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IMPLEMENTATION OF GENETIC ALGORITHMS IN FPGA-BASED RECONFIGURABLE COMPUTING SYSTEMS

A Thesis
Presented to
the Graduate School of
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In Partial Fulfillment
of the Requirements for the Degree
Master of Science
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by
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ABSTRACT

Genetic Algorithms (GAs) are used to solve many optimization problems in science and engineering. GA is a heuristics approach which relies largely on random numbers to determine the approximate solution of an optimization problem. We use the Mersenne Twister Algorithm (MTA) to generate a non-overlapping sequence of random numbers with a period of $2^{19937}-1$. The random numbers are generated from a state vector that consists of 624 elements. Our work on state vector generation and the GA implementation targets the solution of a flow-line scheduling problem where the flow-lines have jobs to process and the goal is to find a suitable completion time for all jobs using a GA. The state vector generation algorithm (MTA) performs poorly in traditional von Neumann architectures due to its poor temporal and spatial locality. Therefore its performance is limited by the speed at which we can access memory. With an approximate increase of processor performance by 60% per year and a drop of memory latency only 7% per year, a new approach is needed for performance improvement. On the other hand, the GA implementation in a general-purpose microprocessor, though performs reasonably well, has scope for performance gain in a parallel implementation. The parallel implementation of the GA can work as a kernel for applications that uses a GA to reach a solution. Our approach is to implement the state vector generation process and the GA in an FPGA-based Reconfigurable Computing (RC) system with the goal of improving the overall performance.

Application design for FPGA-based RC systems is not trivial and the performance improvement is not guaranteed. Designing for RC systems requires algorithmic
parallelism in order to exploit the inherent parallelism of the FPGA. We are using a high-level language that provides a level of abstraction from the lower-level hardware in the RC system making it difficult to fully exploit some of the architectural benefits of the FPGA. Considering these factors, we improve the state vector generation process algorithmically. Our implementation generates state vectors 5X faster than the previous implementation in an Intel Xeon microprocessor of 2GHz. The modified algorithm is also implemented in a Xilinx Virtex-4 FPGA that results in a 2.4X speedup. Improvement in this preprocessing step accelerates GA application performance as random numbers are generated from these state vectors for the genetic operators. We simulate the basic operations of a GA in an FPGA to study its behavior in a parallel environment and analyze the results. The initial FPGA implementation of the GA runs about 7X slower than its microprocessor counterpart. The reasons are explained along with suggestions for improvement and future work.
DEDICATION

I dedicate this work to all of them who continually encourage me strive for the best.
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CHAPTER 1

INTRODUCTION

Genetic Algorithms have important applications in problems related to optimization, machine learning, game theory, design automation, evolvable hardware, distributed systems, network security, bioinformatics and many more. Genetic algorithms are iterative procedures that work on groups of solution representations called chromosomes. Each chromosome is composed of smaller segments of data called genes. A set of chromosomes together form a population. We generally initialize each gene in each chromosome randomly. The basic iterative work of the genetic algorithm is evolution from one population say $t$, to the next population, $t+1$. This evolution is done through the application of genetic operators – Selection, Crossover and Mutation, which introduce many random elements from one population to the next. Through this iterative procedure, the solution of the optimization problem evolves toward a better one.

This research is based on the work of Kurz [1] on scheduling industrial flow-lines. These flow-lines have sequence-dependent setup time, i.e. setup times depend on the order jobs are scheduled to the machines. The flow-line has several stages in series. Each stage contains a different number of machines and each machine has different jobs. Machines in parallel are identical in capability and processing rate. The flow-line is flexible in the sense that jobs may skip stages. Given the above conditions, the problem is to find a schedule that will result in an acceptable completion time of all jobs. The sequence dependent setup time makes this a general case of the Travelling Salesman Problem (TSP), and thereby an NP-Hard optimization problem. The solution
representation of this flow-line scheduling problem is analogous to the chromosomes in
the GA representation.

For the purpose of this research, chromosomes of a GA represent the order in
which jobs are processed in one stage. There are other versions of the algorithm where
chromosomes represent jobs in more than one stage. But for this algorithm, the order of
jobs in the remaining stages depends on the order of jobs at the first stage only. That is,
randomness comes into play only at the first stage and all other stages are deterministic.
Each gene of a chromosome has a value that is generated randomly. Based on these
values, jobs are sorted and assigned to machines. The goal is to find a combination of
jobs in stages that will result in a satisfactory makespan for the flow-line, where
makespan is the max completion time of all jobs. Through various genetic operations like
Crossover and Mutation, the GA tries to reach this goal. These genetic operations
introduce randomness in the GA process.

After each iteration of the GA, a specific set of genetic operators and parameters
known as a configuration is obtained. To arrive at a better solution for the optimization
problem, we must determine which configuration is better. That is, which configuration
results in the lowest makespan for the flow-line scheduling. To obtain different
configurations, we need an independent set of random numbers. If one iteration of a GA
uses upto 600 million random numbers, 600 million random numbers are needed to
produce one configuration. In order to facilitate appropriate statistical analysis, the sets of
random numbers should be non-overlapping, so that the assumption of independent sets
of random numbers can be made. The Mersenne Twister Algorithm [2] facilitates this, as
it has a period of $2^{19937}-1$, meaning it can provide sufficient random numbers before repeating. This period is in contrast to the basic `rand` function in the standard C library, which has a period of just over 32,600.

For solving the flow-line scheduling problem, researchers generally consider three different versions of the GA. We call each of them an algorithm. In order to determine reasonable performance measures, most GA research requires each algorithm to be executed many times [3,4], such as 50 times, per input data set. The input data set for the flow-line scheduling problem consists of number of stages, number of machines per stage, number of jobs and setup, and ready and processing time for each job. The flow-line scheduling may have different problem types, i.e. scheduling may be for different industries leading to different requirements. The input values of the data set may vary, resulting in different input files. For our optimization problem, there are 180 different problem types, 10 different input files for each with 3 different algorithms, totaling 5,400 files. If we consider only the simplest algorithm, then one replication (180×10=1,800 files per algorithm) requires 45 hours in a single core Pentium IV 3GHz HT machine which would scale to 80 days of run time to complete the necessary 50 replications of a single data set. Kurz abandoned this research approach due to the immense computational time until discovering the task parallelism potential of Condor Grid computing.

However, while the introduction of the grid environment of Condor removed the barriers of excessive computational time, managing the random number usage became problematic. Though the use of MTA solves the problem of generating a non-overlapping sequence of random numbers, ensuring that each run uses a non-overlapping
stream of random numbers generated from the same seed for replicability must be considered. For example, we could use the first set of 600 million numbers for iteration 1, then the second set for iteration 2. In a traditional computing environment, we could just allow the 2\textsuperscript{nd} iteration to start when the 1\textsuperscript{st} iteration left off. But in grid computing, iteration 1 and 2 may be running simultaneously. In that case, iteration 2 must first generate and throw away 600 million random numbers and then begin its work. This approach, while functionally correct, requires over 4,000 days to burn through the 600 million numbers before reaching the second set on the 250,000\textsuperscript{th} iteration. Each iteration requires approximately 45 hours of computation time making the overhead unacceptable.

In contrast, we could generate the random numbers offline and store them as an additional input file for each run. However, storage becomes an issue as the file size for 600 million numbers requires over 3 GB. The 50 replications required for just one data set equates to 150 GB of storage. Again, while the idea is nominally feasible for a small experiment, the storage requirements render this approach infeasible for the general case.

Fortunately, the MTA has an internal state, which is exposed in a structure composed of one integer and 624 values of unsigned integer or unsigned long. So, while we echo the sentiment of generating many random numbers offline, we only need to store the algorithm state, in a state file, at set intervals. Then, we can read in the state information and begin the new generation from that point, reducing the storage space requirement. In previous work, Kurz has generated and saved 360,000 state files that are 1 billion random numbers apart. This generation took about 22 days to complete on a dual core AMD Opteron 885 @ 2.6 GHz.
Due to their inherent parallelism, FPGAs are well suited for applications that have some form of parallelism in their characteristics. If an application can be designed in a way so that it can exploit the parallelism of an FPGA, we can have a significant performance gain over its general-purpose microprocessor counterpart. As FPGAs run at a much lower clock frequency, any performance gain is achieved at much lower power. But these gains are not free of cost. The price is paid in terms of resource utilization. FPGAs are equipped with on chip resources like Block RAM, DSP units and on-board memories like SRAM, SDRAM etc. An application must maximize the utilization of these resources to maximally exploit the inherent parallelism of an FPGA.

In this thesis, we present an improved state file generation algorithm which is 5X faster than its previous implementation on an Intel Xeon 5130@2GHz. Porting this code to an FPGA gives a modest 2.4X speedup due to several conditional statements that limit the performance. For our purpose, we must save state vectors at one billion number intervals, meaning we need to iterate through the original MTA algorithm one billion times before saving one state vector. We modify the algorithm such that it does not need to iterate one billion times. Also we eliminate the random number tempering portion of the original MTA algorithm as those are not required when generating state files. These two factors provide the speedup while generating state vectors. The previous GA implementation of this flow-line scheduling problem was designed for a traditional von Neumann architecture. After profiling the original code, hardware suitable functions were implemented on the FPGA. We implement the basic computations of the GA in an FPGA and study its performance while generating and feeding the random numbers to the GA.
process inside the FPGA. The performance is compared to its original implementation in
a general-purpose microprocessor. A comprehensive analysis of result is given along with
directions for future improvements.

