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Abstract

The most popular multithreaded languages based on the fork-join concurrency model (Cilk-Plus, OpenMP) are currently being extended to support other forms of parallelism (vectorization, pipelining and single-instruction-multiple-data (SIMD)). In the SIMD case, the objective is to execute the corresponding code on a many-core device, like a GPGPU, for which the CUDA language is a natural choice. Since the programming concepts of CilkPlus and OpenMP are very different from those of CUDA, it is desirable to automatically generate optimized CUDA-like code from CilkPlus or OpenMP.

In this thesis, we propose an accelerator model for annotated C/C++ code together with an implementation that allows the automatic generation of CUDA code. One of the key features of this CUDA code generator is that it supports the generation of CUDA kernel code where program parameters (like number of threads per block) and machine parameters (like shared memory size) are treated as unknown symbols. Hence, these parameters need not to be known at code-generation-time: machine parameters and program parameters can be respectively determined when the generated code is installed on the target machine.

In addition, we show how these parametric CUDA programs can be optimized at compile-time in the form of a case discussion, where cases depend on the values of machine parameters (e.g. hardware resource limits) and program parameters (e.g. dimension sizes of thread-blocks).

This generation of parametric CUDA kernels requires to deal with non-linear polynomial expressions during the dependence analysis and tiling phase of the input code. To achieve these algebraic calculations, we take advantage of techniques from computer algebra, in particular in the RegularChains library of Maple. Various illustrative examples are provided together with performance evaluation. Our preliminary implementation uses LLVM, Maple and PPCG; moreover, it successfully processes a variety of standard test-examples.

Keywords: Many-core machine model; Parametric CUDA code generation; Polynomial arithmetic; Compiler optimization
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Dream my life, live my dream. To a better self.
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Chapter 1

Introduction

It is well known that the impact of the high-performance software for both numerical and exact linear algebra on engineering and scientific computing is tremendous, worthy of the great efforts that have been put for more than thirty years. The most significant results are software projects like BLAS [76], LAPACK [3], ATLAS [121] and LinBox [43]. The same remark should be extended to the processing of linear transforms, in particular, Fast Fourier Transforms, where notable works are the SPIRAL [98] and FFTW [48] projects, as well as to computer algebra, see the recent proceedings of the international workshop on parallel symbolic computation (PASCO) [44, 88, 89].

The successful techniques employed in those software projects include implementation of highly efficient algorithms for basic routines, block-based algorithms for better exploiting the memory hierarchies, parallel and distributed processing, as well as automatically optimizing and tuning code on different platforms. This trend has been stimulated by the advent of hardware acceleration technologies (multicore processors, cell processors, general-purpose graphics processing units (GPGPUs), field programmable gate arrays (FPGAs)) provide vast opportunities for innovation in computing. In particular, GPGPUs combined with low-level heterogeneous programming models, such as CUDA (the Compute Unified Device Architecture, see [95, 73]), brought super-computing to the level of the desktop computer.

However, these low-level programming models carry notable challenges, even to expert programmers. Indeed, fully exploiting the power of hardware accelerators by writing CUDA code often requires significant code optimization efforts. While such efforts can yield high performance, it is desirable for many programmers to avoid the explicit management of the hardware accelerator, e.g. data transfer between the host (or CPU) and the device (or GPGPU) or between memory levels of the device. For this reason, the most popular multithreaded languages, in particular CilkPlus [10, 79] and OpenMP [41, 13, 9], are based on the fork-join concurrency model targeting multi-core architectures rather than GPGPUs. Currently, these multithreaded languages are being extended to support other forms of parallelism, such as vectorization, pipe-lining and single-instruction-multiple-data (SIMD). In the SIMD case, the case we are interested in this thesis, the objective is to execute the corresponding code on a many-core GPGPU, for which the CUDA language is a natural choice.

To overcome the challenge of developing software targeting many-core GPGPUs, high-level models for accelerator programming, notably OpenMP and OpenACC [113, 56], have become an important research direction. With these models, programmers only need to annotate
their C/C++ (or FORTRAN) code to indicate which portion of code is to be executed on the device, and how data is mapped between the host and the device.

In OpenMP and OpenACC, the work distributed among the processors of the device can be expressed in a loose manner or even ignored. This implies that code optimization techniques must be applied in order to derive efficient CUDA code. Moreover, existing software packages (e.g. PPCG [115], C-to-CUDA [6], hiCUDA [57], CUDA-CHiLL [101]) for generating CUDA code from annotated C/C++ programs, either let the user choose, or make assumptions on the characteristics of the targeted hardware and on how the work is divided among the processors of that device. These choices and assumptions limit code portability as well as opportunities for code optimization. This latter fact will be illustrated with the following example for reversing a one-dimensional array.

**Example 1** Consider the C program on Figure 1.1 for reversing a one-dimensional array, where each element of the input vector In is moved to the appropriate position of the output vector Out. Figure 1.2 gives a corresponding CUDA program, where the host code, shown on Figure 1.2(a), launches a kernel on the device, shown on Figure 1.2(b). Note that the kernel code makes a few assumptions:

1. The array length N is less than the maximum number of threads that the hardware architecture supports.
2. 32 divides N.

In order to launch a kernel function, one must specify the grid and thread-block formats (introduced in Section 2.2). In our example of Figure 1.2 32 threads per thread-block and N/32 thread-blocks per grid are specified for kernel0. Observe also that the program explicitly handles the data placement on the device, for instance, on Figure 1.2 allocating shared memory for array In and using the global memory for array Out.

```c
int N, In[N], Out[N];
// Initializing
for (int i = 0; i < N; i++)
    In[i] = i+1;

// Reversing the array In
for(int i = 0; i < N; i++)
    Out[N-1-i] = In[i];
```

Figure 1.1: A C program for reversing a one-dimensional array

The C-to-CUDA code generation introduced in [6] is known to be the first source-to-source polyhedral framework (see Section 2.5 for the term polyhedral) that translates serial C programs into CUDA programs. However, manual post-processing, such as placing synchronization statements in the kernel code, is required for generating a compilable CUDA kernel.

PPCG [115] automatically generates CUDA code from a given serial C program. In particular, Feautrier’s and PLUTO’s algorithms, respectively presented in [46] and [15], are adopted by the PPCG designers for exploring parallelization opportunities in serial programs.
int N, In[N], Out[N];

// Initializing
for (int i = 0; i < N; i++)
    In[i] = i+1;

int *dev_In, *dev_Out;

// Allocating memory spaces on the device
cudaMalloc((void **) &dev_In, (N)*sizeof(int));
cudaMalloc((void **) &dev_Out, (N)*sizeof(int));

// Copying the data from host to device
cudaMemcpy(dev_In, In, (N)*sizeof(int), cudaMemcpyHostToDevice);
cudaMemcpy(dev_Out, Out, (N)*sizeof(int), cudaMemcpyHostToDevice);

// Launching the kernel
dim3 dimBlock(32);
dim3 dimGrid(N/32);
kernel0 <<<dimGrid, dimBlock>>> (dev_In, dev_Out, N);

// Copying the data from device to host
cudaMemcpy(Out, dev_Out, (N)*sizeof(int), cudaMemcpyDeviceToHost);

// Freeing the memory spaces on the device
cudaFree(dev_In);
cudaFree(dev_Out);

(a) The host code

__global__ void kernel0(int *In, int *Out, int N) {
    int idx = blockIdx.x * 32 + threadIdx.x;
    __shared__ int shared_In[32];
    if (idx < N) {
        shared_In[threadIdx.x] = In[idx];
        __syncthreads();
        Out[N-1-idx] = shared_In[threadIdx.x];
    }
}

(b) The device code

Figure 1.2: The CUDA program for reversing a one-dimensional array

However, by default, PPCG uses 32 (resp. 16 × 32) as the thread-block format in the generated one-dimensional (resp. two-dimensional) kernels; meanwhile, PPCG also allows the user to pass a numerical value as the thread-block format.

hiCUDA (acronym for high-level CUDA) [57] is defined as a directive-based language for generating CUDA programs from C programs. To be precise, hiCUDA relies on the pragma mechanism provided by the C/C++ standards to indicate how the work is distributed among
threads and how the data is mapped on the device. However, the grid and thread-block formats of a kernel must be specified as integer expressions.

CUDA-CHiLL [101] presents a script-based compiler framework for transforming annotated C programs to CUDA programs. Moreover, in [72], a transformation strategy generator is introduced with CUDA-CHiLL, such that several candidate CUDA kernels are generated, by varying data placement (shared memory or register) and numerical values of parameters (like thread-block sizes). Then, empirical evaluation of those candidate CUDA kernels is performed so as to select the CUDA kernel with the best performance by the end.

In summary, for those CUDA kernels generated by PPCG, C-to-CUDA, hiCUDA and CUDA-CHiLL, the program parameters (e.g. the number of threads per thread-block) and the machine parameters (e.g. the shared memory size) are numerical values instead of unknown symbols.

In this thesis, we proposes an accelerator model for annotated C/C++ code, together with an implementation, that allows the automatic generation of CUDA code. One of the key features of this CUDA code generator is that it supports the generation of CUDA kernel code where program parameters (like number of threads per block) and machine parameters (like shared memory size) are treated as unknown symbols. Thus, machine parameters can be specialized to actual values when the generated CUDA code is compiled on the targeted hardware. The program parameters can be either optimized by techniques developed in this thesis or automatically tuned at run-time. As an illustration, following up on Example 1, a CUDA kernel code depending on one program parameter, along with a kernel function call from the host, is shown in Figure 1.3 where the variable B specifies the thread-block format. Observe that kernel1 takes the program parameter B as an argument, whereas the kernel0 in Figure 1.2 takes data parameters In, Out and N only.

```c
__global__ void kernel1(int *In, int *Out, int N, int B) {
    int idx = blockIdx.x * B + threadIdx.x;
    // BLOCK_0 should be pre-defined as a constant
    // and be equal to B
    __shared__ int shared_In[BLOCK_0];

    if (idx < N) {
        shared_In[threadIdx.x] = In[idx];
        __syncthreads();
        Out[N-1-idx] = shared_In[threadIdx.x];
    }
}

// The kernel function call from the host
dim3 dimBlock(B);
dim3 dimGrid(N/B);
kernell <<<dimGrid, dimBlock>>> (dev_In, dev_Out, N, B);
```

Figure 1.3: Parametric CUDA kernel for reversing a one-dimensional array

Furthermore, given machine parameters represented as unknown symbols as well, we aim at optimizing CUDA kernels, such that for each possible value of the machine and program
parameters, we provide a correspondingly optimal CUDA kernel. Of course, the meaning of optimal has to be clearly defined, which will be done in Chapter 7.

To continue with our example, consider two machine parameters for the targeted GPGPU device:

- \( Z \): the maximum number of shared memory words per processor supported by the hardware architecture,
- \( R \): the maximum number of registers per thread supported by the hardware architecture.

For the C program in Example 1, we generate two CUDA kernels based on the possible values of the machine and program parameters, see Figure 1.4.

Observe that each thread in `kernel2` moves two elements of array \( \text{In} \) to the corresponding positions of array \( \text{Out} \), whereas `kernel1` is identical to the kernel in Figure 1.3. By doing so, `kernel2` increases arithmetic intensity so as to hide the data transfer time between the global and shared memories for array \( \text{In} \); however, this increases both register usage (from 6 to 8) and shared memory usage (from \( B \) to \( 2^*B \)) machine words. Therefore, `kernel2` works correctly and is optimal under the system of the constraints \( C_2 \) but not under the system of the constraints \( C_1 \).

### 1.1 Contributions of this thesis

One of main objectives of this thesis is to generate CUDA kernels with program and machine parameters represented by unknown symbols. We call such kernels parametric. These parameters need not be known at code-generation time. Machine parameters (e.g. shared memory size) and program parameters (e.g. number of threads per thread-block) can be, respectively, determined when the generated CUDA code is installed on the targeted hardware. The challenge here is that any manipulation of non-linear expressions to be generated in the CUDA kernel requires specific code generation techniques relying on algebraic computation.

A second objective is to optimize parametric CUDA programs at compile-time, in the form of a case discussion, where cases depend on the possible values of machine parameters and program parameters. This leads us to the concept of comprehensive parametric CUDA kernels. To be more precise, given an input annotated C code, this is a decision tree, where each edge holds a Boolean expression (given by polynomial constraints) and each leaf is either a CUDA program such that for each leaf \( K \) we have:

1. \( K \) works correctly under the conjunction of the Boolean expressions located between the root node and the leaf, and
2. \( K \) is semantically equivalent to \( C \).

In each Boolean expression, the unknown variables represent machine parameters and program parameters. This case discussion can be handled by techniques from symbolic computation. Automatic parametric kernel code generation can, then, be achieved by means of combining an optimizing compiler and a computer algebra system.

A third objective of this thesis is to measure the performance of parametric algorithms or programs targeting many-core devices like GPGPUs. To this end, a model of multithreaded computation targeting many-core architectures is introduced, such that one can either tune a program parameter to determine a value range minimizing parallelism overheads, or compare different multi-threaded algorithms solving the same problem.
Figure 1.4: Two CUDA kernels based on possible values of machine and program parameters for reversing a one-dimensional array, where \( Z \) is the maximum number of shared memory words per processor supported by the hardware architecture and \( R \) is the maximum number of registers per thread supported by the hardware architecture.

We observe that classical models of parallel computation, namely the fork-join concurrency model [11] and the parallel random access machine (PRAM) model [110, 51], do not distinguish between the task-based and data-based parallelism. Thus, those models are too simplistic for analyzing algorithms targeting GPGPUs.

1.2 Outline of this thesis

In Chapter 2, various topics related to our work are reviewed. We first discuss briefly classical models of computation for concurrency platforms. In particular, we review the CUDA programming model and some important features of modern GPU architectures [36, 37]. We also
1.2. Outline of this thesis

review the MetaFork language [29], which is a linguistic extension of C/C++ with high-level parallel programming constructs. Similarly to OpenMP and OpenACC, the MetaFork language offers a high-level model for accelerator programming. We stress the fact that this thesis does not deal with dependence analysis and the computation of schedules in automatic parallelization; this can be done via the polyhedral model [46][15] that we briefly review in Chapter 2. Finally, in that same chapter, we give an overview of the algorithmic tools [4][25][23][24][112][22] for dealing with systems of non-linear polynomial equations and inequalities.

In Chapter 3 we present the Basic Polynomial Algebra Subprograms (BPAS) library for arithmetic operations with univariate and multivariate polynomials, dense or sparse. The BPAS library is written in CilkPlus targeting multi-core architectures. The original goal of this thesis was to investigate how to integrate code targeting GPGPUs (for instance, from the CUMODP library) within the BPAS library. These two libraries, BPAS and CUMODP, are developed in our research group; moreover, this leads us to the idea of developing frameworks, such as MetaFork, for translating programs between different concurrency platforms. We also note that, the BPAS library, as a computer algebra library targeting high-performance, could also be used, in the future, for improving the efficiency of our algebraic tools for generating parametric CUDA kernels. The work reported in Chapter 3 is an extended version of [20] as well as a joint project with Changbo Chen, Svyatoslav Covano, Farnam Mansouri, Robert Moir, Marc Moreno Maza and Yuzhen Xie. Appendix A lists sample codes from the BPAS library.

Chapter 4 presents a model of multithreaded computation with an emphasis on estimating the parallelism overheads of programs targeting many-core machines. To evaluate the performance of a given CUDA-like program, we consider three complexity measures, namely work, span and parallelism overhead. We also extend the Graham-Brent theorem so as to estimate the running time of a CUDA-like program on a given number of streaming multiprocessors. We evaluate the benefits of our model on six fundamental algorithms, including the Euclidean algorithm for univariate polynomial GCDs, two fast Fourier transform algorithms, the plain and FFT-based univariate polynomial multiplication algorithms, and radix sort [103]. Each studied algorithm is either implemented in the CUMODP library or reported in [103]. We observe that experimentation is coherent with the theoretical analysis based on our model. The work in Chapter 4 is an extended version of [61] as well as a joint project with Sardar Anisul Haque and Marc Moreno Maza.

Chapter 5 reports on a preliminary implementation of the C-to-CUDA code generator discussed above. Generating parametric CUDA kernels implies dealing with non-linear polynomial expressions, particularly during the tiling phase. To achieve these algebraic calculations, we take advantage of quantifier elimination (QE) and its implementation of the RegularChains library of MAPLE [27]. In order to illustrate the merits of parametric CUDA kernels, we use our code generator on eight test cases: array reversal, 1D Jacobi, 2D Jacobi, LU decomposition, matrix transposition, matrix addition, matrix vector multiplication and matrix matrix multiplication. The performance evaluation of the generated CUDA programs of each test case is provided as well. Chapter 5 is related to [19] and is joint work with Changbo Chen, Xiaohui Chen and Marc Moreno Maza. Appendix C provides documentation of our MetaFork-to-CUDA code generator, while Appendix D collects the generated CUDA programs by PPCG for those same eight test cases.

In Chapter 6 based on the experimental results conducted in Chapter 5, we study advanced optimization techniques, such as the controlling granularity of threads and loop unrolling, We
use six test cases so as to verify whether these techniques can, in general, further improve the performance of parametric CUDA kernels. For each test case, since we use the MetaFork language as the high-level accelerator programming model, the portion of the code that should be translated into CUDA kernels is annotated manually. Along with the previous chapter, experimentation shows that the generation of parametric CUDA kernels can lead to significant performance improvement w.r.t. approaches based on the generation of CUDA kernels that are not parametric. Moreover, for certain test cases, our experimental results show that the optimal choices for program parameters may depend on the problem size.

In Chapter 7, we propose an algorithm for comprehensive optimization of an annotated C program, depending on parameters treated as symbols at compile-time. We use this algorithm to generate optimized parametric CUDA kernels in the form of a case discussion based on the possible values of the machine and program parameters. In our preliminary implementation of the comprehensive optimization algorithm, we consider two machine parameters: register usage per thread and required shared memory per thread-block; meanwhile, we apply four code optimization strategies: caching data in local or shared memory, reducing register usage per thread, controlling thread granularity and eliminating common sub-expressions. This is a proof-of-concept implementation written in Maple and dedicated to the comprehensive generation of optimized MetaFork programs from an input MetaFork program. For each of the six test cases: array reversal, matrix vector multiplication, 1D Jacobi, matrix addition, matrix transpose and matrix matrix multiplication, three optimized MetaFork programs are generated with systems of constraints. Chapter 7 is joint work with Xiaohui Chen and Marc Moreno Maza. Appendix E provides the pseudo-codes of the implemented algorithms.

Figure 1.5: Overview of the thesis

To summarize, Figure 1.5 gives an overview of this thesis. The BPAS library is freely

In the future, we plan to apply our MetaFork-to-CUDA code generator to efficiency-critical routines of the BPAS library code. Meanwhile, we plan to improve the run-time performance of our MetaFork-to-CUDA code generator by using BPAS instead of MAPLE for dealing with the non-linear expressions arising in the generation of parametric CUDA code.
Chapter 2

Background

In this chapter, we review background materials related to our work. Section 2.1 is dedicated to classical models of parallel computation. Section 2.2 discusses briefly the features of modern GPU architectures and the programming model CUDA. Section 2.3 gives an overview of the CUMODP library, which is used in the experimentation reported in Chapter 4. In Section 2.4, we present the MetaFork language, which serves as a high-level parallel programming model for automatic CUDA code generation in Chapters 5, 6 and 7. In Section 2.5, we study Feautrier’s and PLUTO’s algorithms in the so-called polyhedral model. Finally, Section 2.6 is an overview of the algorithmic tools used for dealing with systems of polynomial equations and inequalities.

2.1 Models of computation

Based on Flynn’s taxonomy \cite{ Flynn46}, single instruction multiple data (SIMD) is defined as one of the categories of parallel computers, where multiple processing units execute the same instructions on multiple data sets. In this scheme, data is distributed across different processing units so as to achieve data parallelism. In contrast, task parallelism distributes tasks (or functions) to different processing units.

As mentioned before, with the pervasive ubiquity of many-core processors, in particular GPUs, models of computation must take into account both task-based and data-based parallelism. In fact, popular concurrency platforms (such as CilkPlus \cite{CilkPlus, CilkPlus2}, CUDA \cite{CUDA, CUDA2} and OpenCL \cite{OpenCL}) offer both forms of parallelism, with parallel constructs specific to each case.

Hereafter, we review two classical models of parallel computation, the fork-join model in Section 2.1.1 and the parallel RAM (PRAM) model in Section 2.1.2. Those models do not distinguish between task-based parallelism and data-based parallelism; thus, those models are too simplistic for analyzing algorithms targeting many-cores.

2.1.1 Fork-join model

In the fork-join model \cite{ForkJoin}, one can consider a multithreaded program (see Chapter 27 in \cite{Skiena}) as a directed acyclic graph \( G = (V, E) \), called computation DAG. Figure 2.1 shows the computation DAG for calculating the 5-th Fibonacci number, namely \( F_4 \), where \( F_n = F_{n-1} + F_{n-2} \) if
2.1. Models of computation

$n \geq 2$ and $F_0 = F_1 = 1$ otherwise. In a multithreaded program, we call a strand a sequence of consecutive instructions without parallel constructs.

Figure 2.1: An example of computation DAG: Fourth Fibonacci

This model defines two complexity measures:

1. the work $T_1$ of a multithreaded computation is the total time to execute the entire computation on one processor, and

2. the span $T_\infty$ is the longest time to execute the strands along any path in the DAG.

Assuming that each strand executes in unit time, the work is the number of vertices, and the span equals the number of vertices on a longest path or critical path in the computation DAG.

Those two measures, work and span, and one machine parameter, the number $P$ of processors, can be combined in results like the Graham-Brent theorem ([11, 53]) or the Blumofe-Leiserson theorem (Theorems 13 & 14 in [12]) in order to give running time estimates. We recall that the Graham-Brent theorem states that the running time $T_P$ on $P$ processors satisfies $T_P \leq T_1/P + T_\infty$. A refinement of this latter theorem actually supports the implementation (on multi-core architectures) of the parallel performance analyzer, called Cilkview [63]. In this context, the running time $T_P$ is bounded in expectation by $T_1/P + 2\hat{T}_\infty$, where $\delta$ is a constant (called the span coefficient) and $\hat{T}_\infty$ is the so-called burdened span.

The fork-join model has become popular with the development of the concurrency platform CilkPlus [10, 79] targeting multi-core architectures. An example of code written in CilkPlus is given in Figure 2.2 for computing matrix multiplication. This CilkPlus code is implemented with a divide & conquer method and the blocking strategy reported in [49] by Frigo, Leiserson, Prokop and Ramachandran.

2.1.2 PRAM model

The PRAM model [110, 51] is defined as a synchronous model of parallel computation. The PRAM machine consists of a number of processors, each of which is a RAM with a private local memory, and a shared memory that processors communicate with. Since the amount of shared memory is limited, it restricts the amount of data that can be communicated between processors in one step.

Moreover, the PRAM model defines four types of accesses to the same shared memory cell with respect to read and write operations. These four types are exclusive read exclusive write
void parallel_dandc(int i0, int i1, int j0, int j1, int k0, int k1, int* A, int* B, int* C, int N, int X) {
    int di = i1 - i0;
    int dj = j1 - j0;
    int dk = k1 - k0;
    if (di >= dj && di >= dk && di >= X) {
        int mi = i0 + di / 2;
        cilk_spawn parallel_dandc(i0, mi, j0, j1, k0, k1, A, B, C, N, X);
        parallel_dandc(mi, i1, j0, j1, k0, k1, A, B, C, N, X);
        cilk_sync;
    }
    else if (dj >= dk && dj >= X) {
        int mj = j0 + dj / 2;
        cilk_spawn parallel_dandc(i0, i1, j0, mj, k0, k1, A, B, C, N, X);
        parallel_dandc(i0, i1, mj, j1, k0, k1, A, B, C, N, X);
        cilk_sync;
    }
    else if (dk >= X) {
        int mk = k0 + dk / 2;
        parallel_dandc(i0, i1, j0, j1, k0, mk, A, B, C, N, X);
        parallel_dandc(i0, i1, j0, j1, mk, k1, A, B, C, N, X);
    }
    else {
        // The base case using the serial, naive matrix multiplication
        for (int i = i0; i < i1; ++i)
            for (int j = j0; j < j1; ++j)
                for (int k = k0; k < k1; ++k)
                    C[i * N + j] += A[i * N + k] * B[k * N + j];
    }
}

Figure 2.2: Matrix multiplication written in CilkPlus

(EREW), concurrent read exclusive write (CREW), exclusive read concurrent write (ERCW) and concurrent read concurrent write (CRCW). At one step, “exclusive” indicates that at most one processor can read from or write to the same memory cell, while “concurrent” indicates that processors can simultaneously read from or write to the same memory cell. The advantage of the PRAM model is that it focuses on designing efficient parallel algorithms in terms of arithmetic operations while ignoring communication issues. In addition, all memory accesses are assumed to take place in constant time. Unfortunately, concrete machines cannot currently scale to large numbers of processors while preserving uniformly fast access time to the shared memory.

An attempt to integrate memory contention into the PRAM model has been made with the queue read queue write (QRQW) PRAM, defined in [52]. This model enhances the Graham-Brent theorem with memory access time. However, both time spent in arithmetic operations
and time spent in read/write accesses are conflated in a single quantity. We believe that this unification is not appropriate for recent many-core processors, such as NVIDIA GPUs, for which the ratio between one global memory read/write access and one floating point operation can be in the 100’s.

A more practical PRAM model is proposed in [51] called asynchronous PRAM. Unlike the PRAM model, the processors of an asynchronous PRAM run asynchronously, that is, each processor executing its instructions independently without interrupting others. It considers subset independent synchronization and fixed communication delays for global reads/writes. However, this family of PRAM is not suitable to modern GPUs, since it is trivial to consider a machine parameter for synchronization, which is not an issue for modern GPUs.

2.2 General-purpose GPU computing

![Overview of a hybrid CPU-GPU system](image)

Figure 2.3: Overview of a hybrid CPU-GPU system

General-purpose GPU computing has evolved from graphics processors to massively parallel many-core multiprocessors with the adoption of co-processing between the CPU and the GPU. In such a hybrid CPU-GPU system, GPUs can be used to accelerate a portion of sequential code by employing task-parallelism and data-parallelism. Figure 2.3 shows an overview of a typical CPU-GPU system architecture. One of the main differences between the CPU and the GPU is the number of cores that are designed for different types of execution. Indeed, a GPU device consists of several streaming multiprocessors (SMs); meanwhile, each SM consists of a large number of processing cores. For this reason, these cores are particularly designed to take advantage of parallel execution in the SIMD (single instruction multiple data) manner. While memory coherence is maintained by the CPU, the programmer needs to explicitly deal with data management on the GPU. Section 2.2.1 introduces the programming model, the Compute Unified Device Architecture (CUDA). In Section 2.2.2, we summarize the characteristics of modern GPU architectures.
2.2.1 The Compute Unified Device Architecture (CUDA)

CUDA is the programming model introduced to implement parallel algorithms on NVIDIA GPU devices \[95, 73\]. A CUDA program consists of one or more blocks of code that are executed on either the host (CPU) or the device (GPU). On the host, the code is implemented with little or no parallelism, while on the device, the code is designed to exhibit a rich amount of task or data parallelism, in particular, in a SIMD fashion.

![Figure 2.4: Execution of a CUDA program](image)

The execution of a typical CUDA program is shown in Figure 2.4. It starts with the serial C/C++ code and then launches a kernel on the device to execute the parallel code. Upon the completion of the kernel, the process resumes to the execution of serial code on the host. When a kernel function is invoked, the number of thread-blocks and the number of threads per thread-block are specified, according to the problem size, by the programmer. The syntax for launching a kernel from the host code extends that of a C function call with kernel execution configuration parameters surrounded by \[\lll\text{and} \rr\rr\], where the execution configuration parameters define the dimensions of the grid and the dimensions of each thread-block. All the threads comprising a kernel during an invocation are collectively called a \emph{grid}.

In addition, the programmer needs to allocate the problem size on the device memory using the keyword \texttt{cudaAlloc} and free the device memory after computation using the keyword \texttt{cudaFree}. To transfer pertinent data between the host and allocated device memories, one can use the keyword \texttt{cudaMemcpy}, with \texttt{cudaMemcpyHostToDevice} and \texttt{cudaMemcpyDeviceToHost} specifying where the data is copied from and to. The function type qualifier \texttt{global1} written prior to a function declaration, indicates that the function is a kernel; meanwhile, on the host, this kernel is called to generate a grid of threads on the device. Moreover, \texttt{threadIdx},

\footnote{See \url{http://docs.nvidia.com/cuda/cuda-c-programming-guide/index.html} for CUDA C programming guide.}
blockIdx and blockDim are built-in variables to identify the thread index, the thread-block index and the thread-block dimension, respectively.

2.2.2 Modern GPU architectures

Unlike CPUs optimized for low latency access to memory systems, modern GPUs are designed for data-parallel, throughput computation and tolerance of memory latency. A GPU typically consists of a global memory (dynamic random-access memory (DRAM)), a number of streaming multiprocessors (SMs) and a global scheduler. Each SM contains a number of cores, a shared memory and a thread scheduler.

Apart from the evolution of graphic techniques on GPUs, we summarize key features of modern GPU architectures with respect to general-purpose GPU computing as below.

1. **Hardware thread scheduling.** Based on the documentation of NVIDIA Fermi [36] and Kepler [37] architectures, there is a two-level, distributed thread scheduler, such that thread-blocks from the same or different kernels are scheduled to various SMs and warps of 32 threads of a thread-block are distributed to execute concurrently.

2. **Memory hierarchy.** This hierarchy offers the benefits of the on-chip local memory for each thread, the on-chip shared memory shared among threads in an SM with low latency, and the global memory for sharing across the GPU with high throughput. In order to declare an array allocated in the shared memory of the SM, one shall use the _shared_ qualifier prior to the declaration of the array (see Figure 1.3 for an example). Moreover, when a warp of threads accesses consecutive memory locations to the global memory, the hardware coalesces its memory accesses into a consolidated access.

3. **Data parallelism.** This is implemented as SIMD processors or in a single instruction multiple thread (SIMT) fashion.

4. **Task parallelism.** This type of parallelism, also called dynamic parallelism, can launch a new kernel from a thread, synchronize on the results and control the scheduling via the hardware paths; moreover, these operations are done independently of the CPU. This allows data-dependent and recursive code to execute in parallel on GPUs.

5. **Latency hidden.** This is done via computation by SIMT (instead of cache). Based on the concept of fast context switching, active warps of threads that wait for memory accesses are switched to those whose data are available. Typically, 128 threads per SM occupy an SM during the computation to hide the global memory access latency.

6. **Synchronization.** Threads are independent of each other, such that there are no synchronization issues among thread-blocks per grid, while threads with a thread-block can be synchronized by issuing _syncthreads()._ Additionally, synchronization of task parallelism is explicitly managed prior to launching a (child) kernel to ensure that all data is ready. If thread divergence occurs, both sides of the branch execute within a warp, and idle threads in a warp wait for others to complete.

To be consistent with the programming framework, a thread-block is mapped to one SM, such that threads within a thread-block communicate via the shared memory with low latency and low throughput. Thread-blocks mapped to different SMs communicate via the global memory with high latency and high throughput.
2.3 Dense arithmetic over finite fields with the CUMODP library

CUMODP [59] is a CUDA library for exact computations with dense polynomials over finite fields. A variety of operations, like multiplication, division, computation of subresultants, multi-point evaluation, interpolation and many others, are provided. These routines are primarily designed to offer GPU support to polynomial system solvers and a bivariate system solver is part of the library. Algorithms combine FFT-based and plain arithmetic, while the implementation strategy emphasizes reducing parallelism overheads and optimizing hardware usage.

Polynomial multiplication and matrix multiplication are at the core of many algorithms in symbolic computation. Expressing, in terms of multiplication time, the algebraic complexity of an operation like univariate polynomial division or the computation of a characteristic polynomial is a standard practice, see for instance the landmark book [120]. At the software level, the motto “reducing everything to multiplication”[^1] is also common, see for instance the computer algebra systems Magma [16], NTL [108] or FLINT [62].

With the advent of hardware accelerator technologies, multi-core processors and Graphics Processing Units (GPUs), this reduction to multiplication is, of course, still desirable, but becomes more complex since both algebraic complexity and parallelism need to be considered when selecting and implementing a multiplication algorithm. In fact, other performance factors, such as cache usage or CPU pipeline optimization, should be taken into account on modern computers, even on single-core processors. These observations guide the developers of projects like SPIRAL [98] or FFTW [48].

The CUMODP library provides arithmetic operations for matrices and polynomials (in a dense representation) primarily with modular integer coefficients, targeting many-core GPUs. Some operations are available for integer or floating point coefficients as well. A large portion of the CUMODP library code is devoted to polynomial multiplication and the integration of that operation into higher-level algorithms.

Typical CUMODP operations are matrix determinant computation, polynomial multiplication (both plain and FFT-based), univariate polynomial division, the Euclidean algorithm for univariate polynomial GCDs, subproduct tree techniques for multi-point evaluation and interpolation, subresultant chain computation for multivariate polynomials and bivariate system solving. The CUMODP library is written in CUDA [95, 73] and its source code is publicly available at [www.cumodp.org](http://www.cumodp.org).

This work is reported in [59] and is a joint project with Sardar Anisul Haque, Xin Li, Farnam Mansouri, Marc Moreno Maza and Wei Pan.

[^1]: Quoting a talk title by Allan Steel, from the Magma Project.
2.4 The MetaFork Language

MetaFork [29] is a high-level programming language extending C/C++, which combines several models of concurrency, including fork-join and pipelining parallelisms. MetaFork is also a compilation framework, which aims at facilitating the design and implementation of concurrent programs through three key features:

1. Perform automatic code translation between concurrency platforms targeting both multi-core and many-core GPU architectures.
2. Provide a high-level language for expressing concurrency as in the fork-join model, the SIMD (single instruction multiple data) paradigm and the pipelining parallelism.
3. Generate parallel code from serial code with an emphasis on code depending on machine or program parameters (e.g. cache size, number of processors, number of threads per thread-block).

As of today, the publicly available and latest release of MetaFork, see [www.metafork.org](http://www.metafork.org), offers the second feature stated above, a preliminary implementation of the third feature as well as the multi-core and many-core portions of the first one. To be more specific, MetaFork is a meta-language for concurrency platforms based on the fork-join model, pipelining parallelism and the SIMD paradigm. This meta-language forms a bridge between actual multithreaded programming languages, and we use it to perform automatic code translation between those languages.

In an earlier work [29], MetaFork was introduced as an extension of both the C and C++ languages into a multithreaded language based on the fork-join concurrency model [11]. Thus, concurrent execution is obtained by a parent thread creating and launching one or more children threads, so that the parent and its children execute a so-called parallel region. An important example of parallel regions are for-loop bodies. MetaFork has four parallel constructs dedicated to the fork-join model: function call spawn, block spawn, parallel for-loop and synchronization barrier. The first two use the keyword metafork, while the other two use, respectively, the keywords metafor and metajoin. Similar to the CilkPlus specifications, the parallel constructs of MetaFork grant permission for concurrent execution but do not command it. Hence, a MetaFork program can execute on a single core machine. We emphasize the fact that metafork allows the programmer to spawn a function call (like in CilkPlus [10, 79, 35]) as well as a block (like in OpenMP [41, 13, 9]). Using the same examples from [29], Figures 2.5, 2.6 and 2.7 illustrate automatic code translation between the OpenMP program and the CilkPlus program via the MetaFork language.

On the other hand, stencil computations are a major pattern in scientific computing. Stencil codes perform a sequence of sweeps (called time-steps) through a given array, and each sweep can be seen as the execution of a pipeline. When expressed with concurrency platforms based on, and limited by, the fork-join model, parallel stencil computations incur excessive
parallelism overheads. This problem is studied by Shirako, Unnikrishnan, Chatterjee, Li and Sarkar [107] together with a solution in the context of OpenMP by proposing new synchronization constructs to enable *do-across parallelism*. These observations have motivated a first extension of the MetaFork language with three constructs to express pipelining parallelism: *meta_pipe*, *meta_wait* and *meta_continue*. Recall that a pipeline is a linear sequence of processing stages through which data items flow from the first stage to the last stage. If each stage can process only one data item at a time, then the pipeline is said to be *serial* and can be depicted by a (directed) path in the sense of graph theory. If a stage can process more than one data item at a time, then the pipeline is said to be *parallel* and can be depicted by a directed acyclic graph (DAG), where each parallel stage is represented by an independent set, that is, a set of vertices of which no pair is adjacent.

In order to generate efficient CUDA code from an input MetaFork program, we introduced a tenth keyword, namely *meta_schedule*, in [19]. This keyword allows its body to be scheduled on a device, such as the NVIDIA GPU, to execute in a SIMD fashion. In Chapter 5 as well as Appendix C, we depict the MetaFork-to-CUDA code generator, which is capable of
2.5. Automatic parallelization in the polyhedral model

Automatic parallelization deals mainly with the transformation of for-loop nests so as to expose parallelism, that is, being able to execute one or more for-loops in a parallel fashion. By executing a for-loop in a parallel fashion, we mean that all iterations of that loop can be executed concurrently without changing the semantics of that loop. For the sake of simplicity, we shall make the following assumption in the sequel of this section.

Automatically generating compilable CUDA programs with program parameters (like number of threads per thread-block) and machine parameters (like shared memory size) allowed at code-generation-time.

This work is summarized in [19].

**Figure 2.7:** Using MetaFork to translate a given OpenMP program into a CilkPlus program

```
int main() {
    int sum_a = 0, sum_b = 0;
    int a[ 5 ] = {0,1,2,3,4};
    int b[ 5 ] = {0,1,2,3,4};
    #pragma omp parallel
    {
        #pragma omp sections
        {
            #pragma omp section
            {
                for(int i=0; i<5; i++)
                    sum_a += a[ i ];
            }
            #pragma omp section
            {
                for(int i=0; i<5; i++)
                    sum_b += b[ i ];
            }
        }
    }
    int sum_a = 0, sum_b = 0;
    int a[ 5 ] = {0,1,2,3,4};
    int b[ 5 ] = {0,1,2,3,4};
    meta_fork shared(sum_a) {
        for(int i=0; i<5; i++)
            sum_a += a[ i ];
    }
    meta_fork shared(sum_b) {
        for(int i=0; i<5; i++)
            sum_b += b[ i ];
    }
    meta_join;
}
```

(a) A given OpenMP program

```
void fork_func0(int* sum_a,int* a) {
    for(int i=0; i<5; i++)
        (*sum_a) += a[ i ];
}
void fork_func1(int* sum_b,int* b) {
    for(int i=0; i<5; i++)
        (*sum_b) += b[ i ];
}
```

(b) The intermediate MetaFork program

```
int main() {
    int sum_a = 0, sum_b = 0;
    int a[ 5 ] = {0,1,2,3,4};
    int b[ 5 ] = {0,1,2,3,4};
    cilk_spawn fork_func0(&sum_a,a);
    cilk_spawn fork_func1(&sum_b,b);
    cilk_sync;
}
```

(c) The translated CilkPlus program

Figure 2.7: Using MetaFork to translate a given OpenMP program into a CilkPlus program
Hypothesis 1 We consider for-loop nests having the format shown on Figure 2.8. Note that

\[
\text{for (i}_1 = 0; i_1 < N_1; i_1++) \\
\text{.} \\
\text{.} \\
\text{for (i}_e = 0; i_e < N_e; i_e++) \
\{
// for-loop nest body
\}
\]

Figure 2.8: For-loop nest in the polyhedral model

Figure 2.8 suggests that the for-loop nest is perfect, that is, only the innermost loop may have statements other than a for-loop.

In fact, imperfect for-loop nests can be reduced to perfect for-loop nests [71] without change of semantics but with possible loss of efficiency in terms of consumption of computer resources (time and space).

Paul Feautrier’s [46] and PLUTO’s [15] algorithms are two procedures for generating parallel code from serial C code automatically. The two procedures focus on parallelizing for-loop nests (including imperfect ones) under the following assumption.

Hypothesis 2 Referring to the notations introduced in Figure 2.8 in every array reference (e.g. $i_1 * N_1 + i_2$ in $a[i_1 * N_1 + i_2]$) in the for-loop nest body,

(i) the index is an arithmetic expression linear in the vector $(i_1, \ldots, i_e, N_1, \ldots, N_e)$, where

(ii) each of $N_1, \ldots, N_e$ is an arithmetic expression linear in all the variables occurring in that expression.

Note that $i_1 * N_1 + i_2$ is linear in each of the variables $i_1$, $N_1$, $i_2$, but not in the vector $(i_1, i_2, N_1)$.

Hypothesis 2 implies that the iterations of the for-loop nest can be represented by the integer points of a polyhedron. For that reason, Feautrier’s and PLUTO’s algorithms are said to be based on the polyhedral model. Before giving a sketch of those algorithms in Sections 2.5.7 and 2.5.8 we review the concepts of a $\mathbb{Z}$-polyhedron in Section 2.5.1, a polyhedral iteration domain in Section 2.5.2, a data dependence graph in Section 2.5.3, the dependence polyhedron in Section 2.5.4, an affine transformation in Section 2.5.5 and Farkas multipliers in Section 2.5.6.

2.5.1 $\mathbb{Z}$-polyhedron

Let $\mathbb{Z}$ be the ring of the integer numbers and $m$ be a positive integer. The set of all vectors $\vec{x} \in \mathbb{Z}^m$ where $h \vec{x} = k$ with $h \in \mathbb{Z}^m$ and $k \in \mathbb{Z}$ is an affine $\mathbb{Z}$-hyperplane. The set of all vectors $\vec{x} \in \mathbb{Z}^m$ where $A \vec{x} + \vec{c} \geq 0$ with an $p \times m$ integer matrix $A$ (for $p > 0$) and $\vec{c} \in \mathbb{Z}^m$ defines a $\mathbb{Z}$-polyhedron. A $\mathbb{Z}$-polytope is a bounded $\mathbb{Z}$-polyhedron.
2.5.2 Polyhedral iteration domain

We observe that every for-loop nest in the format shown on Figure 2.8 naturally defines a \( \mathbb{Z} \)-polytope \( D \) in \( \mathbb{Z}^e \), called iteration domain of the for-loop nest; moreover, the points of \( D \) are called iteration vectors. Let \( \vec{I} = (I_1, \ldots, I_e) \) and \( \vec{J} = (J_1, \ldots, J_e) \) be two iteration vectors in \( D \). We write \( \vec{J} \leq_{\text{lex}} \vec{I} \) whenever \( \vec{I} \) is lexicographically greater than \( \vec{J} \). We write \( \vec{J} \preceq_{\text{lex}} \vec{I} \) if either \( \vec{J} \prec_{\text{lex}} \vec{I} \) or \( \vec{J} = \vec{I} \) holds. We call defining system of \( D \) the system of linear inequalities:

\[
\begin{align*}
0 &\leq i_1 < N_1 \\
& \vdots \\
0 &\leq i_e < N_e
\end{align*}
\]

Let \( \vec{I} = (I_1, \ldots, I_e) \) and \( S \) be respectively an iteration vector of \( D \) and a statement of the body of our perfect loop nest. We denote by \( S(\vec{I}) \) the execution of \( S \) during the \( \vec{I} \)-th iteration of the for-loop nest.

2.5.3 Data dependence graph

The following definition is restricted to our context of a perfect for-loop nest, while it could be stated for an arbitrary program. Let \( \vec{I} \) and \( \vec{J} \) be two iteration vectors in \( D \). Let \( S_s \) and \( S_t \) be two statements of the for-loop nest body. We say that there is a data dependence from \( S_s(\vec{I}) \) to \( S_t(\vec{J}) \) (alternatively, we say that \( S_t(\vec{J}) \) depends on \( S_s(\vec{I}) \)), and we write \( S_s(\vec{I}) \Rightarrow S_t(\vec{J}) \), whenever the following conditions hold simultaneously:

1. both \( S_s(\vec{I}) \) and \( S_t(\vec{J}) \) access the same memory location and at least one of them stores data into this memory location, and
2. we have \( \vec{I} \prec_{\text{lex}} \vec{J} \).

The data dependence graph (DDG) \( G = (V, E) \) is a labelled directed-graph, where

1. the set \( V \) of the vertices is the set of the statements of the for-loop nest body,
2. \( (S_s, S_t) \) is an edge if there exist two iteration vectors \( \vec{I} \) and \( \vec{J} \) such that we have \( S_s(\vec{I}) \Rightarrow S_t(\vec{J}) \),
3. each edge \( (S_s, S_t) \) is labelled with the set of the pairs \( (\vec{I}, \vec{J}) \) of iteration vectors such that we have \( S_s(\vec{I}) \Rightarrow S_t(\vec{J}) \).

We classify the data dependence from statement \( S_s(\vec{I}) \) to statement \( S_t(\vec{J}) \) into three categories, based on the sequences of read and write operations to the same memory location \( X \):

- **True dependence** (also known as a flow dependence): \( X \) is written in \( S_s \) before it is read in \( S_t \).

\[
\begin{align*}
S_s : & \quad X = \ldots \\
S_t : & \quad \ldots = X
\end{align*}
\]

- **Anti-dependence**: \( X \) is read in \( S_s \) before it is written in \( S_t \).

\[
\begin{align*}
S_s : & \quad \ldots = X \\
S_t : & \quad X = \ldots
\end{align*}
\]

- **Output dependence**: \( X \) is written in \( S_s \) before it is written in \( S_t \).

\[
\begin{align*}
S_s : & \quad X = \ldots \\
S_t : & \quad X = \ldots
\end{align*}
\]


Example 2 Consider the source program given below:

```c
for (int i = 0; i < N; ++i)
    for (int j = 1; j < N; ++j)
```

Figure 2.9 shows the data dependence graph of the source program in Example 2. Observe that data dependence occurs for each of the following:

- from \(A[i][j-1]\) in statement \(S\) to \(A[i'][j']\) in statement \(S\) when \(i' = i, j' = j\)
- \((\text{flow dependence})\)
  - from \(A[j][i]\) in statement \(S\) to \(A[i'][j']\) in statement \(S\) when \(i' = j, j' = i, i < j\)
  - \((\text{flow dependence})\)
  - from \(A[i][j]\) in statement \(S\) to \(A[j'][i']\) in statement \(S\) when \(j' = i, i' = j, j < i\)
  - \((\text{anti-dependence})\)

where \((i, j)\) and \((i', j')\) are two iteration vectors of the for-loop nest; thus, \(i, j, i', j'\) are non-negative integers satisfying \(i < N, 0 < j, j < N, i' < N, 0 < j', j' < N\).

![Data Dependence Graph](image)

Figure 2.9: An example of the data dependence graph of the source program

2.5.4 Dependence polyhedron

Let \(e := (S_s, S_t)\) be an edge of the data dependence graph. Let \(\vec{I}\) and \(\vec{J}\) be two iteration vectors such that we have \(S_s(\vec{I}) \implies S_t(\vec{J})\). We call dependence polyhedron associated with \(e\) the \(\mathbb{Z}\)-polyhedron defined by the conjunction of the inequalities on \(\vec{I}, \vec{J}\) expressing the fact that the following conditions hold simultaneously:

1. \(\vec{I} \in D\)
2. \(\vec{J} \in D\)
3. \(S_s(\vec{I}) \implies S_t(\vec{J})\)
2.5. Automatic parallelization in the polyhedral model

In Example 2 for the flow dependence from \(A[i][j-1]\) to \(A[i'][j']\) in Figure 2.9 we observe that the dependence polyhedron is

\[
\begin{bmatrix}
1 & 0 & 0 & 0 & 0 & 0 \\
-1 & 0 & 0 & 0 & 1 & -1 \\
0 & 1 & 0 & 0 & 0 & -1 \\
0 & -1 & 0 & 0 & 1 & -1 \\
0 & 0 & 1 & 0 & 0 & 0 \\
0 & 0 & -1 & 0 & 1 & -1 \\
0 & 0 & 0 & 1 & 0 & -1 \\
0 & 0 & 0 & -1 & 1 & -1 \\
1 & 0 & -1 & 0 & 0 & 0 \\
0 & 1 & 0 & -1 & 0 & -1 \\
\end{bmatrix}
\begin{bmatrix}
i \\
j \\
i' \\
j' \\
N \\
1 \\
\end{bmatrix} \geq 0
\]  
\(= 0\)  \hspace{1cm} (2.1)

2.5.5 Affine transformation

Consider a for-loop nest and a statement \(S\) in the body of this loop nest. Let \(x_1, \ldots, x_{m_s}\) be the loop iteration counters occurring in the indices of the array references in the statement \(S\); we write \(\vec{x} = [x_1 \cdots x_{m_s}]\). By definition, a one-dimensional affine transformation for the statement \(S\) maps \(\vec{x}\) to \(\phi_S(\vec{x})\) defined by

\[
\phi_S(\vec{x}) = [t_1 \cdots t_{m_s}] (\vec{x}) + t_0,
\]

where \(t_0 \in \mathbb{Z}\) and \([t_1 \cdots t_{m_s}]\) is a row-vector over \(\mathbb{Z}\). In the literature, the map \(\phi_S\) is sometimes referred as an affine hyperplane. An \(n\)-dimensional affine transformation for statement \(S\) is given by

\[
\phi_S(\vec{x}) = T (\vec{x}) + \vec{t},
\]

where \(T\) is a constant matrix with \(n\) rows (and \(m_s\) columns) and \(\vec{t}\) is a constant vector of size \(n\), the coefficients of both \(T\) and \(\vec{t}\) belonging to \(\mathbb{Z}\). In the literature, the map \(\phi_S\) is also called a scattering function.

In the polyhedral model, the notions of one-dimensional affine transformations and \(n\)-dimensional affine transformation are used to perform an affine change of coordinate system for the loop iteration counters so as to exhibit for-loop nests whose iterations can be executed concurrently. Note that, because of the inequality constraints that appear in the polyhedron, this cannot simply be done via linear algebra and requires polyhedral operations, like projection, which can be done by the Fourier-Motzkin elimination algorithm\footnote{https://en.wikipedia.org/wiki/Fourier-Motzkin_elimination}.

To be more precise, one-dimensional affine transformations and \(n\)-dimensional affine transformations are meant to be used as time coordinates and any other other coordinates are regarded as space coordinates. A tuple of space coordinates can be interpreted as a processor and the transformed for-loop can be interpreted as a schedule for each processor. Therefore, if such one-dimensional affine transformations exist, it can be understood as a way of executing the original for-loop in a parallel manner. In the sequel, we will call parallelization such a one-dimensional affine transformation.
For the source program of Example 2, one can obtain such new coordinates: \( t = i + j \) and \( p = i \), where \( t \) and \( p \) represent time and processor, respectively. The transformed code is shown in Figure 2.10 where each iteration of the \( p \)-loop can be executed concurrently.

```cpp
// 1 <= t = i + j < 2N
for (int t = 1; t < 2*N; t++)
  // 0 <= p = i < N & 1 <= t - p = j < N
  parallel for (int p = max(0, t-N-1), p < min(t-2, N); p++)
```

**Figure 2.10:** The transformed code based on time and processor coordinates

### 2.5.6 Farkas multipliers

Let \( \mathcal{D} \subset \mathbb{Z}^m \) be a nonempty \( \mathbb{Z} \)-polyhedron defined by \( p \) affine inequalities:

\[
A_k \vec{x} + b_k \geq 0, \quad 1 \leq k \leq p,
\]

where \( A_k \in \mathbb{Z}^m \) and \( b_k \in \mathbb{Z} \). It follows from Farkas’ Lemma \([46, 105]\) that an affine form \( \phi : \vec{x} \in \mathbb{Z}^m \mapsto \phi(\vec{x}) \in \mathbb{Z} \) is non-negative in \( \mathcal{D} \) if there exist non-negative integers \( \mu_0, \mu_1, \ldots, \mu_p \), called Farkas multipliers, such that for all \( \vec{x} \in \mathbb{Z}^m \) we have:

\[
\phi(\vec{x}) = \mu_0 + \sum_{k=0}^p \mu_k (A_k \vec{x} + b_k).
\]

### 2.5.7 Feautrier’s algorithm

In \([46]\), Paul Feautrier introduced an algorithm, based on Farkas’ Lemma, that, given a for-loop nest, determines a parallelization of that for-loop nest, if such affine transformation exists. We give a brief sketch of Feautrier’s algorithm.

Consider a for-loop nest. Let us call program parameter any scalar variable read but not written in that for-loop nest. Write \( \vec{n} = (n_1, \ldots, n_d) \) a vector of all those program parameters. Typically, \( \vec{n} \) includes for-loop upper bounds, dimension sizes of arrays, etc. Consider a statement \( S_i \) in this loop nest and denote by \( \vec{x} \) a vector of the loop iteration counters occurring in the indices of the array references in \( S_i \). Then, the iteration domain \( \mathcal{D}_{S_i} \) for \( S_i \) is given by a system of \( p \) linear inequalities of the following form:

\[
A_{S_{i,k}} \begin{pmatrix} \vec{x} \\ \vec{n} \end{pmatrix} + b_{S_{i,k}} \geq 0, \quad 1 \leq k \leq p,
\]

where \( A_{S_{i,k}} \) is a row-vector over \( \mathbb{Z} \) and \( b_{S_{i,k}} \in \mathbb{Z} \) holds. Applying Farkas’ lemma, an one-dimensional affine transformation \( \vec{x} \mapsto \phi_{S_i}(\vec{x}) \) is non-negative in the domain \( \mathcal{D}_{S_i} \) if there exist non-negative integers \( \mu_{S_{i,k}} \) with \( 0 \leq k \leq p \), such that

\[
\phi_{S_i}(\vec{x}) = \mu_{S_{i,0}} + \sum_k \mu_{S_{i,k}} \left( A_{S_{i,k}} \begin{pmatrix} \vec{x} \\ \vec{n} \end{pmatrix} + b_{S_{i,k}} \right).
\]
In the sequel, we view $\phi$ as a function that, given a statement $S_i$ and an iteration vector $\bar{x}$ (of all the loop iteration counters occurring in the indices of the array references in $S_j$), maps $(S_i, \bar{x})$ to a $\phi_{S_i}(\bar{x})$, where $\phi_{S_i}$ is a one-dimensional affine transformation.

We say that $\phi$ is a schedule whenever for all statements $S_i, S_j$ and all iteration vectors $\bar{x}$ (resp. $\bar{y}$) of $S_i$ (resp. $S_j$) such that $S_i(\bar{x}) \Rightarrow S_j(\bar{y})$ holds, we have:

\[
\phi_{S_j}(\bar{y}) \geq \phi_{S_i}(\bar{x}) + 1.
\]

One can interpret the number 1 as the time unit for executing a statement.

We associate each edge $e := (S_i, S_j)$ of the DDG with a delay denoted by $\Delta_e$ and defined by

\[
\Delta_e = \phi_{S_j}(\bar{y}) - \phi_{S_i}(\bar{x}) - 1 \geq 0.
\]

Recall that the dependence polyhedron associated with $e$ (see Section 2.5.4) is a system of $q$ linear inequalities of the form:

\[
C_{e,k} \begin{pmatrix} \bar{x} \\ \bar{y} \\ \bar{n} \end{pmatrix} + d_{e,k} \geq 0, \quad 1 \leq k \leq q,
\]

where $C_{e,k}$ is a row-vector over $\mathbb{Z}$ and $d_{e,k} \in \mathbb{Z}$.

