# Software Pipelined Execution of Stream Programs on GPUs

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Abstract—The StreamIt programming model has been proposed to exploit parallelism in streaming applications on general purpose multi-core architectures. This model allows programmers to specify the structure of a program as a set of filters that act upon data, and a set of communication channels between them. The StreamIt graphs describe task, data and pipeline parallelism which can be exploited on modern Graphics Processing Units (GPUs), as they support abundant parallelism in hardware.

In this paper, we describe the challenges in mapping StreamIt to GPUs and propose an efficient technique to software pipeline the execution of stream programs on GPUs. We formulate this problem — both scheduling and assignment of filters to processors — as an efficient Integer Linear Program (ILP), which is then solved using ILP solvers. We also describe a novel buffer layout technique for GPUs which facilitates exploiting the high memory bandwidth available in GPUs. The proposed scheduling utilizes both the scalar units in GPU, to exploit data parallelism, and multiprocessors, to exploit task and pipeline parallelism. Further it takes into consideration the synchronization and bandwidth limitations of GPUs, and yields speedups between 1.87X and 36.83X over a single threaded CPU.

Index Terms—CUDA; GPU Programming; Software Pipelining; Stream Programming

## I. INTRODUCTION

Graphics Processing Units (GPUs) have emerged from being fixed function pipelines to massively parallel Turing complete machines, capable of performing general purpose computation. The latest generation of GPUs, consisting of hundreds of stream processing units are capable of supporting thousands of concurrently executing threads, with zero-cost hardware controlled context switching between threads. For instance the ATI Radeon 4870 has 800 stream processors connected by a 256-bit wide bus to 512 MB of memory, while the GeForce 8800 GTS 512 consists of 128 stream processors [1], organized as 16 multiprocessors, with a similar memory configuration. Each of the multiprocessors in the NVIDIA GPUs can be conceptually viewed as a very wide SIMD processor. While CPUs have traditionally added hierarchies of caches to tolerate memory latency, GPUs address the problem by providing a high bandwidth processor-memory link and supporting high degrees of Simultaneous Multithreading (SMT) within the processing elements, which switches to another set of threads, while the current set of threads is waiting for data from memory. This combination of SIMD and

SMT enables these devices to achieve a peak throughput of 400 GFLOPs [2].

The high performance of GPUs, however, comes at the cost of reduced flexibility and greater programming effort from the user. For instance, in the NVIDIA GPUs, threads executing on different multiprocessors can neither synchronize in an efficient manner, nor can they communicate in a reliable and consistent manner through the device memory, within a kernel invocation [1]. Also, the GPU cannot access the memory of the host system, requiring the CPU to initiate any transfer from the host memory to device memory or vice-versa. Finally, while the memory bus is capable of delivering very high bandwidth, accesses to device memory by threads executing on a multiprocessor, need to be *coalesced* in order to actually achieve the high bandwidth. All these factors translate to a greater programming effort from the user.

ATI and NVIDIA have proposed CTM [3] and CUDA [4] frameworks, respectively, for developing general purpose applications targeting the GPUs. However, both of these frameworks still require the programmer to express the program as data-parallel computations, that can be executed efficiently on the GPU. Also, programming with these frameworks tie the application to the platform. Any change in the platform would require significant rework in porting the applications. Further, these frameworks provide only low-level synchronization primitives, which the programmer would need to adapt to suit the application's requirements. All these frameworks.

On the other hand, the StreamIt programming language, proposed to express streaming applications in a platform independent manner, aims to expose task, data and pipeline parallelism in the application in a natural way [5]. The advantage of this approach is that an optimizing compiler for a target platform can exploit parallelism in the most efficient manner for each target platform, without requiring significant effort from the programmer. A StreamIt program is expressed as a hierarchical composition of simple stream structures, which may then be *flattened* [6] into a set of filters connected by FIFO channels. Apart from providing a natural way to express streaming applications, this frees the programmer from orchestrating the communication between filters. Brook [7] and Accelerator [8] are two prior works which use GPUs for streaming and general purpose computations. However, these

provide a single kernel execution as the base abstraction, with the programmer having to manually handle communication between kernels [7], or require that the programmer express the program using special data structures like the data parallel array [8]. These techniques still require considerable effort by the programmer in order to transform the streaming application to execute efficiently on the GPU. A recent proposal maps StreamIt onto the CellBE platform [9], whose programming model is vastly different from that of GPUs. Figure 1 illustrates the differences between various paradigms.

To efficiently harness the compute and bandwidth resources of the CPU, parallelism must be exploited at various levels:

- 1) The data parallelism across threads executing on a multiprocessor.
- 2) The SMT parallelism among threads on the same multiprocessor needs to be managed to provide optimum performance. Higher levels of SMT do not automatically translate to higher performance, since the number of registers in each multiprocessor is fixed. Exceeding the number of available registers, in a blind attempt to increase the level of SMT parallelism, causes spills into the longer latency device memory, which could degrade performance.
- Parallelism offered by having multiple multiprocessors on a GPU should be used to exploit the task and pipeline level parallelism in the StreamIt program.

Further, accesses to device memory need to be coalesced as far as possible to ensure optimal usage of available bandwidth. Clearly, these challenges have created a gap between being able to express streaming applications naturally and efficiently and targeting the GPUs to execute these streaming applications. Our work attempts to bridge this gap.

This paper makes the following contributions:

- We describe a software pipelining framework for efficiently mapping StreamIt programs onto GPUs.
- We present a buffer mapping scheme for StreamIt programs to make efficient use of the memory bandwidth of the GPU.
- 3) We describe a profile based approach to decide on the optimal number of threads assigned to a multiprocessor, henceforth called *execution configuration* for StreamIt filters.
- We implement our scheme in the StreamIt compiler and demonstrate a 1.87X to 36.83X speedup over a single threaded CPU on a set of streaming applications.

