EXPERIMENTAL RESULTS ON MULTIPLE PATTERN MATCHING ALGORITHMS FOR BIOLOGICAL SEQUENCES

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Abstract: With the remarkable increase in the number of DNA and proteins sequences, it is very important to study the performance of multiple pattern matching algorithms when querying sequence patterns in biological sequence databases. In this paper, we present a performance study of the running time of well known multiple pattern matching algorithms on widely used biological sequence databases containing the building blocks of nucleotides (in the case of nucleic acid sequence databases) and amino acids (in the case of protein sequence databases).

1 INTRODUCTION

Multiple pattern matching is the computationally intensive kernel of many applications including information retrieval, intrusion detection systems, web filtering, virus scanners and spam filters. In recent years however, an immediate interest in string-matching problems as a powerful tool in locating nucleotide or amino acid sequence patterns in biological sequence databases has been witnessed. For example, when proteomics data is used for genome annotation in a process called proteogenomic mapping (Jaffe et al., 2004), a set of peptide identifications obtained using mass spectrometry is matched against a target genome translated in all six reading frames.

The multiple pattern matching problem can be defined as follows: Given a sequence database (or text) \( T = t_1t_2...t_n \) of length \( n \) and a finite set of \( r \) patterns \( P = p^1, p^2, ..., p^r \), where each \( p^i \) is a string \( p^i = p^i_1p^i_2...p^i_m \) of length \( m \) over a finite character set \( \Sigma \), the task is to find all occurrences of any of the patterns in the sequence database.

The naive solution to this problem is to perform \( r \) separate searches with a sequential algorithm (Navarro and Raffinot, 2002). While frequently used in the past, this technique is not efficient when a large pattern set is involved. The aim of the multiple pattern matching algorithms is to scan the input string \( T \) in a single pass to locate the occurrences of all patterns. These algorithms are based on single-pattern matching algorithms, with some of their functions generalized to process multiple patterns simultaneously during the preprocessing phase, generally with the use of trie structures and hashing.

The multiple pattern matching is widely used in computational biology for a variety of pattern matching tasks. For example, Brundo and Morgenstern use a simplified version of the Aho-Corasick algorithm to identify anchor points in their CHAOS algorithm for fast alignment of large genomic sequences (Brudno and Morgenstern, 2002; Brudno et al., 2004). Hyyro et al. demonstrate that Aho-Corasick outperforms other algorithms for locating unique oligonucleotides in the yeast genome (Hyyro et al., 2005). The SITEBLAST algorithm (Michael et al., 2005) employs the Aho-Corasick algorithm to retrieve all motif anchors for a local alignment procedure for genomic sequences that makes use of prior knowledge. Buhler et al use Aho-Corasick to design simultaneous seeds for DNA similarity search (Buhler et al., 2005).

This paper presents experiments for the running time of the well known Commentz-Walter, Wu-Manber, Set Backward Oracle Matching and Salmela-Tarhio-Kytöjoki multiple pattern matching algorithms for biological sequences. A detailed analysis of the multiple pattern matching algorithms presented in this paper, additional experiments on different types of data as well as a study on the preprocessing time and the memory requirements of the algorithms can be found in (Kouzinopoulos and Margaritis, 2010).
The aim of this experimental study is to identify a suitable and preferably fast multiple pattern matching algorithm for several problem parameters such as a given biological database, the size of the pattern set, and the length of the patterns.

2 EXPERIMENTAL METHODOLOGY

The experiments were executed locally on an Intel Core 2 Duo CPU with a 3.00GHz clock speed and 2 Gb of memory, 64 KB L1 cache and 6 MB L2 cache. The Ubuntu Linux operating system was used and during the experiments only the typical background processes ran. To decrease random variation, the time results were averages of 100 runs. All algorithms were implemented using the ANSI C programming language and were compiled using the GCC 4.4.3 compiler with the “-O2” and “-funroll-loops” optimization flags.

To compare the pattern matching algorithms, the practical running time was used as a measure. Practical running time is the total time in seconds an algorithm needs to find all occurrences of a pattern in an input string including any preprocessing time and was measured using the MPI_Wtime function of the Message Passing Interface since it has a better resolution than the standard clock() function.

The data set was similar to the ones used in (Sheik et al., 2005) and (Kalsi et al., 2008). It consisted of the SWISS-PROT Amino Acid sequence database with a size of \( n = 182,116,687 \) characters and an alphabet of size 20, the FASTA Amino Acid (FAA) sequence of the A-thaliana genome with a size of \( n = 11,273,437 \) characters and an alphabet of size 20 and the FASTA Nucleic Acid (FNA) sequence of the A-thaliana genome with a size of \( n = 118,100,062 \) characters and an alphabet of size 4.

3 EXPERIMENTAL RESULTS

In this section, the performance of the algorithms is evaluated according to their running time for different biological databases.

Figures 1 to 3 present the running time of the algorithms including preprocessing for the SWISS-PROT amino acid sequence database and for the FASTA amino acid and nucleic acid databases of the A-thaliana genome respectively for a pattern length of \( m = 8 \) and \( m = 32 \) and for 100 to 100,000 patterns. As can generally be seen from the Figures, by varying different parameters such as the size of the pattern set, the length of the patterns and the size of the alphabet can affect the performance of the algorithms in different ways.