We summarize the results as follows:

• Speedup in state vector generation using the Mersenne Twister Algorithm:
  5X in general-purpose microprocessor and 2.4X in an FPGA.
• A comprehensive study of the simulation results and measured data of basic
  GA operations implemented in an FPGA.

The remainder of this thesis is organized as follows. Chapter II provides background
information, which includes a general description of Genetic Algorithms, how it is used
to solve the optimization problem of flow-line scheduling, justification for using MTA,
and the systems and tools used to conduct the experiments. Chapter III discusses how
different components of these experiments were modeled to fit within the constraints of
the FPGA–based systems used and also discusses the limitations of our design. Chapter
IV analyzes the performance and results of the random number generation and Genetic
Algorithm simulation process. And finally Chapter V offers conclusions and directions
for potential solutions of the limitations of our design.
CHAPTER 2

BACKGROUND

This chapter discusses how the solution of the flow-line scheduling problem is analogous to Genetic Algorithms, some previous work on MTA and GA implementations on FPGAs, and how our solution differs from them.

2.1 Genetic Algorithms and Flow Line Scheduling

In this research, the solution of a flow line scheduling problem is represented in terms of chromosomes and genes of a Genetic Algorithm. In this section, we will discuss GA details, how a flow-line scheduling is mapped to a GA, and why the Mersenne Twister Algorithm is used in this research.

2.1.1 Anatomy of a Genetic Algorithm

Genetic Algorithm, a heuristic based approach for solving optimization problems, was introduced by Holland [9]. A typical GA has two steps [5]: a representation of the solution domain that reflects the genetic representation in a genome and a fitness function to evaluate the fitness of the current representation. The solution representations are generally in bits but may vary based on the application. For example, for our flow-line scheduling problem, we have a double-precision floating-point representation. GAs employ the following general steps: Initialization, Selection, Crossover, and Mutation. The algorithm starts with the random initialization of the initial population. Each population has a number of chromosomes and each chromosome has a number of genes. Each gene is also initialized by a random number. In this research, we generate the random numbers using the Mersenne Twister Algorithm [2], which is further explained in
section 2.1.3. After initialization, two parents are selected to generate their successor during the Selection stage. This selection is based on a fitness function [5] as parents of higher fitness values are expected to produce a better next generation. To generate the successor, the GA uses the Crossover operation where a crossover point is selected randomly. In the successor, solutions from the first parent are selected before the crossover point. Solutions after the crossover point are taken from the second parent. After Crossover, the Mutation operation is applied to increase the probability of the fitness of the solution. In Mutation, a random gene of the successor chromosome is changed with some probability. This process continues until the stopping criteria are satisfied. The probability of Mutation is a constant that is dependent on the application. Theoretically, the best set of chromosomes is expected to survive eventually. The overall GA process is shown in Figure 2.1. The Crossover and Mutation operations are shown in detail in Figure 2.2 and 2.3.

Figure 2.1: Anatomy of a Genetic Algorithm
In this research, the Mutation operation of the GA is replaced by the Immigration operation. In Mutation, one specific gene of a chromosome is changed with some probability. But in Immigration, a fresh new set of chromosomes are immigrated into the next generation of the population. That is, all genes of those chromosomes are replaced with a random value. How many chromosomes will be immigrated depends on a predefined constant and is generally determined by the given optimization problem.

2.1.2 Modeling Optimization Problems into Genetic Algorithms

Our target optimization problem is a flow-line scheduling problem that is very common in industrial manufacturing. These manufacturing systems have taken many forms with the added complexity of limited resources, time constraints, complicated process plans etc. For example, flow-lines of the semiconductor industry have multiple
machines in each stage and jobs revisit previous stages multiple times [1]. Another example is in the printed circuit board industry where jobs may skip stages depending on the circuit board specification. Each of these industries has different scheduling objectives but minimizing the overall completion time of all jobs, i.e. makespan can be considered a generic goal. These common goals are why operation researchers have focused on the makespan criterion for optimization.

Figure 2.4: Flow-Line Scheduling Problem.

Figure 2.4 shows a simple representation of our target flow-line scheduling problem. The flow-line has a number of stages. Each stage has machines and each machine has some number of jobs. This flow-line is also “hybrid” since multiple identical machines can run in parallel at some stages. Jobs are processed at exactly one machine per stage if they do not skip that stage. Also we call this flow-line “flexible” since jobs may skip stages. A job may not revisit a stage that it has already visited. We make the following assumptions [1] for the purpose of this research:

- All input data are known deterministically.
- Machines are available continuously with no breakdowns and no scheduled or
unscheduled maintenance.

• Jobs are non-preemptive, processed without error, and have no associated priority.

• Jobs are available for processing at a stage as soon as they have finished processing at the previous stage.

• The ready time for a job is the maximum time it takes to complete processing in the previous stages.

• Non-anticipatory sequence dependent setup times exist between jobs at a stage.

• Machines cannot be blocked because the current job has nowhere to go, i.e. infinite buffer exists before, after, and between stages.

• Machines in parallel are identical in capabilities and processing rate.

We are given the number of stages, number of machines in each stage, number of jobs, setup and ready time of each job, etc. as input data for the problem. The goal is to find a schedule that is suitable as a solution. The solution representation for this flow-line scheduling problem is analogous to a GA where the genes inside a chromosome represent the order in which jobs are processed in a stage. One important point to consider is that there is no direct mapping of input data for the problem to the GA representation; the GA is only for the solution representation. The only thing we can directly map from a given problem to the solution are jobs: a job represents a gene in the solution space. A closer mapping of the problem data to solution space is shown in Figure 2.5.
To express the problem and the stopping criteria addressed in this research, we use the following definitions [1]:

- $n$: number of jobs to be scheduled
- $g$: number of serial stages
- $m^t$: number of machines at stage $t$
- $p^t_i$: processing time for job $i$ at stage $t$ (assumed to be integral)
- $s^t_{ij}$: setup time from job $i$ to job $j$ at stage $t$
- $S_t$: set of jobs that visit stage $t = \{i: p^t_i > 0\}$
- $z$: makespan

For this flow-line scheduling problem, we apply the restriction that each stage must be visited by at least as many jobs as there are machines in that stage. If this is not
true then there is no problem for job scheduling. The goal is to find a feasible solution subject to many constraints. We can formulate the problem [1] as:

$$ P: \min z $$

(1)

That is, we want to minimize makespan. So eq. (1) is our objective function.

For this flow-line scheduling problem, we also make the following assumptions [1]:

- Stages are independent except that stage $t$’s completion time is stage $t+1$’s ready time.
- Setup times are such that an optimal solution will always exist.

Each chromosome is evaluated to check whether it satisfies some stopping criteria, i.e. whether the current schedule results in an optimal solution. The optimal solution for a specific problem needs to meet the lower bound requirement defined by eq. (2) and (3).

$$ LB^{(1)} = \max_{i=1,\ldots,n} \left\{ \sum_{t \in S_i} \left( p_i^t + \min_{j=0,\ldots,n} s_{ji}^t \right) \right\} $$

(2)

$$ LB^{(2)} = \max_{i=1,\ldots,n} \left\{ \min_{i \in S} \sum_{t=1}^{l-1} \left( p_i^t + \min_{j=0,\ldots,n} s_{ji}^t \right) + \sum_{t \in S} \left( p_i^t + \min_{j=0,\ldots,n} s_{ji}^t \right) + \min_{i \in S} \sum_{t=1}^{l-1} \left( p_i^t + \min_{j=0,\ldots,n} s_{ji}^t \right) \right\} $$

(3)

$LB^{(1)}$ is a job based bound and $LB^{(2)}$ is machine based [1]. For the job based bound, every job must be setup and processed at every stage. Setup requires minimal amount of time for setting up job $i$. For the machine based bound, every stage $t$ needs time for processing job 0. It also needs time for preemptive processing and a minimal setup time for the rest of the jobs. We can also consider minimal time to get to the stage and minimal time after finishing that stage. In our implementation, we consider the larger of
these two lower bounds as our stopping criteria. Once the best chromosome’s makespan hits that lower bound, we stop our iterations of the GA. Otherwise we continue evaluating all the chromosomes with basic GA operations until we exhaust all of them or hit a lower bound, whichever comes first.

2.1.3 Mersenne Twister Algorithm

As evident from the GA process, its operation largely depends on random numbers. For this implementation, the random numbers are generated using the Mersenne Twister Algorithm [2]. MTA is a uniform pseudo random number generator. It has a period of $2^{19937} - 1$ and 623-dimensional equidistribution up to 32 bit accuracy [2]. Such a long period implies that it generates $2^{19937} - 1$ random numbers before repeating. This non-overlapping sequence is large enough for our problems of intent. Also the very high order of dimensional equidistribution implies that there is very negligible correlation between successive values of output sequence [10]. MTA passes diehard tests [22] and numerous other tests of randomness [23]. This algorithm is designed specifically for Monte Carlo and other statistical simulations, but it is not suitable for Cryptography as observing a sufficient number of iterations (624 in this case) will lead one to predict the rest of the iterations. In this research, MTA is chosen because of its long non-overlapping period.

