Modification of Valiant's Parsing Algorithm for the String-Searching Problem*

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Abstract. Some string-matching problems can be reduced to parsing: verification whether some sequence can be derived in the given grammar. To apply parser-based solutions to such area as bioinformatics, one needs to improve parsing techniques so that the processing of large amounts of data was possible. The most asymptotically efficient parsing algorithm that can be applied to any context-free grammar is a matrix-based algorithm proposed by Valiant. This paper presents a modification of the Valiant's algorithm, which facilitates efficient utilization of modern hardware in highly-parallel implementation. Moreover, the modified version significantly decreases the number of excessive computations, accelerating the search of substrings.

Keywords: Context-free grammar \cdot Parsing \cdot Valiant's algorithm \cdot Stringmatching \cdot Secondary structure.

1 Introduction

The secondary structure of RNA is tightly related to biological functions of organisms and plays an important role in classification and recognition problems. One of the approaches to analyze the secondary structure of RNA is based on formal language methods. Namely, one can process RNA sequence as a string over 4-letter alphabet $\{G, A, C, U\}$ and use formal language methods to describe properties of this string and then analyze strings w.r.t. described properties by parsing them.

Context-free languages (CFL-s) are the most prominent in this area, while probabilistic context-free grammars are widely used for secondary structure description and related tasks [3,8]. But more expressive language classes are required to describe some important features of secondary structure. For example,

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pseudoknots cannot be expressed in terms of a context-free grammar, but can be expressed with a conjunctive grammar [14] which were proposed by Alexander Okhotin [10] and are the natural extension of CFG.

For some problems, it is necessary to find all derivable substrings of a given string [4]. This case is the string-matching problem also known as a string-searching problem. The classical example of it is to find substrings matched by the given regular expression (or regular template). But if one tries to find a substring with a specific secondary structure, then it is necessary to use at least context-free template (context-free grammar) and, as a result, utilize an appropriate parsing algorithm.

Most CFG-based approaches suffer the same issue: the computational complexity is poor. Traditionally used CYK [7, 13] runs with a cubic time complexity and demonstrates poor performance on long strings or big grammars [9]. We argue that more efficient algorithms are needed in fields such as bioinformatics where large amount of data is common.

Asymptotically most efficient parsing algorithm is Valiant's algorithm [12] which is based on matrix multiplication. Okhotin generalized this algorithm to conjunctive and boolean grammars [11]. Moreover, in comparison to CYK, Valiant's algorithm simplifies the utilization of parallel techniques to improve performance by offloading critical computations onto matrices multiplication. However, this algorithm is not suitable for the string-matching problem because it computes a huge amount of redundant information.

In this paper we present the modification of Valiant's algorithm which improves the utilization of GPGPU and parallel computations by processing some submatrices products concurrently. Also, the proposed algorithm can be easily utilized for the string-matching problem. We also prove the correctness of our algorithm and analyze its time complexity. The proposed solution was implemented using fast matrix multiplication algorithms and parallel techniques. Evaluation of the implementation shows that GPGPU version is up to 20 times faster then GPGPU-based implementation of the Valiant's algorithm.

2 Background

We start by introducing some basic definitions from the formal language theory. Using these definitions, we describe Valiant's parsing algorithm on which we base our modification.

2.1 Formal Languages

An alphabet Σ is a finite nonempty set of symbols. Σ^* denotes the set of all finite strings over Σ . A context-free grammar G_S is a quadruple (Σ, N, R, S) , where Σ is a finite set of terminals, N is a finite set of nonterminals, R is a finite set of productions of the form $A \to \beta$, where $\Sigma \cap N = \emptyset$, $A \in N$, $\beta \in V^*$, $V = \Sigma \cup N$ and $S \in N$ is a start nonterminal. Context-free grammar $G_S = (\Sigma, N, R, S)$ is said to be in Chomsky normal form if all productions in R are of the form:

 $A \to BC, \ A \to a$, or $S \to \varepsilon$, where $A, B, C \in N, a \in \Sigma, \varepsilon$ is an empty string. For each context-free grammar G of length N one can construct an equivalent grammar in Chomsky normal form with length N^2 [6].