Due to the fact that a schedule does exist, we can find Farkas multipliers $\lambda_{ek}$ for the delay $\Delta_e$ such that we have

\[
\phi_{S_j}(\bar{y}) - \phi_{S_i}(\bar{x}) - 1 = \lambda_{e0} + \sum_k \lambda_{ek} \left( C_{e,k} \begin{pmatrix} \bar{x} \\ \bar{y} \\ \bar{n} \end{pmatrix} + d_{e,k} \right).
\]

Equations 2.2 and 2.3 yield a system of linear equations and inequalities including the fact all $\mu$’s and $\lambda$’s are non-negative integers. Using techniques from integer programming, in particular from the landmark paper [45] of P. Feautrier, one can compute values for the $\mu$’s and finally deduce $\phi_{S_i}(\bar{x})$.

### 2.5.8 PLUTO’s algorithm

In [15], Bondhugula, Hartono, Ramanujam and Sadayappan proposed PLUTO’s algorithm for automatically generating the OpenMP [41, 13, 9] code from a given serial C program targeting multi-cores. We describe PLUTO’s algorithm as the following. Consider a loop nest with statements $S_1, \ldots, S_\ell$. Let $\phi_{S_i}$ be a one-dimensional affine transformation for statement $S_i$, for $i = 1 \cdots \ell$. The sequence $\{\phi_{S_1}, \phi_{S_2}, \ldots, \phi_{S_\ell}\}$ is called a tiling hyperplane. We say that $\{\phi_{S_1}, \phi_{S_2}, \ldots, \phi_{S_\ell}\}$ is legal (or statement-wise) if the following holds for each dependence edge $e$ from statement $S_i$ to statement $S_j$:

\[
\phi_{S_j}(\bar{t}) - \phi_{S_i}(\bar{s}) \geq 0,
\]

where $\bar{s}$ and $\bar{t}$ are the vectors of loop iteration counters occurring in the indices of array references in $S_i$ and $S_j$, respectively. A cost function for the dependence edge $e$ is defined as the following affine form:

\[
\delta_e(\bar{s}, \bar{t}) = \phi_{S_j}(\bar{t}) - \phi_{S_i}(\bar{s}).
\]
Since the loop iteration counters themselves can be bounded by the affine functions of the parameters, one can always find an affine form \( v(\vec{n}) \) in the program parameters \( \vec{n} \), that is, \( v(\vec{n}) = \vec{u} \cdot \vec{n} + w \), where \( \vec{u} \) is a row-vector over \( \mathbb{Z} \) and \( w \in \mathbb{Z} \). Thus, for all \( \vec{s}, \vec{t} \), we have
\[
\delta_e(\vec{s}, \vec{t}) \leq v(\vec{n}).
\]  
(2.5)

Given a statement \( S \) and its iteration counters \( \vec{x} \), we define the affine transformation \( \phi_S(\vec{x}) = \vec{c} \cdot \vec{x} + c_0 \), where \( \vec{c} \) is a row-vector over \( \mathbb{Z} \) and \( c_0 \in \mathbb{Z} \). Our unknowns are \( \vec{u}, w, c_0, \vec{c} \). We determine them as the solution of a minimization problem defined as follows.

Assume that the polyhedron associated with the dependence edge \( e \) having \( p \) linear inequalities is given by \( A_{e,k} \begin{bmatrix} \vec{s} \\ \vec{t} \\ \vec{n} \end{bmatrix} + b_{e,k} \geq 0 \), where \( A_{e,k} \) is a row-vector over \( \mathbb{Z} \) and \( b_{e,k} \in \mathbb{Z} \) for \( 1 \leq k \leq p \). Then, applying the Farkas Lemma, we rewrite the legality in Equation [2.4] as
\[
\phi_s(\vec{t}) - \phi_s(\vec{s}) = \mu_{e0} + \sum_{k} \mu_{e_k} \begin{bmatrix} A_{e,k} \begin{bmatrix} \vec{s} \\ \vec{t} \\ \vec{n} \end{bmatrix} + b_{e,k} \end{bmatrix}, \quad 1 \leq k \leq p,
\]
where \( \mu_{e0}, \ldots, \mu_{ep} \) are the Farkas multipliers, and the cost function in Equation [2.5] as
\[
v(\vec{n}) - \delta_e(\vec{s}, \vec{t}) = v_{e0} + \sum_{k} \lambda_{e_k} \begin{bmatrix} A_{e,k} \begin{bmatrix} \vec{s} \\ \vec{t} \\ \vec{n} \end{bmatrix} + b_{e,k} \end{bmatrix}, \quad 1 \leq k \leq p,
\]
where \( \lambda_{e0}, \ldots, \lambda_{ep} \) are the Farkas multipliers.

We obtain constraints on \( \vec{u}, w, c_0, \vec{c} \), from a process of identification between the multipliers \( \mu \) and \( \lambda \). Then, the best solution for the dependence edge \( e \) is the lexicographic minimal solution:
\[
\text{minimize}_<\{\vec{u}, w, c_0, \vec{c}\}.
\]
Recall that given two posets \( A \) and \( B \), the lexicographical order on the Cartesian product \( A \times B \) is defined as \( (a, b) \leq (a', b') \) iff \( a < a' \) or \( (a = a' \text{ and } b \leq b') \). Solving this minimization problem can be handled by the parametric integer programming (PIP) [84] software.

Due to the fact that \( v(\vec{n}) \) gives a maximum delay to schedule the original program, if we could find an affine form with \( \vec{u} = 0 \) and \( w > 0 \), then we have a small delay to execute the statement. In the case \( \vec{u} = 0 \) and \( w = 0 \), there is no dependence in the transformed loop nest; thus the transformed loop nest can be executed in parallel. Once solutions of \( \vec{c} \) and \( c_0 \) to the affine transformation \( \phi \) are obtained by PLUTO’s algorithm, one can parallelize the for-loops iff \( \vec{u} = 0 \) and \( w \geq 0 \).

### 2.6 Solving systems of polynomial equations and inequalities

This section is an overview of the algorithmic tools used in this thesis for dealing with systems of polynomial equations and inequalities. We rely on the theory of regular chains and its implementation in the RegularChains library.
2.6. Solving systems of polynomial equations and inequalities

The notion of a regular chain (introduced independently in [70] by Kalkbrener, and in [82] by Yang & Zhang) is closely related to that of a triangular decomposition of a polynomial system. Broadly speaking, a triangular decomposition [22] of a polynomial system $S$ is a set of simpler (in a precise sense) polynomial systems $S_1, \ldots, S_e$ such that

$$p \text{ is a solution of } S \iff \exists i : p \text{ is a solution of } S_i. \quad (2.6)$$

When the purpose is to describe all the solutions of $S$, whether their coordinates are real numbers or not, in which case $S$ is said to be algebraic, those simpler systems are required to be regular chains [5]. If the coefficients of $S$ are real numbers and if only the real solutions are required, in which case $S$ is said to be semi-algebraic, then those real solutions can be obtained by a triangular decomposition into so-called regular semi-algebraic systems, a notion introduced in [22]. In both cases, each of these simpler systems has a triangular shape and remarkable properties, which justifies the terminology. We refer to [41, 25] for a formal presentation on the concepts of a regular chain and a triangular decomposition of a polynomial system. We recall, however, the concept of a regular semi-algebraic system since it is at the core of the present paper. A regular semi-algebraic system is a triple $[T, Q, P]$ where $T$ is a regular chain, $Q$ is a quantifier-free formula involving only the free variables of $T$ and $P$ is a set of polynomial inequalities; moreover $[T, Q, P]$ must satisfy the following properties.

(i) $Q$ defines a non-empty open set in the space of the free variables of $T$,

(ii) $[T, P]$ specializes well at any point $\beta$ defined by $Q$,

(iii) At any point $\alpha$ defined by $Q$, the specialized system $[T_\alpha, P_\alpha]$ admits at least one real solution $\beta$, in the sense that every polynomial in $T_\alpha$ is zero at $\beta$, and every polynomial in $P_\alpha$ is positive at $\beta$.

Consider semi-algebraic systems $[T_1, Q_1, P_1], \ldots, [T_e, Q_e, P_e]$ forming a triangular decomposition of the polynomial system $S$. A consequence of the above relation (2.6) is that, if all the $T_i$ have the same free variables, i.e. parameters, then the disjunction $Q_1 \lor \cdots \lor Q_e$ defines the set of the parameter values for which the input system possesses real solutions. However, it is not necessary for all the $T_i$ to have the same free variables and the example below given by (2.7) illustrates this fact.

An important property of any regular semi-algebraic system $[T, Q, P]$ is the fact that it is a parametrization of its zero set. Therefore, a triangular decomposition of a semi-algebraic system $S$ decomposes the zero set of $S$ into components, with each of them given by a parametric representation. This type of representation of the solutions of $S$ is very useful to compute geometrical quantities such as dimension. As a first illustration let us consider the following semi-algebraic system

$$\begin{cases} (x-1)(y^2 + t^2) + (x-2)(y^2 - t) = 0 \\ (x-1)(x-2) = 0, \end{cases} \quad (2.7)$$

and solve it with RealTriangularize command of the RegularChains library, leading to the computations on Figure 2.11.

---

http://en.wikipedia.org/wiki/Triangular_decomposition

More generally, a triangular decomposition into regular chains of a polynomial system $S$ with coefficients in an arbitrary field $K$ describes the solutions of $S$ whose coordinates are in the algebraic closure of $K$.

This means that at any point $u$ defined by $Q$, the specialized set $T(u)$ is a squarefree regular chain with the same rank as $T$ and each specialized polynomial $P_i(u)$ is invertible modulo $\langle T(u) \rangle$. 
Chapter 2. Background

The above triangular decomposition consists of three regular semi-algebraic systems. Let us denote them respectively by \([T_1, Q_1, P_1]\), \([T_2, Q_2, P_2]\), \([T_3, Q_3, P_3]\). The first and the third ones consist simply of a regular chain, thus we have \(P_1 = P_3 = \emptyset\) and \(Q_1 = Q_3 = \text{true}\). In fact each of \([T_1, Q_1, P_1]\), \([T_3, Q_3, P_3]\) simply encodes a point, that is, a zero-dimensional component. For the second one, we have \(P_2 = \emptyset\) and \(Q_2 = 0 < t\), thus \(T_2 = \{y^2 - t, x - 1\}\). Therefore, \([T_2, Q_2, P_2]\) is a parametrization of a one-dimensional component.

```maple
> R := PolynomialRing([y, x, t]);
   \textit{R:= polynomial\_ring}
> F := [(x-1)*(x-2)=0, (x-1)*(y^2+t^2) + (x-2)*(y^2-t)=0];
   \textit{F:= \{x-1\} (x-2) = 0, (x-1) (y^2 + t^2) + (x-2) (y^2 - t) = 0}
> RealTriangularize(F, R, output=record);
   \begin{align*}
   y - 0 & | \quad y^2 - t = 0 & | \quad y - 0 \\
   x - 2 - 0 & | \quad x - 1 = 0 & | \quad x - 1 - 0 \\
   t - 0 & | \quad t > 0 & | \quad t - 0 
   \end{align*}
```

Figure 2.11: A triangular decomposition into semi-algebraic systems computed with the \texttt{RealTriangularize} command

```
> R := PolynomialRing([x, y, z]); F := [5*x^2+2*x*z+5*y^4+15*y^5-5*y^3-15*y^5-5*y^3+3];
   \textit{F:= \{5*x^2 + 2*x*z + 5*y^4 + 15*y^5 - 5*y^3 - 15*y^5 - 5*y^3 + 3\}}
> RealTriangularize(F, R, output=record);
   \begin{align*}
   5 x^2 + 2 z^2 x + 5 y^6 + 15 y^4 - 5 y^3 - 15 y^5 + 5 z^2 - 0 \\
   25 y^6 - 75 y^5 + 15 y^4 - 25 y^3 + 25 z^2 < 0 \\
   5 x + z^2 = 0 \\
   25 y^6 - 75 y^5 + 15 y^4 - 25 y^3 - z^4 + 25 z = 0 \\
   64 z^3 - 1600 z^2 + 25 > 0 \\
   z \neq 0 \\
   z - 5 = 0 \\
   z + 5 = 0 \\
   x + 5 = 0 & | \quad x + 0 & | \quad x + 5 - 0 \\
   y = 0 & | \quad y - 1 = 0 & | \quad y = 0 \\
   z - 5 = 0 & | \quad z + 5 = 0 & | \quad z + 5 = 0 \\
   5 x + z^2 = 0 \\
   2 y - 1 = 0
\end{align*}
```

Figure 2.12: Output of the \texttt{RealTriangularize} command for the \textit{EVE} surface

Figure 2.12 contains a second and more advanced example, where the purpose of the MAPLE session is to obtain a description of the real points of the hypersurface \textit{EVE} from the \textit{Algebraic Surface Gallery\footnote{This is a collection of algebraic surfaces, well-known in the mathematical literature and available at \url{http://www.freigeist.cc/gallery.html}}} and whose equation is

\[5 x^2 + 2 x z^2 + 5 y^6 + 15 y^4 + 5 z^2 - 15 y^5 - 5 y^3 = 0.\]
The solutions of the above are all \((x, y, z)\) where \(x, y, z\) are complex numbers satisfying this equation. The output of `RealTriangularize` consists of 9 regular semi-algebraic systems for which the variables are ordered as \(x > y > z\). The first regular semi-algebraic system represents a two-dimensional component. Indeed, it defines \(x\) as the solution of a parametric equation of degree 2, where \(y, z\) are regarded as parameters subject to an inequality (defined by the discriminant of the equation) which ensures the existence of two \(x\)-values for each valid \((y, z)\)-value. The second regular semi-algebraic system represents a one-dimensional component: the two equations define \((x, y)\) as functions of \(z\), which is subject to various inequalities. Each of the other 7 regular semi-algebraic systems encodes a zero-dimensional component, that is, a finite set of points.

Triangular decompositions into regular semi-algebraic systems are an interesting representation of semi-algebraic sets for the following reasons. First, triangular decompositions into regular chains are a space-efficient encoding of algebraic sets. This fact is formally established in [42] and experimentally verified with the `RegularChains` library in [25]. Secondly, triangular decompositions into regular chains (or regular semi-algebraic systems) reveal important geometrical properties (dimensions of the irreducible components, fibration structure, etc.) of the input algebraic sets (or semi-algebraic sets), as illustrated by the previous examples. These can be used to design efficient algorithms for the operations manipulating (semi-)algebraic sets. For instance, performing set-theoretic operations (in particular, set theoretical difference) can be done very efficiently on both constructible sets and semi-algebraic sets as reported in [24] and [23], respectively. Last but not least, triangular decompositions and related techniques, such as dynamic evaluation, are well suited for supporting weaker solving specifications. Two examples of that are triangular decomposition of algebraic systems in the sense of Kalkbrener [70] and lazy triangular decomposition of semi-algebraic systems [22]. These types of decompositions provide a description of the “generic solutions” plus a continuation mechanism for obtaining the other solutions, if necessary. One major supporting argument for those decompositions is the existence of favorable algebraic complexity estimates. Both of them can indeed be computed in singly exponential time with respect to the number of variables, as established in [112] and [22], respectively.
Chapter 3

The Basic Polynomial Algebra Subprograms

The Basic Polynomial Algebra Subprograms (BPAS) provide arithmetic operations (multiplication, division, root isolation, etc.) for univariate and multivariate polynomials over prime fields or with integer, rational number or complex rational number coefficients. The code is mainly written in CilkPlus [10, 79] targeting multi-core processors. The current distribution focuses on dense polynomials and the sparse case is work in progress. A strong emphasis is put on adaptive algorithms as the library aims at supporting a wide variety of situations in terms of problem sizes and available computing resources. One of the purposes of the BPAS project is to take advantage of hardware accelerators in the development of polynomial system solvers. The BPAS library is publicly available in source at www.bpaslib.org.

We describe the design and specification in Section 3.1. In Section 3.2, we introduce the user interface of our library. In Sections 3.3 and 3.4, we demonstrate the core algorithms, like FFT and polynomial multiplication, and the experimental results, respectively. Applications to solve real root isolation and symbolic-numeric integration are summerized in Section 3.5. Appendix A shows sample code in the BPAS library.

This chapter is an extended version of [20] and contains joint work with Changbo Chen, Svyatoslav Covanov, Farnam Mansouri, Robert Moir, Marc Moreno Maza and Yuzhen Xie.

3.1 Design and specification

Inspired by the Basic Linear Algebra Subprograms (BLAS) [77], BPAS functionalities are organized into three levels. At Level 1, one finds basic arithmetic operations that are specific to a polynomial representation or specific to a coefficient ring. Examples of Level-1 operations are
multi-dimensional FFTs/TFTs and univariate real root isolation. At Level 2, arithmetic operations are implemented for all types of coefficients rings that BPAS supports (prime fields, ring of integers, field of rational numbers). Level 3 gathers advanced arithmetic operations taking as input a zero-dimensional regular chain, e.g. normal form of a polynomial, multivariate real root isolation.

Level 1 functions are highly optimized in terms of data locality and parallelism. In particular, the underlying algorithms are nearly optimal in terms of cache complexity [49]. This is the case, for instance, for our modular multi-dimensional FFTs/TFTs [94], modular dense polynomial arithmetic [23] and Taylor shift [28] algorithms.

At Level 2, users can choose between algorithms that either minimize work (at the possible expense of decreasing parallelism) or maximizes parallelism (at the possible expense of increasing work). For instance, five different integer polynomial multiplication algorithms are available, namely: Schönhage-Strassen [104], 8-way Toom-Cook [14], 4-way Toom-Cook [14], divide-and-conquer plain multiplication and the two-convolution method [21]. The first one has optimal work (i.e. algebraic complexity) but is purely serial due to the difficulties of parallelizing 1D FFTs on multi-core processors. The next three algorithms are parallelized but their parallelism is static, that is, independent of the input data size; these algorithms are practically efficient when both the input data size and the number of available cores are small, see [87] for details. The fifth algorithm relies on modular 2D FFTs which are computed by means of the row-column scheme; this algorithm delivers high scalability and can fully utilize the hardware on fat multi-core nodes.

Another example of Level 2 functionality is parallel Taylor shift computation for which four different algorithms are available: the two plain algorithms presented in [28], Algorithm (E) of [119] and an optimized version of Algorithm (F) of [119]. The first two are highly effective when both the input data size and the number of available cores are small. The third algorithm creates parallelism by means of a divide-and-conquer procedure and relies on polynomial multiplication; this approach is effective when 8-way Toom-Cook multiplication is selected. The fourth algorithm reduces a Taylor shift computation to a single polynomial multiplication; this latter approach outperforms the other three, as soon as the two-convolution multiplication dominates its counterparts, that is, when either input data size and the number of available cores become large.

This variety of parallel solutions leads, at Level 3, to adaptive algorithms which select appropriate Level 2 functions depending on available resources (number of cores, input data size). An example is parallel real root isolation. Many procedures for this purpose are based on a subdivision scheme. However, on many examples, this scheme exposes only a limited amount of opportunities for concurrent execution, see [28]. It is, therefore, essential to extract as much as parallelism from the underlying routines, such as Taylor shift computations.

### 3.2 User interface

Inspired by computer algebra systems like AXIOM [69] and Magma [16], the BPAS library makes use of type constructors [96] so as to provide genericity. For instance, SparseUnivariatePolynomial (SUP) can be instantiated over any BPAS ring. On the other hand, for efficiency consideration, certain polynomial type constructors, like DistributedDenseMultivariateMod-
ularPolynomial (DDMMP), are only available over finite fields in order to ensure that the data encoding a DDMMP polynomial consists only of consecutive memory cells. For the same efficiency consideration, the most frequently used polynomial rings, like DenseUnivariateIntegerPolynomial (DUZP) and DenseUnivariateRationalNumberPolynomial (DUQP) are primitive types. Consequently, DUZP and SUP<Integer> implement the same functionalities; however, the implementation of the former is further optimized. Figure 3.1 shows a subset of BPAS’s tree of algebraic data structures. Each class with its name starting with “BPAS” is defined as an abstract class, while others are defined as concrete classes. Abstract classes are used to indicate the public members that their children must implement. Each concrete derived class overrides the pure virtual member functions of its parent(s).

With the support of C++ operator overloading, we define the basic arithmetic operations ‘+’, ‘-’, ‘*’ and ‘/’ for each BPAS polynomial to execute addition, subtraction, multiplication and exact division, respectively. In addition, the insertion (‘<<’) operator for output streams (to the screen and files) is overloaded for each BPAS polynomial as well as each BPAS field. Particularly for BPAS univariate polynomials, we redefine shift operators ‘<<’ and ‘>>’ to represent multiplying and dividing, respectively, by a power of the variable. Several member functions, such as monicDivide, pseudoDivide, squareFree and gcd, are extended for BPAS univariate polynomials as well.

While the concrete class Integer (resp. RationalNumber) inherits from BPASRing and mpz_class (resp. mpq_class) (from the GMP library), arithmetic operations (addition, subtraction, multiplication and division) rely on the GMP library. Since the sparse univariate or multivariate polynomial can take an arbitrary ring as the coefficient ring, the programmer shall realize the coefficient ring with the pure virtual functions defined by BPAS ring class, such as isZero() and zero(). In order to identify the type of the coefficient ring in the sparse univariate polynomial, we add static attributes, such as characteristic and isPrimeField, to each BPAS ring, so that SUP<Ring> (and later SMP<Ring>) can invoke an efficient implementation of the functionality from those polynomial classes in a dense representation.
Moreover, we implement a `ComplexRationalNumber` class based on two rational numbers from the GMP library to represent the real part and the imaginary part, respectively. With this support as well as the definition of the BPAS field, we expand the BPAS ring with a univariate rational function so as to solve symbolic numeric integration. Moreover, the conversion between BPAS univariate polynomials, `Integer`, `RationalNumber` and `ComplexRationalNumber` can be done via the construction of each concrete class.

![Figure 3.2: Another subset of BPAS algebraic data structures](image)

Last but not least, BPAS is capable of isolating real roots given a regular chain over $\mathbb{Q}$. Figure 3.2 shows another subset of BPAS data structures to support triangular sets and regular chains of polynomial systems. BPAS also counts other classes, for instance, `Intervals`.

```cpp
#include <bpas.h>

int main(int argc, char *argv[]) {
    int d = 4095;
    /* Univariate Integer Polynomial Multiplication */
    DUZP a(d+1), b(d+1);
    a.read("a_input.dat"); b.read("b_input.dat");
    DUZP c = a * b;
    std::cout << "c = " << c << std::endl;
    /* Real Root Isolation */
    DUQP p;
    p = (p + mpq_class(1) << d) + mpq_class(d); // Cn,d(x) = x^d + d
    Intervals boxes = p.realRootIsolate(mpq_class(1, 20));
    std::cout << "boxes = " << boxes << std::endl;
    /* Symbolic Numeric Integration */
    SparseUnivariatePolynomial<RationalNumber> f, g;
    f.one(); f.setVariableName("x");
    g = (polynomialParser("1+2*x+2*x^2")^4);
    UnivariateRationalFunction<SparseUnivariatePolynomial<RationalNumber>, RationalNumber> h (f, g);
    h.realSymbolicNumericIntegrate(53);
    return 0;
}
```

![Figure 3.3: A snapshot of BPAS code](image)

The snapshot of the BPAS code in Figure 3.3 first shows how two dense univariate polynomials over $\mathbb{Z}$ are read from files and how their product is computed. Then, on the same code
fragment, a dense univariate polynomial over $\mathbb{Q}$ is assigned by operators ‘+’ and ‘<<’ and its real roots are isolated. Finally, it has a polynomial parser of $\text{SUP<RationalNumber>}$ and symbolic numeric integration of a univariate rational function.

### 3.3 Implementation techniques

Modular FFTs are at the core of asymptotically fast algorithms for dense polynomial arithmetic operations. A substantial body of code of the BPAS library is, therefore, devoted to the computation of one-dimensional and multi-dimensional FFTs over finite fields. In the current release, the characteristic of those fields is of machine word size, while larger characteristics are a work in progress.

The techniques used for the multi-dimensional FFTs are described in [94, 93] while those for one-dimensional FFTs are inspired by the design of the FFTW [48].

BPAS’ one-dimensional FFTs code is optimized in terms of cache complexity and register usage. To achieve this, the FFT of a vector of size $n$ is computed in a divide-and-conquer manner until the vector size is smaller than a threshold, at which point FFTs are computed using a tiling strategy [124]. This threshold can be specified by the user through an environment variable $\text{HTHRESHOLD}$ or determined automatically when installing the library. At compile time, this threshold is used to generate and optimize the code. For instance, the code of all FFTs of size less than or equal to $\text{HTHRESHOLD}$ is decomposed into blocks (typically performing FFTs on 8 or 16 points) for which straight-line program (SLP) [5] machine code is generated. Instruction level parallelism (ILP) is carefully considered: vectorized instructions are explicitly used (SSE2, SSE4) and instruction pipeline usage is highly optimized. Other environment variables are available for the user to control different parameters in the code generation.

#### Table 3.1: One-dimensional modular FFTs: Modpn vs BPAS

<table>
<thead>
<tr>
<th>Input size</th>
<th>Modpn</th>
<th>BPAS</th>
<th>Speedup</th>
</tr>
</thead>
<tbody>
<tr>
<td>16777216</td>
<td>6.232</td>
<td>1.391</td>
<td>4.48</td>
</tr>
<tr>
<td>33554432</td>
<td>12.987</td>
<td>2.957</td>
<td>4.392</td>
</tr>
<tr>
<td>67108864</td>
<td>26.783</td>
<td>6.266</td>
<td>4.274</td>
</tr>
<tr>
<td>134217728</td>
<td>55.329</td>
<td>13.235</td>
<td>4.181</td>
</tr>
<tr>
<td>268435456</td>
<td>113.8</td>
<td>27.901</td>
<td>4.079</td>
</tr>
</tbody>
</table>

Table 3.1 compares running times (in sec. on Intel Xeon 5650) of one-dimensional modular FFTs computed by the Modpn library [80] and BPAS, both using serial C code in this case. The first column of Table 3.1 gives the size of the input vector; coefficients are in a prime field whose characteristic is a 57-bit prime.

Modular FFTs support the implementation of several algorithms performing dense polynomial arithmetic. As an example, we consider parallel multiplication of dense polynomials with integer coefficients by means of the two-convolution method [21] and which is illustrated on Figure 3.4. Given two univariate polynomials $a(y), b(y)$ with integer coefficients, their product $c(y)$ is computed as follows.

(S1) Convert $a(y), b(y)$ to bivariate integer polynomials $A(x, y), B(x, y)$ s.t. $a(y) = A(\beta, y)$ and
3.4. Experimental evaluation

\[ b(y) = B(\beta, y) \] hold at \( \beta = 2^M \), \( K = \deg(A, x) = \deg(B, x) \), where \( M \) is essentially the maximum bit size of a coefficient in \( a \) and \( b \).

(S2) Consider \( C^+(x, y) \equiv A(x, y) \ B(x, y) \mod (x^K + 1) \) and \( C^-(x, y) \equiv A(x, y) \ B(x, y) \mod (x^K - 1) \). Compute \( C^+(x, y) \) and \( C^-(x, y) \) modulo machine-word primes so as to use modular 2D FFTs.

(S3) Consider \( C(x, y) = \frac{C^+(x, y)}{2} (x^K - 1) + \frac{C^-(x, y)}{2} (x^K + 1) \) and evaluate \( C(x, y) \) at \( x = \beta \), which finally gives \( c(y) = a(y) \ b(y) \).

![Figure 3.4: Multiplication scheme for dense univariate integer polynomials](image)

The conversions from the univariate polynomials \( a(y), b(y) \) to the bivariate polynomials \( A(x, y), B(x, y) \) in Step (S1) as well as the conversions from the bivariate polynomials \( C^+(x, y) \) and \( C^-(x, y) \) in Step (S3) require only additions and shift operations on machine words. Moreover, the polynomials \( C^+(x, y) \) and \( C^-(x, y) \) are reconstructed from their modular images (in practice two modular images are sufficient) within Step (S3). Consequently, the data produced by 2D FFT computations is converted in a single pass into the final result \( c(y) \). Similarly the bivariate polynomials \( A(x, y), B(x, y) \) are obtained from \( a(y), b(y) \) (here again by means of additions and shift operations on machine words) in a single pass. Since BPAS’ 2D FFT computations are optimal in terms of cache complexity \[93\], the whole multiplication procedure is optimal for that same complexity measure. Last but not least, BPAS’ 2D FFTs are computed by the row-column scheme which provides lots of parallelism with limited overheads on multi-core architectures. As a result, our multiplication code, based on this two-convolution method scales well on multi-cores as illustrated hereafter.

3.4 Experimental evaluation

As mentioned above, one of the main purposes of the BPAS library is to take advantage of hardware accelerators and support the implementation of polynomial system solvers. With this
goal, polynomial multiplication plays a central role. Moreover, both sparse and dense representations are important. Indeed, input polynomial systems are often sparse while many algebraic transactions, like substitution, tend to densify data. Parallel sparse polynomial arithmetic has been studied by Gastineau and Laskar in [50] and by Monagan and Pearce in [91].

To the best of our knowledge, BPAS is the first publicly available library for parallel dense integer polynomial arithmetic. For this reason, we compare BPAS’ parallel dense polynomial multiplication against its state-of-the-art counterpart implementation in FLINT 2.5.2 and Maple 2015. In Figure 3.5, the input of each test case is a pair of polynomials of degree $d$ where each coefficient has bit size $N$. Two plots are provided: one for which $d = N$ holds and one for which $d$ is much smaller than $N$.

The BPAS library is implemented with the multithreaded language CilkPlus [79] and we compile our code with the CilkPlus branch of GCC. Our experimental results were obtained on an 48-core AMD Opteron 6168, running at 900Mhz with 256 GB of RAM and 512KB of L2 cache. Figure 3.6 shows that using an interactive process viewer for Unix systems, namely htop, BPAS can use all available cores when multiplying two large integer polynomials.

Figure 3.5: Dense integer polynomial multiplication: BPAS vs FLINT vs Maple

http://gcc.gnu.org/svn/gcc/branches/cilkplus/
3.5. Application

Table 3.2 shows that the work overhead (measured by Cilkview, the performance analysis tool of CilkPlus) of the BPAS method w.r.t. to a method based on Schönhage & Strassen algorithm (KS) is only around 2, whereas BPAS provides a large amount of parallelism, see the last column.

Table 3.2: Cilkview analysis of BPAS and KS (* shows the number of instructions)

<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>*</td>
<td></td>
<td>*</td>
<td></td>
</tr>
<tr>
<td>2048</td>
<td>795,549,545</td>
<td>1,364,160,088</td>
<td>41,143,119</td>
<td>33.16</td>
<td>1.715</td>
</tr>
<tr>
<td>4096</td>
<td>4,302,927,423</td>
<td>5,663,423,709</td>
<td>96,032,325</td>
<td>58.97</td>
<td>1.316</td>
</tr>
<tr>
<td>8192</td>
<td>16,782,031,611</td>
<td>23,827,123,688</td>
<td>292,735,521</td>
<td>81.39</td>
<td>1.420</td>
</tr>
<tr>
<td>16384</td>
<td>63,573,232,166</td>
<td>100,688,072,711</td>
<td>1,017,726,160</td>
<td>98.93</td>
<td>1.584</td>
</tr>
<tr>
<td>32768</td>
<td>269,887,534,779</td>
<td>425,149,529,176</td>
<td>3,804,178,563</td>
<td>111.76</td>
<td>1.575</td>
</tr>
</tbody>
</table>

3.5 Application

Turning to parallel univariate real root isolation, given a squarefree univariate polynomial \( f(x) = a_d x^d + \cdots + a_1 x + a_0 \in \mathbb{Q}[x] \), we compute the real roots of \( f(x) = 0 \). We represent real roots as a list of pairwise disjoint intervals \([a_1, b_1], \ldots, [a_e, b_e]\) with rational number endpoints, such that (1) each \([a_i, b_i]\) contains one and only one real root of \( f(x) \); (2) if \( a_i = b_i \), the real root \( x_i = a_i (b_i) \); (3) otherwise, the real root \( a_i < x_i < b_i \) (\( f(x) \) does not vanish at either endpoint).

In [28], Chen, Moreno Maza and Xie presents a parallel algorithm (CMY) directly using the Vincent-Collins-Akritas (VCA) [32] algorithm for univariate real root isolation targeting multi-cores. Authors observe that the most consuming time is the operation of so-called Taylor shift [119], that is, the map \( f(x) \mapsto f(x + 1) \). Among six classic Taylor shift algorithms reported in [119], the authors in [28] parallelize the Horner’s method, which represents the computation in a Pascal’s triangle. This method has an optimal cache complexity [49] but runs in quadratic time.

BPAS performs the Taylor shift operation by means of a divide and conquer method, that is, the Algorithm (E) in [119]. It reduces calculations to integer polynomial multiplications in large degrees and continues to use the algorithm of [28] in small degrees. This adaptive
Taylor shift algorithm combines FFT-based arithmetic (via Algorithm (E)) and plain arithmetic (via [28]).

We run two parallel univariate real root algorithms, BPAS and CMY [28], which are both implemented in CilkPlus, against MAPLE 18 serial realroot command. Table 3.3 shows the running times (in sec.) of four well-known test problems, including Cnd, Chebycheff, Laguerre and Wilkinson. Moreover, for each test problem, the degree of the input polynomial varies in a range. The results reported in Table 3.3 show that integrating parallel integer polynomial multiplication into our real root isolation code has substantially improved the performance of the latter.

Table 3.3: Univariate real root isolation running times (in secs.) for four examples

<table>
<thead>
<tr>
<th></th>
<th>Size</th>
<th>BPAS</th>
<th>CMY [28]</th>
<th>realroot</th>
<th>#Roots</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cnd</td>
<td>32768</td>
<td>18.141</td>
<td>125.902</td>
<td>816.134</td>
<td>1</td>
</tr>
<tr>
<td></td>
<td>65536</td>
<td>66.436</td>
<td>664.438</td>
<td>7,526.428</td>
<td>1</td>
</tr>
<tr>
<td>Chebycheff</td>
<td>2048</td>
<td>608.738</td>
<td>594.82</td>
<td>1,378.444</td>
<td>2047</td>
</tr>
<tr>
<td></td>
<td>4096</td>
<td>8,194.06</td>
<td>10,014</td>
<td>35,880.069</td>
<td>4095</td>
</tr>
<tr>
<td>Laguerre</td>
<td>2048</td>
<td>1,336.14</td>
<td>1,324.33</td>
<td>3,706.749</td>
<td>2047</td>
</tr>
<tr>
<td></td>
<td>4096</td>
<td>20,727.9</td>
<td>23,605.7</td>
<td>91,668.577</td>
<td>4095</td>
</tr>
<tr>
<td>Wilkinson</td>
<td>2048</td>
<td>630.481</td>
<td>614.94</td>
<td>1,031.36</td>
<td>2047</td>
</tr>
<tr>
<td></td>
<td>4096</td>
<td>9,359.25</td>
<td>10,733.3</td>
<td>26,496.979</td>
<td>4095</td>
</tr>
</tbody>
</table>

Consider a system of $n$-variable polynomials over $\mathbb{Q}$ with $n$ equations:

\[
\begin{align*}
    f_1(x_1, x_2, \ldots, x_n) &= 0, \\
    f_2(x_1, x_2, \ldots, x_n) &= 0, \\
    \vdots \\
    f_n(x_1, x_2, \ldots, x_n) &= 0.
\end{align*}
\]

We compute the common real roots of $f_i(x_1, x_2, \ldots, x_n) = 0$ for each $1 \leq i \leq n$. In this case, each real root is the Cartesian product of $n$ intervals (one for each of the coordinates $x_1, \ldots, x_n$) with two endpoints $[a, b] \in \mathbb{Q}$.

After applying the Triangularize algorithm [25] to the above system, one obtains a description of the solution set in the form of a list of regular chains. A regular chain is a particular type of polynomial systems with a triangular shape:

\[
\begin{align*}
    g_n(x_1, x_2, \ldots, x_n) &= 0, \\
    \vdots \\
    g_2(x_1, x_2) &= 0, \\
    g_1(x_1) &= 0.
\end{align*}
\] (3.1)

In addition, for all $2 \leq i \leq n$, the leading coefficient of $g_i$ does not vanish at any root $(x_1, \ldots, x_{i-1})$ of $g_1 = \cdots = g_{i-1} = 0$.

One can write each multivariate polynomial as

\[
g_i = h_i(x_1, \ldots, x_{i-1}) x_i^f + \cdots + h_1(x_1, \ldots, x_{i-1}) x_i + h_0(x_1, \ldots, x_{i-1}),
\]
where \( h_\ell, h_{\ell-1}, \ldots, h_1, h_0 \in \mathbb{Q}[x_1, \ldots, x_{\ell-1}] \) hold and \( \ell \) is the degree of \( g_i \) w.r.t. its leading variable \( x_i \).

For each isolation box \( X = ([a_1, b_1], \ldots, [a_{\ell-1}, b_{\ell-1}]) \) of a real root of \( g_1 = \cdots = g_{\ell-1} = 0 \), where \([a_j, b_j] \in \mathbb{Q}\) for \( 1 \leq j \leq \ell - 1 \), we evaluate \( g_i \) by \( h_\ell(X) x_\ell^\ell + \cdots + h_0(X) \), using the interval arithmetic [126]: \( [\alpha, \beta] + [\gamma, \delta] = [\alpha + \gamma, \beta + \delta] \) and \( [\alpha, \beta] \times [\gamma, \delta] = [\min(\theta), \max(\theta)] \), where \( \theta = [\alpha \gamma, \alpha \delta, \beta \gamma, \beta \delta] \) and \([\alpha, \beta], [\gamma, \delta] \) are intervals.

Then we obtain
\[
g_i(X, x_i) = [c_\ell, d_\ell] x_\ell^\ell + \cdots + [c_0, d_0], \quad \text{where} \quad [c_j, d_j] \in \mathbb{Q}, \quad \text{for} \quad 0 \leq j \leq \ell.
\]

Univariate polynomials \( g_j = c_\ell x_\ell^\ell + \cdots + c_0 \) and \( \overline{g}_i = d_\ell x_\ell^\ell + \cdots + d_0 \) are defined as the *sleeve bound polynomials* [83] of \( g_i \) in \( X \). For \( x_i > 0 \), we have
\[
 g_j(x_i) \leq g_i(X, x_i) \leq \overline{g}_i(x_i).
\]

Now, we can apply the univariate real root isolation routine to compute the real roots of \( g_j(x_i) \) and \( \overline{g}_i(x_i) \), so as to obtain two lists of real root intervals, namely \( List_1 \) and \( List_2 \), respectively. The sequence of the intervals from \( List_1 \) and \( List_2 \) is *matchable*, with respect to a range value defined by the user, whenever both the following conditions are satisfied (see Figure 3.7 for an example):

1. The sequence of the real root intervals from \( List_1 \) and \( List_2 \) takes one of the following forms (‘1’ denotes an interval belonging to \( List_1 \) and ‘2’ denotes an interval of \( List_2 \)):
   (a) 1, 2, 2, 1, 1, 2, 2, 1, \ldots, 1, 2, 2, 1;
   (b) 1, 2, 2, 1, 1, 2, 2, 1, \ldots, 1, 2, 2, 1, 1, 2;
   (c) 2, 1, 1, 2, 2, 1, 1, 2, \ldots, 2, 1, 1, 2;
   (d) 2, 1, 1, 2, 2, 1, 1, 2, \ldots, 2, 1, 1, 2, 2, 1.

2. In each interval from \( List_1 \) and \( List_2 \), the considered polynomial \( g_i(X, x_i) \) is *monotonic*.

Once such a matchable sequence of intervals is obtained, we say the real roots of \( g_i \) are found.

![Figure 3.7: An example of matchable interval lists](image)

For each real root isolation box \( X = x_1, \ldots, x_{\ell-1} \), we parallelize the computation of isolating \( x_i \) of \( g_i(X, x_i) \) by using the algorithm described above. In general, after Triangularize an \( n \)-variable polynomial system, a list of regular chains is obtained. Thus, we can parallelize the real root isolation for each regular chain. Given a regular chain following the format of Equation 3.1, one can use \texttt{realRootIsolate} from the \texttt{RationalRegularChain} class in the \texttt{BPAS} library to isolate real roots of polynomial systems.

We compare our implementation with MAPLE 17 \texttt{RealRootIsolate} and \texttt{Isolate} written in C. In Table 3.4, we collect the running times (in secs.) and speedup factors. We observe that for 12 of 17 examples, \texttt{BPAS} outperforms the others and that the speedup factor varies from 1 to 16. To the best of our knowledge, this is the first parallel implementation of multivariate real root isolation.
Table 3.4: Running times (in secs.) of multivariate real root isolation: BPAS vs Maple 17 RealRootIsolate vs C (with Maple 17 interface) Isolate

<table>
<thead>
<tr>
<th>Example</th>
<th>BPAS (CilkPlus)</th>
<th>RealRootIsolate</th>
<th>Isolate</th>
<th>Speedup factors</th>
</tr>
</thead>
<tbody>
<tr>
<td>4-Body-Homog</td>
<td>0.402</td>
<td>0.608</td>
<td>0.382</td>
<td></td>
</tr>
<tr>
<td>Arnborg-Lazard</td>
<td>0.146</td>
<td>0.299</td>
<td>0.066</td>
<td></td>
</tr>
<tr>
<td>Caprasse</td>
<td>0.018</td>
<td>0.14</td>
<td>0.154</td>
<td>7.778</td>
</tr>
<tr>
<td>Circles</td>
<td>0.051</td>
<td>0.894</td>
<td>0.814</td>
<td>15.961</td>
</tr>
<tr>
<td>Cyclic-5</td>
<td>0.021</td>
<td>0.147</td>
<td>0.206</td>
<td>9.810</td>
</tr>
<tr>
<td>Czapor-Geddes-Wang</td>
<td>0.2</td>
<td>0.135</td>
<td>0.184</td>
<td></td>
</tr>
<tr>
<td>D2v10</td>
<td>0.029</td>
<td>0.075</td>
<td>177.999</td>
<td>2.586</td>
</tr>
<tr>
<td>D4v5</td>
<td>0.037</td>
<td>0.044</td>
<td>49.09</td>
<td>1.189</td>
</tr>
<tr>
<td>Fabfaux</td>
<td>0.192</td>
<td>0.231</td>
<td>0.071</td>
<td></td>
</tr>
<tr>
<td>Katsura-4</td>
<td>0.171</td>
<td>0.416</td>
<td>0.044</td>
<td></td>
</tr>
<tr>
<td>L-3</td>
<td>0.02</td>
<td>0.252</td>
<td>0.12</td>
<td>6.0</td>
</tr>
<tr>
<td>Neural-Network</td>
<td>0.029</td>
<td>0.332</td>
<td>0.131</td>
<td>4.517</td>
</tr>
<tr>
<td>R-6</td>
<td>0.014</td>
<td>0.048</td>
<td>20.612</td>
<td>3.429</td>
</tr>
<tr>
<td>Rose</td>
<td>0.026</td>
<td>0.336</td>
<td>0.599</td>
<td>12.923</td>
</tr>
<tr>
<td>Takeuchi-Lu</td>
<td>0.027</td>
<td>0.16</td>
<td>0.031</td>
<td>1.148</td>
</tr>
<tr>
<td>Wilkinsonxy</td>
<td>0.023</td>
<td>0.165</td>
<td>0.046</td>
<td>2.0</td>
</tr>
<tr>
<td>Nld-10-3</td>
<td>1.249</td>
<td>8.993</td>
<td>707.334</td>
<td>7.20</td>
</tr>
</tbody>
</table>

Furthermore, for symbolic-numeric integration of symbolic functions focusing on rational functions, we rely on BPAS polynomial arithmetic operations (multiplication, division, root isolation, etc.) for univariate polynomials with integer, rational number or complex rational number coefficients. Our method proceeds by using the Lazard-Rioboo-Trager algorithm [78], while it relies on efficient numerical computation of isolating polynomial roots followed by symbolic post-processing. The symbolic computation is done within BPAS, while the numerical computation is done via using a highly optimized multiprecision rootfinding package, namely MPSolve.

The implementation of our algorithm is integrated in the BPAS library; it can be called through the realSymbolicNumericIntegrate method of the UnivariateRationalFunction template class. The following output formats are available: approximate (either floating point number or rational number) and symbolic expression (in either Maple or MATLAB syntax); see Figure 3.8 for a combination of floating point and Maple output formats.

![Figure 3.8: A sample output of realSymbolicNumericIntegrate](image-url)
Chapter 4

A Many-Core Machine Model for Designing Algorithms with Minimum Parallelism Overheads

We present a model of multithreaded computation with an emphasis on estimating the parallelism overheads of programs written for modern many-core architectures. We establish a Graham-Brent theorem so as to estimate execution time of programs running on a given number of streaming multiprocessors. We evaluate the benefits of our model on fundamental algorithms from scientific computing. For three case studies, our model is used to minimize parallelism overheads by determining an appropriate value range for a given program parameter. For other cases, our model is used to compare different algorithms solving the same problem. In each case, the studied algorithms were implemented and the results of their experimental comparison are coherent with the theoretical analysis based on our model.

This chapter is an extended version of [61] and contains joint work with Sardar Anisul Haque and Marc Moreno Maza.

4.1 Introduction

Designing efficient algorithms targeting hardware accelerators (multi-core processors, graphics processing units (GPUs) and field-programmable gate arrays) creates major challenges for computer scientists. A first difficulty is to define models of computation retaining the computer hardware characteristics that have a dominant impact on program performance. In addition to specifying the appropriate complexity measures, those models must consider the relevant parameters characterizing the abstract machine executing the algorithms to be analyzed. A second difficulty is, for a given model of computation, to combine its complexity measures so as to determine the “best” algorithm among different possible solutions to a given algorithmic problem.

In the fork-join concurrency model [11], two complexity measures, the work $T_1$ and the span $T_\infty$, and one machine parameter, the number $P$ of processors, are combined into a running time estimate, namely the Graham-Brent theorem [11, 53], which states that the running time $T_P$ on $P$ processors satisfies $T_P \leq T_1/P + T_\infty$. A refinement of this theorem supports the imple-
mentation (on multi-core architectures) of the parallel performance analyzer Cilkview [63]. In this context, the running time $T_P$ is bounded in expectation by $T_1/P + 2\delta T_\infty$, where $\delta$ is a constant (called the *span coefficient*) and $T_\infty$ is the burdened span, which captures parallelism overheads due to scheduling and synchronization.

The well-known PRAM (parallel random-access machine) model [110, 51] has also been enhanced [1] so as to integrate communication delay into the computation time. However, a PRAM abstract machine consists of an unbounded collection of RAM processors, whereas a many-core GPU holds a collection of streaming multiprocessors (SMs). Hence, applying the PRAM model to GPU programs fails to capture all the features (and thus the impact) of data transfer between the SMs and the global memory of the device.

Ma, Agrawal and Chamberlain [85] introduce the TMM (Threaded Many-core Memory) model, which retains many important characteristics of GPU-type architectures as machine parameters, like memory access bandwidth and hardware limit on the number of threads per core. In TMM analysis, the running time of an algorithm is estimated by choosing the maximum quantity among work, span and the number of memory accesses. Such running time estimates depend on the machine parameters. Hong and Kim [67] present an analytical model to predict the execution time of an actual GPU program. No abstract machine is defined in this case. Instead, a few metrics are used to estimate the CPI (cycles per instruction) of the considered program.

Many works, such as [86, 81], targeting code optimization and performance prediction of GPU programs are related to our work. However, these papers do not define an abstract model in support of the analysis of algorithms.

In this chapter, we propose a many-core machine (MCM) model with two objectives: (1) tuning program parameters to minimize parallelism overheads of algorithms targeting GPU-like architectures, and (2) comparing different algorithms independently of the targeted hardware device. In the design of this model, we insist on the following features:

1. **Two-level DAG programs.** Defined in Section 4.2, they capture the two levels of parallelism (fork-join and single instruction, multiple data) of heterogeneous programs (like a CilkPlus program using #pragma simd [100] or a CUDA program with the so-called dynamic parallelism [37]).

2. **Parallelism overhead.** We introduce this complexity measure in Section 4.2.3 with the objective of capturing communication and synchronization costs.

3. **A Graham-Brent theorem.** We combine three complexity measures (work, span and parallelism overhead) and one machine parameter (data transfer throughput) in order to estimate the running time of an MCM program on $P$ streaming multiprocessors, see Theorem 4.2.1. However, as we shall see through a case study series, this machine parameter has no influence on the comparison of algorithms.

Our model extends both the fork-join concurrency and PRAM models, with an emphasis on parallelism overheads resulting from communication and synchronization.

We sketch below how, in practice, we use this model to tune a program parameter so as to minimize parallelism overheads of programs targeting many-core GPUs. Consider an MCM program $\mathcal{P}$, that is, an algorithm expressed in the MCM model. Assume that a program parameter $s$ (like the number of threads per thread-block running on an SM) can be arbitrarily chosen within some range $S$ while preserving the specifications of $\mathcal{P}$. Let $s_0$ be a particular value of $s$
4.2. A many-core machine model

The model of parallel computations presented in this Chapter aims at capturing communication and synchronization overheads of programs written for modern many-core architectures. One of our objectives is to optimize algorithms by techniques like reducing redundant memory accesses. The reason for this optimization is that, on actual GPUs, global memory latency is approximately 400 to 800 clock cycles. This memory latency, when not properly taken into account, may have a dramatically negative impact on program performance. Another objective of our model is to compare different algorithms targeting implementation on GPUs without taking hardware parameters into account.

As specified in Sections 4.2.1 and 4.2.2, our many-core machine (MCM) model retains many of the characteristics of modern GPU architectures and programming models, like CUDA or OpenCL. However, in order to support algorithm analysis with an emphasis on parallelism overheads, as defined in Section 4.2.3 and 4.2.4, the MCM abstract machines admit a few simplifications and limitations with respect to actual many-core devices.

4.2.1 Characteristics of the abstract many-core machines

Architecture. An MCM abstract machine shown in Figure 4.1 possesses an unbounded number of streaming multiprocessors (SMs), which are all identical. Each SM has a finite number of processing cores and a fixed-size private memory. An MCM machine has a two-level memory that corresponds to an instance $\mathcal{P}_0$ of $\mathcal{P}$, which, in practice, is seen as an initial version of the algorithm to be optimized.

We consider the ratios of work, span and parallelism overhead given by $W_{\mathcal{P}_0}/W_{\mathcal{P}}$, $S_{\mathcal{P}_0}/S_{\mathcal{P}}$ and $O_{\mathcal{P}_0}/O_{\mathcal{P}}$. Assume that, when $s$ varies within $\mathcal{S}$, the work and span ratios stay within $O(s)$ (in fact, $\Theta(1)$ is often the case), but the ratio of the parallelism overhead reduces by a factor in $\Theta(s)$. Thereby, we determine a value $s_{\min} \in \mathcal{S}$ maximizing the parallelism overhead ratio. Next, we use our version of the Graham-Brent theorem (more precisely, Corollary 4.2.2) to check whether the upper bound for the running time of $\mathcal{P}(s_{\min})$ is less than that of $\mathcal{P}(s_0)$. If this holds, we view $\mathcal{P}(s_{\min})$ as a solution of our problem of algorithm optimization (in terms of parallelism overheads).

To evaluate the benefits of our model, we applied it successfully to six fundamental algorithms in scientific computing, see Sections 4.3 to 4.6. These six algorithms are the Euclidean algorithm, the Cooley & Tukey and Stockham fast Fourier transform algorithms, the plain and FFT-based univariate polynomial multiplication algorithms, and radix sort [103]. The former five algorithms are implemented in CUDA and publicly available with benchmarking scripts from http://www.cumodp.org/

Following the strategy described above for algorithm optimization, our model is used to tune a program parameter in the case of the Euclidean algorithm, the plain multiplication algorithm and radix sort. Next, our model is used to compare the two fast Fourier transform algorithms and then the two univariate polynomial multiplication algorithms. In each case, work, span and parallelism overhead are evaluated so as to obtain running time estimates via our version of the Graham-Brent theorem and then select a proper algorithm.
hierarchy, comprising an unbounded global memory with high latency and low throughput and fixed size private memories with low latency and high throughput.

**Figure 4.1: Overview of an abstract many-core machine**

**Programs.** An MCM program is a directed acyclic graph (DAG), whose vertices are kernels (defined hereafter) and edges indicate serial dependencies; moreover, this DAG is similar to the instruction stream DAGs of the fork-join concurrency model. A kernel is an SIMD (single instruction, multiple data) program capable of branches and decomposed into a number of thread-blocks. Each thread-block is executed by a single SM, and each SM executes a single thread-block at a time. Similar to a CUDA program, an MCM program specifies for each kernel the number of thread-blocks and the number of threads per thread-block. Figure 4.2 depicts the different types of components of an MCM program.

**Figure 4.2: Overview of a many-core machine program**

**Scheduling and synchronization.** At run time, an MCM machine schedules thread-blocks (from the same or different kernels) onto SMs, based on the dependencies specified by the edges of the DAG and the hardware resources required by each thread-block. Threads within a thread-block can cooperate with each other via the private memory of the SM running the thread-block.
4.2. A many-core machine model

Meanwhile, thread-blocks interact with each other via the global memory. In addition, threads within a thread-block are executed physically in parallel by an SM. Moreover, the programmer cannot make any assumptions on the order in which thread-blocks of a given kernel are mapped to the SMs. Hence, an MCM program runs correctly on any fixed number of SMs.

**Memory access policy.** All threads of a given thread-block can access simultaneously any memory cell of the private memory or the global memory: read/write conflicts are handled by the CREW (concurrent read, exclusive write) policy. However, we assume that read/write requests to the global memory by two different thread-blocks cannot be executed simultaneously. In case of simultaneous requests, one thread-block is chosen randomly and served first, then the other is served.

Toward analyzing program performance, we define two *machine parameters*:

- $U$: time (expressed in clock cycles) to transfer one machine word between the global memory and the private memory of any SM; hence, we have $U > 0$, and
- $Z$: size (expressed in machine words) of the private memory of any SM, which sets up an upper bound on several program parameters.

The private memory size $Z$ sets several characteristics and limitations of an SM and, thus, of a thread-block. Indeed, each of the following quantities is at most equal to $Z$: the number of threads of a thread-block and the number of words in a data transfer between the global memory and the private memory. The quantity $1/U$ is a throughput measure and has the following property. If $\alpha$ and $\beta$ are the maximum numbers of words, respectively, read and written to the global memory by one thread of a thread-block $B$, and $\ell$ is the number of threads per thread-block, then the total time $T_D$ spent in data transfer between the global memory and the private memory of an SM executing $B$ satisfies:

$$T_D \leq (\alpha + \beta) U, \text{ if coalesced accesses occur, or } \ell (\alpha + \beta) U, \text{ otherwise.} \tag{4.1}$$

On actual GPU devices, some hardware characteristics may reduce data transfer time, for instance, fast context switching between warps executed by an SM. Other hardware characteristics, like partition camping [102], may increase data transfer time. As an abstract machine, the MCM aims at capturing either the best or the worst scenario for data transfer time of a thread-block, which leads us to Relation (4.1).