MTA is a twisted feedback generalized shift register [11], the algorithm is based on the recurrence relation eq. (4):

$$x_{k+n} := x_{k+m} \oplus (x'_k | x'_{k+1}) A$$  \hspace{1cm} (4)
Here,

\[ n \] degree of recurrence
\[ w \] word width (in number of bits)
\[ m \] middle word or the number of parallel sequences, \( 1 \leq m \leq n \)
\[ u,l \] Mersenne Twister tempering bit shift
\[ x \] a word of width \( w \)
\[ x',x'' \] \( x \) with lower and upper mask applied
\[ A \] matrix that contains twist information
\[ k \] constant with values 0,1,…

Figure 2.7 shows the MTA algorithm that generates 32-bit random numbers. As mentioned earlier, we require a maximum of 600 million random numbers.

One interesting advantage of MTA is that random numbers can be generated from a state vector that was saved before actually generating the numbers. These state vectors work as an entry point for a specific sequence of random numbers. State vectors are the specific states of the MTA after a sequence of random numbers and can be later used to regenerate the same sequence of random numbers. Though for this project, we need a maximum of 600 million random numbers, vectors are saved after one billion for more general-purpose use. Although saving state vectors at one billion number intervals adds to the poor performance of state vector generation process, this has the benefit of lower storage requirement (fewer state vectors). Figure 2.6 shows a snapshot of this process.
1. int[0..623] MT
2. function initializeGenerator(int seed) {
   3. MT[0] := seed
   4. for i from 1 to 623 {
      5. MT[i] := last 32 bits of(1812433253 * (MT[i-1] xor (right shift by 30 bits(MT[i-1]))) + i) }
11. function extractNumber() {
   12.  if index == 0 {generateNumbers()}
   13.  int y := MT[index]
   14.  y := y xor (right shift by 11 bits(y))
   15.  y := y xor (left shift by 7 bits(y) and (2636928640))
   16.  y := y xor (left shift by 15 bits(y) and (4022730752))
   17.  y := y xor (right shift by 18 bits(y))
   18.  index := (index + 1) mod 624
   19.  return y}
21. function generateNumbers() {
   22. for i from 0 to 623 {
      23. int y := 32nd bit of(MT[i]) + last 31 bits of(MT[(i+1) mod 624])
      24. MT[i] := MT[(i + 397) mod 624] xor (right shift by 1 bit(y))
      25. if (y mod 2) == 1 {
         26. MT[i] := MT[i] xor (2567483615) }}

Figure 2.6: Saving State Vectors of MTA

Figure 2.7: Mersenne Twister Algorithm - MT19937
2.2 Related Research

There are a number of publications that discuss implementing the Mersenne Twister Algorithm and Genetic Algorithms in FPGAs. Ishaan et al. [6] did parallel implementations of 32, 64, 128–bit SIMD MTA on Xilinx Virtex–II Pro FPGAs. They used interleaved and chunked parallelism and showed how the ‘jump ahead’ technique can produce multiple independent sequences to yield higher throughput. Shrutisagar et al. [12] worked on partial pipelining and sub-expression elimination to increase the throughput per clock cycle on the RC1000 FPGA Development platform that is equipped with Xilinx XCV2000E FPGAs. Both FPGA implementations of MTA used VHDL whereas ours is implemented in High-Level Language DIME-C [18]. Hossam et al. [7] implemented the basic GA modules along with the random number generator module in three different types of Xilinx FPGAs: XC4005, SPARTAN2 XC2S100-5-tq144, and Virtex XCV800 using VHDL and Mentor Graphics tools. They tested their design in applications ranging from thermistor data processing, linear function interpolation, and computation of vehicle lateral interpolation to test how the design performs with respect to producing the optimal solutions. Tatsuhiro et al. [8] designed two tools to facilitate the hardware design of GAs to predict the synthesis results based on input parameters, number of parallel pipelines, etc. Edson et al. [13] implemented a parallel and reconfigurable architecture for synthesizing combinational circuits using GAs. Paul and Brent [14] implemented a parallel GA for optimizing symmetric Traveling Salesman Problem (TSPs) using Splash 2. Emam et al. [15] introduced an FPGA-based GA for blind signal separation.
2.3 Our Approach

In all the previous research, the MTA or GA is a customized implementation specifically targeting the architecture, in this case FPGAs. Our approach significantly differs as we try to accelerate an existing application originally designed for von Neumann architectures. Both approaches have their own advantages and disadvantages. In the previous research, though they have achieved a performance gain in GA process, they do not consider how it performs when the GA works as a part of a larger application. In our approach, the probability of overall application acceleration is low as the original application design never considered exploiting parallelism. But this approach demonstrates what can happen when the GA is a small part of an application which was not originally designed for a parallel architecture. Also it shows us the necessity of designing and implementing an application specifically to take advantage of the parallel architecture. Our approach uses the high-level language DIME-C, but to the best of our knowledge, all the previous work used hardware description languages such as VHDL or Verilog.
CHAPTER 3

DESIGN AND IMPLEMENTATION

Our design and implementations are divided into two main parts. The first step is to design and implement state vector generation process in the FPGA. These state vectors are an integral part of the GA as they are used to generate the required random numbers. The second step is to design and implement the basic GA operations in an FPGA along with the generation of random numbers from the previously stored state vectors. Before designing the system for FPGA implementation, we conducted function profiling of the existing GA implementation that was written for a general-purpose microprocessor. From the profile data, we identified critical code segments for possible implementation in the FPGA and analyzed the issues related to hardware/software partitioning. We designed an improved algorithm for generating state vectors using the MTA which is 5X faster than its previous implementation in a general-purpose microprocessor and 2.4X faster than the previous FPGA implementation. We also implemented the basic GA operations in an FPGA. This chapter discusses and justifies our hardware/software partitioning approach, systems and tools used, implementation model, and design and implementation techniques for the state vector generation and GA operations. Finally we discuss the limitations of our approach.

3.1 Hardware/Software Partitioning

For a given application, a hardware/software partition maps each region of the application onto hardware (ASIC or Reconfigurable Logic) or software (microprocessor).
That is, a partition is a complete mapping of an application to either hardware or software. The goal of the partition is to maximize performance within the constraint of limited resources, in this case one Xilinx Virtex-4 LX100 FPGA.

There are several issues to consider for hardware/software partitioning. Some of them are listed below:

- **Granularity**: types of regions to consider.
- **Partition evaluation**: determining the goodness of the partition.
- **Alternative region implementation**: alternatives of hardware implementation.
- **Implementation model**: interfacing between microprocessor and FPGA.
- **Exploration**: finding good solution quickly.

Granularity is of two types: coarse and fine. If we partition based on tasks, functions and loops, that is called coarse-grained partitioning. On the other hand, fine-grained partitioning partitions regions based on code blocks, statements and operations. Both approaches have their own advantages and disadvantages. Therefore a heterogeneous granularity may be considered to take advantage of both extremes. The most intuitive approach to partitioning an application is based on its functions, i.e. coarse-grained partitioning. Also, coarse-grained partitioning may result in more accurate estimations during partition evaluation as it does not require the combination of several small regions and their communication overhead. An important disadvantage of coarse-grained partitioning is that it often has less inter-partition communication. That means, more data communication occurs between the host processor and FPGA than among different
Processing Elements (PEs) inside the FPGA. This situation may outweigh the benefit of implementing regions in hardware as the hardware/software communication is generally expensive. On the other hand, fine-grained partitioning gives more control over the exploitation of parallelism and less communication overhead between host processor and FPGA. But it is not intuitive and so generally takes longer to find a good partition. Also, estimation during partition evaluation is more difficult in this case due to their inter-partition communication.

We use gprof for function profiling of the original GA implementation that was targeted for a general-purpose microprocessor. After profiling, we have two types of profile data: flat profile and call graph.