 $L_G(S) = \{\omega \mid S \xrightarrow{*}_{G_S} \omega\}$ is a language specified by the grammar $G_S = (\Sigma, N, R, S)$, where $S \xrightarrow{*}_{G_S} \omega$ means that ω can be derived in a finite number of rules applications from the start symbol S.

2.2 Valiant's Parsing Algorithm

Tabular parsing algorithms construct a matrix T, cells of which are filled with nonterminals from which the corresponding substring can be derived. These algorithms usually work with the grammar in Chomsky normal form. For $G_S = (\Sigma, N, R, S)$, $T_{i,j} = \{A \mid A \in N, a_{i+1} \dots a_j \in L_G(A)\} \quad \forall i < j$.

The parsing matrix T are filled successively starting with diagonal elements $T_{i-1,i} = \{A \mid A \to a_i \in R\}$. Then, $T_{i,j} = f(P_{i,j})$, where $P_{i,j} = \bigcup_{\substack{i < k < j \\ i < k < j}} T_{i,k} \times T_{k,j}$ (here, \times is the Cartesian product of two sets) and $f(P) = \{A \mid \exists A \to BC \in R : (B,C) \in P\}$. Finally, the input string $a_1a_2 \dots a_n$ belongs to $L_G(S)$ iff $S \in T_{0,n}$. If all cells are filled sequentially, the time complexity of this algorithm is $O(n^3)$. Valiant proposed to offload the most intensive computations to the Boolean matrix multiplication. The most time-consuming is computing $\bigcup_{\substack{i < k < j \\ i < k < j}} T_{i,k} \times T_{k,j}$ and Valiant's idea is to compute $T_{i,j}$ by multiplication of submatrices of T.

Multiplication of two submatrices of parsing table T is defined as follows. Let $X \in (2^N)^{m \times l}$ and $Y \in (2^N)^{l \times n}$ be two submatrices of the parsing table T. Then, denote $X \times Y = Z$, where $Z \in (2^{N \times N})^{m \times n}$ and $Z_{i,j} = \bigcup_{1 \le k \le l} X_{i,k} \times Y_{k,j}$.

Note that the computation of $X \times Y$ can be replaced by the $|N|^2$ multiplication of |N| Boolean matrices (for each nonterminal pair). Denote the matrix corresponding to the pair $(B,C) \in N \times N$ as $Z^{(B,C)}$, then $Z^{(B,C)}_{i,j} = 1$ iff $(B,C) \in Z_{i,j}$. It should also be noted that $Z^{(B,C)} = X^B \times Y^C$. Each Boolean matrix multiplication can be computed independently. Following these changes, time complexity of this algorithm is $O(|G|\operatorname{BMM}(n)\log(n))$ for an input string of length n, where $\operatorname{BMM}(n)$ is the number of operations needed to multiply two Boolean matrices of size $n \times n$.

Valiant's algorithm written as described by Okhotin is presented in Listing 1. All elements of T and P are initialized by empty sets. Then, the elements of these two table are successively filled by two recursive procedures.

The procedure compute(l, m) computes values of $T_{i,j}$ for all $l \leq i < j < m$. The procedure complete(l, m, l', m') constructs the submatrix $T_{i,j}$ for all $l \leq i < m$, $l' \leq j < m'$. This procedure assumes $T_{i,j}$ for all $l \leq i < j < m$, $l' \leq i < j < m'$ are already constructed and the current value of

$$P_{i,j} = \{(B,C) \mid \exists k, (m \le k < l'), a_{i+1} \dots a_k \in L(B), a_{k+1} \dots a_j \in L(C)\}$$

for all $l \le i < m, l' \le j < m'$. The submatrix partition during the procedure call is shown in Figure 1.

