Accelerating the Execution of Matrix Languages on the Cell Broadband Engine Architecture

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November 14, 2009

Abstract

Matrix languages, including MATLAB and Octave, are established standards for applications in science and engineering. They provide interactive programming environments that are easy to use due to their scripting languages with matrix data types. Current implementations of matrix languages do not fully utilise high-performance, special-purpose chip architectures such as the IBM PowerXCell processor (Cell), which is currently used in the fastest computer in the world.

We present a new framework that extends Octave to harness the computational power of the Cell. With this framework the programmer is relieved of the burden of introducing explicit notions of parallelism. Instead the programmer uses a new matrix data-type to execute matrix operations in parallel on the synergistic processing elements (SPEs) of the Cell. We employ lazy evaluation semantics for our new matrix data-type to obtain execution traces of matrix operations. Traces are converted to data dependence graphs; operations in the data dependence graph are lowered (split into sub-matrices), scheduled and executed on the SPEs. Thereby we exploit (1) data parallelism, (2) instruction level parallelism, (3) pipeline parallelism and (4) task parallelism of matrix language programs. We conducted extensive experiments to show the validity of our approach. Our Cell-based implementation achieves speedups of up to a factor of 12 over code run on recent Intel Core2 Quad processors.

1 Introduction

Matrix languages including MATLAB [36] and Octave [17] are established standards for rapid-prototyping in scientific and engineering domains. One of the main reasons for the widespread adoption of these languages is their ease of use. They provide interactive execution of code and simple, high-level syntax for matrix calculations. Complex scientific and engineering problems are solved with a few lines of code because there exists a cornucopia of commercial and open source libraries for standard mathematical problems.

Despite their ease of use, matrix languages traditionally have sequential execution semantics and utilise a single thread of execution only [38]. While the performance growth of single-core processors is reaching its limits [51], scientists and engineers have increasingly large data-sets which must be processed efficiently [21]. Thus, the use of matrix languages will plateau in the near future if not adapted to modern parallel computer architectures.

With the advent of hardware accelerators for high-performance computing such as General Purpose Graphics Processors (GPGPUs) [40] and the Cell Broadband Engine [19], significant performance boosts over single-core architectures are possible. However, harnessing their computational power is challenging in the context of matrix languages. Hardware accelerators for high-performance computing are attributed to having non-uniform memory accesses and complex parallel programming patterns. Extending matrix languages to execute on high-performance, accelerator architectures can be achieved

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by adopting either (1) an explicit parallel programming model which requires users to manually introduce a notion of parallelism in their matrix language program, or (2) an implicit parallel programming model in which parallelism is elicited from a matrix language program with little user intervention. The explicit model contradicts the initial design goals of matrix languages as the users of these languages are most often untrained in concurrent programming. Hence, it is of paramount importance to the continued success of matrix languages that an implicit model is adopted which is capable of fully utilising the computational power of modern accelerator architectures for high-performance computing.

A large body of research \cite{8, 30, 14, 53, 52, 47, 11, 12} already exists on how matrix languages can be parallelised for distributed parallel architectures that were popular before the turn of last century. Some of the parallel extensions developed are reported to offer good performance, however this performance gain was often paid at the expense of ease of use by the programmer \cite{8}. Little research has been conducted in how matrix languages can be parallelised for modern accelerator architectures which present very different challenges in achieving good performance. Current state of the art techniques (e.g. those employed in MATLAB) for parallelisation on multicore CPUs involve using parallel math libraries \cite{55} to exploit data parallelism within matrix operations. Several emerging projects \cite{1, 37} have investigated using simple bindings to execute MATLAB functions on GPGPUs, again exploiting data parallelism in matrix operations. However, these projects offer a naïve approach, neglecting other types of parallelism that exist in matrix language programs and resulting in under-utilisation of their target architectures.

In this paper, we introduce a new framework for the automatic parallelisation of matrix languages which is specifically targeted towards modern hardware accelerators for high-performance computing. We have implemented this framework as an extension to the Octave interpreter running on the Cell Broadband Engine. The Cell Broadband Engine is a heterogeneous multicore architecture, which is currently deployed in the fastest computer in the world \cite{6}, as well as the Sony Playstation 3 computer games console. The Cell consists of a PowerPC core that is connected to several Synergistic Processing Elements (SPEs) via a high-speed interconnect double ring bus. The PowerPC unit is an in-order RISC processor with two hyper-threads, whereas the SPEs are small-sized vector (i.e. SIMD) machines with 256kB of local memory that is shared for data and instructions. Each SPE delivers approximately 25 GFLOPs peak performance for fused multiply-add operations \cite{24}.

GNU Octave is an open source alternative to MATLAB (a commercial product developed by MathWorks). Octave mimics MATLAB’s syntax and has been used in our work because MATLAB does not support PowerPC-based architectures such as Cell Broadband Engine.

Our framework exploits several types of parallelism in an Octave program to obtain high utilisation of the Cell processor:

1. **Data parallelism** is exploited by partitioning large matrices into sub-matrices and executing operations on the sub-matrices in parallel,
2. **Instruction level parallelism** is exploited by executing matrix operations of an execution trace in parallel if there is no data dependence between them,
3. **Pipeline parallelism** is exploited by overlapping communication between cores with computation of matrix operations,
4. **Task parallelism** is exploited by overlapping execution of the Octave interpreter, construction of the schedule, and execution of the matrix operations on the SPEs.

Lazy evaluation is used to generate execution traces of matrix operations by deferring their computation until a result is required. A key feature of our framework is that partitioned matrix operations are scheduled among the parallel execution elements of the Cell processor in a way that satisfies inter-operation data dependencies, prior to execution of the trace. Using estimates of the execution time for each operation, operations can be scheduled such that the utilisation of parallel elements is improved and the total execution time (makespan) is reduced.
We perform an extensive evaluation of our framework using 9 Octave benchmark programs. The execution time of these benchmarks running on our framework with the Cell processor is compared with their execution time on a default installation of Octave on an Intel Core2 Quad processor. Comparisons are also made with several other configurations. Furthermore, we perform experiments to evaluate the benefits of scheduling matrix operations and to determine the extent to which the Cell architecture is utilised.

The contributions of this work are as follows:

1. We introduce a framework for the execution of matrix languages on modern parallel architectures. Our framework exploits both data parallelism and instruction level parallelism of matrix programs. Instruction level parallelism is achieved through lazy evaluation of matrix operations.

2. We provide a ≈7000 line C/C++ implementation of this framework for the Octave programming language on the Cell Broadband Engine architecture.

3. We provide a novel, efficient technique for partitioning matrix operations to maximise the parallelism available within an execution trace.

4. We develop accurate time models for estimating the execution times of matrix operations through multivariate regression analysis.

5. We introduce a new heuristic scheduling for scheduling matrix operations among parallel processing elements. The algorithm takes into account the estimated execution times of operations and the pipelined nature of processing elements.

6. We formulate the scheduling problem as an integer linear program and compare the obtained optimal solution with the solution produced by the heuristic scheduling algorithm.

This paper is organised as follows: In section 2 we survey related work on (1) parallelising matrix languages and (2) the scheduling problem for precedence constrained tasks on multiprocessors. In section 3 we give an overview of our framework and describe each of the major components. In section 4 we describe a motivating example that illustrates how an Octave program is executed in parallel with our framework. In section 5 the lowering process is explained which decomposes operations on large matrices into operations on smaller matrices. In section 6 we describe the scheduling problem and provide a heuristic algorithm as well as an integer linear programming formulation which yields the optimal solution. In section 7 we give an overview of the Cell Broadband Engine architecture and we discuss the implementation of our framework on this architecture. In section 8 we describe details of the development, testing and optimisation of our framework. In section 9 we present the experimental results and discuss the observed performance of our framework. We summarise our work and draw our conclusions in section 10.

2 Related Work

2.1 Parallel MATLAB

There are a variety of extensions for MATLAB designed to utilise parallel computers. The methods of achieving parallelism, the target architecture and the extent to which the parallelisation process is automated vary from extension to extension. In our work we present a new parallel system for a matrix language with two identifying goals:

1. It allows automatic parallelisation of code with no intervention from the programmer.

2. It is designed specifically to achieve high performance on modern hardware accelerator architectures.
Choy and Edelman provide a survey [8] of 27 projects that extend matrix languages with parallel features. The survey classifies the projects in four main categories:

- **Embarrassingly parallel**: These projects make use of multiple MATLAB processes running simultaneously. There is only communication involved when a new process is spawned or a process has completed its task. These parallel extensions for MATLAB are limited to applications that can adopt an embarrassingly parallel programming scheme.

- **Message passing**: These projects provide message passing functionality between MATLAB processes. The complexity of these extensions varies from simple wrappers for MPI functions [49] to higher level abstractions. The programmer has to express the parallelism explicitly.

- **MATLAB compilers**: These projects compile MATLAB scripts into an executable form, either directly or through the use of an intermediate language such as Fortran or C. Some of these projects link their executables with parallel math libraries while others generate code that utilises MPI.

- **Backend support**: These projects use a single MATLAB process as a front-end, which generates jobs that are submitted to a computation engine and executed in parallel, often using numerical libraries like ScaLAPACK.

We now examine the key parallel MATLAB systems in each of these categories. We have chosen to include systems that we believe are representative of the current state of research including several which are more recent than the survey provided by Choy and Edelman.

### 2.1.1 Message Passing and Embarrassingly Parallel Extensions

MatlabMPI [30] is an implementation of the Message Passing Interface (MPI) for MATLAB, developed at MIT. MatlabMPI works by spawning several MATLAB processes that communicate via a shared file system. A sender process writes a variable to a data file on the file system and touches (creates) a lock file when the send is complete. A receiver polls for the existence of the lock file and when it exists, reads in the data file and does any necessary cleanup. The system consists of 350 lines of pure MATLAB code which makes it very portable. There are several similar projects in existence which aim to provide pure message passing functionality in MATLAB including the MPI Toolbox from the University of Granada [14]. They provide flexibility and control in parallelising a program. The performance of MatlabMPI is also reported to approach that of equivalent C MPI code for large messages. However, these systems do not reduce the complexity of concurrent programming for a user and can perform very poorly for certain workloads.

In recognition of the need to reduce this complexity for the typical user, Bliss and Kepner developed pMATLAB [55] which built upon MatlabMPI. Rather than requiring users to perform communication between MATLAB processes explicitly, pMATLAB allows users to declare distributed numerical arrays (or matrices) and an associated mapping of these arrays to available processors in the style of High Performance Fortran (HPF) [29]. A map consists of a grid specifying the partitioning of the array as well as a list of processor IDs that define the processors that will hold the data. Given this data partitioning the program is automatically parallelised. Overloaded MATLAB functions, which take distributed arrays as arguments, automatically perform the required message passing to coordinate computation on these arrays in parallel. Bliss and Kepner report comparable performance to C MPI code for some benchmarks, with a greatly reduced amount of code. A user study they have performed indicates that there is not a steep learning curve for converting a MATLAB application to a pMATLAB application and less than 1% of code requires modification. The HPF programming model limits the applications which can benefit from this type of parallelisation to those which have regular data access patterns. Irregular access patterns on distributed matrices can result in significant slow downs.

The team at MIT build upon pMATLAB again with pMapper [52]. pMapper provides fully automated parallelisation of MATLAB code by generating the array distribution maps for pMATLAB. A
A heuristic approach is taken to produce maps at run-time. One feature of pMapper is that it is designed to be independent of any single parallel architecture or parallel library. Instead, when pMapper is installed on a system it performs an initialisation phase which generates a performance model for the system. This performance model contains timing information for parallel functions with different input sizes. The performance model is then used to generate maps specific to an architecture. pMapper is designed specifically for signal processing applications and the benchmark results are largely due to simulations.

The company MathWorks (the vendor of MATLAB) provides parallel extensions for MATLAB in the form of a commercial Parallel Computing Toolbox \cite{47}. The toolbox has been designed with a heavy focus on an explicit parallel programming model. They achieve this by introducing several extensions to the MATLAB language including parallel loops, distributed arrays and message passing functions. The basic structure of the system uses several worker processes which can communicate with each other and the client MATLAB process. The client formulates computations as a series of jobs which are submitted to a scheduler, executed on workers and return a result. The toolbox allows up to 8 workers to be running locally on a single machine and can be scaled to multiple computers in a cluster with the use of MATLAB Distributed Computing Server. The toolbox draws from several other pieces of software such as MPI libraries and parallel math libraries. It is worth mentioning that MathWorks conducted a survey and found that “reusability of existing MATLAB code was cited as the most important feature of any parallel computing toolset", however they have introduced several new constructs to the language which must be used in order to achieve high levels of parallelism.

With reference to the goals of our framework, they are largely unaddressed by message passing approaches to parallel MATLAB. Firstly, existing systems are targeted toward distributed parallel architectures. Although they can achieve parallelism on some modern accelerator architectures for high-performance computing, communication overheads can restrict even moderate utilisation of the processing elements. In MatlabMPI, for example, inter-process communication is achieved through the file system which is going to be many orders of magnitude slower than specific communication means on modern accelerator hardware. Secondly, as noted most message passing systems require a large amount of intervention from the programmer in order to achieve parallelism.

2.1.2 Compilers

FALCON \cite{11,12} is a programming environment developed at the University of Illinois designed to support the development of optimised numerical applications and libraries. The input language is MATLAB which is translated to the target language, Fortran 90, in three stages. The first stage, program analysis, constructs an Abstract Syntax Tree (AST) and determines the type and shape of variables. As with any untyped language, in order to compile the code, the types of variables must first be inferred. This can not always be done statically, so FALCON utilises both static and dynamic analysis, combined with user input to determine variable types. The next stage uses a collection of transformation rules to restructure the code in order to perform optimisations. This phase is fully interactive with optimisations suggested by FALCON and selected by the programmer. Finally, the code generation stage uses information collected during the analysis phase to produce Fortran 90 code. This code is annotated with compiler directives which allow automatic parallelisation by Polaris \cite{7}, a parallelising Fortran compiler. Despite achieving speedups of up to 1000 times over the MATLAB interpreter the work presented in FALCON focuses mainly on producing high-performance sequential Fortran code. Parallelisation of this code is left largely uninvestigated and as future work with no results reported for benchmarks on multiprocessor machines. Unfortunately this project is now seemingly dormant.

Work from FALCON continued with a MATLAB Just-In-Time (JIT) compiler called MsJiC \cite{4}. In JIT compilation no static analysis of the code is done. Instead, portions of the code are compiled at run time in order to achieve better performance than that of purely interpreted code. Although this project did not explore program parallelisation in the compilation process it remains interesting since, as far as the authors are aware, it is the only research project that uses JIT compilation for
MATLAB. JIT compilation is desirable for two reasons. Firstly, it allows MATLAB to remain an interpreted (and untyped) language which is important in facilitating rapid prototyping. Secondly, it means that optimisations can be applied at run time that may not be apparent at compile time. In our work we adopt JIT compilation techniques to efficiently schedule matrix operations on the PowerXCell architecture.

Otter \cite{44, 45} is another MATLAB compiler developed at Oregon State University. Unlike FALCON which translates MATLAB scripts first to sequential Fortran code and then parallelises that code, Otter translates MATLAB code directly to parallelised C code with calls to MPI and existing parallel numerical libraries. The translation process used in Otter to generate C code is based on the translation process developed for FALCON. However, extra compilation passes are used to introduce parallelism. Calls to the Otter run-time library are inserted to provide a means of communication between processors or utilise existing numerical libraries, such as ScaLAPACK when possible. Otter is advantageous as it provides completely automatic parallelisation of general code and is portable between all architectures supporting MPI. A limitation, as with pMATLAB, is that it again achieves parallelism through distributed arrays which require regular data access patterns to achieve good performance. Also, it is bound to the usage of numerical libraries and thus the matrix distributions associated with those libraries. While these compilers provide some automated parallelisation of code, they are once again focused toward parallelism on distributed architectures.

MATCH \cite{5} is another MATLAB compiler however, unlike FALCON and Otter, it is targeted toward heterogeneous parallel architectures. These architectures consist of an interconnection of various processing components such as embedded processors, digital signal processors and commercial off-the-shelf components, each of which can perform certain types of computation very efficiently. Despite being developed nearly a decade ago, MATCH is relevant in the context of our research because the heterogeneous architectures presented resemble modern parallel architectures in some ways. In particular these heterogeneous architectures use a single general purpose processor (the MicroSPARC-II in this case) to handle the control flow of the program and specialised units perform computation-intensive portions of the program more efficiently. This can be compared to the Cell processor which has a PowerPC processor to handle control flow and several Synergistic Processing Elements (SPEs) optimised for computation. MATCH works by first producing an abstract syntax tree, as is done in FALCON and Otter. The AST is then partitioned into sub-trees whose nodes correspond to library functions in MATLAB. Each sub-tree is mapped to a given processing resource depending on how efficiently that portion of the program can be executed on the resource. This mapping can be produced automatically by MATCH using timing information and a mixed integer linear programming approach, however user guided mappings are also possible. Code is then generated for each partition of the AST, depending on the architecture to which the partition is mapped. Built-in MATLAB functions, such as matrix multiplies, are pre-compiled in architecture specific libraries. Distributed arrays are again used to generate parallel code on target architectures for user defined functions.

2.1.3 Backend Support

Our framework falls into the category of backend support as it has a computation engine that receives matrix operations from the Octave interpreter, executes the operations and returns the result to the client.