### 3.1.1 MTA Partitioning Analysis

The hardware/software partitioning for state vector generation using MTA is straightforward. We did not profile the MTA implementation as the main computation occurs in a single function called `genrand_32`. Therefore it is obvious that we have to implement that function in the FPGA. The partitioning of the MTA is a coarse-grained partitioning. We did not explore any other alternative regions for the FPGA implementation, as there were no other compute intensive functions. Therefore we had no options for partition evaluation. The implementation model of this partition interfaces between the microprocessor and FPGA using the PCI-X communication bus.
3.1.2 GA Partitioning Analysis

Unlike the MTA implementation, the GA implementation has about 2000 lines of code. Therefore we performed function profiling before deciding on hardware/software partitioning. The flat profile data for the GA application is shown in the pie-chart in Figure 3.1. The profile data looks challenging for hardware acceleration as it is not concentrated in a single (or few) function(s) that largely dominate the execution time, making the hardware/software partitioning decision difficult. The most time consuming function in the GA is \textit{InStage} (23.46\%), which checks if a specific job has entered any of the stages of the flow-line. Based on Amdahl’s law [16], we can state that the speedup of an application is limited by the portion of the program not being parallelized. So if we go by the rule of thumb, that is implementing the regions that contribute to the highest execution time, the maximum theoretical speedup we can achieve is:

\[
\text{Speedup} = \frac{1}{1-p} \quad \text{(5)}
\]

\[
= \frac{1}{1-0.2346} \\
= \frac{1}{0.7654} \\
= 1.3X
\]

Here, \(p\) is the portion of the code that is implemented in hardware and therefore parallelized. To find the maximum speedup, we assume infinite parallelism by implementing a code snippet in hardware. We see that even with these assumptions, the speedup is insignificant. Also there are other significant constraints when functions are implemented in hardware. For example, topology of the call graph to the function,
communication overhead between host and FPGA, amount of data passed, actual bandwidth of the communication interface, etc. Examining the *InStage* function, it is not ideally suitable for implementation in an FPGA as it has mainly conditional statements and no significant computation. Also it is called from many calling functions. So we must consider a different approach to the hardware/software partitioning.

Figure 3.1: Flat Profile of GA Process.

One approach is to implement more functions such that the cumulative execution
time of all the functions implemented in FPGA is closer to 80%. In that case, we can expect a theoretical speedup of 5X according to eq. (5). But this partition exposes some practical limitations mainly due to the topology of the call graph. Figure 3.2 shows a snapshot of the overall call graph of the GA process. Figures 3.3 and 3.4 are the two alternative partitions we evaluated.

Figure 3.2: Snap Shot of the Overall Call Graph
Figure 3.3: Call Graph for 1st Partition Evaluation

Figure 3.4: Call Graph for 2nd Partition Evaluation
InSage, SiftUp, GenerateDiscUnif, BestCompleteReadyOrder, GetStartComplete, and CreateSchedFromChrom is a set of functions that have a cumulative execution time of 83.35%. As evident from Figure 3.3, the biggest disadvantage of the GA code for hardware implementation is that functions are called from many different places and many times. So if we want to implement a function in an FPGA by minimizing the host-to-FPGA communication overhead, we need to implement at least some of its predecessors. A call graph of this nature exposes additional problems. For example, even if a function is suitable for implementation in an FPGA, its predecessor may not be. The predecessors may have many conditional and branch statements. These statements are very ill suited for FPGAs as FPGAs do not have branch prediction units whereas general purpose microprocessors are equipped with efficient branch prediction units. Also if we continue to implement the predecessors in the FPGA, at some point we will run out of resources. These characteristics of the existing GA implementation make the hardware/software partitioning more difficult. Especially the coarse-grained partitioning is very hard in this case. As mentioned before, the fine-grained partitioning is not intuitive. Considering these issues, we have implemented the basic GA operations (InitChrom, ParaUnifCross and Immigrate as shown in Figure 3.4) in the FPGA. In Figure 3.2, 3.3 and 3.4, the numbers above each function indicate the number of times it is called by its calling function. Our implementation follows a coarse-grained partitioning as we partition based on the functions.

The partition evaluation approach we have followed is based on an estimation of the overall application performance with the constraint of using one user FPGA. We
consider the cumulative execution time of the functions implemented in the FPGA, the amount of data passed to and from the FPGA, and the resources they consume as our estimation criteria. We have only explored the two possibilities shown in the two call graphs of Figure 3.3 and 3.4 and decided to go on with Figure 3.4. As Figure 3.4 incorporates the basic GA functions, it supports our claim of an implementation of the GA in an FPGA. Also the call graph associated with these GA operations is more contained than the call graph of Figure 3.3, minimizing the host-to-FPGA communication overhead.

Our implementation uses only one FPGA for two reasons. First, as we are using coarse-grained partitioning, there is less opportunity for inter-partition communication, so it is very unlikely that the computation result of one PE will be used by another. Therefore even if we chose to use more than one FPGA, it is highly probable that the results of one FPGA are not needed by the second one. In that case, the second FPGA depends on the data passed to it by the host which has higher communication cost than data passed from the first FPGA. Secondly, the more functions we implement in FPGA, the more complicated the overall application call tree becomes. Because of these factors, we have only implemented those functions in the FPGA that account for a cumulative execution time of 2.46%. Therefore the maximum theoretical speedup we can expect according to eq. (5) is 1.025X.
3.2 Systems and Tools Used

Our implementation was developed in a 2GHz Intel Xeon CPU populated with 2 Nallatech H101 PCI-XM FPGA accelerator boards. The specification [17] of this board is as follows:

- 1 user FPGA – Virtex 4 LX100 (XC4VLX100-10FF1148C)
- 4 banks of DDR2 SSRAM
  - Each has 4MB of memory, totaling 16 MB
  - Total bandwidth: 6.4GB/s
- 1 bank of DDR2 SDRAM
  - 512MB of memory
  - Bandwidth: 3.2GB/s
- 4 channel serial communication (board-to-board)
  - Bandwidth: 2.5 Gb/s
  - Latency: 340 ns
- PCI-X connectivity with host
  - 64-bit, 133MHz

Figure 3.5 shows the functional diagram of H101-PCIXM [17].
Figure 3.5: Functional Diagram of H101 PCI-XM

During the development process, we used DIME-C [18] which is a high-level language, rather than a hardware description language such as VHDL or Verilog. DIME-C [18] is a subset of ANSI-C and provides the programmer with the flexibility of writing code for the FPGA without having to focus on the hardware in detail. DIME-C compiles code to VHDL automatically. This VHDL module is then used as a Processing Element (PE) that works inside an FPGA. Then we use DIMETalk [19], a tool for designing the network that interfaces with the host. DIMETalk defines how the PEs are connected to
other on-chip or on-board resources. Using DIMETalk, we generate the bitstream and the sample host code to be used in the host application. The host code is then modified according to the application and the data that must be transferred. The host code is responsible for reading from and writing to the FPGA, switching on and off the FPGA, and other housekeeping operations.

3.3 Implementation Models

We will discuss four methods that comprise our implementation model. The methods are:

- Communication Methods
- Execution Methods
- Implementation Methods
- Configuration Methods

The Reconfigurable Processing Fabric (RPF), or FPGA, is used as a coprocessor in our system. Figure 3.6 shows the location of FPGA in the memory hierarchy.
Using the FPGA as a coprocessor has advantages and disadvantages [20]. Integration of the FPGA as a coprocessor to an existing computing system is simplified compared to the tightly coupled or loosely coupled RPF architectures. But the primary limitations are the restricted communication bandwidth and increased latency. Therefore this type of coupling is well suited for applications where the coprocessor can compute despite limited communication to and from the host. For example data streaming applications like digital signal processing and image processing are suitable for this type of coupling. This topology is one of the most important limiting factors in accelerating our GA application since it is not a streaming application.

There are two types of execution methods [20]: mutually exclusive and parallel. In the former, the FPGA and microprocessor never execute simultaneously whereas in the later, they can execute simultaneously to improve parallelism and performance. Our
implementation model is mutually exclusive, which also limits the achievable performance of the overall GA application. One advantage of the mutually exclusive execution method is that it makes the partition evaluation estimation easier. But in this case we are only exploring a few hardware/software partitions.

Implementation methods are of two types: separate datapath and fused datapath. In separate datapath, the flow of data is independent whereas in fused, the paths are fused to reduce the area overhead. Although in the fused datapath approach, the performance may suffer due to a longer circuit path. As we are using a high-level language to program the FPGA, we do not have control over these methods. Whether our implementation will use separate or fused datapaths depends on the FPGA platforms and tools we are using.

Dynamic reconfiguration and partial runtime reconfiguration are two configuration methods that increase the effective size of the FPGAs. Dynamic reconfiguration allows tasks to be time shared [21]. We do not use this feature as the logic resources are not the main limiting factor in our approach. Partial runtime reconfiguration allows configuration of a portion of the FPGA while other portions continue to operate. This approach can improve performance as it can execute code in a portion of the FPGA without interrupting the other segments of the FPGA that are running. But this feature is generally not supported in currently available high-performance RC systems.

3.4 Accelerating State Vector Generation

The original MTA algorithm is shown in Figure 3.7. For generating state vectors,
this algorithm iterates one billion times since we are saving state vectors at intervals of one billion numbers. That is why there is an outer loop (line 3 of Figure 3.7) that iterates one billion times. This outer loop limits the performance in the FPGA implementation since DIME-C can only unroll the innermost loop. Even with the two innermost loops (lines 6 and 9 shown in Figure 3.7), there are challenges in finding and exploiting parallelism. The primary limitation is the memory access pattern of the MTA algorithm. Inside the for-loop, the same location of \( mt \) array is read and written back. This array is an input to the FPGA passed from the host. In our implementation, we use BRAM to store that array. Therefore according to the DIME-C specification [18], an array stored in a BRAM is passed to the PE module. For this reason, even if the BRAMs are dual ported, the consecutive read/write inside the for-loop cannot occur in the same cycle. So even if the two for loops in Figure 3.7 are the innermost loops, they cannot be fully unrolled as they would violate this condition. As a result, these two innermost loops cannot be pipelined. Since we loop through the code body one billion times, this overhead is multiplied and causes poor performance of the MTA algorithm in the FPGA.