Listing 1: Parsing by Matrix Multiplication: Valiant's Algorithm

```
Input: Grammar G = (\Sigma, N, R, S), w = a_1 \dots a_n, n \ge 1, a_i \in \Sigma, where n + 1 = 2^p
     main():
        compute(0, n + 1);
 2
        accept iff S \in T_{0,n}
 3
 4 compute(l, m):
 5 if m-l \geq 4 then
            compute(l, \frac{l+m}{2});
    compute(\frac{l+m}{2}, m)
complete(l, \frac{l+m}{2}, \frac{l+m}{2}, m)
complete(l, m, l', m'):
10 if m - l = 4 and m = l' then T_{l,l+1} = \{A \mid A \to a_{l+1} \in R\};
11 else if m - l = 1 and m < l' then T_{l,l'} = f(P_{l,l'});
     else if m-l>1 then
           \begin{array}{l} \textit{leftgrounded} = (l, \frac{l+m}{2}, \frac{l+m}{2}, m), \textit{rightgrounded} = (l', \frac{l'+m'}{2}, \frac{l'+m'}{2}, m'), \\ \textit{bottom} = (\frac{l+m}{2}, m, l', \frac{l'+m'}{2}), \textit{left} = (l, \frac{l+m}{2}, l', \frac{l'+m'}{2}), \end{array}
13
14
           right = (\frac{l+m}{2}, m, \frac{l'+m'}{2}, m'), top = (l, \frac{l+m}{2}, \frac{l'+m'}{2}, m');
15
            complete(bottom);
16
            P_{left} = P_{left} \cup (T_{leftgrounded} \times T_{bottom});
17
            complete(left);
18
            P_{right} = P_{right} \cup (T_{bottom} \times T_{rightgrounded});
19
            complete(right);
20
            P_{top} = P_{top} \cup (T_{leftgrounded} \times T_{right});
\mathbf{21}
            P_{top} = P_{top} \cup (T_{left} \times T_{rightgrounded});
\mathbf{22}
            complete(top)
23
```

A simple example of the several first steps of Valiant's algorithm execution is presented in Figure 3. Only several steps are shown, but it is enough to compare our version with the original algorithm.

3 Modified Valiant's Algorithm

In this section we propose a way to rearrange submatrices handling in the algorithm. The different order improves the independence of submatrices handling and facilitates the implementation of parallel submatrix processing.

3.1 Layered Submatrices Processing

We propose to divide the parsing table into layers of disjoint submatrices of the same size (see Figure 2). Such division is possible because the derivation of a substring of the fixed length does not depend on either left or right contexts. Each layer consists of square matrices which size is a power of 2. The layers are computed successively in the bottom-up order. Each matrix in the layer can be handled independently, which facilitates parallelization of layer processing.

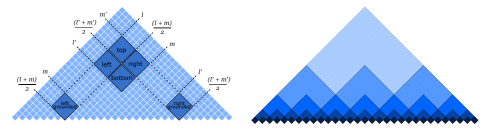


Fig. 1. Matrix partition used in procedure complete(l, m, l', m')

Fig. 2. Matrix partition on V-shaped layers used in modification

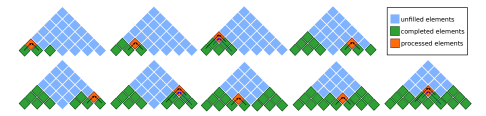


Fig. 3. An example of the first steps of Valiant's algorithm

Figure 4 demonstrates the modified algorithm. The lowest layer (submatrices of size 1) has already been computed. The second layer is filled in during steps 1–2. So the same part of parsing matrix (as in Figure 3) can be computed in just two steps using parallel computation of submatrix products.

The modified version of Valiant's algorithm is presented in Listing 2. The procedure main() computes the lowest layer $(T_{l,l+1})$, and then divides the table into layers, and computes them with the completeVLayer() function. Thus, main() computes all elements of parsing table T.

We define right grounded(subm), left grounded(subm), left (subm), right (subm), top(subm) and bottom(subm) functions which return the submatrices for matrix subm = (l, m, l', m') according to the original Valiant's algorithm (Figure 2).

The procedure complete V Layer(M) takes an array of disjoint submatrices M which represents a layer. For each $subm = (l, m, l', m') \in M$ this procedure computes left(subm), right(subm), top(subm). It is assumed in the procedure that the elements of bottom(subm) and $T_{i,j}$ for all i and j such that $l \leq i < j < m$ and $l' \leq i < j < m'$ are already constructed. Also it is assumed that the current value of $P_{i,j} = \{(B,C) \mid \exists k, (m \leq k < l'), a_{i+1} \dots a_k \in L_G(B), a_{k+1} \dots a_j \in L_G(C)\}$ for all i and j such that $l \leq i < m$ and $l' \leq j < m'$.