Jacket \cite{1} is a MATLAB backend that runs on General Purpose Graphic Processing Units (GPGPUs). It is a commercial product developed by Accelereyes and few details are available about its design. It is one of several systems that have emerged recently for the acceleration of MATLAB code on GPGPUs \cite{37, 56}. In Jacket, the programmer casts matrices into GPU matrices, which are transferred to GPU memory. Operations on GPU matrices are executed on the GPGPU by compiling code on-the-fly with the NVIDIA/CUDA \cite{40} infrastructure. Jacket also provides some syntax extensions for executing for-loops with parallel semantics. Though Jacket provides a high level of abstraction from the details of parallelism, the programmer still requires knowledge of the underlying GPGPU architecture to write efficient Jacket/MATLAB code as they must understand the overheads involved.
in transferring data between main memory and the device memory of the GPGPU. Furthermore, it appears that Jacket essentially provides MATLAB bindings to CUDA functions and as such, instruction level parallelism is not exploited by the system.

Star-P [27] is another commercial product which began as a research project at MIT in 1998 and was commercialised in 2004. It involved the same authors who conducted the parallel MATLAB survey discussed previously and incorporates ideas from several existing parallel MATLAB systems. From a user’s perspective, Star-P behaves in a similar way as Jacket. Programmers denote a distributed variable by tagging it with the characters \(*\_p\), converting it to a custom data type. Star-P then provides overloaded versions of standard MATLAB functions which operate on the custom data type in parallel. Star-P targets a distributed computing environment. The main purpose of the MATLAB client is to distribute tasks to a cluster running the Star-P server, which will run computations and deliver results. The server software is a general purpose computation engine which is not specific to MATLAB and supports clients running many different languages, including Python. It utilises existing math libraries to perform parallel computations on each server.

2.2 Task Scheduling

In our work we attempt to improve utilisation of modern accelerator architectures by scheduling matrix operations among parallel execution elements. We wish to schedule operations such that (1) the data dependencies between operations are satisfied, i.e. an operation can only begin execution after its operands have been computed, and (2) the total time of execution (makespan) is minimised. This problem is well explored in literature [34], with two main formulations — the delay model [22] and the malleable tasks model [54].

2.2.1 The Delay Model

In the delay model we are given a precedence graph \(G(V, E)\) as the input to the problem along with a number of available processors \(p\). \(G\) is a directed acyclic graph (DAG) whose vertices \(V\) represent the tasks and directed edges \(E\) represent the dependencies between tasks. There is an arc from a node \(s \in V\) to a node \(t \in V\) if task \(t\) depends on task \(s\). Vertices are annotated with the cost of executing the task. An edge from \(s\) to \(t\) is annotated with the cost of communication between the tasks \(s\) and \(t\).

The goal is to find a legal schedule that minimises the makespan. A schedule consists of a start time for each task and the processor to which it is assigned. Note that a scheduled task must be executed completely on the processor it has been assigned without interruption.

Our problem of scheduling matrix operations on SPEs resembles this problem. Matrix operations become the vertices in the precedence graph and the data dependencies between operations form the edges. The costs of operations can be estimated by using a time model that is obtained by profiling and regression analysis.

It has been proven that finding an optimal solution to the scheduling problem with the delay model is NP-complete [16]. As such, work has been done on finding optimal, polynomial time solutions for simplified versions of the problem. Hu [22] addresses a variation of the problem in which the task precedence graph is assumed to have a tree structure, tasks are assumed to have unit execution costs and communication costs are ignored. A simple, list scheduling algorithm is proposed to find the optimal solution in polynomial time. List scheduling algorithms are a common approach to the scheduling problem that order tasks in a list according to some heuristic. The heuristic typically ensures a topological ordering of the task precedence graph. Tasks are then iteratively removed from the list in order and assigned to the processor allowing the earliest start time. Hu uses a heuristic in which tasks are ordered by their distance from the root of the tree in the task precedence graph.

Coffman and Graham [9] also propose a polynomial time, optimal algorithm with the simplifying assumptions that there are only 2 processors, unit execution costs and no communication costs. Papadimitriou and Yannakakis [42] propose a polynomial time, optimal algorithm with the simplifying assumption that there are unit execution costs and the precedence graph is interval ordered. In
our work we must schedule operations that have arbitrary precedence constraints over more than 2 processors. Hence, the above mentioned algorithms cannot be used.

Recognising the difficulty of finding an optimal solution for the general problem, many heuristic algorithms [34] have arisen which are usually variations of the list scheduling approach. Adam et al. [2] performed a simulated analysis of these algorithms and found that a Highest Level First with Estimated Times (HLFET) heuristic gave the best results for the chosen benchmarks. In this approach, tasks are ordered by the cost of the longest path to an exit task (a sink node) in the precedence graph. This value is known as the \( b \)-level of the task. Tasks with a larger \( b \)-level are scheduled first. Using this approach ensures that tasks along the critical path of the graph will be scheduled first. The critical path is the longest path through the precedence graph from an entry node (source) to an exit node (sink) and is important as it represents the minimum length of an optimal schedule. It was shown that this approach can produce near-optimal schedules and has a run-time of \( O(n^2) \) for \( n \) tasks. Graham [18] showed that the schedule length, \( SL \), generated using level-based, list-scheduling algorithms such as HLFET was no more than twice the length of the optimal schedule, \( SL_{opt} \), such that \( SL \leq (2 - \frac{1}{p})SL_{opt} \) where \( p \) is the number of processors.

When communication costs are allowed to be arbitrary, several other algorithms have been proposed. A well known algorithm in this category is the Insertion Scheduling Heuristic (ISH) [33]. It works in a similar way to HLFET by using the \( b \)-level of a task as its priority in a list scheduling algorithm. However, while HLFET may leave gaps in the schedule, ISH seeks to fill these gaps and thus reduce the schedule length. Every time a gap is introduced in the schedule, ISH examines unscheduled tasks which are ready to be scheduled and attempts to use them to legally fill the gap. The time complexity of ISH is \( O(n^2) \).

Another category of algorithms are proposed for scheduling tasks with arbitrary precedence constraints which allow task duplication. These algorithms recognise that by redundantly duplicating tasks, the time waiting for a parent task to complete might be reduced which may reduce the overall time of the schedule. The Duplication Scheduling Heuristic (DSH) [32] is a representation of such algorithms. It works by iteratively attempting to duplicate a tasks ancestors on a processor if it allows the task to be scheduled at an earlier start time. The time complexity of the algorithm is \( O(n^4) \) which may be unsatisfactory for applications with a limited time budget for scheduling.

More recently, polynomial time approximation algorithms have been proposed for the general case of the delay model scheduling problem, with communication costs considered. These yield a solution of bounded quality. One such algorithm [39] gives an approximation guarantee of \( 2.33 - 1.33m \) where \( m \) is the number of edges in the task precedence graph. The algorithm is based on rounding of a relaxed linear programming solution to find a schedule with minimum makespan on an unbounded number of processors. A list scheduling algorithm is then used to produce an optimal schedule for a finite number of processors. Due to the use of linear programming, these algorithms can again be impractical for time-critical applications.

2.2.2 The Malleable Tasks Model

The malleable tasks model is a more recent model than the delay model and as such, there is less work on it. It is similar to the delay model but with two main differences. Firstly, communication costs are incorporated in the execution cost of each task. Secondly, tasks in this model do not have to execute on a single processor but can be divided up and executed over several processors with a reduced execution time. As such vertices in the precedence graph are not merely labelled with a single execution cost but instead a cost function, \( c(k) \), which is dependent on the number of processors, \( k \), which the task is assigned to. A schedule that satisfies the problem consists of a start time for each task and a number of processors over which each task is to execute.

We could apply the malleable task scheduling problem to our framework because the tasks in our framework are matrix operations. These can be divided into operations on sub-matrices which can be executed across several processing elements.

It has been proven that the scheduling problem for malleable tasks with arbitrary precedence
constraints is NP-hard with only 3 processors [13]. Even the problem of finding an optimal schedule for malleable tasks without precedence constraints has been shown to be NP-hard with 5 or more processors [13].

Despite the hardness of the problem there has been some recent work on approximation algorithms. Lepre et al. [35] present a polynomial time approximation algorithm for scheduling malleable tasks under precedence constraints with an approximation ratio of $3 + \sqrt{5} \approx 5.23006$. Their algorithm is developed by identifying the relationship between the scheduling problem and the allotment problem, which is approximated by rounding the solutions of a linear programming relaxation [48]. This is combined with a list scheduling algorithm to provide the final approximation algorithm. Jansen and Zhang [28] improve on the work of Lepre et al., with an approximation ratio of $100/43 + 100(\sqrt{4349} - 7)/2451 \approx 4.730598$. Existing approximation algorithms for the malleable tasks scheduling problem may not be viable in time-critical situations. This is due to their use of linear programming which has polynomial time complexity, but high overheads in practice.

3 System Overview

Our framework is a system extension for Octave. It takes the form of a shared library that is loaded by the Octave interpreter at runtime. The framework automatically parallelises matrix instructions for the Cell Broadband Engine and only requires minimal changes to existing Octave code when deployed. These changes consist of casting all matrix declarations to a new Octave data type called \texttt{p-matrix}. Standard operators, such as $+$, $-$ and $\ast$, as well as built-in functions, such as \texttt{sin} and \texttt{round}, have been overloaded to operate on the \texttt{p-matrix} data type (see section 8).

The underlying idea of our framework is to execute several matrix instructions at the same time to optimally harness the computational power of the Cell Broadband Engine. However, the sequential execution semantics of matrix languages do not provide the notion of concurrent execution of matrix instructions, besides dividing matrix instructions into sub-operations which are distributed among parallel processing elements. To further increase the parallelism in Octave programs, we employ lazy evaluation of matrix instructions. Lazy evaluation delays the execution of matrix instructions until the result of an instruction is required. This concept is heavily used in functional programming languages and has numerous applications there, including avoiding unnecessary computations and error conditions, being able to operate on infinite data structures, and defining control flow structures in the language itself [23].

Our framework uses lazy evaluation to collect a trace of matrix instructions. The overloaded functions of the new data type facilitate the construction of the trace which is then analysed to determine the data dependencies between operations. The data dependencies in the trace loosen the strict sequential ordering of instructions to a partial ordering that allows independent matrix instructions to be executed in parallel. A data dependence graph $G(I, E)$ is constructed for the trace where $I$ is the set of nodes in the graph representing instructions in the trace, and $E$ is the set of data dependencies between pairs of matrix instructions. For example, the lazily evaluated statement $A = B \ast C$ imposes two directed edges $(B, A)$ and $(C, A)$ because the result $A$ of the matrix multiplication depends on the matrix operands $B$ and $C$, as shown in Figure 1. The source nodes in this graph (i.e. in-degree of 0) are typically constant or computed matrices, whose value is already available.

Our framework constructs the data dependence graph on the fly when matrix instructions are lazily evaluated. Note that the constructed graph is acyclic even for loops. A matrix instruction that is executed multiple times inside a loop is represented by a set of nodes in the graph. For each execution instance of the matrix instruction there exists exactly one node in the graph.

A trace will continue to grow in length as the program is executed until either (1) a statement is reached that requires the result of an unexecuted matrix operation in the trace and cannot be lazily evaluated, e.g., displaying the value of a matrix, or (2) the length of the trace has reached a certain threshold, i.e., it becomes opportunistic to execute the matrix operations in parallel. If either of these criteria are met, execution of the data dependence graph is triggered.
Figure 1: A data dependence graph for the statement $A = B \ast C$. $A$ depends on the values of matrices $B$ and $C$ so there is a directed edge in the graph from $B$ to $A$ and from $C$ to $A$.

Figure 2: Pipelined execution of matrix instructions. There are 3 stages in the pipeline: data fetch (df), execution (ex) and write back (wb), each of which can be overlapped.

The first step in execution of the data dependence graph is lowering the graph. The memory of the SPEs on the Cell architecture is at a premium, i.e., code and data share the same memory which is limited to 256kB. To be able to compute larger matrices, the framework decomposes matrix instructions into matrix instructions that operate on sub-matrices. This decomposition of the instructions not only enables the execution of matrix instructions on the parallel processing elements of the Cell but also exposes data parallelism in the matrix instructions. We refer to this process of decomposing the instructions of the data dependence graph into instructions that operate on sub-matrices as lowering (see section 5). The lowering process rewrites the original data dependence graph into a lowered data dependence graph, which has an increased number of operations and dependencies.

The lowered data dependence graph is then scheduled among the parallel processing elements in the underlying architecture (see section 6). Scheduling assigns each parallel processing element a subset of the lowered operations which have a specified order in which they are to be executed. The scheduling is performed in a way that satisfies the data dependencies between operations and minimises the total execution time (makespan) of the trace. Execution times of matrix operations are estimated using time-models constructed from profiling data. This ensures that an accurate schedule is produced. Since the scheduling of operations happens at run-time it is also important that a schedule is produced quickly.

In the final step, the lowered matrix instructions are executed on the parallel processing elements according to the schedule. This component of the framework is referred to as the computation engine. The computation engine is abstracted from the details of the underlying architecture and instead viewed only as an asynchronous, pipelined, Multiple-Instruction/Multiple-Data (MIMD) architecture with a shared memory. The architecture executes matrix instructions concurrently in an asynchronous fashion. To hide the communication between memory and the processing element, the architecture utilises a pipeline. The pipeline stages of a single matrix instruction are assumed to be timely interleaved as depicted in Figure 2, which is an idealised scenario assuming that the durations of the stages have the same duration and there are no “bubbles” or gaps in the pipeline. We employ the following pipeline stages in our computation engine:

1. **Data Fetch (df):** The operands of the matrix instruction are loaded from main memory into the memory of the parallel processing element,

2. **Execute (ex):** The matrix instruction is executed on the parallel processing element,
Figure 3: The main software components of the framework. These are: (1) the p_matrix data type extension to the Octave interpreter, (2) the lowerer for partitioning matrix operations, (3) the scheduler for scheduling matrix operations among the parallel execution elements of the architecture, and (4) the computation engine which executes the partitioned matrix operations.

3. **Write Back (wb):** the result of the matrix instruction is written back to main memory.

In contrast to a super-pipelined, super-scalar CPU [43], the matrix instructions are not assumed to be synchronised, i.e., there is no global clock that triggers a new step with a constant period.

This abstraction from the details of the underlying parallel architecture allows the framework to be easily ported to many different architectures (such as GPGPUs or multi-core CPUs) by customising the computation engine. In this work, we implement a computation engine for the Cell Broadband Engine architecture (see section 7). Each of the SPEs in the Cell processor acts as a processing element and executes a sequence of matrix instructions. We call the program that runs on the SPEs a *Matrix Execution Unit* (MEU). The MEUs need to be synchronised globally. An execution control mechanism guarantees that a matrix instruction on an MEU is only executed if its operands are already available in main memory. The execution control is run on the PowerPC Processing Element (PPE) of the Cell.

After the completion of the execution of lowered operations, the results are made available for use by the Octave interpreter.

The software components of our framework are depicted in Figure 3. The first component is a data type extension to the Octave interpreter called p_matrix. The operators for the new data type are overloaded to perform lazy evaluation and to obtain the execution trace. From this, the data dependence graph of operations in the trace is computed, on the fly. The data dependence graph is passed on to the lowerer which decomposes matrix instructions into instructions that operate on sub-matrices. The scheduler computes the schedule for the computation engine. Finally, the matrix instructions are executed on the computation engine according to the schedule. All four components can be executed in parallel, i.e., the Octave interpreter, the Lowerer, the Scheduler and the computation engine are executed in separate execution threads allowing overlapped execution of all four components.

4 Motivating Example

Assume we want to compute the value of $B = A^i$ using an Octave script, where $A$ is a square matrix of dimensions $n \times n$ and $i$ is a positive integer. For the purpose of this example we let $A$ be a random
A = rand(100);
i = 3;
B = A;
for k = 1:i-1
    B = B * A;
end
disp(B);

A = p_matrix(rand(100));
i = 3;
B = A;
for k = 1:i-1
    B = B * A;
end
disp(B);

Figure 4: Octave script for the computation of $B = A^i$.

Figure 5: Parallel data type modifications for $B = A^i$.

100 \times 100$ matrix and $i = 3$. A naïve Octave implementation for calculating matrix $B$ and displaying its value to the screen is given in Figure 4.

To use our framework, the Octave programmer converts matrix declarations to the custom $p$-matrix Octave data type by wrapping them with the $p$-matrix function. In our example program, the only matrix declaration is for the matrix $A$ in line 1 of Figure 4. The modified code that uses our framework is shown in Figure 5. Note that a user does not have to decide which matrices should be converted to this new data type (all matrices can be safely converted), and the need for this additional data type could be completely eliminated with alterations to the Octave interpreter.

When the script in Figure 5 is executed in the Octave interpreter, matrix operations involving operands of the type $p$-matrix are lazily evaluated by our framework. In the example program, the only such matrix operation is the multiplication $B = B \ast A$, in line 6 of Figure 5. The result of this operation (the matrix $B$) is never required inside the for-loop. Hence, rather than eagerly executing this operation each time it is reached inside the loop (as would be the case when executing in a default installation of Octave), lazy evaluation defers execution of the operation. Instead, the operation is recorded in an execution trace (shown in Figure 6). The trace is kept as an internal data structure and a data dependence graph of the trace is constructed on the fly. The data dependence graph shows which operations depend on the results of other operations and determines a partial order in which operations must be executed to yield a correct result. This partial order enables the parallel execution of matrix operations in contrast to strict sequential execution of matrix languages.

Let $B_k$ denote the value of $B$ in the $k$'th iteration of the for-loop. Execution of the example program from Figure 5, construction of the trace (Figure 6) and construction of the data dependence graph (Figure 7) proceeds as follows:

1. Matrix $A$ in line 1 is declared. The conversion of the random matrix to type $p$-matrix causes our framework to add the matrix to the data dependence graph as a constant matrix (Figure 7(a)). Note that constant matrices (like $A$) are not true matrix operations as their result is already available and they do not need to be executed. Instead they are added to the data dependence graph to denote the dependence of another operation on that matrix.

2. Matrix $A$ is assigned to variable $B_0$ in line 4. Again this causes a constant matrix to be added
to the data dependence graph (Figure 7(b)). Note that a deep copy of matrix $A$ is not made upon assignment to $B_0$. For the sake of simplicity, we represent them separately in the data dependence graph.