To avoid the problems associated with the memory access pattern, there are a couple of options to consider. Instead of using BRAM to store the data, one option is to use SRAM. We have four banks of SRAM available in the system. Data can be duplicated in those banks allowing read and write to the same location in different memory banks. This option, while it enables the loop unrolling, has the associated overhead of copying the modified \( mt \) back to the SRAM before the next iteration of the outermost loop. Performing this copy operation one billion times outweighs the benefits
of unrolling the innermost loops. Another option is to stripe the data across the SRAMs. For example, array location from 0 to $N-M-1$ can be stored in one SRAM and the remainder of the array in another. But if we use any language of sequential nature like DIME-C, we need to be careful in read/write operations so that appropriate order is maintained. The whole process becomes easier if we use a language which is inherently parallel. Therefore, rather than trying to improve the performance of the state vector generation process using the techniques mentioned above, our approach seeks to improve it algorithmically.

```c
1. unsigned int y;
2. static unsigned int mag01[2]={0x0U, 0x9908b0dfU};
3. for (i=1000000000-1;i>0;i--){
4.     if (mti >= N) {
5.         int kk;
6.         for (kk=0;kk<N-M;kk++) {
7.             y = (mt[kk]&UPPER_MASK) | (mt[kk+1]&LOWER_MASK);
8.             mt[kk] = mt[kk+M] ^ (y >> 1) ^ mag01[y & 0x1U];
9.         } for (;kk<N-1;kk++) {
10.            y = (mt[kk]&UPPER_MASK) | (mt[kk+1]&LOWER_MASK);
11.            mt[kk] = mt[kk+(M-N)] ^ (y >> 1) ^ mag01[y & 0x1U];
12.        } y = (mt[N-1]&UPPER_MASK) | (mt[0]&LOWER_MASK);
13.        mt[N-1] = mt[M-1] ^ (y >> 1) ^ mag01[y & 0x1U];
14.        mti = 0;
15.    } y = mt[mti++];
16.    y ^= (y >> 11);
17.    y ^= (y << 7) & 0x9d2c5680U;
18.    y ^= (y << 15) & 0xefc60000U;
19.    y ^= (y >> 18);
20.    return y;
21. }
```

Figure 3.7: Original MTA Algorithm

```c
1. unsigned int mag01[2]={0x0U, MATRIX_A};
2. unsigned int y;
```
3. int kk, I, if1=0, if2=0, else1=0, flag=0;
4. for (i=1000000000-1; i>0; i--){
5. if (mti>=N){
6.   flag=1;
7.   for (kk=0; kk<N-M; kk++) {
8.     y = (mt[kk] & UPPER_MASK) | (mt[kk+1] & LOWER_MASK);
9.     mt[kk] = mt[kk+M] ^ (y >> 1) ^ mag01[y & 0x1U];
10.  for (kk=N-M; kk<N-1; kk++) {
11.    y = (mt[kk] & UPPER_MASK) | (mt[kk+1] & LOWER_MASK);
12.    mt[kk] = mt[kk+(M-N)] ^ (y >> 1) ^ mag01[y & 0x1U];
13.   }
14.   y = (mt[N-1] & UPPER_MASK) | (mt[0] & LOWER_MASK);
15.   mt[N-1] = mt[M-1] ^ (y >> 1) ^ mag01[y & 0x1U];
16.   if (i-624>=0){
17.     mti=624;
18.     i=i-623; }
19.  else{
20.     mti=i;
21.     i=1; }
22. } else if( mti<624 && flag==0) {
23.   flag=1;
24.   i=i-(624-mti)+1;
25.   mti=624; }
26. }

Figure 3.8: Improved MTA for State Vector Generation

Examining Figure 3.7 more closely, we find that there are many unnecessary computations if we are only generating state vectors. First, outside the if statement, generation and tempering of random numbers (the shaded box in Figure 3.7, lines 16-21) are not necessary since we do not retain the random numbers while generating the state vectors. Another important issue is the if loop of line 4 is executed only when $mti$ is greater than or equal to $N$, therefore we do not need to loop through it one billion times. Considering these observations, the state vector generation algorithm is modified as shown in Figure 3.8. The modification occurs between lines 15-25 to reflect the
observations mentioned above.

![Figure 3.9: MTA Implementation in an FPGA](image)

Implementation of this algorithm in an FPGA is straightforward. The host sends the required input data to the BRAMs and collects the output data from BRAMs using the PCI-X communication bus. The MTA block in Figure 3.9 is the Processing Element (PE) in this case. The PE does the computations required to generate the state vectors using the MTA algorithm. The three BRAMs indicate the three parameters passed to the PE module as arrays, one for input $mt$ and the other two for output $mt$ and $mti$. The initial $mti$ is passed to the FPGA as a standalone integer. After the host receives one set of the output (state vector) from the FPGA, it writes the results in a file called the state file. One state file is used in one iteration of GA process. For our purposes, we need a maximum of six-hundred million random numbers. According to our algorithm, one state vector can generate one billion non-overlapping random numbers. Therefore we can generate sufficient random numbers for one GA iteration form one state vector. We generate 32-
bit unsigned integers as the state vectors though the flow-line scheduling research used 64-bit unsigned long numbers. This smaller data type is used because DIME-C does not support long integers. A workaround for generating 64-bit state vectors will be discussed in section 3.6.

3.5 Implementing Genetic Algorithm in FPGA

The design of the basic GA functions for FPGA implementation is similar to the MTA implementation in the sense that the current implementation of the GA only uses BRAMs, no external memory. First, the amount of data is small enough to fit in the BRAMs and second the GA is not a streaming application. The basic block diagram of the design used to implement the core GA functions is shown in Figure 3.10. Three PEs are implemented in the FPGA that correspond to the four main operations of GA. Each of the PEs requires random numbers that are generated using the previously saved state vectors. In our implementation, we generate the random numbers inside each of the PEs as they are required. In our implementation, different PEs are called from the host at different times. As each of the PEs generate random numbers from the state vector inside the FPGA, the mti value is updated inside them. The mti value along with the mt array is always passed back to the host from the PEs so that in the next call to PEs, the updated ones are used.

The PE labeled InitChrom initializes the population. In our implementation, the size of the population is 100, i.e. the population consists of 100 chromosomes. Each of the chromosomes is initialized with random gene values. These gene values are double-
precision floating-point numbers. The PE labeled *ParaUnifCross* works on selection and crossover steps of the GA. Therefore this PE needs data for all of the chromosomes. Random parents are selected for the crossover operation along with the random crossover point. After the crossover, the successors are generated, i.e. gene values of the chromosomes are updated. These updated values are passed back to the host to be used in other required computations of the flow-line scheduling problem. The PE labeled *Immigrate* performs the immigration operation discussed in section 2.1.1. The starting point of the immigration along with the gene values are the inputs to the PE. The gene values of the chromosomes are updated during immigration and passed back to the host via the PCI-X bus. The FPGA implementation of the basic GA operations is shown in Figure 3.10. As indicated by the dashed box outside the PEs, we have one single bitfile for the overall GA application, although each of the PEs are called at different times from the host to find a solution for our problem.

![Figure 3.10: Design of Basic GA Functions in the FPGA](image)

The overall GA implementation flow diagram is shown in Figure 3.11. Chromosomes
are initialized randomly. These chromosomes are then evaluated in host to see if it satisfies the optimization criteria. If not then the next three steps of the GA starts executing in the FPGA - selection of qualified parents for generating a better successor, crossover to produce the successor, and mutation (replaced by immigration in this implementation) to increase the fitness of the successor.

3.6 Limitations of Our Design

Since the GA is not a streaming application, it is not ideally suited for our system (where FPGA is a coprocessor) as discussed in section 3.3. There are a couple of options that can potentially improve the performance within the constraints of the system and tools. Our current design does not consider those options, but they are discussed below with justifications for not supporting them.

The original GA implementation for solving the flow-line scheduling problem was targeted for a general-purpose microprocessor. We take pieces of that code to implement in a FPGA based on profile data. This approach limits the overall performance
of the FPGA implementation as the original implementation never considered a specialized architecture. The topology of the call graph indicates a significant amount of communication overhead between the basic GA processes and other modules of the flow-line scheduling implementation. This communication overhead, though insignificant for a general-purpose microprocessor, is a limiting factor in an FPGA implementation. This bottleneck in performance results from the hardware/software partition, which separates the basic GA operations in the FPGA implementation from the overall application running in the host. That means, if we do not consider redesigning the application targeting the FPGA, we have to pay the price for communication overhead between the host and the FPGA. Therefore while targeting this application for the FPGA implementation, better performance is possible if the entire application were redesigned specifically for an FPGA-based platform.

We improved the state vector generation process algorithmically, but we have not incorporated the suggested memory access pattern discussed in section 3.4, which could further improve performance.