The procedure completeLayer(M) takes an array of disjoint submatrices M, but unlike the previous one, it computes $T_{i,j}$ for all $(i,j) \in subm$. This procedure requires the same assumptions on $T_{i,j}$ and $P_{i,j}$ as in the original algorithm.

In other words, completeVLayer(M) computes the entire layer M and $completeLayer(M_2)$ is a helper function which is necessary for computation of smaller square submatrices $subm_2 \in M_2$, where M_2 is a sublayer of M.

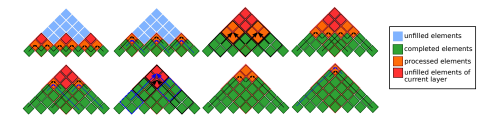


Fig. 4. Steps of the modification of Valiant's algorithm

Finally, the procedure performMultiplication(tasks), where tasks is an array of triples of submatrices, performs the basic step of the algorithm: matrix multiplication. It is worth mentioning that in this procedure $|tasks| \geq 1$ and each task can be computed independently, while the original algorithm handles one task per step sequentially. Thus, the practical implementation of this procedure can easily utilize different techniques of parallel array processing.

3.2 Correctness and Complexity

We provide the proof of correctness and time complexity of the proposed modification in this section.

Lemma 1. Let M be a layer. If for all $(l, m, l', m') \in M$:

- 1. $T_{i,j} = \{A \mid a_{i+1} \dots a_j \in L_G(A)\}$ for all i and j such that $l \leq i < j < m$ and $l' \leq i < j < m'$;
- 2. $P_{i,j} = \{(B,C) \mid \exists k, (m \leq k < l') : a_{i+1} \dots a_k \in L_G(B), a_{k+1} \dots a_j \in L_G(C)\}$ for all $l \leq i < m$ and $l' \leq j < m'$.

Then the procedure complete Layer(M), returns correctly computed sets of $T_{i,j}$ for all $l \leq i < m$ and $l' \leq j < m'$ for all $(l, m, l', m') \in M$.

Proof. We only sketch the proof of this lemma, as completeLayer() and completeVLayer() are two mutually recursive functions, so this lemma and the theorem 1 overlap and is proven similarly.

As basis we consider the matrix of size m-l=1 (and then the elements of layer M are correctly computed in lines 10-11) and thereafter the lemma can be proved by induction on m-l.

Theorem 1. Algorithm from listing 2 correctly computes $T_{i,j}$ for all i and j, thus an input string $a = a_1 a_2 \dots a_n \in L_G(S)$ if and only if $S \in T_{0,n}$.

Proof. Primarily to prove the theorem, we show by induction that all layers of the parsing table T are computed correctly.

Basis: layer of size 1×1 . Parsing table T consists of one layer of size 1 and its elements are correctly computed in lines 2-3 in listing 2.

<u>Inductive step:</u> assume any layer of size less than or equal to $2^{r-2} \times 2^{r-2}$ are computed correctly.

Listing 2: Parsing by Matrix Multiplication: Modified Version

```
Input: G = (\Sigma, N, R, S), w = a_1 \dots a_n, n \ge 1, n + 1 = 2^p, a_i \in \Sigma
   1 main():
   2 for l \in \{1, ..., n\} do T_{l,l+1} = \{A | A \rightarrow a_{l+1} \in R\};
   3 for 1 \le i \le p-1 do
                    layer = constructLayer(i);
                     complete VLayer(layer)
   6 accept iff S \in T_{0,n}
   7 constructLayer(i):
   8 \{(k \cdot 2^i, (k+1) \cdot 2^i, (k+1) \cdot 2^i, (k+2) \cdot 2^i) \mid 0 \le k < 2^{p-i} - 1\}
   9 completeLayer(M):
10 if \forall (l, m, l', m') \in M \quad (m - l = 1) then
                    for (l, m, l', m') \in M do T_{l,l'} = f(P_{l,l'});
11
12 else
13
                     completeLayer(\{bottom(subm) \mid subm \in M\});
                     complete VLayer(M)
15 completeVLayer(M):
16 multiplication Task_1 =
             \{left(subm), leftgrounded(subm), bottom(subm) \mid subm \in M\} \cup \{left(subm), leftgrounded(subm), bottom(subm), botto
             \{right(subm), bottom(subm), rightgrounded(subm) \mid subm \in M\};
17 multiplication Task_2 = \{top(subm), leftgrounded(subm), right(subm) \mid subm \in M\};
18 multiplicationTask_3 = \{top(subm), left(subm), rightgrounded(subm) \mid subm \in M\};
19 performMultiplications(multiplicationTask_1);
20 completeLayer(\{left(subm) \mid subm \in M\} \cup \{right(subm) \mid subm \in M\});
21 performMultiplications(multiplicationTask<sub>2</sub>);
22 performMultiplications(multiplication Task_3);
23 completeLayer(\{top(subm) \mid subm \in M\})
24 performMultiplications(tasks):
25 for (m, m1, m2) \in tasks do P_m = P_m \cup (T_{m1} \times T_{m2});
```

