Advantages and challenges of programming the Micron Automata Processor

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Advantages and challenges of programming the Micron Automata Processor

by

Christopher R. Sabotta

A thesis submitted to the graduate faculty
in partial fulfillment of the requirements for the degree of
MASTER OF SCIENCE

Major: Computer Engineering

Program of Study Committee:
Srinivas Aluru, Major Professor
Chris Chu
Akhilesh Tyagi

Iowa State University
Ames, Iowa
2013

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DEDICATION

I would like to dedicate this thesis to my family. Their unconditional care for me has been at the heart of all my accomplishments.

I would also like to thank my friends for their care and support during the writing of this work.
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I would also like to thank Michael Leventhal, Matt Tanner, and all of their colleagues at Micron for their assistance. Their persistent communication and collaboration has been essential.
Non-Von Neumann computer architectures are being explored for acceleration of difficult problems. The Automata Processor is a unique non Von Neumann architecture capable of efficient modeling and execution of non-deterministic finite automata. The Automata Processor is shown to be excellent in string comparison operations, specifically with regard to bioinformatics problems. A greatly accelerated solution for Prosite pattern matching using the Automata Processor called PROTOMOTA is presented. Furthermore, a developers’ guide detailing the lessons learnt while designing and implementing PROTOMOTA is provided. It is hoped that the developers’ guide would aid future developers to avoid critical pitfalls, while exploiting the capabilities of the Automata Processor to the fullest.

**Keywords:** Automata Processor, Reconfigurable computing, Pattern matching.
CHAPTER 1  INTRODUCTION

Today, a lot is known about the world around us. Ironically, relatively little is known about what governs what is going on inside us. This has been changing of late with the continued introduction of extremely fast and increasingly accurate biological instruments and computing methods which has enabled the field of biology to grow in ways that might otherwise been impossible. It is the combination of this computer engineering and biology which brings us now the study of bioinformatics, perhaps one of the most prolific fields of the twenty-first century.

Many important bioinformatics problems are complex and often take a lot of time and resources. In order to truly extend our knowledge in this field, efficient solutions to these problems need to be developed so that many instances of the problem can be solved providing the necessary insight to understand the underlying biology. Therefore we need to accelerate the processes. Many researchers are looking at accelerating these using Von Neumann architectures like multicore, many core systems, GPUs etc. This thesis examines whether certain problems could be better solved using a novel non-Von Neumann architecture called the Micron Automata Processor.

Conventional computer processors are extremely quick at performing individual operations in linear succession. With these CPUs billions of individual commands can be executed one after another every second. Unfortunately, solutions of many problems in bioinformatics contain a high density of conditional branching, wherein even this seemingly fast processor can take unacceptable amounts of time to find the result. Non-deterministic Finite Automata (NFA) is an intuitive mechanism for modeling such problems, but the author is unaware of any hardware architecture which can directly execute parallel paths in an NFA concurrently. In CPU based
architectures, execution of an NFA may lead to a state-space explosion leading to exponential run-time complexity.

The Automata Processor (AP) is the first semiconductor device where the execution of an NFA can be directly emulated on the device. Besides, the large capacity of the processor allows the execution of many NFAs in parallel, leading to large scale speedup. Thus the Automata Processor can work as an effective accelerator for many applications in bioinformatics.

An intuitive use of the Automata Processor is to aid in the characterization of protein sequences through the identification of motif-patterns present in them. Prosite is a large annotated database of known protein motifs. These motifs are represented either as patterns defined using a syntax very similar to that of regular expressions or on Hidden Markov Model based profiles. In the rest of this document we would only be considering only motifs expressed as patterns in the database, of which there are currently 1308 entries. Each of these patterns are converted into an automaton and defined using the Automata Network Markup Language (ANML, pronounced as ANiMaL) in order to enable their execution on the Automata Processor. After the automata are loaded onto the Automata Processor, the protein sequence(s) are streamed to check for the occurrence of any motifs in them. The occurrence of a motif is signaled by the reaching of an accept state of the corresponding automaton. The ending position of the occurrence of the motif in a protein sequence is marked by offset in the protein sequence when the accept state was matched. The large capacity of the Automata Processor allows the automata for the entire database of Prosite patterns to be loaded on a single chip simultaneously. By executing all the atomata concurrently, large speedups can be obtained.

It goes without saying that the Automata Processor is a unique device. Utilizing the power of the Automata Processor means modeling solutions to problems in unconventional ways. Even skilled developers may initially find it challenging to think of solutions in terms of NFA. Besides, the developers need to learn the nuances of using this device effectively and the restrictions that it places on the automata for their efficient execution. In this thesis, the Developers Note section contains all the lessons learnt while designing and implementing PROTOMOTA. This section details the discovered brilliance and pitfalls of developing for the
Automata Processor and seeks to lower the barriers of entry to programming on the Automata Processor.

Although adjusting to the novelty of the Automata Processor was at times arduous, it proved to be rewarding. The implementation of the PROSITE patterns in the Automata Processor is intuitive, and the acceleration provided is exemplary. For such direct, regular-expression-like string matching the Automata Processor seems unparalleled. The Automata Processor stands to show itself as an impressive boon for the world of bioinformatics and will be watched with anticipation as future generations of it are developed.

The rest of this thesis is organized as follows. First, chapter 2 provides an introduction to the Automata Processor. Further, chapter 3 then discusses programming model and ways of programming the automata processor. In chapter 4, an application to accelerate the search for motif-patterns in protein sequences using the Automata Processor called PROTOMATA is presented. This is followed by the detailed developers’ notes in Chapter 5. Finally, this thesis culminated with conclusion and scope of future work in Chapter 6.
CHAPTER 2 UNDERSTANDING THE MICRON AUTOMATA PROCESSOR

2.1 Introduction

The Micron Automata Processor (AP) is a highly parallel reconfigurable non-von Neumann architecture designed for the emulation of Non-Deterministic Finite Automata (NFA). The Automata Processor is effective in modeling complex regular expressions as well as other more complex automata. The architecture allows for effective solutions for pattern matching problems found in computational biology, cyber security, graphics processing, and more. Its purpose-built design exceeds the capabilities of similar FPGA-based solutions while also significantly outperforming them. The Automata Processor connects to a system through PCI Express and is designed to supplement a CPU as an acceleration device rather than a standalone utility.

2.2 Understanding the Automata Processor

The Automata Processor is purpose-built hardware designed specifically for the acceleration of processing Non-Deterministic Finite Automata (NFA). The processing of NFA has become an important topic as they offer an intuitive model for regular expression matching problems, which can be extended to numerous fields.

2.2.1 Sequential processor implementations of NFA

A von Neumann sequential processor (CPU) may take a direct approach of modeling an NFA by defining a set of states and connections. By maintaining a list of which states are active, and given the next token of input data, the next set of states can be computed and
updated. Unfortunately, the CPU must make considerations for each of its active states and their connections sequentially, and such state processing time is increased with increased activity across the graph. In the worst case, the entire NFA may be active resulting in the entire graph needing to be considered for a given character of input - yielding a processing complexity $O(n^2)$. Although such direct solutions offer a simple storage cost $O(n)$, the large processing complexity makes them unrealistic for large problems.

