G-BLASTN: accelerating nucleotide alignment by graphics processors
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ABSTRACT
Motivation: Since 1990, the Basic Local Alignment Search Tool (BLAST) has become one of the most popular and fundamental bioinformatics tools for sequence similarity searching, receiving extensive attention from the research community. The two pioneering papers on BLAST have received over 96,000 citations. Given the huge population of BLAST users and the increasing size of sequence databases, an urgent topic of study is how to improve the speed. Recently, graphics processing units (GPUs) have been widely used as low-cost, high-performance computing platforms. The existing GPU-BLAST is a promising software tool that uses a GPU to accelerate protein sequence alignment. Unfortunately, there is still no GPU-accelerated software tool for BLAST-based nucleotide sequence alignment.

Results: We developed G-BLASTN, a GPU-accelerated nucleotide alignment tool based on the widely used NCBI-BLAST. G-BLASTN can produce exactly the same results as NCBI-BLAST, and it has very similar user commands. Compared with the sequential NCBI-BLAST, G-BLASTN can achieve an overall speedup of 14.80X under megablast mode. More impressively, it achieves an overall speedup of 7.15X over the multithreaded NCBI-BLAST running on 4 CPU cores. When running under blastn mode, the overall speedups are 4.32X (against 1-core) and 1.56X (against 4-core). G-BLASTN also supports a pipeline mode that further improves the overall performance by up to 44% when handling a batch of queries as a whole.

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1 INTRODUCTION
BLAST is one of the most fundamental software tools in bioinformatics for matching biological sequences (Altschul et al., 1990; Altschul et al., 1997). Due to the explosive growth of sequence data, improving the speed of BLAST has become increasingly critical. In the last decade, many attempts have been made to design and develop new BLAST software tools for specific hardware (Jacob et al., 2007; Sotiriades and Dallas, 2007; Fei et al., 2008) or even parallel supercomputers (Lin et al., 2008). Unfortunately, most researchers do not have access to these hardware platforms. Following the popularity of multicore processors, several BLAST software tools using multiple CPU cores for increased speed have been developed. One good example is the widely used National Center for Biotechnology Information (NCBI) BLAST, which supports multithreading in the preliminary stage of the BLAST algorithm (Camacho et al., 2009). Our experiments on a server with two quad-core Intel Xeon CPUs show that the multithreaded NCBI-BLAST can achieve an average speedup of 3~4X over the sequential version. NCBI-BLAST also supports an indexed MegaBLAST module, which uses the database index to achieve an approximate speedup of 2~4X (Morgulis, 2008). PLAST is a parallel implementation of BLAST (Nguyen and Lavenier, 2009) that applies a new indexing technique together with SSE instructions and multithreading to achieve better alignment speed. At present, PLAST only supports protein sequence alignment.

In recent years, Graphics Processing Units (GPUs) have been widely accepted as low-cost, high-performance computing platforms (Owens et al., 2008). Compared with traditional multi-core CPUs, GPUs have much higher computational horsepower and memory bandwidth. Many bioinformatics tools have been accelerated by GPUs in recent years (Manavski and Valle, 2008; Dematte and Prandi, 2010; Liu et al., 2012; Lu et al., 2012; Lu et al., 2013). The significant difference between GPU and CPU architectures has created many challenges in developing highly efficient GPU software (Nickolls, 2007). Without the development of carefully designed parallel algorithms and sophisticated optimizations, the huge potential of GPUs may not be fully realized.

Some GPU-based software tools have been developed for protein sequence alignment. Ling’s GPU-based BLAST software can achieve a speedup of 1.7~2.7X, compared with NCBI-BLAST (Ling and Benkrid, 2010). Recently, Vouzis and Sahinidis developed GPU-BLAST, which can typically achieve acceleration speedup of 3~4X relative to the sequential NCBI-BLAST (Vouzis and Sahinidis, 2011). The major advantage of GPU-BLAST is that it can produce the same results as NCBI-BLAST. Remarkably, this work was one of the top-ten highest downloaded articles published in Bioinformatics in 2011.