An important design strategy of most FPGA implementations is passing data to and from FPGA only when it is absolutely necessary. It is crucial to move data judicially as the corresponding overheads are significant. In our design, we pass data back to host whenever we have enough results for creating one state file. Though it does not have a significant impact for generating one state file, it has a cumulative impact on how quickly we can generate multiple state files.

Using a C-like programming language (such as DIME-C), though has the
advantage of higher productivity from the programmer’s point of view, makes it harder to take full advantage of the FPGA. Since DIME-C is still sequential in nature, the programmer needs to code explicitly to take advantage of the inherent parallelism of an FPGA. Conversely, as VHDL is inherently parallel, it is easier to exploit the parallelism of an FPGA.

The biggest disadvantage of this design comes when we consider the design of the GA algorithm in the FPGA. The random number generation is independent of the basic GA operations since they are produced from the previously generated state files, using variables that are independent of the basic GA operations. But we are not fully taking advantage of this available parallelism. Using the MTA algorithm and the previously generated state vectors, random numbers can be generated and stored in a FIFO from where these GA operators can access them as needed. This concept is shown in Figure 3.12. Our current implementation does not incorporate this approach and is considered for future work.

![Figure 3.12: Random Number Generation in Parallel Using a FIFO](image-url)
3.7 Summary

Our design and implementation techniques achieve a performance gain in the state vector generation but performance decreases in the overall GA process. Chapter 4 discusses the results and analyzes them within the constraints of the RC systems and tools used.
CHAPTER 4

PERFORMANCE AND RESULT ANALYSIS

This chapter presents and analyzes the performance results from the system’s perspective: the underlying hardware architecture, memory hierarchy, and communication interface. We also discuss how the algorithm interacts with the system and impacts the final results and performance.

4.1 Performance Improvement of State Vector Generation

Compared to the original implementation, the improved MTA algorithm for state vector generation, introduced in Chapter 3, performs 5X faster in the general-purpose microprocessor and 2.4 times faster than its original in the FPGA implementation. The original FPGA implementation executes at a clock frequency of 139MHz while the improved one runs at 157MHz. The clock frequency increase is due to the improvement in the algorithm. As shown in Figure 3.8, the improved algorithm does not iterate one billion times through the code body. Therefore it does not access the state vector array and the random numbers that many times. As a result, outputs are generated with fewer memory accesses resulting in a higher clock frequency. Table 4.1 shows a summary of the performance data of state vector generation algorithm.
Table 4.1: Performance Data of State Vector Generation Using MTA Algorithm

<table>
<thead>
<tr>
<th>Property</th>
<th>Original Algorithm</th>
<th>Improved Algorithm</th>
<th>Improvement</th>
</tr>
</thead>
<tbody>
<tr>
<td>S/W time for one state file (microprocessor)</td>
<td>45.3 sec</td>
<td>9 sec</td>
<td>5.03X</td>
</tr>
<tr>
<td>S/W time for one state file (FPGA implementation)</td>
<td>350 sec</td>
<td>146.44 sec</td>
<td>2.4X</td>
</tr>
<tr>
<td>H/W Cycles</td>
<td>12,724,078</td>
<td>9,892,294</td>
<td>1.488X</td>
</tr>
</tbody>
</table>

An important point to note from Table 4.1 is that the improved algorithm has about 5X speedup over its original general-purpose microprocessor implementation while only about 2.4X speedup in the FPGA implementation. The improved algorithm of Figure 3.8 has more conditional statements than the original, which is suitable for the von Neumann architecture but not ideal for an FPGA. As general-purpose microprocessors are equipped with efficient branch prediction units, efficient execution of the improved algorithm is not a problem. Since FPGAs do not have similar branch prediction units, a significant performance bottleneck results while executing conditional and branch statements. Without these branch prediction units, FPGAs may implement all the cases of conditional statements in the data path to improve performance. Whenever a conditional statement occurs, assessing that condition requires extra cycles that are not necessary in a general-purpose microprocessor. For most modern microprocessors, the branch prediction accuracy is more than 95%; meaning, in 95% cases, they do not execute the conditional
statements. They can safely assume the prediction result from Prediction History Table (PHT), a table where the previous conditional statement check results are stored. Based on the results of PHT, they move to the later stages of instruction execution. Only in less than 5% of the cases, the miss-prediction occurs and requires extra cycles to bring the proper set of instructions in the pipeline stages. As general-purpose microprocessors typically run 10X to 20X faster than FPGAs, the miss-prediction penalty of extra cycles is insignificant. In our design, the FPGA runs at around 15X slower clock frequency than the microprocessor. The cumulative effect of these issues results in the poor performance of an FPGA while executing conditional and branch statements.

Another important issue is the potential increase in area overhead when conditional statements are implemented in FPGAs. In lieu of branch prediction units, FPGAs can implement both data path conditions increasing the area requirement.

The number of hardware cycles required in the improved algorithm implementation is 9,892,294 whereas the original implementation requires 14,724,078 cycles, a 1.488X improvement. The hardware cycle count only considers the cycles required for an algorithm to execute in the FPGA. Whereas the runtimes discussed before also include the data transfers and other communication. 19.5KB of data is passed between the host and the FPGA and the communication overhead is 16 microseconds.

We use the following terms to define speedup of an algorithm in an FPGA compared to its general-purpose microprocessor implementation:

\[ S \quad \text{CPU clock cycles} \]

\[ H \quad \text{Hardware (FPGA in this case) cycles} \]
\[ F \quad \text{FPGA clock frequency} \]

\[ M \quad \text{Microprocessor clock frequency} \]

\[ Speedup = \frac{S}{H} \times \frac{F}{M} \quad (6) \]

For our system, \( M \) is 2 GHz.

For the original algorithm, \( F = 139 \text{ MHz}, H = 12,724,078 \text{ cycles} \) and for the improved algorithm, \( F = 157 \text{ MHz}, H = 9,892,294 \text{ cycles} \). To calculate \( S \), we will use eq. (7).

\[ \text{CPU clock cycles} = \text{CPU execution time} \times \text{Clock frequency} \quad (7) \]

CPU execution time is 45.3 sec for the original and 9 sec for the improved algorithm. Clock frequency is 2GHz. Therefore using eq. (7), \( S \) for original and improved algorithm are 45,300,000,000 cycles and 9,000,000,000 cycles respectively. Now using eq. (6), the speedup for the original algorithm in the FPGA implementation is 494.87X. The FPGA implementation of the original algorithm runs 494.87X faster than the general-purpose microprocessor if we consider the fact that the clock frequency of the FPGA is about 20X (in our design 14.388X) slower than the microprocessor. Conversely, the normalized speedup for the improved algorithm is 142.84X in the FPGA implementation. Even though the improved design runs at a higher clock frequency (157MHz), the performance is lower than the speedup of the original algorithm. As previously mentioned, the improved algorithm (improved for the microprocessor implementation) is not well suited for an FPGA as it has several conditional statements. Therefore the number of hardware cycles is not low enough to raise the speedup value based on eq. (6).

The resource utilization for both the original and improved algorithms are shown in Table 4.2. As compared to the other resources, the LUT requirement is significantly
reduced (30.44%) for the improved algorithm. Referring to the algorithm shown in Figure 3.8, the reduction in resource utilization results from not performing the random number tempering on the variable $y$. In other words, we are using the variable $y$ fewer times than in the original algorithm of Figure 3.7 contributing to this reduction in the resource requirements.

Table 4.2: Resource Utilization of State Vector Generation Algorithm

<table>
<thead>
<tr>
<th>Resource name</th>
<th>Original algorithm</th>
<th>Improved algorithm</th>
<th>% resource reduction</th>
</tr>
</thead>
<tbody>
<tr>
<td>BRAMs</td>
<td>51/240 (21%)</td>
<td>37/240 (15%)</td>
<td>27.45%</td>
</tr>
<tr>
<td>Slices</td>
<td>5451/49152 (11%)</td>
<td>4124/49152 (8%)</td>
<td>24.34%</td>
</tr>
<tr>
<td>4-input LUTs</td>
<td>8351/98304 (8%)</td>
<td>5809/98304 (5%)</td>
<td>30.44%</td>
</tr>
</tbody>
</table>

Both implementations pass 19.5KB of data between host and FPGAs through the PCI-X bus. This PCI-X bus is capable of passing 64-bit data with a maximum of 133MHz theoretical speed [24]. Therefore the maximum data transfer rate is 8.3125Gb/s. This data transfer rate is the theoretical rate and the effective rate is actually lower due to the overhead and other system design tradeoffs. The overheads include data transfer overhead, transaction layer packet overhead, flow control overhead, etc. [25]. Considering these factors, the sustainable host bandwidth is 400 MB/s [24]. As the data
passed between the host and FPGA is only about 19.5KB for one state file, the communication overhead is insignificant, i.e. 16 microseconds to be specific. But considering that we must generate many state files, this overhead can be significant. For example, the flow line scheduling requires generating 360K state files. With 16 microsecond communication overhead per file, the cumulative overhead is 5.76 seconds for all 360K files. One possible improvement of this design would be to store state vectors in on-board memory (like SRAM) and send them at set intervals or at the conclusion of the file generation taking maximum advantage of the PCI-X bus bandwidth. This approach will require analysis of the tradeoff between SRAM size and PCI-X bus bandwidth and when it is most efficient to send data back to the host. As the size of each SRAM is 4MB and the size of one state file is 20KB (625 unsigned integers), we can store a maximum of 200K state files in a SRAM. But the time to generate 4MB data is too long. With our improved algorithm, the generation of 4MB data will require about 339 days (200K files, each with 146.44 seconds). That means, if results are returned to the host once the SRAM is filled up, the user will have to wait for 339 days before she can see the first state file. Therefore there should be a tradeoff between the maximal use of PCI-X bandwidth and throughput rate based on the application requirement.
4.2 GA Performance Analysis

In this section we discuss the implementation results and performance of basic GA operations. Table 4.3 shows the performance results of each of the PEs of the GA process.