Relation (4.1) calls for another comment. One could expect the introduction of a third machine parameter, say $V$, which would be the time to execute one *local operation* (arithmetic operation, read/write in the private memory), such that, if $\sigma$ is the maximum number of local operations performed by one thread of a thread-block $B$, then the total time $T_A$ spent in local operations by an SM executing $B$ would satisfy

$$T_A \leq \sigma V. \tag{4.2}$$

Therefore, for the total running time $T$ of the thread-block $B$, we would have

$$T = T_A + T_D \leq \sigma V + \epsilon (\alpha + \beta) U,$$

where $\epsilon$ is either 1 or $\ell$. Instead of introducing this third machine parameter $V$, we let $V = 1$. Thus, $U$ can be understood as the ratio of the time to transfer a machine word to the time to execute a local operation.
4.2.2 Many-core machine programs

Recall that each MCM program $P$ is a DAG $(K, E)$, called the kernel DAG of $P$, where each node $K \in K$ represents a kernel, and each edge $E \in E$ records the fact that a kernel call must precede another kernel call. In other words, a kernel call can be executed once all its predecessors in the DAG $(K, E)$ have completed their execution.

**Synchronization costs.** Recall that each kernel decomposes into thread-blocks and that all threads within a given kernel execute the same serial program, but with possibly different input data. In addition, all threads within a thread-block are executed physically in parallel by an SM. It follows that the MCM kernel code needs no synchronization statement. Consequently, the only form of synchronization taking place among the threads executing a given thread-block is implied by code divergence [58]. This latter phenomenon can be seen as parallelism overhead. Furthermore, an MCM machine handles code divergence by eliminating the corresponding conditional branches via code replication [106]; thereby, the corresponding cost will be captured by the complexity measures (work, span and parallelism overhead) of the MCM model.

**Scheduling costs.** Since an MCM abstract machine has infinitely many SMs and since the kernel DAG defining an MCM program $P$ is assumed to be known when $P$ starts to execute, scheduling $P$’s kernels onto the SMs can be done in time $O(\Gamma)$, where $\Gamma$ is the total length of $P$’s kernel code. Thus, we neglect those costs in comparison to the costs of data transfer between SMs’ private memories and the global memory. We also note that assuming that the kernel DAG is known when $P$ starts to execute allows us to focus on parallelism overheads resulting from this data transfer.

Extending MCM machines to program DAGs unfolding dynamically at run time and integrating the resulting scheduling costs are a work in progress. This key observation helps understand the complexity measures introduced in Section 4.2.3.

**Thread-block DAG.** Since each kernel of the program $P$ decomposes into finitely many thread-blocks, we map $P$ to a second graph, called the thread-block DAG of $P$, whose vertex set $B(P)$ consists of all thread-blocks of the kernels of $P$, such that $(B_1, B_2)$ is an edge if $B_1$ is a thread-block of a kernel preceding the kernel of the thread-block $B_2$ in $P$. This second graph defines two important quantities:

- $N(P)$: number of vertices in the thread-block DAG of $P$, and
- $L(P)$: critical path length (where length of a path is the number of edges in that path) in the thread-block DAG of $P$.

4.2.3 Complexity measures for the many-core machine model

Consider an MCM program $P$ given by its kernel DAG $(K, E)$. Let $K \in K$ be a kernel of $P$ and $B$ be a thread-block of $K$. We define the work of $B$, denoted by $W(B)$, as the total number of local operations performed by all threads of $B$. We define the span of $B$, denoted by $S(B)$, as the maximum number of local operations performed by a thread of $B$. As before, let $\alpha$ and $\beta$ be the maximum numbers of words read and written (from the global memory) by a thread of $B$, and $\ell$ be the number of threads per thread-block. Then, we define the overhead of $B$, denoted
4.2. A many-core machine model

by O(B), as

\[(\alpha + \beta) U, \text{if memory accesses can be coalesced or}
\]
\[\ell (\alpha + \beta) U, \text{otherwise.}\]  

Next, the work (resp. overhead) W(K) (resp. O(K)) of the kernel K is the sum of the works (resp. overheads) of its thread-blocks, while the span S(K) of the kernel K is the maximum of the spans of its thread-blocks. We consider now the entire program P. The work W(P) of P is defined as the total work of all its kernels.

\[W(P) = \sum_{K \in \mathcal{K}} W(K).\]

Regarding the graph \((\mathcal{K}, \mathcal{E})\) as a weighted-vertex graph, where the weight of a vertex \(K \in \mathcal{K}\) is its span \(S(K)\), we define the weight \(S(\gamma)\) of any path \(\gamma\) from the first executing kernel to a terminal kernel (that is, a kernel with no successors in \(P\)) as \(S(\gamma) = \sum_{k \in \gamma} S(K)\). Then, we define the span \(S(P)\) of \(P\) as

\[S(P) = \max_{\gamma} S(\gamma),\]

the longest path, counting the weight (span) of each vertex (kernel), in the kernel DAG. Finally, we define the overhead \(O(P)\) of the program \(P\) as the total overhead of all its kernels.

\[O(P) = \sum_{K \in \mathcal{K}} O(K).\]

Observe that, according to Mirsky’s theorem [90], the number \(\pi\) of parallel steps in \(P\) (which form a partition of \(\mathcal{K}\) into anti-chains in the DAG \((\mathcal{K}, \mathcal{E})\) regarded as a partially ordered set) is greater or equal to the maximum length of a path in \((\mathcal{K}, \mathcal{E})\) from the first executing kernel to a terminal kernel.

4.2.4 A Graham-Brent theorem with parallelism overhead

**Theorem 4.2.1** The running time \(T_P\) of the program \(P\) executed on \(P\) SMs satisfies the inequality:

\[T_P \leq \frac{(N(P)}{\lambda} + L(P)C(P),\]

where \(C(P) = \max_{B \in \mathcal{B}(P)} (S(B) + O(B))\).

The proof is similar to that of the original result [11, 53], while the proof of the following corollary follows from Theorem 4.2.1 and from the fact that costs of scheduling thread-blocks onto SMs are neglected.

**Corollary 4.2.2** Let \(K\) be the maximum number of thread-blocks along an anti-chain of the thread-block DAG of \(P\). Then the running time \(T_P\) of the program \(P\) satisfies:

\[T_P \leq \frac{(N(P)}{K} + L(P)C(P).\]  

(4.4)

The analysis of the running time estimate is based on the thread-block DAG. In Figure 4.3, we show an example of a thread-block DAG with the assumption that each kernel executes in a coalesced manner.

As we shall see in Sections 4.3 through 4.5, Corollary 4.2.2 allows us to estimate the running time of an MCM program as a function of the number \(\ell\) of threads per thread-block, the single machine parameter \(U\) and the thread-block DAG of \(P\). However, the dependence on the machine parameter \(Z\) (the size of a private memory) is only through inequalities specifying upper bounds for \(\ell\). In addition, in each of the case studies, there is no need to make any assumptions (like inequality constraints) on the machine parameter \(U\).
4.3 The Euclidean algorithm

Our first application of the MCM model deals with a multithreaded algorithm for computing the greatest common divisor (GCD) of two univariate polynomials. To specify notations, let $\mathbb{K}$ be a field of coefficients (like the finite field $\mathbb{Z}/p\mathbb{Z}$ of prime characteristic $p$) and $\mathbb{K}[X]$ be the set of all univariate polynomials with coefficients in $\mathbb{K}$.

Our approach is based on the Euclidean algorithm, which can be reviewed in Chapter 4 of [74]. Given a positive integer $s$, we proceed by repeatedly calling a subroutine, see Algorithm 1. This subroutine takes as input a pair $(a, b)$ of polynomials in $\mathbb{K}[X]$ with $\deg(a) \geq \deg(b) > 0$ and returns another pair $(a', b')$ of polynomials in $\mathbb{K}[X]$, such that $\gcd(a, b) = \gcd(a', b')$ and, either $b' = 0$ (in which case we have $\gcd(a, b) = a'$), or we have $\deg(a') + \deg(b') \leq \deg(a) + \deg(b) - s$. We will take advantage of our MCM model to tune the program parameter $s$ in order to obtain an optimized multithreaded version of the Euclidean algorithm.

Let $n$ and $m$ be positive integers such that $\deg(a) = n - 1$ and $\deg(b) = m - 1$, assuming $n \geq m$. We use a dense representation for encoding the polynomials $a$ and $b$. Thus, $n$ and $m$ are the number of coefficients (zero or not) of $a$ and $b$, respectively. Algorithm 1 uses two one-dimensional arrays of size $n$ and $m$, respectively, to represent $a$ and $b$, where the coefficient of the term of $a$ (resp. $b$) in degree $i$ is stored in $a[i]$ (resp. $b[i]$). Observe that Algorithm 1 updates $a$ and $b$. We denote by $d_a$ and $d_b$ the degrees of these updated polynomials, while we reserve $n$ and $m$ for the initial values of the sizes of $a$ and $b$.

Algorithm 1 is implemented as a kernel, which proceeds as follows. While $d_a \geq d_b$ holds, the polynomial $a$ is replaced by $a - cX^d b$, where $c$ is the leading coefficient of $a$ divided by that of $b$ and $d = d_a - d_b$. If this process reduces $a$ to zero, or decreases $d_a$ by $s$, in case $d_a \geq d_b$ still holds, then the updated pair $(a, b)$ is returned. If the condition $d_a \geq d_b$ becomes false before $d_a$ can decrease by $s$, then the roles of $a$ and $b$ are exchanged; thus, $a$ becomes the divisor.

Hence, each call to this kernel either computes the GCD of its input polynomials, or makes the sum of their degrees decreased at least by $s$. Since this may require performing $s$ division
steps (using either \(a\) or \(b\) as divisor), each thread-block must compute the leading coefficients of \(a\) and \(b\) at each division step.

From the specifications of Algorithm 1 it follows that computing \(\gcd(a, b)\) requires at most \(\left\lceil \frac{n+m}{s} \right\rceil\) kernel calls. With the goal of optimizing the use of computing resources and sharpening the analysis of our multithreaded GCD computation, we observe that these kernel calls can be

### Algorithm 1: \(\text{OptGcdKer}(a, b, s, da, db)\)

**Input:** \(a, b \in \mathbb{K}[X]\), an integer \(s \geq 1\) and \(da, db\) store the current degrees of \(a\) and \(b\) respectively.

**Output:** Either one of \(a, b\) was set to \(\gcd(a, b)\) (and the other to 0), or \(da + db\) reduced at least by \(s\).

1. Let \(Alc, Blc, A, B\) be arrays of size \(s, s, \ell + s, \ell + s\) respectively with coefficients in \(\mathbb{K}\), allocated on the private memory of the device; local integers \(u = v = w = e = 0\);
2. \( j = \text{blockID-blockDim + threadID}; t = \text{threadID}; /* \text{copying from global memory} */
3. \(\text{if } t < s \text{ then}
   4. \quad Alc[t] = a[da-t]; \quad Blc[t] = b[db-t];
4. \(\text{if } t \geq s \text{ then}
   5. \quad A[t-s] = a[da-s \text{ blockID}-t]; \quad B[t-s] = b[db-s \text{ blockID}-t];
   /* \text{computing next remainders} */
6. \(\text{for } (k = 0; k < s; k = k + 1) \text{ do}
   7. \quad \text{if } (da \geq db \text{ and } db \geq 0) \text{ then}
      8. \quad \quad \text{if } (u + t < s) \text{ and } (v + t < s) \text{ then}
         9. \quad \quad \quad Alc[u+t] := Blc[v+t]-Alc[u]*Blc[v]^{-1};
      10. \quad \quad \text{if } (u + t \geq s) \text{ and } (v + t \geq s) \text{ then}
            11. \quad \quad \quad A[w+t-s] := B[e+t-s]-Alc[u]*Blc[v]^{-1};
      12. \quad \quad \quad \text{if } t == 0 \text{ then}
            13. \quad \quad \quad \quad \text{while } Alc[u] = 0 \text{ do}
                14. \quad \quad \quad \quad \quad u = u + 1; \quad w = w + 1; \quad da = da - 1;
      16. \quad \quad \text{if } (db \geq da) \text{ and } (da \geq 0) \text{ then}
            17. \quad \quad \quad \text{if } (u + t < s) \text{ and } (v + t < s) \text{ then}
                18. \quad \quad \quad \quad Blc[v+t] := Alc[u+t]*Blc[v]-Alc[u]^{-1};
            19. \quad \quad \quad \text{if } (u + t \geq s) \text{ and } (v + t \geq s) \text{ then}
                20. \quad \quad \quad \quad B[e+t-s] := A[w+t-s]*Blc[v]-Alc[u]^{-1};
            21. \quad \quad \quad \quad \text{if } t == 0 \text{ then}
                22. \quad \quad \quad \quad \quad \text{while } Blc[v] = 0 \text{ do}
                    23. \quad \quad \quad \quad \quad \quad v = v + 1; \quad e = e + 1; \quad db = db - 1;
      24. \quad \quad \text{if } t \geq s \text{ then}
            25. \quad \quad \quad /* \text{writing to global memory} */
                26. \quad \quad \quad a[da-s \text{ blockID}-t] = A[t-s];
            27. \quad \quad \quad b[db-s \text{ blockID}-t] = B[t-s];
      28. \quad \quad \text{if } j = \min(da, db) \text{ then}
            29. \quad \quad \quad \text{Update } da, db \text{ with the new degrees of } a \text{ and } b;\)
separated into two computational phases, which we call ping-pong and ping-pong. During the ping-pong phase, the inequality $d_a \geq d_b$ remains true, and thus this phase amounts to at most $\lceil \frac{n-m}{s} \rceil$ kernel calls. The subsequent kernel calls form the ping-pong phase, during which the role of the divisor alternates between $a$ and $b$; thus, there are at most $\lceil \frac{2m}{s} \rceil$ kernel calls in this second phase.

Denoting by $\ell$ the number of threads per thread-block, we observe that each kernel call requires $\lceil \frac{m}{2} \rceil$ thread-blocks in the ping-pong phase. During the ping-pong phase, for a kernel called on polynomials with current degrees $d_a$ and $d_b$, the number of required thread-blocks becomes $\min(\lceil \frac{d_a+1}{\ell} \rceil, \lceil \frac{d_b+1}{\ell} \rceil)$.

After executing a kernel call in the ping-pong phase, the $s$ largest-degree coefficients of $a$ have been set to zero; meanwhile, $b$ is left unchanged. Each thread-block updates $\ell$ other coefficients of $a$. After executing a kernel call in the ping-pong phase, $s$ largest-degree coefficients among $a$ and $b$ have been set to zero; meanwhile, each thread-block has updated $2 \ell$ other coefficients of $a$ and $b$. See Figure 4.4 for an illustration of both scenarios.

To ensure that every kernel call (in either ping-pong or ping-pong phase) can perform (at most) $s$ division steps correctly, each thread-block reads the $s$ largest-degree coefficients from both $a$ and $b$, as well as $\ell + s$ other consecutive coefficients from both $a$ and $b$. Thereby, $4s + 2\ell$ coefficients must fit into the private memory of each streaming multiprocessor; hence, we have $4s + 2\ell \leq Z$.

We note that two consecutive thread-blocks have $s$ common coefficients of both $a$ and $b$. Moreover, all thread-blocks of a given kernel call read the $s$ largest-degree coefficient of both $a$ and $b$. Despite this duplication of data access and the corresponding duplication of work, this
multithreaded GCD algorithm is practically effective and is optimized for a relatively large $s$, as we shall see.

The work, span and parallelism overhead are given\(^1\) respectively, by

\[
W_s = 3 m^2 + 6 n m + 3 s + \frac{3(5 m + 4 n s + 14 m + n s^2 + 6 s)}{8 \ell},
\]

\[
S_s = 3 n + 3 m \quad \text{and}
\]

\[
O_s = \frac{4 n m U (2 n + m + s)}{s \ell}.
\]

To determine a value range for $s$ that minimizes the parallelism overhead of our multithreaded algorithm, we choose $s = 1$ as a starting point. Let $W_1$, $S_1$ and $O_1$ be the work, span and parallelism overhead at $s = 1$. The work ratio $W_1/W_s$ is asymptotically equivalent to

\[
\left(\frac{16 \ell + 8}{16 \ell + 4 s + 4} + \frac{19}{(8 \ell + 5 s + 14) m}\right)
\]

when $n$ (and thus $m$) escapes to infinity. The span ratio $S_1/S_s$ is 1, and the parallelism overhead ratio $O_1/O_s$ is

\[
\frac{(2 n + m + 1) s}{2 n + m + s}.
\]

We observe that when $s \in \Theta(\ell)$, the work is increased only by a constant factor, while the parallelism overhead is reduced by a factor in $\Theta(s)$.

Hence, choosing $s \in \Theta(\ell)$ seems a good choice. To verify this, we apply Corollary 4.2.2. One can easily check that the quantities characterizing the thread-block DAG of the computation are

\[
N_s = \frac{2 n m + m^2 + m s}{2 s \ell}, \quad L_s = \frac{n + m}{s} \quad \text{and} \quad C_s = 3 s + 8 U.
\]

Then, applying Corollary 4.2.2 we estimate the running time on $\Theta(\frac{m}{\ell})$ SMs as

\[
T_s = \frac{4 n + 3 m + s}{2 s} (3 s + 8 U).
\]

Denoting by $T_1$ the estimated running time when $s = 1$, the running time ratio $R = T_1/T_s$ on $\Theta(\frac{m}{\ell})$ SMs is given by

\[
\left(\frac{4 n + 3 m + 1}{4 n + 3 m + s}\right) \frac{(3 + 8 U) s}{(3 s + 8 U)}.
\]

When $n$ and $m$ escape to infinity, the latter ratio asymptotically becomes

\[
\frac{(3 + 8 U) s}{3 s + 8 U},
\]

which is greater than 1 if and only if $s > 1$. Thus, the algorithm with $s = \Theta(\ell)$ performs better than that with $s = 1$. Figure 4.5 shows the experimental results with $s = \ell = 256$ and $s = 1$ on an NVIDIA Kepler architecture, which confirms our theoretical analysis.

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\(^1\)See the detailed analysis in the form of executable MAPLE worksheets of all applications: [http://www.csd.uwo.ca/~nxie6/projects/mcm/](http://www.csd.uwo.ca/~nxie6/projects/mcm/) or the PDF version of these worksheets in Appendix B.
Figure 4.5: Running time on GeForce GTX 670 of our multithreaded Euclidean algorithm for univariate polynomials of sizes $n$ and $m$ over $\mathbb{Z}/p\mathbb{Z}$, where $p$ is a 30-bit prime, whereas the program parameter takes values $s = 1$ and $s = 256$.

4.4 Fast Fourier Transform

Let $f$ be a univariate polynomial over the prime field of characteristic $p$, namely $\mathbb{F}_p := \mathbb{Z}/p\mathbb{Z}$, where $p$ is a prime number greater than 2. Let $n$ be the smallest power of 2, such that the degree of $f$ is less than $n$, that is, $n = \min\{2^e \mid \deg(f) < 2^e \text{ and } e \in \mathbb{N}\}$. We assume that $n$ divides $p - 1$, which guarantees that the field $\mathbb{F}_p$ admits an $n$-th primitive root of unity. Hence, let $\omega \in \mathbb{F}_p$ such that $\omega^n = 1$ holds, while for all $0 \leq i < n$, we have $\omega^i \neq 1$. The $n$-point Discrete Fourier Transform (DFT) at $\omega$ is the linear map from the $\mathbb{F}_p$-vector space $\mathbb{F}_p^n$ to itself, defined by $x \mapsto \text{DFT}_n x$ with the $n$-th DFT matrix given by

$$\text{DFT}_n = [\omega^{ij}]_{0 \leq i, j < n}. \quad (4.5)$$

Since $\omega$ is an $n$-th primitive root of unity, this map is invertible and its inverse is $1/n$ times the DFT at $\omega'$, where $\omega'$ is the inverse of $\omega$ in $\mathbb{F}_p$.

A fast Fourier transform (FFT) is an algorithm to compute the DFT and its inverse. Two algorithms of the most commonly used FFTs are that of Cooley & Tukey [33] and that of Stockham [109]. Before reviewing and analyzing those algorithms on an MCM machine, we introduce a few notations. Given an input vector $v$ of length $nm$, we define the stride permutation $L_{nm}$ as the permutation that maps the entry of $v$ at position $in + j$ to position $jm + i$ for $0 \leq j \leq (m-1)$ and $0 \leq i \leq (n-1)$. We denote by $I_x$ the identity matrix of order $x$. The symbols $\oplus$ and $\otimes$ are used for the direct sum and Kronecker product\footnote{Matrix multiplications are not commutative. Throughout, the product $\prod_{i=1}^s M_i$ stands for $M_1 M_2 \cdots M_s$, while $\prod_{i=1}^s M_i$ means $M_s M_{s-1} \cdots M_1$.} of matrices. We define the twiddle matrix $D_{x,y}$ as the diagonal matrix of order $xy$ given by $\oplus_{j=0}^{x-1} \text{diag}(1, w^j, \ldots, w^{j(y-1)})$.

4.4.1 Cooley & Tukey algorithm

Let $k = \log_2(n)$ and fix $r$ for $0 < r < k$. Define $m = 2^r$ and thus $m$ divides $n$. The algorithm of Cooley & Tukey is based on the following factorization\footnote{https://en.wikipedia.org/wiki/Kronecker_product} of the matrix DFT$_n$

$$\text{DFT}_n = M_{k,r} M'_{k,r} M''_{k,r}, \quad (4.6)$$

Let $s = 1$ and $s = 256$.
where $M_{k,r} = \prod_{i=0}^{k-r-1} (I_2 \otimes \text{DFT}_2 \otimes I_{2^i})(I_2 \otimes D_{2,2^{k-i-1}}), M'_{k,r} = I_{2^r} \otimes \text{DFT}_m$ and $M''_{k,r} = \prod_{i=k-r-1}^{0} I_2 \otimes L_2^{2^i}$. Each of $M_{k,r}, M'_{k,r}, M''_{k,r}$ is a structured square matrix of order $n$ representing a computational step of the algorithm.

As we shall see, the multiplications by the matrices $M''_{k,r}$ and $M'_{k,r}$ are difficult to implement on a GPU-like architecture. Each factor of the matrix $M''_{k,r}$ is a Kronecker product of the form $I_2 \otimes L_2^{2^i}$, that is, a diagonal matrix, which has order $2^i$ and is a permutation matrix. One may assume that, for a kernel implementing the multiplication by $I_2 \otimes L_2^{2^i}$, all thread-blocks perform coalesced reads. But, for a small enough $i$, several non-coalesced memory accesses to the global memory may occur when writing back within one thread-block. Turning our attention now to the multiplication by $I_2 \otimes D_{2,2^{k-i}}$ operates on a sub-vector of length $2^{k-i+1}$, the powers $\{1, \omega^{2^i}, (\omega^{2^i})^2, \ldots, (\omega^{2^i})^{2^{k-i}-1}\}$ need to be computed. A thread-block operating on a sub-vector of size $x$ may need to compute $x$ of those consecutive powers, which can be done in time $O(\log_2(x))$. Finally, multiplication by $M'_{k,r}$ causes no difficulty as long as $m$ is large enough so as to avoid non-coalesced memory accesses. In our implementation, $m = 16$ is appropriate.

Let $\ell$ be the number of threads per thread-block, then $M_{k,r}$ and $M'_{k,r}$ are computed by $\log_2(n) - \log_2(m)$ kernel calls, respectively, requiring $\frac{n}{\ell}$ thread-blocks per kernel, and $M''_{k,r}$ is computed by one kernel with $\frac{n}{m\ell}$ thread-blocks.

We compute the work, span and parallelism overhead, respectively, as

\begin{align*}
W_{ct} &= n (34 \log_2(n) \log_2(\ell) + 47 \log_2(n) + 32 m + 9 - 34 \log_2(m) \log_2(\ell) - 47 \log_2(m)), \\
S_{ct} &= 34 \log_2(n) \log_2(\ell) + 47 \log_2(n) + 32 m \log_2(m) + 30 m + 11 - 34 \log_2(\ell) \log_2(m) \\
&\quad - 79 \log_2(m) \quad \text{and} \\
O_{ct} &= \frac{2nU}{\ell} (4 \log_2(n) + \ell \log_2(\ell) + 1 - \log_2(\ell) - 4 \log_2(m)).
\end{align*}

To apply Corollary 4.2.2, one can easily check that those three quantities characterizing the thread-block DAG are

\begin{align*}
N_{ct} &= \frac{n}{m\ell} (2 m \log_2(n) + 1 - 2 m \log_2(m)), \\
L_{ct} &= 2 \log_2(n) - 2 \log_2(m) + 1 \quad \text{and} \\
C_{ct} &= 2 U \ell + 34 \log_2(\ell) + 2 U + 27.
\end{align*}

Thus, we estimate that the running time on $\Theta(\frac{n}{\ell})$ streaming multiprocessors is

\[ T_{ct} = (4 \log_2(n) + 1 + \frac{1}{m} - 4 \log_2(m)) (2 U \ell + 34 \log_2(\ell) + 2 U + 27). \]

### 4.4.2 Stockholm algorithm

The Stockholm algorithm is based on the following factorization of the matrix $\text{DFT}_n$

\[ \text{DFT}_n = \prod_{i=0}^{k-1} (\text{DFT}_2 \otimes I_{2^i})(D_{2,2^{k-i-1}} \otimes I_2)(L_2^{2^i} \otimes I_2), \]

where $k = \log_2(n)$ as before. For each $0 \leq i < k$, one performs three matrix-vector multiplications:

\[ A_{k,i} : x \mapsto (L_2^{2^i} \otimes I_2) x, \quad A'_{k,i} : x \mapsto (D_{2,2^{k-i-1}} \otimes I_2) x, \quad \text{and} \quad A''_{k,i} : x \mapsto (\text{DFT}_2 \otimes I_{2^i}) x. \]
Thus, the Stockham algorithm can be implemented as \( \log_2(n) \) calls to a kernel performing successively the matrix-vector multiplications defined by \( A_{k,i} \), \( A'_{k,i} \) and \( A''_{k,i} \). Each of the corresponding matrices has a structure permitting coalesced read/write memory accesses. Indeed, for each of these matrices, all coefficients are null except along a few segments parallel to the diagonal. See [92] for details.

Let \( \ell \) be the number of threads per thread-block; thus, each kernel requires \( \frac{n}{\ell} \) thread-blocks. We compute the work, span and parallelism overhead, respectively, as

\[
W_{sh} = n \cdot 43 \log_2(n) + \frac{n}{4\ell} + 12 \ell + 1 - 30n, \\
S_{sh} = 43 \log_2(n) + 16 \log_2(\ell) + 3 \quad \text{and} \\
O_{sh} = \frac{5nU \log_2(n)}{\ell} + \frac{5nU}{4\ell^2}.
\]

Applying Corollary 4.2.2, the quantities characterizing the thread-block DAG are

\[
N_{sh} = n \cdot \frac{(8 \log_2(n) - 5)}{4 \ell}, \quad L_{sh} = 3 \log_2(n) + 1 \quad \text{and} \quad C_{sh} = 8 \log_2(\ell) + 4U + 17.
\]

Hence, the running time estimate on \( \Theta \left( \frac{n}{\ell} \right) \) SMs is

\[
T_{sh} = \log_2(n) \cdot (40 \log_2(\ell) + 20U + 85) - 2 \log_2(\ell) - U - \frac{17}{4}.
\]

### 4.4.3 Comparison of running time estimates

The work ratio \( W_{ct}/W_{sh} \) is asymptotically equivalent to

\[
\frac{4n \cdot (47 \log_2(n) \ell + 34 \log_2(n) \ell \log_2(\ell))}{172n \log_2(n) \ell + n + 48\ell^2}.
\]

when \( n \) escapes to infinity. Since \( \ell \in O(Z) \), the quantity \( \ell \) is bounded over on a given machine. Thus, the work ratio is asymptotically equivalent to \( \log_2(\ell) \) when \( n \) escapes to infinity, while the span ratio \( S_{ct}/S_{sh} \) is asymptotically equivalent to

\[
\frac{34 \log_2(n) \log_2(\ell) + 47 \log_2(n)}{43 \log_2(n) + 16 \log_2(\ell)},
\]

which is also in \( \Theta(\log_2(\ell)) \). In other words, both the work and span of the algorithm of Cooley & Tukey are increased by \( \Theta(\log_2(\ell)) \) factor w.r.t their counterparts in the Stockham algorithm. Next, we compute the parallelism overhead ratio \( O_{ct}/O_{sh} \) as

\[
\frac{8n \cdot (4 \log_2(n) + \ell \log_2(\ell) - \log_2(\ell) - 15)}{20n \log_2(n) + 5n - 4\ell}.
\]

Applying Corollary 4.2.2, we obtain the running time ratio \( R = T_{ct}/T_{sh} \) on \( \Theta \left( \frac{n}{\ell} \right) \) SMs as

\[
R \sim \frac{\log_2(n)(2U \ell + 34 \log_2(\ell) + 2U)}{5 \log_2(n)(U + 2 \log_2(\ell))},
\]

when \( n \) escapes to infinity. This latter ratio is greater than 1 if and only if \( \ell > 1 \).

Hence, the Stockham algorithm outperforms the Cooley & Tukey algorithm on an MCM machine. Table 4.1 shows the experimental results comparing the Cooley & Tukey and Stockham algorithms with \( \ell = 128 \) on an NVIDIA Kepler architecture. The experimentation confirms our theoretical analysis.
4.5 Polynomial multiplication

Multithreaded algorithms for polynomial multiplication are our third application of the MCM model. As in Section 4.3, we denote by \(a\) and \(b\) two univariate polynomials with coefficients in the prime field \(\mathbb{F}_p\), and we write their degrees \(\text{deg}(a) = n - 1\) and \(\text{deg}(b) = m - 1\), for two positive integers \(n \geq m\). We compute the product \(f = a \times b\) in two ways: plain multiplication and FFT-based multiplication. We describe the plain multiplication approach in Section 4.5.1; moreover, we analyze it so as to tune a program parameter \(s\) and thus obtain an optimized algorithm. In Section 4.5.2, we analyze an FFT-based multiplication algorithm that uses the Stockham FFT algorithm. Finally, we compare the plain and FFT-based multiplication algorithms in Section 4.5.3 via the MCM model and experimentally.

4.5.1 Plain multiplication

Our first multithreaded polynomial multiplication algorithm is based on the well-known long multiplication\(^4\) and consists of two phases. During the multiplication phase, every coefficient of \(a\) is multiplied with every coefficient of \(b\), and the resulting coefficient products are accumulated in an auxiliary array \(M\). Then, during the addition phase, these accumulated products are added together to form the polynomial \(f\). The top level algorithm shown in Algorithm 2 performs the multiplication phase once via Algorithm 3, and the addition phase by repeated calls to Algorithm 4. We consider as a program parameter the number \(s > 0\) of coefficients that each thread writes back to the global memory at the end of each (multiplication or addition) phase.

We denote by \(\ell\) the number of threads per thread-block. In the multiplication phase, each thread-block reads \(s\ell + s - 1\) coefficients of \(a\) and \(s\) coefficients of \(b\), and then computes \(\ell s^2\) products followed by \(\ell s (s - 1)\) additions. Thus, each thread-block contributes \(s\ell\) partial sums to a two-dimensional array \(M\), whose format is \(x \cdot y\) with \(x = \frac{m}{s}\) and \(y = n + s - 1\). This multiplication phase, illustrated by Figure 4.6, loads \(s\ell + s - 1\) coefficients of \(a\) to guarantee the correctness of the results in \(M\). Thereby, \(2s\ell + 2s - 1\) coefficients must fit into the private memory, that is, \(2s\ell + 2s - 1 \leq Z\), and the kernel requires \(\frac{xy}{\ell s} = \frac{(n + s - 1)m}{\ell s^2}\) thread-blocks.

In the addition phase, the \(x\) rows of the auxiliary array \(M\) are added pairwise in \(\log_2(x)\) parallel steps. After each step, the number of rows in \(M\) is reduced by half, until we obtain

\(^4\text{http://en.wikipedia.org/wiki/Multiplication_algorithm}\)

Table 4.1: Running time (in secs) of the Cooley & Tukey and Stockham FFT algorithms with the input size \(n\) on GeForce GTX 670

<table>
<thead>
<tr>
<th>(n)</th>
<th>Cooley &amp; Tukey</th>
<th>Stockham</th>
</tr>
</thead>
<tbody>
<tr>
<td>(2^{14})</td>
<td>0.583296</td>
<td>0.666496</td>
</tr>
<tr>
<td>(2^{15})</td>
<td>0.826784</td>
<td>0.7624</td>
</tr>
<tr>
<td>(2^{16})</td>
<td>1.19542</td>
<td>0.929632</td>
</tr>
<tr>
<td>(2^{17})</td>
<td>2.07514</td>
<td>1.24928</td>
</tr>
<tr>
<td>(2^{18})</td>
<td>4.66762</td>
<td>1.86458</td>
</tr>
<tr>
<td>(2^{19})</td>
<td>9.11498</td>
<td>3.04365</td>
</tr>
<tr>
<td>(2^{20})</td>
<td>16.8699</td>
<td>5.38781</td>
</tr>
</tbody>
</table>
only one row, that is, \( f = a \times b \). To be more specific, at a parallel step \( k \) \( (0 \leq k < \log_2(n)) \), adding rows \( i \) and \( j \) (for \( i < j \)) as shown in Figure 4.7, the kernel requires \( \frac{xy}{2^{m+1}r_1} = \frac{(n+1)x}{2^{m+1}r_1n^2} \) thread-blocks, while each thread-block loads \( s \ell \) elements of \( M[i] \) and \( M[j] \), respectively, and then adds \( M[j] \) to \( M[i] \).

We compute the work, span and parallelism overhead of the plain multiplication with an
4.5. Polynomial multiplication

**Algorithm 4: AddKer**($M, f, y, s, x, i$)

**Input:** $M, f, y, s, x, i$ are positive integers.

1. $j = \text{blockID} \cdot \text{blockDim} + \text{threadID};$
2. $h = s \cdot j \mod y;$
3. $k = 2^i - 1 + 2^{i+1} \lfloor s \cdot j/y \rfloor;$
4. if $h < 2^i \cdot s$ then
5. for $(e = 0; e < s; e = e + 1)$ do
6. $f[k \cdot s + h + e] += M[k \cdot x + h + e];$
7. else
8. for $(e = 0; e < s; e = e + 1)$ do
9. $M[(k + 2^i) \cdot x + h - 2^i \cdot s + e] += M[k \cdot x + h + e];$

![Figure 4.6: Multiplication phase: illustration of a thread-block reading coefficients from $a, b$ and writing to the auxiliary array $M$](image)

![Figure 4.7: Addition phase: illustration of a thread-block reading and writing to the auxiliary array $M$](image)

We also obtain the quantities characterizing the thread-block DAG that are required in order to apply Corollary 4.2.2:

$$W_s = (2m - \frac{1}{2})(n + s - 1), \quad S_s = 2s^2 + s \log_2 \frac{m}{s} - s \quad \text{and} \quad O_s = \frac{(n + s - 1)(5ms + 2m - 3s^2)}{s^2 \ell} U.$$
We set \( s = 1 \) in the above two phases, and denote its work, span and parallelism overhead as \( W_1, S_1 \) and \( O_1 \), respectively. The work ratio
\[
\frac{W_1}{W_s} = \frac{n}{n + s - 1}
\]
is asymptotically constant as \( n \) escapes to infinity. The span ratio
\[
\frac{S_1}{S_s} = \frac{\log_2(m) + 1}{s(\log_2(m/s) + 2s - 1)}
\]
shows that \( S_s \) grows asymptotically with \( s \). The parallelism overhead ratio is
\[
\frac{O_1}{O_s} = \frac{n s^2 (7m - 3)}{(n + s - 1)(5ms + 2m - 3s^2)}.
\]
We observe that, as \( n \) and \( m \) escape to infinity, this latter ratio is asymptotically equivalent to \( s \). Applying Corollary 4.2.2, the estimated running time on \( \Theta(\frac{(n+s-1)m}{s^2}) \) SMs is
\[
T_s = \left( \frac{2m - s}{m} + \log_2\left(\frac{m}{s}\right) + 1 \right) (2U s + 2s^2 + 2U - s).
\]
Let \( R \) be the ratio of the running time estimate between the algorithm with \( s = 1 \) and that for an arbitrary \( s \), we obtain
\[
R = \frac{(m \log_2(m) + 3m - 1)(1 + 4U)}{(m \log_2\left(\frac{m}{s}\right) + 3m - s)(2U s + 2U + 2s^2 - s)}.
\]
which is asymptotically equivalent to
\[
\frac{2U \log_2(m)}{s(s + U) \log_2(m/s)}.
\]
This latter ratio is smaller than 1 for \( s > 1 \). In other words, increasing \( s \) worsens the algorithm performance. In practice, as shown in Figure 4.8, setting \( s = 4 \) (where \( \ell = 256 \)) performs best, while with larger \( s \), the running time becomes slower. This practical observation is coherent with our theoretical analysis.

### 4.5.2 FFT-based multiplication

Let \( \omega \) be an \( n \)-th root primitive of unity \( w \), as defined in Section 4.4. We assume that the successive powers \( \{1, w, w^2, \ldots, w^{n-1}\} \) are pre-computed via a parallel prefix sum. The product \( f = a \times b \) is computed as follows:

1. FFT computations: \( a' = \text{DFT}_w(a) \) and \( b' = \text{DFT}_w(b) \);
2. Point-wise multiplication: \( f' = a' \times b' \);
3. FFT computation: \( f'' = \text{DFT}_{w^{-1}}(f') \);
4. Scale the vector: \( f = \frac{1}{n} f'' \) and return \( f \);
4.5. Polynomial multiplication

A kernel for computing the Stockham FFT algorithm has been described in Section 4.4. Two other kernels perform, respectively, the point-wise multiplication and vector scaling.

Let \( \ell \) be the number of threads per thread-block. The analysis of the Stockham FFT algorithm is described in Section 4.4.3. To compute the point-wise multiplication, the kernel requires \( n \ell \) thread-blocks with each thread reading two data items and writing one data item back to the global memory. The final kernel, for vector scaling, also requires \( n \ell \) thread-blocks with each thread moving one data item between the global memory and the private memory. Hence, data movement in each kernel call is coalesced.

We compute the work, span and parallelism overhead of the overall FFT-based polynomial multiplication, respectively, as

\[
W_{fft} = 129n \log_2(n) - 94n, \quad S_{fft} = 129 \log_2(n) - 94 \quad \text{and} \quad O_{fft} = \frac{nU(15 \log_2(n) - 4)}{\ell}.
\]

Applying Corollary 4.2.2, we obtain the quantities characterizing the thread-block DAG:

\[
N_{fft} = \frac{12n \log_2(n) - 5}{2\ell}, \quad L_{fft} = 9 \log_2(n) - 4 \quad \text{and} \quad C_{fft} = 4U + 25.
\]

Thus, the running time estimate on \( \Theta\left(\frac{n^2}{\ell}\right) \) SMs is

\[
T_{fft} = (15 \log_2(n) - \frac{13}{2})(4U + 25).
\]

4.5.3 Comparison of running time estimates

Back to the plain multiplication, using \( s = 4 \) obtained from experimental results and setting \( m = n \), we compute

\[
W_{plain} = 2n^2 + \frac{11}{2}n - \frac{3}{2}, \quad S_{plain} = 4 \log_2(n) + 20 \quad \text{and} \quad O_{plain} = \frac{U(n + 3)(11n - 24)}{8\ell}.
\]

We observe that \( W_{plain} / W_{fft} \) is in \( O\left(\frac{n}{\log_2(n)}\right) \), \( S_{plain} / S_{fft} \) is asymptotically constant, and \( O_{plain} / O_{fft} \) is in \( O\left(\frac{n}{\log_2(n)}\right) \). Next, the estimated running time of the plain multiplication on \( \Theta\left(\frac{n(n+3)}{16\ell}\right) \) SMs is

\[
T_{plain} = \left(\frac{2(n - 2)}{n} + \log_2(n) - 1\right)(10U + 28).
\]
The estimated running time ratio $T_{\text{plain}}/T_{\text{fft}}$ is essentially constant when $n$ escapes to infinity, while the plain multiplication performs more work and parallelism overhead.

However, the estimated running time of the plain multiplication on $\Theta(n/\ell)$ SMs, in the case that we have limited resources, is

$$T'_{\text{plain}} = \left(\frac{(n + 3)(n - 2)}{8n} + \log_2(n) - 1\right) (10U + 28),$$

whereas the estimated running time of the FFT-based multiplication is also based on $\Theta(n/\ell)$ SMs. Thus, when $n$ escapes to infinity, the estimated running time ratio $T'_{\text{plain}}/T_{\text{fft}}$ on $\Theta(n/\ell)$ SMs is asymptotically equivalent to

$$\frac{5U(n + 8 \log_2(n))}{240U \log_2(n)},$$

and thus in $\Theta(n)$. Hence, the FFT-based multiplication outperforms the plain multiplication for a large enough $n$.

Figure 4.9 shows the experimental results comparing the plain and FFT-based multiplication algorithms with $\ell = 256$ on an NVIDIA Kepler architecture. We observe that for relatively small $n$, the plain multiplication performs better, but with $n$ growing, the FFT-based multiplication becomes faster. Both phenomena were predicted by our theoretical analysis.

![Figure 4.9: Running time of the plain and FFT-based multiplication algorithms with the input size $n$ on GeForce GTX 670](image)

4.6 Radix sort

In [103], Satish, Harris and Garland present a CUDA implementation of the radix sort algorithm. Assuming that all entries are non-negative integers of bit-size $c$, this CUDA implementation sorts $n$ entries by performing $\frac{c}{s}$ *passes*, where $s$ is a program parameter. In each pass, each thread-block first loads and sorts its tile using $s$ iterations of 1-bit split, and writes back its $2^s$-entry digit histogram and sorted data. Then, it performs a prefix sum over the histogram stored in a column-major order. Finally, each thread-block copies its elements to the correct output position.

Let $\ell$ be the number of threads per thread-block. Following [103], we assume that each thread deals with 4 elements. Then, for each thread-block, $4\ell$ original elements, $4\ell$ sorted elements and a $2^s$-entry digit histogram must fit into the private memory; hence, $8\ell + 2^s \leq Z$. 

Thus, the maximum parallelism overhead per thread-block is $9U$ (loading 4 elements, writing back 4 sorted elements and 1 value of the histogram).

We compute the work, span and parallelism overhead, respectively, as

$$W_s = c\left(\frac{22}{4s}\ell + s\frac{12}{s} + \frac{2^s + 20}{16s}\ell^2 + 1\right)n + \frac{c(16+192\ell)}{16s},$$

$$S_s = c\left(8\log_2(\ell) + \frac{16}{s}\log_2(\ell) + 41 + \frac{24}{s}\right)$$

and

$$O_s = cU\left(\frac{9}{2s}\ell n + \frac{172}{16s}\ell^2 n - \frac{1}{s}\right).$$

We view the case $s = 1$ as a naive radix sort algorithm, with the work, span and parallelism overhead given by $W_1$, $S_1$ and $O_1$, respectively. Letting $n$ escaping to infinity, the work ratio $W_1/W_s$ is asymptotically equivalent to:

$$\frac{W_1}{W_s} \sim \frac{104s\ell^2 + 92s\ell + 2s}{88s\ell^2 + 16\ell^2 + 20s\ell + 4s\ell + 48\ell + 2s^2}.$$

Similarly, the span ratio

$$\frac{S_1}{S_s} = \frac{s(24\log_2(\ell) + 95)}{(8s + 16)\log_2(\ell) + 41s + 54}$$

is asymptotically constant; meanwhile, the parallelism overhead ratio is asymptotically equivalent to:

$$\frac{O_1}{O_s} \sim \frac{s(72\ell + 34)}{72\ell + 172s}.$$

We notice that if $2^s = \Theta(\ell)$ holds, we can reduce the parallelism overhead by a factor of $s$, while increasing the work only by a constant factor. However, with $2^s = \Theta(\ell^2)$, we increase both the work and the parallelism overhead by a non-constant factor. In this latter scenario, we could not optimize the naive algorithm in any sense.

To apply Corollary 4.2.2, we compute the three quantities characterizing the thread-block DAG:

$$N_s = c\left(\frac{1}{2s\ell} + \frac{2s}{8\ell^2}\right)n, \quad L_s = \frac{5c}{s} \quad \text{and} \quad C_s = s(41 + 8\log_2(\ell)) + 12 + 9U.$$

We also denote these three quantities $N_1$, $L_1$ and $C_1$ for the naive algorithm when $s = 1$. Then, the ratio $R$ of the running time estimates on $\Theta(\frac{n}{4\ell})$ SMs between the naive algorithm with $s = 1$ and that for an arbitrary $s$ is:

$$R = \frac{(14\ell + 2)(8\log_2(\ell) + 9U + 53)s}{(14\ell + 2^s)(8s\log_2(\ell) + 41s + 9U + 12)}.$$

We then replace $s$ by $\log_2(\ell)$, since, in this case, we would like to determine whether the estimated overall running time is better or worse. The quotient of the leading terms of $\ell\log_2(\ell)$ in $R$ becomes

$$\frac{112\log_2(\ell) + 126U + 742}{120\log_2(\ell) + 615}.$$

This ratio is greater than 1 for $\ell < 2^{15.75U + 15.875}$, which is realistic since $U > 0$. Therefore, letting $2^s = \Theta(\ell)$, the data transfer overhead can be reduced by a factor of $s$ and lead to an optimized algorithm. This is consistent with the empirical results of [103].
4.7 Conclusion

We have presented a model of multithreaded computation combining the fork-join and SIMD parallelisms, with an emphasis on estimating parallelism overheads of GPU programs, so as to reduce communication and synchronization costs in GPU programs. In practice, our model determines a trade-off among work, span and parallelism overhead by checking the estimated overall running time so as to either (1) tune a program parameter or, (2) compare different algorithms independently of the hardware details.

Several applications illustrate the effectiveness of our model. With the Euclidean algorithm, plain multiplication and radix sort, we determine a range of values for a program parameter in order to optimize the corresponding algorithm in terms of parallelism overheads. Particularly for the Euclidean algorithm, our running time estimate matches that obtained with the systolic VLSI array model [17]. Moreover, our CUDA code in [60] implementing this optimized Euclidean algorithm runs in linear time w.r.t to the input polynomials up to degree 10,000. With FFT algorithms and polynomial multiplication algorithms, our model successfully compares, in each case, two different algorithms to determine the trend of their running time when the input data size grows. In all cases, experimentation validates the model analysis.
Chapter 5

MetaFork-to-CUDA: Generation of Parametric CUDA Kernels

In the past decade, the introduction of low-level heterogeneous programming models, in particular CUDA, has brought supercomputing to the level of the desktop computer. However, these models bring notable challenges, even to expert programmers. Indeed, fully exploiting the power of hardware accelerators with CUDA-like code often requires significant code optimization efforts. While this development can indeed yield high performance, it is desirable for some programmers to avoid the explicit management of device initialization and data transfer between memory levels. To this end, high-level models for accelerator programming have become an important research direction. With these models, programmers only need to annotate their C/C++ code to indicate which code portion is to be executed on the device and how data maps between host and device.

As of today, OpenMP [41, 13, 9] and OpenACC [113, 56] are among the most developed accelerator programming models. Both OpenMP and OpenACC are built on a host-centric execution model. The execution of the program starts on the host and may offload target regions to the device for execution. The device may have a separated memory space or may share memory with the host, so that memory coherence is not guaranteed and must be handled by the programmer. In OpenMP and OpenACC, the division of the work between thread-blocks within a grid and between threads within a thread-block can be expressed in a loose manner or even be ignored. This implies that code optimization techniques can be applied in order to derive efficient CUDA [95, 73]-like code.

In this chapter, we present the accelerator model of MetaFork (introduced in Section 2.4) together with the software framework that allows automatic generation of CUDA code from annotated C/C++ programs. One of the key properties of this CUDA code generator is that it supports the generation of CUDA kernel code where program parameters (like number of threads per thread-block) and machine parameters (like shared memory size) are allowed. These parameters need not to be known at code-generation-time: machine parameters and program parameters can be, respectively, determined and optimized when the generated code is installed on the targeted hardware.

The need for CUDA programs (more precisely, kernels) depending on program parameters and machine parameters is argued in Section 5.1. In Section 5.2, following the work reported in [55], we observe that generating parametric CUDA kernels requires the manipulation of sys-
tems of non-linear polynomial equations and the use of techniques like quantifier elimination (QE). To this end, we take advantage of the RegularChains library of Maple [27] and its QuantifierElimination command, which has been designed to efficiently support the non-linear polynomial systems coming from automatic parallelization.

Section 5.3 reports a preliminary implementation of parametric CUDA kernel code generation by the MetaFork compilation framework extending PPCG [115]. Finally, Section 5.4 gathers experimental data demonstrating the performance of our generated parametric CUDA code. These results show not only that the generation of parametric CUDA kernels helps optimize code independently of the values of the machine parameters of the targeted hardware, but also that automatic generation of parametric CUDA kernels can discover better values for the program parameters than those computed by a tool generating non-parametric CUDA kernels.

This chapter is related to the work reported in [19] and contains joint work with Changbo Chen, Xiaohui Chen and Marc Moreno Maza.

5.1 Optimizing CUDA kernels depending on program parameters

In Chapter 4, we propose a many-core machine (MCM) model for multithreaded computation combining the fork-join and SIMD parallelisms; meanwhile, a driving motivation in this work is to estimate the parallelism overheads (data communication and synchronization costs) of GPU programs. In practice, the MCM model determines a trade-off among work, span and parallelism overhead by checking the estimated overall running time so as to either (1) tune a program parameter or, (2) compare different algorithms independently of the hardware details.

The MCM model retains many of the characteristics of modern GPU architectures and programming models, like CUDA [95, 73] and OpenCL [111]. However, in order to support algorithm analysis with an emphasis on parallelism overheads, the MCM abstract machines admit a few simplifications and limitations with respect to actual many-core devices.

To further discuss CUDA kernel performance, let us consider now the programming model of CUDA itself and its differences w.r.t. the MCM model. In CUDA, instructions are issued per warp, and a warp consists of a fixed number $S_{warp}$ of threads. Typically, $S_{warp}$ is 32; thus, executing a thread-block on an SM means executing several warps in turn. If an operand of an executing instruction is not ready, then the corresponding warp stalls and context switching happens between warps running on the same SM.

Registers and shared memory are allocated for a thread-block as long as that thread-block is active. Once a thread-block is active, it stays active until all threads in that thread-block have completed. Context switching is very fast because registers and shared memory do not need to be saved and restored. The intention is to hide the latency (of data transfer between the global memory and the private memory of an SM) by having more memory transactions in fly. There is, of course, a hardware limitation to this, characterized by (at least) two numbers:

1. The maximum number of active warps per SM, denoted here by $M_{warp}$; a typical value for $M_{warp}$ is 48 on a Fermi NVIDIA GPU card, leading to a maximum number of $32 \times 48 = 1536$ active threads per SM.
2. The maximum number of active thread-blocks per SM, denoted here by $M_{block}$; a typical
value for $M_{\text{block}}$ is 8 on a Fermi NVIDIA GPU card.

One can now define a popular performance counter of CUDA kernels, the *occupancy* of an SM, given by $A_{\text{warp}}/M_{\text{warp}}$, where $A_{\text{warp}}$ is the number of active warps on that SM. Since resources (registers, shared memory, thread slots) are allocated for an entire thread-block (as long as that thread-block is active), there are three potential limitations to occupancy: register usage, shared memory usage and thread-block size.

As in our discussion of the MCM model, we denote the thread-block size by $\ell$. Regarding the possible values of $\ell$, we observe that (1) the total number of active threads is bounded over by $\ell M_{\text{block}}$; hence, a small value for $\ell$ may limit occupancy, and that (2) a larger value for $\ell$ reduces the number of registers and shared memory words available per thread; thus, this may limit data reuse within a thread-block and potentially increase the amount of data transfer between global memory and the private memory of an SM. Overall, this suggests again that generating the kernel code with $\ell$ and other program parameters considered as input arguments is a desirable goal. With such parametric code at hand, one can optimize, at run-time, the values of those program parameters (like $\ell$), once the machine parameters (like $S_{\text{warp}}$, $M_{\text{warp}}$, $M_{\text{block}}$, $Z$ (private memory size) and the size of the register file) are known.

### 5.2 Automatic parametric CUDA kernel generation

The general purpose of automatic parallelization is to convert sequential computer programs into multithreaded or vectorized code. Following the discussion of Section 5.1, we are interested here in the following more specific question.

Given a theoretically good parallel algorithm (e.g. divide-and-conquer matrix multiplication) and given a type of hardware that depends on various parameters (e.g. a GPGPU with $Z$ words of private memory per SM and a maximum number $M_{\text{warp}}$ of warps supported by an SM, etc.), we aim at automatically generating CUDA kernels that depend on the hardware parameters ($Z$, $M_{\text{warp}}$, etc.) as well as program parameters (e.g. the number $\ell$ of threads per thread-block). Thus, (1) those parameters need not to be known at compile-time, and (2) are encoded as symbols in the generated kernel code. For this reason, we call such CUDA kernels *parametric*.

In contrast, current technology requires that machine and program parameters are specialized to numerical values at the time of generating the GPGPU code, see [57, 6, 66, 115]. The *polyhedron model* described in Section 2.5 is a powerful geometrical tool for analyzing the relation (w.r.t. data locality or parallelization) between the iterations of nested for-loops. Once the polyhedron representing the *iteration space* of a loop nest is calculated, techniques of linear algebra and linear programming can transform this polyhedron into another polyhedron encoding the loop steps into a coordinate system based on time and space (processors). From there, a parallel program can be generated.

For example, for the following code computing the product of two univariate polynomials $a$ and $b$, both of degree $n$, and writing the result to $c$,

```c
for(int i = 0; i <= n; i++) { c[i] = 0; c[i+n] = 0; }
for(int i = 0; i <= n; i++) {
  for(int j = 0; j <= n; j++)
```
For efficient code generation, one should avoid a fine-grained parallelization, and this is achieved by grouping loop steps into so-called *tiles*, which are generally trapezoids [65]. It is also desirable for the generated code to depend on parameters such as tile, cache sizes and number of processors, etc. These extensions lead, however, to the manipulation of systems of non-linear polynomial equations and the use of techniques like quantifier elimination (QE). This was noticed by Größlinger, Griebl and Lengauer in [55] who also observed that work remained to be done for adapting QE tools to the needs of automatic parallelization.