While CPUs struggle with the processing of multiple simultaneous states in NFA, Deterministic Finite Automata (DFA) require singularity in state. This property allows for the direct processing $O(1)$ of DFA, and encourages a translation from NFA to DFA. For an NFA to be modeled as a DFA, the equivalent DFA must be able to model any combination of simultaneous states possible in the NFA. For this reason while NFA can be reliably converted into equivalent DFA [2], it comes with the challenge of significantly inceased storage cost.

The storage cost reduction of NFA-derived DFA is the target of much research, and has been viewed from many different perspectives. Some solutions such as Campeanu et al. [3] focus on simplifying the NFA as much as possible by combining excessive states. Other solutions attempt to reduce the scope of the problem to allow for simplifications. Chia-Hsiang et al. [4] require compressed NFA specifically taken from regular expressions for their solution. Alternately, Holub [13] attempts storage and simulation simplifications through dynamic programming. While many unique solutions have been developed however, no solution escapes the exponential DFA storage cost. [25, 24, 23, 21]

2.2.2 Automata Processor implementations of NFA

The Automata Processor allows for a physical implementation of NFA. It has many configurable elements which act as individual states for a modeled NFA. These elements can be dynamically connected to one another and configured to respond to different stimuli. In this way the Automata Processor is capable of literally modeling NFA directly. Because of this the Automata Processor has a direct NFA storage cost of $O(n)$.

The physical design of the Automata Processor allows for processing to be done at an
### Table 2.1 Processing and Storage Complexity for NFA Implementation

<table>
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<th>Implementation</th>
<th>Processing Complexity</th>
<th>Storage Cost</th>
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<tr>
<td>Sequential Processor NFA</td>
<td>$O(n^2)$</td>
<td>$O(n)$</td>
</tr>
<tr>
<td>Sequential Processor Equivalent DFA</td>
<td>$O(1)$</td>
<td>$O(\sum^n)$</td>
</tr>
<tr>
<td>Automata Processor NFA</td>
<td>$O(1)$</td>
<td>$O(n)$</td>
</tr>
</tbody>
</table>

element level, and gives the input to all elements simultaneously. Because of this, all state processing is done in parallel. This means that regardless of how many states are active in an Automata Processor modeled NFA, the next-state computation is constant. This gives it an $O(1)$ processing cost, and an impressive advantage over generic CPU solutions shown in Table 2.1.

#### 2.2.3 Gaining parallelism

A common target of NFA solutions is pattern matching. An NFA can be used for pattern matching rather directly. For every character in a target string, a state is created to look for that character. With the states connected in the order in which they appear in the target string, the automata is complete. For the state of the final character to have been entered, it must have passed through all previous states consecutively. This way we know that if the final state has been entered, the pattern has been matched.

The Automata Processor is capable of simultaneously processing action for each of its active states for each token of input. The time step in which an input character is processed and the states are updated is known as a symbol cycle. The effective parallelism of the Automata Processor is in part derived from this simultaneous inspection of multiple active states within an NFA. Parallelism for the Automata Processor is further derived from its ability to simultaneously process multiple separate NFA. A simple example of this parallelism is shown with two string matching automata processing an input stream of 'COCO...' in Figure 2.1. Note that unique possibilities are explored within a single automata, as well as within different automata simultaneously.
2.3 Automata Processor implementation

The entire Automata Processor exists as a conglomeration of 6 distinct ranks connected to a system through PCI Express. An Automata Processor rank consists of 8 distinct Automata Processor cores on a single chip. Each core consists of two half-cores containing 24K elements each, where no connections can be made between elements of unique half-cores. It is the elements of these half-cores which are used to directly model the states of NFA.

With a section of the Automata Processor configured to model a desired NFA, input can be streamed. The modeled NFA views each streamed-in character as the stimulus for a potential state transition. An Automata Processor core processes data at a rate of 1 Gbps. Because one byte characters are the fundamental unit of the Automata Processor this can be better viewed as $128 \times 10^6$ characters per second. This means that the state of a modeled NFA can be updated every $7.45 \times 10^{-9}$ s. This time is also known as a symbol cycle.

Cores can be associated among their rank, in groups of 1, 2, 4, 8 cores. Grouped cores will receive data from the same stream of data, while cores of different groups can concurrently process different streams of data. With all cores associated in one group of 8 the Automata Processor has an effective throughput of 1 Gbps. With all cores grouped individually with their own data streams an effective throughput of 8 Gbps is achieved.


2.4 Competing NFA hardware

While NFA are difficult to emulate with a sequential processor, Field Programmable Gate Arrays (FPGAs) are capable of NFA modeling with the use of look up tables [24, 23, 21]. Similarly, GPU based solutions have been devised [25]. Due to its specialization, the Automata Processor is able to show decisive advantage over such FPGA and GPU based solutions. Furthermore, the Automata Processor remains the only such specialized non-FPGA, non-GPU hardware.

2.4.1 Scope of ability

FPGA and GPU based competing NFA solutions focus directly on implementation of Perl Compatible Regular Expressions (PCRE)[1]. For the sake of direct comparison, the Automata Processor is assessed in terms of such regular expressions. It is important to note however that the Automata Processor maintains the unique ability to model non-PCRE NFA through its own configuration language. This configuration language is known as Automata Network Markup Language (ANML) and allows for simple directed implementation of NFAs with an XML structure.

The Automata Processor uniquely allows for scaling across multiple chips with balancing for capacity and throughput. The architecture also allows for new automata to be added to dynamically to the chip without recompiling the existing automata. This process is known as incremental update and is only possible with the Automata Processor, whereas competing FPGA and GPU solutions require a new compilation [25, 23, 21, 25]. Similarly, the Automata Processor allows for the dynamic reconfiguration of match values and path pruning without reproducing the layout of the chip. Outside of the Automata Processor, dynamic reconfiguration is only present in competing solutions using restricted RegEx [8].

2.4.2 Performance and capacity

Although there are 6 ranks in one instance of the Automata Processor, because they cannot cooperate only 1 rank is used for comparison against other solutions. While statistics on
throughput and capacity could be realistically increased by a factor of 6 in most cases, multiple instances of other solutions could be used to the same effect. Still, the Automata Processor maintains the advantage of being able to connect these 6 ranks to the rest of a system through only a singular PCI Express port.

The Automata Processor is shown to be competitive with FPGA based solutions [24, 23, 21] in terms of throughput. Raw comparisons of throughput are difficult to produce for a number of reasons. First, various FPGA solutions show data consumption rates ranging anywhere from 1 to 8 characters per cycle. While this increases the technical throughput, it does not necessarily model the rate at which the NFA can transition between states. It must also be considered however that larger per-cycle data consumption potentially introduces a higher level of control. Wang et al. [23] boast a derived 2.57Gbps throughput but offer significantly lower capacity than the Automata Processor, requiring 6 chips to fit roughly half the capacity of an Automata Processor rank. Yang et al. [24] are capable of a 10Gbps result, but this is with a consumption of 8 characters per cycle. With slightly better efficiency they also present a 3.5Gbps result consuming 2 characters per cycle, though all solutions present significantly lower capacity than the Automata Processor at less than one fourth of one rank.