The rest of the paper is organized as follows: Section II provides an overview of the StreamIt language and the organization of the NVIDIA GeForce 8800 series of GPUs. Sections III and IV detail the ILP formulation used for scheduling the stream program across multiprocessors and the code generation for the GPU, respectively. In Section V we report the experimental results. Section VI discusses related work and Section VII provides concluding remarks.



Fig. 1. Differences between various paradigms, with the execution of a saxpy kernel as an example. The saxpy kernel computes the function  $z[i] = s \times x[i] + y[i]$ , where s is a scalar. Each vertical arrow represents one thread of execution. The flow of data is from top to bottom. (a) Execution in a unithreaded CPU, which processes elements in a loop. (b) Execution in a CPU with a 4-wide SIMD unit, 4 elements are processed in one iteration, reducing the trip-count of the loop. (c) Execution in a GPU, where thousands of iterations of the kernel are carried out in parallel, greatly reducing the trip-count of the loop. (d) Approach taken in [9]



Fig. 2. Organization of the NVIDIA GeForce 8800 series of GPUs

#### II. BACKGROUND

## A. Organization of the GPU

Figure 2(a) shows The architecture of the GeForce 8800 series of GPUs [1] which consists of 16 Streaming Multiprocessors (labelled SMn in Figure 2(a)), which share readonly texture and constant caches and a global device memory. Unlike a constant, a texture can be bound to any address in the device memory by the host processor prior to kernel invocation. The program executing on the GPU can access the data referred to by the texture in a cached fashion by issuing *texture fetches*. The architecture of an individual SM consists of 8 Scalar Units (labelled SU0 – SU7 in Figure 2(b)), with a common instruction fetch and thread control mechanism. The shared memory is akin to a software managed cache accessible by all the SUs in an SM. Each SM also has a large partitioned 32-bit register file, with 8192 registers. The



Fig. 3. The StreamIt constructs. (a) Pipeline, (b) Split-Join and (c) Feedback Loop

memory bus connecting the SMs and the device memory is 256 or 384 bits wide (depending on the model).

Within each SM, threads execute in a Single Instruction Multiple Data (SIMD) fashion. The hardware periodically switches between threads to hide the latency of memory accesses. The basic hardware schedulable entity is a *warp*, which consists of 32 threads with consecutive threadids. Threads in a warp execute in lockstep. The number of threads that may be active simultaneously depends on the register requirements of each thread. The CUDA compiler allows the programmer to restrict the number of registers allocated per thread at compile time, generating spill code if necessary. From the programmer's point of view, threads are grouped into thread blocks, each of which consists of a maximum of 512 threads. All threads of a thread block are assigned to exactly one SM by the CUDA runtime. A kernel call dispatched to the GPU through the CUDA runtime consists of exactly one grid, which is a group of thread blocks. Every thread has access to its threadID and the blockID of the thread block it belongs to by means of local variables named threadIdx and blockIdx, respectively, that are passed by the runtime. Further details on the organization of the NVIDIA GPUs and the CUDA programming model can be found in [1].

Efficient usage of the available memory bandwidth requires that simultaneous accesses to the device memory by the threads of a warp be to contiguous addresses, with the first warp addressing the first bank. Formally, thread N of a warp must access an address of the form WarpBaseAddress + N, with  $WarpBaseAddress \equiv 0 \mod Number of Banks$ . Such accesses by all threads can then be *coalesced* into a single access [1].

## B. The StreamIt Language

A StreamIt program consists of filters (also called nodes) connected by communication channels. Hierarchical composition of stream graphs is achieved using *split–join*, *pipeline* and *feedback loop* constructs [6], shown in Figure 3. A splitter can either be a *duplicate* splitter, in which case it copies every element in its input FIFO to each of its outputs, or a *round robbin* splitter, in which case it copies each element from its input FIFO into exactly one output FIFO, based on its weights. A filter may consume one or more tokens (values) from its input FIFO by executing the pop() method, and may produce one or more tokens into its output FIFO by executing the push() method. A filter may also inspect its input FIFO without consuming a token by executing the peek (n) method, where n is the depth into the FIFO at which the value is inspected. These are the only primitives by which filters may manipulate their input and output FIFOs. The tokens in the FIFO buffers can either be objects of primitive types like int, float, etc, or can be objects of user defined types. The push rate and the pop rate of a filter are defined to be the number of tokens produced and consumed, respectively, during each execution of the filter. Also, the peek depth of a filter is defined as the depth up to which a filter can look into the input FIFO during each execution. Note that *peek depth* is always greater than or equal to *pop rate*. A filter is allowed to execute subject to its firing rule, which is simply that it should have at least peek depth tokens in its input FIFO and enough space for at least *push rate* tokens in its output FIFO.

Filters may be *stateful* or *stateless*. Stateful filters are those which have some internal state that is persistent across firings. The state may be updated at each firing as a function of its inputs and the previous values of the state variables [10]. This necessitates a serial ordering of the different instances of stateful filters. On the other hand, *stateless* filters carry no state from one firing to another. Hence different instances of stateless filters can be executed concurrently. We consider only programs with stateless filters in this work.

The push and pop rates of each node in a StreamIt program can be non-unity and non-identical, but are fixed and known at compile time. In order to ensure that such graphs can execute an infinite number of times with finite buffer requirements, the number of firings of different nodes can be different. The firing rate of each node can be computed by solving the steady state rate equations [11]. We call each firing of a particular filter an *instance* of that filter in the schedule. A schedule in which the execution counts (firing rates) of different filters are governed by the steady state equations is called a *steady* state schedule. A schedule in which the execution counts of each of the nodes are mutually prime when taken pairwise is called a primitive steady-state schedule. Two primitive steadystate schedules may differ in the order in which the instances of filters are invoked and hence may have different buffer requirements. Single Appearance Schedules (SAS) [12] [11] require the maximum buffering, while Minimum Latency Schedules require the minimum [13]. A steady state iteration of a stream graph is defined as one execution of the steady state schedule.