To illustrate these observations, we return to the above example and use a tiling approach: we consider a one-dimensional grid of thread-blocks where each thread-block is in charge of updating at most $B$ coefficients of the polynomial $c$. Therefore, we introduce three variables $B$, $b$ and $u$, where the latter two represent a thread-block index and a thread index (within a thread-block). This brings the following additional relations:

$$
\begin{cases}
0 \leq b \\
0 \leq u < B \\
p = bB + u,
\end{cases}
$$

(5.1)

to the previous system

$$
\begin{cases}
0 < n \\
0 \leq i \leq n \\
0 \leq j \leq n \\
t = n - j \\
p = i + j.
\end{cases}
$$

(5.2)

To determine the target program, one needs to eliminate the variables $i$ and $j$. In this case, Fourier-Motzkin elimination (FME) does not apply any more, due to the presence of non-linear constraints. If all the non-linear constraints appearing in a system of relations are polynomial constraints, the set of real solutions of such a system is a semi-algebraic set. The celebrated Tarski theorem [8] tells us that there always exists a quantifier elimination algorithm to project a semi-algebraic set of $\mathbb{R}^n$ to a semi-algebraic set of $\mathbb{R}^m$, $m \leq n$. The most popular method for conducting quantifier elimination (QE) of a semi-algebraic set is through cylindrical algebraic decomposition (CAD) [31]. Implementation of QE and CAD can be found in software such as QEPCAD [18], REDUCE [64], MATHEMATICA [125] as well as the RegularChains library of

$$
c[i+j] += a[i] * b[j];
$$

}
The MetaFork-to-CUDA code generator

With the goal to generate efficient CUDA code from an input MetaFork program, the keyword meta_schedule is introduced to the MetaFork language such that one can annotate the Meta-
Chapter 5. MetaFork-to-CUDA: Generation of Parametric CUDA Kernels

Fork code targeting many-cores in a precise manner. The semantic of the meta_schedule statement is an indication to the MetaFork-to-CUDA code generator that every meta_for-loop nest of the meta_schedule statement must translate to a CUDA kernel call. In Section 5.2 we illustrated the process of parametric CUDA kernel generation from a sequential C program using MetaFork as an intermediate language. In this section, we assume that, from a C program, one has generated a MetaFork program, which contains one or more meta_schedule statements.

A meta_schedule statement generating a one-dimensional grid with one-dimensional thread-blocks has the following structure

```c
meta_schedule {
    // only for loops are supported here
    meta_for (int i = 0; i < blockDim.x; i++)
    // only for loops are supported here
    meta_for (int j = 0; j < blockDim.x; j++) {
        ... // nested for-loop body
    }
}
```

where the grid (resp. thread-block) dimension size is extracted from the outer (resp. inner) meta_for loop upper bound. Similarly, a meta_schedule statement generating a two-dimensional grid with two-dimensional thread-blocks has the following structure

```c
meta_schedule {
    // only for loops are supported here
    meta_for (int u = 0; u < blockDim.y; u++)
    meta_for (int i = 0; i < blockDim.x; i++)
    // only for loops are supported here
    meta_for (int v = 0; v < blockDim.y; v++)
    meta_for (int j = 0; j < blockDim.x; j++) {
        ... // nested for-loop body
    }
}
```

where the first two outer meta_for loops correspond to the grid and the inner meta_for loops to the thread-blocks.

To generate those parametric CUDA kernels, we rely on PPCG \[115\], a C-to-CUDA compilation framework, which we have modified, in order to generate compilable CUDA kernels. Figure 5.1 illustrates the software architecture of our C-to-CUDA code generator, based on PPCG. As shown in Figure 5.1, one can write a tiled MetaFork program directly or use QE to tile each loop nest in the MetaFork code. After that, one shall pass this tiled MetaFork code to our code generator, in particular, taking advantage of the sophisticated CUDA code generator in PPCG.

The original PPCG framework generates CUDA kernels with the number of threads per thread-block specified as a constant, explicitly given either by the user or by default using 32 for a one-dimensional kernel or 16 × 32 for a two-dimensional kernel. Our code generator focuses on generating parametric CUDA kernels and extends to support identifying meta_for
5.3. The **MetaFork**-to-CUDA code generator

![Diagram of the implementation of the MetaFork-to-CUDA code generator](image)

Figure 5.1: Overview of the implementation of the MetaFork-to-CUDA code generator

loops, such that CUDA code generation is fully automatic for using the global memory as well as, under some assumptions (demonstrated in Appendix C.1), for using the shared memory in the kernel code. Note that, for both data dependence and memory accessing pattern cases, the analysis involving non-linear expressions is not performed by PPCG or MetaFork. Consequently, either the programmer avoids such analysis by writing a proper MetaFork program, or a post-processing phase may be required.

Since the MetaFork code is obtained after computing affine transformation and tiling (via quantifier elimination), we have to bypass the process of computing affine transformation and tiling that PPCG performs. Thus, we modify the internal data structure, namely the schedule tree [117], which is used to represent the execution order of the input code in the polyhedral model and can be converted to the abstract syntax tree (AST). When we pass the MetaFork code to the software PET (polyhedral extraction tool) [116] for parsing, a schedule tree is initiated with the iteration space; however, we mark each `meta for` as “permutable,” which indicates that later, it corresponds to a thread index or a thread-block index. More details on how we use the schedule tree to represent MetaFork and parametric CUDA programs are described in Appendix [C.2]. Furthermore, to generate the parametric variables, such as the grid and thread-block sizes, as symbols, we extend relevant data structures of PPCG to store those variable symbols so as to pass them from the schedule tree to the AST for CUDA code generation. Last but not least, we assign the thread-block indices to variables `b0`, `b1` and the thread indices to variables `t0`, `t1`.

Consider an example, a one-dimensional stencil computation, namely Jacobi. The original (and naive) C version is shown in Figure 5.2, where initialization statements have been
for (int t = 0; t < T; ++t) {
    for (int i = 1; i < N-1; ++i)
        b[i] = (a[i-1] + a[i] + a[i+1]) / 3;

    for (int i = 1; i < N-1; ++i)
        a[i] = b[i];
}

Figure 5.2: One-dimensional stencil computation, namely Jacobi, written in C

removed. From this C code fragment, we apply the tiling techniques mentioned in Section 5.2 and obtain the MetaFork code shown in Figure 5.3. Observe that the meta_schedule statement has two meta_for loop nests yielding two CUDA kernels.

    int ub_v = (N - 2) / B;
    meta_schedule {
        for (int t = 0; t < T; ++t) {
            meta_for (int v = 0; v < ub_v; v++) {
                meta_for (int u = 0; u < B; u++) {
                    int p = v * B + u;
                    b[p+1] = (a[p] + a[p+1] + a[p+2]) / 3;
                }
            }
            meta_for (int v = 0; v < ub_v; v++) {
                meta_for (int u = 0; u < B; u++) {
                    int w = v * B + u;
                    a[w+1] = b[w+1];
                }
            }
        }
    }

Figure 5.3: 1D Jacobi written in MetaFork

Our MetaFork-to-CUDA code generator produces two kernel functions, a header file (for those two kernels) and a host code file where those kernels are called. Those two kernel functions are shown in Figure 5.4. In each kernel, we use the shared memory for those arrays read and use the global memory for those arrays written only once. Observe that kernel0 and kernel1 take a program parameter, the thread-block format B, as an argument, whereas non-parametric CUDA kernels usually take parameters a, b, c, N, T, c0 only. Correspondingly, the generated host code replacing meta_schedule and its body is shown in Figure 5.5. Data transfers between the CPU and GPU global memories are done before those two kernels launched and after those two kernels completed, respectively. In the case that the number of thread-blocks per grid, aka ub_v in the MetaFork code, exceeds the hardware limit, which is 32768 shown in the host code, each kernel uses 32768 as the grid dimension size, while inside the kernel code, the amount of work per thread-block is incremented via a serial loop.
5.3. The MetaFork-to-CUDA Code Generator

```c
__global__ void kernel0(int *a, int *b, int N, int T, int ub_v,
                         int B, int c0) {
    int b0 = blockIdx.x;
    int t0 = threadIdx.x;
    int private_p;
    __shared__ int shared_a[BLOCK_0+2]; // BLOCK_0 = B
    for (int c1 = b0; c1 < ub_v; c1 += 32768) {
        for (int c2 = t0; c2 <= min(B + 1, N - B * c1 - 1); c2 += B)
            shared_a[c2] = a[B * c1 + c2];
        __syncthreads();
        private_p = (((c1) * (B)) + (t0));
        b[private_p + 1] = (((shared_a[private_p - B * c1] +
                       shared_a[private_p - B * c1 + 1]) +
                       shared_a[private_p - B * c1 + 2]) / 3);
        __syncthreads();
    }
}

__global__ void kernel1(int *a, int *b, int N, int T, int ub_v,
                         int B, int c0) {
    int b0 = blockIdx.x;
    int t0 = threadIdx.x;
    int private_w;
    __shared__ int shared_b[BLOCK_0]; // BLOCK_0 = B
    for (int c1 = b0; c1 < ub_v; c1 += 32768) {
        if (N >= t0 + B * c1 + 2)
            shared_b[t0] = b[t0 + B * c1 + 1];
        __syncthreads();
        private_w = (((c1) * (B)) + (t0));
        a[private_w + 1] = shared_b[private_w - B * c1];
        __syncthreads();
    }
}
```

Figure 5.4: Generated parametric CUDA kernel for 1D Jacobi

In order to enable using the shared memory, we enforce PPCG to declare arrays in the shared memory anyway, while PPCG could decide not to use the shared memory because it fails to analyze non-linear expressions. However, this causes another issue, that is, how to calculate the index expressions of these arrays allocated in the shared memory. Recall that the generated kernel code should first copy the data from the global memory to its shared memory counterpart, and then use this shared memory counterpart to compute. In other words, to generate a copy statement, we shall insert into the schedule tree new nodes with the tiling size and affine transformation; however, these new nodes should be inserted to the schedule tree prior to where is the node referring to the statement with the arrays for computation. Using
if (T >= 1 && ub_v >= 1 && B >= 0) {
#define cudaCheckReturn(ret) \ 
  do { \ 
    cudaError_t cudaCheckReturn_e = (ret); \ 
    if (cudaCheckReturn_e != cudaSuccess) { \ 
      fprintf(stderr, "CUDA error: %s\n", \ 
        cudaGetErrorString(cudaCheckReturn_e)); \ 
      fflush(stderr); \ 
    } \ 
    assert(cudaCheckReturn_e == cudaSuccess); \ 
  } while(0)
#define cudaCheckKernel() \ 
  do { \ 
    cudaCheckReturn(cudaGetLastError()); \ 
  } while(0)

int *dev_a;
int *dev_b;

cudaCheckReturn(cudaMalloc((void **) &dev_a, (N) * sizeof(int)));
cudaCheckReturn(cudaMalloc((void **) &dev_b, (N) * sizeof(int)));
if (N >= 1) {
  cudaCheckReturn(cudaMemcpy(dev_a, a, (N) * sizeof(int), cudaMemcpyHostToDevice));
cudaCheckReturn(cudaMemcpy(dev_b, b, (N) * sizeof(int), cudaMemcpyHostToDevice));
}
for (int c0 = 0; c0 < T; c0 += 1) {
  dim3 k0_dimBlock(B);
  dim3 k0_dimGrid(ub_v <= 32767 ? ub_v : 32768);
  kernel0 <<<k0_dimGrid, k0_dimBlock>>> (dev_a, dev_b, N, T, ub_v, B, c0);
cudaCheckKernel();

  dim3 k1_dimBlock(B);
  dim3 k1_dimGrid(ub_v <= 32767 ? ub_v : 32768);
  kernel1 <<<k1_dimGrid, k1_dimBlock>>> (dev_a, dev_b, N, T, ub_v, B, c0);
cudaCheckKernel();
} if (N >= 1) {
  cudaMemcpy(a, dev_a, (N) * sizeof(int), cudaMemcpyDeviceToHost);
cudaCheckReturn(cudaCheckReturn(cudaFree(dev_a)));
cudaCheckReturn(cudaFree(dev_b));
}

Figure 5.5: Generated host code for 1D Jacobi
5.4. Experimentation

In this section, we present experimental results on an NVIDIA Tesla M2050. Most of them were obtained by running times of CUDA programs generated with our preliminary implementation of our MetaFork-to-CUDA code generator described in Section 5.3 and the original version of the PPCG C-to-CUDA code generator \[115\]. We use eight simple test cases: array reversal (Figure 5.6, Table 5.1), 1D Jacobi (Table 5.2), 2D Jacobi (Figure 5.7, Table 5.3), LU decomposition (Figure 5.8, Table 5.4), matrix transposition (Figure 5.9, Table 5.5), matrix addition (Figure 5.10, Table 5.6), matrix vector multiplication (Figure 5.11, Table 5.7) and matrix matrix multiplication (Figure 5.13, Table 5.8). In all cases, we use dense representations for our matrices and vectors.

For both the PPCG C-to-CUDA and our MetaFork-to-CUDA code generators, Tables 5.1, 5.2, 5.3, 5.4, 5.5, 5.6, 5.7 and 5.8 give the speedup factors of the generated code, as the timing ratio of the generated code to their untiled C code. Since PPCG determines a thread-block format, the timings in those tables corresponding to PPCG depend only on the input data size. Meanwhile, since the CUDA kernels generated by MetaFork are parametric, the MetaFork timings are obtained for various formats of thread-blocks and various input data sizes. Indeed, recall that our generated CUDA code admits parameters for the dimension sizes of the thread-blocks. This generated parametric code is then specialized with the thread-block formats listed in the first column of those tables.
Figures 5.6, 5.3 (with 5.2 and 5.4), 5.7, 5.8, 5.9, 5.10, 5.11 and 5.13 show the MetaFork code of eight examples with their untiled serial C programs and automatically generated CUDA kernels. In order to allocate unit sizes of shared memory in the kernel code, we predefine BLOCK_0 and BLOCK_1 (if applicable) as macros and specify their values at compile time. The tiled code for each MetaFork program is done by the quantifier elimination (QE) from the RegularChains library of Maple. These eight examples generated by PPCG are shown in Appendix D.

**Array reversal.** Both MetaFork and PPCG generate CUDA code that uses a one-dimensional kernel grid and the shared memory. We specialize the MetaFork generated parametric code successively to the thread-block size \( B = 16, 32, 64, 128, 256, 512 \); meanwhile, PPCG by default chooses 32 as the thread-block size. As we can see in Table 5.1 based on the generated parametric CUDA kernel, one can tune the thread-block size to be 256 to obtain the best performance.

**Table 5.1: Speedup comparison of reversing a one-dimensional array between PPCG and MetaFork kernel code**

<table>
<thead>
<tr>
<th>Speedup (kernel)</th>
<th>Input size</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>( 2^{23} )</td>
</tr>
<tr>
<td>PPCG</td>
<td></td>
</tr>
<tr>
<td>32</td>
<td>8.312</td>
</tr>
<tr>
<td>MetaFork</td>
<td></td>
</tr>
<tr>
<td>16</td>
<td>4.035</td>
</tr>
<tr>
<td>32</td>
<td>7.612</td>
</tr>
<tr>
<td>64</td>
<td>13.183</td>
</tr>
<tr>
<td>128</td>
<td>19.357</td>
</tr>
<tr>
<td>256</td>
<td>20.451</td>
</tr>
<tr>
<td>512</td>
<td>18.768</td>
</tr>
</tbody>
</table>

Serial code

```c
for (int i = 0; i < N; i++)
    Out[N - 1 - i] = In[i];
```

MetaFork code

```c
__global__ void kernel0(int *In, int *Out, int N, int ub_v, int B)
{
    int b0 = blockIdx.x;
    int t0 = threadIdx.x;
    int private_inoffset;
    int private_outoffset;
    __shared__ int shared_In[BLOCK_0]; \( \text{BLOCK}_0 = B \)
    for (int c0 = b0; c0 < ub_v; c0 += 32768) {
        if (N >= t0 + B * c0 + 1)
            shared_In[t0] = In[t0 + B * c0];
        __syncthreads();
        private_inoffset = (((c0) * (B)) + (t0));
        private_outoffset = (((N) - 1) - private_inoffset);
        Out[private_outoffset] = shared_In[private_inoffset - B * c0];
        __syncthreads();
    }
}
```

Figure 5.6: Serial code, MetaFork code and generated parametric CUDA kernel for array reversal

**1D Jacobi.** Our second example is a one-dimensional stencil computation, namely 1D Jacobi. The kernel generated by MetaFork uses a 1D kernel grid and the shared memory,
while the kernel generated by PPCG uses a 1D kernel grid and the global memory. PPCG by default chooses a thread-block format of 32, while **MetaFork** preferred format is 64.

<table>
<thead>
<tr>
<th>Speedup (kernel)</th>
<th>Input size</th>
<th>Thread-block size</th>
</tr>
</thead>
<tbody>
<tr>
<td>PPCG</td>
<td>1.416</td>
<td>2</td>
</tr>
<tr>
<td></td>
<td>2.424</td>
<td>2</td>
</tr>
<tr>
<td></td>
<td>5.035</td>
<td>2</td>
</tr>
<tr>
<td><strong>MetaFork</strong></td>
<td>1.274</td>
<td>16</td>
</tr>
<tr>
<td></td>
<td>3.660</td>
<td>16</td>
</tr>
<tr>
<td></td>
<td>5.268</td>
<td>16</td>
</tr>
<tr>
<td>32, 32</td>
<td>2.122</td>
<td>64</td>
</tr>
<tr>
<td></td>
<td>4.020</td>
<td>64</td>
</tr>
<tr>
<td></td>
<td>7.309</td>
<td>64</td>
</tr>
<tr>
<td>128, 128</td>
<td>1.787</td>
<td>256</td>
</tr>
<tr>
<td></td>
<td>3.234</td>
<td>256</td>
</tr>
<tr>
<td></td>
<td>6.168</td>
<td>256</td>
</tr>
<tr>
<td>256, 256</td>
<td>1.789</td>
<td>512</td>
</tr>
<tr>
<td></td>
<td>3.516</td>
<td>512</td>
</tr>
<tr>
<td></td>
<td>6.218</td>
<td>512</td>
</tr>
<tr>
<td>512, 512</td>
<td>2.193</td>
<td>128</td>
</tr>
<tr>
<td></td>
<td>3.518</td>
<td>128</td>
</tr>
<tr>
<td></td>
<td>6.070</td>
<td>128</td>
</tr>
</tbody>
</table>

**2D Jacobi.** Our next example is a two-dimensional stencil computation, namely 2D Jacobi. Both the CUDA kernels generated by **MetaFork** and PPCG use a 2D kernel grid and the global memory. PPCG by default chooses a thread-block format of 16 × 32, while **MetaFork** preferred format varies based on input size.

<table>
<thead>
<tr>
<th>Speedup (kernel)</th>
<th>Input size</th>
<th>Thread-block size</th>
</tr>
</thead>
<tbody>
<tr>
<td>PPCG</td>
<td>11.230</td>
<td>16 * 32</td>
</tr>
<tr>
<td></td>
<td>11.303</td>
<td>16 * 32</td>
</tr>
<tr>
<td></td>
<td>9.785</td>
<td>16 * 32</td>
</tr>
<tr>
<td><strong>MetaFork</strong></td>
<td>5.000</td>
<td>8 * 4</td>
</tr>
<tr>
<td></td>
<td>5.256</td>
<td>8 * 4</td>
</tr>
<tr>
<td></td>
<td>4.666</td>
<td>8 * 4</td>
</tr>
<tr>
<td>16 * 4</td>
<td>7.867</td>
<td>32 * 4</td>
</tr>
<tr>
<td></td>
<td>8.724</td>
<td>32 * 4</td>
</tr>
<tr>
<td></td>
<td>7.962</td>
<td>32 * 4</td>
</tr>
<tr>
<td>8 * 8</td>
<td>11.607</td>
<td>16 * 8</td>
</tr>
<tr>
<td></td>
<td>11.143</td>
<td>16 * 8</td>
</tr>
<tr>
<td></td>
<td>9.726</td>
<td>16 * 8</td>
</tr>
<tr>
<td>32 * 8</td>
<td>7.209</td>
<td>32 * 8</td>
</tr>
<tr>
<td></td>
<td>7.776</td>
<td>32 * 8</td>
</tr>
<tr>
<td></td>
<td>6.704</td>
<td>32 * 8</td>
</tr>
<tr>
<td>8 * 16</td>
<td>10.499</td>
<td>8 * 16</td>
</tr>
<tr>
<td></td>
<td>10.502</td>
<td>8 * 16</td>
</tr>
<tr>
<td></td>
<td>7.442</td>
<td>8 * 16</td>
</tr>
<tr>
<td>16 * 16</td>
<td>12.236</td>
<td>32 * 16</td>
</tr>
<tr>
<td></td>
<td>11.487</td>
<td>32 * 16</td>
</tr>
<tr>
<td></td>
<td>9.182</td>
<td>32 * 16</td>
</tr>
<tr>
<td>8 * 32</td>
<td>8.859</td>
<td>8 * 32</td>
</tr>
<tr>
<td></td>
<td>8.825</td>
<td>8 * 32</td>
</tr>
<tr>
<td></td>
<td>5.637</td>
<td>8 * 32</td>
</tr>
<tr>
<td>16 * 32</td>
<td>10.774</td>
<td>16 * 32</td>
</tr>
<tr>
<td></td>
<td>10.709</td>
<td>16 * 32</td>
</tr>
<tr>
<td></td>
<td>7.694</td>
<td>16 * 32</td>
</tr>
<tr>
<td>32 * 32</td>
<td>11.969</td>
<td>32 * 32</td>
</tr>
<tr>
<td></td>
<td>11.442</td>
<td>32 * 32</td>
</tr>
<tr>
<td></td>
<td>10.469</td>
<td>32 * 32</td>
</tr>
</tbody>
</table>

**LU decomposition.** **MetaFork** and PPCG both generate two CUDA kernels: one with a 1D grid and one with a 2D grid, both using the shared memory. The default selected thread-block formats for PPCG are 32 and 16 × 32; meanwhile, the preferred formats by **MetaFork** are 128 and 16 × 16. Tuning the number of threads per thread-block in our parametric code allows **MetaFork** to outperform PPCG.

**Matrix transpose.** Both the CUDA kernels generated by **MetaFork** and PPCG use a 2D grid and the shared memory. PPCG by default chooses a thread-block format of 16 × 32, while
MetaFork preferred format is $8 \times 32$. For MetaFork, the allocation unit size of shared memory for the input matrix is the same as thread-block format. However, for PPCG, the allocation unit size of shared memory for the input matrix is $32 \times 32$, while the thread-block format is $16 \times 32$. Thus, PPCG code transposes two coefficients of the matrix within each thread.

**Matrix addition.** Both the CUDA kernels generated by MetaFork and PPCG use a 2D grid and the global memory. The default chosen thread-block format for PPCG is $16 \times 32$, while MetaFork preferred format is $32 \times 8$.

**Matrix vector multiplication.** For both MetaFork and PPCG, the generated kernels use a 1D grid and the shared memory. The thread-block size chosen by PPCG is $32$, while MetaFork preferred thread-block size varies based on different input sizes. PPCG has a slight advantage due to its ability here to analyze shared and local memory usage for the 2D array in a 1D kernel. For MetaFork, to enhance some components of the PPCG infrastructure with symbolic computation is required, which is a work in progress.

Within the current framework, we post-process the kernel code, such that coalesced accesses occur for copying the 2D array from the global memory to the shared memory. Fig-
Table 5.4: Speedup comparison of LU decomposition between PPCG and MetaFork kernel code

<table>
<thead>
<tr>
<th>Speedup (kernel)</th>
<th>Input size</th>
<th>2^{10} \times 2^{10}</th>
<th>2^{11} \times 2^{11}</th>
</tr>
</thead>
<tbody>
<tr>
<td>Thread-block size</td>
<td>kernel0, kernel1</td>
<td>PPCG</td>
<td>MetaFork</td>
</tr>
<tr>
<td></td>
<td></td>
<td>32, 16 * 32</td>
<td>10.712</td>
</tr>
<tr>
<td></td>
<td></td>
<td>128, 4 * 4</td>
<td>3.063</td>
</tr>
<tr>
<td></td>
<td></td>
<td>256, 4 * 4</td>
<td>3.077</td>
</tr>
<tr>
<td></td>
<td></td>
<td>512, 4 * 4</td>
<td>3.095</td>
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<tr>
<td></td>
<td></td>
<td>32, 8 * 8</td>
<td>10.721</td>
</tr>
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<td></td>
<td></td>
<td>64, 8 * 8</td>
<td>10.604</td>
</tr>
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<td></td>
<td></td>
<td>128, 8 * 8</td>
<td>10.463</td>
</tr>
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<td></td>
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<td>256, 8 * 8</td>
<td>10.831</td>
</tr>
<tr>
<td></td>
<td></td>
<td>512, 8 * 8</td>
<td>10.416</td>
</tr>
<tr>
<td></td>
<td></td>
<td>32, 16 * 16</td>
<td>14.533</td>
</tr>
<tr>
<td></td>
<td></td>
<td>64, 16 * 16</td>
<td>14.457</td>
</tr>
<tr>
<td></td>
<td></td>
<td>128, 16 * 16</td>
<td>14.877</td>
</tr>
<tr>
<td></td>
<td></td>
<td>256, 16 * 16</td>
<td>14.803</td>
</tr>
<tr>
<td></td>
<td></td>
<td>512, 16 * 16</td>
<td>14.479</td>
</tr>
</tbody>
</table>

Table 5.5: Speedup comparison of matrix transpose between PPCG and MetaFork kernel code

<table>
<thead>
<tr>
<th>Speedup (kernel)</th>
<th>Input size</th>
<th>2^{13} \times 2^{13}</th>
<th>2^{14} \times 2^{14}</th>
</tr>
</thead>
<tbody>
<tr>
<td>Thread-block size</td>
<td>PPCG</td>
<td>16 * 32</td>
<td>62.656</td>
</tr>
<tr>
<td></td>
<td>MetaFork</td>
<td>8 * 4</td>
<td>28.626</td>
</tr>
<tr>
<td></td>
<td></td>
<td>16 * 4</td>
<td>40.381</td>
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<tr>
<td></td>
<td></td>
<td>32 * 4</td>
<td>28.728</td>
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<tr>
<td></td>
<td></td>
<td>8 * 8</td>
<td>51.889</td>
</tr>
<tr>
<td></td>
<td></td>
<td>16 * 8</td>
<td>44.759</td>
</tr>
<tr>
<td></td>
<td></td>
<td>32 * 8</td>
<td>37.586</td>
</tr>
<tr>
<td></td>
<td></td>
<td>8 * 16</td>
<td>70.716</td>
</tr>
<tr>
<td></td>
<td></td>
<td>16 * 16</td>
<td>64.812</td>
</tr>
<tr>
<td></td>
<td></td>
<td>32 * 16</td>
<td>36.109</td>
</tr>
<tr>
<td></td>
<td></td>
<td>8 * 32</td>
<td>77.327</td>
</tr>
<tr>
<td></td>
<td></td>
<td>16 * 32</td>
<td>62.268</td>
</tr>
</tbody>
</table>

ure[5.12] shows the post-processed CUDA kernel code. On line 16 for copying the input matrix from the global memory to the array, namely shared_a, allocated in the shared memory, we exchanged the row index with the column index of this array shared_a, so that threads within a warp access to the corresponding global memory in a coalesced manner. Table[5.7] shows the speedup factors obtained with the kernels generated by PPCG, MetaFork and MetaFork with
Figure 5.8: Serial code, MetaFork code and generated parametric CUDA kernel for LU decomposition
Serial code
for (int v0 = 0; v0 < n; v0++)
for (int v1 = 0; v1 < n; v1++)
c[v0][v1] = a[v1][v0];

MetaFork code
int dim0 = n / B0, dim1 = n / B1;
meta_schedule {
  meta_for (int v0 = 0; v0 < dim0; v0++)
    meta_for (int v1 = 0; v1 < dim1; v1++)
      meta_for (int u0 = 0; u0 < B0; u0++)
        meta_for (int u1 = 0; u1 < B1; u1++)
        {
          int i = u0 + v0 * B0;
          int j = u1 + v1 * B1;
          c[j][i] = a[i][j];
        }
  }
}

__global__ void kernel0(int *a, int *c, int n, int dim0,
int dim1, int B0, int B1) {
  int b0 = blockIdx.y, b1 = blockIdx.x;
  int t0 = threadIdx.y, t1 = threadIdx.x;
  int private_i;
  int private_j;
  // BLOCK_0 = B0, BLOCK_1 = B1
  __shared__ int shared_a[BLOCK_0][BLOCK_1];

  for (int c0 = b0; c0 < dim0; c0 += 256)
    for (int c1 = b1; c1 < dim1; c1 += 256) {
      if (n >= t0 + B0 * c0 + 1 && n >= t1 + B1 * c1 + 1)
        shared_a[t0][t1] = a[(t0 + B0 * c0) * n + (t1 + B1 * c1)];
      __syncthreads();
      private_i = ((t0) + ((c0) * (B0)));
      private_j = ((t1) + ((c1) * (B1)));
      c[private_j * n + private_i] =
        shared_a[private_i - B0 * c0][private_j - B1 * c1];
      __syncthreads();
    }
}

Figure 5.9: Serial code, MetaFork code and generated parametric CUDA kernel for matrix transpose

Table 5.6: Speedup comparison of matrix addition between PPCG and MetaFork kernel code

<table>
<thead>
<tr>
<th>Speedup (kernel)</th>
<th>Input size</th>
</tr>
</thead>
<tbody>
<tr>
<td>Thread-block size</td>
<td>2^12</td>
</tr>
<tr>
<td>PPCG</td>
<td>16 * 32</td>
</tr>
<tr>
<td>MetaFork</td>
<td>8 * 4</td>
</tr>
<tr>
<td></td>
<td>16 * 4</td>
</tr>
<tr>
<td></td>
<td>32 * 4</td>
</tr>
<tr>
<td></td>
<td>8 * 8</td>
</tr>
<tr>
<td></td>
<td>16 * 8</td>
</tr>
<tr>
<td></td>
<td>32 * 8</td>
</tr>
<tr>
<td></td>
<td>8 * 16</td>
</tr>
<tr>
<td></td>
<td>16 * 16</td>
</tr>
<tr>
<td></td>
<td>32 * 16</td>
</tr>
</tbody>
</table>

Fork preferred thread-block size varies based on input sizes. For MetaFork, the allocation unit size of shared memory for each input matrix is the same as thread-block format. However, for PPCG, the allocation unit size of shared memory for each input matrix is 32 × 32, while the thread-block format is 16 × 32. In fact, PPCG code computes two coefficients of the output matrix within each thread, thus increasing index arithmetic amortization and occupancy.

We conclude this section with timings (in seconds) for the quantifier elimination (QE) required to generate MetaFork tiled code, see Table 5.9. Our tests are based on the latest version of the RegularChains library of Maple, available at [www.regularchains.org](http://www.regularchains.org). These results show that the use of QE is not a bottleneck in our C-to-CUDA code translation process, despite the theoretically high algebraic complexity of quantifier elimination.
Serial code

for (int v0 = 0; v0 < n; v0++)
  for (int v1 = 0; v1 < n; v1++)
    c[v0][v1] = a[v0][v1] + b[v0][v1];

MetaFork code

int dim0 = n / B0, dim1 = n / B1;
meta_schedule {
  meta_for (int v0 = 0; v0 < dim0; v0++)
    meta_for (int v1 = 0; v1 < dim1; v1++)
      meta_for (int u0 = 0; u0 < B0; u0++)
        meta_for (int u1 = 0; u1 < B1; u1++)
        {
          int i = u0 + v0 * B0;
          int j = u1 + v1 * B1;
          c[i][j] = a[i][j] + b[i][j];
        }
}

__global__ void kernel0(int *a, int *b, int *c, int n,
  int dim0, int dim1, int B0, int B1) {
  int b0 = blockIdx.y, b1 = blockIdx.x;
  int t0 = threadIdx.y, t1 = threadIdx.x;
  int private_i;
  int private_j;
  for (int c0 = b0; c0 < dim0; c0 += 256)
    for (int c1 = b1; c1 < dim1; c1 += 256) {
      private_i = ((t0) + ((c0) * (B0)));
      private_j = ((t1) + ((c1) * (B1)));
      c[private_i * n + private_j] =
        (a[private_i * n + private_j] +
        b[private_i * n + private_j]);
    }
  __syncthreads();
}

Figure 5.10: Serial code, MetaFork code and generated parametric CUDA kernel for matrix addition

Table 5.7: Speedup comparison of matrix vector multiplication among PPCG kernel code, MetaFork kernel code and MetaFork kernel code with post-processing

<table>
<thead>
<tr>
<th>Speedup (kernel)</th>
<th>Input size</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>2^11</td>
</tr>
<tr>
<td>Thread-block size</td>
<td></td>
</tr>
<tr>
<td>PPCG</td>
<td></td>
</tr>
<tr>
<td>32</td>
<td>3.954</td>
</tr>
<tr>
<td>MetaFork</td>
<td></td>
</tr>
<tr>
<td>16</td>
<td>3.108</td>
</tr>
<tr>
<td>32</td>
<td><strong>4.116</strong></td>
</tr>
<tr>
<td>64</td>
<td>2.955</td>
</tr>
<tr>
<td>128</td>
<td>2.658</td>
</tr>
<tr>
<td>256</td>
<td>2.215</td>
</tr>
<tr>
<td>MetaFork with post-processing</td>
<td></td>
</tr>
<tr>
<td>16</td>
<td>4.976</td>
</tr>
<tr>
<td>32</td>
<td><strong>8.698</strong></td>
</tr>
<tr>
<td>64</td>
<td>4.260</td>
</tr>
</tbody>
</table>

5.5 Conclusion

In this chapter, we have presented enhancements of the MetaFork language so as to provide the model of concurrency for SIMD on GPUs. Our objective is to facilitate automatic code translation of high-level programming models supporting hardware accelerator (like OpenMP and OpenACC) to low-level heterogeneous programming models (like CUDA).

As illustrated in Sections 5.3, MetaFork has language constructs to help generate efficient CUDA code. Moreover, the MetaFork framework relies on advanced techniques (quantifier elimination in non-linear polynomial expressions) for code optimization, in particular tiling. The experimentation reported in Section 5.4 shows the benefits of generating parametric CUDA kernels. Not only does this feature provide more portability, but it also helps to improve per-
5.5. Conclusion

Serial code

```c
for (int i = 0; i < n; i++)
    for (int j = 0; j < n; j++)
        c[i] += a[i][j] * b[j];
```

MetaFork code

```c
int dim = n / B;
meta_schedule {
    for (int i = 0; i < n / 16; i++)
        for (int j = 0; j < 16; j++)
            if (n >= i16 + u) {
                int p = v * B + u;
                c[p] += a[p][i16 + j] * b[i16 + j];
            }
}
```

__global__ void kernel0(int *a, int *b, int *c, int n, int dim, int B) {
    int b0 = blockIdx.x;
    int t0 = threadIdx.x;
    int private_p;
    // BLOCK_0 = B
    __shared__ int shared_a[BLOCK_0][16];
    __shared__ int shared_b[16];
    __shared__ int shared_c[BLOCK_0];
    for (int c0 = b0; c0 < dim; c0 += 32768) {
        if (n >= t0 + B * c0 + 1)
            shared_c[t0] = c[t0 + B * c0];
        for (int c1 = 0; c1 < n / 16; c1 += 1) {
            if (n >= t0 + B * c0 + 1)
                for (int c3 = 0; c3 <= 15; c3 += 1)
                    shared_a[t0][c3] =
                        a[(t0 + B * c0) * n + (16 * c1 + c3)];
            if (t0 <= 15)
                shared_b[t0] = b[t0 + 16 * c1];
            __syncthreads();
            for (int c3 = 0; c3 <= 15; c3 += 1)
                private_p = (((c0) * (B)) + (t0));
                shared_c[private_p - B * c0] +=
                    (shared_a[private_p - B * c0][c3] * shared_b[c3]);
            __syncthreads();
        }
    if (n >= t0 + B * c0 + 1)
        c[t0 + B * c0] = shared_c[t0];
    __syncthreads();
}
```

Figure 5.11: Serial code, MetaFork code and generated parametric CUDA kernel for matrix vector multiplication

Table 5.8: Speedup comparison of matrix multiplication between PPCG and MetaFork kernel code

<table>
<thead>
<tr>
<th>Speedup (kernel)</th>
<th>Input size</th>
<th>2^{10} * 2^{10}</th>
<th>2^{11} * 2^{11}</th>
</tr>
</thead>
<tbody>
<tr>
<td>Thread-block size</td>
<td>16 * 32</td>
<td>129.853</td>
<td>393.851</td>
</tr>
<tr>
<td>PPCG</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>16 * 4</td>
<td>32.157</td>
<td>96.652</td>
<td></td>
</tr>
<tr>
<td>16 * 8</td>
<td>54.578</td>
<td>171.621</td>
<td></td>
</tr>
<tr>
<td>32 * 4</td>
<td>53.399</td>
<td>156.493</td>
<td></td>
</tr>
<tr>
<td>32 * 8</td>
<td>60.358</td>
<td>182.557</td>
<td></td>
</tr>
<tr>
<td>16 * 16</td>
<td>87.919</td>
<td>287.002</td>
<td></td>
</tr>
<tr>
<td>32 * 16</td>
<td>84.057</td>
<td>247.220</td>
<td></td>
</tr>
<tr>
<td>MetaFork</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>8 * 16</td>
<td>100.521</td>
<td>299.228</td>
<td></td>
</tr>
<tr>
<td>16 * 16</td>
<td>100.264</td>
<td>330.965</td>
<td></td>
</tr>
</tbody>
</table>

Performance with automatically generated code.
__global__ void kernel0(int *a, int *b, int *c, int n, int dim, int B) {
    int b0 = blockIdx.x;
    int t0 = threadIdx.x;
    int private_p;
    // BLOCK_0 = B
    __shared__ int shared_a[BLOCK_0][BLOCK_0];
    __shared__ int shared_b[BLOCK_0];
    __shared__ int shared_c[BLOCK_0];

    for (int c0 = b0; c0 < dim; c0 += 32768) {
        if (n >= t0 + B * c0 + 1)
            shared_c[t0] = c[t0 + B * c0];
        for (int c1 = 0; c1 < n / BLOCK_0; c1 += 1) {
            if (n >= t0 + B * c0 + 1)
                for (int c3 = 0; c3 < BLOCK_0; c3 += 1)
                    shared_a[c3][t0] = a[(c3 + B * c0) * n + (B * c1 + t0)];
            shared_b[t0] = b[t0 + B * c1];
            __syncthreads();
            for (int c3 = 0; c3 < BLOCK_0; c3 += 1)
                private_p = (((c0) * (B)) + (t0));
            shared_c[private_p - B * c0] +=
                (shared_a[private_p - B * c0][c3] * shared_b[c3]);
            __syncthreads();
        }
    }
    __syncthreads();
}

if (n >= t0 + B * c0 + 1)
    c[t0 + B * c0] = shared_c[t0];
__syncthreads();
}

Figure 5.12: Post-processing CUDA kernel with coalesced accesses for matrix vector multiplication

<table>
<thead>
<tr>
<th>Example</th>
<th>Timing</th>
</tr>
</thead>
<tbody>
<tr>
<td>Array reversal</td>
<td>0.072</td>
</tr>
<tr>
<td>1D Jacobi</td>
<td>0.948</td>
</tr>
<tr>
<td>2D Jacobi</td>
<td>7.735</td>
</tr>
<tr>
<td>LU decomposition</td>
<td>4.416</td>
</tr>
<tr>
<td>matrix transposition</td>
<td>1.314</td>
</tr>
<tr>
<td>matrix addition</td>
<td>1.314</td>
</tr>
<tr>
<td>matrix vector multiplication</td>
<td>0.072</td>
</tr>
<tr>
<td>matrix matrix multiplication</td>
<td>2.849</td>
</tr>
</tbody>
</table>
5.5. Conclusion

Serial code

```c
for (int i = 0; i < n; i++)
    for (int j = 0; j < n; j++)
        for (int k = 0; k < n; ++k)
            c[i][j] += a[i][k] * b[k][j];
```

MetaFork code

```c
int dim0 = n / B0, dim1 = n / B1;
meta_schedule {
    meta_for (int i = 0; i < dim0; i++)
        meta_for (int j = 0; j < dim1; j++)
            meta_for (int v = 0; v < B0; v++)
                meta_for (int u = 0; u < B1; u++)
                {
                    int p = i * B0 + v;
                    int w = j * B1 + u;
                    for (int z = 0; z < 4; z++)
                        c[p][w] +=
                            a[p][4*k+z] * b[4*k+z][w];
                }
}
```

MetaFork code

```c
__global__ void kernel0(int *a, int *b, int *c, int n, int dim0,
                        int dim1, int B0, int B1) {
    int b0 = blockIdx.y, b1 = blockIdx.x;
    int t0 = threadIdx.y, t1 = threadIdx.x;
    int private_p;
    int private_w;
    // BLOCK_0 = B0, BLOCK_1 = B1
    __shared__ int shared_a[BLOCK_0][4];
    __shared__ int shared_b[4][BLOCK_1];
    __shared__ int shared_c[BLOCK_0][BLOCK_1];
    for (int c0 = b0; c0 < dim0; c0 += 256)
        for (int c1 = b1; c1 < dim1; c1 += 256) {
            if (n >= t0 + B0 * c0 + 1 && n >= t1 + B1 * c1 + 1)
                shared_c[t0][t1] =
                    a[(t0 + B0 * c0) * n + (t1 + B1 * c1)];
            for (int c2 = 0; c2 < n / 4; c2 += 1) {
                if (t1 <= 3 && n >= t0 + B0 * c0 + 1)
                    shared_a[t0][t1] =
                        a[(t0 + B0 * c0) * n + (t1 + 4 * c2)];
                if (t0 <= 3 && n >= t1 + B1 * c1 + 1)
                    shared_b[t0][t1] =
                        b[(t0 + 4 * c2) * n + (t1 + B1 * c1)];
            }
            __syncthreads();
            private_p = (((c0) * (B0)) + (t0));
            private_w = (((c1) * (B1)) + (t1));
            for (int c5 = 0; c5 < 3; c5 += 1)
                shared_c[private_p - B0 * c0][private_w - B1 * c1] +=
                    (shared_a[private_p - B0 * c0][c5] *
                     shared_b[c5][private_w - B1 * c1]);
            __syncthreads();
            if (n >= t0 + B0 * c0 + 1 && n >= t1 + B1 * c1 + 1)
                c[(t0 + B0 * c0) * n + (t1 + B1 * c1)] =
                    shared_c[t0][t1];
        }
}
```

Figure 5.13: Serial code, MetaFork code and generated parametric CUDA kernel for matrix matrix multiplication
Chapter 6

Generation of Optimized CUDA Kernel Code

In the previous chapter, we observed that, for several test cases, such as matrix multiplication and matrix transposition, the parametric CUDA kernels generated by MetaFork could not outperform those that are generated by PPCG, for which thread-block formats are given by numerical values. This observation motivates the work reported in this chapter. In Section 6.1, we revisit a particular test case, matrix multiplication, by exploring various optimization techniques, such as controlling the granularity of threads and using the local memory of the device. Since accessing shared (resp. local) memory has low latency, one can consider to declare an array allocated in the shared (resp. local) memory for temporarily storing the corresponding elements of the array in the global memory. In the sequel of this chapter, we refer such an array allocated in the shared (resp. local) memory as the shared (resp. local) memory counterpart of the array in the global memory. Furthermore, we apply those optimization techniques to five additional test cases in Section 6.2, so as to evaluate the benefits of those optimization techniques. All experimental results are collected on an NVIDIA Tesla M2050.

6.1 Case study: matrix multiplication

The MetaFork code for matrix multiplication, shown in Figure 5.13 of Chapter 5, follows the naive implementation described in [73]. To be more precise, each thread-block computes
6.1. Case study: matrix multiplication

a block of the output matrix, while each thread computes one element of the output matrix. Figure 6.1 shows the formula to calculate the multiplication of two matrices with each block of size 1. Recall that when each input matrix is of order $2^{10}$, the best speedup factor that MetaFork achieves is 100.521 for the thread-block format of size $8 \times 16$, while PPCG achieves the speedup factor of 129.853 for the thread-block format of size $16 \times 32$. Our generated kernel code uses the shared memory for each of the input and output matrices. However, we observe that PPCG generates a local memory counterpart of size 2 for the output matrix and two shared memory counterparts, each of size $32 \times 32$, for input matrices. This means that for each thread, the kernel code generated by PPCG computes two elements of the output matrix.

Some studies \cite{73, 118} demonstrate that better performance can be obtained, when one tunes the granularity of threads to increase the arithmetic intensity and hide the data transfer time. This suggests that the amount of work that each thread executes can become a program parameter. We can tune this parameter as we do with the thread-block format parameters. Thus, we introduce a granularity loop into the original MetaFork code as shown in Figure 6.2, where the $w$-loop on Line (13) increases the work per thread by a parameter $s$. Let each thread-block of the original MetaFork code compute $B_0 \times B_1$ elements of the output matrix. Then, for the new code with the granularity loop, each thread-block computes the same number of elements of the output matrix, while each thread computes $s$ elements and the number of threads per thread-block reduces to $B_0 \times B_1 / s$ (that is, $B_0 \times ub_1$ as shown in Figure 6.2). In addition, we guarantee that the data, which threads access to, has a good alignment during each iteration of the $w$-loop, such that coalesced accesses from a warp of threads occur.

```cpp
// n * n matrices
// Program parameters: B0, ub1, s
assert(BLOCK == min(B0, ub1 * s));
int dim0 = n / B0, dim1 = n / (ub1 * s);

meta_schedule {
   meta_for (int i = 0; i < dim0; i++)
      meta_for (int j = 0; j < dim1; j++)
         for (int k = 0; k < n / BLOCK; ++k)
            meta_for (int v = 0; v < B0; v++)
               meta_for (int u = 0; u < ub1; u++)
                  // Each thread computes BLOCK*s outputs
                  for (int w = 0; w < s; ++w) {
                     int p = i * B0 + v;
                     int q = j * ub1 * s + w * ub1 + u;
                     for (int z = 0; z < BLOCK; z++)
                        c[p][q] += a[p][BLOCK*k+z] * b[BLOCK*k+z][q];
                  }
   }
}
```

Figure 6.2: The MetaFork code with the granularity loop and good data alignment for matrix multiplication

Introducing the granularity loop leads to a non-linear expression

$$ j \times ub1 \times s + w \times ub1 + u $$
on Line (15) in Figure 6.2, which is used as a column index of the two-dimensional arrays b and c. The current MetaFork-to-CUDA code generator does not support this format of non-linear expressions yet. In order to see the benefits of using the granularity loop with good data alignment, we manually process the generated kernel code to obtain the new kernel code shown in Figure 6.3. In the first scenario, we continue to use the shared memory for input and output matrices in the generated kernel code. During the post-processing phase, we modify the indices of arrays allocated in the global and shared memories, respectively. Then we collect the speedup factors and achieved occupancies in Table 6.1 for various values of the number of threads per thread-block and the granularity of threads. For the best case, when the thread-block format is $8 \times 32$ and the granularity of threads is 4, the kernel code uses 23 registers and 8448 bytes of shared memory. We observe that this kernel achieves better performance, when compared to those kernels automatically generated by PPCG and MetaFork.

Table 6.1: Experimental results of matrix multiplication for the CUDA kernel with the shared memory for the output matrix and the granularity of threads

(a) speedup factors

<table>
<thead>
<tr>
<th>Thread-block \ Granularity</th>
<th>2</th>
<th>4</th>
<th>8</th>
</tr>
</thead>
<tbody>
<tr>
<td>(8, 8)</td>
<td>99.422</td>
<td>92.612</td>
<td>100.406</td>
</tr>
<tr>
<td>(16, 8)</td>
<td>130.053</td>
<td>143.995</td>
<td>130.213</td>
</tr>
<tr>
<td>(32, 8)</td>
<td>138.445</td>
<td><strong>152.369</strong></td>
<td>106.153</td>
</tr>
<tr>
<td>(64, 8)</td>
<td>124.984</td>
<td>128.164</td>
<td>84.301</td>
</tr>
</tbody>
</table>

(b) achieved occupancies

<table>
<thead>
<tr>
<th>Thread-block \ Granularity</th>
<th>2</th>
<th>4</th>
<th>8</th>
</tr>
</thead>
<tbody>
<tr>
<td>(8, 8)</td>
<td>0.332</td>
<td>0.330</td>
<td>0.326</td>
</tr>
<tr>
<td>(16, 8)</td>
<td>0.660</td>
<td>0.654</td>
<td>0.407</td>
</tr>
<tr>
<td>(32, 8)</td>
<td>0.823</td>
<td><strong>0.813</strong></td>
<td>0.332</td>
</tr>
<tr>
<td>(64, 8)</td>
<td>0.662</td>
<td>0.658</td>
<td>0.333</td>
</tr>
</tbody>
</table>

Input size $2^{10} \times 2^{10}$

In the second scenario, we use the local memory, instead of the shared memory, for the output matrix. Thereby, we replace the code

```c
__shared__ int shared_c[BLOCK_0][BLOCK_1]
```

on Line (11) in Figure 6.3 by

```c
int private_c [1][STRIDE],
```

where STRIDE is predefined as a macro and is equal to s. For the array references

```c
shared_c[t0][c5*ub1+t1],
```

which appeared at Lines (19), (30) and (38) on Figure 6.3 we replace each array reference by

```c
private_c[0][c5].
```
6.1. Case study: matrix multiplication

```c
__global__ void kernel0(int *a, int *b, int *c, int n, int dim0, int dim1,
    int B0, int ub1, int s) {
    int b0 = blockIdx.y, b1 = blockIdx.x;
    int t0 = threadIdx.y, t1 = threadIdx.x;
    int private_p;
    int private_q;
    assert (BLOCK == min(B0, ub1 * s));
    assert (BLOCK_0 == B0); assert (BLOCK_1 == ub1 * s);
    __shared__ int shared_a[BLOCK_0][BLOCK];
    __shared__ int shared_b[BLOCK][BLOCK_1];
    __shared__ int shared_c[BLOCK_0][BLOCK_1];

    for (int c0 = b0; c0 < dim0; c0 += 256)
        for (int c1 = b1; c1 < dim1; c1 += 256) {
            private_p = ((c0) * (B0)) + (t0);
            private_q = (c1) * (ub1 * s) + (t1);
            for (int c5 = 0; c5 < s; c5 += 1)
                if (n >= private_p + 1 && n >= private_q + (c5) * (ub1) + 1)
                    shared_c[t0][c5*ub1+t1] =
                        c[(private_p) * n + (private_q + (c5) * (ub1))];
            for (int c2 = 0; c2 < n / BLOCK; c2 += 1) {
                if (t1 < BLOCK && n >= private_p + 1)
                    shared_a[t0][t1] = a[(private_p) * n + (t1 + BLOCK * c2)];
                for (int c5 = 0; c5 < s; c5 += 1)
                    if (t0 < BLOCK && n >= private_q + (c5) * (ub1) + 1)
                        shared_b[t0][(c5) * (ub1) + t1] =
                            b[(t0 + BLOCK * c2) * n + (private_q + (c5) * (ub1))];
            __syncthreads();
            for (int c6 = 0; c6 < BLOCK; c6 += 1)
                shared_c[t0][c5*ub1+t1] +=
                    (shared_a[t0][c6] * shared_b[c6][c5 * ub1 + t1]);
        __syncthreads();
    }
    __syncthreads();
}
```

Figure 6.3: Post-processing the generated CUDA kernel code for matrix multiplication with the granularity loop

We collect the experimental results in Table 6.2, which contains the speedup factors and achieved occupancies for various values of the number of threads per thread-block and the granularity of threads. For the best case, when the thread-block format is $8 \times 32$ and the granularity of threads is $4$, the kernel code uses $16$ bytes of stack frame, $22$ registers and $2176$ bytes of shared memory. However, the performance is worse than that of the kernel using the shared memory for the output matrix.
Table 6.2: Experimental results of matrix multiplication for the CUDA kernel with the local memory for the output matrix and the granularity of threads

(a) speedup factors

<table>
<thead>
<tr>
<th>Thread-block \ Granularity</th>
<th>2</th>
<th>4</th>
<th>8</th>
</tr>
</thead>
<tbody>
<tr>
<td>(8, 8)</td>
<td>87.652</td>
<td>93.717</td>
<td>95.032</td>
</tr>
<tr>
<td>(16, 8)</td>
<td>127.22</td>
<td>133.620</td>
<td>127.204</td>
</tr>
<tr>
<td>(32, 8)</td>
<td>133.312</td>
<td>142.351</td>
<td>125.563</td>
</tr>
<tr>
<td>(64, 8)</td>
<td>114.836</td>
<td>124.092</td>
<td>116.646</td>
</tr>
</tbody>
</table>

(b) achieved occupancies

<table>
<thead>
<tr>
<th>Thread-block \ Granularity</th>
<th>2</th>
<th>4</th>
<th>8</th>
</tr>
</thead>
<tbody>
<tr>
<td>(8, 8)</td>
<td>0.332</td>
<td>0.331</td>
<td>0.327</td>
</tr>
<tr>
<td>(16, 8)</td>
<td>0.661</td>
<td>0.655</td>
<td>0.647</td>
</tr>
<tr>
<td>(32, 8)</td>
<td>0.824</td>
<td>0.815</td>
<td>0.804</td>
</tr>
<tr>
<td>(64, 8)</td>
<td>0.662</td>
<td>0.658</td>
<td>0.651</td>
</tr>
</tbody>
</table>

Input size $2^{10} \times 2^{10}$

One key reason explaining the relatively low performance of our kernel code using the local memory is the fact that the upper bound of the granularity loop is a variable instead of a constant. Indeed, this prevents the compiler from applying loop unrolling, which is an essential trick for exposing instruction level parallelism (ILP). Hence, we shall use a constant as the w-loop’s upper bound at Line (13) on Figure 6.2. Correspondingly, we modify the generated kernel code at Line (24) on Figure 6.3 by changing the variable $s$ to the constant $\text{STRIDE}$. Note that $\text{STRIDE}$, $\text{BLOCK}_0$ and $\text{BLOCK}_1$ are constants that are passed by the compilation flags -DSTRIDE, -DBLOCK_0 and -DBLOCK_1, respectively. Then we collect the speedup factors for various values of the granularity size and the thread-block format in Table 6.3. Due to the loop unrolling, the usage of registers per thread increases from 22 to 27. At this point, the performance of the modified kernel has been further improved, compared to the kernel in the first scenario, that is, using the shared memory for the output matrix.

Table 6.3: For input matrices of order $2^{10}$, speedup factors of the matrix multiplication kernel unrolling the computation

<table>
<thead>
<tr>
<th>Thread-block \ Granularity</th>
<th>2</th>
<th>4</th>
<th>Resource usage</th>
</tr>
</thead>
<tbody>
<tr>
<td>(16, 4)</td>
<td>79.038</td>
<td>89.233</td>
<td>26 registers, 2112 bytes of shared memory</td>
</tr>
<tr>
<td>(32, 4)</td>
<td>104.898</td>
<td>123.262</td>
<td></td>
</tr>
<tr>
<td>(64, 4)</td>
<td>106.147</td>
<td>111.481</td>
<td></td>
</tr>
<tr>
<td>(8, 8)</td>
<td>116.090</td>
<td>137.318</td>
<td></td>
</tr>
<tr>
<td>(16, 8)</td>
<td>165.192</td>
<td>194.326</td>
<td>27 registers, 2304 bytes of shared memory</td>
</tr>
<tr>
<td>(32, 8)</td>
<td>166.310</td>
<td>180.835</td>
<td></td>
</tr>
<tr>
<td>(64, 8)</td>
<td>137.937</td>
<td>161.536</td>
<td></td>
</tr>
</tbody>
</table>

With the intention to maximize ILP, we unroll the loop for the copy-in and copy-out phases of the data transfer between the global memory and the local memory. Then, we modify the upper bounds on Lines (17) and (35) by changing the variable $s$ to the constant $\text{STRIDE}$, so as to obtain the final modified kernel code as shown in Figure 6.4. Then we collect the experimental results in Table 6.4 for various values of the granularity size and the thread-block format. We
6.1. Case study: matrix multiplication

__global__ void kernel0(int *a, int *b, int *c, int n, int dim0, int dim1,
    int B0, int ub1, int s) {
    int b0 = blockIdx.y, b1 = blockIdx.x;
    int t0 = threadIdx.y, t1 = threadIdx.x;
    int private_p;
    int private_q;
    assert (BLOCK == min(B0, ub1 * s));
    assert (BLOCK_0 == B0); assert (BLOCK_1 == ub1 * s);
    assert (STRIDE == s);
    __shared__ int shared_a[BLOCK_0][BLOCK];
    __shared__ int shared_b[BLOCK][BLOCK_1];
    int private_c[1][STRIDE];

    for (int c0 = b0; c0 < dim0; c0 += 256)
        for (int c1 = b1; c1 < dim1; c1 += 256) {
            private_p = (c0) * (B0) + (t0);
            private_q = (c1) * (ub1 * s) + (t1);
            for (int c5 = 0; c5 < STRIDE; c5 += 1)
                if (n >= private_p + 1 && n >= private_q + (c5) * (ub1) + 1)
                    private_c[0][c5] =
                    c[(private_p) * n + (private_q + (c5) * (ub1))];
            for (int c2 = 0; c2 < n / BLOCK; c2 += 1) {
                if (t1 < BLOCK && n >= private_p + 1)
                    shared_a[t0][t1] = a[(private_p) * n + (t1 + BLOCK * c2)];
                for (int c5 = 0; c5 < s; c5 += 1)
                    if (t0 < BLOCK && n >= private_q + (c5) * (ub1) + 1)
                        shared_b[t0][(c5) * (ub1) + t1] =
                        b[(t0 + BLOCK * c2) * n + (private_q + (c5) * (ub1))];
                    __syncthreads();
            for (int c6 = 0; c6 < BLOCK; c6 += 1)
                for (int c5 = 0; c5 < STRIDE; c5 += 1)
                    private_c[0][c5] +=
                    (shared_a[t0][c6] * shared_b[c6][c5 * ub1 + t1]);
            __syncthreads();
        }
    for (int c5 = 0; c5 < STRIDE; c5 += 1)
        if (n >= private_p + 1 && n >= private_q + (c5) * (ub1) + 1)
            c[(private_p) * n + (private_q + (c5) * (ub1))] =
            private_c[0][c5];
    __syncthreads();
}

Figure 6.4: CUDA kernel with unrolling the granularity loop for matrix multiplication

observe that each thread, based on the value of $B_0$, uses 38 or 34 registers, which are more than the default allowance, that is, 32 per thread, set by the compiler. In this case, register spilling [95] occurs, which causes the data to be accessed in the L2 cache, and slows down the execution time. Meanwhile, the nvcc [38] compiler for CUDA programs allows the programmer to set a number for the number of registers that each thread can use, as long as this number does not exceed the hardware limit (which is 64 on the NVIDIA Tesla M2050). Thus, we pass
a compilation flag --maxrregcount=40 to allow each thread to use 40 registers. In the end, we obtain a significant improvement in performance.

Table 6.4: For input matrices of order $2^{10}$, speedup factors of the matrix multiplication kernel unrolling the copy-in, computation and copy-out phases with a compilation flag --maxrregcount=40

<table>
<thead>
<tr>
<th>Thread-block \ Granularity</th>
<th>2</th>
<th>4</th>
<th>Resource usage</th>
</tr>
</thead>
<tbody>
<tr>
<td>(16, 4)</td>
<td>117.032</td>
<td>151.667</td>
<td>38 registers, 2112 bytes of shared memory</td>
</tr>
<tr>
<td>(32, 4)</td>
<td>151.555</td>
<td><strong>184.812</strong></td>
<td></td>
</tr>
<tr>
<td>(64, 4)</td>
<td>132.588</td>
<td>171.805</td>
<td></td>
</tr>
<tr>
<td>(8, 8)</td>
<td>145.972</td>
<td>178.111</td>
<td></td>
</tr>
<tr>
<td>(16, 8)</td>
<td>196.625</td>
<td><strong>230.870</strong></td>
<td>34 registers, 2304 bytes of shared memory</td>
</tr>
<tr>
<td>(32, 8)</td>
<td>193.753</td>
<td>226.934</td>
<td></td>
</tr>
<tr>
<td>(64, 8)</td>
<td>111.214</td>
<td>169.528</td>
<td></td>
</tr>
</tbody>
</table>

Moreover, we realize that we compared our kernel code with the naive, serial matrix multiplication (shown in Figure 5.13 of Chapter 5), which lacks good data locality. Thus, using a blocking strategy, we implement the version of the serial C code as shown in Figure 6.5, which has a nearly optimal cache complexity. Finally, we collect speedup factors for the two kernels generated by PPCG and MetaFork against this latter C code. Of course, the MetaFork-generated kernel is the one described above. In particular, a post-processing phase is required for unrolling the granularity loop and using the local memory. Experimental results are collected in Table 6.5. For the best case, the MetaFork-generated kernel outperforms by a factor of 1.7 (resp. 1.9) the PPCG-generated kernel for matrices of order $2^{10}$ (resp. $2^{11}$).