In the case of the SWISS-PROT database and for a pattern length of \( m = 8 \), the SOG and BG algorithms had the best performance when up to 10,000 patterns were used while the SBOM algorithm was faster for more than 10,000 patterns. When a pattern length of \( m = 32 \) was used, the SOG and BG algorithms had the fastest running time for up to 20,000 patterns, while SBOM was faster for more than 20,000 patterns. The HG and Wu-Manber algorithms had an average performance for either \( m = 8 \) or \( m = 32 \) while Commentz-Walter was consistently the slowest algorithm in terms of running time.

For the FASTA amino acid database, and for a pattern length of \( m = 8 \), the SOG and BG algorithms were faster when up to 10,000 patterns were used while for more than 10,000 patterns, the Wu-Manber algorithm had the best performance, followed by SBOM. When a pattern length of \( m = 32 \) was used, the SOG and BG algorithms had the fastest running time for up to 30,000 patterns. For bigger pattern sets, Wu-Manber was the fastest algorithm. Commentz-Walter was the algorithm with the worst performance when \( m = 8 \) was used while for \( m = 32 \), the Commentz-Walter and the SBOM algorithms had the worst performance.

In the case of the FASTA nucleic acid database, SBOM was the algorithm that worked consistently faster for a pattern length of \( m = 8 \). When up to 2,000 patterns were used, Commentz-Walter was the slowest algorithm while for more than 2,000 patterns, HG was the algorithm with the worst performance. For a pattern length of \( m = 32 \) the Wu-Manber was the fastest algorithm, especially when more than 20,000 patterns were used.

Specific performance comments on the algorithms follow. Commentz-Walter was the algorithm with the fastest running time when used on the FASTA nucleic acid database with a pattern length of \( m = 32 \), especially when more than 10,000 to 50,000 patterns were used. The algorithm had the worst performance when used on the SWISS-PROT and the FASTA amino acid databases and thus its use is not recommended in general on large alphabet sizes such as amino acid databases. Wu-Manber was the fastest algorithm on the FASTA amino acid database when more than 10,000 patterns were used and on the FASTA nucleic acid for a pattern length of \( m = 32 \) together with the Commentz-Walter algorithm. On the SWISS-PROT database and for a pattern length of \( m = 8 \), the algorithm had a good performance with
Figure 1: Running time of the algorithms for the SWISS-PROT Amino Acid database.

Figure 2: Running time of the algorithms for the FASTA Amino Acid database.

a running time close to that of the other algorithms. The SBOM algorithm outperformed the rest of the algorithms when the SWISS-PROT database was used and for a pattern set size of more than 10,000 to 20,000 patterns. It had also the best performance on the FASTA nucleic acid text when a pattern length of \( m = 8 \) was used. On the FASTA amino acid had an average performance for \( m = 8 \) and the worst performance comparing to the rest of the algorithms for \( m = 32 \). Among the Salmela-Tarhio-Kytöjoki algorithms, SOG and BG had the best performance in practice when used on the SWISS-PROT database together with the SBOM algorithm and on the FASTA amino acid database for fewer than 5,000 to 20,000 patterns. On the FASTA nucleic acid database the Salmela-Tarhio-Kytöjoki algorithms had the slowest running time comparing to the rest of the algorithms and so their used is not recommended on small alphabet sizes such as DNA-type databases.

4 CONCLUSIONS

In this paper, experimental results of the well-known Commentz-Walter, Wu-Manber, SBOM and the Salmela-Tarhio-Kytöjoki algorithms were presented. The algorithms were compared in terms of running time for the SWISS-PROT Amino Acid sequence database and the FASTA Amino Acid (FAA) and FASTA Nucleic Acid (FNA) sequences of the A-thaliana genome and for sets of size between 100 and 100,000 patterns with a length of \( m = 8 \) and \( m = 32 \). The experimental study proved that no algorithm is the best for all values of the problem parameters. Instead it was shown that for different databases, different algorithms are preferable: Commentz-Walter had the best performance on the FASTA nucleic acid database for more than 10,000 patterns. Wu-Manber was the fastest algorithm for the FASTA amino acid database for more than 10,000 to 50,000
patterns. The Wu-Manber algorithm also outperformed the rest of the algorithms on the FASTA nucleic acid database for a pattern length of \( m = 32 \) and for more than 20,000 patterns. The SBOM algorithm had the best performance on the SWISS-PROT database for \( m = 8 \) and for more than 10,000 patterns and on the FASTA nucleic acid database, especially when a pattern length of \( m = 8 \) was used. Among the Salmela-Tarhio-Kytöjoki algorithms, HG did not perform well on any biological database while the SOG and BG algorithms had the fastest running time on the SWISS-PROT database when up to 10,000 to 20,000 patterns were used, and on the FASTA amino acid database when up to 10,000 to 30,000 patterns were used.

Since biological databases and pattern sets are usually inherently parallel in nature, the work presented in this paper could be extended with a focus on the speed up of the existing algorithms when parallel processed on traditional parallel architectures like cluster environments and multicore systems as well as on modern parallel systems like GPU architectures.

REFERENCES