To the best of our knowledge, we are the first to provide an open-source GPU solution, namely G-BLASTN, for nucleotide sequence alignment that can produce the same results as NCBI-BLAST. G-BLASTN is developed on top of the NCBI-BLAST source code. It currently supports the megablast and blastn modes of NCBI-BLAST. Hereafter, we use BLASTN to refer to the nucleotide blast module of NCBI-BLAST, for simplicity. The major idea behind G-BLASTN is to store a small hash table in the fast GPU cache memory and then scan the DNA database in parallel using all of the available GPU cores. We have overcome several challenges to fully use the GPU horsepower. To achieve significant speedup, some other parts of BLASTN have also been optimized. We evaluate G-BLASTN’s performance by running a set of exper-
imments on human and mouse genome databases, as well as a partial of the NCBI nucleotide collection (nt) database. Using a contemporary NVIDIA GTX780 GPU with a cost of $650, G-BLASTN under megablast mode can achieve significant speedups over the multithreaded BLASTN running on 4-core or 8-core CPUs. When running under the more sensitive blastn mode, G-BLASTN also achieves reasonable speedups. When processing a batch of queries, G-BLASTN supports a pipeline mode that can further improve the performance by up to 44%. We believe that G-BLASTN is an attractive and cost-effective option for bioinformatics practitioners. The remainder of the article is organized as follows. In Section 2, we briefly review the main algorithms of BLASTN and present the design for G-BLASTN. In Section 3, we present the detailed implementation of G-BLASTN. In Section 4, we present the experimental results. We conclude the article in Section 5.

2 METHODS

2.1 BLASTN algorithms

BLASTN is designed to efficiently search nucleotide databases using a nucleotide query sequence (Camacho et al., 2009). BLASTN’s high level pseudocode is given in Figure 1, which consists of four stages. The setup stage prepares search options, reads and prepares the query sequence and database sequence and builds the lookup table. The scanning stage performs a preliminary search comprising three steps: seeding, ungapped extensions and gapped extensions. The seeding step scans the database for hits (i.e., a match with some word in the lookup table). The hits are then extended by ungapped alignment. The alignments that exceed a threshold score will go through the gapped extensions. Only the gapped alignments that exceed another threshold score will be saved as “preliminary” matches. The trace-back stage takes the preliminary matches as input, considering ambiguous nucleotides, and finds the locations of insertions and deletions. The output stage displays the alignment results to the user.

1 [Setup] prepare the BLASTN options, query, database, lookup table
2 [Scanning] for each of N threads {
3 2.1 while the database still has unsearched sequences {
4 2.2 Retrieve a group of sequences from the database
5 2.3 Seeding: find exact word matches
6 2.4 Ungapped extensions
7 2.5 Gapped extensions
8 2.6 }
9 2.7 }
10 [Trace-back] for each database sequence containing alignments, perform trace-back
11 [Output] print the alignment results

Fig. 1. High Level Pseudocode of BLASTN

BLASTN’s efficiency relies on the assumption that any alignment of interest between the query and the database will contain at least one W-gram (i.e., a subsequence of length W), where W is a parameter known as “BLASTN word size”. In practice, for any given query sequence BLASTN will construct a lookup table that stores the offsets into the query where each possible w-gram occurs, where w is a parameter known as the “lookup table word size” which is less than or equal to W. Because each letter can be one of \{A, C, G, T\}, the lookup table has 4w entries. The seeding procedure walks through the database sequence to find hits. In each round, it fetches a w-gram, calculates its hash value, looks into the lookup table and records all matched offset pairs (i.e., the pair of offsets of the matched w-gram in the query and database, respectively) if there are any. When W is larger than w, it is not necessary to scan the database letter by letter; instead, BLASTN scans the database in strides. The maximum stride size without missing any match is W-w+1. For extremely long nucleotide databases, the seeding procedure is usually the most time-consuming step.

After all w-gram hits have been found, we must determine whether each hit belongs to a W-gram match. This is done through the mini-extension procedure (a.k.a. exact match extension), which extends each w-gram in both the left and right directions to check the existence of exact W-gram matches. The mini-extension step can be time consuming if millions of w-gram hits must be extended. Once we find all of the W-gram hits, the ungapped extension step begins, allowing for mismatches. Ungapped alignments that exceed a threshold score are stored for gapped extension. In the scanning stage, gapped extension returns only the score and extent of the alignment while the number and position of insertions, deletions and matching letters are not stored. During the whole scanning stage, BLASTN processes the sequence in NCBI-NA2 format, in which each nucleic acid is represented by two bits. Hence, ambiguities cannot be handled. In the trace-back stage, ambiguous nucleotides are restored by converting NCBI-NA2 format into NCBI-NA8, and more sensitive heuristic parameters are used for the final gapped alignment. Finally, the output step formats the results according to the user options and prints the results for the user.