Define layer of size $2^{r-1} \times 2^{r-1}$ as M. Hereinafter subm = (l, m, l', m') is a typical element of layer M.

Consider completeVLayer(M) call.

First, $performMultiplications(multiplicationTask_1)$ adds to each $P_{i,j}$ all pairs (B,C) such that $\exists k, \ (\frac{l+m}{2} \leq k < l'), \ a_{i+1} \dots a_k \in L_G(B), \ a_{k+1} \dots a_j \in L_G(C)$ for all $(i,j) \in leftsublayer(M)$ and (B,C) such that $\exists k, \ (m \leq k < \frac{l'+m'}{2}), \ a_{i+1} \dots a_k \in L_G(B), \ a_{k+1} \dots a_j \in L_G(C)$ for all $(i,j) \in rightsublayer(M)$. Now $completeLayer(leftsublayer(M) \cup rightsublayer(M))$ can be called and it returns the correctly computed $leftsublayer(M) \cup rightsublayer(M)$.

Then performMultiplications called with arguments multiplication Task₂ and multiplication Task₃ adds pairs (B, C) such that $\exists k, (\frac{l+m}{2} \leq k < m), a_{i+1} \dots a_k \in$

 $L_G(B), a_{k+1} \dots a_i \in L_G(C)$ and pairs (B, C) such that $\exists k, (l' \leq k < \frac{l'+m'}{2}),$ $a_{i+1} \ldots a_k \in L_G(B), \ a_{k+1} \ldots a_j \in L_G(C)$ to each element $P_{i,j}$ for all $(i,j) \in$ topsublayer(M). So as m=l' (from the construction of the layer), condition for elements of matrix P are fulfilled. Now completeLayer(topsublayer(M)) can be called and it returns the correctly computed topsublayer(M).

All $T[i,j] \ \forall (i,j) \in M$ are computed correctly.

Thus, complete V Layer(M) returns correct $T_{i,j}$ for all $(i,j) \in M$ for any layer M of parsing table T and lines 4-6 in listing 2 return all $T_{i,j} = \{A \mid A \in N, \}$ $a_{i+1} \dots a_i \in L_G(A)$.

Lemma 2. Let calls, be a number of the calls of complete VLayer(M) where for all $(l, m, l', m') \in M$ with $m - l = 2^{p-r}$.

- for all $r \in \{1, \dots, p-1\}$ $\sum_{n=1}^{calls_r} |M|$ is exactly $2^{2r-1} 2^{r-1}$; for all $r \in \{1, \dots, p-1\}$ products of submatrices of size $2^{p-r} \times 2^{p-r}$ are calculated exactly $2^{2r-1} 2^r$ times.

Proof. Prove the first statement by induction on r.

<u>Basis:</u> r = 1. $calls_1$ and |M| = 1. So, $2^{2r-1} - 2^{r-1} = 2^1 - 2^0 = 1$.

Inductive step: assume that $\sum_{n=1}^{call_{r}} |M|$ is exactly $2^{2r-1} - 2^{r-1}$ for all $r \in \{1, ..., q\}.$

Let us consider r = q + 1.