Competing GPU based solutions in that of Zu et al. [25] show results as high as nearly 14Gbps. In exchange for this rapid processing both routing and capacity are found to be significantly weaker than with the Automata Processor. This GPU solution only allows for 4 connections from each element, where the Automata Processor allows for 16. Furthermore the GPU solution can support only one sixth of the elements of a rank of the Automata Processor. Although the GPU solution boasts a slightly higher throughput, multiple ranks of the Automata Processor can match such throughput much more quickly than multiple instances of the GPU can match the Automata Processors capacity. Even with matched capacity, the Automata Processor remains superior in routing.

A rank of the Automata Processor is capable of producing up to 8Gbps throughput if the problem can be dissolved into 8 unique streams. For large automata that must span multiple cores and require the same input, the effective throughput is reduced relative to the
group size. This notion of chip scaling makes definative throughput comparisons difficult. Though the Automata Processor cannot be determined to have higher throughput than all GPU and FPGA solutions in all cases, it is shown to be competitive. For more information on the comparative capabilities of the Automata Processor view *Supplementary Material for An Efficient and Scalable Semiconductor Architecture for Parallel Automata Processing* [8]
CHAPTER 3 USING THE MICRON AUTOMATA PROCESSOR

The Automata Processor proves to be exceptionally effective for NFA implementation on account of its reconfigurability and unique modeling capabilities. To fully utilize the strengths of the Automata Processor its components and automata definition mechanisms must be understood. This chapter begins by defining the building blocks of Automata Processor NFA, and continues to explain how such NFA can be programmed into the device.

3.1 Automata Processor components

The definition of NFA using the Automata Processor can be divided into two major components: elements and connections. Individual states of NFA can be modeled using Automata Processor elements. The directional connections between states are modeled by defining routing characteristics for these elements. Finally, input and output characteristics can be defined for the automata are defined to complete a solution.

Elements of an automata can be either active or inactive at any given time. An active element models an active state within the modeled automata. For each input character in the stream whether or a not a state is active is determined by the following two queries:

1. The given element is on the receiving end of a connection with a currently active element

2. The given element is configured to accept or match the given input character

If both of these queries are true, the given element enters an active state for the next processed input character. If either or both of these queries are false, the given element enters an inactive state, regardless of its current state as a default action. Any element may become
permanently active after it is first activated. This process is known as latching. If an element is in an active state, it is said to be driving any elements to which it connects.

### 3.1.1 Types of element

Elements provide the building blocks for any graph modeled by the Automata Processor. Some elements are used for modeling state, while other elements exist for extended capabilities such as basic logic and counting. Each type of element has unique configurable properties for NFA modeling.

#### 3.1.1.1 State transition element

The state transition element (STE) is the fundamental component of the automata. Each STE holds a configurable symbol set which defines its matching characteristics. This symbol set can be defined to be any subset contained within the full set of 256 possible input characters. An STE directly models an NFA state and is processed for each character of input in what is known as a symbol cycle. It is by far the most plentiful and most important type of element for the Automata Processor.

#### 3.1.1.2 Counter element

Counter elements serve to supplement the work of STEs, and add higher levels of modeling capability to the Automata Processor. The counter element does not define a symbol set, nor any other form of direct comparison. Instead, counter elements activate when a configurable count is reached. Counter elements update an internal counter by one during each symbol cycle they are driven by an active element. When the defined target is reached, the counter drives its outgoing connections. This output driving can be configured to be either for a single pulse or a latched, continuous operation. Counter elements also define an input port for resetting the count. If a connection is driven to the reset port the count is returned to its original value. Finally, it is important to note that counter elements driven by STEs are processed in the same symbol cycle as the driving STE.
The counter element can be used to consolidate a large number of duplicate states. An example is given showing two implementations for an automata designed to find 5 occurrences of the character 'A' before a final character 'B'. Figure 3.1 shows a solution with exclusively STEs. Figure 3.2 shows a solution using the counter element.

![Counting machine using exclusively STEs](image1)

![Counting machine using a Counter Element](image2)

### 3.1.1.3 Logical element

The final class of element for constructing graphs on the Automata Processor is the logical element. A logical element is similar to a counter element in that its action is determined by input signals, and it is processed in the same symbol cycle as the STEs which drive it. Logical elements are configured to view a series of incoming connections as a set of Boolean variables. If a signal is being driven, it is considered to be a 1, and if it is not actively being driven it is considered to be 0. The variable representation of these signals are then considered as a programmable digital logic expression. The logical element is capable of simulating basic AND, OR, NAND and NOR gates as well as Sum of Product (SoP) and Product of Sum (PoS) expressions. Given a set of input signals and its programmed logic, output signals can be
driven based on the result.

3.1.2 Input and output

After defining the core implementation of an NFA, its beginning and ending states must be considered. At the beginning of each problem instance for an Automata Processor solution, the automata has no active states. Because the activation of further states is dependent on being driven by currently active states, designated elements configured with the start property are required. If an element is a start element it is considered to be driven by the data stream, and does not require being driven by another element. This start characteristic can be specified either for only the first character of the data stream, or for all character of the data stream.

Output can be generated as a result of entering a designated state. If an element is configured to generate output upon its activation the element is said to be reporting. A reporting element can be configured to generate output data immediately upon its activation. Alternately, a reporting element can be configured to wait until the end of the data stream and report if it had been activated. A word of report data consists of the element which reported it as well as the cycle in which the data has been matched. Current implementations of the Automata Processor suffer large time penalties for repeated quick reporting. Unfortunately if an element does not report until the end of data the knowledge of the matching symbol cycle is lost. A quality Automata Processor solution will reasonably balance the use of reporting elements.

3.2 Modeling automata

Programmed place and route executable solutions for the Automata Processor exist in uniquely defined finite state machine (.fsm) automata files. Such solutions can be generated in one of two ways using the Automata Processor compiler. 1) Regular Expressions can be directly converted into programmable automata, or 2) Automata may be directly defined (and go beyond regular expressions) with the XML based Automata Network Markup Language (ANML). Generating automata directly from regular expressions is trivial, however it is very
limiting in terms of its capabilities. For this reason the remainder of this section focuses on the implementation of automata using ANML. Although the general form of ANML is discussed, it is important to review the most recent schema for the most up-to-date syntax and component definition information.

### 3.2.1 Using Automata Network Markup Language

The Automata Network Markup Language (ANML, pronounced ANiMaL) provides a simple yet comprehensive language for describing automata in the vein of XML. ANML serves to define elements and their connections within the scope of an automata network. Start and reporting characteristics as well as potential latching and other configurable aspects of elements can be defined for each defined component. ANML also offers the notion of a macro for the compartmentalization of sub graphs as well as simple reuse of functional graph components.

#### 3.2.1.1 ANML XML layout

All specifications in ANML are made within matched XML tags. All defined components of an automata must be described between matched automata-network tags. Beyond this the definitions may be totally flat. The only other requirement for an ANML file is that all components are given a unique id in their opening tag.

