Sequence analysis

Improved BLAST searches using longer words for protein seeding

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Received on August 8, 2007; revised on September 13, 2007; accepted on September 19, 2007

Advance Access publication October 6, 2007

Associate Editor: Thomas Lengauer

ABSTRACT

Motivation: The blastp and tblastn modules of BLAST are widely used methods for searching protein queries against protein and nucleotide databases, respectively. One heuristic used in BLAST is to consider only database sequences that contain a high-scoring match of length at most 5 to the query. We implemented the capability to use words of length 6 or 7. We demonstrate an improved trade-off between running time and retrieval accuracy, controlled by the score threshold used for short word matches. For example, the running time can be reduced by 20-30% while achieving ROC (receiver operator characteristic) scores similar to those obtained with current default parameters.

Availability: The option to use long words is in the NCBI C and C++ toolkit code for BLAST, starting with version 2.2.16 of blastall. A Linux executable used to produce the results herein is available at: ftp://ftp.ncbi.nlm.nih.gov/pub/agarwala/protein_longwords

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1 INTRODUCTION

BLAST is a widely used set of programs that produce local alignments for input query sequences by searching a database of subject sequences. In this note, we consider the blastp module where the query is a protein and the database also contains proteins, and the tblastn module where the query is a protein and the database contains DNA sequences that are hypothetically translated. The BLAST algorithm (Altschul et al., 1990, 1997) consists of three major stages:

(1) Seeding stage: subject sequences are scanned to find locations that resemble the query sequence. The output is a list of offset pairs called seeds. Each seed represents coordinates on the query and subject sequences.

(2) Ungapped extension stage: an initial extension for each seed is performed by comparing the amino acids on both sides of the seed. A decision is made whether the location is of interest or not. At this stage, the majority (more than 99%) of seeds are typically discarded.

(3) Gapped extension stage: each high-enough-scoring ungapped alignment becomes the starting point for a local gapped alignment that attempts to improve the score further. Gapped alignments for which the calculated expect-value is below the threshold (default is 10) are considered for the final result set.

Each stage takes ~30% of the total running time.

This note presents improvements in the seeding stage. In BLAST, seeding is implemented via a lookup table approach. When we scan a sequence, for each location we group several consecutive letters into a word. A numerical representation of this word becomes the entry’s offset in the lookup table. The lookup table was populated in advance using words from the query sequence, so when we scan the subject and find that the word being considered corresponds to a non-empty entry in the lookup table, we have a seed. An example of seeds of length 6 is shown below:

Query: 127 THRHMTEFTGLDMEMAF
T_RH+ E +D_EMAF

To improve lookup performance, BLAST utilizes CPU caches by employing a small bit array structure that is consulted prior to accessing the lookup table (Cameron et al., 2006). The bit array is used to avoid unnecessary (and expensive) lookups in the lookup table for entries known to be empty.

2 METHODS

To improve the performance of the blastp and tblastn modules, we modified the implementation of the seeding stage to reduce the number of seeds generated, thus reducing time spent in later stages. The primary concern was not with the speed of the seeding stage (although we did spend some time optimizing it), but with the number and quality of seeds produced.

An obvious way to reduce the number of seeds is to increase the number of amino acids considered, which we call word size, when deciding if we have a seed or not. The baseline code from which we started supports word size up to 5, with the default of 3 and an additional requirement that there be two seeds in close proximity (called two-hit in BLAST). An early version used size 4 (Altschul et al., 1990). A previous study showed that the two-hit requirement with word size 3 has a better time-sensitivity trade-off than using the older single-hit rule (Altschul et al., 1997). Since the current code supports word size 5, it was natural to ask whether lookup tables for word sizes 6 and 7 could be designed, and if so, how they would perform.

A simple increase in word size is not practical because of exponential growth in the size of the lookup table, e.g. for word size 7 we would have at least 20^7 = 1.3e9 lookup table entries! To tackle this explosion,

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we decided to use compressed alphabets that group similar letters together into disjoint sets (Edgar, 2004). Grouping letters decreases memory requirements, and if the compressed alphabet is chosen carefully, the loss of information is less than the smaller alphabet size would suggest.