```c
for (int i = 0; i < n; i += 32)
for (int j = 0; j < n; j += 32)
for (int k = 0; k < n; k += 32)
    for (int i0 = i; i0 < min(i + 32, n); i0++)
        for (int j0 = j; j0 < min(j + 32, n); j0++)
            for (int k0 = k; k0 < min(k + 32, n); k0++)
                c[i0][j0] += a[i0][k0] * b[k0][j0];
```

Figure 6.5: The serial C code with good data locality for matrix multiplication

### 6.2 Experimentation

In this section, we experiment with the granularity loop on the MetaFork code of five simple test cases: array reversal (Figure 6.6, Table 6.6), 1D Jacobi (Figure 6.7, Table 6.7), matrix addition (Figure 6.8, Table 6.8), matrix transposition (Figure 6.9, Table 6.9) and matrix vector multiplication (Figure 6.10, Table 6.10). For each test case, we compare the kernel code with the granularity loop against the kernel code without the granularity loop, that is, the code automatically generated in Chapter 5.

For matrix addition, the kernel uses the global memory for the input and output matrices. For matrix vector multiplication, we generate a local memory counterpart for the output vector.
Table 6.5: Speedup factors obtained with kernels generated by PPCG and MetaFork with post-processing, respectively, w.r.t. the serial C code with good data locality for matrix multiplication

<table>
<thead>
<tr>
<th>Speedup (kernel)</th>
<th>Input size</th>
</tr>
</thead>
<tbody>
<tr>
<td>Thread-block size</td>
<td>$2^{10} \times 2^{10}$</td>
</tr>
<tr>
<td>PPCG</td>
<td>109</td>
</tr>
<tr>
<td><strong>MetaFork with post-processing</strong></td>
<td></td>
</tr>
<tr>
<td>Granularity</td>
<td>2</td>
</tr>
<tr>
<td>(16, 32)</td>
<td>95</td>
</tr>
<tr>
<td>(32, 4)</td>
<td>128</td>
</tr>
<tr>
<td>(64, 4)</td>
<td>111</td>
</tr>
<tr>
<td>(8, 8)</td>
<td>131</td>
</tr>
<tr>
<td>(16, 8)</td>
<td>164</td>
</tr>
<tr>
<td>(32, 8)</td>
<td>163</td>
</tr>
<tr>
<td>(64, 8)</td>
<td>94</td>
</tr>
</tbody>
</table>

and two shared memory counterparts for both the input matrix and the input vector. For the other test cases, we generate shared memory counterparts for the input vectors or matrices and continue to use the global memory for the output. In the case that vectors or matrices are required to copy from/to the arrays allocated in the shared or local memory, post-processing is required. Figures 6.6, 6.7, 6.8, 6.9 and 6.10 show the MetaFork code and its kernel code for each test case. Tables 6.6, 6.7, 6.8, 6.9 and 6.10 collect the speedup factors for various values of one thread-block dimension and its granularity of threads.

**Array reversal.** For the input vector of length $2^{25}$, the kernel code generated in Chapter 5 achieves the best speedup factor of 22.965 for the thread-block format of size 256, while for the same thread-block format but with the granularity 4, the kernel with the granularity loop performs slightly better.

Table 6.6: Speedup factors of reversing a one-dimensional array for input vector of length $2^{25}$

<table>
<thead>
<tr>
<th>Thread-block \ Granularity</th>
<th>2</th>
<th>4</th>
<th>8</th>
</tr>
</thead>
<tbody>
<tr>
<td>16</td>
<td>5.239</td>
<td>5.561</td>
<td>5.743</td>
</tr>
<tr>
<td>32</td>
<td>7.867</td>
<td>7.959</td>
<td>8.176</td>
</tr>
<tr>
<td>128</td>
<td>20.354</td>
<td>20.872</td>
<td>21.036</td>
</tr>
<tr>
<td>256</td>
<td>23.554</td>
<td>24.511</td>
<td></td>
</tr>
</tbody>
</table>

**1D Jacobi.** For the input vector of length $2^{15} + 2$, the parametric kernels generated in Chapter 5 achieve the best speedup factor of 7.309 with the thread-block format of size 64, while for the thread-block format of size 128 and the granularity 2, the kernels with the granularity loop perform slightly better.

**Matrix addition.** For the input matrix of order $2^{12}$, the kernel code generated in Chapter 5 achieves the best speedup factor of 56.650 for the thread-block format of size $32 \times 8$, while
// Array with of size N  
// Program parameters: B, s  
int ub_v = N / B, ub_u = B / s;  

meta_schedule  
  meta_for (int v = 0; v < ub_v; v++)  
    meta_for (int u = 0; u < ub_u; u++)  
      for (int w = 0; w < s; ++w)  
        int x = v * B + w * ub_u + u;  
        int y = N - 1 - x;  
        Out[y] = In[x];  
  
}

global void kernel0(int *In, int *Out, int N, int ub_v,  
                    int ub_u, int s, int B) {  
  int b0 = blockIdx.x;  
  int t0 = threadIdx.x;  
  int private_x;  
  int private_y;  
  _shared_ int shared_In[BLOCK_0]; // BLOCK_0 = B  
  for (int c0 = b0; c0 < ub_v; c0 += 32768) {  
    for (int c2 = 0; c2 < s; c2 += 1) {  
      private_x = (c0) * (B) + (c2) * (ub_u) + (t0);  
      if (N >= private_x + 1)  
        shared_In[(c2) * (ub_u) + t0] = In[private_x];  
        _syncthreads();  
      private_y = (((N) - 1) - private_x);  
      Out[private_y] = shared_In[(c2) * (ub_u) + t0];  
    }  
    _syncthreads();  
  }

Figure 6.6: The MetaFork code and its kernel code with the granularity loop for reversing a one-dimensional array

Table 6.7: Speedup factors of 1D Jacobi for time iteration 4 and input vector of length $2^{15}+2$

<table>
<thead>
<tr>
<th>Thread-block \ Granularity</th>
<th>2</th>
<th>4</th>
<th>8</th>
</tr>
</thead>
<tbody>
<tr>
<td>16</td>
<td>3.340</td>
<td>4.357</td>
<td>4.975</td>
</tr>
<tr>
<td>32</td>
<td>4.785</td>
<td>5.252</td>
<td>5.206</td>
</tr>
<tr>
<td>64</td>
<td>5.927</td>
<td>6.264</td>
<td>6.412</td>
</tr>
<tr>
<td>128</td>
<td><strong>10.400</strong></td>
<td>8.952</td>
<td>5.793</td>
</tr>
<tr>
<td>256</td>
<td>6.859</td>
<td>6.246</td>
<td></td>
</tr>
</tbody>
</table>

for the thread-block format of size 64 x 8 and the granularity 2, the kernel with the granularity loop performs almost the same.

Table 6.8: Speedup factors of matrix addition for input matrix of order $2^{12}$

<table>
<thead>
<tr>
<th>Thread-block \ Granularity</th>
<th>2</th>
<th>4</th>
<th>8</th>
</tr>
</thead>
<tbody>
<tr>
<td>(4, 8)</td>
<td>34.144</td>
<td>37.196</td>
<td>34.187</td>
</tr>
<tr>
<td>(8, 8)</td>
<td>45.906</td>
<td>43.287</td>
<td>41.956</td>
</tr>
<tr>
<td>(16, 8)</td>
<td>55.219</td>
<td>52.543</td>
<td>51.304</td>
</tr>
<tr>
<td>(32, 8)</td>
<td><strong>56.815</strong></td>
<td>55.843</td>
<td>54.402</td>
</tr>
<tr>
<td>(64, 8)</td>
<td>39.128</td>
<td>44.819</td>
<td></td>
</tr>
</tbody>
</table>

**Matrix transpose.** For the input matrix of order $2^{14}$, the kernel code generated in Chapter 5 achieves the best speedup factor of 93.051 for the thread-block format of size 8 x 32, while for the same thread-block format but with the granularity 2, the kernel with the granularity loop outperforms those two kernels automatically generated by PPCG and MetaFork.

**Matrix vector multiplication.** For the input matrix of order $2^{13}$ and the input vector of length $2^{13}$, the kernel code generated in Chapter 5 achieves the best speedup factor of 10.340 for the thread-block format of size 32. In this test case, since we use the local memory for the output vector, a constant is used as the upper bound of the granularity loop so as to allow loop
6.2. Experimentation

// Array with of size N + 2
// Program parameters: B, s
int ub_v = (N - 2) / B, ub_u = B / s;

meta_schedule {
    for (int t = 0; t < T; ++t) {
        for (int v = 0; v < ub_v; v++)
            for (int u = 0; u < ub_u; u++)
                int p = v * B + i * ub_u + u;
                b[p+1] = (a[p]+a[p+1]+a[p+2])/3;
        for (int v = 0; v < ub_v; v++)
            for (int u = 0; u < B; u++)
                int w = v * B + u;
                a[w+1] = b[w+1];
    }
}

Figure 6.7: The MetaFork code and its kernel code with the granularity loop for 1D Jacobi

// n * n matrices
// Program parameters: B0, ub_u1, s
int dim0 = n/B0, dim1 = n/B1, ub_u1 = B1/s;

meta_schedule {
    for (int v0 = 0; v0 < dim0; v0++)
        for (int v1 = 0; v1 < dim1; v1++)
            for (int u0 = 0; u0 < B0; u0++)
                for (int i = 0; i < s; ++i)
                    int j = v1 * B1 + w*ub_u1 + u1;
                    c[i][j] = a[i][j] + b[i][j];
    }
}

Figure 6.8: The MetaFork code and its kernel code with the granularity loop for matrix addition

unrolling for the copy-in, computation and copy-out phases. For the thread-block format of size 32 and the granularity 2, this kernel uses 22 registers and 8320 bytes of shared memory, while for the same thread-block format but without the granularity loop, the kernel automatically
Table 6.9: Speedup factors of matrix transpose for input matrix of order $2^{14}$

<table>
<thead>
<tr>
<th>Thread-block \ Granularity</th>
<th>2</th>
<th>4</th>
<th>8</th>
</tr>
</thead>
<tbody>
<tr>
<td>(4, 32)</td>
<td>103.281</td>
<td>96.284</td>
<td>75.211</td>
</tr>
<tr>
<td>(8, 32)</td>
<td><strong>111.971</strong></td>
<td>90.625</td>
<td>85.422</td>
</tr>
<tr>
<td>(16, 32)</td>
<td>78.476</td>
<td>68.894</td>
<td>48.822</td>
</tr>
<tr>
<td>(32, 32)</td>
<td>45.084</td>
<td>46.425</td>
<td>32.824</td>
</tr>
</tbody>
</table>

Figure 6.9: The MetaFork code and its kernel code with the granularity loop for matrix transpose.

generated by MetaFork uses 29 registers and 4352 bytes of shared memory. However, the kernel with the granularity loop performs worse.

Table 6.10: Speedup factors of matrix vector multiplication for input matrix of order $2^{13}$ and input vector of length $2^{13}$ (An error indicates that the total amount of required shared memory exceeds the hardware limit.)

<table>
<thead>
<tr>
<th>Thread-block \ Granularity</th>
<th>2</th>
<th>4</th>
<th>8</th>
</tr>
</thead>
<tbody>
<tr>
<td>8</td>
<td>2.975</td>
<td>2.677</td>
<td>1.974</td>
</tr>
<tr>
<td>16</td>
<td>4.281</td>
<td>3.424</td>
<td>3.377</td>
</tr>
<tr>
<td>32</td>
<td><strong>5.477</strong></td>
<td>1.916</td>
<td>0.988</td>
</tr>
<tr>
<td>64</td>
<td>2.286</td>
<td>Error</td>
<td>Error</td>
</tr>
</tbody>
</table>

6.3 Conclusion

We have experimented with thread granularity control for six test cases. For two of these test cases, matrix multiplication and matrix vector multiplication, we use the local memory for the
output matrix or vector in order to expose ILP by helping the compiler to apply loop unrolling. We believe that these techniques could help the performance of CUDA kernels. Although matrix transposition and matrix multiplication show a significant improvement, matrix vector multiplication performs worse, and other test cases behave slightly better or almost the same as the original kernel without the granularity loop control. The question whether we should apply those techniques in process of automatic code generation remains unanswered. This leads to the next chapter where we discuss an algorithm to generate a case distinction depending on available hardware resources.

However, adding a granularity loop in the MetaFork code violates those assumptions listed in Appendix C. Thus, for using the shared memory, post-processing is required to obtain correct index expressions of the shared memory counterparts. This manual modification is difficult and painful, especially for non-CUDA experts. To fully support any non-linear expressions in the current MetaFork-to-CUDA code generator, simply hacking into PPCG source code is not enough to solve the issue. We leave it to future work.

Moreover, our automatic code generator described in Chapter 5 does not support using the local memory. We could extend our code generator to allow the user to choose whether to use the local memory in the kernel code. Alternatively, we could automatically detect the situations when the local memory should be used. To be more specific, based on those test cases that we experienced, we can use the local memory for the output array when each element in this array is accessed more than once by each thread.

```c
void kernel0(int *a, int *b, int *c, int N, int ub_v, int ub_u, int s, int B) {
    int b0 = blockIdx.x;
    int t0 = threadIdx.x;
    // BLOCK = B0, STRIDE = s
    __shared__ int shared_a[BLOCK][BLOCK/STRIDE];
    __shared__ int shared_b[BLOCK/STRIDE];
    int private_c[STRIDE];

    for (int c0 = b0; c0 < ub_v; c0 += 32768) {
        for (int c4 = 0; c4 < STRIDE; c4 += 1)
            private_c[c4] = c[ub_u * c4 + t0];
        for (int c1 = 0; c1 < N / ub_u; c1 += 1) {
            for (int c3 = 0; c3 < ub_u; c3 += 1)
                shared_a[c4 * ub_u + c3][t0] =
                    a[(ub_u * c4 + c0 * B + c3) * N + (ub_u * c1 + t0)];
            shared_b[t0] = b[t0 + ub_u * c1];
            __syncthreads();
            for (int c3 = 0; c3 < ub_u; c3 += 1)
                for (int c4 = 0; c4 < STRIDE; c4 += 1)
                    private_c[c4] +=
                        (shared_a[c4 * ub_u + t0][c3] * shared_b[c3]);
            __syncthreads();
        }
    }
}
```
Chapter 7
Towards Comprehensive Parametric CUDA Kernel Generation

In Chapter 5, we demonstrated that, from an annotated C code, it was possible to generate CUDA kernels that depend on program parameters considered unknown at compile-time. Our experimental results in Chapters 5 and 6 suggest that those parametric CUDA kernels could help with increasing portability and performance of CUDA code.

In the present chapter, we enhance this strategy as follows. First, we propose an algorithm for comprehensive optimization allowing us to optimize C code (and in particular CUDA code) depending on unknown machine and program parameters. Then, we use this algorithm to generate optimized parametric CUDA kernels, in the form of a case distinction based on the possible values of the machine and program parameters. We call comprehensive parametric CUDA kernels the resulting CUDA kernels, see Section 7.2.

In broad terms, this is a decision tree, where each edge holds a Boolean expression, given by a conjunction of polynomial constraints, and each leaf is either a CUDA kernel or the symbol $\emptyset$, such that for each leaf $K$, with $K \neq \emptyset$, we have:

1. $K$ works correctly under the conjunction of the Boolean expressions located between the root node and the leaf, and
2. $K$ is semantically equivalent to a common input annotated C code $P$.

In each Boolean expression, the unknown variables represent machine parameters (like hardware resource limits), program parameters (like dimension sizes of thread-blocks) or data parameters (like input data size). The symbol $\emptyset$ is used to denote a situation (in fact, value ranges for the machine and program parameters) where no CUDA kernel equivalent to $P$ is provided.

The intention, with the concept of comprehensive parametric CUDA kernels, is to automatically generate optimized CUDA kernels from an annotated C code without knowing the numerical values of some, or all, of the machine and program parameters. This naturally yields a case distinction depending on the values of those parameters. Indeed, some optimization techniques (like loop unrolling) can only be applied when enough computing resources are available, while other optimization techniques (like common sub-expression elimination) can be applied to reduce computing resource consumption. These case distinctions can be handled by techniques from symbolic computation, for which software libraries are available, in particular, the RegularChains library freely available at [www.regularchains.org](http://www.regularchains.org).

Let us illustrate, with an example, the notion of comprehensive parametric CUDA kernels,
along with a procedure to generate them. For computing the sum of two matrices \(a\) and \(b\) of order \(N\), our input is the meta_for-loop nest within the meta_schedule statement on the right-hand portion of Figure 7.1, whereas the serial code without tiling is provided on the left-hand portion of Figure 7.1.

```c
for (int i = 0; i < N; i++)
    for (int j = 0; j < N; j++)
        c[i][j] = a[i][j] + b[i][j];
```

```c
int dim0 = N/B0, dim1 = N/(2*B1);
meta_schedule {
    meta_for (int v = 0; v < dim0; v++)
        meta_for (int p = 0; p < dim1; p++)
            meta_for (int u = 0; u < B0; u++)
                meta_for (int q = 0; q < B1; q++)
                {
                    int i = v * B0 + u;
                    int j = p * B1 + q;
                    if (i < N && j < N/2) {
                        c[i][j] = a[i][j] + b[i][j];
                        c[i][j+N/2] = a[i][j+N/2] + b[i][j+N/2];
                    }
                }
}
```

(a) Before tiling, the C program    (b) After tiling, the MetaFork program

Figure 7.1: Matrix addition written in C (the left-hand portion) and in MetaFork (the right-hand portion) with a meta_for loop nest, respectively.

We make the following simplistic assumptions for the translation of this meta_for-loop nest to a CUDA program.

1. The target machine has two parameters: the maximum number \(R_1\) of registers per thread, and the maximum number \(R_2\) of threads per thread-block; moreover, all other hardware limits are ignored.
2. The generated kernels depend on two program parameters, \(B_0\) and \(B_1\), which define the format of a 2D thread-block.
3. The optimization strategy (w.r.t. register usage per thread) consists in reducing the work per thread via removing the 2-way loop unrolling.

The possible comprehensive parametric CUDA kernels are given by the pairs \((C_1, K_1)\) and \((C_2, K_2)\), where \(C_1, C_2\) are two sets of algebraic constraints on the machine and program parameters and \(K_1, K_2\) are two CUDA kernels that are optimized under the constraints, respectively, given by \(C_1, C_2\), see Figure 7.2. The following computational steps yield the pairs \((C_1, K_1)\) and \((C_2, K_2)\). (S1) Tiling techniques, based on quantifier elimination (QE), are applied to the meta_for loop nest of Figure 7.1 in order to decompose the matrices into tiles of format \(B_0 \times B_1\), see [19] for details.

(S2) The tiled MetaFork code is mapped to an intermediate representation (IR) say that of LLVM\(^1\) or alternatively, to PTX\(^2\) code.

---

1 Quoting Wikipedia: “The LLVM compiler infrastructure project (formerly Low Level Virtual Machine) is a framework for developing compiler front ends and back ends”.
2 The Parallel Thread Execution (PTX) is the pseudo-assembly language to which CUDA programs are
(S3) Using this IR (or PTX) code, one can estimate the number of registers that a thread requires; thus, using LLVM IR on this example, we obtain an estimate of 14.  
(S4) Next, we apply the optimization strategy, yielding a new IR (or PTX) code, for which register pressure reduces to 10. Since no other optimization techniques are considered, the procedure stops with the result shown in Figure 7.2.  
Based on the above steps, Figure 7.3 shows the decision tree for generating these two pairs, each of them consisting of a system of polynomial constraints and a CUDA kernel for matrix addition. Note that, strictly speaking, the kernels $K_1$ and $K_2$ on Figure 7.2 should be given by PTX code. But for simplicity, we are presenting them by the CUDA code counterpart.  

**Figure 7.2: Comprehensive translation of MetaFork code to two kernels for matrix addition**

One can observe that loop unrolling is applied to $K_1$ so as to increase arithmetic intensity. This code transformation increases register pressure, which is possible under the constraints of $C_1$ but not under those of $C_2$.  
In general, to achieve a comprehensive translation of the annotated C program $\mathcal{P}$ into CUDA kernels, one could be tempted to proceed as follows:  
(S1) Perform a comprehensive optimization of $\mathcal{P}$, as a MetaFork program, by applying the comprehensive optimization algorithm demonstrated in Section 7.1.  
(S2) Apply the MetaFork-to-CUDA code generator introduced in Chapter 5 to each MetaFork program generated in the first step.  
However, since the system of polynomial constraints associated with each optimized MetaFork program is determined by an IR representation of that program, this system would not be accurate for the CUDA code generated by a source-to-source MetaFork-to-CUDA code generator.  

compiled by NVIDIA’s nvcc compiler. PTX code can also be generated from (enhanced) LLVM IR, using nvptx back-end [68], following the work of [99].
7.1 Comprehensive optimization

We consider a code fragment written in the C language or in one of its linguistic extensions targeting a computer device, which can be, for instance, a hardware accelerator or a desktop CPU. We assume that some, or all, of the hardware characteristics of this device are unknown at compile time. However, we would like to optimize our input code fragment w.r.t prescribed resource counters (e.g., memory usage) and performance counters (e.g., clock-cycle per instruction). To this end, we treat the hardware characteristics of this device as symbols and generate polynomial constraints (with those symbols as indeterminate variables) ensuring when such...
and such code transformation is valid.

Section 7.1.1 states the hypotheses made on the input code fragment. Section 7.1.2 specifies the notations for the hardware characteristics of the targeted device. In Section 7.1.3 we describes the evaluation of resource and performance counters. In Section 7.1.4 we define the optimization strategies that can reduce resource counters or increase performance counters. Section 7.1.5 formally gives the definition of comprehensive optimization of an input code fragment. Section 7.1.6 specifies the data structures that are used in our algorithm. Finally, in Section 7.1.7 we demonstrate our algorithm for comprehensive optimization.

7.1.1 Hypotheses on the input code fragment

We consider a sequence $S$ of statements from the C programming language and introduce the following.

**Definition 1** We call parameter of $S$ any scalar variable that is
(i) read in $S$ at least once and
(ii) never written in $S$.

We call data of $S$ any non-scalar variable (e.g. array) that is not initialized but possibly overwritten within $S$. If a parameter of $S$ gives a dimension size of a data of $S$, then this parameter is called a data parameter; otherwise, it is simply called a program parameter.

**Notation 1** We denote by $D_1, \ldots, D_u$ and $E_1, \ldots, E_v$ the data parameters and program parameters of $S$, respectively.

**Hypothesis 3** We make the following assumptions on $S$.

(H1) All parameters are assumed to be non-negative integers.

(H2) We assume that $S$ can be viewed as the body of a valid C function having the parameters and data of $S$ as unique arguments.

**Example 3** $S$ can be the body of a kernel function in CUDA. Recall that the kernel code for computing matrix vector multiplication in Figure 5.12 of Chapter 5. This kernel code multiplies a square matrix $a$ of order $n$ with a vector $b$ of length $n$ and stores the result to a vector $c$ of length $n$. We note that $a$, $b$, and $c$ are the data, and that $n$ is the data parameter. Moreover, the grid and thread-block dimensions of this kernel are specified as dim and B, respectively, which are then the program parameters.

7.1.2 Hardware resource limits and performance measures

We denote by $R_1, \ldots, R_s$ the hardware resource limits of the targeted hardware device. Examples of these quantities for the NVIDIA Kepler micro-architecture are:
- the maximum number of registers to be allocated per thread,
- the maximum number of shared memory words to be allocated per thread-block,
- the maximum number of threads in a thread-block.

We denote by $P_1, \ldots, P_t$ the performance measures of a program running on the device. These are dimensionless quantities typically defined as percentages. Examples of these quantities for the NVIDIA Kepler micro-architecture are:
7.1. Comprehensive optimization

- the ratio of the actual to the maximum number of words that can be read or written per unit of time from the global memory,
- the ratio of the actual to the maximum number of floating point operations that can be performed per unit of time by all streaming processors (SMs),
- the SM occupancy, that is, the ratio of active warps to the maximum number of active warps,
- the cache hit rate in an SM.

For a given hardware device, \( R_1, \ldots, R_s \) are positive integers, and each of them is the maximum value of a hardware resource. Meanwhile, \( P_1, \ldots, P_t \) are rational numbers between 0 and 1. However, for the purpose of writing code portable across a variety of devices with similar characteristics, the quantities \( R_1, \ldots, R_s \) and \( P_1, \ldots, P_t \) will be treated as unknown and independent variables. These hardware resource limits and performance measures will be called the machine parameters.

Each function \( K \) (and, in particular, our input code fragment \( S \)) written in the C language for the targeted hardware device has resource counters \( r_1, \ldots, r_s \) and performance counters \( p_1, \ldots, p_t \) corresponding, respectively, to \( R_1, \ldots, R_s \) and \( P_1, \ldots, P_t \). In other words, the quantities \( r_1, \ldots, r_s \) are the amounts of resources, corresponding to \( R_1, \ldots, R_s \), respectively, that \( K \) requires for executing. Similarly, the quantities \( p_1, \ldots, p_t \) are the performance measures, corresponding to \( P_1, \ldots, P_t \), respectively, that \( K \) exhibits when executing. Therefore, the inequalities \( 0 \leq r_1 \leq R_1, \ldots, 0 \leq r_s \leq R_s \) must hold for the function \( K \) to execute correctly. Similarly, \( 0 \leq p_1 \leq 1, \ldots, 0 \leq p_t \leq 1 \) are satisfied by the definition of the performance measures.

**Remark 1** We note that \( r_1, \ldots, r_s, p_1, \ldots, p_t \) may be numerical values, which we can assume to be non-negative rational numbers. This will be the case, for instance, for the minimum number of registers required per thread in a thread-block. The resource counters \( r_1, \ldots, r_s \) may also be polynomial expressions whose indeterminate variables can be program parameters (like the dimension sizes of a thread-block or grid) or data parameters (like the input data sizes). Meanwhile, the performance counters \( p_1, \ldots, p_t \) may further depend on the hardware resource limits (like the maximum number of active warps supported by an SM). To summarize, we observe that \( r_1, \ldots, r_s \) are polynomials in \( \mathbb{Q}[D_1, \ldots, D_u, E_1, \ldots, E_d] \) and \( p_1, \ldots, p_t \) are rational functions where numerators and denominators are in \( \mathbb{Q}[D_1, \ldots, D_u, E_1, \ldots, E_d, R_1, \ldots, R_s] \). Moreover, we can assume that the denominators of those rational functions are positive.

**Example 4** For computing the product of a dense square matrix of order \( N \) by a dense vector of length \( N \), consider the serial C code on the left-hand portion of Figure 7.4 and the corresponding MetaFork code on the right-hand portion of Figure 7.4. The meta_schedule statement yields the CUDA kernel shown in Figure 5.12 of Chapter 5. The grid of this kernel is one-dimensional of size \( \text{dim} \), and its thread-blocks are also one-dimensional, each counting \( B \) threads.

Observe that, in the process of generating this CUDA kernel, some optimization techniques (like using the shared memory for arrays \( a, b \) and \( c \) in Figure 5.12 of Chapter 5) are applied, while other optimization techniques remain to be applied (like the granularity of threads as used in Figure 6.10 of Chapter 6). If \( R_2 \) and \( R_3 \) are two machine parameters that define, respectively, the maximum number of threads in a thread-block and the maximum number of shared memory words per thread-block, then the following constraints must hold:

\[
B \leq R_2 \quad \text{and} \quad B^2 + 2B \leq R_3.
\]
for (int p = 0; p < N; p++)
    for (int q = 0; q < N; q++)
        c[p] += a[p][q] * b[q];

int dim = N / B;

meta_schedule {
    meta_for (int v = 0; v < dim; v++)
        for (int i = 0; i < dim; ++i)
            meta_for (int u = 0; u < B; u++)
                for (int j = 0; j < B; ++j) {
                    int p = v * B + u;
                    int q = i * B + j;
                    c[p] += a[p][q] * b[q];
                }
}

(a) Before tiling, the C program
(b) After tiling, the MetaFork program

Figure 7.4: Matrix vector multiplication written in C (the left-hand portion) and in MetaFork (the right-hand portion), respectively.

Note that, in Figure 5.12 of Chapter 5, each thread-block allocates a unit size $B^2$ of shared memory words for the array $a$, and a unit size $B$ of shared memory words, respectively, for each of arrays $b$ and $c$. Thus, this leads to the inequality $B^2 + 2B \leq R_3$.

### 7.1.3 Evaluation of resource and performance counters

Let $G_C(S)$ be the control flow graph (CFG) [123] of $S$. Hence, the statements in the basic blocks of $G_C(S)$ are C statements, and we call such a CFG the source CFG. We also map $S$ to an intermediate representation, which, itself, is encoded in the form of a CFG, denoted by $G_L(S)$, and we call it the IR CFG. Here, we refer to the landmark textbook [2] for the notion of the control flow graph and that of intermediate representation.

We observe that $S$ can trivially be reconstructed from $G_C(S)$; hence, the knowledge of $S$ and that of $G_C(S)$ can be regarded as equivalent. In contrast, $G_L(S)$ depends not only on $S$ but also on the optimization strategies that are applied to the IR of $S$.

Equipped with $G_C(S)$ and $G_L(S)$, we assume that we can estimate each of the resource counters $r_1, \ldots, r_s$ (resp. performance counters $p_1, \ldots, p_t$) by applying functions $f_1, \ldots, f_s$ (resp. $g_1, \ldots, g_t$) to either $G_C(S)$ or $G_L(S)$. We call $f_1, \ldots, f_s$ (resp. $g_1, \ldots, g_t$) the resource (resp. performance) evaluation functions.

For instance, when $S$ is the body of a CUDA kernel and $S$ reads (resp. writes) a given array, computing the total amount of elements read (resp. written) by one thread-block can be determined from $G_C(S)$. Meanwhile, computing the minimum number of registers to be allocated to a thread executing $S$ requires the knowledge of $G_L(S)$.

### 7.1.4 Optimization strategies

In order to reduce the consumption of hardware resources and increase performance counters, we assume that we have $w$ optimization procedures $O_1, \ldots, O_w$, each of them mapping either a source CFG to another source CFG, or an IR CFG to another IR CFG. Of course, we assume the code transformations performed by $O_1, \ldots, O_w$ preserve semantics.
We associate each resource counter \( r_i \), for \( i = 1 \cdots s \), with a non-empty subset \( \sigma(r_i) \) of \( \{O_1, \ldots, O_w\} \), such that we have

\[
f_i(O(S)) \leq f_i(S) \quad \text{for } O \in \sigma(r_i).
\] (7.1)

Hence, \( \sigma(r_i) \) is a subset of the optimization strategies among \( O_1, \ldots, O_w \) that have the potential to reduce \( r_i \). Of course, the intention is that for at least one \( O \in \sigma(r_i) \), we have \( f_i(O(S)) < f_i(S) \).

A reason for not finding such \( O \) would be that \( S \) cannot be further optimized w.r.t. \( r_i \). We also make a natural idempotence assumption:

\[
f_i(O(O(S))) = f_i(O(S)) \quad \text{for } O \in \sigma(r_i).
\] (7.2)

Similarly, we associate each performance counter \( p_i \), for \( i = 1 \cdots t \), with a non-empty subset \( \sigma(p_i) \) of \( \{O_1, \ldots, O_u\} \), such that we have

\[
g_i(O(S)) \geq g_i(S) \quad \text{and} \quad g_i(O(O(S))) = g_i(O(S)) \quad \text{for } O \in \sigma(p_i).
\] (7.3)

Hence, \( \sigma(p_i) \) is a subset of the optimization strategies among \( O_1, \ldots, O_u \) that have the potential to increase \( p_i \). The intention is, again, that for at least one \( O \in \sigma(p_i) \), we have \( g_i(O(S)) > g_i(S) \).

### 7.1.5 Comprehensive optimization

Let \( C_1, \ldots, C_e \) be semi-algebraic systems with \( P_1, \ldots, P_t, R_1, \ldots, R_s, D_1, \ldots, D_u, E_1, \ldots, E_v \) as indeterminate variables. Let \( S_1, \ldots, S_e \) be fragments of C programs such that the parameters of each of them are among \( D_1, \ldots, D_u, E_1, \ldots, E_v \).

**Definition 2** We say that the sequence of pairs \( (C_1, S_1), \ldots, (C_e, S_e) \) is a comprehensive optimization of \( S \) w.r.t.

- the resource evaluation functions \( f_1, \ldots, f_s \),
- the performance evaluation functions \( g_1, \ldots, g_t \), and
- the optimization strategies \( O_1, \ldots, O_w \)

if the following conditions hold:

(i) [constraint soundness] Each of the semi-algebraic systems \( C_1, \ldots, C_e \) is consistent, that is, admits at least one real solution.

(ii) [code soundness] For all real values \( h_1, \ldots, h_t, x_1, \ldots, x_s, y_1, \ldots, y_{1t}, z_1, \ldots, z_v \) of \( P_1, \ldots, P_t, R_1, \ldots, R_s, D_1, \ldots, D_u, E_1, \ldots, E_v \) respectively, for all \( i \in \{1, \ldots, e\} \) such that \( (h_1, \ldots, h_t, x_1, \ldots, x_s, y_1, \ldots, y_{1t}, z_1, \ldots, z_v) \) is a solution of \( C_i \), then the code fragment \( S_i \) produces the same output as \( S \) on any data that makes \( S \) execute correctly.

(iii) [coverage] For all real values \( y_1, \ldots, y_{1t}, z_1, \ldots, z_v \) of \( D_1, \ldots, D_u, E_1, \ldots, E_v \) respectively, there exist \( i \in \{1, \ldots, e\} \) and real values \( h_1, \ldots, h_t, x_1, \ldots, x_s \) of \( P_1, \ldots, P_t, R_1, \ldots, R_s \) such that \( (h_1, \ldots, h_t, x_1, \ldots, x_s, y_1, \ldots, y_{1t}, z_1, \ldots, z_v) \) is a solution of \( C_i \) and \( S_i \) produces the same output as \( S \) on any data that makes \( S \) execute correctly.

(iv) [optimality] For every \( i \in \{1, \ldots, s\} \) (resp. \( \{1, \ldots, t\} \)), there exists \( \ell \in \{1, \ldots, e\} \) such that for all \( O \in \sigma(r_i) \) (resp. \( \sigma(p_i) \)) we have \( f_i(O(S_i)) = f_i(S_i) \) (resp. \( g_i(O(S_i)) = g_i(S_i) \)).

To summarize Definition[2] in non technical terms:
- Condition (i) states that each system of constraints is meaningful.
- Condition (ii) states that as long as the machine, program and data parameters satisfy $C_i$, the code fragment $S_i$ produces the same output as $S$ on whichever data that makes $S$ execute correctly.
- Condition (iii) states that as long as $S$ executes correctly on a given set of parameters and data, there exists a code fragment $S', \lambda$ for suitable values of the machine parameters, such that $S_i$ produces the same output as $S$ on that set of parameters and data.
- Condition (iv) states that for each resource counter $r_i$ (performance counter $p_i$), there exists at least one code fragment $S_i$ for which this counter is optimal in the sense that it cannot be further optimized by the optimization strategies from $\sigma(r_i)$ (resp. $\sigma(p_i)$).

### 7.1.6 Data-structures

The algorithm presented in Section 7.1.7 computes a comprehensive optimization of $S$ w.r.t. the evaluation functions $f_1, \ldots, f_i, g_1, \ldots, g_i$ and optimization strategies $O_1, \ldots, O_w$.

Hereafter, we define the main data-structure used during the course of the algorithm. We associate $S$ with what we call a quintuple, denoted by $Q(S)$ and defined as follows:

$$Q(S) = (G_C(S), \lambda(S), \omega(S), \gamma(S), C(S))$$

where

1. $\lambda(S)$ is the sequence of the optimization procedures among $O_1, \ldots, O_w$ that have already been applied to the IR of $S$; hence, $G_C(S)$ together with $\lambda(S)$ defines $G_L(S)$; initially, $\lambda(S)$ is empty.
2. $\omega(S)$ is the sequence of the optimization procedures among $O_1, \ldots, O_w$ that have not been applied so far to either $G_C(S)$ or $G_L(S)$; initially, $\omega(S)$ is $O_1, \ldots, O_w$.
3. $\gamma(S)$ is the sequence of resource and performance counters that remain to be evaluated on $S$; initially, $\gamma(S)$ is $r_1, \ldots, r_s, p_1, \ldots, p_t$.
4. $C(S)$ is the sequence of the constraints (polynomial equations and inequalities) on $P_1, \ldots, P_i, R_1, \ldots, R_s, D_1, \ldots, D_u, E_1, \ldots, E_v$ that have been computed so far; initially, $C(S)$ is $1 \geq P_1 \geq 0, \ldots, 1 \geq P_i \geq 0, R_1 \geq 0, \ldots, R_s \geq 0, D_1 \geq 0, \ldots, D_u \geq 0, E_1 \geq 0, \ldots, E_v \geq 0$.

We say that the quintuple $Q(S)$ is *processed* whenever $\gamma(S)$ is empty; otherwise, we say that the quintuple $Q(S)$ is *in-process*.

**Remark 2** For the above $Q(S)$, each of the sequences $\lambda(S), \omega(S), \gamma(S)$ and $C(S)$ is implemented as a stack in Algorithms 5 and 6. Hence, we need to specify how operations on a sequence are performed on the corresponding stack. Let $s_1, s_2, \ldots, s_N$ be a sequence.
- Popping one element out of this sequence returns $s_1$ and leaves that sequence with $s_2, \ldots, s_N$.
- Pushing an element $t_1$ on $s_1, s_2, \ldots, s_N$ will update that sequence to $t_1, s_1, s_2, \ldots, s_N$.
- Pushing a sequence of elements $t_1, t_2, \ldots, t_M$ on $s_1, s_2, \ldots, s_N$ will update that sequence to $t_M, \ldots, t_2, t_1, s_1, s_2, \ldots, s_N$.

### 7.1.7 The algorithm

Algorithm 5 is the top-level procedure. If its input is a processed quintuple $Q(S)$, then it returns the pair $(G_C(S), \lambda(S))$ (such that, after optimizing $S$ with the optimization strategies in $\lambda(S)$,
one can generate the IR of the optimized S) together with the system of constraints C(S).
Otherwise, Algorithm 5 is called recursively on each quintuple returned by Optimize(Q(S)).

The pseudo-code of the Optimize routine is given by Algorithm 6.

Algorithm 5: ComprehensiveOptimization (Q(S))

Input: The quintuple Q(S)
Output: A comprehensive optimization of S w.r.t. the resource evaluation functions f₁, ..., fₙ, the performance evaluation functions g₁, ..., gₜ and the optimization strategies O₁, ..., Oᵢ₁

1 if γ(S) is empty then
  2 return (⟨Gₖ(S), λ(S), C(S)⟩);
3 The output stack is initially empty;
4 for each Q(S′) ∈ Optimize(Q(S)) do
  5 Push ComprehensiveOptimization(Q(S′) on the output stack;
6 return the output stack;

Remark 3 We make a few observations about Algorithm 6.

(R1) Observe that at Line (5), a deep copy of the input Q(S′) is made, and this copy is called Q(S′′). This duplication allows the computations to fork. Note that at Line (6), Q(S′) is modified.

(R2) In this forking process, we call Q(S′) the accept branch and Q(S′′) the refuse branch. In the former case, the relation 0 ≤ vᵢ ≤ Rᵢ holds thus implying that enough Rᵢ-resources are available for executing the code fragment S′. In the latter case, the relation Rᵢ < vᵢ holds thus implying that not enough Rᵢ-resources are available for executing the code fragment S′′.

(R3) Observe that vᵢ is either a numerical value, a polynomial in Q[D₁, ..., Dᵦ, E₁, ..., Eᵣ] or a rational function where its numerator and denominator are in Q[D₁, ..., Dᵦ, E₁, ..., Eᵣ, R₁, ..., Rₙ].

(R4) At Lines (18-20), a similar forking process occurs. Here again, we call Q(S′) the accept branch and Q(S′′) the refuse branch. In the former case, the relation 0 ≤ vᵢ ≤ Pᵢ implies that the Pᵢ-performance counter may have reached its maximum ratio; hence, no optimization strategies are applied to improve this counter. In the latter case, the relation Pᵢ < vᵢ ≤ 1 holds thus implying that the Pᵢ-performance counter has not reached its maximum value; hence, optimization strategies are applied to improve this counter if such optimization strategies are available. Observe that if this optimization strategy does make the estimated value of Pᵢ larger then an algebraic contradiction would happen and the branch will be discarded.

(R5) Line (30) in Algorithm 6 requires non-trivial computations with polynomial equations and inequalities. The necessary algorithms can be found in [22] and are implemented in the RegularChains library of MAPLE.

Remark 4 We make a few remarks about the handling of algebraic computation during the execution of Algorithm 6.

(R6) Each system of algebraic constraints C is updated by adding a polynomial inequality to it at either Lines (6), (7), (19) or (20). This incremental process can be performed by the RealTriangularize algorithm [22] and implemented in the RegularChains library.
Algorithm 6: Optimize

Input: A quintuple $Q(S')$

Output: A stack of quintuples

1. Initialize an empty stack, called `result`;
2. Take out from $\gamma(S')$ the next resource or performance counter to be evaluated, say $c$;
3. Evaluate $c$ on $S'$ (using the appropriate functions among $f_1, \ldots, f_s, g_1, \ldots, g_t$) thus obtaining a value $v_i$, which can be either a numerical value, a polynomial in $\mathbb{Q}[D_1, \ldots, D_u, E_1, \ldots, E_v, R_1, \ldots, R_s]$;
4. if $c$ is a resource counter $r_i$ then
   - Make a deep copy $Q(S'')$ of $Q(S')$, since we are going to split the computation into two branches: $R_i < v_i$ and $0 \leq v_i \leq R_i$;
   - Add the constraint $0 \leq v_i \leq R_i$ to $C(S')$ and push $Q(S')$ onto `result`;
   - Add the constraint $R_i < v_i$ to $C(S'')$ and search $\omega(S'')$ for an optimization strategy of $\sigma(r_i)$;
   - if no such optimization strategy exists then
     - return `result`;
   - else
     - Apply such an optimization strategy to $Q(S'')$ yielding $Q(S''')$;
     - Remove this optimization strategy from $\omega(S''')$;
     - if this optimization strategy is applied to the IR of $S''$ then
       - Add it to $\lambda(S'')$;
     - Push $r_1, \ldots, r_{i-1}, r_i$ onto $\gamma(S'')$;
     - Make a recursive call to Optimize on $Q(S''')$ and push the returned quintuples onto `result`;
   - if $c$ is a performance counter $p_i$ then
     - Make a deep copy $Q(S'')$ of $Q(S')$, since we are going to split the computation into two branches: $0 \leq v_i \leq P_i$ and $P_i < v_i \leq 1$;
     - Add the constraint $0 \leq v_i \leq P_i$ to $C(S')$ and push $Q(S')$ onto `result`;
     - Add the constraint $P_i < v_i \leq 1$ to $C(S'')$ and search $\omega(S'')$ for an optimization strategy of $\sigma(p_i)$;
     - if no such optimization strategy exists then
       - return `result`;
     - else
       - Apply such an optimization strategy to $Q(S'')$ yielding $Q(S''')$;
       - Remove this optimization strategy from $\omega(S''')$;
       - if this optimization strategy is applied to the IR of $S''$ then
         - Add it to $\lambda(S'')$;
       - Push $r_1, \ldots, r_s, p_i$ onto $\gamma(S''')$;
       - Make a recursive call to Optimize on $Q(S''')$ and push the returned quintuples onto `result`;
5. if $c$ is a performance counter $p_i$ then
   - return `result`;
6. if $c$ is a resource counter $r_i$ then
   - return `result`;
7. Remove from `result` any quintuple with an inconsistent system of constraints;
8. return `result`;
(R7) Each of these inequalities can be either strict (using $>$) or large (using $\leq$); the left-hand side is a polynomial of either $\mathbb{Q}[D_1, \ldots, D_u, E_1, \ldots, E_v]$ or $\mathbb{Q}[D_1, \ldots, D_u, E_1, \ldots, E_v, R_1, \ldots, R_s]$, and the right-hand side is either one of the variables $R_1, \ldots, R_s$ or a polynomial of $\mathbb{Q}[D_1, \ldots, D_u, E_1, \ldots, E_v, R_1, \ldots, R_s]$ times one of the variables $P_1, \ldots, P_t$.

(R8) Because of the recursive calls at Lines (16) and (29) several inequalities involving the same variable among $R_1, \ldots, R_s$, $P_1, \ldots, P_t$ may be added to a given system $C$. As a result, $C$ may become inconsistent. For instance if $10 \leq R_1$ and $R_1 < 10$ are both added to the same system $C$. Note that inconstancy is automatically detected by the RealTriangularize algorithm.

(R9) When using RealTriangularize, variables should be ordered. We choose a variable ordering such that
- any of $P_1, \ldots, P_t$ is greater than any of the other variables,
- any of $R_1, \ldots, R_s$ is greater than any of $D_1, \ldots, D_u, E_1, \ldots, E_v$.

(R10) Then, the RealTriangularize represents the solution of $C$ as the union of the solution sets of finitely many regular semi-algebraic systems. Each such regular semi-algebraic system $\mathcal{T}$ consists of
- polynomial constraints involving $D_1, \ldots, D_u, E_1, \ldots, E_v$ only,
- polynomial constraints involving $D_1, \ldots, D_u, E_1, \ldots, E_v, R_1, \ldots, R_s$ only and of positive degree in at least one of $R_1, \ldots, R_s$,
- constraints that are linear in $P_1, \ldots, P_t$ and polynomial in $D_1, \ldots, D_u, E_1, \ldots, E_v, R_1, \ldots, R_s$.

Let us denote by $\mathcal{T}_{D,E}$, $\mathcal{T}_{D,E,R}$ and $\mathcal{T}_{D,E,R,P}$ these three sets of polynomial constraints, respectively. It follows from the properties of regular semi-algebraic systems that
(a) $\mathcal{T}_{D,E}$ is consistent,
(b) for all real values $y_1, \ldots, y_u, z_1, \ldots, z_v$ of $D_1, \ldots, D_u, E_1, \ldots, E_v$ such that $(y_1, \ldots, y_u, z_1, \ldots, z_v)$ solves $\mathcal{T}_{D,E}$, there exists real values $x_1, \ldots, x_s$ of $R_1, \ldots, R_s$ such that $(y_1, \ldots, y_u, z_1, \ldots, z_v, x_1, \ldots, x_s)$ solves $\mathcal{T}_{D,E,R}$ and real values $h_1, \ldots, h_t$ of $P_1, \ldots, P_t$ such that $(y_1, \ldots, y_u, z_1, \ldots, z_v, x_1, \ldots, x_s, h_1, \ldots, h_t)$ solves $\mathcal{T}_{D,E,R,P}$.

Notation 2 We associate the execution of Algorithm 5 applied to $Q(S)$, with a tree denoted by $\mathcal{T}(Q(S))$ and where both nodes and edges of $\mathcal{T}(Q(S))$ are labelled. We use the same notations as in Algorithm 6. We define $\mathcal{T}(Q(S))$ recursively as follows:
(T1) We label the root of $\mathcal{T}(Q(S))$ with $Q(S)$.
(T2) If $\gamma(S)$ is empty, then $\mathcal{T}(Q(S))$ has no children; otherwise, two cases arise:
(T2.1) If no optimization strategy is to be applied for optimizing the counter $c$, then $\mathcal{T}(Q(S))$ has a single subtree, which is the tree associated with Optimize($Q(S')$) where $Q(S')$ is obtained from $Q(S)$ by augmenting $C(S)$ either with $0 \leq v_i \leq R_i$ if $c$ is a resource counter or with $0 \leq v_i \leq P_i$ otherwise.
(T2.2) If an optimization strategy is applied, then $\mathcal{T}(Q(S))$ has two subtrees:
(T2.2.1) The first one is the tree associated with Optimize($Q(S'')$) (where $Q(S'')$ is defined as above) and is connected to its parent node by the accept edge, labelled with either $0 \leq v_i \leq R_i$ or $0 \leq v_i \leq P_i$; see Figure 7.5.
(T2.2.2) The second one is the tree associated with Optimize($Q(S''')$) (where $Q(S''')$ is obtained by applying the optimization strategy to the deep copy of the input.
quintuple $Q(S)$) and is connected to its parent node by the refuse edge, labelled with either $R_i < v_i$ or $P_i < v_i \leq 1$; see Figure 7.5. Observe that every node of $T(Q(S))$ is labelled with a quintuple and every edge is labelled with an inequality constraint.

**Remark 5** Figure 7.5 illustrates how Algorithm 6, applied to $Q(S')$, generates the associated tree $T(Q(S'))$. The cases for a resource counter and a performance counter are distinguished in the sub-figures (a) and (b), respectively. Observe that, in both cases, the accept edges go south-east, while the refuse edges go south-west.

**Lemma 7.1.1** The height of the tree $T(Q(S))$ is at most $w(s + t)$. Therefore, Algorithm 5 terminates.

**Proof** Consider a path $\Gamma$ from the root of $T(Q(S))$ to any node $N$ of $T(Q(S))$. Observe that $\Gamma$ counts at most $w$ refuse edges. Indeed, following a refuse edge decreases by one the number of optimization strategies to be used. Observe also that the length of every sequence of consecutive accept edges is at most $s + t$. Indeed, following an accept edge decreases by one the number of resource and performance counters to be evaluated. Therefore, the number of edges in $\Gamma$ is at most $w(s + t)$.

**Lemma 7.1.2** Let $U := \{U_1, \ldots, U_z\}$ be a subset of $\{O_1, \ldots, O_w\}$. There exists a path from the root of $T(Q(S))$ to a leaf of $T(Q(S))$ along which the optimization strategies being applied are exactly those of $U$. 

Figure 7.5: The decision subtree for resource or performance counters

(a) The decision subtree for resource counter $r_i$

(b) The decision subtree for performance counter $p_i$
Proof Let us start at the root of $T(Q(S))$ and apply the following procedure:
1. follow the refuse edge if it uses an optimization strategy from $\{U_1, \ldots, U_z\}$,
2. follow the accept edge, otherwise.
This creates a path from the root of $T(Q(S))$ to a leaf with the desired property.

Definition 3 Let $i \in \{1, \ldots, s\}$ (resp. $\{1, \ldots, t\}$). Let $N$ be a node of $T(Q(S))$ and $Q(S_N)$ be the quintuple labelling this node. We say that $r_i$ (resp. $p_i$) is optimal at $N$ w.r.t. the evaluation function $f_i$ (resp. $g_i$) and the subset $\sigma(r_i)$ (resp. $\sigma(p_i)$) of the optimization strategies $O_1, \ldots, O_w$, whenever for all $O \in \sigma(r_i)$ (resp. $O \in \sigma(p_i)$) we have $f_i(O(S_N)) = f_i(S_N)$ (resp. $g_i(O(S_N)) = g_i(S_N)$).

Lemma 7.1.3 Let $i \in \{1, \ldots, s\}$ (resp. $\{1, \ldots, t\}$). There exists at least one leaf $L$ of $T(Q(S))$ such that $r_i$ (resp. $p_i$) is optimal at $L$ w.r.t. the evaluation function $f_i$ (resp. $g_i$) and the subset $\sigma(r_i)$ (resp. $\sigma(p_i)$) of the optimization strategies $O_1, \ldots, O_w$.

Proof Apply Lemma 7.1.2 with $U = \sigma(r_i)$ (resp. $U = \sigma(p_i)$).

Lemma 7.1.4 Algorithm 5 satisfies its output specifications.

Proof From Lemma 7.1.1 we know that Algorithm 5 terminates. So let $(C_1, S_1), \ldots, (C_e, S_e)$ be its output. We shall prove $(C_1, S_1), \ldots, (C_e, S_e)$ satisfies the conditions (i) to (iv) of Definition 2. Condition (i) is satisfied by the properties of the RealTriangularize algorithm, see Section 2.6. Condition (ii) follows clearly from the assumption that the code transformations performed by $O_1, \ldots, O_w$ preserve semantics. Observe that each time a polynomial inequality is added to a system of constraints, the negation of this inequality is also to the same system in another branch of the computations. By using a simple induction on $s + t$, we deduce that Condition (iii) is satisfied. Finally, we prove Condition (iv) by using Lemma 7.1.3.

7.2 Comprehensive translation of an annotated C program into CUDA kernels

Given a high-level model for accelerator programming (like OpenCL [111], OpenMP [41,13,9], OpenACC [113,56] or MetaFork [29]), we consider the problem of translating a program written for such a high-level model into a programming model for GPGPU devices, such as CUDA [95,73]. We assume that the numerical values of some, or all, of the hardware characteristics of the targeted GPGPU device are unknown. Hence, these quantities are treated as symbols. Similarly, we would like that some, or all, of the program parameters remain symbols in the generated code.

In our implementation, we focus on one high-level model for accelerator programming, namely MetaFork. However, we believe that an adaptation to another high-level model for accelerator programming would not be difficult. One supporting reason for that claim is the fact that automatic code translation between the MetaFork and OpenMP languages can already be done within the MetaFork compilation framework, see [29].

The hardware characteristics of the GPGPU device can be the maximum number of registers to be allocated per thread in a thread-block and the maximum number of shared memory
words to be allocated per thread in a thread-block. Similarly, the program parameters can be the number of threads per thread-block and the granularity of threads. For the generated code to be valid, hardware characteristics and program parameters need to satisfy constraints in the form of polynomial equations and inequalities. Moreover, applying code transformation (like optimization techniques) requires a case distinction based on the values of those symbols, as we saw with the example in the introduction.

In Section 7.2.1, we specify the required properties of the input code fragment $S$ from the given MetaFork program, so that the comprehensive optimization algorithm, demonstrated in Section 7.1 can handle this MetaFork program. Section 7.2.2 discusses the procedure of comprehensive translation of the MetaFork program into parametric CUDA kernels, which yields the definition of comprehensive parametric CUDA kernels.

### 7.2.1 Input MetaFork code fragment

Consider a `meta_schedule` statement $M$ and its surrounding MetaFork program $P$. In this process of code analysis and transformation, we focus on the `meta_schedule` statement $M$ and assume that the rest of the program $P$ is serial C code. Hence, our examples, like the matrix vector multiplication and matrix addition examples, consist simply of a `meta_schedule` statement $M$ together with a few (possibly none) statements located before $M$ and initializing variables used in $M$.

Consider the `meta_schedule` statement $M$, that is, a statement of the form