2.2 Design of G-BLASTN

GPUs have become mature, many-core processors with much higher computational power and memory bandwidth than today’s CPUs. A GPU consists of a scalable number of streaming multiprocessors (SMs), each containing some streaming processors (SPs), special function units (SFUs), a multithreaded instruction fetch and issue unit, registers and a read/write shared memory. CUDA is currently the most popular programming model for general purpose GPU computing. The best way to use the hundreds to thousands of GPU cores is to generate a large number of CUDA threads that can access data from multiple memory spaces during their execution, as illustrated in Figure 2. Each thread has its private registers and local memory. Each GPU kernel function generates a grid of threads that are organized into thread blocks. Each thread block has shared memory visible to all threads within the block and with the same lifetime as the block. All threads have access to the same global memory. Two additional read-only memory spaces are accessible by all threads: the constant and texture memory spaces, both of which have limited caches.

Due to the complexity of BLASTN software, exploiting GPUs to accelerate BLASTN is a non-trivial task. The main challenge is that not all of the steps involved in BLASTN are suitable to be parallelized by GPUs. To identify which steps should be parallelized, we conducted a profiling study by running 300 different queries with a broad range of lengths against the human build 36 genome database to analyze the time distribution of differ-
ent BLASTN steps under megablast mode (Figure 3). We mainly observed the following details. The scanning stage is the most time-consuming and accounts for 69-93% of the total execution time. Surprisingly, BLASTN spends 5-25% of the total execution time in the setup stage, mainly initializing the mask database. The trace-back stage takes negligible time for most queries, but can occasionally take a very long time.

To achieve a good overall speedup, we designed G-BLASTN as follows. Its major component is a set of CUDA kernel functions that run on GPUs to significantly accelerate the seeding and mini-extension steps in the scanning stage. It is designed to initialize the mask database once and then serve a large number of queries. Therefore, the time spent in database initialization can be largely removed. We optimized the two most time consuming functions in the trace-back stage and further designed a pipeline mode under which the trace-back, output and scanning stages can run simultaneously. The general framework of G-BLASTN is shown in Figure 4.

3 IMPLEMENTATION

We use CUDA C language to implement G-BLASTN based on NCBI BLAST 2.2.28 software package. It supports both Windows and Linux platforms. In the following, we present the detailed implementation of the major modules of G-BLASTN.

3.1 Accelerating the seeding step by GPU

The main task of the seeding step is to scan the database sequences and identify all w-gram matches. Due to the large database sizes, the seeding step is the most time consuming in BLASTN. Fortunately, there is a good chance that the seeding step can be parallelized due to the independence of the tasks at different offsets of the database. G-BLASTN first loads the database sequences to GPU global memory. Then for each query sequence, it stores a copy of the lookup table in GPU texture memory to achieve ultrafast table lookup. For each database sequence, it invokes a GPU kernel function that generates a large number of GPU threads to scan the database sequence in parallel; and hence the large number of GPU cores can be fully utilized to speed up the seeding step.

The implementation of the seeding step on the GPU is a major challenge, however. In CUDA, each thread block is organized as a number of warps, and each warp of threads is executed by a single instruction, multiple data (SIMD) hardware. When threads within a warp take different execution paths, the SIMD hardware will take multiple runs to go through these divergent paths, which will significantly decrease the utilization of GPU cores. In the case of BLASTN, the w-grams at different offsets of the database sequence may have no match or many matches to the query sequence, which can lead to severe thread branch divergence that decreases the GPU performance significantly. To conquer this challenge, we divide the seeding step into two sub-steps: scan and lookup. In the scan sub-step, we go through the whole database sequence in parallel and record all offsets of the database that have at least one match to the query. Notice that we do not need to know how many matches have been found and where they are for each offset. Thus, each GPU thread can perform almost the same execution path and the effect of thread branch divergence can be minimized. In the lookup sub-step, we use another GPU kernel function to recheck all matched offsets and construct the complete set of matched offset pairs. This strategy works very well because the scan sub-step dominates the time of the seeding step.

There is yet another challenge in efficiently implementing the scan sub-step on the GPU. Once a thread finds a w-gram match, it has to increase a global counter and then write the matched offset into a global array. There are two negative consequences: (1) increasing the global counter must be an atomic operation, which means only one among all threads can operate while others have to wait; and (2) writing a single offset pair into the global array can waste a lot of GPU memory bandwidth. To overcome this challenge, we use a local counter and a local array for each thread block as temporary storage for the global counter and global array. The local counter and array are held in GPU shared memory, which is very fast. Now all thread blocks can operate on their own local counters and arrays simultaneously, boosting the overall performance. Once a local array becomes full, the set of offset pairs is written into global array as a whole and the global counter is updated by an atomic operation. We exploit coalesced memory write operations to achieve very high memory bandwidth. Meanwhile, the number of atomic operations on the global counter can be significantly reduced. The framework of the scan sub-step on GPU is shown in Figure 5.