## **III. ILP FORMULATION**

In this section, we formulate the scheduling of the stream graph across the multiprocessors (SMs) of the GPU as an Integer Linear Program (ILP). We consider each instance of each filter to be the fundamental schedulable entity.

## A. Definitions

V denotes the set of all nodes (filters) in the stream graph. N is the cardinality of V. E denotes the set of all (directed) edges in the stream graph.  $k_v$  represents the number of firings of a filter  $v \in V$ , as dictated by the steady state rate equations.  $P = \{0, 1, ... P_{max} - 1\}$  is the set of all SMs in the GPU. Trepresents the Initiation Interval (II) of the software pipelined loop. d(v) is the delay or execution time of filter  $v \in V$ .  $I_{uv}$ and  $O_{uv}$  represent the number of elements consumed by filter v on each firing of v, and the number of filters produced by filter u on each firing of u respectively, for an edge  $(u, v) \in E$ .  $m_{uv}$  denotes the number of elements initially present on the edge  $(u, v) \in E$ .

#### B. Resource Constraints

We define 0-1 integer variables  $w_{k,v,p}$ ,  $\forall k \in [0, k_v)$ ,  $\forall v \in V$ ,  $\forall p \in [0, P_{max})$ , such that,  $w_{k,v,p} = 1$  implies that the  $k^{th}$  instance of the filter v has been assigned to SM p. We now model various resource constraints as follows:

$$\sum_{p=0}^{P_{max}-1} w_{k,v,p} = 1, \ \forall k \in [0, k_v), \ \forall v \in V$$
(1)

The above constraint ensures that each instance of each filter is assigned to *exactly one* SM. We model the fact that all the filter instances assigned to an SM can be scheduled in the given II, using the constraint:

$$\sum_{k=0}^{k_v-1} \sum_{v \in V} (w_{k,v,p} \times d(v)) \le T, \ \forall p \in P$$
(2)

## C. Dependence Constraints

Before modeling the dependence constraints, we first need to ensure that the execution of any instance of any filter cannot wrap around into the next II. Consider the linear form of a software pipelined schedule [14] given by:  $\sigma(j,k,v) = T \times j + A_{k,v}$  where  $\sigma(j,k,v)$  represents the time at which the execution of the  $k^{th}$  instance of filter v in the  $j^{th}$  steady state iteration of the stream graph is scheduled and  $A_{k,v}$  are integer variables with  $A_{k,v} \ge 0$ ,  $\forall k \in [0, k_v)$ ,  $\forall v \in V$ . We write each  $A_{k,v}$  as  $A_{k,v} = T \times \lfloor A_{k,v}/T \rfloor + A_{k,v} \mod T$  We define  $f_{k,v} = \lfloor A_{k,v}/T \rfloor$  and  $o_{k,v} = A_{k,v} \mod T$  Thus, the linear form of the schedule is given by:

$$\sigma(j,k,v) = T \times (j+f_{k,v}) + o_{k,v} \tag{3}$$

In the equation (3), the  $o_{k,v}$  indicate the time instant in the software pipelined kernel at which each filter must be scheduled to fire; The  $f_{k,v}$  serve to set up the iteration numbers of the instances of the filters, in the software pipelined schedule.  $(f_{k,v}-f_{k',v'})$  denotes the number of iterations that the instance k of filter v and the instance k' of filter v' are separated by. Note that  $f_{k,v}$  are integer variables greater than or equal to 0. We now constrain the starting time of filters  $o_{k,v}$  as follows:

$$o_{k,v} + d(v) < T, \ \forall v \in V, \ \forall k \in [0, k_v)$$

$$\tag{4}$$

This constraint ensures that *every* firing of a filter is scheduled so that it can complete within the same II, preventing the execution of a filter from wrapping around an II.

We now model the dependence constraints. In order to ensure that the *firing rule* for each instance of each filter is



Fig. 4. (a) A multirate stream graph. Filter A pushes 2 tokens on each firing and filter B pops 3 tokens on each firing. (b) The dependency graph on each instance of each filter, with the edges indicating the number of tokens transmitted with each dependence.

satisfied by a schedule, the *admissibility* of a schedule has been given in [15] as:

$$\begin{split} \sigma(i,v) &\geq \sigma\left(\left\lceil \frac{(i+1)\times I_{uv}-m_{uv}-O_{uv}}{O_{uv}}\right\rceil, u\right) + d(u), \\ \forall (u,v) \in E, \; \forall i \geq 0 \end{split}$$

where  $\sigma(i, v)$  denotes the time of the  $i^{th}$  firing of filter v. This admissibility condition makes an implicit assumption that the firings of the instances of filter v are *serialized*. However this assumption does not hold for the model we are building, where instances of each filter *could* execute out of order, or in parallel across different SMs, as long as the *firing rule* is satisfied. So we must ensure that the schedule is admissible with respect to *all* the predecessors of the  $i^{th}$  firing of the filter v. Figure 4 describes this scenario. Figure 4(a) shows a multirate stream-graph where filter A pushes two elements and filter B pops three elements on each firing. Figure 4(b) shows the dependencies that exist between the various instances of filter A and B. The label represents the filter and the subscript represents the instance number, which can range from 0 to  $k_v - 1$  for a given filter v. To model this behavior for each edge (u, v), we need to analytically determine which instances of a filter u produce data that is consumed by the  $i^{th}$  firing of a filter v. Any firing of a filter v depends on at most  $\left[\frac{I_{uv}}{O_{uv}}\right] + 1$ consecutive firings of filter  $u, \forall (u, v) \in E$ . Intuitively, the  $i^{th}$ firing of a filter v must wait for all the  $I_{uv}$  tokens produced after its  $(i-1)^{th}$  firing. We model this as:

$$\sigma(i,v) \ge \sigma\left(\left\lceil \frac{i \times I_{uv} + l - m_{uv} - O_{uv}}{O_{uv}} \right\rceil, u\right) + d(u),$$
  
$$\forall l \in [1, I_{uv}], \ \forall (u,v) \in E, \ \forall i \ge 0$$
(5)

Although it appears that there are  $I_{uv}$  constraints for each edge, there are in fact *at most*  $\left[\frac{I_{uv}}{O_{uv}}\right] + 1$  constraints, since some constraints are repeated.