```
meta_schedule A
```

where $A$ is a compound statement of the form $\{A_0A_1\cdots A_\ell\}$ and each of $A_0, A_1, \ldots, A_\ell$ is a `for`-loop nest, such that:

1. each `for`-loop nest contains 2 or 4 `meta_for` loops; hence, it can be executed in a parallel fashion,
2. the body of the innermost loop can be any valid sequence of C statements; in particular, such a statement can be a `for`-loop, and

In practice, a parameter (in the sense of Definition 1) of the `meta_schedule` statement $M$ is either a `data parameter` (that is, related to data being processed, like a number of rows or columns in a matrix) or a `program parameter` (that is, related to the division of the work among the threads executing the parallel `for`-loops). In Example 4, the variable $N$ is a data parameter, whereas the variables $\text{dim}$ and $B$ are the program parameters.

Moreover, for the sake of clarity, we shall assume that the `meta_schedule` statement $M$ counts a single `meta_for` loop nest $A$. Extending the present section to the case where $M$ counts several `meta_for` loop nests can be done by existing techniques as we briefly explain now. Indeed, each `meta_for` loop nest can be handled separately. Then, “merging” the corresponding results can be done by techniques from symbolic computation, see [26, 27]. Therefore, we consider the serial elision (as defined in [29]) of $A$ in $M$ as the code fragment $S$. Turning our attention back to Example 4, Figure 7.6 shows the serial elision of the MetaFork program on the right-hand portion of Figure 7.4.
7.3. Implementation details

```c
int dim = N / B, v, u;
// v is corresponding to the thread-block index
// u is corresponding to the thread index in a thread-block

// The following code is the serial elision
for (int i = 0; i < dim; ++i)
    for (int j = 0; j < B; ++j) {
        int p = v * B + u;
        int q = i * B + j;
        c[p] += a[p][q] * b[q];
    }
```

Figure 7.6: The serial elision of the MetaFork program for matrix vector multiplication

7.2.2 Comprehensive translation into parametric CUDA kernels

Now, applying the comprehensive optimization algorithm (described in Section 7.1) on the serial elision $S$ of the `meta_schedule` statement $\mathcal{M}$ (with prescribed resource evaluation functions, performance evaluation functions and optimization strategies), we obtain a sequence of processed quintuples of `meta_schedule` statements $Q_1(\mathcal{M}), Q_2(\mathcal{M}), \ldots, Q_\ell(\mathcal{M})$, which forms a comprehensive optimization in the sense of Definition 2.

If, as mentioned in the introduction of this chapter, PTX is used as intermediate representation (IR) then, for each $i = 1, \ldots, \ell$, under the constraints defined by the polynomial system associated with $Q_i(\mathcal{M})$, the IR code associated with $Q_i(\mathcal{M})$ is the translation in assembly language of a CUDA counterpart of $\mathcal{M}$. In our implementation, we also translate to CUDA source code the MetaFork code in each $Q_i(\mathcal{M})$, since this is easier to read for a human being.

Therefore, in broad terms, a comprehensive translation of the `meta_schedule` statement $\mathcal{M}$ into parametric CUDA kernels is a decision tree, where each edge holds a Boolean expression (given by a polynomial constraint) and each leaf is either a CUDA program in PTX form, or the symbol $\emptyset$, such that for each leaf $K$, with $K \neq \emptyset$, we have:

1. $K$ works correctly under the conjunction of the Boolean expressions located between the root node and the leaf, and
2. $K$ is semantically equivalent to $\mathcal{P}$.

The symbol $\emptyset$ is used to denote a situation (in fact, value ranges for the machine and program parameters) where no CUDA program equivalent to $\mathcal{P}$ is provided.

7.3 Implementation details

In this section, we present a preliminary implementation of the comprehensive optimization algorithm demonstrated in Section 7.1. This implementation takes a given MetaFork program as input and is dedicated to the optimization of `meta_schedule` statements in view of generating parametric CUDA kernels.

For the algorithm stated in Section 7.1.7 to satisfy its specifications, one should use the PTX language for the IR. However, for simplicity, in our proof-of-concept implementation
here, we use the IR of the LLVM compiler infrastructure [75], since the MetaFork compilation framework is based on Clang [30].

Two hardware resource counters are considered: register usage per thread and local/shared memory allocated per thread-block. No performance counters are specified; however, by design, the algorithm tries to minimize the usage of hardware resources. Four optimization strategies are used: (i) reducing register pressure, (ii) controlling thread granularity, (iii) common sub-expression elimination (CSE), and (iv) caching data in local/shared memory. Details are given hereafter.

Figure 7.7 gives an overview of the software tools that are used for our implementation. Appendix E shows the implemented algorithms with these two resource counters and these three optimization strategies.

Conversion between source code and CFG. Clang [30] is used as the front-end for parsing the MetaFork source code and generating the source CFG. This latter is converted into a Maple DAG in order to take advantage of Maple’s capabilities for algebraic computation. Conversely, given a Maple version of a CFG, we can generate the corresponding MetaFork code by traversing this CFG.

Register pressure. Given the MetaFork source code, we use LLVM to generate the low-level machine instructions in the intermediate representation (IR), which are in a static single assignment (SSA) form [40]. A benefit of using the SSA form is that one can calculate the liveness set[123] without data flow analysis [123]. Once the lifetime information is computed, we use the classical linear scan algorithm [97] to estimate the register usage.

Thread granularity. A common method to improve arithmetic intensity and instruction level parallelism (ILP) is through controlling the granularity of threads [114], that is, each thread computing more than one of the output results. One can achieve this goal by adding a serial for loop within a thread. However, this method may increase the register usage as well as the amount of required shared memory (see the matrix multiplication example in Section 6.1).

Figure 7.7: The software tools involved for the implementation

[3]In the MetaFork language, the keyword cache is used to indicate that every thread accessing a specified array a must copy in local/shared memory the data it accesses in a.

7.4. Experimentation

In the case that adding the granularity loop causes the needed resources to exceed the hardware limits, our algorithm applies an optimized strategy, “Granularity set to 1,” to remove that serial loop from the generated kernel code. We implement this strategy during the translation phase from the CFG to the MetaFork code by not generating this loop.

**Common sub-expression elimination (CSE).** For each basic block in the CFG, built from the MetaFork source code, we consider the basic blocks with more than one statement. Then, we use the codegen[optimize] package of Maple, such that CSE is applied to those statements and a sequence of new statements is generated. Finally, we update each basic block with those new statements. Moreover, the optimization technique has two levels: one using Maple’s default CSE algorithm and the other using the try-harder option of codegen[optimize].

**Cache amount.** We take advantage of the PET (polyhedral extraction tool) [116] to collect information related to the index expression of an array: occurring program parameters (defined as in Section 7.2), loop iteration counters and inequalities (that give the lower and upper bounds of those loop iteration counters). We now illustrate with an example for computing the amount of words that a thread-block requires. Consider the MetaFork program with the granularity for reversing a 1D array as shown on the left-hand portion of Figure 7.8. Note that w.r.t the MetaFork code, iteration counters i, j and k of for- (and meta_for-) loops are counted as neither program nor data parameters; thus, we shall consider them as bounded variables. However, in order to calculate the amount of words required per thread-block in the Regular-Chains library of Maple, we treat the iteration counter i, which indicates the thread-block index of the CUDA kernel, as a program parameter. The function call ValueRangeWithConstraintsAndParameters to the RegularChains library, as shown on the right-hand portion of Figure 7.8 is used to calculate the required words per thread-block for vector c. As a result, a range [1, s*B+1] is returned, such that we can determine that vector c accesses s*B words per thread-block.

(a) MetaFork program with the granularity (b) function call to RegularChains library

Figure 7.8: Computing the amount of words required per thread-block for reversing a 1D array

7.4 Experimentation

We present experimental results for the implementation described in Section 7.3. We consider six simple test cases: array reversal, matrix vector multiplication, 1D Jacobi, matrix addition, matrix transpose and matrix matrix multiplication. Recall that we consider two machine
parameters: the amount $Z_B$ of shared memory per streaming multiprocessor (SM) and the maximum number $R_B$ of registers per thread in a thread-block.

Three scenarios of optimized MetaFork programs based on different systems of constraints are generated by our implementation of the comprehensive optimization algorithm. The first case of optimized MetaFork programs uses the shared memory and a granularity parameter $s$. The second case uses the shared memory but sets the granularity parameter $s$ to 1. The third case removes the cache\(^5\) keyword and sets $s$ to 1. In this latter case, the amount of words read and written per thread-block is more than the maximum amount $Z_B$ of shared memory per SM. However, the cache keyword is not implemented in the MetaFork compilation framework yet, so that we manually process editing for this keyword. For each of these optimization strategies, we use a shortened code shown in Table 7.1.

<table>
<thead>
<tr>
<th>Strategy name</th>
<th>Its code</th>
<th>Strategy name</th>
<th>Its code</th>
</tr>
</thead>
<tbody>
<tr>
<td>“Accept register pressure”</td>
<td>(1)</td>
<td>“CSE applied”</td>
<td>(2)</td>
</tr>
<tr>
<td>“No granularity reduction”</td>
<td>(3a)</td>
<td>“Granularity set to 1”</td>
<td>(3b)</td>
</tr>
<tr>
<td>“Accept caching”</td>
<td>(4a)</td>
<td>“Refuse caching”</td>
<td>(4b)</td>
</tr>
</tbody>
</table>

Array reversal. Our comprehensive optimization algorithm applies to the source code the following optimization strategy codes (1) (4a) (3a) (2) (2). For the first case, it generates the optimized MetaFork code shown in Figure 7.9:

```c
int N, s, B, dim = N/(s*B);
int a[N], c[N];
meta_schedule cache(a, c) {
    Constraints:
    meta_for (int i = 0; i < dim; i++)
    meta_for (int j = 0; j < B; j++)
        for (int k = 0; k < s; ++k) {
            int x = (i*s+k)*B+j;
            int y = N-1-x;
            c[y] = a[x];
        }
}
```

Figure 7.9: The first case of the optimized MetaFork code for array reversal.

Applying optimization strategy codes in a sequence either (1) (3b) (4a) (3a) (2) (2) or (2) (2) (3b) (1) (4a) (3a), the second case generates the optimized MetaFork code shown in Figure 7.10.

Applying optimization strategy codes in a sequence either (1) (3b) (2) (2) (4b) or (2) (2) (3b) (1) (4b) (3a), the third case generates the optimized MetaFork code shown in Figure 7.11.

Matrix vector multiplication. Applying optimization strategy codes in a sequence either (1) (4a) (3a) (2) (2) or (2) (1) (4a) (3a) (2), the first case generates the optimized MetaFork code shown in Figure 7.12.

---

\(^5\) In the MetaFork language, the intention of using the keyword `cache` is to indicate that every thread accessing a specified array a must copy in local/shared memory the data it accesses in a.
7.4. Experimentation

Constraints:
\[
\begin{align*}
2B & \leq Z_B < 2sB \\
3 & \leq R_B
\end{align*}
\]

strategies (1) (3b) (4a) (3a) (2) (2) applied or
\[
\begin{align*}
2B & \leq Z_B < 2sB \\
3 & \leq R_B < 4
\end{align*}
\]

strategies (2) (2) (3b) (1) (4a) (3a) applied

Figure 7.10: The second case of the optimized MetaFork code for array reversal

Constraints:
\[
\begin{align*}
Z_B & < 2B \\
3 & \leq R_B
\end{align*}
\]

strategies (1) (3b) (2) (2) (4b) applied or
\[
\begin{align*}
Z_B & < 2B \\
3 & \leq R_B < 4
\end{align*}
\]

strategies (2) (2) (3b) (1) (4b) applied

Figure 7.11: The third case of the optimized MetaFork code for array reversal

Constraints:
\[
\begin{align*}
sB^2 + sB + B & \leq Z_B \\
8 & \leq R_B
\end{align*}
\]

strategies (1) (4a) (3a) (2) (2) applied or
\[
\begin{align*}
sB^2 + sB + B & \leq Z_B \\
8 & \leq R_B < 9
\end{align*}
\]

strategies (2) (1) (4a) (3a) (2) applied

Figure 7.12: The first case of the optimized MetaFork code for matrix vector multiplication

Applying the optimization strategy codes in a sequence either (1) (3b) (4a) (3b) (2) (2), (2) (1) (3b) (4a) (3a) (2) or (2) (2) (3b) (1) (4a) (3a), the second case generates the optimized MetaFork code shown in Figure 7.13.

Applying the optimization strategy codes in a sequence either (1) (3b) (2) (2) (4b), (2) (1) (3b) (2) (4b) or (2) (2) (3b) (1) (4b), the third case generates the optimized MetaFork code shown in Figure 7.14.

1D Jacobi. Given 1D Jacobi source code written in MetaFork, shown in Figure 7.15, the CSE strategy is applied successfully for all cases of optimized MetaFork programs. This example requires post-processing for calculating the total amount of required shared memory.
Chapter 7. Towards Comprehensive Parametric CUDA Kernel Generation

Constraints:
\[
\begin{align*}
& B^2 + 2B \leq Z_B < sB^2 + sB + B \\
& 7 \leq R_B
\end{align*}
\]

strategies (1) (3b) (4a) (3a) (2) (2) applied

or

\[
\begin{align*}
& B^2 + 2B \leq Z_B < sB^2 + sB + B \\
& 7 \leq R_B < 9
\end{align*}
\]

strategies (2) (1) (3b) (4a) (3a) (2) applied

or

\[
\begin{align*}
& B^2 + 2B \leq Z_B < sB^2 + sB + B \\
& 7 \leq R_B < 8
\end{align*}
\]

strategies (2) (2) (3b) (1) (4a) (3a) applied

Figure 7.13: The second case of the optimized MetaFork code for matrix vector multiplication

```
int N, s = 1, B, dim0 = N/(s*B), dim1 = N/B;
int a[N][N], b[N], c[N];
meta_schedule cache(a, b, c) {
    for (int i = 0; i < dim1; i++)
        for (int j = 0; j < B; ++j) {
            int p = v*B+u;
            int q = i*B+j;
            c[p] = a[p][q]*b[q]+c[p];
        }
}
```

Figure 7.14: The third case of the optimized MetaFork code for matrix vector multiplication

```
int N, s = 1, B, dim0 = N/(s*B), dim1 = N/B;
int a[N][N], b[N], c[N];
meta_schedule {
    for (int i = 0; i < dim1; i++)
        for (int j = 0; j < B; ++j) {
            int p = v*B+u;
            int q = i*B+j;
            c[p] = a[p][q]*b[q]+c[p];
        }
}
```

Matrix addition. Due to the limitation in the codegen[optimize] package of Maple, the CSE optimizer could not handle a two-dimensional array on the left-hand side of assignments. Thus, we use a one-dimensional array to represent the output matrix. Applying the optimization strategy codes in a sequence (1) (4a) (3a) (2) (2), the first case generates the optimized MetaFork code shown in Figure 7.16.

Applying the optimization strategy codes in a sequence (1) (3b) (4a) (3a) (2) (2), the second case generates the optimized MetaFork code shown in Figure 7.17.

Applying the optimization strategy codes in a sequence (1) (3b) (2) (2) (4b), the third case generates the optimized MetaFork code shown in Figure 7.18.

Matrix addition. Due to the limitation in the codegen[optimize] package of Maple, the CSE optimizer could not handle a two-dimensional array on the left-hand side of assignments. Thus, we use a one-dimensional array to represent the output matrix. Applying the optimization strategy codes in a sequence (1) (4a) (3a) (2) (2), the first case generates the optimized MetaFork code shown in Figure 7.19.
7.4. Experimentation

```c
int T, N, s, B, dim = (N-2)/(s*B);
int a[2*N];
for (int t = 0; t < T; ++t)
meta_schedule {
    meta_for (int i = 0; i < dim; i++)
    meta_for (int j = 0; j < B; j++)
    for (int k = 0; k < s; ++k) {
        int p = i * s * B + k * B + j;
        int p1 = p + 1;
        int p2 = p + 2;
        int np = N + p;
        int np1 = N + p + 1;
        int np2 = N + p + 2;
        if (t % 2)
            a[p1] = (a[np] + a[np1] + a[np2]) / 3;
        else
            a[np1] = (a[p] + a[p1] + a[p2]) / 3;
    }
}
```

Figure 7.15: The MetaFork source code for 1D Jacobi

```c
int T, N, s, B, dim = (N-2)/(s*B);
int a[2*N];
for (int t = 0; t < T; ++t)
meta_schedule cache(a) {
    meta_for (int i = 0; i < dim; i++)
    meta_for (int j = 0; j < B; j++)
    for (int k = 0; k < s; ++k) {
        int p = j+(i*s+k)*B;
        int p1 = p+1;
        int p2 = p+2;
        int np = N+p;
        int np1 = N+p+1;
        int np2 = N+p+2;
        if (t % 2)
            a[p1] = (a[np]+a[np1]+a[np2])/3;
        else
            a[np1] = (a[p]+a[p1]+a[p2])/3;
    }
}
```

Constraints:

\[
\begin{align*}
2sB + 2 & \leq Z_B \\
9 & \leq R_B
\end{align*}
\]

strategies (1) (4a) (3a) (2) (2) applied

Figure 7.16: The first case of the optimized MetaFork code for 1D Jacobi

Applying the optimization strategy codes in a sequence either (1) (3b) (4a) (3a) (2) (2) or (2) (2) (3b) (1) (4a) (3a), the second case generates the optimized MetaFork code shown in Figure 7.20.

Applying the optimization strategy codes in a sequence either (1) (3b) (2) (2) (4b) or (2) (2) (3b) (1) (4b), the third case generates the optimized MetaFork code shown in Figure 7.21.

Matrix transpose. Applying the optimization strategy codes in a sequence (1) (4a) (3a)
Towards Comprehensive Parametric CUDA Kernel Generation

Constraints:
\[
\begin{align*}
2B + 2 & \leq Z_B < 2sB + 2 \\
9 & \leq R_B
\end{align*}
\]
strategies (1) (3b) (4a) (3a) (2) (2) applied

Figure 7.17: The second case of the optimized \texttt{MetaFork} code for 1D Jacobi

(2) (2), the first case generates the optimized \texttt{MetaFork} code shown in Figure 7.22.

Applying the optimization strategy codes in a sequence either (1) (3b) (4a) (3a) (2) (2) or (2) (2) (3b) (1) (4a) (3a), the second case generates the optimized \texttt{MetaFork} code shown in Figure 7.23.

Applying the optimization strategy codes in a sequence either (1) (3b) (2) (2) (4b) or (2) (2) (3b) (1) (4b), the third case generates the optimized \texttt{MetaFork} code shown in Figure 7.24.

\textbf{Matrix matrix multiplication.} Applying the optimization strategy codes in a sequence (1)
7.4. Experimentation

Constraints:

\[
\begin{align*}
3sB_0B_1 & \leq Z_B \\
7 & \leq R_B
\end{align*}
\]

strategies (1) (4a) (3a) (2) applied

Figure 7.19: The first case of the optimized MetaFork code for matrix addition

Constraints:

\[
\begin{align*}
3B_0B_1 & \leq Z_B < 3sB_0B_1 \\
6 & \leq R_B < 7
\end{align*}
\]

strategies (1) (3b) (4a) (3a) (2) (2) applied

Figure 7.20: The second case of the optimized MetaFork code for matrix addition

Constraints:

\[
\begin{align*}
Z_B & < 3B_0B_1 \\
6 & \leq R_B < 7
\end{align*}
\]

strategies (2) (2) (3b) (1) (4a) (3a) applied

Figure 7.21: The third case of the optimized MetaFork code for matrix addition

(4a) (3a) (2) (2), the first case generates the optimized MetaFork code shown in Figure 7.25.

Applying the optimization strategy codes in a sequence either (1) (3b) (4a) (3a) (2) (2) or (2) (2) (3b) (1) (4a) (3a), the second case generates the optimized MetaFork code shown in Figure 7.26.

Applying the optimization strategy codes in a sequence either (1) (3b) (2) (2) (4b) or (2)
Chapter 7. Towards Comprehensive Parametric CUDA Kernel Generation

Constraints:
\[
\begin{align*}
2sB_0B_1 & \leq Z_B \\
6 & \leq R_B
\end{align*}
\]
strategies (1) (4a) (3a) (2) (2) applied

Figure 7.22: The first case of the optimized MetaFork code for matrix transpose

Constraints:
\[
\begin{align*}
2B_0B_1 & < 2sB_0B_1 \\
5 & \leq R_B
\end{align*}
\]
strategies (1) (3b) (4a) (3a) (2) (2) applied or

Figure 7.23: The second case of the optimized MetaFork code for matrix transpose

Constraints:
\[
\begin{align*}
Z_B & < 2B_0B_1 \\
5 & \leq R_B
\end{align*}
\]
strategies (1) (3b) (2) (2) (4b) applied or

Figure 7.24: The third case of the optimized MetaFork code for matrix transpose

(2) (3b) (1) (4b), the third case generates the optimized MetaFork code shown in Figure 7.27
7.5 Conclusion

In this chapter, we proposed a comprehensive optimization algorithm that optimizes the input code fragment depending on unknown machine and program parameters; meanwhile, we realized a proof-of-concept implementation for generating comprehensive parametric MetaFork programs, in the form of a case distinction based on the possible values of the machine and program parameters.
The comprehensive optimization algorithm that we proposed takes optimization strategies, resource counters and performance counters into account; we implemented two resource counters and four optimization strategies in this comprehensive optimization algorithm.

With this preliminary implementation, experimentation shows that given a MetaFork program, three scenarios of optimized MetaFork programs are generated, each of them with a system of constraints specifying when the corresponding code is valid.

In addition, from the experimental results, we observe that different sequences of the optimization strategies yield the same optimized MetaFork program. However, since some optimization strategies are applied to the intermediate representation of the source code, the corresponding improvements are not shown in Section 7.4.
Chapter 8

Conclusion and Future Work

Our ultimate goal was to generate optimized CUDA kernels, where machine and program parameters are unknown symbols and can be, respectively, determined and optimized when the CUDA code is installed on the targeted hardware.

In our route to this goal, we developed a model of multithreaded computation (MCM) targeting many-core machines so as to measure the work, span and parallelism overhead of a CUDA-like program as well as to estimate the overall execution time of such program. Our experimentation shows that this model was able to determine the value ranges of some program parameters so that one can obtain a program with reduced parallelism overheads.

We acknowledge the fact that the MCM abstract machines admit a few simplifications and limitations with respect to actual many-core devices. Some factors, such as bank conflicts of shared memory and register spilling, increase memory access time because of the hardware limits of a streaming multiprocessor; however, these factors are ignored by our model. It is desirable to enhance our model with such features so as to obtain more accurate complexity estimates of a CUDA-like program. We leave this task for future work.

Taking advantage of the sophisticated CUDA code generator in PPCG [115], we realized a preliminary implementation of automatic code translator from MetaFork [29] programs to CUDA programs. In our automatic code generator, the number of threads per thread-block is generated as a symbol, whereas this number is generated as a numerical value by other tools, in particular PPCG.

In order to obtain optimized CUDA kernels, we also tuned a program parameter representing the granularity of a thread in a thread-block. Experimental results indicate that the CUDA kernels with these program parameters can lead to significant performance improvement and that the optimal choices for program parameters may depend on the problem size.

In addition, the non-linear expressions supported by our automatic code generator may involve parameters related to thread-block formats but not to the granularity of threads. Relaxing this limitation is a work in progress. On the other hand, our BPAS library provides efficient, parallel algorithms dealing with polynomial arithmetic operations; hence, it is desirable to integrate support from the BPAS library into our MetaFork-to-CUDA code generation framework.

Ultimately, we have proposed a comprehensive optimization algorithm that, given unknown machine and program parameters, optimizes the input code fragment in the form of a case discussion based on the possible values of the machine and program parameters. In the preliminary implementation, we considered optimizing MetaFork programs using LLVM as inter-
mediate representation. In the future, this algorithm should be implemented within the Meta-Fork-to-CUDA and should use PTX as intermediate representation.
Bibliography


Appendix A

Sample Code in the BPAS Library

We collect sample code from the BPAS library, including two adaptive algorithms shown in Appendix A.1 and some user interface code for BPAS rings and polynomials in Appendix A.2.

A.1 Adaptive algorithms

/*************************************************************************
 * Internal multiplication function for integer univariate polynomials *
 * It is an adaptive algorithm based on input sizes and available cores. *
 * @param mul, a vector of coefficients of the product polynomial *
 * @param a, a vector of coefficients over Z representing a polynomial *
 * @param n, the size of the above vector *
 * @param b, a vector of coefficients over Z representing a polynomial *
 * @param m, the size of the above vector *
 **********************************************************************/
void univariateMultiplication(mpz_class* mul, mpz_class* a, int n,
                             mpz_class* b, int m) {
    int size = n + m - 1;
    // UnivariateIntegerPolynomial: internal data structure
    // for integer univariate polynomials
    UnivariateIntegerPolynomial aPoly(n, a);
    UnivariateIntegerPolynomial bPoly(m, b);
    UnivariateIntegerPolynomial rPoly(size, mul);

    if (size < 64) { // Naive multiplication; serial
        MulNaive naive;
        naive.multiply(&aPoly, &bPoly, &rPoly);
    } else if (size < 2048) { // Kronecker substitution method; serial
        MulKS ks;
        ks.multiply(&aPoly, &bPoly, &rPoly);
    } else if (size < 4096) { // Toom Cook method; parallel
        int worker = __cilkrts_get_nworkers();
    }
if (worker <= 6) { // 4-way
    MulToom4 toom4;
    toom4.multiply(&aPoly, &bPoly, &rPoly, 4);
} else { // 8-way
    MulToom8 toom8;
    toom8.multiply(&aPoly, &bPoly, &rPoly, 8);
}
else { // Two convolution method; parallel
    MulSSA ssa;
    ssa.multiply(&aPoly, &bPoly, &rPoly);
}
}

// The adaptive Taylor shift algorithm combining a divide and conquer method, *
// aka Algorithm (E) in [119], with the algorithm in [28] as the base case. *
// *param a, a vector of coefficients over Z representing a polynomial *
// *param m, the size of the above vector, power of 2 *
// *param bi, a vector of coefficients representing (x+1)^k, for all 1<k<=m/2 *
// *param B, the size determining whether to switch to Algorithm (E) *
******************************************************************************/
void taylorShiftBasePower2(mpz_class* a, int m, mpz_class* bi, int B) {
    int d = (B > m)? m : B;

    // Base case, calling to the algorithm in [28]
    cilk_for (int i = 0; i < m/d; ++i)
        taylorShiftIncrementalCilkFor(&a[i*B], d, 16);

    mpz_class* mul = new mpz_class[m];
    for (int i = B; i < m; i <<= 1)
        cilk_for (int k = 0; k < m/(2*i); ++k)
            univariateMultiplication(&mul[2*k*i], &a[2*k*i+i], i, &bi[i-1], i);
    for (int k = 0; k < m; ++k)
        a[k] += mul[k];
    delete [] mul;
}

A.2 User interfaces

/******************************************************************************
* A child class of abstract BPASRing implements a ring *
* structure in the sense of commutative algebra. *
******************************************************************************/
class BPASRing {

public:
static int characteristic;
static bool isPrimeField;
static bool isComplexField;
virtual bool isZero() = 0;
virtual void zero() = 0;
virtual bool isOne() = 0;
virtual void one() = 0;
virtual bool isNegativeOne() = 0;
virtual void negativeOne() = 0;
virtual int isConstant() = 0;
};

typedef int DataType; // Different data type for each concrete polynomial

class BPASPolynomial : public BPASRing {
public:
    BPASPolynomial& operator= (BPASPolynomial&);
    BPASPolynomial& operator+ (BPASPolynomial&);
    BPASPolynomial& operator+= (BPASPolynomial&);
    BPASPolynomial& operator- (BPASPolynomial&);
    BPASPolynomial& operator- ();
    BPASPolynomial& operator-= (BPASPolynomial&);
    BPASPolynomial& operator* (BPASPolynomial&);
    BPASPolynomial& operator*= (BPASPolynomial&);
    BPASPolynomial& operator/ (BPASPolynomial&);
    BPASPolynomial& operator/= (BPASPolynomial&);
    BPASPolynomial& operator^ (int);
    bool operator== (BPASPolynomial&);
    bool operator!= (BPASPolynomial&);
friend std::ostream& operator<< (std::ostream&, BPASPolynomial&);
};

class BPASUnivariatePolynomial : public BPASPolynomial {
public:
    // A child class of abstract BPASUnivariatePolynomial inherits from BPASPolynomial.
    // Moreover, it implements a univariate polynomial ring in which a notion of Greatest Common Divisor (GCD) makes sense.
    // Arithmetic operations (addition, subtraction, multiplication, exact division, exponentiation, and equality test) are provided.
}
* division) can take a polynomial and a coefficient as input arguments. *
* Arithmetic operations also include shift left and shift right (that *
* is, multiplication or division by a power of the variable) as well as *
* divisions (monic, pseudo and lazy), differentiation, evaluation, GCD *
* computation and square-free factorization. One can query the degree, *
* the leading coefficient, the coefficient of a prescribed monomial, *
* the variable name, the content of a polynomial. *
*************************************************************************/
class BPASUnivariatePolynomial : public BPASPolynomial {
public:
    BPASUnivariatePolynomial& operator+ (DataType);
    BPASUnivariatePolynomial& operator+= (DataType);
    BPASUnivariatePolynomial& operator- (DataType);
    BPASUnivariatePolynomial& operator-= (DataType);
    BPASUnivariatePolynomial& operator* (DataType);
    BPASUnivariatePolynomial& operator*= (DataType);
    BPASUnivariatePolynomial& operator/ (DataType);
    BPASUnivariatePolynomial& operator/= (DataType);
    BPASUnivariatePolynomial& operator<< (int);
    BPASUnivariatePolynomial& operator<<= (int);
    BPASUnivariatePolynomial& operator>> (int);
    BPASUnivariatePolynomial& operator>>= (int);
    BPASUnivariatePolynomial& monicDivide(BPASUnivariatePolynomial&);
    BPASUnivariatePolynomial& monicDivide(BPASUnivariatePolynomial&,
        BPASUnivariatePolynomial*);
    BPASUnivariatePolynomial& lazyPseudoDivide(BPASUnivariatePolynomial&,
        DataType*, DataType*);
    BPASUnivariatePolynomial& lazyPseudoDivide(BPASUnivariatePolynomial&,
        BPASUnivariatePolynomial*, DataType*, DataType*);
    BPASUnivariatePolynomial& pseudoDivide(BPASUnivariatePolynomial&,
        DataType*);
    BPASUnivariatePolynomial& pseudoDivide(BPASUnivariatePolynomial&,
        BPASUnivariatePolynomial*, DataType*);
    virtual void differentiate(int) = 0;
    DataType content();
    BPASUnivariatePolynomial& gcd(BPASUnivariatePolynomial&);
    std::vector<BPASUnivariatePolynomial&> squareFree();
    virtual int degree() = 0;
    DataType leadingCoefficient();
    DataType coefficient(int);
    void setCoefficient(int, DataType);
    virtual void setVariableName (std::string) = 0;
    virtual std::string variable() = 0;
    virtual bool isTrailingCoefficientZero() = 0;
    DataType evaluate(DataType);
};
Appendix B

Theoretical Analysis of Fundamental Algorithms Using the MCM Model

With respect to using our MCM model to analyze fundamental algorithms in Chapter 4, we attach the PDF versions of the executable MAPLE worksheets of all applications.

Euclidean

This document is to demonstrate the analysis of the Euclidean algorithm

Given two polynomials $a$ and $b$ over a finite field, with $\text{deg}(a) = n - 1$, and $\text{deg}(b) = m - 1$, where $n \geq m$, return the GCD of $(b, r)$, where $r$ is the remainder in the division of $a$ by $b$

- Computing 1 coefficient in the intermediate remainder (in one division step) needs 3 steps along the span
  1. multiplication factor
  2. multiplication
  3. subtraction
- Assumptions:
  1. At each step, the degree of $a$ or $b$ decreases by 1
  2. $GCD(a, b) = 1$

```maple
> restart;
> terms_a := n;
terms_a := n

(1)

> terms_b := m;
terms_b := m

(2)
```
```
> steps_for_one_coefficient := 3;
steps_for_one_coefficient := 3

\textbf{Optimized Algorithm}

\begin{itemize}
\item Phase 1, for the degree of \(a\) and \(b\) will decrease by \((n-1, m-1),\)
\((n-s-1, m), (n-2s-1, m-1), \ldots, (m-1, m-1),\) such that
there are \(\frac{n-m}{s}\) kernel calls, and each kernel call has \(\frac{m}{l}\) blocks (\(l\)
divides \(m\), and \(l\) is even).
\item Phase 2, for the first kernel call, we use \(\frac{m}{l}\) blocks, for the second
kernel call, we use \(\left\lfloor \frac{m-s}{2l} \right\rfloor\) blocks, for the third kernel call, we use
\(\frac{m-s}{l}\) blocks, and so on, until the last kernel call, we use 1 block
with \(\frac{s}{2}\) terms. There are \(2\frac{m}{s}\) kernel calls.
\item For each block, it performs \(s\) division steps, and we use \(l\) threads,
computing 1 leading coefficient and 2 other coefficients
\end{itemize}

\begin{align*}
> \text{optimized\_param} & := s; \\
& \text{optimized\_param} := s \quad (1.1)
> \text{optimized\_threads} & := l; \\
& \text{optimized\_threads} := l \quad (1.2)
> \text{optimized\_euclidean\_kernels\_phase1} & := (\text{terms\_a-terms\_b})/\text{optimized\_param}; \\
& \text{optimized\_euclidean\_kernels\_phase1} := \frac{n-m}{s} \quad (1.3)
> \text{optimized\_euclidean\_kernels\_phase2} & := 2^s \text{terms\_b}/\text{optimized\_param}; \\
& \text{optimized\_euclidean\_kernels\_phase2} := \frac{2m}{s} \quad (1.4)
> \text{optimized\_euclidean\_kernels} & := \text{optimized\_euclidean\_kernels\_phase1} + \\
& \text{optimized\_euclidean\_kernels\_phase2}; \\
& \text{optimized\_euclidean\_kernels} := \frac{n-m}{s} + \frac{2m}{s} \quad (1.5)
> \text{optimized\_euclidean\_first\_blocks} & := \text{terms\_b}/\text{optimized\_threads}; \\
& \text{optimized\_euclidean\_first\_blocks} := \frac{m}{l} \quad (1.6)
\end{align*}
```
# Work

\[
\text{optimized\_euclidean\_phase1\_workperblock} := \text{normal}\left(\frac{3}{2} \cdot s^2 + \frac{3}{2} \cdot s + 6 \cdot l \cdot s\right) \quad (1.1.1)
\]

\[
\text{optimized\_euclidean\_phase2\_workperblock} := \text{normal}\left(\frac{9}{4} \cdot s^2 + \frac{9}{2} \cdot s + 6 \cdot l \cdot s\right) \quad (1.1.2)
\]

Phase 1: Work per block is the same as optimized division algorithm.

Phase 2: For the last kernel call, it uses \(\frac{s}{2}\) threads to compute the leading coefficients and \(l\) threads to compute other coefficients.

Phase 2: For the rest \(\frac{2m}{s} - 1\) kernel calls, it uses \(l\) threads to compute the leading coefficients \((s + s - 1 + s - 1 + ... + \frac{s}{2} + \frac{s}{2})\) for 1 block, and \(l\) threads to compute other coefficients.

\[
\text{optimized\_euclidean\_work} := \text{normal}\left(\text{optimized\_euclidean\_kernels\_phase1} \times \text{optimized\_euclidean\_first\_blocks} \times \text{optimized\_euclidean\_phase1\_workperblock} + 2 \times \sum\left((\text{terms\_b} - i \times \text{optimized\_param}/2) / \text{optimized\_threads}, i = 0 \ldots (\text{terms\_b} / \text{optimized\_param}) - 1\right) \times \text{optimized\_euclidean\_phase2\_workperblock}\right) \quad (1.1.3)
\]

\[
\text{optimized\_euclidean\_work} := \frac{3}{8} m (81m + 161n + 81s + 5ms + 4ns + 3s^2 + 14m + 4n + 6s) / l
\]

# Span

\[
\text{optimized\_euclidean\_span} := \text{normal}(\text{normal}(\text{optimized\_euclidean\_kernels} \times \text{steps\_for\_one\_coefficient} \times \text{optimized\_param})) \quad (1.2.1)
\]

\[
\text{optimized\_euclidean\_span} := 3n + 3m
\]

# Overhead

For each kernel calls, each block needs \(s\) leading coefficients and
other coefficients of a and same of b, then it takes time to load data and time to write back
\[
\text{optimized\_euclidean\_overhead\_perblock} := 8^*U; \\
\text{optimized\_euclidean\_overhead\_perblock} := 8U 
\] (1.3.1)

\[
\text{optimized\_euclidean\_overhead\_a} := \\
\text{optimized\_euclidean\_kernels\_phase1} * \\
\text{optimized\_euclidean\_first\_blocks} * \\
\text{optimized\_euclidean\_overhead\_perblock} \\
\text{optimized\_euclidean\_overhead\_a} := \frac{8(n - m) m U}{s l} 
\] (1.3.2)

\[
\text{optimized\_euclidean\_overhead\_b} := \text{normal}(2^*\text{sum}((\text{terms\_b} - i^* \\
\text{optimized\_param}/2)/\text{optimized\_threads}, i=0.. \\
\text{terms\_b}/\text{optimized\_param}-1) * \\
\text{optimized\_euclidean\_overhead\_perblock}) \\
\text{optimized\_euclidean\_overhead\_b} := \frac{4 m (3 m + s) U}{l s} 
\] (1.3.3)

\[
\text{optimized\_euclidean\_overhead} := \text{normal} \\
(\text{optimized\_euclidean\_overhead\_a} + \\
\text{optimized\_euclidean\_overhead\_b}) \\
\text{optimized\_euclidean\_overhead} := \frac{4 m U(2 n + m + s)}{l s} 
\] (1.3.4)

\[
\text{Tp} \\
\text{optimized\_euclidean\_workperthread} := \\
\text{steps\_for\_one\_coefficient}^*\text{optimized\_param} \\
\text{optimized\_euclidean\_workperthread} := 3s 
\] (1.4.1)

\[
\text{optimized\_euclidean\_overheadperthread} := 8^*U; \\
\text{optimized\_euclidean\_overheadperthread} := 8U 
\] (1.4.2)

\[
\text{optimized\_euclidean\_C} := \text{optimized\_euclidean\_workperthread} + \\
\text{optimized\_euclidean\_overheadperthread}; \\
\text{optimized\_euclidean\_C} := 3s + 8U 
\] (1.4.3)

\[
\text{optimized\_euclidean\_N} := \text{normal} \\
(\text{optimized\_euclidean\_kernels\_phase1}^* \\
\text{optimized\_euclidean\_first\_blocks} + \\
2^*\text{sum}((\text{terms\_b} - i^*\text{optimized\_param}/2)/\text{optimized\_threads}, i=0. \\
\text{terms\_b}/\text{optimized\_param}-1)) \\
\text{optimized\_euclidean\_N} := \frac{1}{2} \frac{m (2 n + m + s)}{l s} 
\] (1.4.4)

\[
\text{optimized\_euclidean\_L} := \text{normal}(\text{optimized\_euclidean\_kernels}) \\
\] (1.4.5)
\[ \text{optimized\_euclidean\_L} := \frac{n+m}{s} \]  
(1.4.5)

\[ \text{optimized\_euclidean\_K} := \frac{m}{l} \]  
(1.4.6)

\[ \text{optimized\_euclidean\_Tp := normal(} \]
\[ \frac{(\text{optimized\_euclidean\_N/optimized\_euclidean\_K+optimized\_euclidean\_L)\star\text{optimized\_euclidean\_C)}}{s} \]
\[ \text{)} \]

\[ \text{optimized\_euclidean\_Tp} := \frac{1}{2} \frac{(4n+3m+s)(3s+8U)}{s} \]  
(1.4.7)

\[ > \text{naive\_euclidean\_work := eval(\text{optimized\_euclidean\_work}, [s=1])}; \]
\[ \text{naive\_euclidean\_work} := \frac{3}{8} \frac{m(8lm+16ln+8l+19m+8n+9)}{l} \]  
(2)

\[ > \text{naive\_euclidean\_span := eval(\text{optimized\_euclidean\_span}, [s=1])}; \]
\[ \text{naive\_euclidean\_span} := \frac{3n+3m}{l} \]  
(3)

\[ > \text{naive\_euclidean\_overhead := eval(\text{optimized\_euclidean\_overhead}, [s=1])}; \]
\[ \text{naive\_euclidean\_overhead} := \frac{4mU(2n+m+1)}{l} \]  
(4)

\[ > \text{naive\_euclidean\_Tp := eval(\text{optimized\_euclidean\_Tp}, [s=1])}; \]
\[ \text{naive\_euclidean\_Tp} := \frac{1}{2} \frac{(4n+3m+1)(3+8U)}{l} \]  
(5)

\[ > \text{work\_ratio := normal(} \]
\[ \frac{\text{naive\_euclidean\_work/optimized\_euclidean\_work)}}{\text{)} \]
\[ \text{work\_ratio} := \frac{8lm+16ln+8l+19m+8n+9}{8lm+16ln+8ls+5ms+4ns+3s^2+14m+4n+6s} \]  
(6)

\[ > \text{span\_ratio := normal(} \]
\[ \frac{\text{naive\_euclidean\_span/optimized\_euclidean\_span)}}{\text{)} \]
\[ \text{span\_ratio} := 1 \]  
(7)

\[ > \text{overhead\_ratio := normal(} \]
\[ \frac{\text{naive\_euclidean\_overhead/optimized\_euclidean\_overhead)}}{\text{)} \]
\[ \text{overhead\_ratio} := \frac{(2n+m+1)s}{2n+m+s} \]  
(8)

\[ > \text{Tp\_ratio := normal(} \]
\[ \frac{\text{naive\_euclidean\_Tp/optimized\_euclidean\_Tp)}}{\text{)} \]
\[ \text{Tp\_ratio} := \frac{(4n+3m+1)(3+8U)s}{(4n+3m+s)(3s+8U)} \]  
(9)

\[ s = O(l) \] on the condition that \( 4s + 2l \) must fit into the local memory
Fast Fourier Transform

This documentation is to demonstrate the analysis of Cooley-Tukey and Stockham FFT algorithm on GPUs

\[
DFT_{\{2^k\}} = \prod_{i=0}^{k-1} \left( DFT_{\{2^i\}} \times L_{\{2^{k-1}\}} \right) \left( D_{\{2, 2^{k-1}\}} \right) \\
\times \left( L_{\{2^i\}} \right) \left( L_{\{2^{k-i}\}} \times L_{\{2^i\}} \right)
\]

\textbf{Cooley-Tukey FFT algorithm}

Phase 1: \( x \rightarrow \prod_{i=0}^{k-1} \left( L_{\{2^{k-i}\}} \times L_{\{2^i\}} \times D_{\{2, 2^{k-1}\}} \right) x \)

Phase 2: \( x \rightarrow \left( L_{\{2^{k-i}\}} \times DFT_{\{m\}} \right) x \)

Phase 3:
\[
x \rightarrow \prod_{i=0}^{k-1} \left( L_{\{2^{k-i}\}} \times DFT_{\{2\}} \times L_{\{2^{k-1}\}} \right) \left( L_{\{2^i\}} \times D_{\{2, 2^{k-1}\}} \right) x
\]

\begin{align*}
&> \texttt{cooleytukey\_param} := m; \\
&\quad \texttt{cooleytukey\_param} := m \quad (1.1) \\
&> \texttt{cooleytukey\_threadsperblock} := l; \\
&\quad \texttt{cooleytukey\_threadsperblock} := l \quad (1.2) \\
&> \texttt{cooleytukey\_size} := n; \\
&\quad \texttt{cooleytukey\_size} := n \quad (1.3)
\end{align*}

First data shuffle:
In list\_transpose2, each thread moves 2 data, and it executes 9 additions, 11 bit operations.
Further, list\_transpose2 is called \( \log(n) - \log(s) \) times

Second list\_fft, each thread moves \( s \) data. For each data, it executes \( \frac{s}{2} + \ldots + 1 \) with 2 modular additions (each 5 arithmetic operations), 2 modular multiplications (each 11 arithmetic operations).
Also for each data, it executes 30 bit operations.
Additionally, 1 modular multiplication.
Third list_butterfly, each trhead moves 2 data and reads 1 data, and it executes 6 additions, 6 bit operations, 2 modular additions.

Or compute jumped powers:
move 2 data, 2 modular additions, 2 \( \cdot (\log(i)) \) calls to
fourier_reduction, with \( i = 1 \ldots l \), 1 additional call to fourier_reduction
\( \ln \) fourier_reduction, 11 bit operations, 5 additions, 1 multiplications

\[
\begin{align*}
\text{cooleytukey\_phase3\_workperblock\_powers} & := 17 \cdot 2 \cdot \text{simplify}\left(\sum \log[2](i), i=1..\text{cooleytukey\_threadsperblock}\right) + \\
& \quad \text{cooleytukey\_threadsperblock} \cdot 27; \\
\text{cooleytukey\_phase3\_workperblock\_powers} & := \frac{34 \ln(\Gamma(l+1))}{\ln(2)} + 27 \ln(l) \tag{1.4} \\
\text{cooleytukey\_phase3\_spanperkernel\_powers} & := 17 \cdot 2 \cdot \log[2] \left(\text{cooleytukey\_threadsperblock}\right) + 27; \\
\text{cooleytukey\_phase3\_spanperkernel\_powers} & := \frac{34 \ln(l)}{\ln(2)} + 27 \tag{1.5} \\
\text{cooleytukey\_phase3\_overheadperblock\_powers} & := 4 \cdot U; \\
\text{cooleytukey\_phase3\_overheadperblock\_powers} & := 4 U \tag{1.6}
\end{align*}
\]

\section*{Work}

\[
\begin{align*}
\text{cooleytukey\_phase1\_kernels} & := \log[2](\text{cooleytukey\_size})-\log[2](\text{cooleytukey\_param}); \\
\text{cooleytukey\_phase1\_kernels} & := \frac{\ln(n)}{\ln(2)} - \frac{\ln(m)}{\ln(2)} \tag{1.1.1} \\
\text{cooleytukey\_phase1\_blocks} & := \frac{\text{cooleytukey\_size}}{\text{cooleytukey\_threadsperblock}}; \\
\text{cooleytukey\_phase1\_blocks} & := \frac{n}{l} \tag{1.1.2} \\
\text{cooleytukey\_phase1\_workperblock} & := 20 \cdot \text{cooleytukey\_threadsperblock}; \\
\text{cooleytukey\_phase1\_workperblock} & := 20 l \tag{1.1.3} \\
\text{cooleytukey\_phase1\_work} & := \text{cooleytukey\_phase1\_kernels} \cdot \text{cooleytukey\_phase1\_blocks} \cdot \text{cooleytukey\_phase1\_workperblock}; \\
\text{cooleytukey\_phase1\_work} & := 20 \left(\frac{\ln(n)}{\ln(2)} - \frac{\ln(m)}{\ln(2)} \right) n \tag{1.1.4} \\
\text{cooleytukey\_phase2\_blocks} & := \frac{\text{cooleytukey\_size}}{\text{cooleytukey\_threadsperblock} \cdot \text{cooleytukey\_param}}; \\
\text{cooleytukey\_phase2\_blocks} & := \frac{n}{lm} \tag{1.1.5} \\
\text{cooleytukey\_phase2\_workperblock} & := \left(\text{cooleytukey\_threadsperblock} \cdot \text{simplify}(\text{cooleytukey\_param}) + \right. \\
& \quad 32 \cdot \sum(\text{cooleytukey\_param}/2^i, i=1..\log[2])
\end{align*}
\]
\begin{align*}
&\text{(cooleytukey\_param)) + 11) + 30 \times \text{cooleytukey\_param);} \\
&\text{cooleytukey\_phase2\_workperblock} := 1 \left(32 \, m^2 + 9 \, m\right) \\
&\text{cooleytukey\_phase2\_work} := \frac{n \left(32 \, m^2 + 9 \, m\right)}{m} \quad (1.1.6) \\
&\text{cooleytukey\_phase3\_kernels} := \log[2](\text{cooleytukey\_size}) - \log[2](\text{cooleytukey\_param}); \\
&\text{cooleytukey\_phase3\_blocks} := \frac{n}{\text{cooleytukey\_threadsperblock}}; \\
&\text{cooleytukey\_phase3\_blocks} := \frac{n}{\text{cooleytukey\_threadsperblock}} \quad (1.1.9) \\
&\text{cooleytukey\_phase3\_work} := 22 \left(\frac{\ln(n)}{\ln(2)} - \frac{\ln(m)}{\ln(2)}\right) n \quad (1.1.10) \\
&\text{cooleytukey\_phase3\_work} := 22 \left(\frac{\ln(n)}{\ln(2)} - \frac{\ln(m)}{\ln(2)}\right) n \quad (1.1.11) \\
&\text{cooleytukey\_phase3\_workowers} := \text{simplify} \\
&\text{(cooleytukey\_phase3\_kernels} \times \text{cooleytukey\_phase3\_blocks} \times \text{cooleytukey\_phase3\_workperblock\_powers}); \\
&\text{cooleytukey\_phase3\_workowers} := \frac{(\ln(n) - \ln(m)) n (27 \, \ln(2) + 34 \, \ln(\Gamma(l+1)))}{\ln(2)^2 \, l} \quad (1.1.12) \\
&\text{cooleytukey\_work} := \text{normal(cooleytukey\_phase1\_work} + \\
&\text{cooleytukey\_phase2\_work} + \text{cooleytukey\_phase3\_work}); \\
&\text{cooleytukey\_work} := \frac{n \left(32 \, \ln(2) \, m + 42 \, \ln(n) + 9 \, \ln(2) - 42 \, \ln(m)\right)}{\ln(2)} \quad (1.1.13) \\
&\text{cooleytukey\_work1} := \text{eval(cooleytukey\_work, [m=16])}; \\
&\text{cooleytukey\_work1} := \frac{n \left(353 \, \ln(2) + 42 \, \ln(n)\right)}{\ln(2)} \quad (1.1.14) \\
&\text{cooleytukey\_work\_powers} := \text{normal(cooleytukey\_phase1\_work} + \\
&\text{cooleytukey\_phase2\_work} + \text{cooleytukey\_phase3\_work\_powers}); \\
&\text{cooleytukey\_work\_powers} := \frac{1}{\ln(2)^2 \, l} \left(\text{n} \left(32 \, \ln(2)^2 \, m \, l + 47 \, \ln(2) \, \ln(n)\right) \quad (1.1.15) \\
&\quad + 9 \, \ln(2)^2 \, l - 47 \, \ln(2) \, \ln(m) + 34 \, \ln(n) \, \ln(\Gamma(l+1)) - 34 \, \ln(\Gamma(l+1)) \, \ln(m))\right) \\
&\text{cooleytukey\_work\_powers} := \text{expand(eval}
\[
\text{cooleytukey\_work1\_powers} := 333 n + \frac{47 n \ln(n)}{\ln(2)} + \frac{34 n \ln(n) \ln(\Gamma(l))}{\ln(2)^2 l} - \frac{136 n \ln(\Gamma(l))}{\ln(2) l} \tag{1.1.16}
\]

\textbf{Span}

\[\text{cooleytukey\_phase1\_span} := 20 \times \text{cooleytukey\_phase1\_kernels};\]

\[\text{cooleytukey\_phase1\_span} := \frac{20 \ln(n)}{\ln(2)} - \frac{20 \ln(m)}{\ln(2)} \tag{1.2.1}\]

\[\text{cooleytukey\_phase2\_span} := \text{simplify}\left(32 \times \log[2]_{(\text{cooleytukey\_param})} \times \sum(\text{cooleytukey\_param}/2^i, i=1..\log[2]_{(\text{cooleytukey\_param})} + 30 \times \text{cooleytukey\_param}) + 11\right);\]

\[\text{cooleytukey\_phase2\_span} := \frac{32 \ln(m) (-1 + m)}{\ln(2)} + 30 m + 11 \tag{1.2.2}\]

\[\text{cooleytukey\_phase3\_span} := 22 \times \text{cooleytukey\_phase3\_kernels};\]

\[\text{cooleytukey\_phase3\_span} := \frac{22 \ln(n)}{\ln(2)} - \frac{22 \ln(m)}{\ln(2)} \tag{1.2.3}\]

\[\text{cooleytukey\_phase3\_span\_powers} := \text{simplify}\left(\text{cooleytukey\_phase3\_kernels} \times \text{cooleytukey\_phase3\_span\_per\_kernel\_powers}\right);\]

\[\text{cooleytukey\_phase3\_span\_powers} := \frac{(\ln(n) - \ln(m)) (34 \ln(l) + 27 \ln(2))}{\ln(2)^2} \tag{1.2.4}\]

\[\text{cooleytukey\_span} := \text{normal(}\text{cooleytukey\_phase1\_span} + \text{cooleytukey\_phase2\_span} + \text{cooleytukey\_phase3\_span});\]

\[\text{cooleytukey\_span} := \frac{30 \ln(2) m + 32 \ln(m) m + 42 \ln(n) + 11 \ln(2) - 74 \ln(m)}{\ln(2)} \tag{1.2.5}\]

\[\text{cooleytukey\_span1} := \text{eval(}\text{cooleytukey\_span}, \left[m=16\right]);\]

\[\text{cooleytukey\_span1} := \frac{2243 \ln(2) + 42 \ln(n)}{\ln(2)} \tag{1.2.6}\]

\[\text{cooleytukey\_span\_powers} := \text{normal(}\text{cooleytukey\_phase1\_span} + \text{cooleytukey\_phase2\_span} + \text{cooleytukey\_phase3\_span\_powers});\]

\[\text{cooleytukey\_span\_powers} := \frac{1}{\ln(2)^2} \left(30 m \ln(2)^2 + 32 m \ln(2) m + 34 \ln(n) \ln(l) - 34 \ln(m) \ln(l) + 47 \ln(n) \ln(2) + 11 \ln(2)^2 - 79 \ln(m) \ln(2)\right) \tag{1.2.7}\]

\[\text{cooleytukey\_span1\_powers} := \text{expand(}\text{eval(}\text{cooleytukey\_span\_powers}, \left[m=16\right]));\]
\[
\text{cooleytukey\_span1\_powers} := 2223 + \frac{34 \ln(n) \ln(l)}{\ln(2)^2} - \frac{136 \ln(l)}{\ln(2)} + \frac{47 \ln(n)}{\ln(2)}
\]

\textbf{Overhead}

\[
\text{cooleytukey\_phase1\_overheadperblocka} := 2 \cdot U + 2 \cdot U \cdot \text{cooleytukey\_threadsperblock};
\]
\[
\text{cooleytukey\_phase1\_overheadperblockb} := 4 \cdot U;
\]
\[
\text{cooleytukey\_phase1\_overhead} := \text{simplify}(\log[2] (\text{cooleytukey\_threadsperblock}) \cdot \text{cooleytukey\_phase1\_blocks} \cdot \text{cooleytukey\_phase1\_overheadperblocka} + (\text{cooleytukey\_phase1\_kernels} - \log[2] (\text{cooleytukey\_threadsperblock})) \cdot \text{cooleytukey\_phase1\_blocks} \cdot \text{cooleytukey\_phase1\_overheadperblockb});
\]
\[
\text{cooleytukey\_phase1\_overhead} := \frac{2 \cdot n \cdot U \cdot (\ln(l) \cdot l - \ln(l) + 2 \ln(n) - 2 \ln(m))}{\ln(2)}
\]

\[
\text{cooleytukey\_phase2\_overheadperblock} := 2 \cdot \text{cooleytukey\_param} \cdot U;
\]
\[
\text{cooleytukey\_phase2\_overheadperblock} := 2 \cdot m \cdot U
\]
\[
\text{cooleytukey\_phase2\_overhead} := \text{cooleytukey\_phase2\_blocks} \cdot \text{cooleytukey\_phase2\_overheadperblock};
\]
\[
\text{cooleytukey\_phase2\_overhead} := \frac{2 \cdot n \cdot U}{l}
\]

\[
\text{cooleytukey\_phase3\_overheadperblock} := 5 \cdot U;
\]
\[
\text{cooleytukey\_phase3\_overheadperblock} := 5 \cdot U
\]
\[
\text{cooleytukey\_phase3\_overhead} := \text{cooleytukey\_phase3\_kernels} \cdot \text{cooleytukey\_phase3\_blocks} \cdot \text{cooleytukey\_phase3\_overheadperblock};
\]
\[
\text{cooleytukey\_phase3\_overhead} := \frac{5 \left( \frac{\ln(n)}{\ln(2)} - \frac{\ln(m)}{\ln(2)} \right) \cdot n \cdot U}{l}
\]
\[
\text{cooleytukey\_phase3\_overhead\_powers} := \text{cooleytukey\_phase3\_kernels} \cdot \text{cooleytukey\_phase3\_blocks} \cdot \text{cooleytukey\_phase3\_overheadperblock\_powers};
\]
\[
\text{cooleytukey\_phase3\_overhead\_powers} := \frac{4 \left( \frac{\ln(n)}{\ln(2)} - \frac{\ln(m)}{\ln(2)} \right) \cdot n \cdot U}{l}
\]
\[
\text{cooleytukey\_overhead} := \text{normal}(\text{cooleytukey\_phase1\_overhead} + \text{cooleytukey\_phase2\_overhead} + \text{cooleytukey\_phase3\_overhead})
\]
\[
\begin{align}
\text{cooleytukey\_phase2\_overhead} + \text{cooleytukey\_phase3\_overhead}; \\
\text{cooleytukey\_overhead} := \frac{n U (2 \ln(l) l - 2 \ln(l) + 9 \ln(n) + 2 \ln(2) - 9 \ln(m))}{l \ln(2)} \\
\text{cooleytukey\_overhead1} := \frac{n U (2 \ln(l) l - 2 \ln(l) + 9 \ln(n) - 34 \ln(2))}{l \ln(2)} \\
\text{cooleytukey\_overhead_powers} := \frac{2 n U (\ln(l) l - \ln(l) + 4 \ln(n) + \ln(2) - 4 \ln(m))}{l \ln(2)} \\
\text{cooleytukey\_overhead1\_powers} := \frac{2 n U \ln(l)}{l \ln(2)} - \frac{2 n U \ln(l)}{l \ln(2)} + \frac{8 n U \ln(n)}{l \ln(2)} - \frac{30 n U}{l} \\
\end{align}
\]
cooleytukey\_Tp := \frac{1}{m \ln(2)^2} ((4 m \ln(n) + \ln(2)) m - 4 \ln(m) m + \ln(2)) (2 U \ln(2) + 2 U \ln(2) + 30 \ln(2) m + 32 \ln(m) m + 11 \ln(2) - 32 \ln(m)) 

> cooleytukey\_Tp1 := eval(cooleytukey\_Tp, [m=16]);

cooleytukey\_Tp1 := \frac{1}{16} (64 \ln(n) - 239 \ln(2)) (2 U \ln(2) + 2 U \ln(2) + 2411 \ln(2)) \ln(2)^2

> cooleytukey\_C\_powers := eval(cooleytukey\_phase3\_spanperkernel\_powers + cooleytukey\_phase1\_overheadperblocka, [s=16]);

cooleytukey\_C\_powers := \frac{34 \ln(l)}{\ln(2)} + 27 + 2 U + 2 U

> cooleytukey\_Tp\_powers := normal((cooleytukey\_N/cooleytukey\_K + cooleytukey\_L) * cooleytukey\_C\_powers);

cooleytukey\_Tp\_powers := \frac{1}{m \ln(2)^2} ((4 m \ln(n) + \ln(2)) m - 4 \ln(m) m + \ln(2)) (2 U \ln(2) + 2 U \ln(2) + 34 \ln(l)) \ln(2)

> cooleytukey\_Tp1\_powers := eval(cooleytukey\_Tp\_powers, [m=16]);

cooleytukey\_Tp1\_powers := \frac{1}{16} (64 \ln(n) - 239 \ln(2)) (2 U \ln(2) + 2 U \ln(2) + 34 \ln(l) + 27 \ln(2))

---

**Cooley-Tukey with vs without pre-computed jumped powers**

> \textbf{Rwork} := normal(cooleytukey\_work1\_powers / cooleytukey\_work1); \quad (1)

\textbf{Rwork} := \frac{333 \ln(2)^2 l + 47 l \ln(2) \ln(m) + 34 \ln(m) \ln(\Gamma(l) l) - 136 \ln(\Gamma(l) l) \ln(2)}{\ln(2) l (353 \ln(2) + 42 \ln(n))}

> \textbf{Rspan} := normal(cooleytukey\_span1\_powers / cooleytukey\_span1); \quad (2)

\textbf{Rspan} := \frac{2223 \ln(2)^2 l + 34 \ln(n) \ln(l) - 136 \ln(2) \ln(l) + 47 \ln(n) \ln(2)}{\ln(2) (2243 \ln(2) + 42 \ln(n))}

> \textbf{Roverhead} := normal(cooleytukey\_overhead1\_powers / cooleytukey\_overhead1); \quad (3)

\textbf{Roverhead} := \frac{2 (\ln(l) l - \ln(l) + 4 \ln(n) - 15 \ln(2))}{2 \ln(l) l - 2 \ln(l) + 9 \ln(n) - 34 \ln(2)}

> \textbf{Rt} := normal(cooleytukey\_Tp1\_powers / cooleytukey\_Tp1); \quad (4)

\textbf{Rt} := \frac{2 U \ln(2) + 2 U \ln(2) + 34 \ln(l) + 27 \ln(2)}{\ln(2) (2 U + 2 U + 2411)}
Stockham FFT algorithm

For each fixed $0 \leq i < k$, there are three computational steps:

Phase 1: $x \rightarrow \left( L(2)^{2^{k-i}} \times L(2^i) \right) x$

Phase 2: $x \rightarrow \left( D(2, 2^{k-i-1}) \times L(2^i) \right) x$

Phase 3: $x \rightarrow \left( DFT(2) \times L(2^{k-1}) \right) x$