To implement the seeding method described in this article, the following subproblems must be solved: (i) choose an alphabet size; (ii) given an alphabet size, determine the compressed alphabet; (iii) given a compressed alphabet, decide how to populate the lookup table; (iv) given a location, decide how to score a candidate seed; (v) decide how to evaluate performance of the algorithm.

2.1 Choice of compressed alphabets sizes
The size of the CPU cache was the primary factor in deciding the size of the alphabet to use for a given word size. On modern CPUs with traditional architecture, the size of the level 2 cache is usually somewhere between 512 KB and 4 MB, so 22-25 bits of address space is an estimate for the ‘working set’ of the bit array and this in turn gives us the number of entries in the lookup table.

Because the access pattern is highly non-uniform (due to the non-uniform frequency distribution of amino acids), the actual size of the bit array can be slightly larger than available cache while still being very effective. For example, words starting with many ‘W’ characters are extremely rare. Thus, we can rely on the CPU’s cache control mechanism to adapt automatically to the incoming stream of letters to keep the most frequent combinations in cache. An additional benefit of a non-uniform access pattern is that performance changes gracefully as the amount of available CPU cache changes.

For word sizes 6 and 7, we chose 15-letter and 10-letter alphabets, respectively. This setup leads to a 1.4 MB array of bits for 6-letter words and a 1.2 MB bit array for 7-letter words.

2.2 Choice of compressed alphabets
Using compressed alphabets in (Edgar, 2004) as the starting point, we tested various compressed alphabets and selected the following ones:

   | J | L | M | V | A | S | T | B | D | E | N | Z | K | R | Q | G | F | Y | P | H | C | W |
   | 1 |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |
   | ST | I | V | L | K | R | E | Q | Z | A | G | B | D | P | N | F | Y | H | C | W |   |

   Each string of consecutive letters without spaces represents one letter set in the compressed alphabet. Alphabet (1) is the ‘SE-V(10)’ alphabet in Edgar (2004) with ambiguity characters added. For the 15-letter alphabet (2), however, we added one more letter set to Edgar’s ‘SE-B(14)’.

   Because the bit array may not fit into CPU cache on some machines, using an alphabet’s matrix entropy (Altschul, 1991) as the sole selection mechanism as used in protein BLAST’s lookup table population. Not only is the exact encoding of a word included in the lookup table, but also all of the word’s neighbors, words that match the original word with a score not lower than a given threshold.

   The matching score between two words is the sum of substitution scores for each pair of corresponding letters. The score for aligning one letter with another is taken from a score matrix.

   Because substitution score usage is central to the algorithm, we cautiously decided not to use BLOSUM62 (Henikoff and Henikoff, 1992) at the standard precision because that precision may be too low.

   Instead we used a scaled version of the integer scores obtained by rounding $C\ln r_{ij} = C\ln(q_i/(p_i p_j))$, where $C$ is a constant, $q_i$ is the probability of aligning amino acid $i$ with amino acid $j$ and $p_i$, $p_j$ are the background probabilities of $i$ and $j$, respectively (Schräffer et al., 2001).

   Using a high-precision threshold allowed us much greater freedom in selecting the sensitivity of the BLAST algorithm. For example, in baseline blastp, going from threshold value 12 to 11 almost doubles the running time and there is no allowed threshold in between. Now, we can select any value and thereby adjust performance to meet time and computing power constraints. The new threshold is entered as a floating point number, treated on the same scale as the current low-precision threshold, and then internally converted to a scaled-up integer.

2.4 Calculating matching score
Converting the query and subject to the compressed alphabet and using a square score matrix (10 x 10 or 15 x 15) produced subpar results. Using a square matrix loses information in both query and subject sequences, leading to a high proportion of seeds that are discarded in later stages.