For performance consideration, BLASTN supports two types of lookup tables for different types of queries: small and megablast\(^1\). Each type of lookup table has its own set of algorithms. Therefore, we have to implement different GPU kernel functions for different types of lookup tables.

\(1\) In current NCBI-BLAST, both types of lookup tables are supported by blastn mode and megablast mode.
The megablast lookup table comprises three arrays: presence vector (PV array), hash table (hashTable[]) and next position (nextPos[]). The PV array is a bit field with one bit for each hash table entry. If a hash table entry contains a query offset, the corresponding bit in the PV array is set. The scanning process first checks the PV array to see whether there are any query offsets in a particular lookup table entry. The hashTable[] array is a thick backbone with one word for each of the lookup table entries. If a lookup table entry has no query offsets, the corresponding entry in hashTable[] is zero; otherwise, it is an offset into nextPos[]. The position in nextPos[] is in fact the query offset, and the actual value at that position is a pointer to the succeeding query offset in the chain. A value of zero means the end of the chain. The pseudocode of our GPU scan and lookup kernel functions using the megablast lookup table is shown in Figures 8 and 9, respectively. The scan kernel function checks the PV array to quickly determine whether there is a match. To achieve the best table lookup performance, the PV array is held in texture memory. The lookup kernel function takes the output of scan function as input and checks the hashTable[] and nextPos[] to find the complete set of matched offset pairs.

3.2 Accelerating the mini-extension step by GPU

It is not uncommon for the scan sub-step to return millions of seed matches. The mini-extension step is designed to verify whether each w-gram match can be extended to a W-gram match when w < W. We can create a huge number of GPU threads to extend those w-gram matches simultaneously. Each GPU thread reads one offset pair from the matched offset pair array, extends on the left side and then extends on the right side. If it finds a W-gram match, this offset pair will be recorded for further gapped extension. Given that the mini-extension algorithm exhibits no big difference from the original BLASTN, we do not provide the pseudocode here. We note that there are two versions of mini-extension, one for the small lookup table and another for the megablast lookup table.
3.3 Optimizing the trace-back step

As mentioned in Section 2.2, occasionally the trace-back step takes quite a long time, which may counteract the speedup achieved by the previous steps. Unfortunately, the trace-back step is not naturally suitable for GPUs. Therefore, we resort to the following optimization techniques. First, function $s\_SeqDBMap\_NA2\_To\_NA8()$ uses a translation table to convert sequence data from NCBI-NA2 to BLAST-NA8 format. BLAST translates the data character by character, which does not fully use the CPU memory bandwidth. In G-BLASTN, we replace four 8-bit memory writes with a single character by character. Therefore, the speedup of 3–4X. The SSE instructions are shown in Figure 10.

3.4 Pipeline mode for multiple queries

Once we have accelerated the scanning stage by GPU, other stages such as trace-back and output may start to occupy a relatively large portion of the total execution time, especially when there are many final hits. G-BLASTN supports a pipeline mode when handling a batch of queries. The main advantage of the pipeline mode is that the GPU and CPU can work on different tasks simultaneously, as shown in Figure 11. In short, when the GPU is busy with seeding or main-extension, the CPU can execute the trace-back or output steps for a previous query. To achieve this purpose, G-BLASTN uses multithreading to maintain four queues: query, job, prelim and result. A master thread reads the queries and puts them into the query queue, and then creates the job queue. The job(s) fetches jobs from the job queue and uses GPU to execute the preliminary search, storing the results in the prelim queue. The trace-back thread(s) reads from the prelim queue, executes the trace-back step and stores the results in the results queue. Finally, the print thread prints the results. This pipeline design can efficiently use both GPU and CPU resources.

4 RESULTS

4.1 General setup and data sets

The GPU experiments were performed on a desktop computer with an Intel quad-core CPU and Nvidia GTX780 GPU. The CPU experiments were performed on two different platforms: a 4-core platform which is the same computer that runs the GPU experiments; and an 8-core platform which is a server with two Intel Xeon CPUs. The detailed system configuration is shown in Table 1.


efficient use both GPU and CPU resources.