```
> stockham_threadsperblock := l;
  stockham_threadsperblock := l               (2.1)
> stockham_size := n;
  stockham_size := n                           (2.2)
```

First compute phase 3 $DFT(2) \times L(2^{k-1})$ once, a thread reading 2 data + writing back the same + executing 3 bit operations, 3 additions, 2 modular add operations (5 arithmetic operations)

Second, for each $0 \leq i < k$, compute phase 1, 2 & 3

In phase 1, if $2^i < l$, a thread reading 1 data and writing back the same + executing 14 bit operations, 10 additions, 1 multiplications.

If $2^i \geq l$, a thread reading 1 data and writing back the same + executing 11 bit operations, 8 additions.

In phase 2, a thread reading 1 data and writing back the same + executing 5 bit operations, 4 additions, 1 modular multiply operations (11 arithmetic operations)

Prefix sum to compute the powers of the n-th primitive root of unity

Each thread deals with 4 data items

```
> roots_blocks := stockham_size/(4*stockham_threadsperblock);
  roots_blocks := \frac{1}{4} \frac{n}{l}               (2.1.1)
```

# Regard to the prefix sum, one block performs the up-sweep phase, with 3 instructions, and then performs the down-sweep phase, with 5 instructions, regard to Listing 2 in scan.pdf. (Except mediate step)
roots_inclusive_workperblock := normal(simplify( 8*(sum (stockham_threadsperblock/2^(i-1),i=1..log[2] (stockham_threadsperblock)))+2)+1 ));

roots_inclusive_workperblock := 1 + 16 l

(2.1.10)

roots_inclusive_overheadperblock := (4+4)*U;

roots_inclusive_overheadperblock := 8 U

(2.1.11)

roots_inclusive_work := roots_blocks* roots_inclusive_workperblock;

roots_inclusive_work := \frac{1}{4} \frac{n(1 + 16 l)}{l}

(2.1.12)

roots_inclusive_span := simplify( 8*(log[2] (stockham_threadsperblock)+2)+1 );

roots_inclusive_span := \frac{8 \ln(l)}{\ln(2)} + 17

(2.1.13)

roots_inclusive_overhead := roots_blocks* roots_inclusive_overheadperblock;

roots_inclusive_overhead := \frac{2 n U}{l}

(2.1.14)

roots_mediate_overheadperblock := (4+4)*U;

roots_mediate_overheadperblock := 8 U

(2.1.15)

roots_mediate_work := normal(simplify( 8*(sum (stockham_threadsperblock/2^(i-1),i=1..log[2] (stockham_threadsperblock)))+2)+1));

roots_mediate_work := 1 + 16 l

(2.1.16)

roots_mediate_overhead := roots_mediate_overheadperblock;

roots_mediate_overhead := 8 U

(2.1.17)

roots_mediate_span := 8*(log[2](stockham_threadsperblock)+2) +1;

roots_mediate_span := \frac{8 \ln(l)}{\ln(2)} + 17

(2.1.18)

roots_exclusive_workperblock := 4*stockham_threadsperblock;

roots_exclusive_workperblock := 4 l

(2.1.19)

roots_exclusive_overheadperblock := (4+1+4)*U;

roots_exclusive_overheadperblock := 9 U

(2.1.20)

roots_exclusive_work := (roots_blocks-1)* roots_exclusive_workperblock;

roots_exclusive_work := 4 \left( \frac{1}{4} \frac{n}{l} - 1 \right) l

(2.1.21)

roots_exclusive_span := 4;

roots_exclusive_span := 4

(2.1.22)

roots_exclusive_overhead := (roots_blocks-1)*
roots_exclusive_overheadperblock;

roots_exclusive_overhead := 9 \left( \frac{1}{4} \frac{n}{l} - 1 \right) U \tag{2.1.15}

# Total work & span in phase 2

> roots_work := normal(roots_inclusive_work +
> roots_mediate_work + roots_exclusive_work);

\[ roots_work := \frac{1}{4} \frac{48 \hat{F} + 20 \ln(l) + 4 l + n}{l} \] \tag{2.1.16}

> roots_span := simplify(roots_inclusive_span +
> roots_mediate_span + roots_exclusive_span);

\[ roots_span := \frac{16 \ln(l)}{\ln(2)} + 38 \] \tag{2.1.17}

> roots_overhead := normal(roots_inclusive_overhead +
> roots_mediate_overhead + roots_exclusive_overhead);

\[ roots_overhead := \frac{1}{4} \frac{U (-17 n + 4 l)}{l} \] \tag{2.1.18}

**Work**

> stockham_phase1_blocks := stockham_size / stockham_threadsperblock;

\[ stockham_phase1_blocks := \frac{n}{l} \] \tag{2.2.1}

> stockham_phase1_workperblock := 25 * stockham_threadsperblock;

\[ stockham_phase1_workperblock := 25 l \] \tag{2.2.2}

> stockham_phase1_work := stockham_phase1_blocks * stockham_phase1_workperblock;

\[ stockham_phase1_work := 25 n \] \tag{2.2.3}

> stockham_phase2_blocks := stockham_size / (2 * stockham_threadsperblock);

\[ stockham_phase2_blocks := \frac{1}{2} \frac{n}{l} \] \tag{2.2.4}

> stockham_phase2_workperblock := 20 * stockham_threadsperblock;

\[ stockham_phase2_workperblock := 20 l \] \tag{2.2.5}

> stockham_phase2_work := stockham_phase2_blocks * stockham_phase2_workperblock;

\[ stockham_phase2_work := 10 n \] \tag{2.2.6}

> stockham_phase3_blocks := stockham_size / (2 * stockham_threadsperblock);

\[ \] \tag{2.2.7}
\[
stockham\_phase3\_blocks := \frac{1}{2} \frac{n}{l} \quad (2.2.7)
\]

\[
stockham\_phase3\_workperblock := 16 \times 
stockham\_threadsperblock;
stockham\_phase3\_workperblock := 16 l \quad (2.2.8)
\]

\[
stockham\_phase3\_work := stockham\_phase3\_blocks \times
stockham\_phase3\_workperblock;
stockham\_phase3\_work := 8 n \quad (2.2.9)
\]

\[
stockham\_work := normal(roots\_work + sum
(stockham\_phase1\_work+stockham\_phase2\_work+
stockham\_phase3\_work, i=0..\log[2](stockham\_size)-2) +
stockham\_phase3\_work);
stockham\_work :=
\frac{1}{4} 172 n \ln(n) + 48 \ln(2) l^2 - 120 n \ln(2) + 4 \ln(2) + n \ln(2)
\ln(2) \quad (2.2.10)
\]

**Span**

\[
stockham\_phase1\_span := 25;
stockham\_phase1\_span := 25 \quad (2.3.1)
\]

\[
stockham\_phase2\_span := 10;
stockham\_phase2\_span := 10 \quad (2.3.2)
\]

\[
stockham\_phase3\_span := 8;
stockham\_phase3\_span := 8 \quad (2.3.3)
\]

\[
stockham\_span := normal(roots\_span + sum
(stockham\_phase1\_span+stockham\_phase2\_span+
stockham\_phase3\_span, i=0..\log[2](n)-2) +
stockham\_phase3\_span);
stockham\_span :=
\frac{16 \ln(l) + 3 \ln(2) + 43 \ln(n)}{\ln(2)} \quad (2.3.4)
\]

**Overhead**

\[
stockham\_phase1\_overheadperblock := 2*U;
stockham\_phase1\_overheadperblock := 2 U \quad (2.4.1)
\]

\[
stockham\_phase1\_overhead := stockham\_phase1\_blocks \times
stockham\_phase1\_overheadperblock;
stockham\_phase1\_overhead := \frac{2 n U}{l} \quad (2.4.2)
\]

\[
stockham\_phase2\_overheadperblock := 2*U;
stockham\_phase2\_overheadperblock := 2 U \quad (2.4.3)
\]

\[
stockham\_phase2\_overhead := stockham\_phase2\_blocks \times
stockham\_phase2\_overheadperblock; 
(2.4.4)
\]
\[ stockham\_phase2\_overhead := \frac{nU}{l} \quad (2.4.4) \]

> \( stockham\_phase3\_overheadperblock := 4U; \)
\[ stockham\_phase3\_overheadperblock := 4U \quad (2.4.5) \]

> \( stockham\_phase3\_overhead := stockham\_phase3\_blocks \times stockham\_phase3\_overheadperblock; \)
\[ stockham\_phase3\_overhead := \frac{2nU}{l} \quad (2.4.6) \]

> \( stockham\_overhead := normal(roots\_overhead + \text{sum} \)
\( stockham\_phase1\_overhead + stockham\_phase2\_overhead + stockham\_phase3\_overhead, \text{for } i = 0..\log_2(stockham\_size) - 2) + \)
\( stockham\_phase3\_overhead); \)
\[ stockham\_overhead := \frac{1}{4} U(20n\ln(n) - 4l\ln(2) + 5n\ln(2)) \quad (2.4.7) \]

\[ Tp \]

> \( stockham\_N := normal(roots\_blocks + \text{sum} \)
\( stockham\_phase1\_blocks + stockham\_phase2\_blocks + stockham\_phase3\_blocks, \text{for } i = 0..\log_2(stockham\_size) - 2) + \)
\( stockham\_phase3\_blocks); \)
\[ stockham\_N := \frac{1}{4} n(-5\ln(2) + 8\ln(n)) \quad (2.5.1) \]

> \( stockham\_L := normal(3 + \text{sum}(3, \text{for } i = 0..\log_2(stockham\_size) - 2)) + 1); \)
\[ stockham\_L := \frac{\ln(2) + 3\ln(n)}{\ln(2)} \quad (2.5.2) \]

> \( stockham\_C := roots\_inclusive\_span + stockham\_phase3\_overheadperblock; \)
\[ stockham\_C := \frac{8\ln(l)}{\ln(2)} + 17 + 4U \quad (2.5.3) \]

> \( stockham\_K := stockham\_phase1\_blocks; \)
\[ stockham\_K := \frac{n}{l} \quad (2.5.4) \]

> \( stockham\_Tp := normal((stockham\_N/stockham\_K + stockham\_L) \times stockham\_C); \)
\[ stockham\_Tp := \frac{1}{4} (-\ln(2) + 20\ln(n)) (4U\ln(2) + 8\ln(l) + 17\ln(2)) \quad (2.5.5) \]

Cooley-Tukey without pre-computed jumped powers vs Stockham
>
\( work\_ratio := normal(cooleytukey\_work1\_powers / stockham\_work); \)
\[ work\_ratio := \quad (5) \]

\[ (4n (333\ln(2)^2 l + 47\ln(2) \ln(n) + 34 \ln(n) \ln(\Gamma(l)) l) \]
\[-136 \ln(\Gamma(l) l \ln(2)) \bigg\rfloor (\ln(2) (172 n l \ln(n) + 48 \ln(2) l^2 - 120 n l \ln(2) \\
+ 4 l \ln(2) + n \ln(2))) / (\ln(2) (16 \ln(l) + 3 \ln(2) + 43 \ln(n))\] 

\(\text{span\_ratio} := \text{normal}(\text{cooley\_tukey\_span1\_powers} / \text{stockham\_span});\)

\[
\text{span\_ratio} := \frac{2223 \ln(2)^2 + 34 \ln(n) \ln(l) - 136 \ln(2) \ln(l) + 47 \ln(n) \ln(2)}{\ln(2) (16 \ln(l) + 3 \ln(2) + 43 \ln(n))}
\]

(6) 

\(\text{overhead\_ratio} := \text{normal}(\text{cooley\_tukey\_overhead1\_powers} / \text{stockham\_overhead});\)

\[
\text{overhead\_ratio} := \frac{8 n (\ln(l) l - \ln(l) + 4 \ln(n) - 15 \ln(2))}{20 n \ln(n) - 4 l \ln(2) + 5 n \ln(2)}
\]

(7) 

\(\text{Tp\_ratio} := \text{normal}(\text{cooley\_tukey\_Tp1\_powers} / \text{stockham\_Tp});\)

\[
\text{Tp\_ratio} := \frac{1}{4} \frac{(64 \ln(n) - 239 \ln(2)) (2 U l \ln(2) + 2 U l \ln(2) + 34 \ln(l) + 27 \ln(2))}{(-ln(2) + 20 \ln(n)) (4 U l \ln(2) + 8 \ln(l) + 17 \ln(2))}
\]

(8)
Plain Polynomial Multiplication Algorithm

This document is to demonstrate the analysis of plain polynomial multiplication algorithm

Given two polynomials $a$ and $b$ over a finite field, with $\text{deg}(a) = n - 1$, and $\text{deg}(b) = m - 1$, where $n \geq m$, compute the product of $a \times b$

```plaintext
> restart;
> terms_a := n;
> terms_b := m;
> multiplication_param := s;
> multiplication_threadsperblock := l;
> multiplication_length := terms_a + multiplication_param - 1;
> multiplication_rows := terms_b/multiplication_param;
```

Phase 1:

- The grid of thread blocks is 2-D. Principle: each thread block computes some of the coefficient products and some of the partial sums toward the polynomial product. Those partial sums are stored in an auxiliary array, which is later processed in Phase 2.
- One row of thread blocks contributes to $y = n + s - 1$ terms of the product of $a \times b$, while there are $x = \frac{m}{s}$ rows of blocks.
- If we use $l$ threads per block, then we have $\frac{x \cdot y}{s \cdot l}$ thread-blocks.
- Each thread in each thread block performs $s^2$ coefficient products and $s \cdot (s - 1)$ additions.
- Each block needs $s$ terms of $b$ and $l \cdot s + s - 1$ terms of $a$. Then it computes $s \cdot l$ elements of the $x \times y$ matrix.

```plaintext
> multiplication_phase1_blocks := multiplication_length* 
multiplication_rows/(multiplication_threadsperblock* 
multiplication_param);
```

```
multiplication_phase1_blocks := \frac{(n + s - 1) m}{s \cdot l}
```
Phase 2:
- The x rows of the auxiliary array M are added pairwise in \( \log(x) \) parallel steps. At each step, the number of rows (to be added) is divided by two.
- At a given parallel step, each thread reads s elements from one row and s from another row.
- Therefore at the i-th parallel step (starting at \( i=0 \) and using \( l \) threads per block) one reads \( 2 \times s \times l \) coefficients per block.
- The total amount of data (coefficients) to be added at the beginning at the i-th parallel is twice less than at the previous step, thus it is \( x \times y \div 2^i \).
- Therefore, at the i-th parallel step, using \( l \) threads per block, we need \( \frac{x \times y}{s \times l \times 2^{i+1}} \) blocks and we have \( \log_2 x \) parallel steps for the whole phase.
- Each thread to compute at most s addition needs s element of one row and s element from another row, then writes back s elements.

\[
\text{multiplication\_phase2\_blocks} := \text{proc}(i) \\
\text{\quad return multiplication\_length} \times \text{multiplication\_rows} / \\
\text{\quad (multiplication\_param} \times \text{multiplication\_threadsperblock} \times 2^i(); \\
\text{end proc;}
\]

\[
\text{multiplication\_phase2\_steps} := \text{simplify}(\log_2(\text{multiplication\_rows}));
\]

\[
\text{multiplication\_phase2\_steps} := \frac{\ln \left( \frac{m}{s} \right)}{\ln(2)} 
\]  

### Work

\[
\text{multiplication\_phase1\_workperblock} := \\
\text{multiplication\_threadsperblock} \times \text{multiplication\_param} \times \\
\text{(multiplication\_param} + \text{multiplication\_param}-1); \\
\text{multiplication\_phase1\_workperblock} := l \times s \times (2 \times s - 1) 
\]  

\[
\text{multiplication\_phase1\_work} := \text{multiplication\_phase1\_blocks} \times \\
\text{multiplication\_phase1\_workperblock}; \\
\text{multiplication\_phase1\_work} := \frac{(n + s - 1) \times m \times (2 \times s - 1)}{s} 
\]  

\[
\text{multiplication\_phase2\_workperblock} := \\
\text{multiplication\_threadsperblock} \times \text{multiplication\_param}; \\
\text{multiplication\_phase2\_workperblock} := l \times s 
\]  

\[
\text{multiplication\_phase2\_work} := \text{simplify(sum} \\
\text{(multiplication\_phase2\_blocks}(i)*) 
\]
\[
\begin{align*}
\text{multiplication\_phase2\_workperblock}, i &= 1.. \text{multiplication\_phase2\_steps}) + \text{multiplication\_length/2)}; \\
\text{multiplication\_phase2\_work} &= \frac{1}{2} 2 m n + 2 m s - n s - s^2 - 2 m + s \\
\text{multiplication\_work} &= \text{normal(multiplication\_phase1\_work + multiplication\_phase2\_work)}; \\
\text{multiplication\_work} &= 2 m n + 2 m s - 2 m - \frac{1}{2} n - \frac{1}{2} s + \frac{1}{2} \\
\text{expand}((2 m - 1/2)(n+s-1)); \\
2 m n + 2 m s - 2 m - \frac{1}{2} n - \frac{1}{2} s + \frac{1}{2}
\end{align*}
\]

\section*{Span}
\[
\begin{align*}
\text{multiplication\_phase1\_span} &= \text{multiplication\_param} * (\text{multiplication\_param} + \text{multiplication\_param-1}); \\
\text{multiplication\_phase1\_span} &= s (2 s - 1) \\
\text{multiplication\_phase2\_span} &= \text{multiplication\_phase2\_steps} * \text{multiplication\_param}; \\
\text{multiplication\_phase2\_span} &= \ln\left(\frac{m}{s}\right) s \ln(2) \\
\text{multiplication\_span} &= \text{expand(simplify (multiplication\_phase1\_span + multiplication\_phase2\_span))}; \\
\text{multiplication\_span} &= 2 s^2 - s + \frac{\ln\left(\frac{m}{s}\right) s}{\ln(2)}
\end{align*}
\]

\section*{Overhead}
\[
\begin{align*}
\text{s from b, l*s+s-1 from a} \\
\text{# s + l\cdot s + s - 1 \leq Z} \\
\text{multiplication\_phase1\_overheadperblock} &= \text{U* (multiplication\_param+1+1+multiplication\_param);} \\
\text{multiplication\_phase1\_overheadperblock} &= U (2 s + 2) \\
\text{multiplication\_phase1\_overhead} &= \text{multiplication\_phase1\_blocks* multiplication\_phase1\_overheadperblock}; \\
\text{multiplication\_phase1\_overhead} &= \frac{(n+s-1) m U (2 s + 2)}{s^2 l} \\
\text{multiplication\_phase2\_overheadperblock} &= \text{U*3* multiplication\_param}; \\
\text{multiplication\_phase2\_overheadperblock} &= 3 U s
\end{align*}
\]
\[
\text{multiplication\_phase2\_overhead} := \text{simplify}(\text{sum} \\
\text{multiplication\_phase2\_blocks(i),i=1..} \\
\text{multiplication\_phase2\_steps})* \\
\text{multiplication\_phase2\_overheadperblock}; \\
\text{multiplication\_phase2\_overhead} := \frac{3(n+s-1)(-s+m)U}{s^3} \\
\text{(3.4)}
\]

\[
\text{multiplication\_overhead} := \text{normal} \\
(\text{multiplication\_phase1\_overhead}+\text{multiplication\_phase2\_overhead}) \\
\text{;} \\
\text{multiplication\_overhead} := \frac{(n+s-1)U(5ms-3s^2+2m)}{s^2} \\
\text{(3.5)}
\]

\text{Tp}

\text{\textbf{Phase 1 N & L & C}}

\[
\text{multiplication\_phase1\_workperthread} := \text{multiplication\_param}^* \\
(\text{multiplication\_param}+\text{multiplication\_param}-1); \\
\text{multiplication\_phase1\_workperthread} := s(2s-1) \\
\text{(4.1.1)}
\]

\[
\text{multiplication\_phase1\_overheadperthread} := \\
\text{multiplication\_phase1\_overheadperblock}; \\
\text{multiplication\_phase1\_overheadperthread} := U(2s+2) \\
\text{(4.1.2)}
\]

\[
\text{multiplication\_phase1\_C} := \\
\text{multiplication\_phase1\_workperthread}+ \\
\text{multiplication\_phase1\_overheadperthread}; \\
\text{multiplication\_phase1\_C} := s(2s-1)+U(2s+2) \\
\text{(4.1.3)}
\]

\[
\text{multiplication\_phase1\_N} := \text{multiplication\_phase1\_blocks}; \\
\text{multiplication\_phase1\_N} := \frac{(n+s-1)m}{s^2} \\
\text{(4.1.4)}
\]

\[
\text{multiplication\_phase1\_L} := 1; \\
\text{multiplication\_phase1\_L} := 1 \\
\text{(4.1.5)}
\]

\[
\text{Tp\_phase\_1} := \text{simplify}((\text{multiplication\_phase1\_N}/\text{P}+ \\
\text{multiplication\_phase1\_L})*\text{multiplication\_phase1\_C}, \text{[P=} \\
\text{multiplication\_phase1\_blocks]})); \\
\text{Tp\_phase\_1} := 4Us+4s^2+4U-2s \\
\text{(4.1.6)}
\]

\text{\textbf{Phase 2 N & L & C}}

\[
\text{multiplication\_phase2\_workperthread} := \text{multiplication\_param}; \\
\text{multiplication\_phase2\_workperthread} := s \\
\text{(4.2.1)}
\]

\[
\text{multiplication\_phase2\_overheadperthread} := \\
\text{multiplication\_phase2\_overheadperblock}; \\
\text{multiplication\_phase2\_overheadperthread} := 3Us \\
\text{(4.2.2)}
\]

\[
\text{multiplication\_phase2\_C} :=
\]
\[ Tp_{\text{phase 2}} = \left( \frac{2(-s+m)}{m} + \frac{\ln\left(\frac{m}{s}\right)}{\ln(2)} \right)(3Us+s) \] (4.2.6)
\[ multiplication\_work1 := 2mn - \frac{1}{2}n \]  \hspace{1cm} (5.1)

\[ multiplication\_span1 := \text{expand}(\text{eval}(\text{multiplication\_span}, [s=1])) \]  \hspace{1cm} (5.2)

\[ multiplication\_overhead1 := \text{normal}(\text{eval}(\text{multiplication\_overhead}, [s=1])) \]  \hspace{1cm} (5.3)

\[ multiplication\_Tp1 := \text{normal}(\text{eval}(\text{multiplication\_Tp}, [s=1])) \]  \hspace{1cm} (5.4)

\[ \text{work\_ratio} := \frac{\text{normal}(\text{multiplication\_work1})}{\text{multiplication\_work}} \]  \hspace{1cm} (9)

\[ \text{span\_ratio} := \frac{\text{normal}(\text{multiplication\_span1})}{\text{multiplication\_span}} \]  \hspace{1cm} (10)

\[ \text{overhead\_ratio} := \frac{\text{normal}(\text{multiplication\_overhead1})}{\text{multiplication\_overhead}} \]  \hspace{1cm} (11)

\[ \text{Tp\_ratio} := \frac{\text{normal}(\text{multiplication\_Tp1})}{\text{multiplication\_Tp}} \]  \hspace{1cm} (12)

\[ s = O(1) \text{ on the condition that } 2ls + 2s \text{ must fit into the local memory} \]
Polynomial Multiplication Algorithm

This document is to demonstrate the analysis of plain vs FFT-based polynomial multiplication algorithm

Given two polynomials $a$ and $b$ over a finite field, with $\deg(a) = n - 1$, and $\deg(b) = m - 1$, where $n \geq m$, compute the product of $a \times b$

```plaintext
> restart;
> terms_a := n;
> terms_b := m;  
```

(1)

Plain multiplication

```plaintext
> plain_param := s;
> plain_threadsperblock := l;
> plain_length := terms_a + plain_param - 1;
> plain_rows := terms_b/plain_param;
> plain_phase1_blocks := plain_length*plain_rows/163
```

Phase 1 multiplication:

- The grid of thread blocks is 2-D. Principle: each thread block computes some of the coefficient products and some of the partial sums toward the polynomial product. Those partial sums are stored in an auxiliary array, which is later processed in Phase 2.
- One row of thread blocks contributes to $y = n + s - 1$ terms of the product of $a \times b$, while there are $x = \frac{m}{s}$ rows of blocks.
- If we use $l$ threads per block, then we have $\frac{x \cdot y}{s \cdot l}$ thread-blocks.
- Each thread in each thread block performs $s^2$ coefficient products and $s \cdot (s - 1)$ additions.
- Each block needs $s$ terms of $b$ and $l \cdot s + s - 1$ terms of $a$. Then it computes $s \cdot l$ elements of the $x \times y$ matrix.

```plaintext
> plain_phase1_blocks := plain_length*plain_rows/163
```
(plain_threadsperblock*plain_param);

\[
\text{plain\_phase1\_blocks} := \frac{(n+s-1)\, m}{s^2\, l} \quad (7)
\]

Phase 2 addition:
- The \(x\) rows of the auxiliary array \(M\) are added pairwise in \(\log(x)\) parallel steps. At each step, the number of rows (to be added) is divided by two.
- At a given parallel step, each thread reads \(s\) elements from one row and \(s\) from another row.
- Therefore, at the \(i\)-th parallel step (starting at \(i=0\) and using \(l\) threads per block) one reads \(2^i\) times \(s\) times \(l\) coefficients per block.
- The total amount of data (coefficients) to be added at the beginning at the \(i\)-th parallel is twice less than at the previous step, thus it is \(x\) times \(y\) divided by \(2^i\).
- Therefore, at the \(i\)-th parallel step, using \(l\) threads per block, we need \(\frac{x\cdot y}{s\cdot l\cdot 2^{i+1}}\) blocks and we have \(\log_2 x\) parallel steps for the whole phase.
- Each thread to compute at most \(s\) addition needs \(s\) element of one row and \(s\) element from another row, then writes back \(s\) elements.

> plain_phase2_blocks := proc(i)
    return plain_length*plain_rows/(plain_param*plain_threadsperblock*2^(i));
end proc:

> plain_phase2_steps := simplify(log[2](plain_rows));

\[
\text{plain\_phase2\_steps} := \frac{\ln\left(\frac{m}{s}\right)}{\ln(2)} \quad (8)
\]

Work

> plain_phase1_workperblock := plain_threadsperblock*plain_param*(plain_param+plain_param-1);

\[
\text{plain\_phase1\_workperblock} := l s \, (2 \, s - 1) \quad (1.1)
\]

> plain_phase1_work := plain_phase1_blocks*
plain_phase1_workperblock;

\[
\text{plain\_phase1\_work} := \frac{(n+s-1)\, m \, (2\, s - 1)}{s} \quad (1.2)
\]

> plain_phase2_workperblock := plain_threadsperblock*plain_param;

\[
\text{plain\_phase2\_workperblock} := l s \quad (1.3)
\]

> plain_phase2_work := simplify(sum(plain_phase2_blocks(i)*
plain_phase2_workperblock,i=1..plain_phase2_steps)+

\[
\]
plain_length/2);

\[
\text{plain\_phase2\_work} := \frac{1}{2} \frac{2 mn+2 ms-n s-s^2-2 m+s}{s}
\]  \hspace{1cm} (1.4)

> plain_work := normal(plain_phase1_work+plain_phase2_work);

\[
\text{plain\_work} := 2 mn+2 ms-2 m-\frac{1}{2} n-\frac{1}{2} s+\frac{1}{2}
\]  \hspace{1cm} (1.5)

> expand((2*m- 1/2)*(n+s-1));

\[
2 mn+2 ms-2 m-\frac{1}{2} n-\frac{1}{2} s+\frac{1}{2}
\]  \hspace{1cm} (1.6)

\section*{Span}

> plain_phase1_span := plain_param*(plain_param+plain_param-1);

\[
\text{plain\_phase1\_span} := s (2 s-1)
\]  \hspace{1cm} (2.1)

> plain_phase2_span := plain_phase2_steps*plain_param;

\[
\text{plain\_phase2\_span} := \frac{\ln\left(\frac{m}{s}\right) s}{\ln(2)}
\]  \hspace{1cm} (2.2)

> plain_span := expand(simplify(plain_phase1_span+
plain_phase2_span));

\[
\text{plain\_span} := 2 s^2 - s + \frac{\ln\left(\frac{m}{s}\right) s}{\ln(2)}
\]  \hspace{1cm} (2.3)

\section*{Overhead}

\# s + \ell \cdot s + s - 1 \leq Z

> plain_phase1_overheadperblock := U*(plain_param+1+plain_param);

\[
\text{plain\_phase1\_overheadperblock} := U (2 s+2)
\]  \hspace{1cm} (3.1)

> plain_phase1_overhead := plain_phase1_blocks*plain_phase1_overheadperblock;

\[
\text{plain\_phase1\_overhead} := \frac{(n+s-1) m U (2 s+2)}{s^2 \ell}
\]  \hspace{1cm} (3.2)

> plain_phase2_overheadperblock := U*3*plain_param;

\[
\text{plain\_phase2\_overheadperblock} := 3 U s
\]  \hspace{1cm} (3.3)

> plain_phase2_overhead := simplify(sum(plain_phase2_blocks(i),i=1..plain_phase2_steps)*plain_phase2_overheadperblock);

\[
\text{plain\_phase2\_overhead} := \frac{3 (n+s-1) (-s+m) U}{s \ell}
\]  \hspace{1cm} (3.4)

> plain_overhead := normal(plain_phase1_overhead+
plain_phase2_overhead);
\[
plain\_overhead := \frac{(n + s - 1) U (5 m s - 3 s^2 + 2 m)}{s^2 l}
\]  

(3.5)

**Phases 1 N & L & C**

- \( plain\_phase1\_workperthread := plain\_param \times (plain\_param + plain\_param - 1) \)
  \[
  plain\_phase1\_workperthread := s (2 s - 1)
  \]  
  (4.1.1)

- \( plain\_phase1\_overheadperthread := plain\_phase1\_overheadperblock \)
  \[
  plain\_phase1\_overheadperthread := U (2 s + 2)
  \]  
  (4.1.2)

- \( plain\_phase1\_C := plain\_phase1\_workperthread + plain\_phase1\_overheadperthread \)
  \[
  plain\_phase1\_C := s (2 s - 1) + U (2 s + 2)
  \]  
  (4.1.3)

- \( plain\_phase1\_N := plain\_phase1\_blocks \)
  \[
  plain\_phase1\_N := \frac{(n + s - 1) m}{s^2 l}
  \]  
  (4.1.4)

- \( plain\_phase1\_L := 1 \)
  \[
  plain\_phase1\_L := 1
  \]  
  (4.1.5)

- \( Tp\_phase\_1 := simplify(eval((plain\_phase1\_N/P + plain\_phase1\_L) \times plain\_phase1\_C, [P=plain\_phase1\_blocks])) \)
  \[
  Tp\_phase\_1 := 4 Us + 4 s^2 + 4 U - 2 s
  \]  
  (4.1.6)

**Phases 2 N & L & C**

- \( plain\_phase2\_workperthread := plain\_param \)
  \[
  plain\_phase2\_workperthread := s
  \]  
  (4.2.1)

- \( plain\_phase2\_overheadperthread := plain\_phase2\_overheadperblock \)
  \[
  plain\_phase2\_overheadperthread := 3 Us
  \]  
  (4.2.2)

- \( plain\_phase2\_C := plain\_phase2\_workperthread + plain\_phase2\_overheadperthread \)
  \[
  plain\_phase2\_C := 3 Us + s
  \]  
  (4.2.3)

- \( plain\_phase2\_N := simplify(sum(plain\_phase2\_blocks(i), i=1..plain\_phase2\_steps)) \)
  \[
  plain\_phase2\_N := \frac{(n + s - 1) (-s + m)}{s^2 l}
  \]  
  (4.2.4)

- \( plain\_phase2\_L := plain\_phase2\_steps \)
  \[
  plain\_phase2\_L := \frac{\ln\left(\frac{m}{s}\right)}{\ln(2)}
  \]  
  (4.2.5)
\[
T_{p\_phase\_2} := \left(\frac{2 \cdot (s + m)}{m} + \frac{\ln\left(\frac{m}{s}\right)}{\ln(2)}\right) (3Us + s) \tag{4.2.6}
\]

\begin{align*}
\textbf{Tp} \\
> \text{plain\_N} := \text{simplify}((\text{plain\_phase1\_N + plain\_phase2\_N}); \\
> \text{plain\_N} := \frac{(n + s - 1)(2m - s)}{s^2l} \tag{4.3.1}
\end{align*}

\begin{align*}
> \text{plain\_L} := \text{plain\_phase1\_L + plain\_phase2\_L}; \\
> \text{plain\_L} := \frac{\ln\left(\frac{m}{s}\right)}{\ln(2)} + 1 \tag{4.3.2}
\end{align*}

\begin{align*}
> \text{plain\_C} := \text{plain\_phase1\_C}; \\
> \text{plain\_C} := s(2s - 1) + U(2s + 2) \tag{4.3.3}
\end{align*}

\begin{align*}
> \text{plain\_Tp} := \text{simplify}((\text{plain\_N/P + plain\_L})\cdot\text{plain\_C}, \{P=\text{plain\_phase1\_blocks}\}); \\
> \text{plain\_Tp} := \left(\frac{2m - s}{m} + \frac{\ln\left(\frac{m}{s}\right)}{\ln(2)} + 1\right) (2Us + 2s^2 + 2U - s) \tag{4.3.4}
\end{align*}

\begin{align*}
\textbf{When} \; s = 4, \; m = n \\
> \text{plain\_work1} := \text{normal}((\text{plain\_work}, \{s=4, m=n\})); \\
> \text{plain\_work1} := 2n^2 + \frac{11}{2}n - \frac{3}{2} \tag{5.1}
\end{align*}

\begin{align*}
> \text{plain\_span1} := \text{expand}((\text{plain\_span}, \{s=4, m=n\})); \\
> \text{plain\_span1} := 20 + \frac{4\ln(n)}{\ln(2)} \tag{5.2}
\end{align*}

\begin{align*}
> \text{plain\_overhead1} := \text{normal}((\text{plain\_overhead}, \{s=4, m=n\})); \\
> \text{plain\_overhead1} := \frac{1}{8} \frac{(n + 3)U(11n - 24)}{l} \tag{5.3}
\end{align*}

\begin{align*}
> \text{plain\_K1} := \text{eval}((\text{plain\_phase1\_blocks}, \{s=4, m=n\})); \\
> \text{plain\_K1} := \frac{(n + 3)n}{16} \tag{5.4}
\end{align*}

\begin{align*}
> \text{plain\_Tp1} := \text{simplify}((\text{plain\_Tp}, \{s=4, m=n\})); \\
> \text{plain\_Tp1} := \left(\frac{2(n - 2)}{n} + \frac{-2\ln(2) + \ln(n)}{\ln(2)} + 1\right) (10U + 28) \tag{5.5}
\end{align*}
FFT-based multiplication

Input \( f, g \in \mathbb{R}[x] \) with degree less than \( n = 2^k \) and a primitive \( n \)-th root of unity \( w \in \mathbb{R} \)

\[
\begin{align*}
\text{compute } &1, w, w^2, \ldots, w^{n-1} \\
\text{let } a &:= DFT\{w\}(f) \\
\text{let } b &:= DFT\{w\}(g) \\
\text{let } c &:= ab \\
\text{return } (DFT\{w\})^{-1}(c) &= \frac{1}{n} DFT\{w^{-1}\}(c)
\end{align*}
\]

Stockham FFT algorithm

\[
DFT\{2^k\} = \prod_{i=0}^{k-1} \left( DFT\{2\} \times I\{2^{k-i-1}\} \right) \left( D\{2, 2^{k-i-1}\} \times I\{2^i\} \right)
\]

For each fixed \( 0 \leq i < k \), there are three computational steps:

**Phase 1:** \( x \to \left( I\{2\} 2^{k-i} \times I\{2^i\} \right) x \)

**Phase 2:** \( x \to \left( D\{2, 2^{k-i-1}\} \times I\{2^i\} \right) x \)

**Phase 3:** \( x \to \left( DFT\{2\} \times I\{2^{k-1}\} \right) x \)

\[> \text{FFT\_threadsperblock} := 1; \]
\[
\text{FFT\_threadsperblock} := l \quad (6.1)
\]

First compute phase 3 \( DFT\{2\} \times I\{2^{k-1}\} \) once, a thread reading 2 data + writing back the same + executing 3 bit operations, 3 additions, 2 modular add operations (5 arithmetic operations)

Second, for each \( 0 \leq i < k \), compute phase 1, 2 & 3

In phase 1, if \( 2^j < l \), a thread reading 1 data and writing back the same + executing 14 bit operations, 10 additions, 1 multiplications.

If \( 2^j \geq l \), a thread reading 1 data and writing back the same + executing 11 bit operations, 8 additions.

In phase 2, a thread reading 1 data and writing back the same + executing 5 bit operations, 4 additions, 1 modular multiply operations (11 arithmetic operations)
**Work**

\[
\text{FFT\_phase1\_blocks} := \frac{n}{\text{FFT\_threadsperblock}}; \\
\text{FFT\_phase1\_blocks} := \frac{n}{l} \\
\] (6.1.1)

\[
\text{FFT\_phase1\_workperblock} := 25 \times \text{FFT\_threadsperblock}; \\
\text{FFT\_phase1\_workperblock} := 25 l \\
\] (6.1.2)

\[
\text{FFT\_phase1\_work} := \text{FFT\_phase1\_blocks} \times \text{FFT\_phase1\_workperblock}; \\
\text{FFT\_phase1\_work} := 25 n \\
\] (6.1.3)

\[
\text{FFT\_phase2\_blocks} := \frac{n}{2 \times \text{FFT\_threadsperblock}}; \\
\text{FFT\_phase2\_blocks} := \frac{n}{2 l} \\
\] (6.1.4)

\[
\text{FFT\_phase2\_workperblock} := 20 \times \text{FFT\_threadsperblock}; \\
\text{FFT\_phase2\_workperblock} := 20 l \\
\] (6.1.5)

\[
\text{FFT\_phase2\_work} := \text{FFT\_phase2\_blocks} \times \text{FFT\_phase2\_workperblock}; \\
\text{FFT\_phase2\_work} := 10 n \\
\] (6.1.6)

\[
\text{FFT\_phase3\_blocks} := \frac{n}{2 \times \text{FFT\_threadsperblock}}; \\
\text{FFT\_phase3\_blocks} := \frac{n}{2 l} \\
\] (6.1.7)

\[
\text{FFT\_phase3\_workperblock} := 16 \times \text{FFT\_threadsperblock}; \\
\text{FFT\_phase3\_workperblock} := 16 l \\
\] (6.1.8)

\[
\text{FFT\_phase3\_work} := \text{FFT\_phase3\_blocks} \times \text{FFT\_phase3\_workperblock}; \\
\text{FFT\_phase3\_work} := 8 n \\
\] (6.1.9)

\[
\text{FFT\_work} := \text{normal(sum(FFT\_phase1\_work+FFT\_phase2\_work+FFT\_phase3\_work, i=0..\log_2(n)-2) + FFT\_phase3\_work);} \\
\text{FFT\_work} := \frac{n (-35 \ln(2) + 43 \ln(n))}{\ln(2)} \\
\] (6.1.10)

**Span**

\[
\text{FFT\_phase1\_span} := 25; \\
\text{FFT\_phase1\_span} := 25 \\
\] (6.2.1)

\[
\text{FFT\_phase2\_span} := 10; \\
\text{FFT\_phase2\_span} := 10 \\
\] (6.2.2)

\[
\text{FFT\_phase3\_span} := 8; \\
\text{FFT\_phase3\_span} := 8 \\
\] (6.2.3)

\[
\text{FFT\_span} := \text{normal(sum(FFT\_phase1\_span+FFT\_phase2\_span+FFT\_phase3\_span, i=0..\log_2(n)-2) + FFT\_phase3\_span);} \\
\text{FFT\_span} := \frac{-35 \ln(2) + 43 \ln(n)}{\ln(2)} \\
\] (6.2.4)
Overhead

\[ \text{FFT\_phase1\_overheadperblock} := 2^*U; \]
\[ \text{FFT\_phase1\_overhead} := \text{FFT\_phase1\_blocks} * \text{FFT\_phase1\_overheadperblock}; \]
\[ \text{FFT\_phase1\_overhead} := \frac{2^n U}{l} \] (6.3.1)

\[ \text{FFT\_phase2\_overheadperblock} := 2^*U; \]
\[ \text{FFT\_phase2\_overhead} := \text{FFT\_phase2\_blocks} \times \text{FFT\_phase2\_overheadperblock}; \]
\[ \text{FFT\_phase2\_overhead} := \frac{n U}{l} \] (6.3.2)

\[ \text{FFT\_phase3\_overheadperblock} := 4^*U; \]
\[ \text{FFT\_phase3\_overhead} := \text{FFT\_phase3\_blocks} \times \text{FFT\_phase3\_overheadperblock}; \]
\[ \text{FFT\_phase3\_overhead} := \frac{2^l U}{l} \] (6.3.3)

\[ \text{FFT\_phase3\_overhead} := \frac{2^l U}{l} \] (6.3.4)

\[ \text{FFT\_phase3\_overhead} := \frac{4^l U}{l} \] (6.3.5)

\[ \text{FFT\_phase3\_overhead} := \frac{4^l U}{l} \] (6.3.6)

\[ \text{FFT\_overhead} := \text{normal}(\text{sum(FFT\_phase1\_overhead+FFT\_phase2\_overhead+FFT\_phase3\_overhead, i=0..log[2](n)-2) + FFT\_phase3\_overhead);} \]
\[ \text{FFT\_overhead} := \frac{n U (-3 \ln(2) + 5 \ln(n))}{\ln(2) l} \] (6.3.7)

Tp

\[ \text{FFT\_N} := \text{normal}(\text{sum(FFT\_phase1\_blocks + FFT\_phase2\_blocks + FFT\_phase3\_blocks, i=0..log[2](n)-2) + FFT\_phase3\_blocks);} \]
\[ \text{FFT\_N} := \frac{1}{2} \frac{n (-3 \ln(2) + 4 \ln(n))}{\ln(2) l} \] (6.4.1)

\[ \text{FFT\_L} := \text{normal}(\text{sum(3, i=0..log[2](n)-2) + 1);} \]
\[ \text{FFT\_L} := \frac{-2 \ln(2) + 3 \ln(n)}{\ln(2)} \] (6.4.2)

\[ \text{FFT\_C} := \text{FFT\_phase1\_span + FFT\_phase3\_overheadperblock;} \]
\[ \text{FFT\_C} := 25 + 4 U \] (6.4.3)

\[ \text{FFT\_Tp} := \text{normal}((\text{FFT\_N}/\text{FFT\_phase1\_blocks} + \text{FFT\_L}) \times \text{FFT\_C);} \]
\[ \text{FFT\_Tp} := \frac{1}{2} \frac{-7 \ln(2) + 10 \ln(n)}{\ln(2)} (25 + 4 U) \] (6.4.4)

Point wise multiplication
PWM_threads := l;  
PWM_blocks := n / PWM_threads; 
PWM_workperblock := 6 * PWM_threads; 
PWM_work := PWM_blocks * PWM_workperblock; 
PWM_span := 6; 
PWM_overheadperblock := 3 * U; 
PWM_overhead := PWM_blocks * PWM_overheadperblock; 
PWM_N := PWM_blocks; 
PWM_L := 1; 
PWM_C := 6 + PWM_overheadperblock; 

PV threads := l; 
PV_blocks := n / PV_threads; 
PV_workperblock := 5 * PV_threads; 
PV_work := PV_blocks * PV_workperblock; 
PV_workperblock := 5 l 
PV_work := PV_blocks * PV_workperblock; 
PV_span := 5; 
PV_overheadperblock := 2 * U; 
PV_overhead := PV_blocks * PV_overheadperblock; 
PV_N := PV_blocks; 
PV_L := 1; 
PV_C := 6 + 3 U 

Scale the vector

SV_threads := l; 
SV_blocks := n / SV_threads; 
SV_workperblock := 5 * SV_threads; 
SV_work := SV_blocks * SV_workperblock; 
SV_span := 5; 
SV_overheadperblock := 2 * U;
\[ SV\_overhead := SV\_blocks \times SV\_overheadperblock; \]
\[ SV\_overhead := \frac{2nU}{l} \]  
(8.7)

\[ SV\_N := SV\_blocks; \]
\[ SV\_N := \frac{n}{I} \]  
(8.8)

\[ SV\_L := 1; \]
\[ SV\_L := 1 \]  
(8.9)

\[ SV\_C := 5 + SV\_overheadperblock; \]
\[ SV\_C := 5 + 2U \]  
(8.10)

**Work, Span & Overhead**

\[ FFT\_based\_work := 3 \times FFT\_work + PWM\_work + SV\_work; \]
\[ FFT\_based\_work := \frac{3n(-35\ln(2) + 43\ln(n))}{\ln(2)} + 11n \]  
(9.1)

\[ FFT\_based\_span := 3 \times FFT\_span + PWM\_span + SV\_span; \]
\[ FFT\_based\_span := \frac{3(-35\ln(2) + 43\ln(n))}{\ln(2)} + 11 \]  
(9.2)

\[ FFT\_based\_overhead := 3 \times FFT\_overhead + PWM\_overhead + SV\_overhead; \]
\[ FFT\_based\_overhead := \frac{3nU(-3\ln(2) + 5\ln(n))}{\ln(2)I} + \frac{5nU}{I} \]  
(9.3)

**Tp**

\[ FFT\_based\_K := FFT\_phase1\_blocks; \]
\[ FFT\_based\_K := \frac{n}{I} \]  
(10.1)

\[ FFT\_based\_N := normal(3 \times FFT\_N + PWM\_N + SV\_N); \]
\[ FFT\_based\_N := \frac{1}{2} \frac{n(-5\ln(2) + 12\ln(n))}{\ln(2)I} \]  
(10.2)

\[ FFT\_based\_L := normal(3 \times FFT\_L + PWM\_L + SV\_L); \]
\[ FFT\_based\_L := \frac{-4\ln(2) + 9\ln(n)}{\ln(2)} \]  
(10.3)

\[ FFT\_based\_C := FFT\_C; \]
\[ FFT\_based\_C := 25 + 4U \]  
(10.4)

\[ FFT\_based\_Tp := normal((FFT\_based\_N + FFT\_based\_K + FFT\_based\_L) \times FFT\_based\_C); \]  
(10.5)
\[ FFTbased_Tp := \frac{1}{2} \frac{(-13 \ln(2) + 30 \ln(n)) (25 + 4 U)}{\ln(2)} \]  \hspace{1cm} (10.5)

Plain vs FFT-based

Rwork := normal(plain_work1 / FFTbased_work);
\[ Rwork := \frac{1}{2} \frac{(4 n^2 + 11 n - 3) \ln(2)}{n (-94 \ln(2) + 129 \ln(n))} \] \hspace{1cm} (9)

Rspan := normal(plain_span1 / FFTbased_span);
\[ Rspan := \frac{4 (5 \ln(2) + \ln(n))}{-94 \ln(2) + 129 \ln(n)} \] \hspace{1cm} (10)

Roverhead := normal(plain_overhead1 / FFTbased_overhead);
\[ Roverhead := \frac{1}{8} \frac{(n+3)(11 n - 24) \ln(2)}{n (-4 \ln(2) + 15 \ln(n))} \] \hspace{1cm} (11)

Rt := normal(plain_Tp1 / FFTbased_Tp);
\[ Rt := \frac{4 (n \ln(n) + n \ln(2) - 4 \ln(2)) (5 U + 14)}{n (-13 \ln(2) + 30 \ln(n)) (25 + 4 U)} \] \hspace{1cm} (12)

plain_Tp1 := simplify(eval(eval((plain_N/P+plain_L)*plain_C, [P=FFTbased_K], [s=4, m=n])));
\[ plain_Tp1 := \left( \frac{1}{8} \frac{(n+3)(n-2)}{n} + \frac{-2 \ln(2) + \ln(n)}{\ln(2)} + 1 \right) (10 U + 28) \] \hspace{1cm} (13)

Rt := normal(plain_Tp1 / FFTbased_Tp);
\[ Rt := \frac{1}{2} \frac{(\ln(2) r^2 + 8 n \ln(n) - 7 n \ln(2) - 6 \ln(2)) (5 U + 14)}{n (-13 \ln(2) + 30 \ln(n)) (25 + 4 U)} \] \hspace{1cm} (14)
Sorting

This document is to demonstrate the analysis of radix sort.

\[ \text{restart;} \]

- For a fixed key size \( c \), radix sort on input size \( n \) is computed by \( \frac{c}{s} \) passes, and each pass sort based on \( s \) ([1..8]) bits.
- We denote \( l \) is the number of threads per block.

\[
\begin{align*}
> \text{radix\_input} & := n; & \quad \text{radix\_input} := n \quad (1) \\
> \text{radix\_keysize} & := c; & \quad \text{radix\_keysize} := c \quad (2) \\
> \text{radix\_threads} & := l; & \quad \text{radix\_threads} := l \quad (3) \\
> \text{radix\_iterations} & := s; & \quad \text{radix\_iterations} := s \quad (4) \\
> \text{radix\_blocks} & := \text{radix\_input}/(4*\text{radix\_threads}); & \quad \text{radix\_blocks} := \frac{n}{4 \ l} \quad (5) \\
> \text{radix\_buckets} & := 2^{\text{radix\_iterations}}; & \quad \text{radix\_buckets} := 2^s \quad (6)
\end{align*}
\]

\textbf{Phase 1: Each block loads and sorts its tile using \( s \) iterations of 1-bit split, and write back its \( 2^s \)-entry digit histogram and sorted data}

- Each of the \( t \) threads deals with 4 elements, so that each thread block needs \( 4 \ l \) elements to load, and writes back \( 4 \ l \) sorted elements and \( 2^s \) entry elements.
- Each thread block computes \( s \) iterations (\( c \) bits), where \( 1 \leq s \leq c \)
- In total, we have \( \frac{n}{4 \ l} \) thread blocks.

\# For each block dealing with one bit, it needs 5 steps to compute the correct position of each input element:
\# 1) Set a "1" in each "0" input;
\# 2) Scan the 1s (using prefix sum algorithm)
3) Compute the total false only once per block
4) Compute the position by 2 additions and 1 comparison
5) Scatter input to the correct position
(Refer to http://cudpp.googlecode.com/svn/trunk/doc/CUDPP_slides.pdf)

To create a $2^S$ bucket, it needs 3 steps
1) Set the bucket to 0
2) Each thread checks 5 elements and update the bucket at most 5 times
3) Each bucket updates its value to the number of elements inside the bucket

Load $4^l$ elements and write back sorted $4^l$ elements and $2^S$ values of the bucket

\[
\text{radix\_phase1\_workperblock} := \text{normal}(\text{radix\_iterations} \cdot (6 \cdot \text{radix\_threads} + 8 \cdot \text{sum}(\text{radix\_threads} / 2^{(i-1)}, i=1..\log_2(\text{radix\_threads})) + 17) + 12); \\
\text{radix\_phase1\_workperblock} := 32 \left(\frac{1}{2}\right)^{\ln^2(l) / \ln^2(2)} l s + 22 l s + 17 s + 12 \quad (1.1)
\]

\[
\text{radix\_phase1\_overheadperblock} := (4+4+1)U; \\
\text{radix\_phase1\_overheadperblock} := 9U \quad (1.2)
\]

\[
\text{radix\_phase1\_work} := \text{simplify(\text{radix\_blocks} \times \text{radix\_phase1\_workperblock})}; \\
\text{radix\_phase1\_work} := \frac{1}{4} n \frac{(22 l s + s + 12)}{l} \quad (1.3)
\]

\[
\text{radix\_phase1\_span} := \text{radix\_iterations} \cdot (4 + (\log_2(\text{radix\_threads}) + 2) \cdot 3 + (\log_2(\text{radix\_threads}) + 2) \cdot 5 + 1 + 3 \cdot 4 + 2 \cdot 4) + 5 + 5 + 1 + 1; \\
\text{radix\_phase1\_span} := s \left(\frac{41 + 8 \ln(l)}{\ln(2)}\right) + 12 \quad (1.4)
\]

\[
\text{radix\_phase1\_overhead} := \text{normal(\text{radix\_blocks} \times \text{radix\_phase1\_overheadperblock})}; \\
\text{radix\_phase1\_overhead} := \frac{9}{4} \frac{n U}{l} \quad (1.5)
\]

Phase 2: Perform a prefix sum over the histogram table, stored in column-major order
• We consider $2^S$ buckets, each having $\frac{n}{4^l}$, the results from phase 1, and perform a prefix sum over the histogram table to compute the
output positions.