3. In the first iteration of the for-loop, $B_0$ is multiplied by $A$ and the result stored in $B_1$. The result of this operation is not yet required in the program. Hence, execution is deferred and a multiplication operation is added to the trace and data dependence graph (Figure 7(c)). The operation depends on 2 values — matrix $A$ and matrix $B_0$. Arcs $(A, B_1)$ and $(B_0, B_1)$ denote these dependencies in the dependence graph.

4. In the second iteration of the for-loop, $B_1$ is multiplied by $A$ and the result stored in $B_2$. Again, the result of the multiplication is not required immediately, so another multiplication operation is added to the trace and data dependence graph (Figure 7(d)). The operation depends on 2 values — matrix $A$ and the result of the previous matrix multiplication operation, $B_1$. Arcs $(A, B_2)$ and $(B_1, B_2)$ denote these dependencies in the dependence graph.

5. The terminating condition of the for-loop is reached.

The final statement of the program to be executed is given in line 9 of Figure 5. It requests that the value of $B_2$ be printed to the screen. However, the value of $B_2$ has not yet been computed due to lazy evaluation. This causes execution of the Octave program to be halted while the data dependence graph from Figure 7(d) is executed to obtain the required result. Note that in this example, execution of the trace was forced by a required value. The other cause of executing a trace is that the length of the trace becomes too large. If that is the case then execution of matrix operations and execution of the Octave interpreter are performed concurrently.

The first step in execution of the data dependence graph is lowering the graph (see section 5). Lowering partitions matrix operations on large matrices into operations on sub-matrices. This is necessary because some matrices may be too large to fit into the memory of parallel execution elements (e.g., with the Cell architecture), but it has the beneficial side-effect of exposing data parallelism in matrix operations. Assume that the parallel elements have enough memory to store operands of dimensions $50 \times 50$, however, the matrices in the example are of dimensions $100 \times 100$. We can partition
the 100 × 100 matrices into four 50 × 50 blocks, and use block matrix multiplication to perform the multiplication operations in our program. Block matrix multiplications works in the same way as regular matrix multiplication, except that instead of multiplying and adding the scalar elements of the two operands, we multiply and add the partitioned sub-matrices of the operands. Figure 8 shows how this partitioning occurs for the first multiplication operation, $B_0A$.

Note that a single matrix operation in our original dependence graph will result in many lowered operations after partitioning. These operations form a new data dependence graph, called the lowered data dependence graph. The lowered data dependence graph for our example is shown in Figure 9.

Once lowering is complete, operations are scheduled among the execution elements of the underlying architecture (see section 6). The aim of scheduling is to assign operations to processors in such a way that minimises the total execution time (makespan) of the operations but also ensures that the data dependencies between operations are enforced: an operation cannot execute before the results of all its operands are available.

Without specific reference to the details of the underlying architecture, it is viewed as a pipelined architecture in which each processor can overlap computation of a matrix operation with data transfers to and from main memory. There are three pipeline stages that are considered — data fetch (df), execution (ex) and write back (wb). The execution stage of an operation can only begin after the completion of the data fetch of that operation and the write back stage can only begin after the completion of the execution stage. To produce an effective schedule, it is necessary to estimate the time that each of these stages will take for each operation that is to be executed.

A heuristic algorithm uses these execution time estimates to schedule the operations among the execution elements. The heuristic works by selecting the unscheduled operation whose operands have been scheduled to complete at the earliest point in time and assigning that operation in the earliest available slot on a processor. A partial schedule of lowered operations from the example is given in Figure 10. Note that the constant matrices (red nodes) are not scheduled as they do not need to be executed (their result is already available). In this case, all of the sub-operations for operation $B_1$ have been scheduled already. By examining the lowered data dependence graph from Figure 9, it can be seen that the next operations that are ready to be scheduled (i.e. whose operands have been scheduled already) are operations: $(B_2)_{1,1a}, (B_2)_{1,1b}, (B_2)_{1,2a}, (B_2)_{1,2b}, (B_2)_{2,1a}, (B_2)_{2,1b}, (B_2)_{2,2a}$ and $(B_2)_{2,2b}$. Each of these operations depends on one of the constant sub-matrices in $A$: $A_{1,1}, A_{1,2}, A_{2,1}, A_{2,2}$; as well as one of the addition sub-operations in $B_1$: $(B_1)_{1,1}, (B_1)_{1,2}, (B_1)_{2,1}, (B_1)_{2,2}$. The addition operation from $B_1$ that has been scheduled to complete at the earliest point in time is $(B_1)_{1,2}$ and we can consider all of the constant matrix operations from $A$ to have been scheduled at time 0. Hence, our heuristic algorithm will choose one of the operations $(B_2)_{1,2a}$ or $(B_2)_{2,2a}$ (which depend on $(B_1)_{1,2}$) to be scheduled next. These operations will complete earliest if scheduled on Processor 2 (as shown in Figure 11) so the heuristic would schedule the operation there. This process continues until all operations have been scheduled.

Once the schedule is produced, the operations are executed on the underlying architecture accordingly. In the implementation of the framework for the Cell Broadband Engine (see section 7), each of the SPEs acts as a processing element which is capable of executing lowered matrix operations. Each SPE runs a specially designed, small virtual machine program called a Matrix Execution Unit which is optimised for computing matrix operations. An execution control mechanism runs on the PPE of the Cell processor. It delivers lowered operations to the MEUs according to the schedule. It also ensures that an operation does not begin execution until all of its operands have finished being computed.

When all operations have completed execution, execution of the Octave program resumes and our example program completes.
Figure 8: Partitioning (lowering) for multiplication operation $B_0A$ from Figure 6. Block-partitioning divides each $100 \times 100$ operand into four $50 \times 50$ sub-matrices. Block matrix multiplication is used to multiply sub-matrices. Each block of the result matrix is computed using two multiplications and one add operation.

\[
B_0A = \begin{bmatrix}
(b_0)_{1,1} & \cdots & (b_0)_{1,50} & (b_0)_{1,51} & \cdots & (b_0)_{1,100} \\
\vdots & & \vdots & & \vdots & \\
(b_0)_{50,1} & \cdots & (b_0)_{50,100} \\
(b_0)_{51,1} & \cdots & (b_0)_{51,100} \\
\vdots & & \vdots & & \vdots & \\
(b_0)_{100,1} & \cdots & (b_0)_{100,50} & (b_0)_{100,51} & \cdots & (b_0)_{100,100}
\end{bmatrix}
\]

\[
= \begin{bmatrix}
(B_0)_{1,1} & (B_0)_{1,2} \\
(B_0)_{2,1} & (B_0)_{2,2}
\end{bmatrix}
\begin{bmatrix}
A_{1,1} & A_{1,2} \\
A_{2,1} & A_{2,2}
\end{bmatrix}
= \begin{bmatrix}
(B_0)_{1,1}A_{1,1} + (B_0)_{1,2}A_{2,1} \\
(B_0)_{2,1}A_{1,1} + (B_0)_{2,2}A_{2,1}
\end{bmatrix}
\begin{bmatrix}
(B_0)_{1,1}A_{1,1} + (B_0)_{1,2}A_{2,1} \\
(B_0)_{2,1}A_{1,1} + (B_0)_{2,2}A_{2,1}
\end{bmatrix}
\]

Figure 9: The data dependence graph of lowered operations for the example program in Figure 6. Red nodes represent constant matrices, blue nodes represent matrix multiplication operations and green nodes represent matrix addition operations. Each of the operations from the original data dependence graph in Figure 7 corresponds to several operations in the lowered graph (which are grouped in the shaded areas of the diagram). A single matrix multiplication operation (e.g. $B_1$) is lowered to a series of multiplications which are summed to produce the final result. Partitioning of matrices is preserved across operations, leading to increased opportunities for parallelism.
Blocks. Allows many processing elements to work on the same matrix operation simultaneously, and it exposes data parallelism that is important for two reasons: (1) it enables us to perform matrix operations on matrices that are too big to fit as a whole into the local stores of processing elements, and (2) it exposes data parallelism that is important for two reasons: (1) it enables us to perform matrix operations on matrices that are too big to fit as a whole into the local stores of processing elements, and (2) it exposes data parallelism.

Matrix partitioning is important for two reasons: (1) it enables us to perform matrix operations on matrices that are too big to fit as a whole into the local stores of processing elements, and (2) it exposes data parallelism.

Figure 10: A partial schedule of the operations from the lowered data dependence graph from Figure 9. The sub-operations of operations \( B_1 \) have all been scheduled. This schedule has been produced by the heuristic scheduling algorithm described in subsection 6.3. The operation whose operands have been scheduled to complete execution at the earliest point in time will be scheduled in the earliest available time-slot on any processor. All three stages of the pipeline must be considered for each processor — data fetch (\( df \)), execution (\( ex \)) and write back (\( wb \)). These stages can have different durations for different operations, such as multiplication and addition, as shown on the right. Red arrows show gaps in execution which we want to avoid to minimise the makespan. \( t_1 \) marks the point in time where the write back of operation \( (B_1)_{1,2} \) finishes. Thus, operation \( (B_2)_{1,2a} \) or \( (B_2)_{2,2a} \) (which depend only on \( (B_1)_{1,2} \) and the constant matrix \( A \)) can be scheduled to begin their data fetch stage any time after this point. The unlabelled blue rectangles on the end of the schedule show the result of scheduling one of these multiplication operations on each of the processors. It can be seen that the operation will complete earlier if scheduled on Processor 2, so the heuristic will choose to schedule it on Processor 2.

## 5 Lowering

Matrices can be partitioned into sub-matrices called blocks. A matrix can be partitioned across rows and columns as depicted by the dashed lines in the following example:

\[
A = \begin{bmatrix}
    a_{1,1} & a_{1,2} & a_{1,3} \\
    a_{2,1} & a_{2,2} & a_{2,3} \\
    a_{3,1} & a_{3,2} & a_{3,3}
\end{bmatrix} = \begin{bmatrix}
    A_{1,1} & A_{1,2} \\
    A_{2,1} & A_{2,2}
\end{bmatrix}.
\]

Matrix \( A \) above is thereby partitioned into a 2 \( \times \) 2 matrix whose entries \( A_{1,1}, A_{1,2}, A_{2,1} \) and \( A_{2,2} \) are blocks.

Operations on partitioned matrices can be performed by treating blocks as numerical entries. A single operation on the original matrix will be converted into several operations on blocks. This technique has been employed in the past in languages such High Performance Fortran [27] for distributing matrix operations among the nodes of a distributed computing environment. Matrix partitioning is important for two reasons: (1) it enables us to perform matrix operations on matrices that are too big to fit as a whole into the local stores of processing elements, and (2) it exposes data parallelism that allows many processing elements to work on the same matrix operation simultaneously.

In general, an \( n \times m \) matrix \( A \) is partitioned into \( p \) rows and \( q \) columns of blocks. \( a_{i,j} \) is the block in the \( i \)’th row and \( j \)’th column and it has \( k_i \) rows and \( l_j \) columns. If two matrices, \( A \) and \( B \) are said to have the same partitioning, it implies that \( p_A = p_B, q_A = q_B, \forall 1 \leq i \leq p_A : k_{i_A} = k_{i_B} \) and \( \forall 1 \leq j \leq q_A : k_{j_A} = k_{j_B} \). Similarly, if the columns of matrix \( A \) are said to have the same partitioning as the row of matrix \( B \), it implies that \( q_A = p_B, \) and \( \forall 1 \leq i \leq q_A : l_{i_A} = k_{i_B} \). We use this notation throughout the remainder of this section.
Here we describe (1) the types of matrix operations which we focus on in our work and how those operations are converted to operate on blocks, (2) the major concerns when developing a partitioning scheme, and (3) a new partitioning scheme we developed called lowering.

5.1 Block Matrix Operations

In our work we focus on three types of matrix operations: (1) unary element-wise operations, (2) binary element-wise operations, and (3) matrix multiplication. We choose to focus on this subset of matrix operations because they represent commonly used operations that fit the scope of the project. For a complete list of the block matrix operations we implemented in our framework, refer to Appendix C.

Unary element-wise operations have a single matrix operand. The result of these operations is computed by applying a mathematical function to every scalar element of the operand. The result matrix has the same dimensions as the operand. Thus, these operations have the form:

$$g(A) = \begin{bmatrix} f(a_{1,1}) & \cdots & f(a_{1,m}) \\ \vdots & & \vdots \\ f(a_{n,1}) & \cdots & f(a_{n,m}) \end{bmatrix}$$

where $g$ is the operation, $f$ is the mathematical function applied by $g$, $A$ is the $n \times m$ operand and $a_{i,j}$ are its elements.

Examples of this type of operation are matrix-scalar addition, rounding the elements of a matrix, finding the sine of each matrix element, and so on. In the case of matrix-scalar addition, the function $f$ simply adds a constant value $c$ to each of the matrix elements, i.e. $f(a_{i,j}) = a_{i,j} + c$.

These operations can be trivially modified to work on partitioned matrices because each element of the result matrix depends only on the corresponding element of the operand. Thus we can partition the matrix in any way we want and apply the operation to each of the sub-matrices. Say we partition matrix $A$ into 3 rows and 2 columns of blocks:

$$A = \begin{bmatrix} A_{1,1} & A_{1,2} \\ A_{2,1} & A_{2,2} \\ A_{3,1} & A_{3,2} \end{bmatrix}$$

operation $g$ can be applied to all 6 sub-matrices separately:

$$g(A) = \begin{bmatrix} g(A_{1,1}) & g(A_{1,2}) \\ g(A_{2,1}) & g(A_{2,2}) \\ g(A_{3,1}) & g(A_{3,2}) \end{bmatrix}$$

Thus, the resulting matrix has the same partitioning as the matrix operand.

Binary element-wise operations are similar to unary element-wise operations but have two matrix operands. The result of the operation is computed by applying a mathematical function to corresponding scalar elements of the two operands. Hence, it is a requirement of these operations that both matrix operands have the same dimensions, and the resulting matrix will also have the same dimensions as the operands. These operations have the form:

$$g(A, B) = \begin{bmatrix} f(a_{1,1}, b_{1,1}) & \cdots & f(a_{1,m}, b_{1,m}) \\ \vdots & & \vdots \\ f(a_{n,1}, b_{n,1}) & \cdots & f(a_{n,m}, b_{n,m}) \end{bmatrix}$$

where $g$ is the operation, $f$ is the mathematical function applied by $g$, $A$ is the first $n \times m$ operand with elements $a_{i,j}$ and $B$ is the second $n \times m$ operand with elements $b_{i,j}$.

Examples of this type of operation are matrix-matrix addition, matrix-matrix subtraction and element-wise multiplication. In the case of matrix-matrix addition, $f$ would be defined as $f(a_{i,j}, b_{i,j}) = a_{i,j} + b_{i,j}$.
These operations can also be trivially modified to work on partitioned matrices because each element of the result matrix depends only on the corresponding elements of the operands. In order for the operands \( A \) and \( B \) to be compatible with this operation they must have the same partitioning so that corresponding blocks have the same dimensions. Say we partition operands \( A \) and \( B \) into 3 rows and 2 columns of blocks:

\[
A = \begin{bmatrix} A_{1,1} & A_{1,2} \\ A_{2,1} & A_{2,2} \\ A_{3,1} & A_{3,2} \end{bmatrix}, \quad B = \begin{bmatrix} B_{1,1} & B_{1,2} \\ B_{2,1} & B_{2,2} \\ B_{3,1} & B_{3,2} \end{bmatrix}
\]

operation \( g \) can be applied to all 6 corresponding sub-matrices separately:

\[
g(A, B) = \begin{bmatrix} g(A_{1,1}, B_{1,1}) & g(A_{1,2}, B_{1,2}) \\ g(A_{2,1}, B_{2,1}) & g(A_{2,2}, B_{2,2}) \\ g(A_{3,1}, B_{3,1}) & g(A_{3,2}, B_{3,2}) \end{bmatrix}
\]

Thus, the resulting matrix has the same partitioning as \( A \) and \( B \).

The final type of operation we consider is matrix multiplication, which is more complex than the previous two types of operation. Matrix multiplication has two matrix operands, \( A \) and \( B \), which are required to have the same inner dimensions. That is, if matrix \( A \) is of dimension \( n \times m \), \( B \) must be of dimensions \( m \times k \). Each element of the result matrix can then be defined by the following summation

\[
g(A, B)_{i,j} = (AB)_{i,j} = \sum_{r=1}^{m} a_{i,r}b_{r,j}
\]

with a dimensions of \( n \times m \).

A technique called block matrix multiplication is used to allow multiplication of partitioned matrices. This is the same technique employed in fast matrix multiplication algorithms such as the Strassen algorithm [50] and the Coppersmith-Winograd algorithm [10] (the fastest known algorithm). Block matrix multiplication works in the same way as regular matrix multiplication but treats blocks of the partitioned operands as scalar elements. In order for the operands \( A \) and \( B \) to be compatible for block matrix multiplication, the column partitioning of matrix \( A \) must match the row partitioning of matrix \( B \). If the operands are partitioned as follows:

\[
A = \begin{bmatrix} A_{1,1} & A_{1,2} \\ A_{2,1} & A_{2,2} \\ A_{3,1} & A_{3,2} \end{bmatrix}, \quad B = \begin{bmatrix} B_{1,1} & B_{1,2} \\ B_{2,1} & B_{2,2} \end{bmatrix}
\]

then block matrix multiplication would multiply these matrices as follows:

\[
C = AB = \begin{bmatrix} A_{1,1}B_{1,1} + A_{1,2}B_{2,1} & A_{1,1}B_{1,2} + A_{1,2}B_{2,2} \\ A_{2,1}B_{1,1} + A_{2,2}B_{2,1} & A_{2,1}B_{1,2} + A_{2,2}B_{2,2} \\ A_{3,1}B_{1,1} + A_{3,2}B_{2,1} & A_{3,1}B_{1,2} + A_{3,2}B_{2,2} \end{bmatrix}
\]

Instead of one matrix multiplication \( AB \) on the individual matrix elements of matrices \( A \) and \( B \), we now have 12 multiplications and 6 addition operations on the blocks of \( A \) and \( B \). In general, each block of the result matrix is defined by

\[
C = (AB)_{i,j} = \sum_{r=1}^{q_A} A_{i,r}B_{r,j}. \quad (1)
\]

The resulting matrix has the same row partitioning as matrix \( A \) and the same column partitioning as matrix \( B \).