We now derive the dependency constraints that in turn determine the constraints on  $f_{k,v}$ . Consider an edge  $(u, v) \in E$ . The  $k^{th}$  instance of filter v, in the  $j^{th}$  steady state iteration is denoted by  $\sigma(j, k, v)$ . Since there are  $k_v$  instances of the filter v in one steady state iteration of the stream graph, using (5), we get:

$$\sigma(j,k,v) \ge \sigma\left(\left\lceil \frac{(j \times k_v + k) \times I_{uv} + l - m_{uv} - O_{uv}}{O_{uv}}\right\rceil, u\right) + d(u),$$
$$\forall l \in [1, I_{uv}], \ \forall j \ge 0, \ \forall k \in [0, k_v), \ \forall (u, v) \in E$$

Since  $I_{uv} \times k_v = O_{uv} \times k_u$  according to the balanced rate equations for multirate stream graphs [15], we can write:

$$\sigma(j,k,v) \ge \sigma\left(\left\lceil (j \times k_u) + \frac{k \times I_{uv} + l - m_{uv} - O_{uv}}{O_{uv}}\right\rceil, u\right) + d(u),$$

$$\forall l \in [1, I_{uv}], \ \forall j \ge 0, \ \forall k \in [0, k_v), \ \forall (u, v) \in E$$

Simplifying further, we get:

$$\sigma(j, k, v) \ge \sigma(j'_l, k'_l, u) + d(u),$$
  
$$\forall l \in [1, I_{uv}], \ \forall j \ge 0, \ \forall k \in [0, k_v), \ \forall (u, v) \in E$$

Where:

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$$\begin{aligned} j'_{l} &= j + j_{lag,l} \\ j_{lag,l} &= j + \left\lfloor \frac{1}{k_{u}} \left\lceil \frac{k \times I_{uv} + l - m_{uv} - O_{uv}}{O_{uv}} \right\rceil \right\rfloor, \; \forall l \in [1, I_{uv}] \\ k'_{l} &= \left\lceil \frac{k \times I_{uv} + l - m_{uv} - O_{uv}}{O_{uv}} \right\rceil \bmod k_{u}, \; \forall l \in [1, I_{uv}] \end{aligned}$$

Note that,  $k \in [0, k_v) \implies j_{lag,l} \le 0, \forall l \in [1, I_{uv}]$ . Now, using the form in (3), we get:

$$\begin{aligned} T\times(j+f_{k,v})+o_{k,v}&\geq T\times(j+j_{lag,l}+f_{k'_{l},u})+o_{k'_{l},u}+d(u),\\ \forall l\in[1,I_{uv}],\;\forall j\geq 0,\;\forall k\in[0,k_{v}),\;\forall (u,v)\in E \end{aligned}$$

Algebraic simplification yields:

$$T \times (f_{k,v} - f_{k'_l,u}) \ge T \times j_{lag,l} + d(u) - (o_{k,v} - o_{k'_l,u}),$$
  
$$\forall l \in [1, I_{uv}], \ \forall k \in [0, k_v), \forall (u, v) \in E$$
(6)

This forms a system of constraints with at most  $\left[\frac{I_{uv}}{O_{uv}}\right] + 1$ constraints for each  $(u, v) \in E$ .

So far, we have assumed that the result of executing a filter u is available d(u) time units after the filter u has started executing. However, the limitations of a GPU imply that this is not the case when the producer and consumer instances are scheduled on different SMs. We define 0 - 1 integer variables  $g_{l,k,u,v}, \forall k \in [0, k_v), \forall (u, v) \in E, \forall l \in [1, I_{uv}]$  as:

$$g_{l,k,u,v} \ge w_{k,v,p} - w_{k'_l,u,p}$$

$$g_{l,k,u,v} \ge w_{k'_l,u,p} - w_{k,v,p}$$

$$\forall k \in [0, k_v), \ \forall (u, v) \in E, \ \forall l \in [1, I_{uv}], \ \forall p \in [0, P_{max}) \ (7)$$

Incorporating the additional constraint into equation (6), and upon algebraic simplifications, we get two systems of constraints:

$$T \times f_{k,v} + o_{k,v} \ge T \times (j_{lag,l} + f_{k'_l,u}) + o_{k'_l,u} + d(u)$$
  

$$T \times f_{k,v} + o_{k,v} \ge T \times (j_{lag,l} + f_{k'_l,u} + g_{l,k,u,v})$$
  

$$\forall (u,v) \in E, \ \forall k \in [0,k_v), \ \forall l \in [1,I_{uv}]$$
(8)

The two constraints are set up such that the former comes into play when  $g_{l,k,u,v} = 0$  and the latter comes into play when  $g_{l,k,u,v} = 1$ , since  $o_{k'_{l,u}} + d(u) \leq T$  at all times (from (4)). The idea is that if *any* predecessor of the instance k of a filter v is scheduled on an SM different from the one where the  $k^{th}$ instance of v is scheduled, then  $g_{l,k,u,v} = 1$  for some l.