   To prioritize potential seeds, we decided to weight the matching score by the probability of a match. For example, the matching score for ‘A’ to the ‘AST’ letter set is calculated based on the frequency ratio of ‘A’ to each one of the three letters weighted by the conditional probability of that letter. This additional weighting is needed because at the time of lookup table population, we do not know which exact letters we will encounter—the compressed alphabet obscures that information. We used background probabilities from Robinson and Robinson (1991) to calculate conditional probabilities, e.g. $p(A|V+S/V+T)$, $p(T|A+S/V+T)$.

   In practice, Robinson-Robinson background probabilities produced good results, sometimes even better than using the observed letter frequencies of the database.

   Mathematically, the implemented scoring matrix is 20 (plus ambiguity characters) rows by 10 or 15 columns. The score for matching amino acid $i$ to letter set $g_k$, denoted by $s_{ik}$, is calculated using the formula:

   $$ s_{ik} = \frac{1}{\lambda} \ln \sum_{j=0}^{20} p(j | g_k) $$

   where $j$ is an amino acid that belongs to the letter set $g_k$, $r_i$ is the frequency ratio defined above, $\lambda$ is a scaling factor and $p(j | g_k)$ is the group conditional probability of amino acid $j$ and is given by $p(j | \sum_{0}^{20} p_i)$.

   The example in the Introduction section has two seeds. The query is dleova2 and the subject is dib8a2z from ASTRAL (Chandonia et al., 2004). Seed1, a 6-letter seed with mismatch at position 1 and 5, illustrates that we can have multiple mismatches at arbitrary positions. Seed1 has a scaled score of 19.1 exceeding the recommended threshold of 18.8.

2.5 Performance evaluation
To evaluate result quality, we used SCOP/ASTRAL (Chandonia et al., 2004). The SCOP/ASTRAL dataset provides a gold standard from which it is possible to identify true and false positives in the output and compute ‘receiver operator characteristic’ (ROC) scores (Gribskov and Robinson, 1996). We report ROC10000 scores, meaning that matches for all queries are combined and ranked by expect-value up to the first 10 000 false positives. Program configuration A is considered to have a higher quality output than configuration B if the ROC score for A is higher than for B. We used sequences of < 40% identity in the current ASTRAL version 1.71. True positives are query-subject pairs in the same superfamily. We ignored self-hits and we used as queries, the 7218 sequences that have at least two sequences in the same superfamily.

   Unfortunately, ASTRAL is not large enough to adequately represent the more commonly used nr and Swiss-Prot databases. We therefore decided to use a hybrid measurement approach. We measured ROC scores produced by using different values of the threshold. We then ran
4 DISCUSSION

Longer seeds would be expected to improve sensitivity of BLAST searches, but memory limitations make longer seeds difficult to implement with a good running time. Our implementation shows that using compressed alphabets can overcome the memory limitations for word sizes 6 and 7, while retaining overall sensitivity of the algorithm. For output quality comparable to the baseline BLAST, the running time decreases by 20–30%. The implementation enables a fine-scale trade-off between running time and output quality. Preliminary experimental results for larger (eight letters or more) word sizes were not promising.

BLAST can take as input multiple queries (e.g. in a single FASTA-formatted file) and can scan the database for several queries simultaneously to save time. Therefore, for any usage with many queries, the effective query size should be large. However, when the effective size of the query or the database is small, our approach cannot yet compete with baseline BLAST or deterministic finite automaton seeding (Cameron et al., 2006) due to additional setup time required for building the lookup table and to slightly slower scanning speed. Our method is targeted for ‘bulk-processing’ applications that align large queries to large databases, for which our implementation outperforms both baseline blastall and the implementation of Cameron et al. (2006).

ACKNOWLEDGEMENTS

This research was supported by the Intramural Research Program of the National Institutes of Health, National Library of Medicine. Thanks to Stephen Altschul, E. Michael Gertz and Aleksandr Morgulis for helpful comments.

Conflict of Interest: none declared.

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