Block matrix multiplication can result in many addition operations on blocks to compute each block of the result matrix (denoted by the summation in Equation [1]). These additions can be performed...
in any order because of the associative property of matrix addition. We use pairwise summation to compute these additions in order maximise parallelism. This allows several addition operations to be computed in parallel. For example, if we had to add four matrix blocks together:

\[
Z = Y_1 + Y_2 + Y_3 + Y_4
\]

we could do it in several ways. A naïve technique would be to compute them in a sequential order as follows. The operations in the innermost parentheses are computed first:

\[
Z = (((Y_1 + Y_2) + Y_3) + Y_4)
\]

However, this neglects parallelism available in the summation. If we instead, add the blocks as follows:

\[
Z = ((Y_1 + Y_2) + (Y_3 + Y_4))
\]

Then, the additions \((Y_1 + Y_2)\) and \((Y_3 + Y_4)\) are computable in parallel, which reduces the execution time of the matrix multiplication. Figure 11 illustrates the improvement in performance that can be gained by using pairwise summation to add 8 matrix blocks together.

Algorithm 1 shows how we construct matrix instructions that compute the blocks of matrix \(C\) as stated in Equation (1) using pairwise summation. Therein statement \(\text{MatrixMult}\) constructs a matrix multiplication operation from two matrix block operands, and \(\text{MatrixAdd}\) constructs a matrix addition operation.

![Diagram](image_url)

**Figure 11**: Sequential and parallel (pairwise) sum computations for matrix block \(C_{i,j} = \sum_{r=1}^{8} A_{i,r} B_{r,j}\). Sequential computation of the sum of products \(C_{i,j}\) as depicted in subfigure (a) requires \(N - 1\) steps for \(N\) operands. Provided we have enough parallel execution elements we can reorder the summations as depicted in subfigure (b) where we sum in pairs, reducing the computation to \(\lceil \log_2 N \rceil\) steps. Algorithm 1 applies pairwise sums to increase parallelism with matrix multiplications on partitioned matrices.

There are known algorithms for more complex matrix operations on blocks, such as matrix inverse [57] however we have not considered these in our work so far.

### 5.2 Partitioning Schemes

A partitioning scheme determines how a matrix is split into blocks. Developing an automatic partitioning scheme can be a non-trivial exercise. We recognise four main concerns when developing a partitioning scheme for modern accelerator architectures (summarised in Figure 14):

1. Operations should be divided into approximately equal sized portions such that the load on processing elements is evenly balanced. This can reduce the makespan of execution, as illustrated in Figure 12.
Algorithm 1 Construction of instructions that compute the sum of products of matrix blocks $C_{i,j}$ using pairwise summation.

```
1   for all $i \in \{1, \ldots, p_A\}$
2     for all $j \in \{1, \ldots, q_B\}$
3       new queue $q$
4       for all $r \in \{1, \ldots, q_A\}$
5         $q$.push_back (new MatrixMult ($A_{i,r}$, $B_{r,j}$))
6     while $q$.size() > 1
7       $s_1 = q$.pop_front()
8       $s_2 = q$.pop_front()
9       $q$.push_back (new MatrixAdd ($s_1$, $s_2$))
```

(a) In this case, the operand $A$ of the operation is partitioned unevenly. This results in an uneven load on the processing elements and an increased makespan.

(b) In this case, the operand $A$ of the operation is partitioned evenly. This results in a balanced load on the processing elements and a reduced makespan.

Figure 12: Partitioning matrices evenly improves the makespan. Assume we are given the Octave code $B = A / 2$. To compute $B$, $A$ is partitioned into two blocks which can be operated on by two different processing elements. Two different partitionings of $A$ and their resulting schedules are given.

2. The small memory available on the parallel execution elements should be maximally utilised. This criteria is specific to architectures like the Cell whose SPEs have a small 256kB local store (see section 7). Peak performance is only obtained when this local store is fully utilised.

Other architectures whose parallel elements have access to a large amount of memory have a different concern which is to determine the number of blocks which a matrix should be partitioned into in order to obtain the best performance. In this case, there is a tradeoff between the number of processors utilised to execute the operation and the execution time of the operation. As a result, the partitioning of operations becomes related the scheduling of operations. This problem has been studied and is known as the problem of scheduling malleable tasks under precedence constraints. A description of the problem and proposed solutions is provided in the related work in section 2 however we do not address this concern in our scheme as we focus on the Cell architecture.

3. The number of synchronisation points in a trace should be minimised. Synchronisation points arise in a trace when two operands of an operation have an incompatible partitioning. For example, the subtraction of two matrices $C = A - B$ requires that both matrices have the same partitioning (as described in subsection 5.1 of this section). Figure 13(a) shows a situation where the matrices $A$ and $B$ were the result of other matrix operations and do not have the same partitioning. Thus, one of them must be re-partitioned in order to make them compatible. In this case, matrix $B$ is re-partitioned into the matrix $B'$ so that its blocks are the same dimensions as $A$. This allows the blocks of $C$ to then be computed in parallel. However, this re-partitioning has a negative side-effect of reducing the amount of parallelism in a trace. Each block in $B'$ contains elements from every block of the matrix $B$. Thus, in order to perform the re-partitioning, we
must wait for all of the blocks of $B$ to be computed. This can introduce gaps in the schedule and result in an increased makespan, as shown in Figure 13(b).

In contrast, if $A$ and $B$ already have the same partitioning (as shown in Figure 13(c)) then no artificial synchronisation points are introduced. In this case, each block of $C$ depends only on the corresponding blocks of $A$ and $B$. For example, $C_1$ depends only on blocks $A_1$ and $B_1$. So we can begin computing the result of $C_1$ as soon as $A_1$ and $B_1$ are available. We do not have to wait until all of the blocks in $A$ or $B$ have been computed. This results in increased parallelism in the trace and a reduced makespan, as shown in Figure 13(d).

In order to avoid synchronisation points in a trace, we want to ensure that the operands of a matrix operation have a compatible partitioning, even if the operand is a result of another operation in the trace.

4. Partitioning should be efficient. Since partitioning occurs at run-time, it is crucial that dividing operations does not incur a large amount of overhead.

These concerns can be in conflict with one another, making it difficult to produce a good partitioning scheme. For example, a naïve solution is to consider each of the matrix operations in a trace separately and determine an individual partitioning for its operands. We can optimise for concerns (1) and (2) of Figure 14 by defining the partitioning problem as a least squares fitting. We want to minimise the difference between the integer-valued block-sizes $k_i \times l_j$ and the real-valued ideal blocks of size $\frac{nm}{pq} \approx S$ as follows:

$$
\min f = \sum_{i,j} \left( \frac{nm}{pq} - k_i l_j \right)^2 \\
\text{s.t.} \quad 1 \leq k_i l_j \leq S \quad i \in [1..p], j \in [1..q] \\
\quad \frac{nm}{pq} \approx S \\
\quad k_i l_j \approx S \\
\quad l_i \geq 0 \quad i \in [1..p] \\
\quad k_j \geq 0 \quad j \in [1..q] \\
\quad \sum_{i=1}^{p} l_i = n \\
\quad \sum_{j=1}^{q} k_j = m.
$$

However, finding choices of $p$, $q$, $k_i$ and $l_j$ that optimally satisfy these criteria for each operation neglects concern (3). That is, it produces an optimal partitioning only for the operands of a single matrix operation meaning that every operation may require operands with a different partitioning. This would cause the need for re-partitioning of matrices resulting in degraded performance. Furthermore, it could be computationally expensive to find these optimal values which neglects concern (4).

Another alternative would be to examine the data dependence graph, assuming that the graph is a tree, and choose a partitioning for the leaf nodes in the graph. This partitioning could then be propagated up the tree. This approach would help to satisfy concern (3) by reducing the need for synchronisation points through the re-use of partitionings. However the problem with this approach is that the partitioning of inner nodes in the graph is determined by the partitioning of their descendants. This could lead to a partitioning for inner nodes which is unbalanced or does not fully utilise the memory of the processing elements (neglecting concerns (1) and (2)). In the worst case, this could lead to a partitioning where each block of an inner node is a single element of a matrix, which would result in very poor performance. Furthermore, in general the data dependence graph is a directed acyclic graph so propagation of partitionings could result in an operation whose partitioning is determined by several different partitionings. In this case a barrier must be introduced to resolve the conflict.
Figure 13: Re-partitioning matrices can degrade performance. A matrix subtraction $C = A - B$ is being computed. Figure 13(a) shows a situation where the operands $A$ and $B$ have a different partitioning and thus are incompatible. $B$ is re-partitioned to $B'$ to match $A$'s partitioning. This allows computation of $C$ to proceed in parallel. However, each block of $B'$ depends on every block of $B$. This acts as a barrier, forcing computation of the entire $B$ matrix before computation of $C$ can begin. This reduces potential parallelism in the trace, as shown in Figure 13(b). In contrast, if $A$ and $B$ have the same partitioning we can avoid re-partitioning and no barriers are introduced, resulting in an improved makespan of the schedule, as shown in Figures 13(c) and 13(d).
5.3 A New Partitioning Scheme

In this work, we introduce a new technique called *lowering* for partitioning matrix operations in an execution trace. The main benefit of our partitioning scheme is that all operands of an operation in a trace are guaranteed to have a compatible partitioning with each other, without ever needing to re-partition matrices. This maximises the amount of parallelism available in the trace. Our partitioning scheme also finds a uniform partitioning of matrices, such that all blocks in a matrix are of approximately the same size. Finally, since partitioning of a matrix relies solely on the dimensions of that matrix, no other operations need to be examined in order to compute the partitioning (i.e. partitioning is local to every matrix). This leads to an efficient algorithm which examines each matrix in the execution trace only once. Thus, we satisfy concerns (1), (3) and (4) from Figure 14. The only disadvantage of our partitioning scheme is that in certain cases it can lead to sub-matrices which are smaller than the capacity of the memory on the SPEs. This can result in under utilisation of the SPEs. Thus there are situations where we fail to address concern (2).

To be applicable to a wider range of architectures, our partitioning scheme abstracts from the underlying hardware, but only to the extent that performance is not sacrificed. We were able to reduce the dependencies of our partitioning scheme to two quantities from the underlying Cell Broadband Engine architecture:

1. The maximum number of matrix elements $S$ we can store in a single matrix block. The local store of a Cell SPE provides 256kB of memory to be shared by program code and data. We use a small ($\approx$26kB) kernel program on each SPE to execute matrix operations. The remaining 230kB of the local store is available for matrix blocks. Because of Cell-related implementation techniques discussed in subsection 7.3 we can devote up to 38kB to a single matrix block, which amounts to $S = 9216$ single precision floating point matrix elements, or $S = 4608$ matrix double precision floating point elements per block.

2. A divisor $\delta$ for the number of rows and columns of a block. For several Cell-related reasons discussed in subsection 7.3 the number of rows and columns in a block must be a multiple of $\delta = 4$ with single precision and a multiple of $\delta = 2$ with double precision. To guarantee that each block has a multiple of $\delta$ rows and columns, it may be necessary to pad rows and columns of a matrix with zero elements. An $n \times m$ matrix may require up to $\delta - 1$ additional rows and/or columns of zeroes, as shown in Figure 15(b). The number of elements of the resulting $n' \times m'$ matrix is bound by $n \times m + (\delta - 1)(n + m) \geq n' \times m'$.

Figure 15(b) shows how we partition an $n \times m$ matrix $A$ into $p$ rows and $q$ columns of blocks. Block $A_{i,j}$ of this partition has $k_i$ rows and $l_j$ columns. The underlying idea of our partitioning scheme is that we limit the shape of a matrix block to a maximum of $\sqrt{S}$ rows and columns. When the divisor $\delta$ is also taken into account, each sub-matrix is limited further to $\delta \lfloor \sqrt{S} \rfloor$ rows and columns. Making this assumption about the maximum number of rows and columns in each block allows us to determine the number of rows of blocks $p$ and columns of blocks $q$ which are required for a matrix:
of blocks in a partitioning. Likewise for the number of columns
rows. Doing the same for the columns results in each block of the matrix having approximately the
⌈rows that fit in a single matrix block. Dividing the former by the latter yields the number of rows

\[ p = \left\lfloor \frac{n}{s} \right\rfloor, \quad \text{and} \quad q = \left\lfloor \frac{m}{\delta} \right\rfloor. \]

Therein \( \left\lfloor \frac{n}{s} \right\rfloor \) is the number of groups-of-δ-rows of a partition, and \( \left\lfloor \frac{m}{\delta} \right\rfloor \) is the number of groups-of-δ-rows that fit in a single matrix block. Dividing the former by the latter yields the number of rows \( p \) of blocks in a partitioning. Likewise for the number of columns \( q \).

Once we have determined the number of rows of blocks \( p \), we divide the rows of the original matrix into those blocks. This ensures that each block in the matrix has approximately the same number of rows. Doing the same for the columns results in each block of the matrix having approximately the same number of elements, leading to a balanced partitioning (satisfying concern (1)). The number of rows \( k_i \) and columns \( l_j \) of a block are given by

\[ k_i = \delta \left( \left\lfloor \frac{n}{p} \right\rfloor + r_i \right) \quad \text{and} \quad l_j = \delta \left( \left\lfloor \frac{m}{q} \right\rfloor + s_j \right), \]
When the number of rows in a matrix does not divide evenly into \( p \), there will be left-over rows. These are distributed among the existing row partitions. If a left-over row is included in a block, it is denoted by the inclusion of a non-zero \( r_i \) term in the calculation of \( k_i \). Likewise for left-over columns. \( r_i \) and \( s_j \) are chosen such that 
\[
\sum_{i=1}^{p} r_i = \left\lceil \frac{n}{\delta} \right\rceil \mod p \quad \text{where} \quad r_i \in \{0, 1\},
\]
and 
\[
\sum_{j=1}^{q} s_j = \left\lceil \frac{m}{\delta} \right\rceil \mod q \quad \text{where} \quad s_j \in \{0, 1\}.
\]
For consistency across matrices we set \( r_i = 1 \) and \( s_j = 1 \) for the smallest indices \( i \) and \( j \) of blocks \( A_{i,j} \) of a matrix partition.

Note that the number of rows in a block, \( k_i \) depends on the divisor \( \delta \), the number of rows in the original matrix \( n \), the number of rows of blocks \( p \) and its position \( i \) in the partitioning (due to the left-over rows). \( p \) depends on \( n \), \( \delta \) and \( S \), the buffer size. Assuming \( \delta \) and \( S \) are constants, this means that \( k_i \) depends only on \( i \) and \( n \) and likewise for the columns. Thus, our partitioning scheme ensures that if two matrices \( A \) and \( B \) have the same number of rows and columns, they will be partitioned in the same way. It also guarantees that if the number of columns in \( A \) is equal to the number of rows \( B \) then the column partitioning of \( A \) will match the row partitioning of \( B \).

With our partitioning scheme, we claim that matrices never have to be re-partitioned. This implies that only the constant matrices in a trace are partitioned. Matrices that result from other operations must be guaranteed to be compatible in any operation in which they are used. Thus, in our example in Figure 3, \( A \) and \( B \) must be guaranteed to have the same partitioning, even if they are the results of other operations in the trace. This is further complicated by the fact that \( A \) and \( B \) might be the operands of more than one operation.

To show this, we first show that if the operands of a matrix operation have been partitioned according to our partitioning scheme then the operands are guaranteed to be compatible. The requirements for compatibility are described in subsection 5.1 of this section. We must do this for each type of operation we consider in our framework:

1. **Unary Element-wise Operations**: Trivially compatible because these operations do not have any requirement on the partitioning of the single operand.

2. **Binary Element-wise Operations**: These operations have two operands \( A \) and \( B \) which are required to have the same partitioning to be compatible. This is ensured for our scheme because \( A \) and \( B \) are of the same dimensions, thus they will have the same partitioning.

3. **Matrix Multiplication**: This operation has two operands, \( A \) and \( B \). The column partitioning of \( A \) must match the row partitioning of \( B \) for these to be compatible. This is ensured for our scheme because the number of columns in \( A \) matches the number of rows in \( B \).

We now show that if the operands of an operation have been partitioned according to our partitioning scheme, then the result of the operation will also be partitioned according to our scheme. The partitioning of the results of operations is also described in subsection 5.1 of this section. We must do this for each type of operation we consider in our framework:

1. **Unary Element-wise Operations**: The result matrix has the same partitioning as the input matrix. Thus it will be partitioned according to our scheme.

2. **Binary Element-wise Operations**: The result matrix has the same partitioning as both input matrices. Thus it will be partitioned according to our scheme.

3. **Matrix Multiplication**: The result matrix has same number of rows and row-partitioning as first operand. It also has same number of columns and column-partitioning as the second operand. Thus it will be partitioned according to our scheme.

Now recall that the data dependence graph of operations is a directed acyclic graph, whose source nodes are constant matrices. Thus, every operation either depends on constant matrices, or on other operations which in turn depend on constant matrices. Since the constant matrices are partitioned with our scheme, the results of other operations will be partitioned according to our scheme. This means that every matrix operand in the trace will be partitioned according to our scheme and so
operands will always be compatible with each other. So re-partitioning is never needed, satisfying concern (3).