Individual elements can be defined using their appropriate definition tags. For each kind of element there are both required fields and optional fields. Specifications for start, reporting, latching and other optional behavior are not required and will use a simple default if unspecified. Intrinsic properties of each element are required and will not compile if left undefined. STEs must define a symbol set, Counter elements must specify a target, and logical elements must specify a logical configuration.

Within the definition of an element, outgoing connections to other elements can be defined. For this the activate-on-match tag is used in combination with the unique id of the target element. With this basic understanding of element definition and connection specification, fully functional NFA can be defined.
3.2.1.2 Using macros

Macros are an effective means for reproducing multiple identical or near identical subgraphs without dramatically expanding the defining ANML. Macros may also be useful for adding a level of organization to an automata by encapsulating complicated subgraphs into a high level block.

As with other ANML components, macro definitions require a unique id for reference. A macro definition is divided into two components labeled port definition and macro body. The port definition is used to characterize the quantity and labels of incoming and outgoing ports for the macro. Within the macro body internal elements are defined and connected. Also within this macro body, a port mappings tag is used to connect internal elements to the predefined ports.

Variables can be introduced into a macro definition for producing unique macro references from the same definition. To accomplish this a parameter declaration field is added to the port definition section of the macro definition. Within this parameter declaration parameter names are defined with default values. These parameters are referenced within the macro body to allow for variability.

Macro references generate instances of a specified macro definition. Macro references require their own unique id, as well as a use field specifying the target macro definition. If the target macro definition has parameters, they must be specified by the macro reference or they are returned to their default values. Connections using macro references can be named by using name of the macro reference followed by the desired port.

3.2.1.3 Generating ANML

While the modeling of NFA in ANML is straightforward and relatively simple it can be difficult to do for sizable solutions. Large and complicated graphs can be particularly difficult to visualize and debug when simply read from a file. To avoid the difficulties of directly coding ANML there are two main classes of abstraction: 1) Develop ANML in a graphical, drag and drop environment or 2) Develop ANML through specialized executable programs.
The AP Workbench has been developed as the pioneer drag and drop graphical environment for the formation of ANML files. It allows for rapid development of small automata, guarantees valid syntax, and requires no formal understanding of ANML whatsoever. It is capable of compiling and simulating its own ANML on the fly and allows for a comprehensive active visual debugging. Unfortunately such graphical applications lack the ability to realistically produce and simulate large applications.

Specialized executable programs offer a powerful alternative to graphical solutions. By using other compiled languages to generate ANML, automata can be generated dynamically with exceptional finesse based on some input parameters. Such an approach lends itself to large, complicated, and automated solutions. Currently, no supporting libraries or API have been released in any such languages. While such conversion applications can be effective, they require significant development work and do not offer the same conveniences of graphical development environments.

For large, complicated, variable automata development ANML conversion scripts or applications may provide the only realistic solution. In such cases graphical environments can still play a beneficial role in debugging small subsections or samples of the entire automata. The AP Workbench may also be used preliminarily to rapidly assess the workings of NFA mechanisms that may be difficult to conceptualize. It is therefore important to maintain an open mind when generating ANML code.
4.1 abstract

Motifs are useful to understand the characteristics and functionalities of a protein by identifying the families and domains of proteins that it belongs to. Currently, 1308 known motifs are represented as patterns in a large annotated database called PROSITE. PROTOMOTA is a hardware-accelerated solution to scan protein sequence(s) for these motifs through the use of a novel semiconductor architecture called the Micron Automata Processor. The Micron Automata Processor is purpose-built to search for thousands of patterns in parallel allowing PROTOMOTA to be many orders of magnitude faster than state-of-the-art CPU based algorithms. For example, all the proteins present in the proteome of Escherichia coli (E.coli) can be scanned for all the motif-patterns present in PROSITE in less than 100 milliseconds in contrast to the several minutes reported by state-of-art CPU based solutions. Besides helping biologists to understand and characterize the ever growing number of newly sequenced proteins, PROTOMOTA is designed to serve as an useful guide for application developers to understand and develop programs using this new accelerator architecture.
4.2 Introduction

PROSITE [17] is a large annotated database of known motif descriptors which can be used to identify families and domains that a protein belongs to. This becomes especially important when the sequence of an unknown protein is too distant to proteins of known protein structures for pairwise sequence alignment to reveal the relationship. A motif specifies a small region in the protein sequence which is conserved in both structure and sequence and plays biologically meaningful functions like binding properties, catalytic sites, enzymatic activity, etc.

In PROSITE, motif descriptors are represented as patterns and profiles. A pattern is expressed using a syntax very similar to regular expressions, whereas a profile is defined using a weight-matrix. Both these methods have their own utilities. While patterns are easy to understand and are effective for identification of short motifs, profiles provide higher sensitivity by identification of larger doamins and allowing higher divergence. Throughout the rest of this paper, the discussion is limited to the scanning of protein sequences for motifs using patterns alone.

ScanProsite [5] is an interactive web access tool which is used to scan protein sequences for motifs. It works in three modes: 1) up to 10 protein sequences can be submitted to be scanned against the PROSITE collection of motifs; 2) a single motif or a combination of motifs can be submitted to be scanned against a protein sequence database; and 3) protein sequences or a protein sequence database and a motif or a combination of motifs can be submitted to be scanned against each other. A PERL-based version of the tool called ps_scan can also be downloaded to scan proteins locally by using pattern entries in PROSITE. Similar programs have been developed by academic groups [9, 18, 15, 16, 12, 22, 6, 11, 10] and commercial companies [19]. A complete repository of these programs can be found in the prosite.prg file distributed by PROSITE.

In this paper, a hardware-accelerated solution to scan for protein motifs using the Micron Automata Processor is presented. The Automata Processor [7] is a purpose-built semiconductor architecture which can be programmed to identify thousands of patterns present in a data stream in parallel. Our software program PROTOMATA automatically converts PROSITE
pattern descriptors to Non-deterministic Finite Automata (NFA) which are used to program the Automata Processor. The data stream comprises of special character-sequences to enable or disable specific patterns and the protein sequence(s) which are to be scanned.

In PROTOMATA, all the 3 modes of operation ScanProsit are supported, and the hardware acceleration provides great performance benefits. Firstly, in PROTOMOTA, proteins can be streamed at 128 MBps, thus allowing even large proteomes to be scanned within milliseconds instead of minutes reported by state-of-the-art CPU-based solutions [14]. Secondly, with such high throughputs, restrictions on the maximum number of protein sequences to be scanned can be overcome. Thirdly, the large capacity of the Automata Processor board allows us to run 48 independent instances of PROTOMATA in parallel, which might be useful in a server environment like PROSITE. Further, the capability of the Automata Processor to execute NFA allows straightforward and easy implementation of matches with insertion and deletion errors. This opens up an interesting biological question. Can longer motifs be defined using longer patterns or combination of patterns allowing insertions and deletions? Such a motif would provide the readability of a pattern while providing the high sensitivity of a profile.

The rest of the paper is organized as follows. Section 4.3 discusses the methodology, whereas Section 4.4 provides the implementation details and results. Finally, we conclude in Section 4.5.