Firstly, note that function costructLayer(r) returns $2^{p-r}-1$ matrices of size 2^r , so in the call of complete VLayer(costructLayer(p-r)) costructLayer(p-r)returns $2^r - 1$ matrices of size 2^{p-r} . Secondly, complete VLayer() is called 3 times for the left, right and top submatrices of size $2^{p-(r-1)}$. Finally, complete VLayer()is called 4 times for the bottom, left, right and top submatrices of size $2^{p-(r-2)}$, except $2^{r-2} - 1$ matrices which were already computed.

Then,

$$\begin{array}{l} \sum_{n=1}^{calls_r} |M| = 2^r - 1 + 3 \times (2^{2(r-1)-1} - 2^{(r-1)-1}) \\ + 4 \times (2^{2(r-2)-1} - 2^{(r-2)-1}) - (2^{r-2} - 1) \\ = 2^{2r-1} - 2^{r-1}. \end{array}$$

Now we know $\sum_{n=1}^{calls_{r-1}} |M|$ is $2^{2(r-1)-1} - 2^{(r-1)-1}$ and we can calculate the number of products of submatrices of size $2^{p-r} \times 2^{p-r}$. During these calls *perfor*mMultiplications runs 3 times, $|multiplicationTask1| = 2 \times 2^{2(r-1)-1} - 2^{(r-1)-1}$ and $|multiplicationTask2| = |multiplicationTask3| = 2^{2(r-1)-1} - 2^{(r-1)-1}$. So. the number of products of submatrices of size $2^{p-r} \times 2^{p-r}$ is

$$4 \times (2^{2(r-1)-1} - 2^{(r-1)-1}) = 2^{2r-1} - 2^r.$$

Theorem 2. Let |G| be the length of the description of the grammar G and let n be a length of an input string. Then algorithm in listing 2 calculates matrix T in $\mathcal{O}(|G|BMM(n)\log n)$ where BMM(n) is the number of operations needed to multiply two Boolean matrices of size $n \times n$.

Proof. The proof is almost identical to the proof of the theorem 1 given by Okhotin [11], because the modified algorithm computes the same number of products of submatrices just like the Valiant's algorithm (lemma 2), so the time complexity of our algorithm is the same as of the original one.

To summarize, we proved the correctness of the modification and shown that the time complexity remained the same as in Valiant's version.

3.3 Algorithm for Substrings

Next, we show how our modification can be applied to the string-matching problem. To find all substrings of size s, which can be derived from the start symbol for an input string of size $n = 2^p - 1$, we need to compute layers with submatrices of size not greater than 2^r , where $2^{r-2} < s \le 2^{r-1}$.

Let r=p-(m-2) and consequently (m-2)=p-r. For any $m \le i \le p$ products of submatrices of size 2^{p-i} are calculated exactly $2^{2i-1}-2^i$ times and each of them imply multiplying $C=\mathcal{O}(|G|)$ Boolean submatrices. Let BMM $=n^{\omega}f(n)$, where $\omega \ge 2$ and $f(n)=n^{o(1)}$. Now we estimate the number of operations needed to find all substrings:

$$C \cdot \sum_{i=m}^{p} 2^{2i-1} \cdot 2^{\omega(p-i)} \cdot f(2^{p-i}) = C \cdot 2^{\omega r} \sum_{i=2}^{r} 2^{(2-\omega)i} \cdot 2^{2(p-r)-1} \cdot f(2^{r-i}) \le C \cdot 2^{\omega r} f(2^r) \cdot 2^{2(p-r)-1} \sum_{i=2}^{r} 2^{(2-\omega)i} = \text{BMM}(2^r) \cdot 2^{2(p-r)-1} \sum_{i=2}^{r} 2^{(2-\omega)i}$$

Thus, time complexity for searching all substrings of size not greater than 2^r is $O(2^{2(p-r)-1} |G| \operatorname{BMM}(2^r)(r-1))$ where the appeared factor meet the number of matrices in the last completed layer, while time complexity for the whole input string is $O(|G|\operatorname{BMM}(2^p)(p-1))$. The Valiant's algorithm completely calculates at least 2 triangle submatrices of size $\frac{n}{2}$, as shown in Figure 5, thus the minimum asymptotic complexity is $O(|G| \operatorname{BMM}(2^{p-1})(p-2))$. Thus we can conclude that the modification is asymptotically faster than the original algorithm for substrings of size $s \ll n$.