The only downside of this partitioning scheme is that for very small-sized matrices and vectors, the partitioning can lead to under-utilisation of the buffer. For example, assume $\delta = 1$ (for simplicity) and we wish to partition a row vector of dimensions $1 \times m$ with a buffer size of $S$. The partitioning occurs as follows:

1. The number of rows $p$ of sub-matrices and columns $q$ of sub-matrices is given by:
   
   $p = \lfloor \frac{1}{\sqrt{S}} \rfloor = 1$
   
   $q = \lfloor \frac{m}{\sqrt{S}} \rfloor$

2. If we ignore the spare rows $r_i$ and spare columns $s_j$, the number of rows per sub-matrix $k_i$ and columns per sub-matrix $l_j$ is given by:
   
   $k_i = \lfloor \frac{1}{p} \rfloor \leq 1$
   
   $l_j = \frac{m}{q} \leq \frac{m}{\sqrt{S}} \leq \sqrt{S}$

Thus, the number of elements in a block is $k_i l_j \leq \sqrt{S}$. So only a square-root of the buffer size is used at most. This results in poorer performance for operations with these kinds of matrices as operands due to the increased number of sub-operations and relative overhead to computation ratio. This could be improved by using a different partitioning for these kinds of matrices, however this would force barriers to be introduced when the partitioning changes. We leave this consideration as future work.

The output of the lowering process is a lowered data dependence graph. This graph is produced through a graph transformation of the original data dependence graph. The transformation algorithm examines each operation in the original graph in a topological order, partitions its matrix operands and produces several new operations which operate on the blocks of the partitioned matrices. Data dependencies are updated according to the partitioned operations. Hence the lowered graph typically has an increased number of nodes (operations) and edges (dependencies) over the original graph.

6 Scheduling

The task of the scheduler is to distribute matrix operations among the processing elements of the parallel architecture. The input of the scheduler is the lowered dependence graph annotated with estimated time durations for each pipeline stage of an operation. The objectives of the scheduler are:

1. to produce a feasible schedule, i.e., operands of a matrix operation must not be scheduled after the operation,

2. to generate the schedule for the MEUs as fast as possible, and
3. to minimise the makespan, i.e., the wall-clock time needed to execute the parallel schedule on the MEUs.

The second and third objectives are hard to achieve because scheduling is an NP-hard problem [16]. Recognising the difficulty of finding an optimal solution for the general problem, many heuristics have arisen, as described in subsection 2.2 of the Related Work.

Our scheduling problem is a variant of the static scheduling problem with arbitrary task precedence constraints, arbitrary execution times for tasks, and uniform workers/processors, as described in subsection 2.2.1. The data dependence graph in our framework resembles the task precedence graph, the workers are the MEUs and the time durations are estimated ahead of time for each lowered matrix instruction. Although there exists approximation algorithms for this problem (see subsection 2.2) they are based on rounding of a relaxed linear programming solution with high practical run-times. Thus, they are not viable for our framework which needs to incur a low overhead. Furthermore, existing algorithms do not take into account the notion of a pipeline for each of the workers where execution of tasks and communication between processors can be overlapped. Thus, using these algorithms would lead to imprecision in the modelling: the communication overheads of matrix instructions would be ignored and the pipeline might stall because of imprecise modelling of matrix operations.

In the following we discuss (1) how to find a time model to accurately estimate the durations of each pipeline stage of a lowered matrix operation, (2) a mathematical program for computing the optimal schedule for the pipelined scheduling problem with task precedence constraints, and (3) a new greedy algorithm for solving the scheduling problem efficiently.

An accurate time model for the scheduling is crucial to find an effective schedule. The mathematical program for finding schedules is not practical but for small problem sizes it gives us a yardstick to compare how good our greedy algorithm performs in comparison to the optimal solution. Because the problem is intractable, we cannot hope for computing schedules with the mathematical program for problems with medium to larger size. Results of the accuracy of time models and the performance of the heuristic scheduling algorithm are provided in section 9.

### 6.1 Time Model

We compute a time model for each type of matrix instruction in our framework, and we use multivariate polynomial functions for the time model that depend on the number of rows and columns of the input and output matrices. The coefficients of the multivariate polynomial are computed using profiling and Ordinary Least Square (OLS) method.

For scheduling we require an accurate time model that estimates the durations of the three phases of a matrix instruction before executing it. Recall from section 6 that we model three pipeline stages for a matrix instruction executing on a processing element:

1. **Data Fetch (df):** The operands of the matrix instruction are loaded from main memory into the memory of the parallel processing element,

2. **Execute (ex):** The matrix instruction is executed on the parallel processing element,

3. **Write Back (wb):** the result of the matrix instruction is written back to main memory.

For computing the time model we instrument the virtual machine that runs on the SPEs (called MEUs). As a side-effect of execution we obtain profiling information. The instrumented version of the MEUs is only executed for profiling purposes to obtain the execution times for each pipeline stage of an instruction. Furthermore, we have a carefully crafted input program that executes each type of matrix instruction for every possible input problem size several times. Note that it is feasible to profile every input problem size since the matrices are limited by the SPE's small data and program memory of 256kB.

The Cell Broadband Engine offers hardware counters [24] for performing the measurements with very high precision. The execution time measurements on the SPEs have little variation due to the lack...
of data caches. However, we observed a higher volatility in the measurements for the data fetch and write back stages which is not surprising since the ring-bus of the Cell that connects SPEs, memory and the PPE, is shared.

The coefficients of the multivariate polynomial are chosen such that the deviation of the polynomial function from the measured time durations is minimised. Assume we have \( k \) time measurements obtained by profiling one type of matrix operation (such as matrix addition). For the \( k \)-th measurement we are given a vector \( \vec{n}_k \) describing the input problem size and the durations \( \Delta_{df}(k) \), \( \Delta_{ex}(k) \), and \( \Delta_{wb}(k) \) for each phase of the pipeline. The input problem size vector typically describes the dimensions of the operands of the instruction. For example a matrix addition has two elements in the input problem size vector describing the number of rows and columns of the two matrices which are to be added together.

We seek a multivariate polynomial for each pipeline phase of each type of matrix instruction:

\[
t_\xi(\vec{n}) = \sum_{k} a_k g_k(\vec{n}) \quad (2)
\]

where \( g_k(\vec{n}) \) is a multivariate term of the multivariate polynomial and \( a_0, \ldots, a_n \) are the coefficients such that the error

\[
R_\xi = \sum_{k=1}^{l} (t_\xi(\vec{n}_k) - \Delta_\xi(k))^2 \quad (3)
\]

becomes minimal (where \( \xi \) is either \( df \), \( ex \) or \( wb \)).

In the following table we give the multivariate polynomials used for estimating the execution times of some types of matrix instructions, the data fetch, and write back duration. Note that the data fetch and write back phases do not depend on the type of operation being executed.

| \( t_\xi(\vec{n}) \) | Operation |
|----------------------|-----------|
| \( a_0 + a_1n_1 + a_2n_2 \) | data fetch and write back |
| \( a_0 + a_1n_1n_2 \) | scalar instr. execution |
| \( a_0 + a_2n_1 + a_1n_1n_2n_3 \) | matrix multipl. execution |

The variables \( n_1, n_2 \) and \( n_3 \) represent the rows and columns of the input matrices which form the problem size vector \( \vec{n} \). For matrix multiplications we do not need to specify the number of rows of the second matrix because it is equal to the number of columns of the first matrix. Furthermore, we add an additional term \( a_2n_1 \) to account for the overhead of the inner-most loop in the matrix calculation to obtain a better fit of the profile data. We used standard methods to obtain the coefficients [31].

For the remainder of the section, we use \( t_\xi(i) \) to denote the function \( t_\xi(\vec{n}) \) where \( \vec{n} \) is the problem size vector of instruction \( i \).

### 6.2 Mathematical Program

We develop an integer linear program that computes an optimal schedule for a given problem instance that consists of the set of matrix instructions \( I = \{1, \ldots, n\} \), their data dependencies \( E \subseteq I \times I \) where \( (i, j) \in E \) denotes that \( j \) depends on \( i \), and the time parameters \( t_{df}(i), t_{ex}(i), \) and \( t_{wb}(i) \) for instructions \( i \in I \). For the model we introduce the following variables,

\[
x_{ij} \in \{0, 1\} \quad i, j \in I
\]

\[
t_i \in \mathbb{R}^+ \quad i \in I
\]

\[
z \in \mathbb{R}^+
\]

where \( z \) is the makespan of the schedule, \( t_i \) is the start time of an instruction and \( x_{ij} \) are elements of an adjacency matrix for the schedule graph which is a directed rooted graph which forms the schedule of operations executed by each processing element.

The structure of a schedule graph is depicted in Figure 16. It is defined to have a dedicated start node \( s \), with at most \( p \) outgoing edges, and no incoming edges. The time parameters of the start node
are all zero and the start node is not a matrix instruction that is executed on a processing element. The remaining nodes in the schedule graph represent matrix instructions that will be executed. Each of these nodes has exactly one predecessor node and at most one successor node. The successor and predecessor node must not be the node itself, i.e., in the graph we do not allow self-loops. The successors nodes of the start nodes (e.g. $v_1$ and $v_3$ from Figure 16) represent the first nodes that will be executed by each processing element. Their successor nodes ($v_2$ and $v_4$) are the second instructions that will be executed, and so on.

We have at most $p$ outgoing edges for the start node $s$ and hence at most $p$ processing elements. The linear constraints that ensure that elements $x_{ij}$ of the adjacency matrix form a schedule graph, are given below:

\[
\begin{align*}
\sum_{i=1}^{n} x_{1i} & \leq p \\
x_{1i} & = 0 \quad i \in I \\
\sum_{j=1}^{n} x_{ij} & \leq 1 \quad i \in I \setminus \{1\} \\
\sum_{j=1}^{n} x_{ji} & = 1 \quad i \in I \setminus \{1\} \\
x_{ii} & = 0 \quad i \in I \setminus \{1\}
\end{align*}
\]

For the task precedence relation $E$ we introduce the time constraints

\[t_i + t_{dy}(i) + t_{exe}(i) + t_{wb}(i) \leq t_j \quad (i, j) \in E\]

that forces an instruction $j \in I$ to be scheduled after its operands $i : (i, j) \in E$ are completed. The time $t_i$ is a global time for all matrix execution units, i.e., operands may be scheduled on different matrix execution units and all matrix execution units have the same wall-clock time.

For two subsequent instructions on a processing element, the durations of the three pipeline stages (data fetch, execute, and write back) cannot overlap. To ensure this, a time constraint is introduced

Figure 16: Schedule Graph: the start node $s$ has at most $p$ outgoing edges and no incoming edges; every other node in the schedule graph has at most one outgoing edge and exactly one incoming edge; self-loops are not permitted. $s$ is a artificial instruction which is not executed. A line of nodes in the schedule graph represents the sequence of instructions that are executed by one of the processing elements. For example, instructions $v_1$ and $v_2$ are executed by the first processing element, in that order. $v_3$ and $v_4$ are executed by the second processing element, and so on.
for each pipeline stage:

$$\sum_{i=1}^{n} (t_i + t_{df}(i)) x_{ij} \leq t_j \quad j \in I$$

$$\sum_{i=1}^{n} (t_i + t_{df-ex}(i)) x_{ij} \leq t_j + t_{df}(j) \quad j \in I$$

$$\sum_{i=1}^{n} (t_i + t_{df-ex-wb}(i)) x_{ij} \leq t_j + t_{df-ex}(j) \quad j \in I$$

where $t_{df-ex}(i) = t_{df}(i) + t_{ex}(i)$ and $t_{df-ex-wb}(i) = t_{df}(i) + t_{ex}(i) + t_{wb}(i)$ for all $i \in I$. The left-hand side of the constraints is a summation of the time duration multiplied with $x_{ij}$ over all predecessors. However, in a schedule graph there exists at most one predecessor modelled by the constraints for the schedule graph.

The expansion of above constraints have a non-linear term $t_i x_{ij}$ which we linearise by introducing a new variable $y_{ij}$. This new variable replaces the quadratic term $t_i x_{ij}$, and we add linear constraints to the program that force the equivalence. The linear constraints are developed by using standard techniques [3], i.e.,

$$y_{ij} \in \mathbb{R}^+ \quad i, j \in I$$

$$y_{ij} \leq U x_{ij} \quad i, j \in I$$

$$t_i + U x_{ij} - U \leq y_{ij} \leq t_i \quad i, j \in I$$

where $U$ is the sum of all time parameters, which is an upper bound for $y_{ji}$ and $t_{ji}$.

The set of constraints for the makespan can only be greater than the completion time of all instructions in $I$, i.e.,

$$t_i + t_{df-ex-wb}(i) \leq z \quad i \in I$$

(4)

and the objective function of the mathematical program is to minimise the makespan $z$. The integer linear program is given at a glance in Appendix A and its implementation as an AMPL [15] script is given in Appendix B.

6.3 Greedy Algorithm

The integer linear programming formulation (described previously) of the task-precedence scheduling problem for an asynchronous super-pipelined, super-scalar matrix engine is intractable. To overcome this problem, we devise a simple heuristic as shown in Algorithm 2 which exhibits a worst-case runtime of $O(n \log n + m)$ where $n$ is the number instructions and $m$ is the number of dependencies in the task precedence graph. In the algorithm, the sets $d^-(i)$ and $d^+(i)$ denote the predecessor and successor sets of an instruction $i \in I$ in the task precedence graph.

The algorithm is based on a list scheduling approach (see subsection 2.2.1 of the related work). The list scheduling algorithm uses a heuristic to order tasks in a list and then greedily distribute them among processors. The underlying idea of the heuristic is to find a topological order for the acyclic data dependence graph $G(I, E)$. This total order $(i_1, \ldots, i_n)$ for the instructions $I$ has the property that for each data dependence $(i, j) \in E$ instruction $j$ is listed after instruction $i$ in the total order. The topological order ensures that instructions never deadlock in its execution, i.e., the operands are either available or the instructions have to wait a finite amount of time for their operands to become available. The topological order is computed with the counter $c_i$ for an instruction $i \in I$. Initially, the counters are set to the number of incoming edges, i.e., $|d^-(i)|$. Only instructions with counter values of zero can be scheduled, and are maintained in a priority queue (min-heap). The priority is determined by the earliest point in time when an instruction can be scheduled. After scheduling an instruction $i$ on one of the processing elements, the counters of its successors $j \in d^+(i)$ in the task precedence graph
Algorithm 2 Heuristic Scheduler: earliest instruction in earliest stream.

1. for all $k \in \{1, \ldots, p\}$
2. \(s_{df}(k) \leftarrow 0\)
3. \(s_{ex}(k) \leftarrow 0\)
4. \(s_{wb}(k) \leftarrow 0\)
5. for all $i \in I$
6. \(c_i \leftarrow |d^{-}(i)|\)
7. \(e_i \leftarrow 0\)
8. if $|d^{-}(i)| = 0$
9. queue $i$ with cost 0
10. while queue not empty
11. dequeue $i$
12. $k \leftarrow \text{minarg}_r\{h(i, r)\}$
13. $t_i \leftarrow h(i, k)$
14. add $i$ to stream $k$
15. $s_{df}(k) \leftarrow t_i + t_{df}(i)$
16. $s_{ex}(k) \leftarrow t_i + t_{df-ex}(i)$
17. $s_{wb}(k) \leftarrow t_i + t_{df-ex-wb}(i)$
18. for all $j \in d^{+}(i)$
19. \(c_j \leftarrow c_j - 1\)
20. \(e_j \leftarrow \max(e_j, t_i + t_{df-ex-wb}(i))\)
21. if $c_j = 0$
22. enqueue $j$ with cost $e_j$

are decremented by one. The successors are scheduled (put into the queue) as soon as their operands have been allocated to a processing element.

From the set of instructions whose operands have been scheduled (or do not have operands) we choose the instruction that can be scheduled the earliest on the time line. The earliest scheduling time $e_i$ of an instruction $i \in I$ is determined by the completion time of the instruction’s operands. When an instruction is scheduled, the earliest scheduling time of its dependent instructions is updated (cf. line 20 in Algorithm 2).

For each instruction we have exactly one enqueue and one dequeue from the priority queue resulting in a worst-case complexity of $O(n \log n)$. We also need to update the earliest scheduling time $e_i$ of every instruction which has a worst-case complexity of $O(m)$. Thus, the worst-case runtime complexity of the algorithm given in Figure 2 is $O(n \log n + m)$. Note that we have the underlying assumption that the number of processors $p$ is constant in this analysis.

Once the earliest instruction is selected we choose to schedule the instruction on the processing element that can execute the instruction at the earliest point in time. For the selection of the processing element, we use a slot function $h(i, k)$ which computes the earliest possible start of instruction $i$ on processing element $k$. The slot function takes into account the completion times of the data fetch, execute, and write back phase of the previous instruction. We keep track of the completion time of the last scheduled instruction on processing element $k$ for each of the pipeline stages. These are denoted by $s_{df}(k)$, $s_{ex}(k)$, and $s_{wb}(k)$ for data fetch, execution and write back, respectively. The slot function is given by

\[
h(i, k) = \max(\max(e_i, s_{df}(k)) + t_{df}(i), s_{ex}(k)) + t_{ex}(i), s_{wb}) - (t_{df}(i) + t_{ex}(i))
\]

and depicted in Figure 17. For an instruction, $i$ being scheduled on processing element $k$, the start time of each of its pipeline stages could be limited by two things. The first is the completion of $i$’s operands, i.e. $i$ cannot begin data fetch until its operands are complete. The second is the completion
Figure 17: Example input for slot function $h(i, k)$. $h(i, k)$ is the earliest point in time when instruction $i$ can be scheduled. $s_{df}(k)$, $s_{ex}(k)$, and $s_{wb}(k)$ denote the last completion times of each phase of the last instruction scheduled on a processing element. The start time of the data fetch of $i$ is not limited by the completion time of its operands $e(i)$ but instead the completion of the data fetch of the previous instruction $s_{df}(k)$. The execution phase is also limited by the completion of the execution of the previous instruction. The start time of the write back of $i$ is limited by the completion of the execution of $i$. From the estimated start time of the write back operation, we subtract $t_{df}(i) + t_{ex}(i)$ to yield $h(i, k)$ which is the earliest time at which $i$ can begin of the execution of $i$ pipeline.

of the previous instruction on processing element $k$, i.e. $i$ cannot begin its data fetch phase until the data fetch of the previous instruction on $k$ is complete (and likewise for the execution and write back phases). The three max functions here are used to determine which of these is the limiting factor in determining the start time of $i$.