4.3 Methodology

The general overview of the working of PROTOMOTA is shown in Figure 4.1. First, all the patterns are downloaded and a c++ program is used to convert the patterns into their equivalent automata expressed in the ANML format. These automata are compiled using the ANML compiler provided with the Automata Processor SDK. At run-time, the image of the compiled automata files are loaded using the loader which is also a part of the SDK. The user provided protein sequence(s) can now be streamed through the Automata Processor and any patterns which are present in the protein sequence(s) are identified and reported to the user.
4.3.1 PROSITE patterns

Prosite patterns are defined as a sequence of pattern elements. Single letter codes as defined by IUPAC are used to represent amino acids. Prosite allows for the following pattern elements using the alphabet of amino acids:

- A single amino acid is denoted by a lone amino acid character. For example: $S$ denotes only the amino acid $S$.

- One of multiple amino acids is denoted by a series of characters within brackets. For example: $[STG]$ denotes $S$ or $T$ or $G$.

- Anything outside of a set of amino acids is denoted by a series of characters within curly brackets. For example: $\{STG\}$ denotes any amino acid that is neither $S$ nor $T$ nor $G$.

- The lower case letter $x$ is used to denote any amino acids.
• A pattern can be restricted to the N-terminal of a protein by beginning the pattern with the character <. For example: < S would only match an S found at the beginning of the sequence.

• A pattern can be restricted to the C-terminal of a protein by ending the pattern with the character >. This can also be found in brackets. For example: T > would only match a T found at the end of the sequence. Similarly [T >] indicates that either a T or the end of the sequence is acceptable.

• The repetition of a pattern element pe can be defined using parenthesis. This can be done for a specific number or a range. For example: pe(m) indicates the repetition of pe m times. Similarly pe(m, n) indicates the repetition of pe anywhere from m to n times inclusive. Here, m can be zero.

• Each pattern element is separated by a concatenation symbol ‘-’. To denote the end of a pattern the period character ‘.’ is used.

An example of a full Prosite pattern is as follows:

PS00008: N-myristoylation site

\[ G − EDRKHPFYW − x(2) − [STAGCN] − P. \]

### 4.3.2 Conversion of PROSITE patterns to ANML automata

For each PROSITE pattern there are 2 automata created: filter-automaton used for identifying the only those patterns which are present in all the protein sequence(s); and location-automaton for identifying the location(s) where these sequence(s) where these patterns occur. This is done to mitigate the amount of output handling and its effect on the run-time of the entire process. The filter-automaton generates output only at the end of the streaming of all protein sequence(s) and the location-automaton generates output only at those positions where a common motif occurs in the sequence(s).

Both the automata are constructed in a modular fashion using macros. The only macro which changes from one motif to another is the pattern macro. The other macros are used
to either enable the search for that motif based on user inputs, or in the case of the filter-
automaton to identify motif common to all the sequences.

A sequential ANML program modeling a pattern can be generated by converting each
item into a set of State Transition Elements (STEs.) Using a conversion executable developed
for this purpose, a unique macro is created for each pattern in an input PROSITE data
file. Each character in a pattern is implemented using a State Transition Element which are
then be connected in the order presented to create the pattern. Single amino acid, one-of-
multiple amino acid, and anything-but amino acid can be directly implementing by shosing
the corresponding character class as the label for the STE. For the wildcard $x$, an STE with
the character set $[A−Z]$ as the label is used.

For small repetitions and ranges, one STE with a given symbol set is produced for the
maximum number of repetitions. These are linked sequentially and can be considered together
as satisfying the pattern element by linking in to the $STE_0$ and out of $STE_n$ where $n$ is the
maximum number of repetitions. In the case of a range $pe(m, n)$, additional links are included
to simultaneously consider all $k$ for $m <= k <= n$. To achieve this the previous pattern
element in the sequence links to $STE_0, STE_1, STE_2...STE_{n−m}$.

For large repetitions it begins with a single regular STE. This STE connects to a counter
element which tallies consecutive occurrences. Introduced with the counter is a second STE
with a complemented symbol set to the target. The complimenting STE resets the counter
to zero any time a non-matching amino acid is read. When the counter reaches its target the
counter activates the next pattern element.

For large ranges a system similar to large repetitions is used. For this, the original counter
is used as a lower bound. An additional counter is introduced to act as an upper bound. When
an upper bound is reached both counters are reset.

With all pattern elements represented and connected in ANML the code is collected as
a Pattern Macro. Other supporting macros are used in conjunction with the unique pattern
macros to support more sophisticated functionality. An enable macro is used as a prefix to
each pattern macro. This macro is programmed with a $16b$ enable code. The beginning of any
input stream is a preamble listing the enable codes of the desired macros. The enable macro if selected will continuously drive the beginning of its pattern macro. If it is an N-terminal macro, the enable macro will only drive the pattern macro for the first cycle of data. The filter-automaton and the location-automaton shown in Figure 4.2 and Figure 4.3 respectively.

The end of the preamble is denoted with the special character 'xff'. After the enable preamble, proteins sequences are streamed. Each protein sequence is delimited by a 'n' character. When all protein sequences have been streamed, a special character 1 is inserted to indicate the end of the stream and to enable reporting. In the filter-automaton, a reporting macro is used as a suffix to each pattern macro to watch for these special characters and reset the pattern macro when necessary. For streaming multiple proteins a reporting macro
will restart the pattern macro when the pattern macro has been found to match the current protein sequence and the $n$ character has been reached. If the pattern is matched in the protein sequence(s), the terminating character 1 is reached, the reporting macro reports a match at the end of the data stream. The reporting macro allows for C-terminal macros by only accepting patterns that have matched directly before the ‘$n$’ character.

4.3.3 Finding patterns in proteins

The modularity of the ANML program allows for multiple different uses without needing to reprogram the Automata Processor. The preamble of the input stream individually enables patterns as their enable codes are read. For easy configuration, groups of enable codes can be loaded from user defined files. To find all of the patterns present in a protein is natural. First the two byte enable code is streamed for each desired pattern, or all. The preamble is ended with the special character ‘\xff’. The protein is then streamed, followed by a ‘\n’ character.
Finally a 1 is streamed to indicate the end of the file. At the point, all enabled patterns which were found in the protein report.

This process can be extended to find all patterns present in all of a series of proteins. For this the enabling mechanism is the same. After this each protein is streamed in order, each delimited by the \(\backslash n\) character. When an enabled macro has not matched and encounters a \(\backslash n\) character it is disabled. When the final protein has streamed only patterns that have matched every protein are enabled and can report.

**4.3.4 Finding pattern locations in proteins**

After finding the patterns common to all proteins with the common-pattern-finding version of the program, the locations of these patterns within the proteins must be found. To accomplish this, the location-finding version of the program is used. The exact same preamble and data are streamed. With the location-finding version, reports are generated for each pattern match for each protein. With this, the matching location for each pattern for each protein of the motif is known.

**4.4 Implementation and results**

Prosite defined patterns are available on their website altogether in a single '.dat' file. For this work this file was used, defining 1308 different patterns. A program has been developed to generically convert patterns defined in this '.dat' form to a single ANML program. The ANML only needs to be generated once and takes 0.87s. The resulting program has 23713 STEs, with 1308 reporting elements or one for each pattern.

