procedure was designed to select domains from different Topology(T)-groups; a T-group is a sublevel that groups together homologous domains within an H-group that are topologically similar. For each ECOD domain, an HMM profile and sequence embedding were calculated using three iterations of HHblits over the UniRef30 database with default settings (Steinegger et al., 2019) and ProtT5 (prot_t5_xl_half_uniref50-enc) (Elnaggar et al., 2021), respectively (note that the usage of the ProtT5 model limits the maximal sequence length to 1000). The embeddings were compared all-against-all with pLM-BLAST and TM-Vec, whereas the HMM profiles were compared with HHsearch. To speed up the embedding comparison, a pre-filtering procedure was used. All the embeddings were individually flattened to a constant size by applying an arithmetic mean and then efficiently compared to each other with a cosine similarity metric. Only pairs that scored 0.4 or higher were passed to the actual pLM-BLAST (the 0.4 cut-off corresponds to the 90th percentile of all scores). The same procedure was used to improve the performance of database searches: first, the database of flattened embeddings is searched with the cosine similarity, which is much faster than the comparison of per-residue embeddings, and then the actual pLM-BLAST comparisons are performed only for matches above the user-provided cosine similarity cut-off (95th percentile by default).

Results

To assess the sensitivity of pLM-BLAST in detecting distant homologs, we evaluated its ability to correctly detect homology between ECOD domains in a benchmark set (see
Methods) and compared its sensitivity in this task to that of BLAST, TM-Vec, and HHsearch. While matches between domains from the same H-group were considered true positives, matches between different H-groups were considered false positives. In our benchmark, HHsearch performed best, pLM-BLAST and TM-Vec performed equally well, and BLAST was the least sensitive. This result was quite unexpected since TM-Vec uses sequence representations from ProtT5 as an input to a specialized deep learning model, trained end-to-end to produce structural alignments, while pLM-BLAST is based only on unsupervised comparisons of these representations. This suggests that the information contained in the raw embeddings is sufficient for homology detection (Figure 1A).

We conjectured that the significantly higher performance of HHsearch in our benchmark may result from it detecting a higher number of true positives and/or fewer false positives. To investigate this systematically, we used US-align (Zhang et al., 2022) to compare the structures corresponding to the true positive and false positive pairs returned by the three tools (Figure 1D). In the case of HHsearch, we found only a few structurally similar false positive pairs, which is to be expected given that ECOD H-groups were defined using HHsearch (Cheng et al., 2014). In contrast to HHsearch, both pLM-BLAST and TM-Vec yielded a number of structurally similar false positive pairs (Supplementary Table 1; Figure 1D), many of which belong to the same X-group, an ECOD level that groups together domains that may be homologous. In addition, both TM-Vec and pLM-BLAST also detected connections between domains of different X-groups that were often hinged on the presence of a subdomain-sized fragment (Alva et al., 2015; Kolodny, 2021). For example, pLM-BLAST found a fragment conserved between domains of two different sandwich folds, namely cupredoxin (ECOD H-group 3156.1) and immunoglobulin (11.1, Figure 1B). A homologous relationship between these two domains has been discussed previously (Gough and Chothia, 2004; Stevens, 2008), and a pLM-BLAST scan of the entire ECOD database with a cupredoxin domain as a query revealed additional connections to Ig-like domains (Figure 1C). We also tested other pLM models such as ESM2 (Lin et al., 2022) and SeqVec (Heinzinger et al., 2019); however, ProtT5 performed best in our tests.

In summary, pLM-BLAST is a sensitive tool for remote homology detection based on comparison of sequence representations obtained from the pLM ProtT5. pLM-BLAST is
available both as a stand-alone package and as an easy-to-use web server within the MPI Bioinformatics Toolkit (Zimmermann et al., 2018), where it can be used to search several precomputed databases (e.g., ECOD, InterPro, and PDB). pLM-BLAST is also quite fast: while the computation of ProtT5 representations for ECOD30, which included 29,330 domains, took 30 minutes on a single NVIDIA RTX A4500 card, a search over it took on average 2 mins on a 20-core CPU (including the steps of loading the database to the memory and formatting the results). Currently, the main limitations of pLM-BLAST are the maximal size of the target database (searching huge, redundant databases containing millions of sequences would be computationally very expensive) and the fact that it typically produces rather short, local alignments (this, however, may be attributed to the used pLM model rather the pLM-BLAST algorithm itself). Nevertheless, we anticipate that pLM-BLAST will prove useful for studying protein function and evolution, also in the cases of singletons/orphan sequences and taxonomically-restricted genes (Barrera-Redondo et al., 2022) for which deep HMM profiles cannot be calculated.

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Figure Legends

**Figure 1.** (A) Benchmark on the task of reconstructing H-groups defined in the ECOD database. False positives are shown in the logarithmic scale. (B) A substitution matrix depicting the similarity between cupredoxin and immunoglobulin folds. Axes numbering denote the consecutive residues. The brighter the color the more similar the sequences in a given area. The highest-scoring local alignment is indicated with a thin red line. (C) (top) pLM-BLAST scan of ECOD database with the sequence of cupredoxin domain. (bottom) Structures of cupredoxin and immunoglobulin folds where homologous regions detected by pLM-BLAST are shown in red. (D) Sequence and structure similarity of false positive pairs.
Sequence similarity was calculated with HHsearch (left panel), pLM-BLAST (middle), and TM-Vec (right).

**Bibliography**


Li, S. et al. (2017) Protein remote homology detection based on bidirectional long short-term memory. BMC Bioinformatics, 18, 443.