Table 4.3: Performance Data of Basic GA Operations.

<table>
<thead>
<tr>
<th>Property</th>
<th>InitChrom</th>
<th>ParaUnifCross</th>
<th>Immigrate</th>
</tr>
</thead>
<tbody>
<tr>
<td>S/W time (general-purpose microprocessor)</td>
<td>50 micro sec.</td>
<td>0.17 sec.</td>
<td>50 micro sec.</td>
</tr>
<tr>
<td>Total time (FPGA)</td>
<td>100 micro sec.</td>
<td>1.21 sec.</td>
<td>100 micro sec.</td>
</tr>
<tr>
<td>H/W Cycles</td>
<td>1612</td>
<td>3123</td>
<td>1116</td>
</tr>
<tr>
<td>FPGA design frequency</td>
<td>101.871MHz</td>
<td>130.162MHz</td>
<td>101.871MHz</td>
</tr>
</tbody>
</table>

As seen in the table, the InitChrom and Immigrate modules run 2X slower in the FPGA than the general-purpose microprocessor whereas ParaUnifCross runs 7.12X slower. The values of $S$ for these three modules are 50, 170 and 50 cycles respectively. Therefore, the speedup for InitChrom, ParaUnifCross and Immigrate based on eq. (6) are $1.58 \times 10^{-3}$, $2.28 \times 10^{-3}$ and $3.54 \times 10^{-3}$ respectively.

Even with the function by function comparison, the FPGA implementation is much less efficient as shown in the execution time comparison. This inefficiency occurs because the implementation of basic GA operations is not well suited for a FPGA. The reasons can be explained by analyzing the pseudocode in Figure 4.1.
1. void InitChrom(struct chrom *Chrom, struct data *PD) {
2.   int j;
3.   for (j=1; j<Chrom->numjobs; j++) {
4.     if (InStage(PD, j, 0)) {
5.       Chrom->StageChroms.keys[j]=GenerateDiscUnif(0,Chrom->StageChroms.nummcs - 1);
6.       Chrom->StageChroms.keys[j]=Chrom->StageChroms.keys[j]*(int)pow(10,Chrom->places);
7.       Chrom->StageChroms.keys[j]=Chrom->StageChroms.keys[j]+GenerateDiscUnif(1, (int)pow(10,Chrom->places)-1);
8.     }}
9.   else Chrom->StageChroms.keys[j]=0;
10. }
11.   Chrom->changed=TRUE;
12. }

Figure 4.1: InitChrom Function in General-Purpose Microprocessor

As shown in lines 4, 5 and 7 of Figure 4.1, different functions are called from this function with different parameters to initialize the chromosome. The InStage function is called with different parameters for each iteration. The GenerateDiscUnif function calls the random number generation process. Even with the in-lining of those functions, code of this nature is not ideally suited for an FPGA implementation. Also the if-else conditions of lines 4 and 9 add to the inefficiencies as discussed earlier. One possible improvement is to execute the InStage function independent of this code segment and store the results in a FIFO. In that case, this code segment can simply read the values from the FIFO and proceed.

If we want performance improvement of an application implemented in an FPGA verses its microprocessor counterpart, the algorithm should have some characteristics that will facilitate exploiting the architectural benefits of an FPGA. They are discussed below.
The algorithm should have some options for exploiting the inherent parallelism of an FPGA. This feature includes independent memory access, options for loop unrolling and loop flattening, independent tasks, etc. Even if the algorithm does not have these inherent characteristics, there should be options available to modify the application to fit to those features. This application is limited in this regard mainly because of the topology of the call graph. The topology led us to a coarse-grained partitioning whereas in most of the cases it would be better if we could do fine-grained partitioning. But fine-grained partitioning would make the partition combination tougher since the functions are called from many different places, and likely their combination overhead would not perform better.

Related to the previous point, an important question is whether the implementation suits the target architecture or not. As discussed in section 4.1, the existence of conditional and branch statements here in the GA process is a bottleneck for performance improvement. The call graph of this algorithm readily reveals the fact that there is significant jumping from one function to another adding to the probability of a performance bottleneck for an FPGA implementation.

Another point is whether the application is streaming or not. As we are using the FPGA as an independent coprocessor, having a streaming application would improve the chances of application acceleration. The GA fails here as it is not a streaming application.

For hardware/software partitioning, we chose coarse-grained partitioning. An important question is whether the fine-grained partitioning would help us in any way for the functions we chose to implement in the FPGA. Examining Figure 4.2, we conclude that
the function *ParaUnifCross* might have shown an improved performance if it were implemented to reflect a finer granularity.

```
1. void ParaUnifCross(struct data *PD, struct population *NewPop, 
    struct population *Pop){
2. int k, j, numcross, start, par1, par2, done;
3. double val;
4. static struct chrom temp1, temp2;
5. numcross = (int) (Pop->popsize * CROSS_PERCENT);
6. start = (int) (Pop->popsize * ELITE_PERCENT);
7. CreateChrom(PD, &temp1);
8. for (k=0; k<numcross; k++) {
9.    par1=GenerateDiscUnif(0, Pop->popsize-1);
10.   par2=GenerateDiscUnif(0, Pop->popsize-1);
11.   done=(par1!=par2);
12.   while (!done) {
13.      par2=GenerateDiscUnif(0, Pop->popsize-1);
14.      done=(par1!=par2);}
15.   for (j=1; j<temp1.numjobs; j++) {
16.      val=genrand_res53();
17.      if (val < PROBPAR1)
18.         temp1.StageChroms.keys[j]=Pop->
19.            Chroms[par1].StageChroms.keys[j];
20.      else
21.         temp1.StageChroms.keys[j]=Pop->
22.            Chroms[par2].StageChroms.keys[j];}
23.   CopyChrom(&NewPop->Chroms[k+start], &temp1); }
```

Figure 4.2: *ParaUnifCross* Function in General-Purpose Microprocessor

There are calls to different functions at lines 7, 9, 10, 13, 16 and 21 of Figure 4.2. A fine-grained partitioning (statement level) is expected to perform better in this case. The reason is the statements of lines 7, 9, 10, 13 and 16 do not depend on their previous statements. They depend only on the random numbers generated by the MTA algorithm using the state vectors. But then again it becomes complicated, as we need to consider
integrating it with the overall GA process and eventually with the flow-line scheduling application.

These reasons add to the inefficient runtime of the GA operators compared to its von Neumann counterpart. But an improved design as shown in Figure 3.12 is expected to improve the performance compared to the current FPGA implementation as in Figure 3.10.

Resource utilization for GA operators (individual resource consumption of PEs) is shown in Table 4.4. It is evident that the ParaUnifCross PE is consuming most of the FPGA resources. ParaUnifCross is actually a combination of selection and crossover operations of the GA so it has much more computation compared to the other PEs currently implemented. The overall resource utilization is almost equal to 100%. An important point to note is that in our current implementation, all the GA operations generate random numbers inside their own module. This approach actually increases the resource requirement. If the implementation reflects the design of Figure 3.12, we can eliminate the need for a random number generator per PE and thereby reduce the resource utilization.

<table>
<thead>
<tr>
<th>Resource Name</th>
<th>InitChrom</th>
<th>ParaUnifCross</th>
<th>Immigrate</th>
</tr>
</thead>
<tbody>
<tr>
<td>BRAMs</td>
<td>54/240 (22%)</td>
<td>109/240 (45%)</td>
<td>54/240 (22%)</td>
</tr>
<tr>
<td>Slices</td>
<td>13,397/49152 (27%)</td>
<td>17890/49152 (36%)</td>
<td>13,428/49152 (27%)</td>
</tr>
<tr>
<td>4-input LUTS</td>
<td>18,592/98304 (18%)</td>
<td>27341/98304 (27%)</td>
<td>18,689/98304 (19%)</td>
</tr>
</tbody>
</table>

Table 4.4: Resource Utilization of GA Operations.
When the three modules are combined to generate a single bitfile, the device utilization summary is:

- Number of Slices: $42634/49152$ (86%)
- Number of 4 input LUTs: $62169/98304$ (63%)
- Number of BRAMs: $200/240$ (83%)
- Number of DSP48s: $45/96$ (46%)

The combination of the three modules consumes 86% resources when all three modules have a random number generation process inside them. If we consider the three modules Initchrom, ParaUnifCross and Immigrate as one single module of the GA, then an important question is how many GA modules can we implement in an FPGA. Clearly, in our current implementation, we can only implement one GA module in a Virtex4-LX100. Our estimation shows that if we implement the design shown in Figure 3.12, we cannot implement more than two GA modules in the same target FPGA. One single GA module takes about 35% of the resource excluding the random number generation portion, so if we implement two GA modules along with one random number generator PE that stores numbers in a FIFO, we will exhaust the resources we have but likely improve the overall performance.