This section continues by breaking down the runtime characteristics of the rest of the process.

**4.4.1 Compile-time overhead**

Due to the modularity of the designed solution, compiling the ANML code into a placed-and-routed '.fsm' automata file must only be done once. This time is the sum of two com-
ponents: 1. The time of compiling the ANML file into an automata file, and 2. The time of
loading the automata file onto the Automata Processor.

Compilation time, automata size and load time vary based on the proteins which are being
represented. This is in the worst case on the order of minutes, and only needs to be performed
once for an indefinite number of instances of the problem. For this reason the compilation and
load times are not considered.

4.4.2 Execution-time overhead

After the programs are in place, the runtime of the application itself must be considered.
This can be divided into three subsections: The preamble, the data streaming, and the read-
out. The time length of the preamble \( t_p \) can be defined as
\[
    t_p = (2ep + 1) \times 7.45 \times 10^{-9}
\]
where \( ep \) is the total number of enabled patterns in each version of the program. If all 1308 patterns
were enabled simultenously for both programs, this yields a maximum combined overhead
cost of \( 3.899 \times 10^{-5}s \).

The data streaming must be done completely one time for each of the two versions of the
program. The total time \( t_d \) of this streaming can be defined as
\[
    t_d = (7.45 \times 10^{-9}) \times \sum_{i=0}^{n-1} lp_i
\]
where \( lp_i \) is the length of the \( i^{th} \) scanned protein, and \( n \) is the total number of scanned proteins.
When scanning 100 proteins of average length 300 this time is \( 2.235 \times 10^{-4}s \) for each program,
or a combined \( 4.470 \times 10^{-4}s \) for both.

Finally the read-out time is considered. For the common-pattern-finding version of the
program, all data is reported simultaneously after the terminating character. The readout cost
is a function of the number of patterns in the program. The time \( t_r \) can be defined as
\[
    t_r = ((\lceil \frac{n}{1000} \rceil \times 40) + 1) \times 7.45 \times 10^{-9}s
\]
where \( n \) is the total number of patterns in the program. For our 1308 pattern example, this time is \( 6.035 \times 10^{-7}s \).

For the location-finding version of the program output is captured as it is generated. Be-
cause each pattern only needs to be matched once to a scanned protein, and because each
scanned protein is of significant length, \( lp \gg 40 \), the readout can be performed in parallel to
the data streaming. The final readout after the data has finished streaming will have proper-
ties identical to the common-pattern-finding readout. For our 1308 pattern example, this time again is \(6.035 \times 10^{-7}\) s.

### 4.4.3 Effective processing rate

The total processing time for an instance of the problem \(t_t\) is then the sum of the components defined for each version of the program. \(t_t = 2 \times (t_p + t_d + t_r)\). Note \(t_p + t_r\) is constant with regard to the number of scanned proteins and their respective lengths. Therefore as the size and quantity of scanned proteins are increased \(t_t\) will approach \(t_d\). The effective data processing rate per character \(R_e\) is the total amount of time spent on a solution divided by the total number of input data characters processed. For a total of \(c\) characters it can then be viewed as \(R_e = (t_p + t_r) / c + 14.7\) ns.

The efficiency of the solution can be measured by how closely the effective processing rate matches the symbol cycle timing. A visual representation of this effective processing rate for an individual version of the program can be seen for two sample problems in Figure 4.4. Here Small Enable refers to a 20 pattern selection while Large Enable refers to the complete set of 1308 patterns.

### 4.4.4 Comparison and results

Because Protomata has been designed to be a general solution, no specific problem is addressed. Even so, the solution shows impressive results and scales exceptionally well for large data streaming. Computation times for sample problems are shown in Table 4.1.

<table>
<thead>
<tr>
<th>Proteins (300B) \ Patterns</th>
<th>1</th>
<th>10</th>
<th>100</th>
<th>1000</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>5.1256 (\times 10^{-6})</td>
<td>5.3938 (\times 10^{-6})</td>
<td>8.0758 (\times 10^{-5})</td>
<td>3.4896 (\times 10^{-5})</td>
</tr>
<tr>
<td>10</td>
<td>4.5356 (\times 10^{-5})</td>
<td>4.5662 (\times 10^{-5})</td>
<td>4.8306 (\times 10^{-5})</td>
<td>7.5126 (\times 10^{-5})</td>
</tr>
<tr>
<td>100</td>
<td>4.4766 (\times 10^{-4})</td>
<td>4.4792 (\times 10^{-4})</td>
<td>4.5061 (\times 10^{-4})</td>
<td>4.7743 (\times 10^{-4})</td>
</tr>
<tr>
<td>1000</td>
<td>4.4707 (\times 10^{-3})</td>
<td>4.4709 (\times 10^{-3})</td>
<td>4.4736 (\times 10^{-3})</td>
<td>4.5004 (\times 10^{-3})</td>
</tr>
<tr>
<td>10000</td>
<td>4.4701 (\times 10^{-2})</td>
<td>4.4701 (\times 10^{-2})</td>
<td>4.4704 (\times 10^{-2})</td>
<td>4.4730 (\times 10^{-2})</td>
</tr>
<tr>
<td>100000</td>
<td>4.4700 (\times 10^{-1})</td>
<td>4.4700 (\times 10^{-1})</td>
<td>4.4700 (\times 10^{-1})</td>
<td>4.4703 (\times 10^{-1})</td>
</tr>
<tr>
<td>1000000</td>
<td>4.4700</td>
<td>4.4700</td>
<td>4.4700</td>
<td>4.4700</td>
</tr>
</tbody>
</table>

Table 4.1  Total Computation Time for Sample Problem Instances (s)
A competing implementation using a Pentium 4, 3.4GHz has been found to map all prosite patterns on the whole Escherichia coli (E. coli) proteome in about 10 minutes [14]. There exist 4339 protein sequences in the E. coli proteome, each averaging a length of 287 amino acids [20]. Given these parameters, the protomata solution is capable of processing the data in $1.8660 \times 10^{-2}$s. With additional time allocated for the CPU Protomata can safely boast an runtime of less than 200ms, resulting in a larger than 3000 times speedup over published results.

### 4.5 Conclusion

In this paper, a greatly accelerated solution to scan for PROSITE patterns called PROTOMATA has been presented. In order to provide this acceleration, PROTOMOTA uses a novel hardware architecture which has been purpose-built to scan thousands of patterns in parallel. The engine serves to find common defined patterns among proteins as quickly as the data can be streamed by the Automata Processor, with minimal overhead and only a singularly
programmed instance.

**Acknowledgment**

The authors would like to thank Paul Dlugosch and his team from Micron Technology, Inc. for providing the Automata Processor boards and sharing valuable insights about how to program the same.
CHAPTER 5 APPLICATION DEVELOPERS’ NOTES

This chapter is dedicated to the discussing the intrinsic details that application developers should be aware of while designing their automata for the Automata Processor. It also contains the lessons learnt while programming PROTOMOTA, so that such pitfalls can be avoided in the future.

