

Introduction

- BLAST, short for Basic Local Alignment Search Tool, is a fundamental algorithm in the life sciences that compares biological sequences [1.2].
- With the advent of next-generation sequencing (NGS), the exponential growth of sequence databases is arguably outstripping our ability to analyze the data.
- The previous studies [3.4.5.6] for accelerating BLAST on GPU used coarse-grained parallel approaches, (i.e. one sequence alignment is mapped to only one thread), which are not adapted to GPU architecture and cannot solve the irregular memory access in BLAST.
- Our goal is to propose a faster GPU-BLAST using the fine-grained multithreaded approach.

BLAST Algorithm

- BLAST is a family of algorithms with variants used for different searching alignments, e.g., BLASTp for protein sequence, BLASTn for nucleotide sequence.
- BLAST is a heuristic method that approximates the Smith-Waterman algorithm, searching for similarities between a short query sequence and a large set of database sequences (subject sequences).
- BLAST algorithm locate high scoring short matches (i.e., hits) between the query sequence and the subject sequences, and extend hits to longer alignments. Four stages of BLAST are presented as below (Fig. 1):
 - **1.** Hit detection identifies high scoring short matches (i.e., hits) with a fixed length between a query sequence and the subject sequences via Deterministic Finite Automaton (DFA) or lookup table.
 - 2. Ungapped extension determines whether multiple hits can form the basis of a local alignment without insertions and deletions of residues. Extensions are triggered only if distances of two neighboring hits are within a threshold.
 - **3.** Gapped extension performs the further extension based on alignments from the previous stage and allows gaps.
 - **4.** Gapped alignment with traceback re-scores all alignments from the previous stage using a traceback algorithm.



Figure 1. First Three Stages of BLAST Execution

• Stage one and two are most time-consuming phases, taking 75% of execution time. Thus, our studies currently focus on these two stages.

cuBLASTP: Fine-Grained Parallelization of Protein Sequence Search on a GPU

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Results

- the overall program execution.
- performance.



Chart 2. Overall speedup of cuBLASTP over FSA-BLASTP. 4t means the program runs with 4 threads; F means the program runs on NVIDIA Fermi GPU; K means the program runs on NVIDIA Kepler GPU.

Conclusions

- We propose cuBLASTP, an efficient fine-grained BLASTP for GPU using the CUDA programming model.
- In cuBLASTP, we decoupled most time-consuming stages hit detection and ungapped extension - into separate kernels to apply multiple strategies on different memory access patterns, and added an additional stage: sorting and filtering, to reorganize intermediate results.
- cuBLASTP has up to 7.8x speedup over FSA-BLAST on a single core and 2.9x speedup over NCBI-BLAST on a quad-core CPU for the critical phases, and up to 4.7x speedup and 3x speedup for the overall performance, respectively.
- Compared with GPU-BLAST, the existing fastest BLAST on GPU: GPU-BLAST, cuBLASTP has up to 2.8x speedup for the critical phases and up to 1.8x speedup for the overall performance.

References

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