Fig. 5. Overview of the compilation process, targeting a StreamIt program onto the GPU

Note that equation (8) is independent of the iteration number j, and forms a system of at most  $\left\lceil \frac{I_{uv}}{O_{uv}} \right\rceil + 1$  constraints for each edge  $(u, v) \in E$ . The inequalities (1), (2), (3), (4), (7) and (8) constitute an ILP formulation for the scheduling of the StreamIt programs on GPUs.

## IV. CODE GENERATION FOR GPUS

In this section, we describe the process of generating a software pipelined executable of a StreamIt program targeted for the GPU. We use the StreamIt compiler, which is a source to source compiler and generates C-like code. This C-like code is then compiled by the native compiler for a specific platform. Figure 5 shows the various phases involved in our method for compiling a StreamIt application for the GPU. The following subsections describe each of these phases in detail.

## A. Profiling and Execution Configuration Selection

As mentioned in Sections I, II and in [2] it is important to determine the optimal execution configuration, specified by the number of threads per block and the number of registers allocated per thread. This is achieved by the profiling each filter independently for different execution configurations. We have modified the StreamIt compiler to generate the CUDA sources along with a driver routine for each filter in the StreamIt program. The generated CUDA code is compiled using the nvcc compiler from NVIDIA, for execution on the GPU. By running multiple profile runs of the same filter, varying the number of threads and the number of registers the filter is compiled for, we identify the optimal number of threads — and hence the optimal level of SMT — for a given filter. The profile runs also enable us to accurately estimate the execution time of each filter on the GPU, for use in the ILP formulation described in Section III.

We generate four versions of the executable for each filter for the profile run, restricting the number of registers to 16, 20, 32 and 64 per thread. These correspond to the register requirements that will allow the kernel to run with 512, 384, 256 and 128 threads respectively. Note that if the per-thread register requirement of a kernel is less than the limit specified, the compilation of the kernel is unaffected. Next, we execute each of these four versions with 128, 256, 384 and 512 threads. For a given execution configuration and the version of the

1: for all $i \in [0, NumFilters -$	· 1	do
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- 2: for all  $numRegs \in \{16, 20, 32, 64\}$  do
- 3: for all  $numThreads \in \{128, 256, 384, 512\}$  do

4:	execute the version of the kernel compiled for
	numRegs registers with numThreads threads for
	numfirings/numThreads iterations.
5:	if kernel fails to execute due to lack of registers then
6:	$runTimes[i][numRegs][numThreads] \leftarrow \infty$
7:	else
8:	$runTimes[i][numRegs][numThreads] \leftarrow t,$

- where, t is the execution time of the kernel.
- 9: end if
- 10: end for
- 11: end for
- 12: end for

#### Fig. 6. Algorithm for Profiling Filters

executable, if the number of registers required per thread is greater than the number of registers the version of the executable was compiled for, then the kernel execution fails and the configuration is infeasible. For all the other feasible configurations we obtain the execution time on the GPU from these runs. To make the comparison fair, we ensure that each execution performs the same amount of work, irrespective of the execution configuration. We accomplish this by changing the number of firings of the filter to be executed in each thread, through a parameter called *numThreads*. This parameter is set to be a multiple of 128, 256, 384 and 512, and is also set to be large enough to amortize the cost of launching the kernel on the GPU over many iterations.

Using the profile data thus obtained, we choose the globally optimum execution configuration for the stream program, as described in Figure 7. The set feasiblePairs used in line 2 of the algorithm consists of all pairs (numRegs, numThreads), such that the configuration is a feasible configuration for all the filters. The CUDA compiler currently does not support extern functions that can be called from a function executing on the device. This restriction imposes that we compile the code for all the filters as a single compilation unit, since calls to these filter work functions will be made by the software pipelined kernel, executing on the device. Thus we need to compile all the filters with the same register usage restrictions. Lines 3 - 6 take into account the numThreads firings of each filter, and recomputes the number of instances of each filter that appears in the steady state schedule. Lines 9 - 13, calculate the candidate initiation interval (II) that can be achieved if the current configuration is chosen. The execution time is scaled in Line 12, to account for the fact that the profile run executes (num firings/k) iterations of the filter, while we require the execution time for only one iteration of the filter. Lines 14 -15 scale the II by the amount of work done. This is necessary, since the work done by the one steady state of the various configurations may not be the same. For example, a particular configuration might yield an II of 20 time units, and produce 200 output tokens, while another might have a higher II of 40, but produce 1000 tokens. Clearly, the latter is a better alternative. The rest of the algorithm, picks the best number of threads for each filter such that the resource constrained II is minimized, and saves the optimal number of threads for

- 1:  $minII = \infty$
- 2: for all  $(numRegs, numThreads) \in feasiblePairs$  do
- 3: for all  $i \in [0, numFilters 1]$  do
- 4: find k < numThreads corresponding to the minimal runTimes[i][numRegs][k]
- 5:  $candidate[i] \leftarrow k$

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6: end for
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- 7: Solve the steady state equations for the execution configuration present in *candidate* and determine the number of instances  $k_v$  for each filter v. Store it in the array *numInstances*.
- 8:  $curII \leftarrow 0$
- 9: **for all**  $i \in [0, NumFilters 1]$  **do** 10:  $bestTime \leftarrow runTimes[i][numRe$ 
  - $bestTime \leftarrow runTimes[i][numRegs][k],$ where, k is chosen as in line 4
- 11:  $bestTime \leftarrow bestTime \times numInstances[i]$
- 12:  $curII \leftarrow curII + bestTime \times (k/numfirings)$