5.1 Automata Processor Board configuration

The Automata Processor board consists of 48 automata processor cores arranged in 6 ranks of 8 processors each. The Automata Processor cores in a single rank share a high speed inter-connect which allows them to be presented with the same data flow concurrently. The processor cores can be logically organized into logical cores and logical core groups to provide maximum throughput while using the device. A large logical core allows the programmer to exploit parallelism in terms of the set of automata executing on a single data flow, whereas the logical core groups allow parallelism through multiple data flows.

A logical core consists of 2, 4 or 8 processor cores from a single rank. All the processor cores in a logical core are presented with the same data flow. The idea is to allow the execution of one large automaton or multiple smaller automata which cumulatively take more resources than what is present on a single core. While designing large automata, one must bear in mind that the physical routing is limited to only a half-core, which as the name suggests contains half of the resources present in a single Automata processor core. When the compiler is presented with an automaton which cannot be fit inside a single half-core, it tries to partition it into multiple logical parts which can be fit into multiple half-cores inside a single logical core. When these half-cores are presented with the same data flow, it behaves exactly like the the
unpartitioned large automaton. However, one must bear in mind, that the partitioning by the compiler might not be very efficient, or in the worst case may not be possible at all. On the other hand, an application developer is cognizant of the design and purpose of the automaton and may be able to produce a much better logical partitioning of the same. Therefore, the programmer is well advised to view a logical core as grouping of half-cores rather than full Automata Processor cores.

A logical core group consists of 2 or more identically programmed logical cores. These cores may come from one or more ranks and are presented with different data-flows. The idea here is to allow the automata logic to be applied to multiple data-flows in parallel for higher throughput. The logical core groups may be handled by one or more CPU thread(s). In the case of multiple CPU threads, some of the resources of the Automata Processor board get distributed amongst the CPU threads. The effects of these partitioning are not clear to the author at the time of writing this document.

5.2 Finer details for designing automata

Having a basic understanding of the Automata Processor and how to configure it will allow any user to begin designing solutions. With a deeper understanding of the device and its configuration, however, users are capable of significantly improving their success in both development and execution.

5.2.1 Element and routing implementation

Along with the STEs, the Automata Processor also contains counter elements and boolean elements. The counter elements are used to compress the size of automata, where as the boolean elements can be used to create automata which have more expressive power than classical NFA. However, programmers should bear two things in mind: 1) counter and boolean elements are much less numerous than STEs and 2) the output event from a counter or boolean element can only be routed to an STE within one cycle.

The counter element is a 12-bit binary down-counter. It is programmed with a value and
every time one of the count enable signals is activated, the counter is decremented by 1. When the count reaches 0, an output event is triggered. The value of a counter can be used but only at the cost of reading out the entire state vector of the Automata Processor. Counters are very effective in identifying whether a sub-expression of a regular expression has been matched for exactly or at least a fixed number of times. Other implementations including those in cellular automata have also been demonstrated.

The boolean element is a function-programmable combinatorial element which can be programmed to perform the following logical functions: OR, AND, NAND, NOR, sum of products, and products of sums. The boolean elements can be used for combination of the results of subexpressions. They can also be programmed with an optional synchronized enable signal which can be used enable all the boolean elements only when the signal is triggered. This feature allows for some aspects of dynamic automata computing.

The routing matrix controls the distribution of signals to and from the automata elements, as programmed by the application. The routing matrix is a complex hierarchical lattice structure of groups of elements. While in an ideal theoretical model of automata every element can potentially be connected to every other element, the actual physical implementation in silicon imposes routing capacity limits related to tradeoffs in clock rate, propagation delay, die size, and power consumption. All programmable elements are subject to these capacity limitations placed by the routing matrix. Elements have a maximum in-degree and out-degree of 16. Automata with larger fan-in or fan-out requirements need duplication of elements. Handling large automata with very high edge densities may create enormous challenges on the efficient mapping of those automata onto the Automata Processor. This could lead to very large compilation times as well as low efficiency of the usage of the elements on the Automata Processor.

5.2.2 Avoiding scaling and routing pitfalls

By definition the Automata Processor reliably allows for the routing of as many as 16 ingoing and 16 outgoing connections for each element of the device. Even if these routing
limitations are met, and the number of utilized elements is small enough to fit on a single half-core, a solution still may not be achievable. Because placing and routing is a difficult problem, the ANML compiler may struggle with particularly dense and large graphs. Even if such a compilation is eventually successful, a large compilation time may not be acceptable for repeatedly configured solutions.

A tree structure is a common example of a difficult to compile graph. A tree structure may be used for the matching of all possible patterns for a small alphabet. Consider an example tree used to map genome bases \{A, C, G, T\} as displayed in Figure 5.1. Although such a tree may use a conservative number of elements and uses only a fraction of maximum outgoing connections for each element, compilation may struggle for large depths of the tree.

![Figure 5.1 Genome mapping tree solution](image)

Because the routing matrix relies in part on locality, the large fanning of a tree structure can explode outgoing connection requirements at a region level rather than an individual element level. To accommodate this, large trees may benefit from being manually decomposed into multiple graphs representing individual paths through the tree. In this way solutions for the Automata Processor may gain easier compilation at the cost of additional STEs. Figure 5.2 shows the decomposition of the previous tree into individual paths or strands.

In addition to a more direct and rapid compilation decomposing complicated graphs allows for simple scaling across multiple half-cores. If an automata is too large to fit in a single half core, the compiler will attempt to decompose the graph itself. While this may work it can lead to larger compile times, and can potentially introduce inefficient or partial use of multiple cores. By defining automata in small strands the compiler is capable of producing quick, accurate,
Even in situations where per-element routing is modest, scale must also be considered. Despite having no directly challenging connection properties, a series of singularly connected elements in a linked list fashion may cause compilation difficulties if it is several thousand elements long. Once again, the decomposition of such large and complicated graphs can aid compilation times at the expense of additional elements. If such difficult automata cannot be decomposed, considering a different approach to the target problem may be in order.

5.3 Designing robust solutions

While understanding the practical capabilities of the Automata Processor in modeling NFA is important, a complete solution may extend beyond the definition of a single automata. In reality a developer must also consider the cost of generating ANML and repeated compilation for exceptionally large tasks, as well as the managing of input and output data streams. Such extended considerations may influence the design of the NFA themselves and can be beneficial in maximizing efficiency across the entire development process.

5.3.1 Multiple automata instances

Often when producing a solution for the Automata Processor, the scale or variability of the problem requires multiple unique instantiations of a rank or core. It is possible that 1) A solution requires dynamically produced NFA based on the properties of the problem instance or 2) A solution requires a set of automata too large to be modeled in one instance of the
Automata Processor, or both. In such situations the generation, compilation, and load times for automata are not simple one-time costs, and must be considered in the total operation time of a solution. Minimizing these costs can contribute tremendously to the overall performance of a solution.

5.3.1.1 Managing multiple compilations

Solutions that require unique automata to be developed, compiled, and loaded onto the Automata Processor for each problem instance should be avoided whenever possible. If a problem absolutely requires the repeated production of such solutions, ANML generation and compilation times become targets for overall runtime reduction.