In the example in Figure 17 the earliest execution time $e(i)$ of instruction $i$ is before the completion of the data fetch stage of the previous instruction $s_{df}$. Thus, the completion of the data fetch stage of the previous instruction is taken as the earliest start time for the data fetch of $i$. We add to this point in time the estimated duration of data fetch for $i$ and compare it with the completion of the execution stage of the previous instruction, $s_{ex}$. This allows us to determine the earliest point in time for commencing the execution stage of $i$. In the example, the completion of the execution stage of the previous instruction takes longer and hence determines the earliest starting time for the execution of $i$. We add to this point in time the duration of the execution phase of $i$ and compare it with the completion of the write back phase of the previous instruction, $s_{wb}$. This allows us to determine at what time the operation can begin its write back. In this case, the write back stage is limited by the completion of the execution phase of $i$. Finally, from the point in time that the write back of $i$ will finish, we subtract the duration of data fetch and execute stage to obtain the earliest point in time to slot instruction $i$ into processing element $k$.

7 Cell Computation Engine

7.1 The Cell Broadband Engine

The Cell Broadband Engine architecture (Cell) is a heterogeneous multicore architecture that was jointly developed by IBM, Sony and Toshiba [19]. The structure of the Cell is shown in Figure 7.1. It consists of a 64-bit PowerPC core (PPE), eight SIMD cores called Synergistic Processing Elements (SPEs), a memory interface controller and an I/O controller. The PPE and SPEs communicate through a high-speed Element Interconnect Bus (EIB). At a 3.2 GHz clock rate, the theoretical peak performance for a single SPE with single-precision floating-point operations is 25.6 GFLOPS, yielding an overall performance of 204.8 GFLOPS for 8 SPEs. For double-precision the theoretical peak performance for a single SPE is 12.8 GFLOPS, and 102.4 GFLOPS aggregate. The EIB supports a
peak bandwidth of 204.8 GB/s for on-chip data transfers between the PPE, SPEs, memory interface controller and I/O controller. The memory interface controller provides 25.2 GB/s peak bandwidth to main memory.

The PPE is the Cell’s main processor, designated to run the operating system, coordinate the SPEs and perform the control-intensive part of applications. The PPE’s memory hierarchy is similar to conventional processors, with 32kB level-1 instruction and data caches and a 512kB level-2 cache.

SPEs are designed for high-performance data-streaming and for data-intensive computations. Their memory hierarchy consists of a 128x128-bit SIMD register file, 256kB of local store memory and the off-chip main memory shared with the PPE. SPEs can run SIMD operations at four different granularities: 16-way 8-bit integers, 8-way 16-bit integers, four-way 32-bit integers, four-way single-precision floating-point numbers, or two-way 64-bit double precision floating point numbers. The 256kB local store of an SPE is shared for code and data. Each SPE can only access the code and data in its own local store. DMA transfers are used to move data between the local store of an SPE and main memory, as well as between the local stores of different SPEs. DMA transfers are asynchronous and enable SPEs to overlap computation with communication. Unlike caches, SPE local stores must be explicitly managed by software.

Additionally, a mailbox mechanism is provided for communication between the PPE and each of the SPEs. This acts as a queue which the PPE can enqueue data items onto and the target SPE can dequeue values from. Each mailbox is capable of holding up to four 32 bit data items at any instant. If a PPE attempts to enqueue more than four data items, one data item will be overwritten or execution will be blocked until an item is dequeued.

### 7.2 Matrix Execution Units

As mentioned in section 3, we implement the computation engine component of our framework for the Cell architecture. At the core of the Cell computation engine are the Matrix Execution Units.
(MEUs). An MEU is a small virtual machine which runs on an individual SPE and executes matrix operations. There is one MEU running on each of the SPEs available in the Cell processor (typically 8). The fact that there are multiple MEUs capable of executing matrix instructions concurrently gives our framework a *superscalar* property. Each MEU idles on its SPE, waiting to be notified by the PPE that there are matrix operations ready for execution. The PPE uses the mailbox mechanism available in the Cell processor to inform an MEU of the memory location of such a ready operation.

After receiving this message, the MEU uses Direct Memory Access (DMA) list commands to transfer the operands involved in the matrix operation from main memory to the local store of the SPE on which the MEU is running. Once the transfer is complete, the MEU executes the matrix operation and again uses DMA list commands to transfer the result of the operation back to main memory. This process repeats for each matrix operation that has been assigned to the MEU.

In order to achieve high performance on the Cell Broadband Engine Architecture, the latency of memory transfers between main memory and an SPE’s local store must be hidden by overlapping them with computation. This technique is known as multi-buffering and is facilitated in the Cell processor by non-blocking DMA commands. When an SPE uses a DMA command to transfer data between its local store and main memory, the transfer is carried out by a separate processor known as the Memory Flow Controller (MFC) and execution can continue on the SPE. The SPE can then issue a further DMA command which will force completion of the transfer, when required.

The Cell Matrix Engine uses a form of multi-buffering known as triple-buffering to hide data transfer latencies. Three buffers are allocated on the local store of each SPE. At any point in time, exactly one of the following data items will be stored in each of the buffers:

1. The operands of the current operation being executed.
2. The result of the current operation being executed.
3. The result of the last operation that was executed as it is transferred out or the operands of the next operation to be executed as they are transferred in.

The use of this triple-buffering technique results in *pipelined* execution of operations on each of the MEUs. These pipeline stages are taken into account by our scheduling algorithm and were described in section 6. However, here we describe them again in the context of the Cell implementation. There are three pipeline stages:

1. **Data Fetch (df):** The operands of an operation are loaded from main memory into the local store of an SPE. We use the notation $df_{start}(op, B_i)$ to denote the initiation of a DMA transfer of operation $op$’s operands into buffer $B_i$ on an SPE. This is a non-blocking call. We use the notation $df_{finish}(op, B_i)$ to denote a blocking call that ensures the completion of the same transfer.

2. **Execute (ex):** The operation is executed on an SPE. We use the notation $ex(op, B_i, B_j)$ to denote the execution of operation $op$ whose operands are contained in buffer $B_i$ and whose result is to be placed in buffer $B_j$.

3. **Write Back (wb):** The result of the operation is transferred from the local store of an SPE back to main memory. We use the notation $wb(op, B_j)$ to denote a non-blocking call to initiate the transfer of $op$’s result which is stored in buffer $B_j$ back to main memory. Note that this call also issues a barrier command which means that the write back transfer must be fully complete before any further DMA commands to buffer $B_j$ can begin. This prevents the chance of race conditions.

Figure 19 describes the triple buffering process for executing three matrix operations: $op_1$, $op_2$ and $op_3$. Algorithm 3 shows pseudocode of the execution loop which runs on the MEU virtual machines and facilitates triple buffering. Therein, the variable $in$ is the number of the buffer containing the operands of the current operation being executed, $out$ is the number of the buffer containing the result
(a) Initially the execution pipeline is empty, and all buffers on the SPE are empty. The first operation, $op_1$ is offered from the PPE and its operands are fetched from main memory into buffer $B_0$. As there is no operation that is ready to execute, the SPE initiates transfer of the next operations ($op_2$’s) operands into the second buffer, $B_1$.

(b) Execution on the SPE blocks until the completion of the data transfer of $op_1$’s operands. When the transfer is completed, execution of $op_1$ begins and the result is placed into buffer $B_2$.

(c) The transfer of the result of $op_1$ back from $B_2$ to main memory is initiated. A DMA barrier command is placed after the write back operation and the data fetch of the next operation ($op_3$) into $B_3$ is also requested. However, this transfer does not begin until the write back of $op_1$ is complete due to the barrier. Execution on the SPE is blocked until transfer of $op_2$ is complete and then the execution of $op_2$ begins, with the result being placed in $B_0$.

(d) Once the write back of $op_1$ is complete, the data fetch of $op_3$ into buffer $B_2$ begins. Execution of $op_2$ continues. We have now filled the execution pipeline and are in the pipeline state as in Figure 19(b).

Figure 19: An illustration of the triple buffering process on the SPEs. Each SPE has 3 buffers labelled $B_0$, $B_1$ and $B_2$. Three matrix operations, $op_1$, $op_2$ and $op_3$ are executed.
Algorithm 3 Execution loop for the MEUs which facilitates triple buffering.

1. \( i \leftarrow 0 \)
2. \( \text{loop} \)
3. \( \text{in} \leftarrow i \mod 3 \)
4. \( \text{next} \leftarrow (i + 1) \mod 3 \)
5. \( \text{out} \leftarrow (i + 2) \mod 3 \)
6. \( \text{op}_{\text{next}} \leftarrow \text{read mailbox} \)
7. \( \text{if } \text{op}_{\text{next}} = \text{terminate} \)
   8. \( \text{exit} \)
9. \( \text{df}_{\text{start}}(\text{op}_{\text{next}}, B_{\text{next}}) \)
10. \( \text{df}_{\text{finish}}(\text{op}_{\text{in}}, B_{\text{in}}) \)
11. \( \text{ex}(\text{op}_{\text{in}}, B_{\text{in}}, B_{\text{out}}) \)
12. \( \text{wb}(\text{op}_{\text{in}}, B_{\text{out}}) \)
13. \( i = i + 1 \)

of the current operation and \( \text{next} \) is the number of the buffer containing the operands of the next operation to be executed.

The matrix operations in the MEUs, such as matrix multiplication and matrix addition, must have implementations that are highly optimised for the SPEs in order to obtain the best performance. There are several considerations when optimising algorithms for the SPEs [24]. SIMD (Single Instruction, Multiple Data) instructions must be used to exploit the 128-bit wide vector processing capabilities of the SPEs. Branches must be eliminated to reduce the chance of costly branch misprediction and loops should be unrolled to expose a maximum amount of instruction-level parallelism and utilise both execution pipelines of the SPEs.

These factors make writing optimised operations for the SPEs a challenging and time-consuming task. Where possible, we utilise the optimised libraries provided with the Cell SDK to perform operations on the SPEs. For example, the Large Matrix Library [25] and SPE BLAS library [25] provide optimised matrix multiplication implementations which are used by our framework.

Where existing library functions were not available, we implemented our own operations. While not heavily optimised, these operations utilise the SIMD instructions through the vector intrinsics provided in the Cell toolchain to yield moderate performance.

7.3 Partitioning, Alignment and Padding of Matrices

The small local stores of SPEs, coupled with the triple buffering technique described in subsection 7.2, limit the size of matrices that can be operated on by the SPEs. As mentioned in subsection 7.2, the local store of an SPE provides 256kB of memory which is shared for code and data. The MEU virtual machine program is approximately 26kB in size, leaving 230kB of space for matrices. Triple buffering divides this space further into 3 buffers of size \( \approx 76kB \), each of which should be capable of storing the operands of a single matrix operation. We assume that each matrix operation can have up to two operands, giving each matrix block a maximum size of \( \approx 38kB \). Given that single and double precision floating point numbers occupy 4 and 8 bytes of memory respectively, the maximum number of matrix elements that will fit in the buffer of an SPE is \( S = 9728 \) single precision elements or \( S = 4864 \) double precision elements. However, the partitioning scheme described in section 5 results in blocks that are up to \( \delta \lfloor \sqrt{S} \rfloor \times \delta \lfloor \sqrt{S} \rfloor \) in dimensions, where \( \delta = 4 \) for single precision and \( \delta = 2 \) for double precision (for reasons described below). This results in an effective buffer size for each operand of \( S = 9216 \) for single and \( S = 4864 \) for double precision elements, as stated in section 5.

Each block of a matrix is transferred to the local store of an SPE using a DMA list transfer (a group of individual DMA commands). Each list entry transfers a single row of the matrix block as shown in Figure 15(b) of section 5. In the Cell architecture, DMA commands can only operate on
memory sizes which are multiples of 16 bytes \[24\]. The partitioning scheme described in section 5 ensures that the number of columns and rows in a block is always a multiple of $\delta$. Hence, we choose the divisor to be $\delta = 4$ for single precision and $\delta = 2$ for double precision which guarantees that each row of a block is a multiple of 16 bytes.

Optimal DMA performance is seen when transfer sizes are multiples of 128 bytes \[24\]. The maximum length of a row in a block (using our partitioning scheme) is 384 bytes. Thus blocks that reach this maximum will achieve optimal transfer performance. A value for the divisor could be chosen such that row lengths are always a multiple of 128 bytes ($\delta = 32$ for single precision and $\delta = 16$ for double precision). Unfortunately, this would result in additional overhead in the extra padding of matrices. For example, if a single precision matrix had 4 columns, a value of $\delta = 32$ would force 28 extra columns of padding to ensure that the length of each row is 32 elements.

Block dimensions which are a multiple of 16 bytes are also necessary in order to easily utilise the SIMD operations on the SPEs which operate on 128 bit (16 byte) vectors. For the same reason, the matrix libraries provided for the SPEs require matrices to be of these dimensions.

A further requirement of DMA commands is that memory is aligned to at least a 16 byte boundary \[24\]. The beginning of matrix arrays are aligned to 128 byte boundaries in main memory using the `malloc_align()` function call provided in the Cell SDK. Since the partitioning scheme results in row lengths guaranteed to be a multiple of 16 bytes (as described above), this means that a partitioned block always begins on a 16 byte boundary in memory. Optimal DMA performance is seen when source and destination addresses are aligned to 128 byte boundaries (one cache-line) \[24\], which is again ensured for all blocks with an appropriate choice for the divisor $\delta$.

### 7.4 Execution Control

A fast and efficient execution control mechanism must be used to deliver scheduled operations from the PPE to the SPEs and track the completion of operations. On the PPE, each SPE is modelled as an execution queue. The PPE enqueues operations on to the end of an execution queue if they are to be executed by the SPE represented by that queue. Once an SPE has finished execution of the operation, the PPE can dequeue the operation from the execution queue and register it as complete. At any point in time, the execution queue contains three kinds of operations (Figure 20):

1. **Unexecuted operations**: A number of operations that the PPE has offered to the SPE but which the SPE has not yet acknowledged.
2. **Executing operations**: A number of operations that the SPE has acknowledged and started data transfer or execution of.
3. **Executed operations**: A number of operations which the SPE has finished executing and whose result has been returned to main memory, but which the PPE has not yet acknowledged as complete.

![Figure 20: Matrix execution queue.](image)

In order to enqueue an unexecuted operation, $op_u$, onto the execution queue, two criteria must be met. Firstly, the number of unexecuted operations already in the execution queue must be less than 4.
This is because the mailbox mechanism of each SPE which is used to deliver operations from the PPE to SPEs can only hold up to four 32 bit values. Secondly, all of \( op_u \)'s operands must be available in main memory. This means that if \( op_u \) depends on the results of other operations, these must be fully computed by an SPE and returned to main memory prior to \( op_u \) being added to the execution queue.

Ideally, the scheduler would produce a schedule which includes the exact wall-clock time at which \( op_u \) can begin executing. However, indeterminisms in the architecture can cause variations in the actual execution times of operations, which cannot be anticipated by the scheduler. To ensure that all operands of an operation are available prior to its execution, we introduce a guard count value to each operation. The guard count is an integer value which represents the number of operands of an operation which have not yet been computed. Before the execution of any operation, every operation’s guard count is equal to the total number of operands that the operation has. For example, a matrix addition operation has 2 operands (the matrices to be added) so its guard count is initially 2. When an executed operation, \( op_e \), has been acknowledged by the PPE as being complete, the guard count of all the operations which depend on \( op_e \) are decremented by 1. This means that it is safe to place an unexecuted operation on the execution queue when its guard count is zero.

In order to signal the PPE to the completion of an operation, the SPE uses DMA commands to write a counter value to a pre-determined location in main memory. The PPE polls this memory location and when it changes, knows that an operation is complete and it is safe to dequeue the instruction from the execution queue. This interprocessor communication technique is chosen over other techniques because it provides better performance \[24\]. It is important that the PPE is notified of completed operations as quickly as possible, as this allows dependant operations to begin execution sooner.

The execution control process consists of the PPE examining each of the execution queues in turn. It enqueues as many unexecuted operations as possible on to the execution queue, in the order that they are specified in the processors schedule. The PPE is notified of these operations through the mailbox mechanism. The PPE then dequeues any complete operations from the execution queue, decrementing the guard count of dependant operations by 1. This process continues until all operations in the current schedule have completed execution.

### 8 Implementation Details and Tuning

The implementation of our framework has \( \approx 7000 \) lines of C and C++ code. The code consists of two separately compiled programs: one written for the PPE and one written for the SPEs. The SPE program consists of the implementation of the Matrix Execution Units (MEUs) as described in section 7. Although this component is smaller (in lines of code) than the PPE program, it is crucial that it achieves high performance and does not consume a significant portion of the small local store of the SPEs. Thus, it is written in C code. The PPE program consists of the implementation of the remainder of the framework, i.e. the Octave extension, the lowerer, scheduler and execution control. It is written in C++ and performance is mostly not as critical as the SPE program.

The development of the software was performed on a conventional x86-based processor running Fedora Core 9 and the latest version of the Cell SDK (3.1.0) \[24\]. The code for the Cell can be compiled on this standard architecture by using cross-compilation tools included in the Cell SDK. Testing was initially performed with the Sony Playstation 3 games console that runs a Cell Broadband Engine processor. However, the Playstation 3 is limited to 256MB of memory which impeded development and limited the size of test cases that could be run. Later in the project we obtained access to a BladeCenter QS22 server, as described in subsection 9.1. Although the QS22 has the same Cell-based processor as the Playstation 3, it offers 32GB of memory allowing large test case to be executed without causing memory swapping and distorting time measurements.