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13: end for
```

- 14: Estimate total work *w* done in one II by the currently chosen execution configuration. (A simple metric for this would be the number of tokens produced at the sink node of the stream graph)
- 15:  $curII \leftarrow curII/w$
- 16: **if** curII < minII **then**
- 17:  $minII \leftarrow curII$
- 18:  $bestThreads \leftarrow numThreads$
- 19:  $bestRegs \leftarrow numRegs$ 
  - for all  $i \in [0, NumFilters 1]$  do
  - $threads[i] \leftarrow k$ , where k is chosen as in line 4
  - $delay[i] \leftarrow runTimes[i][numRegs][threads[i]]$

```
23: end for
```

```
24: end if
```

```
25: end for
```

20: 21:

22:

Fig. 7. Algorithm for Finding the Best Execution Configuration

each filter in an array called *threads*, and the corresponding delays for each filter in the array called *delay*.

## B. Software Pipelining

For the optimal execution configuration selected as described in Section IV-A, the firing rates of the filters in a primitive steady state schedule are different from the corresponding firing rates in the primitive steady state schedule in the original StreamIt program. Each thread of the filter executing on the GPU corresponds to one firing in the primitive steady state schedule of the original StreamIt program. Thus the *push* and *pop* rates of the filter executing on the GPU is the base *push* rate multiplied by the number of threads chosen to execute the filter. These *scaled* push and pop rates correspond to the *unrolled* stream graph, which is identified as the near optimal execution configuration by our heuristic algorithm.

We generate the ILP constraints for software pipelining the *unrolled* stream graph corresponding to the optimal execution configuration. We then use CPLEX, an industrial strength ILP solver, to solve the ILP formulation. Since our ILP formulation is a constraint problem rather than an optimization problem, the time required for solving the ILP formulation is expected to be rather low.

## C. Code Generation

The solution to the ILP obtained is used to orchestrate the execution of the filters across the SMs on the GPU. Although



Fig. 8. A sequential buffer layout, which causes bank conflicts

all the SMs are required to have a single kernel entry point, the individual SMs may diverge with no performance penalty. Once the  $w_{k,v,p}$ , the  $o_{k,v}$ , and the  $f_{k,v}$  have been determined from the ILP solution, we generate a single software pipelined kernel, with the parts required to execute on individual SMs separated using a switch statement. Within each SM, the instances of filters are orchestrated to execute in increasing order of the  $o_{k,v}$  variables corresponding to them. Ties are broken arbitrarily, since it implies that the two instances of filters are not related by a producer-consumer relationship. We use the predicated kernel-only code generation schema described in [16], within each case of the switch statement, with the staging predicate implemented as an array, similar to the scheme described in [9].

## D. Optimizing Buffer Layout

We mentioned in Section II that to achieve the high bandwidths that GPUs are capable of delivering, the accesses to device memory need to be *coalesced*. We propose a buffer layout scheme to ensure that accesses to device memory are coalesced. Figure 8 illustrates the problem, with an example of a filter which has a *pop rate* of 4, executing with 4 threads on a device with memory organized as 8 banks. The buffer is organized sequentially, in the natural FIFO ordering of elements. Thus,  $thread_0$  pops the first four elements,  $thread_1$ , the next four and so on. But the accesses by  $thread_0$  and  $thread_2$ , which occur at time instant  $t_0$ , both hit bank  $B_0$ , causing these accesses to be serialized. Similar collisions occur with respect to  $thread_1$  and  $thread_3$ . Clearly, these collisions occur at every memory access, reducing the utilization of the memory bus and degrading performance. Further, the problem affects both memory reads and writes.

The problem can be alleviated by streaming the entire working set of all the threads into the shared memory in each SM. The shared memory is also organized as banks, and collisions still degrade performance on shared memory, but the degradation is small, since the shared memory has an access latency of 1 cycle [1] as compared to the 400 – 600 cycle latency of the global memory. However, this approach does not scale well for filters with large working sets. For example, a filter that pushes 64 ints and pops the same number of ints at each firing, can then be executed with a maximum of 32 threads, since each SM has only 16KB of shared memory. The low number of threads could now cause performance degradation due to the memory latency that has been exposed with the low levels of SMT. Clearly, the buffer layout scheme needs to be chosen carefully to minimize bank conflicts.



Fig. 9. The optimized buffer layout. The numbers at the top of each position in the buffer show the natural FIFO ordering index that the optimized buffer layout index maps to. We can see that all bank conflicts are avoided, since threads access contiguous addresses. The access pattern of each warp is exactly WarpBaseAddress + tid.

We leverage the fact that the *production* and *consumption* rates on the channels between filters are the same across one entire steady state in a steady state schedule. Consider the stream graph shown in Figure 9. Filter A has a push and pop rate of 2 and filter B has a push and pop rate of 1. Also, filter A and B execute with 256 threads and 128 threads respectively. So there would be 1 instance of A in the steady state schedule and 4 instances of B. We organize the buffers such that the first 128 elements of the buffer contains the first popped elements for each of the 128 threads. This is shown in Figure 9. We group threads into clusters of 128 threads, since this is the gcd of the thread block sizes that we consider. Each thread of each cluster of 128 threads, pops (pushes) the first token it consumes (produces) from contiguous locations in memory. Thus, it is guaranteed that no bank conflicts can occur between accesses of threads in the same block. It is sufficient if the very first input buffer to the stream graph is shuffled, since after that, we can modify the push () and pop () routines to ensure that the data values in the buffers internal to the stream graph are communicated in a consistent manner. Formally, we define the shuffle function in terms of the number of tokens pushed or popped in one entire steady state execution of the stream graph as follows:

$$\begin{aligned} \text{shuff}[i] = & \text{buff}\left[\left\lfloor \frac{i}{128} \right\rfloor + (i \mod 128) \times steady\_pop\_rate\right], \\ & \forall i \in [0, steady\_pop\_rate - 1] \end{aligned}$$

where  $steady\_pop\_rate = o_c \times k_c \times \frac{t_c}{128}$ . Here, c is the first filter of the stream graph in a topological sort ordering.  $o_c$  is the number of elements pushed by *one* thread of filter c.  $k_c$  is the number of steady state firings of c in the primitive schedule.  $t_c$  is the number of threads that c is executed with.

The index j of the  $n^{th}$  element popped by a thread whose thread index is *tid* in an instance of a filter with pop rate o is given by:

$$j = 128 \times n + \left\lfloor \frac{tid}{128} \right\rfloor \times 128 \times o + (tid \text{ mod } 128), \ n < o$$

Similarly the index k of the  $m^{th}$  element pushed by a thread whose thread index is *tid* in an instance of a filter with push rate u is given by:

$$k = 128 \times m + \left\lfloor \frac{tid}{128} \right\rfloor \times 128 \times u + (tid \bmod 128), \ m < u$$

The first term in both equations sets up the offset for the push or pop. The second term serves to organize the pushes or pops into groups of 128, and the final term indicates exactly where in the current block of 128, the element needs to be pushed to or popped from.

With this buffer layout scheme, we totally avoid all bank conflicts and obviate the need to use shared memory. Further, the efficiency of the scheme is oblivious to the push and pop rates of the individual filters.

## V. EXPERIMENTAL EVALUATION

We have implemented the proposed compilation methodology on the StreamIt compiler tool-chain. In this section, we demonstrate the effectiveness of our methodology and compare it with two alternative approaches to solve the problem:

- 1) A software pipelined version that does not coalesce accesses to device memory.
- 2) A serialized execution model, where all filters execute a Single Appearance Schedule (SAS) [12], the execution spanning across all SMs available, with as much data and SMT parallelism as supported by the GPU. However, since a SAS schedule typically has the maximum buffer requirement among all steady state schedules, the buffer requirements of the SAS schedule are constrained to be less than or equal to the buffer requirements of the software pipelined schedule to ensure a fair comparison between the two approaches.

#### A. Experimental Methodology

In order to evaluate our scheme, we use benchmarks distributed along with the StreamIt compiler toolkit version 2.1.1 [17]. The details of each benchmark and a brief description are provided in Table I. We compile each benchmark as described in Figure 5. The nvcc compiler is then used to compile the resulting sources to an executable form. We choose to target 16 blocks to match the 16 SMs available. We only consider programs with stateless filters in this study. Each benchmark was compiled and executed on a machine with dual quad-core Intel Xeon processors, running at 2.83 GHz, with 16 GB of FB-DIMM main memory. The machine runs Linux, with kernel version 2.6.18, and the NVIDIA driver version 173.14.09. We have used a GeForce 8800 GTS 512 GPU with 512 MB of device memory for our experiments. In the results presented later, we do not report execution times, but report only the speedups relative to a single threaded CPU executing the same program. We define  $speedup = \frac{t_{host}}{t_{gpu}}$ , where  $t_{host}$  is the time taken for executing same program on the host CPU mentioned, with a single thread of execution and  $t_{qpu}$  is the time taken for executing the same program on the GPU. The CPU version of the program was compiled using the

Benchmark	Filters	Peeking Filters	Description	
Bitonic	58	0	Bitonic sorting network for sorting 8 integers.	
BitonicRec	61	0	Recursive implementation of the bitonic sorting network.	
DCT	40	0	8x8 Discrete Cosine Transform.	
DES	55	0	Implementation of the DES encryption algorithm.	
FFT	26	0	Fast Fourier Transform	
Filterbank	53	16	Filter bank to perform multirate signal processing.	
FMRadio	67	22	Software FM Radio with equalizer.	
MatrixMult	43	0	Blocked matrix multiply.	

TABLE I Benchmarks Evaluated

uniprocessor backend of the StreamIt compiler suite. The C sources produced by the uniprocessor backend were compiled with gcc using optimization level -03.

The timing information for the GPU version was collected using the event mechanism provided by the CUDA framework, while the timing information for the CPU version was collected using the standard UNIX syscall gettimeofday(). The timing measurements are reported as speedups over the base CPU version, with both versions doing the same amount of work.

**B.** Experimental Results

We evaluate the following aspects of our software pipelining implementation in our experiments:

- 1) The effect of coarsening the granularity of the software pipelined schedule on the execution time.
- Comparison of our methodology with a software pipelining solution that does not coalesce accesses to device memory.
- Comparison with a serialized (one filter at a time, but fully data parallel SAS schedule, which coalesces accesses to device memory.

Figure 10 shows the effect of coarsening the granularity on the execution time. The results for each benchmark are plotted along with the geometric mean as the last bar. In the SWP*n* schedule, each instance of a filter is iterated *n* times to increase the granularity of the GPU kernel. This does not affect the optimality of the schedule, since the delay of each filter is increased by the same proportion, thereby leaving the work distribution still uniform. This has the effect of increasing the amount of work done per kernel invocation, amortizing the cost of launching kernels on the GPU over more iterations. We observe from Figure 10 that the optimized SWP schemes achieve a speedup of 1.87X to 36.83X for the benchmark programs. Further, the gains start to plateau between SWP4 and SWP8 for all benchmarks.

We now compare the performance of our scheme with two other schemes: (i) SWPNC is the same as our implementation, except that memory access coalescing is not done. For this, the profile runs are also executed without memory access coalescing. However, if the number of threads with which the filter is to be executed is such that the working set (the *push* and the *pop* set) can fit into shared memory, then we bring

Bitonic	BitonicRec	DCT	DES
5308416	4472832	29360128	59768832
FFT	Filterbank	FMRadio	MatrixMult
25165824	7471104	1671168	92602368

TABLE II The buffer requirements for each benchmark. All sizes are in bytes.