![Fig. 11. The pipeline mode of G-BLASTN](image)

![Table 1 System configuration](table)

![Diagram](diagram)
We used the following two command lines to run NCBI BLASTN and G-BLASTN, respectively. More details about the command options of G-BLASTN are found on our website.

```bash
$blastn -db <database> -query <query> -task megablast|blastn -outfmt 7 -out <file> -dust yes -window_masker_db <masker_db> -num_threads <1|2|4|8>

$gblastn -db <database> -query_list <query list> -task megablast|blastn -outfmt 7 -out <file> -dust yes -window_masker_db <masker_db> -use_gpu true -mode <1|2> -num_threads <1|4|8>
```

We used `gettimeofday()` functions to measure the program execution time. Each experiment was run 10 times and the average results are reported in this paper.

**Databases.** We chose human build 36 and mouse build 36 genome databases for the experiments of megablast mode. In addition, we constructed a database to test the blastn mode by selecting all sequences with length no less than 2 million from NCBI nt database. This partial NCBI nt database has a raw size of 8.4 GB and can well fit into a single GPU card with 3 GB memory after compression. All databases were masked with WindowMasker (Morgulis et al., 2006a), including low-complexity filtering by DUST (Morgulis et al., 2006b). The length information of all database sequences can be found in Figure S2 of the Supplementary.

**Queries.** To test the megablast mode, we chose queries from the NCBI ftp server: ftp://ftp.ncbi.nlm.nih.gov/pub/agrawal/indexed_megablast/queries (Morgulis 2008). Six query sets, each containing 100 queries, were used, which are referred to as Qsmall (~500 bases, range: 501-506), Qmedium (~10 Kbases, range: 10000-10446) and Qlarge (~100 Kbases, range: 100001-102087). To test the blastn mode, we chose the first 500 bacterial sequences from the NCBI server: http://www.ncbi.nlm.nih.gov/sra/SRX338063, namely Qbac. The length information of all query sequences can be found in Figure S1 of the Supplementary.

### 4.2 Experimental results

#### 4.2.1 Performance under normal mode

Under normal mode, G-BLASTN handles the queries one at a time. We first present the experimental results of megablast. The speedups over 8-core platform on human genome database are shown in Figure 12. The speedups of other experiments are shown in Figures S3-S6 of the Supplementary. We also show the average speedup of each query set in Table 2. The overall speedups are calculated as the average of all 1600 query experiments for each hardware setting. As compared with 4-core Intel i7-3820, G-BLASTN achieves an overall speedup of 7.15X. As compared with the 8-core platform, G-BLASTN achieves an overall speedup of 13.76X. There are several reasons why BLASTN runs much faster on i7-3820 than on Xeon E5620. Firstly, i7-3820 has a much higher working frequency than E5620. Secondly, the memory bandwidth of i7-3820 is twice of E5620. Thirdly, the memory module of our i7-3820 platform is faster than that of E5620 platform. Based on the results of E5620, we can notice that the speedups achieved using 8 cores are only slightly better than using 4 cores.

We also notice that the speedups on the human database are less than those on the mouse database. This is mainly because the human build 36 database consists of 367 sequences, more than 200 of which are short sequences (less than 1 million bps). In contrast, the mouse build 36 database consists of 21 very long sequences. We will discuss more on this issue in Section 5.

<table>
<thead>
<tr>
<th>Database</th>
<th>Query</th>
<th>1-core</th>
<th>4-core</th>
<th>1-core</th>
<th>4-core</th>
<th>8-core</th>
</tr>
</thead>
<tbody>
<tr>
<td>Human</td>
<td>Qsmall</td>
<td>10.47</td>
<td>5.11</td>
<td>26.90</td>
<td>12.88</td>
<td>10.52</td>
</tr>
<tr>
<td></td>
<td>Qmedium</td>
<td>11.49</td>
<td>4.53</td>
<td>32.68</td>
<td>11.98</td>
<td>8.50</td>
</tr>
<tr>
<td></td>
<td>Qlarge</td>
<td>9.22</td>
<td>3.37</td>
<td>21.07</td>
<td>7.54</td>
<td>5.25</td>
</tr>
<tr>
<td></td>
<td>Qbac</td>
<td>10.80</td>
<td>5.37</td>
<td>26.04</td>
<td>12.71</td>
<td>10.47</td>
</tr>
<tr>
<td>Mouse</td>
<td>Qsmall</td>
<td>18.50</td>
<td>9.37</td>
<td>44.12</td>
<td>21.87</td>
<td>18.28</td>
</tr>
<tr>
<td></td>
<td>Qmedium</td>
<td>17.84</td>
<td>7.39</td>
<td>49.16</td>
<td>19.11</td>
<td>14.32</td>
</tr>
<tr>
<td></td>
<td>Qlarge</td>
<td>10.44</td>
<td>4.14</td>
<td>23.16</td>
<td>9.14</td>
<td>6.92</td>
</tr>
<tr>
<td></td>
<td>Qbac</td>
<td>20.97</td>
<td>10.73</td>
<td>49.28</td>
<td>24.71</td>
<td>20.80</td>
</tr>
</tbody>
</table>