Here we describe implementation issues with regard to (1) debugging and testing our framework, (2) details of the implementation of the Octave extension, and (3) details of the performance tuning we performed.
8.1 Debugging and Testing

Debugging on the Cell processor is difficult for several reasons. Firstly, concurrent programming on any architecture is challenging and the Cell processor is no exception. Our framework has several threads of execution running simultaneously on the PPE, as well as each of the 8 SPEs running a separate thread. There is no easy method for finding race conditions or deadlocks between these threads. Secondly, standard memory debugging tools such as Valgrind cannot be used on the Cell architecture because DMA memory transfers between the SPEs and main memory interfere with their operation. This makes debugging memory issues difficult, even when confined to the PPE. Furthermore, the low-level programming model employed on the Cell processor makes programming a highly error-prone exercise. This is particularly true when programming for the SPEs.

We used a few techniques to alleviate this issue. Firstly, we wrote our C++ framework in a platform independent way. Only one component of the framework, the computation engine, contained hardware-specific code (see the system overview in section 3). We wrote versions of the computation engine for the Cell processor, but also for a standard x86 architecture. The x86 implementation is sequential and was not designed for performance but instead for debugging purposes. It contains very simple, naïve implementations of matrix operations. This allowed us to test our framework on a more conventional architecture before having to deal with the idiosyncrasies of the Cell architecture. We were able to use standard debugging tools, including Valgrind, on the x86 to eliminate errors in all components of our framework, besides the Cell computation engine. If we experienced anomalies when running the framework with the Cell architecture that were not experience on the x86 architecture, we could narrow the bug to the Cell implementation of the computation engine. This dual-platform testing technique greatly reduced the time spent on debugging.

As a precursor to the integration of our framework with Octave, we developed a more simple interface to our framework. This was in the form of a simple trace language which allowed a user to enter a trace of matrix operations to be executed. The language does not have any control-flow mechanisms (like loops or if-statements). An example of a script for this language is shown in Figure 21. This script simply defines two matrices and adds them together. This interface allowed us to concentrate on the implementation of the back-end components of our framework prior to the integration with Octave. However, this also served as another debugging tool, allowing us to determine whether there was an issue with the Octave extension or with a different component of the framework. An automatic test generator was written to generate random test traces for this front end based on a number of parameters such as trace length and matrix dimensions. This further assisted us in debugging and testing our framework, particularly with larger traces which would take too long to write by hand.

Debugging tools provided by the Cell SDK were also used to assist in finding bugs in Cell specific components. We used the GNU debugger (gdb) designed for the Cell processor that facilitates tracing the execution of the program on both the PPE and SPEs simultaneously.

Regression testing was employed throughout the project to reduce the chance of introducing bugs and also to reduce debugging time. For every matrix operation that was implemented in the framework, several tests were written. Initially these tests were written for our simple trace language interface, but later they were crafted as Octave scripts. Octave contains several functions designed specifically for testing purposes, such as the assert function. An example of a simple Octave script for testing a matrix addition operation is included in Figure 22. The debugging and testing techniques as mentioned above, gave us a high level of confidence in the quality and correctness of our implementation.

8.2 Octave Extension

GNU Octave [17] is a scripting language and programming environment designed for scientists and engineers. The programming environment consists of a command line interface in which users execute scripts or individual statements interactively. As Octave is an open-source project, its C++ source
code is freely available for download and compilation.

Octave has been designed in a way that facilitates the development of extensions without the need to re-compile the Octave interpreter. This is achieved by using shared libraries for extensions, which can be loaded dynamically by the Octave interpreter at run-time. Extensions are called Oct-Files and can be written in C/C++ or FORTRAN. They are compiled by a utility program included with Octave called `mkoctfile`. This utility is a wrapper for the `gcc` compiler which ensures that the necessary compilation flags and linker options are provided.

Octave permits extensions in the form of user-defined functions as well as user-defined data types. An example of a simple user-defined function for printing a message to the interpreter console is shown in Figure 23. The Oct-File is pure C++ code however numerous macros are provided by the Octave system in order to simplify access to data types and the declaration of functions.

We extended Octave by introducing a user-defined data type called `p_matrix`. This is a custom matrix data type which can be used as a replacement for the default Octave matrix data type. In order to cast standard Octave matrices to the `p_matrix` type, users call a function called `p_matrix()` with the standard matrix as an argument. For example, line 1 of the Octave code shown in Figure 24

---

Figure 21: A script written for our simple trace language interface. It declares two matrices $A$ and $B$ and adds them together, storing the result in $C$.

```cpp
#define matrix A
A = [ 1, 2, 3, 4;
     5, 6, 7, 8;
     9, 10, 11, 12;
    13, 14, 15, 16];

#define matrix B
B = [ 1, 2, 3, 4;
     5, 6, 7, 8;
     9, 10, 11, 12;
    13, 14, 15, 16];

#define add both matrices
C = MADD(A, B);
```

Figure 22: An Octave test script for matrix addition. The `assert` function is built into Octave and checks whether a condition is satisfied or that two results match. The `has_result` function is provided by our framework to determine whether a result has been computed by lazy evaluation or whether its execution is still outstanding.

```octave
% Matrix addition
x = p_matrix([1, 2; 3, 4]);
y = p_matrix([1, 2; 3, 4]);
z = x + y;
assert(!has_result(z));
assert(z, p_matrix([2, 4; 6, 8]));
assert(has_result(z));
```
```cpp
#include <octave/oct.h>

DEFUN_DLD (helloworld, args, nargout, ) {
  int nargin = args.length();
  octave_stdout << "Hello\nWorld\nhas\n" << nargin
    << "input\narguments\nand\n" << nargout << ".\n"
  return octave_value_list();
}
```

Figure 23: An example of a user-defined function written as an extension for Octave. Although this is pure C++ code, macros and variables are provided by Octave to assist with proper integration of the extension. Here, a function called `helloworld` is defined which merely prints the number of input and output arguments to the interpreter console. The `DEFUN_DLD` macro inserts the appropriate C++ code to allow `helloworld` to be called from inside of Octave.

```
1 A = [10 20 30; 40 50 70; 70 80 90];
2 A = p_matrix(A);
3 B = A + 5;
4 C = B;
5 disp(B(5));
```

Figure 24: Simple Octave script which uses our framework.

creates a $3 \times 3$ standard Octave matrix with the given elements. A user could then call the `p_matrix()` function, as shown in line 2, to convert the matrix to our custom data type. This is the only user intervention necessary to use our framework. By replacing the standard Octave matrix data type with our own, no user intervention would be required but this would necessitate modifying and re-compiling the Octave source code which we wanted to avoid.

User defined data types in Octave are implemented as C++ classes which inherit from the `octave::base_value` class [41]. Operators such as `+` and `*`, as well as built-in functions such as `sin` and `round` can be overloaded to behave as required for custom data types.

Our `p_matrix` data type is implemented in a C++ class called `octave_p_matrix`. For a complete list of the Octave operators and functions that were implemented for the `p_matrix` data type refer to Appendix [C]. When a variable of type `p_matrix` is created in Octave, a corresponding C++ `octave_p_matrix` object is generated. In line 2 of the example code in Figure 24, the `p_matrix()` function is called to cast the matrix $A$ to our custom data type. Thus, an `octave_p_matrix` object is generated by Octave. Our framework also constructs a `ConstantMatrix` object to represent $A$. This merely has a pointer to the memory location of the matrix $A$. The `ConstantMatrix` object is added to the data dependence graph in our framework. This is illustrated in Figure 25.

When a user performs an operation on a `p_matrix` variable, the overloaded operators for the type `p_matrix` defer execution of the operation. For example, when a user executes line 3 of the Octave code in Figure 24, the value of $B$ is not computed immediately. Instead, we have overloaded the `+` operator for `p_matrix` to create a new `ScalarAddition` C++ object in our framework to represent the addition of $5$ to $B$. The `ScalarAddition` object contains a pointer to the `ConstantMatrix` to denote that $A$ is the operand of $B$, as shown in Figure 25. This addition operation is also added to the data dependence graph in our framework, with a data dependency on the matrix, $A$. Another `octave_p_matrix` object is generated by Octave to represent the $B$ variable. This contains a pointer to the `ScalarAddition` object in our framework, as depicted in Figure 25.
Figure 25: The memory layout of C++ objects in Octave and our framework. This shows the state of memory after executing line 4 of the Octave script in Figure 24. Every time a `p_matrix` variable is created in Octave, an object of the type `octave_p_matrix` is constructed. An object is also constructed by our framework to represent that operation. When matrix `A` is cast to the `p_matrix` type in line 2, a `ConstantMatrix` object is created in our framework. This merely has a pointer to the memory location of the matrix `A`. The operation is also added to a data dependence graph on the fly. When the variable `B` is declared in line 3, another `octave_p_matrix` object is constructed by Octave. The operation to compute `B` is not executed immediately. Instead, a `ScalarAddition` object is created by our framework representing the addition of the scalar value 5 to the matrix `A`. This contains a pointer to the matrix `A`, the operand of the operation. `B` is added to the data dependence graph, with a dependency on `A`. In line 4, `B` is assigned to `C`. Copy-on-write is used to prevent the execution of `B` being forced and also to prevent making an unneeded deep copy of `B`. Instead, `C` has a pointer to the same `ScalarAddition` object which computes its result.

To improve performance of the framework, we employ copy-on-write semantics. If a matrix of the type `p_matrix` is assigned to another variable, no deep copy of that matrix is made. For example, in line 4 of Figure 24, the variable `B` is assigned to the variable `C`. Producing a deep copy of matrix `B` here would negatively impact performance in two ways. Firstly, we do not need to make a copy of `B` because `B` and `C` point to the exact same matrix value. Thus, making a copy would introduce unnecessary overhead in memory copying operations. Secondly, if we were to make a deep copy of `B` here, it would require the the value of `B` be evaluated. Because of lazy evaluation, `B` has not been computed at this point. Forcing the computation of `B` would unnecessarily shorten the length of the trace of matrix operations and reduce the amount of parallelism to be exploited by our framework. Instead of making a deep copy, the `octave_p_matrix` object representing `C` is assigned a reference to the same `ScalarAddition` operation as `B` in our framework. This is shown in Figure 25. If the matrices `B` or `C` are modified, for example by changing the value of an element, then the operation is forced to be computed and a deep copy of the matrix is made.

Our framework continues to accumulate matrix operations until the result of one of those operations is needed. Results of operations are typically needed when a user wants to print a matrix to the screen or access particular elements of a matrix. Thus, we have overloaded the subscript operators on matrices, as well as the `disp` function such that these trigger the execution of outstanding operations in the framework. For example, in the last line of code in Figure 24, the the 5’th element of `B` is printed to the screen. A call to our overloaded subscript operator is made. This first determines whether the `ScalarAddition` operation to yield the value of `B` has been computed yet. In this case it has not so execution of the outstanding trace of operations in the framework is initiated, and continues as described in section 3. Once the result of `B` has been computed, it is made available to the Octave
script and execution of the script can continue. This technique gives our framework its lazy evaluation semantics.

Octave automatically handles garbage collection of its variables but this must be tied to our framework so we can do the appropriate de-allocation of resources. When a `p_matrix` variable is garbage collected in Octave, the destructor of the corresponding `octave_p_matrix` object is called. This will de-allocate the matrix operation object in our framework which is tied to that variable. In the example code in Figure 24 garbage collection of the variable `B` will result in the de-allocation of the `ScalarAddition` operation tied to `B`. However, we must be careful not to garbage collect operations or matrices that are still needed. If the matrix `A` was garbage collected prior to the evaluation of `B` then our framework would not de-allocate `B`. This could occur, for example, if the code `clear A` was called before line 5 of Figure 24. Our framework recognises that the matrix `A` is still needed to compute `B`. Thus, it will not be de-allocated until execution of the next trace is complete.

8.3 Performance Tuning

It was necessary to optimise our framework to obtain the high performance reported in the experimental subsection (see section 9). In order to identify bottlenecks in the framework, we profiled the code using the performance counters available in the Cell processor [24]. These are highly accurate timers which operate at the time-base frequency of the architecture implementation (26664325Hz for the BladeCenter QS22 implementation). All components of the framework were instrumented with profiling code in both the PPE and SPE programs. A breakdown of time spent in each component of the framework was obtained through use of this profiling and is shown in section 9. A compile time option was used in our framework to enable or disable profiling as required.

A large bottleneck we discovered through profiling was in the time spent making virtual function calls. For example, matrix operation classes in our framework are implemented with an inheritance hierarchy. There is a virtual method implemented by each operation for obtaining the estimated data fetch, execution and write back times for that operation. The heuristic scheduler made many calls to this virtual function for each operation in order to compute the schedule. We were able to reduce the number of calls to once for each operation in a trace which resulted in an improvement in the time spent scheduling operations. This change alone led to a performance improvement of 20% in the scheduling process.

Using appropriate data structures had a significant impact on performance. Profiling allowed us to determine that the scheduling of operations was initially dominating the execution time of a trace for some benchmarks. That is, scheduling was taking longer than the actual execution time. After investigating this, we found that it was due to the use of an array for storing ready operations in the scheduler. This array has to be iterated over to find the earliest operation that could be scheduled and this has to be done for every operation in the schedule (see section 6). By using a min-heap priority queue instead of the array, we were able to reduce the $O(n^2)$ complexity of this portion of the algorithm to $O(n \log n)$. This had a large impact when there were many operations in the trace and reduced the average time spent scheduling operations to less than 15% of the total execution time of the trace.

Another area where we made improvements was with memory fragmentation. Originally, we just used the standard `malloc` function to allocate memory for each matrix operation object in our framework. However, as can be seen in section 9, benchmark traces could contain over 600000 matrix operations. This led to high fragmentation of memory and had a negative impact on performance. We used memory pooling to pre-allocate large contiguous blocks of memory at once. From these large blocks, matrix operation objects were allocated.

Memory pooling was also used to improve the performance of the allocation of matrices. During the execution of matrix multiplication operations, many temporary matrices are used to perform the summation of blocks (described in section 5). Allocation of these temporary matrices degraded performance. Instead of allocating each matrix with a separate `malloc` call, we again used a pool of contiguous memory. Not only did this improve the time spent allocating matrices, but also deallocating
them because returning memory to the pool is a very cheap operation.

Finally, the execution control mechanism (described in section 7) was another area where improvements were made. By instrumenting the SPEs with profiling code, we were able to compute the idle time spent by each SPE. By simplifying the execution control loop, which delivers operations to the SPEs, it was able to service them more quickly. This resulted in up to a 30% decrease in the idle time of SPEs for some benchmarks.

9 Experimental Results

In the experimental evaluation of our framework we focused on the performance improvements that users can expect from running Octave application code with our framework running on the Cell processor. We compared our framework with three other typical system configurations:

1. A default installation of Octave on a contemporary Intel Core2 Quad processor.
2. A default installation of Octave on the Cell processor.
3. A default installation of MATLAB on an Intel Core2 Quad processor.

In addition to these comparisons, we also investigated our approach with respect to:

1. Its scalability to a larger number of SPEs.
2. The accuracy of our time models for estimating the execution time of matrix operations.
3. The quality of schedules derived by our heuristic scheduling algorithm.
4. The extent to which our framework was able to utilise the Cell SPEs.

9.1 Experimental Setup and Benchmark Suite

For the experiments, a BladeCenter QS22 server was used to run our framework. The details of its configuration are summarised in Table 1. The BladeCenter QS22 is comprised of two IBM PowerXCell 8i processors (3.2GHz/1MB L2) and has 32GB DDR2 memory. We used only one out of the two Cell processors available on the BladeCenter due to non-uniform-memory-access (NUMA) issues which complicated use of the second processor (though our framework could be configured to work with an arbitrary number of SPEs). We utilise all 8 of the SPEs as well as the PPE available in the Cell processor (refer to section 7).

Originally, we intended to perform the experiments on the Cell processor found in the Playstation 3 games console. Despite both the BladeCenter QS22 and Playstation 3 being based on the same Cell Broadband Engine architecture with the same capabilities, the Playstation 3 has only \( \approx 200\)MB of accessible RAM. Thus the BladeCenter QS22 was used to alleviate this limitation on the size of benchmark inputs.

| Blade model: | IBM BladeCenter QS22 |
|-------------|----------------------|
| Number Cell processors: | 1 (8 SPEs, 1 PPE) |
| Memory: | 32GB DDR2 |
| Linux distribution: | Red Hat EL Server 5.4 |
| Linux kernel: | 2.6.18 |
| IBM Cell SDK: | 3.1.0 |
| GCC: | 4.1.1 |
| GCC optimisation flags: | -O3 |
| Octave: | 3.0.3 |

Table 1: Experimental setup.
The following benchmarks were selected to evaluate the performance of our framework:

1. **dft** Computation of the Discrete Fourier Transform (DFT) of a series of signals.
2. **synth** A synthetically constructed benchmark with many small, independent matrix multiplications.
3. **hill** Encryption and decryption using Hill ciphers.
4. **hits** Computation of the Hyperlink-Induced Topic Search (HITS) algorithm for estimating the importance of a web-page.
5. **kmeans** Computation of the k-means clustering of a 2D point set.
6. **leontief** Computation of a Leontief input-output model, used to predict performance of economies.
7. **markov** Computation of a Markov chain.
8. **neural** Training of a single-layer neural network.
9. **reachability** Computation of the reachability matrix of a graph.

These benchmarks were chosen to represent commonly used kernel programs from scientific, engineering and computer science domains.