If unique ANML must be written for each problem instance, automation is essential. Although ANML generation applications may bear a large upfront development cost, they are the only reasonable solution for such repeated code production. While graphical editors may provide a convenient way to update automata, any such user-dependant modification introduces delays and potential for error across repeated solutions.

After ANML generation times are addressed, ANML compilation times must be addressed. Although compilation times can be difficult to predict, they are heavily dependent on the size and routing of the desired automata. It is strongly advised that if a solution requires such repeated ANML generation and compilation that the automata are as small and sparsely connected as possible.

5.3.1.2 Avoiding additional compilation

Although compilation time can be lowered significantly with careful graph design, where possible it can be best to design solutions that avoid compilation altogether. To avoid defining unique automata for each instance of a problem, singularly compiled automata can be implemented using one of two approaches: 1) The automata is designed to have configurable functionality based on some input metadata or 2) A generic automata can be defined for a problem where element place and route is consistant for all problem instances.
A reconfigurable functionality automata allows for different effective automata implementations without having to change any connections or values of any elements on the Automata Processor. Such a solution can be beneficial when a large amount of functionality is predefined and only a subsection of it is relevant for a given problem implementation. Supporting subgraphs can be added to a solution to enable or disable sections of the main graph based on metadata. The introduction of such extra streamed input data can reduce the effective processing rate of an automata, and should be minimized by design. Although metadata may extend the data stream, reconfigurable solutions will remain significantly faster than recompiled solutions.

If instances of a problem require uniquely configured automata that share a common element placement and routing, a different approach can be taken. Compiled and loaded automata can have some or all of their non-routing element properties redefined without compilation. This process is known as reflashing, and is an important strength of the Automata Processor. While slower than a reconfigurable functionality approach, reflashing is significantly faster than loading new automata and allows for the disabling of existing sections of the target automata. An example of basic reflashing can be seen in Figure 5.3. Here, a simple matching automata is reflashed to find different words of the same size.

### 5.3.2 Managing input and output data

While input and output data approaches may seem rigid with respect to a given problem, such properties are often quite flexible and can contribute significantly to the overall performance of a solution. Both input data and output reporting introduce unique and important considerations for the design of a comprehensive solution.

#### 5.3.2.1 Input data streams

For any given problem, a result or partial result is achieved when an automata instance reaches the end of the input data stream. This data stream in most cases is already in its simplest, most direct form and cannot be reduced. In some cases, automata may require additional
While metadata can allow singularly programmed automata to achieve a large variety of functionality, it is important to limit such additions to a reasonable amount. Note that if an automata requires a pumping delimiter character every third cycle, this metadata has effectively increased the data stream and consequently runtime by 50%. For optimal throughput metadata should be implied by state where possible, and minimized elsewhere. In an extreme case, specialized automata may be carefully designed to assume implied data that has been removed from the original data stream.

Although input data streams cannot be practically shrunk, in many cases they can be divided. Where possible, solutions can benefit from breaking an input stream into multiple
smaller streams that can be processed in parallel. These processing automata across multiple cores may be identical or may be specialized for their individual section of the original data stream. When designing solutions it is important to consider the possibility of such stream division as the large gains in throughput will often outweigh potential increased resource consumption.

### 5.3.2.2 Output handling

Output for the Automata Processor is an exceptionally important consideration for the design of efficient solutions. Each half of an Automata Processor core is divided into three separate output regions for a total of six across the core. When an STE from a given region reports a match, it is sent to the output event memory of that region. From there, each of the regions output events can be consolidated into the main output event buffer and finally reported. The process of transferring data from the output event memory to the main output event buffer costs roughly 40 input symbol cycles. Furthermore, this cost must be paid serially. If all six output regions report simultaneously, their collective results cannot be reported for $40 \times 6 = 240$ input symbol cycles.

From this there are two important lessons. First, it is important to consolidate output to as few regions as possible. Since each region can only support 1024 reporting lines, a solution may benefit from having fewer or more selective reporting cases. It can be advantageous to have multiple reporting elements with the same goal feed into a smaller representative set of reporting elements.

Secondly, it is important to minimize the amount of reporting in general. Where possible it can be advantageous to have multiple elements wait to report simultaneously or at the end of data. Solutions that require constant small reports of data with only a small number of cycles between each event will perform poorly. A cohesive solution must recognize that the CPU must still handle all of the data that is returned by the Automata Processor. A naive or greedy solution may even produce more data than it takes in, and such solutions should be considered with extreme caution.
CHAPTER 6 CONCLUSION

The Automata Processor is shown to be an extremely promising new processing architecture for large scale pattern matching applications. In the field of Bioinformatics, search for motif patterns in protein sequences provides one such application. PROSITE is a large annotated database of known protein motifs and currently contains 1308 known protein motifs expressed as patterns. Due to the large capacity of the Automata Processor, the automata for all these motif-patterns can be loaded and executed in the Automata Processor in parallel. This gives rise to large scale speed-up with respect to execution times of existing software. For example, all proteins present in the proteome of Escherichia coli (E.coli) can be scanned against all motif-patterns in less than 100 milliseconds. This is more than a 6000 times speed-up over published execution times. Furthermore, when the automata for all these motif-patterns are loaded onto the Automata Processor chip, they collectively take up about half of the resources of the chip. Since a single rank contains 8 Automata Processor chips and the automata for all motif-patterns which can fit into a single rank can be executed in parallel, the execution time for any given protein sequence will remain constant for a motif-database which is up to 16 times the current size.

Though PROTOMOTA currently provides a command-line interface only, it supports all the modes of operation provided by the PROSITE website, and more. In the first mode of operation, PROTOMOTA supports the scanning for all known motifs in a given protein sequence; or all common motifs in multiple protein sequences. Here PROTOMOTA not only provides a more fine-grained control over motifs being searched for, but also does not place an upper limit of 8 on the number of protein sequences as in the case of the PROSITE website. In the second mode of operation the presence of a particular motif can be detected in all proteins
in the database.

PROTOMATA was designed not only to aid biologists in the characterization of a vastly growing number of newly sequenced proteins, but also to serve as a guide for application developers in the field to develop programs on this new accelerator architecture. Therefore, the automaton for any motif is defined in a modular and easy to understand manner using the macros. The script to generate the automaton from a motif-pattern is also provided. This also makes it easy to handle new motifs which will be added to the PROSITE database or are under the investigation of a user.

Future work in this direction consists of executing this software on the real Automata Processor board and making the same available to everybody through a web-based interface. This interface may complement or be part of the PROSITE website. Further, the lessons learnt while developing the PROTOMOTA application (and mentioned in the application developers notes) would play a more significant role while developing larger and more complex bioinformatics problems in the future.

In general, more and more applications in various fields are being successfully designed and implemented on the Automata Processor. The processor has already been shown to considerably accelerate applications in the field of network security, image processing and machine learning, etc. Future generations of the processor which place lower restrictions on the resource requirements of the automata design are already in the works. Though thinking in terms of NFA can be difficult at first, but it paves the way for more natural exploration of difficult problems. The advent of the Automata Processor will provide a way to execute these NFA directly on the processor. With its large capacity to execute thousands of automata in parallel, it will a promising architecture for which to watch out.
Bibliography

[1] Perl compatible regular expressions.