**Table 2** Average speedup of G-BLASTN under megablast mode

We evaluate the performance of blastn mode using Qbac query set against human, mouse, and the partial NCBI nt databases. We show the average speedups of each database and the overall speedups in Table 3. G-BLASTN achieves an overall speedup of 1.56 and 2.95 as compared with 4-core i7-3820 and 8-core E5620, respectively. As compared with megablast mode, blastn mode has a much smaller value of stride size, which results in more scanning workload and more seed hits. Therefore the ungapped extension step under blastn mode takes much longer time than under megablast mode, as shown in Figure S7 of the Supplementary. Since the ungapped extension is sequentially executed on CPU, the speedups achieve under blastn mode are much less than megablast.
4.2.2 Performance under pipeline mode

To evaluate the performance of pipelined G-BLASTN, we use all queries in each data set as a single input to G-BLASTN. The speedups against the NCBI BLAST on Intel i7-3820 are shown in Table 4. If we compare Table 4 with Tables 2 & 3, we can observe a significant improvement on the speedups for many data sets. For small and medium queries under megablast, the trace-back and output steps account for a very small portion of the total time, and hence the pipeline design does not offer much of an advantage. For large queries using megablast, however, the trace-back and output steps take a much longer time due to a greater number of final hits, and thus the pipeline design hides a significant portion of time. Under blastn mode, the pipeline design can further improve the speedups of the Qbac query set by 19-44%.

Table 3 Speedup of G-BLASTN under blastn mode using Qbac query set

<table>
<thead>
<tr>
<th>Database</th>
<th>Intel i7-3820</th>
<th>Intel Xeon E5620</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>1-core</td>
<td>4-core</td>
</tr>
<tr>
<td>Human</td>
<td>4.58</td>
<td>1.57</td>
</tr>
<tr>
<td>Mouse</td>
<td>5.02</td>
<td>1.83</td>
</tr>
<tr>
<td>NCBI nt</td>
<td>3.37</td>
<td>1.29</td>
</tr>
<tr>
<td>Overall</td>
<td>4.32</td>
<td>1.56</td>
</tr>
</tbody>
</table>

4.2.3 Discussion and Conclusions

In this paper, we describe our design and implementation of G-BLASTN, an open source software tool for nucleotide alignment based on the widely used NCBI-BLAST. G-BLASTN exploits the power of GPUs to accelerate nucleotide alignments. Compared with a contemporary quad-core Intel CPU running at 3.6GHz, G-BLASTN on a single $650$ GPU card can achieve overall speedups of $14.8X$ and $4.32X$ under megablast mode and blastn mode respectively. When compared with multithreaded NCBI-BLAST that uses four CPU cores, G-BLASTN can still achieve overall speedups of $7.15X$ (megablast) and $1.56X$ (blastn). G-BLASTN also supports a pipeline mode that further improves the overall performance by up to $44\%$ when handling multiple queries.

G-BLASTN can be improved in the following directions. At present, G-BLASTN invokes a kernel function for each database sequence, which is not efficient when the length of the database sequence is shorter than one million bps. Besides the constant overhead of invoking a kernel function, another reason is that short database sequence only generates a small number of hits during the scanning stage, and hence the effect of our optimization strategy on the atomic operations will diminish. There are several possible solutions to this problem. One possibility is to aggregate short database sequences into longer ones. Another solution is to process multiple database sequences in each kernel function call. G-BLASTN is also limited by the GPU memory size. We plan to extend G-BLASTN to support multiple GPU cards. Doing so can not only support much larger databases, but also achieve better speedups. A more challenging task is to accelerate other steps such as ungapped extension, gapped extension, and trace-back, which will improve the performance of blastn mode significantly. Finally, we also plan to support discontiguous megablast mode in our future work.

Conflicts of Interest: none declared.

REFERENCES