The maximum amount of data passed to the FPGA is for the ParaUnifCross PE, which is about 110 KB. Initchrom, and Immigrate require about 30KB and 16KB of data respectively. Unlike section 4.1, we do not have an option to pass data back to the host in a combined packet as these PEs are activated only when needed by the host application.
As discussed in section 3.1.2, the overall GA application speedup should ideally be 1.025X. But it is readily understandable that with the communication overhead and the design problems discussed in section 4.2 and 3.6, practically it would not be possible to achieve any speedup.

For our research, we have two separate bitfiles for state vector generation and the GA. Though the state vector generation is an initial step necessary for the operation of the GA, we consider the GA our overall application. Therefore the resource utilization in the state vector generation does not count towards the total resource utilization of the application. If we consider the resource utilization from Table 4.2 and Table 4.4, we find that the total percentage of BRAM usage is more than 100%. Since the bitfiles are separate and we do not run them simultaneously, this is not a problem.

The results for our current implementation, though not impressive, open the door for some interesting and seminal work in the future. We have exposed the issues and understand the requirements necessary to design and implement a high-performance computing application for a specialized architecture to gain maximum throughput within the given constraints. We will discuss these in chapter 5.
CHAPTER 5

CONCLUSION AND FUTURE WORK

The focus of our work has been the acceleration of the GA in an FPGA. We have shown that though there are challenges with the use of RC in scientific computing applications like GAs, these challenges can be overcome when certain conditions are met. Our contribution in this field is summarized in section 5.1. Later in section 5.2, we discuss in detail the ideas for future work that will make RC a better fit for GAs.

5.1 Conclusion

We have presented an improved algorithm for state vector generation using MTA. The algorithm eliminates some steps from MTA that are unnecessary for state vector generation. Though there are potential approaches to improve performance by using the FPGA resources more efficiently, the one billion iterations required before generating one state vector limits the achievable performance of an FPGA implementation. Therefore we chose to improve the performance by improving the algorithm. The improved algorithm runs 5X faster in a general-purpose microprocessor than its previous implementation and 2.4X faster in the FPGA than its previous FPGA implementation.

We have designed and implemented the basic GA operations in an FPGA. There are four basic operations of the GA [5] and they are represented in three different PEs in the FPGA. These three PE’s are combined to form a single GA module, meaning our implementation has a single bitstream that represents the GA operations although the three PE’s are called from the host at different times; there is only one active PE at any given time during the FPGA execution. Each of the basic GA operation requires random
numbers for its operation. These random numbers are generated from the state vectors inside each of the PEs. Our FPGA implementation of the GA runs about 7X slower than its microprocessor counterpart for several reasons. The original GA implementation was targeted for a general-purpose microprocessor and does not consider any specialized architectures like FPGAs. Therefore the implementation is not ideal for an FPGA. We do not design the application from scratch targeting our systems, rather we chose some portion of the original implementation based on the profiling results and perform coarse-grained partitioning to implement that in the FPGA. In some cases fine-grain partitioning may perform better, but due to the topology of the call graph, it will not give us an overall performance gain. Also we have not separated the random number generation process from the basic GA operation, which is expected to improve performance in general and allow for implementation of multiple GAs in a single FPGA. Overall, we have inefficiencies in the FPGA implementation of the GA. The reasons are discussed in detail in section 3.6, 4.1, and 4.2 along with suggestions for improvement.
5.2 Future Work

Possible future work involves both the state vector generation and the GA implementations. First, we can implement the application in hardware description languages (HDL) like VHDL or Verilog rather than using a high-level language (HLL) like DIME-C. As HDLs are inherently parallel, we will need less attention to explicitly specifying the parallelism in our implementation. As discussed in section 2.2, all previous work on MTA and GA are in HDLs. One reason is that HDLs allow the designer to exploit the parallelism of the FPGA in a more direct manner (not relying on the compiler to infer the parallelism). Another interesting future direction would be a comparison between the two implementation techniques: HDL and HLL. This comparison will show us how the programming languages and their compilation techniques impact the performance of an application targeted to a FPGA.

As discussed in section 3.6, the original implementation of the GA was targeted for a von Neumann architecture with no consideration of an accelerator or specialized hardware such as an FPGA-based RC platform. Most of the design constraints for our approach relate to the inability to expose parallelism or compute density in the original application. To exploit the parallelism of the RC system, we can design the application from the scratch targeting the FPGA-based systems. The key consideration of that design will be to reduce the communication overhead between the basic GA operations and other portions of the application. Therefore the design should support calling the basic GA module as few times as possible. In that case we will implement only the basic GA operations in the FPGA and the host will use the outputs for other computations. To
reduce the host-to-FPGA communication overhead, we can perform GA operations on the input data multiple times. This number will depend on the application and the host can pass an upper limit of that number to the FPGA. When the output for the first GA operation is available, the host can begin reading those results from the FPGA and proceed with its computation while the FPGA is computing and storing the GA outputs in an on-chip or on-board memory. This design would effectively support parallel execution of the host and the FPGA. As soon as the stopping criterion is satisfied, i.e. the solution is acceptable; the host will halt the execution of the FPGA. The reduced communication overhead along with the parallel execution of the host and the FPGA will account for a performance increase compared to our current implementation and most likely verses the von Neumann counterpart as well. Once we have a GA module like the one described above, we can use it to solve other optimization problems like placement and routing of an FPGA and others that use a GA to find their solutions.

Our current implementation does not take advantage of overlapping communication with computation. This limitation is a significant bottleneck for our current implementation of the state vector generation. Also we need to incorporate this overlapping if we want to implement the design techniques of Figure 3.12 or the technique described in the paragraph above. To hide these data transfer overheads, we could use on-board memories like SRAM or SDRAM or on-chip interfaces like FIFOs to store data temporarily before the host reads them.
Another technique of parallelizing the state vector generation process is to stripe data across memory banks so that at least some of the memory accesses, shown in Figure 3.8 (line 8, 9, 11 and 12) can happen simultaneously.

As discussed in section 3.6, allowing the random number generation to occur in parallel with the GA process by incorporating a FIFO will improve the performance of the application and is considered an important future direction.

Implementing the GA in other architectures like the Cell Broadband Engine (Cell BE) or a Graphics Processing Unit (GPU) may be an interesting area to explore. Each of these systems has its own advantages and disadvantages that may or may not suit an application. GPUs are suitable for streaming applications. As GA is not a streaming application, most likely we will not be able to achieve any significant performance gain by implementing the GA in a GPU. On the other hand, the 6 different Synergistic Processing Elements (SPE) of the Cell BE can be used to run 6 GA modules independently, thereby extracting parallelism out of the multicore architecture of the Cell BE.

Our future work also includes studying the behavior of a GA in a large-scale cluster. The primary focus is to work on a large-scale cluster of FPGAs but a cluster of Cell BE may also be considered once we have the implementation in one Cell BE. With the growing number of computing resources, performance and scalability of the GA implementation in these clusters are important metrics of analysis.

A relatively newer concept related to the GA is Evolvable Hardware (EH). The concept of EH arises from the analogy between a living being and a circuit. The DNA
(deoxyribonucleic acid) of a living being is a string of symbols from a quaternary alphabet (A, C, G, T). Similarly reconfigurable logic devices are configured by a bitstream that constitutes of binary symbols (0,1). This analogy suggests the possibility of applying the concepts of GA into circuit design. The traditional circuit design task is vulnerable to human error and the optimality of a solution cannot be guaranteed for larger circuits. Design automation is challenging for tool designers and with the increasing complexity of circuits, higher abstraction levels are needed [20]. EH arises as a promising solution of this problem since from a behavioral specification of a circuit, the GA will search for a bitstream to describe the circuit. Therefore the designer’s job is reduced to constructing the GA setup (specifying the circuit requirements, the basic elements of GA operations like cross and mutation percentage, etc.) and testing schemes for the fitness function [20].

Thus far, the implementations of the GA using reconfigurable hardware, including the one presented in this thesis, are focused on accelerating existing applications with existing systems and techniques. Accelerating an application in an RC system requires analysis and optimizations by the designers and these are not always easy to do due to the complexity and the size of the problem. The complexity of the design arises from the characteristics of the application that will suit it to a specific architecture and how the tools are going to facilitate the design methods. The fact that we did not achieve performance gain in the GA process is related to this complex interaction among various architectures, application characteristics, and the tools and techniques used to solve the problems. An open problem in this field is the creation of new tools and techniques for
reconfigurable hardware that will make these problems more tractable. Creating these new tools and techniques will however require multidisciplinary efforts between mathematicians, computational scientists, computer scientists, and computer engineers.
REFERENCES