Fig. 5. Valiant's algorithm: it is necessary to calculate at least 2 triangle submatrices of size $\frac{n}{2}$ even for short substrings finding

4 Evaluation

In this section we present the results of experiments whose purpose is to demonstrate the practical applicability of the proposed algorithm. All tests were run on a PC with the following characteristics: OS: Linux Mint 19.1, CPU: Intel i5-8250U, 3400 Mhz, RAM: 8 GB, GPU: NVIDIA GeForce GTX 1050 MAX-Q.

We implement two different versions of Valiant's algorithm and its modification in C++ programming language³:

- CPU-based solutions (valCPU and modCPU). One of the most efficient implementations of the "Method of the Four Russians" [2] from the library M4RI [1] is used for Boolean matrix multiplication.
- GPU-based solutions (valGPU and modGPU). A naive Boolean matrix multiplication in CUDA C with Boolean values treated as bits and packed into uint_32 is implemented.

We evaluate these implementations on context-free grammars D_2 :

$$s \rightarrow s \; s \; | \; A \; s \; U \; | \; C \; s \; G \; | \; \varepsilon$$

and BIO:

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\begin{array}{rcl} s \rightarrow & \operatorname{stem}\langle s0 \rangle \\ \operatorname{any\_str} \rightarrow & \operatorname{any\_smb} * [2..10] \\ s0 \rightarrow & \operatorname{any\_str} \mid \operatorname{any\_str} \operatorname{stem}\langle s0 \rangle \; s0 \\ \operatorname{any\_smb} \rightarrow & A \mid U \mid C \mid G \\ \operatorname{stem1}\langle s1 \rangle \rightarrow & A \; s1 \; U \mid G \; s1 \; C \mid U \; s1 \; A \mid C \; s1 \; G \\ \operatorname{stem2}\langle s1 \rangle \rightarrow & \operatorname{stem1}\langle \operatorname{stem1}\langle s1 \rangle \rangle \\ \operatorname{stem}\langle s1 \rangle \rightarrow & A \; \operatorname{stem}\langle s1 \rangle \; U \mid U \; \operatorname{stem}\langle s1 \rangle \; A \mid C \; \operatorname{stem}\langle s1 \rangle \; G \\ & \mid G \; \operatorname{stem}\langle s1 \rangle \; C \mid \operatorname{stem1}\langle \operatorname{stem2}\langle s1 \rangle \rangle \end{array}
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We want to look at two grammars of different sizes. Grammar D_2 is chosen because grammars that describe well-balanced sequences of brackets are often used in string analysis in bioinformatics. Grammar BIO applies to the tRNA classification problem in paper [5]. We test both synthetic strings and real RNA subsequences with length n up to 8191 and search substrings with length subs up to 2040. The results of the evaluation are summarized in the tables 1 and 2. Time is measured in milliseconds.

The comparative analysis (table 1) shows that the performance of Valiant's and modified algorithms is the same for the CPU-based solutions. The GPU-based implementation of Valiant algorithm is slower for grammar D_2 than the CPU-based one. It is probably because of processing a large amount of small matrix multiplication which cannot be computed concurrently. But it also shows that the modified algorithm is more efficient to use parallel techniques. In other cases, GPU-based solution provides significant performance improvement, especially for our modification, where utilization of parallelism facilitates parallel multiplication of the matrices itself and matrices in a layer.

³ The source code is available on GitHub: https://github.com/SusaninaJulia/PBMM.

	Grammar D_2				Grammar BIO			
n	valCPU	modCPU	valGPU	modGPU	valCPU	modCPU	valGPU	modGPU
127	78	76	195	105	1345	1339	193	106
255	289	292	523	130	5408	5488	525	140
511	1212	1177	1909	250	21969	22347	1994	256
1023	4858	4779	7878	540	88698	90318	7890	598
2047	19613	19379	33508	1500	363324	374204	34010	1701
4095	78361	78279	140473	4453	1467675	1480594	141104	5472
8191	315677	315088	-	13650	-	-	-	18039

Table 1. Comparison of the Valiant's algorithm and the modification

To adapt our algorithm for the string-matching problem the, main() function takes an additional argument sub — the maximum length of strings we want to find, so the modification has no need to compute all layers as shown in section 3.3. The corresponding implementations are named as adpCPU and adpGPU.