```plaintext
> radix_phase2_elements := radix_blocks*radix_buckets;
  radix_phase2_elements := \( \frac{1}{4} \frac{n2^{s}}{l} \) (2.1)

> radix_phase2_blocks := radix_phase2_elements/(4*radix_threads);
  radix_phase2_blocks := \( \frac{1}{16} \frac{n2^{s}}{l^{2}} \) (2.2)

# Regard to the prefix sum, one block performs the up-sweep phase,
with 3 instructions, and then performs the down-sweep phase, with 5
instructions, regard to Listing 2 in scan.pdf. (Except mediate step)

> radix_phase2_inclusive_workperblock := normal(simplify( 8*(sum
  (radix_threads/2^(i-1),i=1..log[2](radix_threads))+2)+1 ));
  radix_phase2_inclusive_workperblock := 1 + 16 l (2.3)

> radix_phase2_inclusive_overheaderblock := (4+4)*U;
  radix_phase2_inclusive_overheaderblock := 8 U (2.4)

> radix_phase2_inclusive_work := radix_phase2_blocks*
  radix_phase2_inclusive_workperblock;
  radix_phase2_inclusive_work := \( \frac{1}{16} \frac{n2^{s}(1+16 l)}{l^{2}} \) (2.5)

> radix_phase2_inclusive_span := simplify( 8*(log[2]
  (radix_threads)+2)+1 );
  radix_phase2_inclusive_span := \( \frac{8 \ln(l)}{\ln(2)} + 17 \) (2.6)

> radix_phase2_inclusive_overhead := radix_phase2_blocks*
  radix_phase2_inclusive_overheaderblock;
  radix_phase2_inclusive_overhead := \( \frac{1}{2} \frac{n2^{s}U}{l^{2}} \) (2.7)

# Store Block Sum to Auxiliary Array, here assume the number of
blocks can fit in one shared memory, say \( \frac{1}{16} \frac{n2^{s}}{l^{2}} \leq Z \).

> radix_phase2_mediate_overheaderblock := (4+4)*U;
  radix_phase2_mediate_overheaderblock := 8 U (2.8)

> radix_phase2_mediate_work := normal(simplify( 8*(sum
  (radix_threads/2^(i-1),i=1..log[2](radix_threads))+2)+1 ));
  radix_phase2_mediate_work := 1 + 16 l (2.9)

> radix_phase2_mediate_span := 8*(log[2](radix_threads)+2)+1;
  (2.10)
```
\[\text{radix\_phase2\_mediate\_span} := \frac{8 \ln(l)}{\ln(2)} + 17 \quad (2.10)\]

\[> \text{radix\_phase2\_mediate\_overhead} := \text{radix\_phase2\_mediate\_overhead}\text{perblock};\]
\[\text{radix\_phase2\_mediate\_overhead} := 8 \ U \quad (2.11)\]

\# Scan Block Sums

\[> \text{radix\_phase2\_exclusive\_work}\text{perblock} := 4 \times \text{radix\_threads};\]
\[\text{radix\_phase2\_exclusive\_work}\text{perblock} := 4 \ l \quad (2.12)\]

\[> \text{radix\_phase2\_exclusive\_overhead}\text{perblock} := (4+1+4)\ U;\]
\[\text{radix\_phase2\_exclusive\_overhead}\text{perblock} := 9 \ U \quad (2.13)\]

\[> \text{radix\_phase2\_exclusive\_work} := (\text{radix\_phase2\_blocks}\ -1) \times \text{radix\_phase2\_exclusive\_work}\text{perblock};\]
\[\text{radix\_phase2\_exclusive\_work} := 4 \left( \frac{1}{16} n \frac{2^s}{l^2} - 1 \right) l \quad (2.14)\]

\[> \text{radix\_phase2\_exclusive\_span} := 4;\]
\[\text{radix\_phase2\_exclusive\_span} := 4 \quad (2.15)\]

\[> \text{radix\_phase2\_exclusive\_overhead} := (\text{radix\_phase2\_blocks}\ -1) \times \text{radix\_phase2\_exclusive\_overhead}\text{perblock};\]
\[\text{radix\_phase2\_exclusive\_overhead} := 9 \left( \frac{1}{16} n \frac{2^s}{l^2} - 1 \right) U \quad (2.16)\]

\# Total work & span in phase 2

\[> \text{radix\_phase2\_work} := \text{normal}(\text{radix\_phase2\_inclusive\_work}+\text{radix\_phase2\_mediate\_work}+\text{radix\_phase2\_exclusive\_work});\]
\[\text{radix\_phase2\_work} := \frac{1}{16} \frac{20 n 2^s l + 192 \ l^3 + n 2^s + 16 \ l^2}{\ l^2} \quad (2.17)\]

\[> \text{radix\_phase2\_span} := \text{simplify}(\text{radix\_phase2\_inclusive\_span}+\text{radix\_phase2\_mediate\_span}+\text{radix\_phase2\_exclusive\_span});\]
\[\text{radix\_phase2\_span} := \frac{16 \ln(l)}{\ln(2)} + 38 \quad (2.18)\]

\[> \text{radix\_phase2\_overhead} := \text{normal}(\text{radix\_phase2\_inclusive\_overhead}+\text{radix\_phase2\_mediate\_overhead}+\text{radix\_phase2\_exclusive\_overhead});\]
\[\text{radix\_phase2\_overhead} := \frac{1}{16} \frac{U \left( 17 n 2^s - 16 \ l^2 \right)}{\ l^2} \quad (2.19)\]

\textbf{Phase 3: Each block copies its elements to the correct output position}

- We have \( \frac{n}{4 \ l} \) blocks, and each block copies \( 4 \ l \) elements to the correct output position
# Each thread needs the update bucket to compute the position of the first element

\[ \text{radix\_phase3\_workperblock} := 4 \times \text{radix\_threads}; \]
\[ \text{radix\_phase3\_workperblock} := 4 l \]

\[ \text{radix\_phase3\_overheadperblock} := (4+1+4) \times U; \]
\[ \text{radix\_phase3\_overheadperblock} := 9 U \]

\[ \text{radix\_phase3\_work} := \text{radix\_blocks} \times \text{radix\_phase3\_workperblock}; \]
\[ \text{radix\_phase3\_work} := n \]

\[ \text{radix\_phase3\_span} := 4; \]
\[ \text{radix\_phase3\_span} := 4 \]

\[ \text{radix\_phase3\_overhead} := \text{radix\_blocks} \times \text{radix\_phase3\_overheadperblock}; \]
\[ \text{radix\_phase3\_overhead} := \frac{9}{4} \frac{n U}{l} \]

\[ \text{Work \& Span \& Overhead \& N \& L \& C} \]

\[ \text{radix\_work} := \text{collect}(\text{radix\_keysize}/\text{radix\_iterations} \times \]
\[ \frac{\left(1 + \frac{22 l s + s + 12}{l} + \frac{1}{16} \frac{2025 l + 2^5}{l^2} + 1 \right) n}{s} \]
\[ + \frac{1}{16} \frac{c (192 l^2 + 16 l^2)}{s l^2} \]

\[ \text{radix\_span} := \text{normal}(\text{radix\_keysize}/\text{radix\_iterations} \times \]
\[ \frac{c (8 s \ln(l) + 41 s \ln(2) + 16 \ln(l) + 54 \ln(2))}{s \ln(2)} \]

\[ \text{radix\_overhead} := \text{normal}(\text{radix\_keysize}/\text{radix\_iterations} \times \]
\[ \frac{c U (17 n^2 - 16 l^2 + 72 n l)}{s l^2} \]

\[ \text{radix\_N} := \text{radix\_keysize}/\text{radix\_iterations} \times (\text{radix\_blocks} + 2 \times \]
\[ \text{radix\_phase2\_blocks} + \text{radix\_blocks}); \]
\[ \text{radix\_N} := \frac{c \left(\frac{1}{2} \frac{n}{l} + \frac{1}{8} \frac{n^2 l^2}{l} \right)}{s} \]

\[ \text{radix\_L} := \text{radix\_keysize}/\text{radix\_iterations} \times (1+3+1); \]
\[
radix_L := \frac{5c}{s} \\
\]
\[
radix_C := \frac{s}{(41 + \frac{8\ln(l)}{\ln(2)})} + 12 + 9U \\
\]
\[
radix_Tp := \text{simplify}\left(\frac{radix_N}{radix_blocks + radix_L} \cdot radix_C\right) \\
\]
\[
radix_Tp := \frac{1}{2} \cdot \frac{c(2^s + 14l)(8s\ln(l) + 9U\ln(2) + 41s\ln(2) + 12\ln(2))}{ls\ln(2)} \\
\]

\textbf{When } s = 1

\[
radix_work_1 := \text{simplify}\left(\text{eval}\left(radix_work, [s=1]\right)\right) \\
\]
\[
radix_work_1 := \frac{c(96\beta + 52\beta^2n + 8\beta^2 + 46ln(n) + n)}{\beta^4} \\
\]
\[
radix_overhead_1 := \text{normal}\left(\text{eval}\left(radix_overhead, [s=1]\right)\right) \\
\]
\[
radix_overhead_1 := -\frac{1}{8} c\left(8\beta^2 - 36\ln(17n)\right) \frac{1}{\beta^2} \\
\]
\[
radix_span_1 := \text{normal}\left(\text{eval}\left(radix_span, [s=1]\right)\right) \\
\]
\[
radix_span_1 := \frac{c(24\ln(l) + 95\ln(2))}{\ln(2)} \\
\]
\[
radix_Tp_1 := \text{simplify}\left(\text{eval}\left(radix_Tp, [s=1]\right)\right) \\
\]
\[
radix_Tp_1 := \frac{c(1 + 7l)(8\ln(l) + 9U\ln(2) + 53\ln(2))}{l\ln(2)} \\
\]

\textbf{Comparison}

\[8l + 2^s \leq Z\]

\[
radix_work_ratio := \text{simplify}\left(\frac{radix_work_1}{radix_work}\right) \\
\]
\[
radix_work_ratio := \frac{2(96\beta + 52\beta^2n + 8\beta^2 + 46ln(n) + n)s}{88s\beta^2n + 20n2^sn + 192\beta^2 + 16\beta^2n + 4lsn + n2^s + 16\beta^2 + 48nl} \\
\]
\[
radix_overhead_ratio := \text{collect}\left(\frac{radix_overhead_1}{radix_overhead}, n\right) \\
\]
\[
radix_overhead_ratio := -\frac{2((-36l - 17)n + 8\beta^2)s}{(172^s + 72l)n - 16\beta^2} \\
\]
\[
radix_span_ratio := \text{simplify}\left(\frac{radix_span_1}{radix_span}\right) \\
\]
\[
radix_span_ratio := \frac{24\ln(l) + 95\ln(2)s}{8s\ln(l) + 41s\ln(2) + 16\ln(l) + 54\ln(2)} \\
\]

Assume \(2^s = O(l)\), we can reduce overhead by \(s\)
\[ \text{radix}_{\text{Tp}}_{\text{R}} := \text{simplify}(\text{radix}_{\text{Tp}}_{\text{1}}/\text{radix}_{\text{Tp}}) \]
\[ \text{radix}_{\text{Tp}}_{\text{R}} = \frac{2 (1 + 7 l) (8 \ln(l) + 9 U \ln(2) + 53 \ln(2)) s}{(2^8 + 14 l) (8 s \ln(l) + 9 U \ln(2) + 41 s \ln(2) + 12 \ln(2))} \quad (6.4) \]
\[ \text{simplify(eval(radix}_{\text{Tp}}_{\text{R}}, [s = \text{log[2](l)]]));} \]
\[ \frac{2}{15} \frac{(1 + 7 l) (8 \ln(l) + 9 U \ln(2) + 53 \ln(2)) \ln(l)}{l (9 U \ln(2)^2 + 8 \ln(l)^2 + 41 \ln(l) \ln(2) + 12 \ln(2)^2)} \quad (6.5) \]
\[ \text{lcN} := \text{collect(expand(2*(1+7*l)*(8*ln(l)+9*U+53)*ln(l)), l)}; \]
\[ \text{lcN} := (112 \ln(l)^2 + 126 \ln(l) U + 742 \ln(l)) l + 16 \ln(l)^2 + 18 \ln(l) U + 106 \ln(l) \quad (6.6) \]
\[ \text{lcD} := 15 l (9 U + 8 \ln(l)^2 + 41 \ln(l) + 12); \]
\[ \text{lcD} := 15 l (9 U + 8 \ln(l)^2 + 41 \ln(l) + 12) \quad (6.7) \]
\[ \text{lcN} := 112 \ln(l)^2 + 126 \ln(l) U + 742 \ln(l); \]
\[ \text{lcN} := 112 \ln(l)^2 + 126 \ln(l) U + 742 \ln(l) \quad (6.8) \]
\[ \text{lcD} := \text{expand}(15*(9*U+8*ln(l)^2+41*ln(l)+12)); \]
\[ \text{lcD} := 135 U + 120 \ln(l)^2 + 615 \ln(l) + 180 \quad (6.9) \]
\[ \text{simplify(lcN/lcD)}; \]
\[ \frac{14}{15} \frac{\ln(l) (8 \ln(l) + 9 U + 53)}{9 U + 8 \ln(l)^2 + 41 \ln(l) + 12} \quad (6.10) \]
\[ \text{expand(14*(8*ln(l))+9*U+53));} \]
\[ 112 \ln(l) + 126 U + 742 \quad (6.11) \]
\[ \text{expand(15*(8*ln(l))+41));} \]
\[ 120 \ln(l) + 615 \quad (6.12) \]
Appendix C

Documentation for MetaFork-to-CUDA Code Generator

Our MetaFork-to-CUDA code generator is based on the version 0.04 of PPCG\(^1\). The code generation follows two algorithms, depending on whether one intends to use the shared memory or not in the kernel code. In the sequel, we refer to these algorithms as the shared memory and global memory modes of the code generator. These two algorithms are available to the user as two different targets of the Makefile for compiling our code generator. By default, the compilation of MetaFork programs, with our extended version of PPCG, uses the global memory only. This mode is compiled by running make at the root of the source tree of PPCG. To enable the use of the shared memory, one should compile the code generator by issuing make mem=mlocal.

In this documentation, we report the assumptions within our MetaFork-to-CUDA code generator in Appendix C.1, and the schedule tree used as a mathematical representation for MetaFork and parametric CUDA code in Appendix C.2.

C.1 Assumptions on the syntax of MetaFork statements

In order to use MetaFork-to-CUDA code generator without post-processing, we make assumptions on the syntax of MetaFork statements as below.

```
meta.schedule { ... }
```

This statement indicates its body will be launched to hardware accelerators, i.e. NVIDIA GPUs. It also transfers the data from CPU to GPU before launching kernels and transfers the data back from GPU to CPU after executing kernels. Furthermore, data transfer between CPU and GPU is automatically detected by this statement.

Statements supported within the body of the meta.schedule statement are a sequence of nested loops. Thus, each nested for loops consist of parallel for loops (only 2 or 4) and/or serial for loops, and will be translated into a kernel call. In the case of parallel for loops, it is identified with the ‘meta_for’ keyword.

\(^{1}\)PPCG’s original code is available at [https://www.openhub.net/p/ppcg](https://www.openhub.net/p/ppcg)
meta_for (initialize; condition; increment) { ...
initialize: 0
condition: < (a variable)
increment: ++ or += 1
For instance, meta_for (int i = 0; i < upper_bound; i++) { ...
}

The upper bound in the condition indicates either the number of threads per thread-block or the number of thread-blocks per grid. Thus, for launching a one-dimension kernel, it requires one outer meta_for loop specifying the grid size and one inner meta_for loop specifying the thread-block size. For a two-dimension kernel, two (immediately) consecutive outer meta_for loops are used to specify the grid sizes, and two (immediately) consecutive inner meta_for loops are used to specify the thread-block sizes. Moreover, the iterators in the first and second outer (resp. inner) meta_for loops correspond to blockIdx.y and blockIdx.x (resp. threadIdx.y and threadIdx.x), respectively, in the generated kernel code. Figure C.1 shows an example of a meta_schedule statement with a 1-D kernel and a 2-D kernel.

meta_schedule {
    // only for loops are supported here
    meta_for (int i = 0; i < gridDim.x; i++)
        // only for loops are supported here
        meta_for (int j = 0; j < blockDim.x; j++) {
            ... // nested for-loop body
        }
    // only for loops are supported here
    meta_for (int u = 0; u < gridDim.y; u++)
        meta_for (int i = 0; i < gridDim.x; i++)
            // only for loops are supported here
            meta_for (int v = 0; v < blockDim.y; v++)
                meta_for (int j = 0; j < blockDim.x; j++) {
                    ... // nested for-loop body
                }
}

Figure C.1: An example of the meta_schedule statement

for (initialize; condition; increment) { ...

For those serial for loops, the upper or lower bound of condition should be a linear expression and increment should be increased or decreased by a constant. This observation is based on the work reported in [116].

array[expression] or array[expression][expression]
C.2 Schedule tree for MetaFork and parametric CUDA code

In the global memory mode, one can only use linear expressions as the indices (expression) of 1D or 2D arrays. However, one can hide non-linear expressions by using a separate statement, which yields a linear expression that can be used as the indices. For instance, int p = i * B + j * s + k; array[p] = ···. Note that non-linear expressions cannot be analyzed by PPCG, such that whether array[p] is reused or coalesced accessed is unknown to PPCG.

In the shared memory mode, due to the lack of analysis of non-linear expressions, we make the following assumptions.

1. If expression is a linear expression, all its variables must refer to the variable counters in the serial for loops of the current loop nest. In this case, we rely on PPCG to analyze the access patterns of the corresponding array.

2. If expression contains one and only one non-linear term, that is, one variable multiplying by another, say i * B, then one variable must refer to the current thread-block index and the other must refer to the corresponding thread-block size. One shall hide this non-linear expression by using a separate statement, while this statement should add a third variable referring to the thread index. For instance, int p = i * B + j; array[p] = ···. Furthermore, adding a constant to the above format of the index of array[] is supported, say array[p] = array[p+1]. For this format, we extend the analysis calculating the total amount of required shared memory per thread-block. Particularly, for a 2D array, the first (resp. second) expression refers to the first (resp. second) dimension of grids and thread-blocks, defined by the first (resp. second) outer and inner meta_for loops, respectively.

3. No other forms of non-linear expressions are accepted in expression.

Moreover, in the shared memory mode, not all arrays occurring in the MetaFork source code will necessarily use the shared memory in the generated CUDA code. In fact, in the addition to the syntax constraints described before, one of the following conditions must also hold:

1. If array[] (resp. array[][][]) is written more than once and threads access it in a coalesced fashion, then a shared memory counterpart of array[] (resp. array[][][]) will be generated.

2. If array[] (resp. array[][][]) is read and threads access it in a coalesced fashion, then a shared memory counterpart of array[] (resp. array[][][]) will be generated.

When none of those conditions is satisfied, then shared memory counterpart of array[] (resp. array[][][]) is not generated.

C.2 Schedule tree for MetaFork and parametric CUDA code

Sven Verdoolaege, et al [117] proposed to explicitly represent schedules in the polyhedral model as a tree structure. We take advantage of this structure with minor modifications, such that it can represent the execution order of a MetaFork program as well as be converted to the AST for parametric CUDA kernel code generation.

For each schedule tree, there are three catalogues of node types, including core, external and convenience nodes. For core nodes, a band node indicates multi-dimensional piecewise quasi-affine partial schedule, a filter node selects statement instances that are executed by descendants, a sequence node defines its children executed in a given order, and a set node has its
children executed in an arbitrary order. For those external node types, a domain node includes a set of statement instances to be scheduled, and a context node specifies external constraints on symbolic constants. For convenience node types, one can attach additional information to subtrees as a mark node.

```c
int ub_v = N / B;
meta_schedule {
    meta_for (int v = 0; v < ub_v; v++)
        meta_for (int u = 0; u < B; u++) {
            int inoffset = v * B + u; // S_1
            int outoffset = N - 1 - inoffset; // S_4
            Out[outoffset] = In[inoffset]; // S_6
        }
}
```

Consider the above MetaFork code for one-dimensional array reversal. Within the nested for loops, there are three statements referred as S_1, S_4, S_6, respectively, in the schedule tree below. This schedule tree is initialized based on the original MetaFork code. Two schedule nodes correspond to two meta_for loops, respectively, while we set “permutable” and “coincident” of each of these two schedule nodes to be 1s. Our goal is to extend this schedule tree so as to obtain another “rich” schedule tree for its corresponding CUDA program.

domain: "[N, ub_v, B] -> { S_6[v, u] : B >= 0 and v >= 0 and v <= -1 + ub_v and u >= 0; S_1[v, u] : B >= 0 and v >= 0 and v <= -1 + ub_v and u >= 0; S_4[v, u] : B >= 0 and v >= 0 and v <= -1 + ub_v and u >= 0 }
child:
    permutable: 1
    coincident: [ 1 ]
child:
    permutable: 1
    coincident: [ 1 ]
child: sequence:
    - filter: "[N, ub_v, B] -> { S_1[v, u] }"
    - filter: "[N, ub_v, B] -> { S_4[v, u] }"
    - filter: "[N, ub_v, B] -> { S_6[v, u] }

The schedule tree shown below is extended from the above schedule tree for generating both the host and device codes based on our MetaFork code. We first insert a mark node, named “kernel,” as on Line 16 below that indicates where the kernel call starts. Then, for each thread-block, we refer variable b0 to the thread-block index blockIdx.x for a one-dimensional kernel, while for a two-dimensional kernel, we refer variable b0 (resp. b1) to blockIdx.y (resp. blockIdx.x), which corresponds to the first (resp. second) outer meta_for loop. Due to
the fact that the number of thread-blocks per grid depends on the problem size, which is an arbitrary number, but that the maximum number of thread-blocks supported by streaming multiprocessors (SMs) is a fixed number, we shall generate a serial loop for each thread-block to execute in sequence, in case the number of required thread-blocks exceeds the hardware limit. Thus, we insert a filter node as on Line 21 of the schedule tree below, which means that there exists an instance for each thread-block starting from $b0$ and incrementing by 32768 (that is, the hardware limit).

```
domain: ",N, ub_v, B] -> \{ kernel0[] : ub_v >= 1 and B >= 0 \}"
```

```
child:
  context: "[N, ub_v, B] -> \{ [] : N <= 2147483647 and N >= 0 and ub_v <= 2147483647 and B <= 2147483647 \}"

child:
  guard: "[N, ub_v, B] -> \{ [] : ub_v >= 1 and B >= 0 and N <= 2147483647 and N >= 0 and ub_v <= 2147483647 and B <= 2147483647 \}"

child:
  contraction: "[N, ub_v, B] -> \{ S_6[v, u] -> kernel0[]; S_1[v, u] -> kernel0[]; S_4[v, u] -> kernel0[] \}"

child:
  expansion: "[N, ub_v, B] -> \{ kernel0[] -> S_6[v, u] : B >= 0 and v >= 0 and v <= -1 + ub_v and u >= 0; kernel0[] -> S_1[v, u] : B >= 0 and v >= 0 and v <= -1 + ub_v and u >= 0; kernel0[] -> S_4[v, u] : B >= 0 and v >= 0 and v <= -1 + ub_v and u >= 0 \}"

child:
  mark: "kernel"

child:
  context: "[N, ub_v, B, b0, t0] -> \{ [] : b0 <= -1 + ub_v and b0 <= 32767 and b0 >= 0 and t0 >= 0 and t0 <= 511 \}"

child:
  filter: "[N, ub_v, B, b0] -> \{ S_1[v, u] : exists (e0 = floor((-b0 + v)/32768): 32768e0 = -b0 + v and b0 >= 0 and b0 <= 32767); S_4[v, u] : exists (e0 = floor((-b0 + v)/32768): 32768e0 = -b0 + v and b0 >= 0 and b0 <= 32767); S_6[v, u] : exists (e0 = floor((-b0 + v)/32768): 32768e0 = -b0 + v and b0 >= 0 and b0 <= 32767) \}"

child:
```

permutable: 1

coincident: [ 1 ]

child:
  extension: "[N, ub_v, B, b0] -> \{ [i0] -> sync0[]; [i0] -> read[[i0] -> In[i0]] : o1 >= B + 1024i0 and o1 <= -1 + N and o1 >= 0; [i0] -> sync1[] \}"

child:
  sequence:
    - filter: "[N, ub_v, B, b0, t0] -> \{ read[[i0] -> In[B + t0 + 1024i0]] \}"
```
child:
  schedule: "[N, ub_v, B, b0] -> shared_In([{ read[[i0] -> In[i]]} -> [(B - 1024i0 + i1)] }]
- filter: "{ sync1[] }
- filter: "[N, ub_v, B, b0] -> { S_1[v, u]; S_4[v, u]; S_6[v, u] }
child:
  filter: "[N, ub_v, B, t0] -> { S_4[v, t0]; S_6[v, t0]; S_1[v, t0] }
child:
  permutable: 1
  coincident: [ 1 ]
  child:
    sequence:
      - filter: "[N, ub_v, B] -> { S_1[v, u] }
      - filter: "[N, ub_v, B] -> { S_4[v, u] }
      - filter: "[N, ub_v, B] -> { S_6[v, u] }
      - filter: "{ sync0[] }

Regarding the thread index variables, we refer variable \( t0 \) to threadIdx.x for a one-dimensional kernel, while for a two-dimensional kernel, we refer variable \( t0 \) (resp. \( t1 \)) to threadIdx.y (resp. threadIdx.x), which corresponds to the first (resp. second) inner MetaFork loop. The maximum number of threads per thread-block is also limited by the hardware architecture. However, when the value of the program parameter \( B \), which indicates the thread-block format, exceeds the hardware limit, the compilation fails when one intends to launch the kernel from the host code. Thus, although in the schedule tree, we explicitly say that \( t0 \) is bounded by 512 (that is, the hardware limit) on the context node as shown on Line 18, this information is ignored by the subtree of that node. The task to control the value of \( B \) within the hardware limit is left to the user.

For this array reversal example, since we would like to use the shared memory for the input array \( \text{In} \) in the kernel, we need to insert relevant nodes in the schedule tree corresponding to copying data statements from the global memory to the shared memory. As shown on Lines 39-40, we insert a filter node that indicates the new statement reading from array \( \text{In}[B + t0 + 1024i0] \). However, this is a trick that we use to encode the non-linear expression \( t0 + B \times i0 \) here, but we later replace it during the phase of translating the AST to the generated code. Then, a schedule node inserted on Lines 42-43 specifies an array \( \text{shared}_\text{In} \) to be allocated in the shared memory. In addition, the allocation unit size of this array \( \text{shared}_\text{In} \) is defined by \( o1 \) on the extension node as on Line 33, while it is bounded by 1024 but later replaced by BLOCK_0 during the kernel code generation phase. Note that BLOCK_0 is predefined as a macro and specified its value at compile time.

Within the kernel mark node, a synchronization filter node is inserted on Line 44 after copy statements as well as on Line 60 after the computation statements. By the end, we insert several nodes, such as domain and guard, to provide the kernel information for the host code. The automatically generated CUDA kernel code for array reversal is shown in Figure 5.6 of
Chapter 5

Turning our attention to the 1D Jacobi example as shown in Figure 5.3 of Chapter 5, we observe that each thread-block of the first kernel shall read $B+2$ elements of the input array, compute the average and update the result to an intermediate array. In the kernel code, we intend to use the shared memory for the input, while PPCG uses the global memory. Thereby, we extend the analysis for using the shared memory, such that when the total amount of required shared memory is based on the number of threads per thread-block and a constant (if applicable), we can allocate a corresponding array in the shared memory. Consequently, the following nodes for copy statements are inserted into the schedule tree for the first CUDA kernel of 1D Jacobi.

- filter: 
  
  \[
  \begin{align*}
  &[[N, T, \text{ub\_v}, B, b0, t0]] \rightarrow \{ \text{read}[[i0, i1]] \rightarrow \text{a}[i2] \} : \exists e0 = \text{floor}((-B - t0 + i2)/1024) : 1024e0 = -B - t0 + i2 \text{ and } t0 \geq 0 \text{ and } t0 \leq 1023) \\
  \end{align*}
  \]

  child:

  schedule: 
  
  \[
  \begin{align*}
  &[[N, T, \text{ub\_v}, B, b0]] \rightarrow \text{shared\_a}[\{ \text{read}[[i0, i1]] \rightarrow \text{a}[i2] \} \\
  &\rightarrow [(-B - 1024\text{i1} + i2)] \\
  \end{align*}
  \]

  The above filter node indicates that a for loop will be generated to read for each thread, since the total amount of required shared memory per thread-block is more than the number of threads per thread-block. Moreover, the upper bound of this for loop and the allocation unit size of the array that is declared in the shared memory are defined by $o2$ on the extension node below. Note that there is one $B$ used to replace 1024 so as to form the non-linear expression $B \times i1$. One can easily calculate from the upper and lower bounds of $o2$ the amount $B+2$, which is used as the upper bound of the for loop for the copy statements and is replaced by BLOCK\_0+2 for allocating the array in the shared memory. The automatically generated CUDA kernel code for 1D Jacobi is shown in Figure 5.4 of Chapter 5.

  extension: 

  \[
  \begin{align*}
  &[[N, T, \text{ub\_v}, B, b0]] \rightarrow \{ \text{[i0, i1]} \rightarrow \text{sync1[]} ; \\
  &\text{[i0, i1]} \rightarrow \text{sync0[]} ; \text{[i0, i1]} \rightarrow \text{read}[[i0, i1]] \rightarrow \text{a}[o2] \} : \\
  &o2 \geq B + 1024i1 \text{ and } o2 \leq 1 + 2B + 1024i1 \text{ and } N \geq 1 \text{ and } \\
  &T \geq 1 \text{ and } \text{ub\_v} \geq 1 \text{ and } B \geq 0 \text{ and } o2 \leq -1 + N \text{ and } o2 \geq 0 \}
  \end{align*}
  \]
Appendix D

Examples Generated by PPCG

We present PPCG code with generated CUDA kernels for eight examples: array reversal (Figure D.1), 1D Jacobi (Figure D.3), 2D Jacobi (Figure D.4), LU decomposition (Figure D.5), matrix transposition (Figure D.7), matrix addition (Figure D.2), matrix vector multiplication (Figure D.6), and matrix matrix multiplication (Figure D.8).

```cpp
#pragma scop
for (int i = 0; i < N; i++)
    Out[N - 1 - i] = In[i];
#pragma endscop

__global__ void kernel0(int *In, int *Out, int N) {
    int b0 = blockIdx.x;
    int t0 = threadIdx.x;
    __shared__ int shared_Out[32];
    #define floord(n,d) (((n)<0) ? -((-(n)+(d)-1)/(d)) : (n)/(d))
    for (int c0 = 32 * b0; c0 < N; c0 += 1048576) {
      __syncthreads();
      if (N >= t0 + c0 + 1)
        shared_Out[-t0 + 31] = In[t0 + c0];
      __syncthreads();
      if (N + t0 >= c0 + 32)
        Out[N + t0 - c0 - 32] = shared_Out[t0];
    }
}

Figure D.1: PPCG code and generated CUDA kernel for array reversal
```

```cpp
#pragma scop
for (int v0 = 0; v0 < n; v0++)
    for (int v1 = 0; v1 < n; v1++)
        c[v0][v1] = a[v0][v1] + b[v0][v1];
#pragma endscop

__global__ void kernel0(int *a, int *b, int *c, int n) {
    int b0 = blockIdx.y, b1 = blockIdx.x;
    int t0 = threadIdx.y, t1 = threadIdx.x;
    #define floord(n,d) (((n)<0) ? -((-(n)+(d)-1)/(d)) : (n)/(d))
    #define min(x,y) ((x) < (y) ? (x) : (y))
    for (int c0 = 32 * b0; c0 < n; c0 += 8192)
        for (int c1 = 32 * b1; c1 < n; c1 += 8192)
            for (int c3 = t1; c3 <= min(31, n - c1 - 1); c3 += 16)
                c[(t0 + c0) * n + (c1 + c3)] =
                    (a[(t0 + c0) * n + (c1 + c3)] +
                    b[(t0 + c0) * n + (c1 + c3)]);
}

Figure D.2: PPCG code and generated CUDA kernel for matrix addition
```
#pragma scop
for (int t = 0; t < T; ++t) {
    for (int i = 1; i < N-1; ++i)
        b[i] = (a[i-1] + a[i] + a[i+1]) / 3;
    for (int i = 1; i < N-1; ++i)
        a[i] = b[i];
}
#pragma endscop

__global__ void kernel0(int *a, int *b, int N, int T, int c0)
{
    int b0 = blockIdx.x;
    int t0 = threadIdx.x;
    #define floord(n,d) (((n)<0) ? -((-(n)+(d)-1)/(d)) : (n)/(d))
    for (int c1 = 32 * b0; c1 < N - 1; c1 += 1048576)
        if (N >= t0 + c1 + 2 && t0 + c1 >= 1)
            b[t0 + c1] = (((a[t0 + c1 - 1] + a[t0 + c1]) +
                a[t0 + c1 + 1]) / 3);
}

__global__ void kernel1(int *a, int *b, int N, int T, int c0)
{
    int b0 = blockIdx.x;
    int t0 = threadIdx.x;
    #define floord(n,d) (((n)<0) ? -((-(n)+(d)-1)/(d)) : (n)/(d))
    for (int c1 = 32 * b0; c1 < N - 1; c1 += 1048576)
        if (N >= t0 + c1 + 2 && t0 + c1 >= 1)
            a[t0 + c1] = b[t0 + c1];
}

Figure D.3: PPCG code and generated CUDA kernel for 1D Jacobi

#pragma scop
for (int t = 0; t < T; t++) {
    for (int i = 0; i < N-2; i++)
        for (int j = 0; j < N-2; j++)
            b[i+1][j+1] = (a[i][j+1] +
                a[i+1][j+1] + a[i+1][j] + a[i+1][j+2]) / 4;
    for (int i = 0; i < N-2; i++)
        for (int j = 0; j < N-2; j++)
            a[i+1][j+1] = b[i+1][j+1];
}
#pragma endscop

__global__ void kernel0(int *a, int *b, int N, int T, int c0)
{
    int b0 = blockIdx.y, b1 = blockIdx.x;
    int t0 = threadIdx.y, t1 = threadIdx.x;
    #define floord(n,d) (((n)<0) ? -((-(n)+(d)-1)/(d)) : (n)/(d))
    #define min(x,y) ((x) < (y) ? (x) : (y))
    for (int c1 = 32 * b0; c1 < N - 2; c1 += 8192)
        if (N >= t0 + c1 + 3)
            for (int c2 = 32 * b1; c2 < N - 2; c2 += 8192)
                for (int c4 = t1; c4 <= min(31, N - c2 - 3); c4 += 16)
                    b[(t0 + c1 + 1) * N + (c2 + c4 + 1)] =
                        (((a[(t0 + c1) * N + (c2 + c4 + 1)] +
                            a[(t0 + c1 + 2) * N + (c2 + c4 + 1)]) +
                            a[(t0 + c1 + 1) * N + (c2 + c4)]) +
                            a[(t0 + c1 + 1) * N + (c2 + c4 + 2)]) / 4;
}

__global__ void kernel1(int *a, int *b, int N, int T, int c0)
{
    int b0 = blockIdx.y, b1 = blockIdx.x;
    int t0 = threadIdx.y, t1 = threadIdx.x;
    #define floord(n,d) (((n)<0) ? -((-(n)+(d)-1)/(d)) : (n)/(d))
    #define min(x,y) ((x) < (y) ? (x) : (y))
    for (int c1 = 32 * b0; c1 < N - 2; c1 += 8192)
        if (N >= t0 + c1 + 3)
            for (int c2 = 32 * b1; c2 < N - 2; c2 += 8192)
                for (int c4 = t1; c4 <= min(31, N - c2 - 3); c4 += 16)
                    a[(t0 + c1 + 1) * N + (c2 + c4 + 1)] =
                        b[(t0 + c1 + 1) * N + (c2 + c4 + 1)];
}

Figure D.4: PPCG code and generated CUDA kernel for 2D Jacobi
Chapter D. Examples Generated by PPCG

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**Figure D.5:** PPCG code and generated CUDA kernel for LU decomposition

```c
#pragma scop
for (int k = 0; k < n; ++k) {
    for (int i = 0; i < n-k-1; i++) {
        // column major representation
        // of L and U
        int p = i + k + 1;
        L[k][p] = U[k][p] / U[k][k];
        for (int j = k; j < n; j++)
            U[j][p] -= L[k][p] * U[j][k];
    }
}
#pragma endscop

__global__ void kernel0(double *L, double *U, int n, int c0)
{
    int b0 = blockIdx.x;
    int t0 = threadIdx.x;
    __shared__ double shared_U_1[1][1];
    #define floord(n,d) (((n)<0) ? -((-(n)+(d)-1)/(d)) : (n)/(d))
    #define min(x,y) ((x) < (y) ? (x) : (y))
    #define max(x,y) ((x) > (y) ? (x) : (y))
    if (n + 30 >= ((32 * b0 + 8191 * c0 + 31) \% 8192) + c0)
        for (int c1 = 32 * b0; c1 < n - c0 - 1; c1 += 8192) {
            if (t0 == 0)
                for (int c3 = t1; c3 <= min(31, n - c0 - c1 - 2); c3 += 16)
                    shared_L[0][c3] = L[c0 * n + (c0 + c1 + c3 + 1)];
            __syncthreads();
            for (int c2 = 32 * b0 + 8192 * ((-32 * b1 + 8160) \% 8192) + c0)
                if (t1 == 0 & & n >= t0 + c2 + 1)
                    shared_U_1[t0][0] = U[(t0 + c2) * n + c0];
            __syncthreads();
            if (n >= t0 + c0 + c1 + 2)
                for (int c4 = max(t1, t1 + 16 * floord(-t1 + c0 - c2 - 1, 16) + 16); c4 <= min(31, n - c2 - 1); c4 += 16)
                    U[(c2 + c4) * n + (t0 + c0 + c1 + 1)] -=
                        (shared_L[0][t0] * shared_U_1[c4][0]);
            __syncthreads();
        }
    __syncthreads();
}
```

---

Figure D.5: PPCG code and generated CUDA kernel for LU decomposition
__global__ void kernel0(int *a, int *b, int *c, int n)
{
    int b0 = blockIdx.x;
    int t0 = threadIdx.x;
    __shared__ int shared_a[32][32];
    __shared__ int shared_b[32];
    int private_c[1];
    #define floord(n,d) (((n)<0) ? -(((n)+(d)-1)/(d)) : (n)/(d))
    #define min(x,y) ((x) < (y) ? (x) : (y))
    for (int c0 = 32 * b0; c0 < n; c0 += 1048576) {
        for (int c1 = 0; c1 < n; c1 += 32) {
            if (n >= t0 + c1 + 1) {
                for (int c2 = 0; c2 <= min(31, n - c0 - 1); c2 += 1)
                    shared_a[c2][t0] = a[(c0 + c2) * n + (t0 + c1)];
                shared_b[t0] = b[t0 + c1];
            }
            __syncthreads();
            if (n >= t0 + c0 + 1 && c1 == 0)
                private_c[0] = 0;
            if (n >= t0 + c0 + 1)
                for (int c3 = 0; c3 <= min(31, n - c1 - 1); c3 += 16)
                    private_c[0] += (shared_a[c3][t0] * shared_b[c3]);
            __syncthreads();
        }
        if (n >= t0 + c0 + 1)
            c[(t0 + c0) * n + (t0 + c0)] = private_c[0];
        __syncthreads();
    }
}

Figure D.6: PPCG code and generated CUDA kernel for matrix vector multiplication

__global__ void kernel0(int *a, int *c, int n)
{
    int b0 = blockIdx.y, b1 = blockIdx.x;
    int t0 = threadIdx.y, t1 = threadIdx.x;
    __shared__ int shared_a[32][32];
    #define floord(n,d) (((n)<0) ? -(((n)+(d)-1)/(d)) : (n)/(d))
    #define min(x,y) ((x) < (y) ? (x) : (y))
    for (int c0 = 32 * b0; c0 < n; c0 += 8192) {
        for (int c1 = 32 * b1; c1 < n; c1 += 8192) {
            if (n >= t0 + c1 + 1)
                for (int c3 = t1; c3 <= min(31, n - c0 - 1); c3 += 16)
                    shared_a[t0][c3] = a[(t0 + c1) * n + (c0 + c3)];
            __syncthreads();
            if (n >= t0 + c0 + 1)
                for (int c3 = t0; c3 <= min(31, n - c1 - 1); c3 += 16)
                    c[(t0 + c0) * n + (c1 + c3)] = shared_a[c3][t0];
            __syncthreads();
        }
    }
}

Figure D.7: PPCG code and generated CUDA kernel for matrix transpose
Figure D.8: PPCG code and generated CUDA kernel for matrix matrix multiplication
Appendix E

The Implementation for Generating Comprehensive MetaFork Programs

In this appendix, we exhibit the pseudocode of the preliminary implementation of the comprehensive optimization algorithm demonstrated in Chapter 7. Algorithm 7 is the implemented algorithm for generating comprehensive MetaFork programs from a given MetaFork program, while Algorithm 8 and Algorithm 9 comprise the implemented Optimize procedure. In this implementation, we consider two resource counters: register usage per thread and data amount per thread-block to be cached in the shared memory; meanwhile, we apply three optimization strategies, including reducing register pressure, controlling thread granularity, and common sub-expression elimination.

Algorithm 7: MultiParametricCodeOptimizer(fileName)

Input: fileName, giving the location of the input program
Output: optimized versions of the input program in the form of a case discussion (depending on the hardware resource limits)

1 plans := [Create_Optimization_Plan(fileName)];
2 results := [];
3 while the number of plans <> 0 do
4     plan := plans[1]; plans := plans[2..-1];
5     task := ExtractTask(plan) [1];
6     new_plan := Optimize(plan, task);
7     for new_plan in new_plan do
8         if IsCompleted(new_plan) then
9             results := [new_plan, op(results)];
10        else
11            plans := [new_plan, op(plans)];
12 return results;
Algorithm 8: Optimize(plan, task)

Input: plan, encoding a program being optimized, and task, an optimization task of plan
Output: A list of new plans obtained by optimizing plan according to task

1. local caching_task, granularity_task, register_task, new_plans, optimized_plans, current_vars, alternative;
2. new_plans := [ ];
3. if task[NAME] = “Register_Pressure_Control” and task[CURRENTLEVEL] <= task[FINALLEVEL] then
   alternative := Copy_Optimization_Plan(plan);
   r := RegisterPressure(plan,task[CURRENTLEVEL]);
   /* Accept case */
   plan[CONSTRAINTS] := ['<='(r, R_B), op(plan[CONSTRAINTS]) ];
   if IsConsistent(plan) then
      register_task := FindTask(plan, “Register_Pressure_Control”);
      register_task[CURRENTLEVEL] := register_task[FINALLEVEL] + 1;
      plan[LOG] := [“Accept register pressure”, op(plan[LOG])];
      caching_task := FindTask(plan, “Caching”);
      if caching_task[CURRENTLEVEL] > caching_task[FINALLEVEL] then
         granularity_task := FindTask(plan, “Granularity_Control”);
         granularity_task[CURRENTLEVEL] := granularity_task[FINALLEVEL] + 1;
         plan[LOG] := [“No granularity reduction”, op(plan[LOG])];
         new_plans := [plan, op(new_plans)];
      /* Refuse case */
      alternative[CONSTRAINTS] := ['<=(R_B, r), op(alternative[CONSTRAINTS]) ];
      if IsConsistent(alternative) then
         register_task := FindTask(alternative, “Register_Pressure_Control”);
         if (register_task[CURRENTLEVEL] < register_task[FINALLEVEL]) then
            register_task[CURRENTLEVEL] := register_task[CURRENTLEVEL] + 1;
            new_plans := [alternative, op(new_plans)];
         else
            optimized_plans := Optimize(alternative,FindTask(alternative, “CSE”));
            if evalb(nops(optimized_plans) <> 0) then
               new_plans := [op(optimized_plans), op(new_plans)];
            else
               optimized_plans := Optimize(alternative,FindTask(alternative, “Granularity_Control”));
               if evalb(nops(optimized_plans) <> 0) then
                  new_plans := [op(optimized_plans), op(new_plans)];
                  /* No ‘else” case since we tried everything we could */
                  /* to reduce register pressure and we failed! */
               new_plans := [plan, op(new_plans)];
            else
               /* To be continued in Algorithm 9 */
         /* No else case since we tried everything we could */
   /* To be continued in Algorithm 9 */
Algorithm 9: Optimize(plan, task)

**Input:** plan, encoding a program being optimized, and task, an optimization task of plan

**Output:** A list of new plans obtained by optimizing plan according to task

```plaintext
/* continuing Algorithm 8 */
if task[NAME] = "Caching" and task[CURRENTLEVEL] <= task[FINALLEVEL] then
  alternative := Copy_Optimization_Plan(plan);
  z := CacheAmount(plan);
  /* Accept case */
  plan[CONSTRAINTS] := [ '<='(z, Z_B), op(plan[CONSTRAINTS]) ];
  current_vars := { op(plan[RING])[variables] };
  current_vars := (current_vars union indets(z)) minus R_B, Z_B;
  plan[RING] := RegularChains:-PolynomialRing([R_B, Z_B, op(current_vars)]);
  if IsConsistent(plan) then
    plan[LOG] := ["Accept caching", op(plan[LOG])];
    task[CURRENTLEVEL] := task[FINALLEVEL] + 1;
    register_task := FindTask(plan, "Register_Pressure_Control");
    if register_task[CURRENTLEVEL] > register_task[FINALLEVEL] then
      granularity_task := FindTask(plan, "Granularity_Control");
      granularity_task[CURRENTLEVEL] := granularity_task[FINALLEVEL] + 1;
      plan[LOG] := ["No granularity reduction", op(plan[LOG])];
      new_plans := [plan, op(new_plans)];
    /* Refuse case */
    alternative[CONSTRAINTS] := [ '<'(Z_B, z), op(alternative[CONSTRAINTS]) ];
    alternative[RING] := plan[RING];
    if IsConsistent(alternative) then
      optimized_plans := Optimize(alternative, FindTask(alternative, "Granularity_Control");
      if evalb(nops(optimized_plans) <> 0) then
        new_plans := [op(optimized_plans), op(new_plans)];
      else
        optimized_plans := Optimize(alternative, FindTask(alternative, "CSE")];
        if evalb(nops(optimized_plans) <> 0) then
          new_plans := [op(optimized_plans), op(new_plans)];
        else
          task := FindTask(alternative, "Caching");
          task[CURRENTLEVEL] := task[FINALLEVEL] + 1;
          new_plan := AbandonCaching(alternative);
          new_plan[LOG] := ["Refuse caching", op(new_plan[LOG])];
          new_plans := [new_plan, op(new_plans)];
        end if;
      end if;
    end if;
  end if;
else if task[NAME] = "CSE" and task[CURRENTLEVEL] <= task[FINALLEVEL] then
  task[CURRENTLEVEL] := task[CURRENTLEVEL] + 1;
  new_plan := ApplyCSE(plan, task[CURRENTLEVEL]);
  new_plan[LOG] := ["CSE applied", op(new_plan[LOG])];
  new_plans := [new_plan, op(new_plans)];
else if task[NAME] = "Granularity_Control" and task[CURRENTLEVEL] <= task[FINALLEVEL] then
  task[CURRENTLEVEL] := task[FINALLEVEL] + 1;
  new_plan := SetGranularityToOne(plan);
  new_plan[LOG] := ["Granularity set to 1", op(new_plan[LOG])];
  new_plans := [new_plan, op(new_plans)];
end if;
return (new_plans);
```

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