in the entire working set into shared memory using coalesced reads. Note that the performance gains from this approach are highly dependent on the *push* and *pop* characteristics of the filter. (ii) The Serial scheme is such that every filter is run as a separate kernel in a SAS schedule. We fix the number of blocks with which a filter executes to 16 - same as in the SWP scheme — and set the number of threads so that the buffer usage is less than or equal to that in SWP8. No buffer sharing is performed in all our schemes; in other words, a buffer allocated for a channel is not shared with another [18]. Table II shows the buffer requirements of the optimized software pipelined schedule which has been coarsened 8 times (SWP8) for each benchmark. Although the serial scheme executes filters in a serial fashion, we have implemented this scheme such that device memory accesses are coalesced.

The optimized software pipelining scheme performs better than the alternative schemes in all benchmarks, except for the MatrixMul and the DCT benchmarks, where the serial version performs slightly better. Firstly, these benchmarks have very large register requirements, causing the local variables to be spilled to device memory, increasing the bandwidth requirements. Secondly, these benchmarks display a phased behavior, with each phase having a splitter which exposes a large amount of task parallelism. However very little work is done in the task parallel branches before a joiner that combines the results of the task parallel branches and passes them on to the next phase. These joiners and splitters are bandwidth hungry by nature, since they only move data around, without any computation. Our scheme does not take these second order effects into account. We believe that this skews the actual execution times of the filters, leading to an imbalanced work distribution. The study of such phased behavior in stream programs, is an interesting area of future research, but is beyond the scope of this work.

SWPNC performs poorly in all benchmarks, yielding a maximum speedup of only 1.22X, except in the Filterbank and FMRadio. The poor performance can be explained as being caused by non-coalesced accesses under-utilizing the memory bus. The high speedups of 11.59 and 31.78 in the Filterbank and FMRadio benchmarks are due to the fact that in these benchmarks, the entire *push* and *pop* working set of all the threads fits into the shared memory and written back from shared memory, using a series of coalesced accesses. Bank conflicts do occur in shared memory, however, these are 1-cycle conflicts and do not impose a severe performance penalty, yielding a net performance gain.



Fig. 10. Impact of Coarsening the granularity: *SWP* - Optimized SWP schedule with no coarsening; *SWP* 4 - Coarsened 4 times; *SWP* 8 - Coarsened 8 times; *SWP* 16 - Coarsened 16 times.



Fig. 11. Performance comparison between: *SWP8* - Optimized SWP, coarsened 8 times; *SWPNC* - SWP implementation with No Coalescing; *Serial* - Serial execution of filters using a SAS schedule.

We conclude the discussion on the experimental results with mention of the efficiency of the compilation process. We have used CPLEX version 9.0, running on Linux to solve the ILP. The machine used was a quad processor Intel Xeon at 3.06GHz, with 4GB of RAM. However, for all the solution times reported, CPLEX was running as a single threaded application and hence did not make use of the SMP available. The ILP solution process which is the most complex step of the compilation trajectory, is quite fast for all the benchmarks. The methodology we used to solve the ILP was to determine<sup>1</sup> the lower bound on the II as max(ResMII, RecMII). Once this was done, the solver was allotted 20 seconds to attempt a solution with this II. If it failed to find a solution in 20 seconds, the II was relaxed by 0.5% and the process was repeated until a feasible solution was found. It should also be noted that the compilation time was dominated by the *nvcc* compiler which takes 1 - 2 orders of magnitude more time than the ILP solver. All of the benchmarks took less than 30 seconds to solve, except for Bitonic, BitonicRec and DCT, which took 161, 122 and 178 seconds respectively. All solutions were found within a 5% relaxation on the II, except for FFT and FMRadio, both of which required a 7% relaxation. Thus the proposed solution is quite efficient.

 $<sup>{}^{1}</sup>RecMII$  was 0 for all the benchmarks, since none of the benchmarks in the set of benchmarks provided with the StreamIt distribution had feedback loops and we have not considered stateful filters.

## VI. RELATED WORK

Stream graphs are fairly well studied in literature, with the early works by Lee, et. al. [12] [11], focusing on the Synchronous Data Flow (SDF) model of computation and the Stream Flow Graphs studied by Gao, et. al. in [19]. Govindarajan, et. al. have studied the software pipelining of Regular Stream Flow Graphs (RSFGs) [15], using a linear programming formulation. The framework was extended to reduce the buffer requirements of rate optimal r-periodic schedules in [20].

More recently, the StreamIt project has revived interest in the dataflow graph model of computing [17]. Although StreamIt introduces a *peek* construct that allows filters to inspect data on the input channels without consuming it, StreamIt graphs form a subset of RSFGs, since the *peek* can be implemented as a filter with internal state. Thus StreamIt graphs have the same schedulability properties as RSFGs. While past research has focussed on software pipelining the execution of StreamIt graphs to target the Raw architecture [21] [5] and the Cell BE architecture [9] [22], to the best of our knowledge we are the first to have proposed a framework for compiling and executing StreamIt programs on GPUs.

Recent work on GPUs has focused on the program optimization space pruning [2]. This method reduces the search space in execution configuration selection and optimization space considerably and could be used in place of the profiling methodology that we have suggested. Other work on GPUs have primarily focused on application performance tuning [23].

## VII. CONCLUSIONS

We have described an efficient framework for mapping StreamIt programs to GPUs. Our framework software pipelines the execution of the filters and performs both scheduling and assignment of filters to processors. We also present a novel buffer layout technique for GPUs which facilitates exploiting the high memory bandwidth available in GPUs. The proposed scheduling exploits both the scalar units in GPU, to exploit data parallelism, and multiprocessors, to exploit task and pipeline parallelism. Further it takes into consideration the synchronization and bandwidth limitations of GPUs, yielding speedups between 1.87X and 36.83X over a single threaded CPU.

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