Table 2. Modified algorithm evaluation on the string-searching for the BIO grammar

sub	n	adpCPU	adpGPU
	1023	2996	242
250	2047	6647	255
250	4095	13825	320
	8191	28904	456
	2047	12178	583
510	4095	26576	653
	8191	56703	884
1020	4095	48314	1590
1020	8191	108382	1953
2040	4095	197324	5100

The results of the second evaluation (table 2) shows that the modified version of algorithm can find all derivable substrings much faster than the Valiant's algorithm, thus it can be efficiently applied to the string-searching problem.

5 Conclusion and Future Works

We presented a modification of the Valiant's algorithm which makes it possible to process each matrix in layer independently and use parallel computations more efficiently. This new algorithm can efficiently handle the problem of finding all substrings of a specified length. The proposed algorithm is accompanied by the proof of correctness and computational complexity analysis. We demonstrated practical applicability of our modification. Concurrent processing of matrices in layer significantly increases the performance of GPU-based solution. Also, the

modification can find all substrings much faster than Valiant's algorithm through the possibility to stop parsing matrix filling.

The directions for future research is to extend the proposed algorithm to handle conjunctive and boolean grammars. It is useful for complex secondary structure features processing.

References

- Albrecht, M., Bard, G., Hart, W.: Algorithm 898. ACM Transactions on Mathematical Software 37(1), 1–14 (Jan 2010). https://doi.org/10.1145/1644001.1644010
- Arlazarov, V.L., Dinitz, Y.A., Kronrod, M., Faradzhev, I.: On economical construction of the transitive closure of an oriented graph. In: Doklady Akademii Nauk. vol. 194, pp. 487–488. Russian Academy of Sciences (1970)
- Dowell, R.D., Eddy, S.R.: Evaluation of several lightweight stochastic context-free grammars for rna secondary structure prediction. BMC bioinformatics 5(1), 71 (2004)
- 4. Durbin, R., Eddy, S., Krogh, A., Mitchison, G.: Biological sequence analysis. Cambridge University Press (1996)
- Grigorev., S., Lunina., P.: The composition of dense neural networks and formal grammars for secondary structure analysis. In: Proceedings of the 12th International Joint Conference on Biomedical Engineering Systems and Technologies - Volume 3 BIOINFORMATICS: BIOINFORMATICS., pp. 234–241. INSTICC, SciTePress (2019). https://doi.org/10.5220/0007472302340241
- Hopcroft, J.E.: Introduction to automata theory, languages, and computation. Pearson Education India (2008)
- 7. Kasami, T.: An efficient recognition and syntax-analysis algorithm for context-free languages. Coordinated Science Laboratory Report no. R-257 (1966)
- 8. Knudsen, B., Hein, J.: Rna secondary structure prediction using stochastic context-free grammars and evolutionary history. Bioinformatics (Oxford, England) 15(6), 446–454 (1999)
- Liu, T., Schmidt, B.: Parallel rna secondary structure prediction using stochastic context-free grammars. Concurrency and Computation: Practice and Experience 17(14), 1669–1685 (2005)
- 10. Okhotin, A.: Conjunctive grammars. J. Autom. Lang. Comb. $\mathbf{6}(4)$, 519–535 (Apr 2001)
- 11. Okhotin, A.: Parsing by matrix multiplication generalized to boolean grammars. Theor. Comput. Sci. **516**, 101–120 (Jan 2014), http://dx.doi.org/10.1016/j.tcs.2013.09.011
- 12. Valiant, L.G.: General context-free recognition in less than cubic time. J. Comput. Syst. Sci. $\bf 10(2)$, 308–315 (Apr 1975), http://dx.doi.org/10.1016/S0022-0000(75)80046-8
- Younger, D.H.: Context-free language processing in time n3. In: Proceedings of the 7th Annual Symposium on Switching and Automata Theory (Swat 1966).
 pp. 7–20. SWAT '66, IEEE Computer Society, Washington, DC, USA (1966), https://doi.org/10.1109/SWAT.1966.7
- Zier-Vogel, R., Domaratzki, M.: Rna pseudoknot prediction through stochastic conjunctive grammars. Computability in Europe 2013. Informal Proceedings pp. 80–89 (2013)