### 9.2 Speedups and Scalability

We first compared our framework on the Cell processor with a default Octave installation on an Intel Core2 Quad Q9550 2.83GHz processor. We chose the Intel Core2 Quad architecture for comparison because it represents a contemporary and frequently used microprocessing architecture which is more modern than the Cell processor. The standard Octave installation on the Intel Core2 Quad uses single-threaded ATLAS BLAS libraries for some matrix operations \[55\]. These libraries utilise the SSE3 multimedia extensions of the processor for performance.

As depicted in Figure 26, our framework achieves speedups of up to a factor of 12 times over standard Octave on the Intel Core2 Quad. For most benchmarks we achieve speedups of over 7. The k-means clustering (“kmeans”) and neural network (“neural”) benchmarks achieve lower speedups. These lower speedups can be attributed to two factors. Firstly, the time models for some matrix operations in these benchmarks may not be accurate. There is further evidence for this provided in subsection 9.3. Inaccurate time models would lead to inaccurate estimation of the execution times of
operations which could result in degraded schedules. Fine-tuning of time-models could see improved performance. The second reason for lower speedups is that these benchmarks contain many matrix operations on vectors. The partitioning scheme employed in our framework results in sub-matrices which are very small, as described in section 5. This results in under-utilisation of the SPEs in the Cell processor and poorer performance. This could be improved by modifying the partitioning scheme for vector operations.

The second configuration that we compared our framework to was a default installation of Octave on the BladeCenter QS22. That is, we compared the runtime of the benchmarks BladeCenter server with and without our extension switched on in Octave. It should be noted that the default Octave installation utilises hardware-specific BLAS libraries which are provided with the IBM Cell SDK. These libraries are highly optimised for the Cell architecture \cite{46} and can utilise both Cell processors available in the QS22 (a total of 16 SPEs).

With this configuration we achieved the speedups depicted in Figure 27. It can be seen that our framework achieves a speedup with all benchmarks (though some are lower than others). One may expect our framework to perform worse than the default installation which uses highly optimised BLAS libraries. However, the BLAS libraries do not accelerate all Octave functions. For example, computation of the sin function of elements in a matrix is not a standard BLAS operation. As a result, the PPE is used to compute such functions, which can be relatively slow. Our framework can utilise the SPEs for all operations that we have implemented and thus see an improvement in performance for these operations.

Furthermore, the Cell BLAS libraries exploit only the data parallelism of matrix operations whereas our framework also exploits instruction level parallelism. This is emphasised with the “synth” benchmark. This benchmark has a large amount of instruction level parallelism because it contains many matrix multiplications which are independent of each other. However, because each matrix multiplication operates only on small matrices, there is no benefit in dividing an operation among SPEs. This causes BLAS to confine computation of the operations to the PPE only. Each operation is computed sequentially by the default installation of Octave leading to a very high runtime with this benchmark. In contrast, our framework recognises that although it is not worth dividing up individual operations among SPEs, the multiplications can be executed concurrently. This allows our framework to obtain high utilisation of the Cell processor and explains the speedup of over 200 times for this benchmark.

The seemingly erratic speedups across benchmarks in this configuration (i.e. speedups vary from 1.1 times to 214 times) can be explained with the same reasoning. That is, the speedup of our framework depends heavily on how much instruction level parallelism is available in a benchmark, as well as the proportion of matrix operations in the benchmark which can be executed by the optimised BLAS libraries in the default Octave installation.
In a variation of this configuration, we forced the default installation of Octave to utilise only the PPE of the Cell processor (and not the SPEs). Our framework achieved speedups of several hundred times in this case.

The third and final configuration that we compared our framework with was a default installation of MATLAB running on an Intel Core2 Quad processor. Since MATLAB is a commercial product, it is likely to be more optimised than Octave, its open-source equivalent. However, we wanted to compare our framework with MATLAB because it is more frequently used than Octave and represents the current state of the art in high-level scientific computing. MATLAB and Octave have very similar syntax and the benchmark programs required no changes for them to run on MATLAB.

In this experiment, we used the latest version of MATLAB (version 7.8) which was found to utilise all four cores of the Intel Core2 Quad processor in the execution of matrix operations. Again, we found we were able to achieve speedups for all benchmarks and up to 8 times in the best case, as shown in Figure 28. However, these speedups were lower than those from the first configuration which confirmed our suspicions that MATLAB provides better performance than Octave.

Again, the speedups of our framework can be attributed to the exploitation of instruction level parallelism and the scheduling of operations among the parallel processing elements of the Cell architecture to improve utilisation. Neither of these are currently performed by Octave or MATLAB on any architecture.

In the next set of experiments, we investigated how the performance of our framework scaled with the number of SPEs utilised on the Cell processor. The results for each of our benchmarks are shown in Figure 29. As can be seen, performance deviates from linear speedup depicted by the dotted line. This can be explained by considering the proportion of sequential and parallelisable work for each benchmark, as shown in Table 2. These values were estimated by measuring the time taken by each component of the framework when utilising only 1 SPE. The proportion of time spent in lowering and scheduling on the PPE could not be parallelised. However the proportion of time spent executing matrix operations on the SPEs could be parallelised by increasing the number of SPEs used.

When the amount of sequential work in each benchmark is taken into consideration, the theoretically achievable speedup with 8 processors can be computed using Amdahl’s law [20]. This states that the maximum speedup achievable is given by $\frac{1}{(1-P)+\frac{P}{N}}$ where $P$ is the parallelisable portion of the program as a fraction and $N$ is the number of processors used (8 in this case). These values for each benchmark, along with the observed speedups are also shown in Table 2. It follows that most benchmarks do not deviate significantly from the maximum theoretical speedup, indicating that to obtain better performance it is necessary to further optimise the sequential portion of the workload, i.e., the lowering and scheduling components.

Any deviations between the maximum theoretical speedups and the observed speedups shown in
Figure 29: Performance scalability of the proposed framework on the IBM BladeCenter QS22. This shows the speedup achieved for each benchmark by varying the number of SPE cores utilised from 1 to 8.

Table 2 are due to deficiencies that arise as the number of processors increases. For example, the Cell’s EIB communications bus will have to cope with a higher load when the number of SPEs increases. The PPE will also face increasing workloads as it has to service more SPEs. Further optimisations to the execution control mechanisms could help reduce this observed deviation from the maximum theoretical speedup by allowing SPEs to be serviced more quickly.

9.3 Time Model

For computing an effective schedule, it is important that the estimated execution times of operations are accurate (see subsection 9.3 of the scheduling section). To verify the accuracy of our time models, we compared the estimated makespan of a trace with the real, run-time makespan of the same trace. The estimated makespan is calculated by the heuristic scheduler using the time models of operations. The real makespan of the trace was measured by timing the execution phase of operations in the trace using the hardware performance counters on the Cell processor. We did this for many different data dependence graphs and computed the deviation between the two makespans.

A histogram of these deviations is shown in Figure 30. The majority of deviations are close to 0%, indicating accurate time modelling. The median of the deviations is 1.3% and the skewness is -0.54, meaning that the distribution is slightly skewed towards the right. The skewness of the distribution is caused by the overhead of the execution control mechanism described in section 7. This mechanism is responsible for synchronising the execution of matrix operations and the overheads involved are not taken into account in the estimated makespan of a trace. Thus the heuristic scheduler underestimates the makespan accordingly.

Note that Figure 30 includes deviations for data dependence graphs generated by all benchmarks except for benchmarks “neural” and “kmeans”. We excluded those benchmarks because there was found to be large deviations between the estimated and actual makespans (up to 300%). This provides evidence that the time models for some operations in these benchmarks are inaccurate and need further fine tuning.

9.4 Scheduling

As a further experiment, we wanted to investigate the quality of our heuristic scheduler by comparing the makespan of the heuristic with the makespan of the optimal solution. The number of operations
Table 2: Sequential and parallelisable portions of benchmark programs are measured by running the benchmarks on a single SPE. The sequential portion is the time that is spent lowering and scheduling operations, allocating memory and performing cleanup. The parallelisable portion is the time spent in the actual execution of operations on the SPEs which can be reduced by utilising more SPEs. The maximum theoretical speedups are computed for 8 SPEs using Amdahl’s law. This states that the maximum speedup achievable is given by \( \frac{1}{(1-P) + \frac{P}{N}} \) where \( P \) is the parallelisable portion of the program as a fraction and \( N \) is the number of processors used (8 in this case). The maximum speedup that each benchmark was observed to achieve on our framework with 8 SPEs is also shown for comparison.

Figure 30: The distribution of deviations between the estimated makespan of the schedule produced by our heuristic algorithm and the observed makespan measured using performance counters on the Cell. Deviations are computed for over 100 traces produced by all benchmarks except “kmeans” and “neural”. Most deviations are close to 0% indicating that our time models for matrix operations are accurate.
Figure 31: Time taken to schedule and execute matrix operations using two scheduling algorithms: our heuristic algorithm and a naïve approach which balances the number of operations on each SPE. Note that although the time spent scheduling operations using our heuristic is much longer than the time spent scheduling using the naïve approach, the total execution time is lower for most benchmarks.

and dependencies in each of the benchmark programs used to compute the speedups of our framework (see subsection 9.2) are shown in Table 3. The heuristic scheduling algorithm was able to compute schedules for these problem sizes in less than a second in all cases. However, these problem sizes were too large to compute an optimal solution using the mathematical program described in subsection 6.2. Thus, we had to reduce the input size of these benchmarks greatly, as listed in Table 4. CPLEX 10.0 [26] was able to solve these smaller problem sizes within an hour.

In all cases, the makespan of the heuristic schedule was within 1% of the makespan of the optimal schedule. However, where the mathematical program took up to 40 minutes to produce a solution, the time to compute the heuristic schedule was too small to be measured (for these problem sizes). This indicates that the heuristic is efficient and effective. Unfortunately, these small problem size instances may not offer enough degrees of freedom to see a large deviation between the heuristic and the optimal solution. We were not able to increase the problem sizes without making the problems intractable to be solved by CPLEX.

Another method we used to evaluate the heuristic scheduling algorithm was to compare it to other heuristic scheduling techniques. Figure 31 shows the time spent scheduling and executing matrix operations for two different scheduling techniques. The first is our heuristic scheduling algorithm and the second is a very naïve scheduling algorithm (simple scheduling) which merely distributes operations evenly and in a topological order among the SPEs. This is done in a round-robin fashion such that each SPE has roughly an equal number of operations to execute. It can be seen that the simple scheduling algorithm spends almost no time scheduling operations. However, the time taken to execute the operations exceeds the sum of the scheduling and execution times using our heuristic algorithm (for most benchmarks). This shows that our scheduling algorithm has a worthwhile impact on the overall execution time of a trace, even though it does incur noticeable overhead. By reducing the overheads of scheduling with further optimisations the benefits of our scheduling algorithm would be even more substantial. It also shows that, in general, scheduling is important in obtaining good performance on modern accelerator architecture. Common, naïve approaches, such as the one described, can lead to poor performance.

Two of the benchmarks execute more quickly with the naïve scheduling as shown in Figure 31. The first is the “synth” benchmark. This is an expected result for this benchmark because every matrix operation in the benchmark is independent and the same size. Hence, as long as operations are distributed evenly among processors, an optimal schedule will be produced. So both scheduling algorithms produce the same schedule, however the naïve algorithm does so much more quickly and a
|                | Original |     |     |     | Lowered |     |     |
|----------------|----------|-----|-----|-----|---------|-----|-----|
|                | Nodes    | Edges | Nodes | Edges |         | Nodes | Edges |
| dft            | 9        | 9    | 72384| 141284|
| synth          | 60003    | 120000| 60003| 120000|
| hill           | 9        | 8    | 73684| 140608|
| hits           | 50       | 98   | 319950| 639450|
| kmeans         | 34       | 23   | 137764| 252743|
| leontief       | 204      | 402  | 129152| 254720|
| markov         | 52       | 100  | 93400| 186580|
| neural         | 10012    | 12489| 231784| 410151|
| reachability   | 960      | 1912 | 119600| 239000|

Table 3: Number of operations (nodes) and dependencies (edges) in the original and lowered dependence graphs for each of the benchmarks used in the experiments described in subsection 9.2.

|                | Nodes | Edges | Time (mins:secs) |
|----------------|-------|-------|------------------|
| dft            | 34    | 44    | 0:02             |
| synth          | 17    | 32    | 1:49             |
| hill           | 28    | 32    | 0:28             |
| hits           | 40    | 72    | 0:06             |
| kmeans         | 79    | 64    | < 0:01           |
| leontief       | 46    | 88    | 0:06             |
| markov         | 48    | 84    | 2:47             |
| neural         | 38    | 35    | 0:04             |
| reachability   | 53    | 104   | 41:19            |

Table 4: Number of operations (nodes) and dependencies (edges) in the largest problem sizes that the integer linear programming solver could compute a solution for within 1 hour. Also included is the time taken by the solver to produce a solution.
corresponding drop in the execution time is observed.

The “neural” benchmark also runs more quickly with the naïve algorithm. This provides further
evidence to support the suggestion that some operations in the benchmark have a time model that
is inaccurate. This would cause a degraded schedule to be produced by our heuristic algorithm
and increase the execution time of the operations as a result.

An online scheduling approach was also experimented with, in which a pool of operations which
were ready to be executed was kept. SPEs selected an operation from this pool arbitrarily. Similar
results to the naïve scheduling algorithm were observed.

9.5 Utilisation of Parallel Execution Elements

In the remaining set of experiments we investigated the extent to which our framework utilised
the parallel execution elements (SPEs) of the Cell. Figure 32 shows a breakdown of execution time spent
for the different phases of our framework. These numbers are averaged across all benchmarks when
run with 8 SPEs. It follows that the majority of time (more than 75%) is spent executing matrix
operations on Cell SPEs. The remaining 24.41% is mainly spent on scheduling (11.4%) and lowering
of matrices into blocks (9.44%). Only 0.83% of time is spent on matrix allocation, and 2.74% on
deallocation and other cleanup.

In another experiment, we measured the time that the SPEs spent idle during the execution phase
of the framework. The results of this experiment are shown in Figure 33. The idle time is composed
of the time the SPEs spent waiting for tasks to be offered by the PPE (“Task”) and the time spent
waiting for the completion of DMA transfers (“DMA”). Again these numbers were determined for
8 SPEs. It follows from Figure 33 that with the “synth” benchmark we approached the Cell’s peak
performance, which is manifested in idle times close to zero.

The remaining benchmarks incur idle times as a result of three main factors. Firstly, operations
such as matrix addition are memory-bound (rather than computationally bound). Although our triple
buffering technique overlaps computation with communication, memory-bound operations can still
induce DMA waiting times because memory transfers for these operations are more expensive than
computation of the result.

The second reason for idle times is that certain traces may contain a large number of data depend-
cyencies which reduce the amount of instruction level parallelism for execution. This can result in idle
time spent waiting for the operands of an operation to become available. Thirdly, if operations are
computed too quickly by the SPEs, the execution control mechanism may not be able to service them.
Figure 33: The percentage of time that SPEs spent idle during each benchmark. 8 SPEs were used for this experiment. Idle time is broken into two components: “DMA” idle time is the time spent waiting for the completion of DMA transfers and “Task” time is the time spent waiting for an operation to be sent by the PPE. The “synth” benchmark exhibits close to 0% idle time and reaches close to the peak performance of the Cell processor.

Figure 34: Speedups achieved by overlapping the scheduling, lowering and execution phases of a trace, versus sequential execution of those phases.

fast enough, resulting in time spent waiting for a task to be offered by the PPE. The impact of this could be reduced by aggregating several operations into a single operation which takes a longer time to execute.

The final experiment we measured the impact of overlapping scheduling of operations, lowering of operations and execution of operations. As described in section 3 if the size of a trace reaches a maximum threshold value, execution of the Octave interpreter will continue while the trace is executed. Scheduling of the trace, lowering of the trace and operation execution are executed in separate threads. Thus it is possible for multiple traces to be in execution concurrently. Figure 34 shows the speedup in the execution time of each benchmark when these separate threads are used versus each of the stages being executed in sequence. Most benchmarks achieve a speedup when compared with sequential execution. Those that do not, such as “kmeans” and “neural” may not produce traces that are large enough to allow any extra parallelism to be exploited and instead, the overhead of using several threads for execution causes a slowdown. Fine tuning of the threshold at which execution of a trace is triggered may result in further speedups from using this technique.
10 Conclusion

In this work we developed a new framework for fully utilising the performance of modern accelerator architectures in the execution of matrix languages. We provide an implementation of the framework for executing Octave programs on the Cell Broadband Engine architecture.

The framework is very easy to use, with only minor changes needed to existing Octave code in order to take advantage of the parallel architecture. As opposed to existing systems, which typically only take advantage of the data parallelism of matrix operations, our framework additionally exploits instruction level parallelism, pipeline parallelism and task parallelism to obtain better performance.

Lazy evaluation is used to extract a trace of matrix operations from a program at run-time. The data dependencies of operations in a trace are examined to elicit instruction level parallelism. A novel partitioning scheme, called lowering, is used to divide operations on matrices in a way that maximises the parallelism available in a trace. A new heuristic scheduling algorithm is then used to schedule operations among the processing elements in a way that improves their utilisation and reduces the execution time of a trace. In order to produce an accurate schedule, we employ time modelling of matrix operations to estimate their execution times.

We performed an extensive evaluation of our framework with positive results. Octave benchmarks executing on our framework for the Cell Broadband Engine architecture are up to a factor of 12 faster than execution on standard Octave on more recent and expensive Intel Core2 Quad processors. Our framework also out-performed an out-of-the box installation of Octave running on the Cell processor, as well as an installation of MATLAB running on an Intel Core2 Quad processor.

We also evaluated a new heuristic scheduling algorithm by comparing the schedules it produced with the optimal schedules produced by a mathematical program we developed. We found that the makespan of the optimal schedule deviated no more than 1% from the makespan of the schedule produced by the heuristic algorithm, for all computable problem sizes.

Further evaluation of our framework revealed that partitioning and scheduling of operations did not incur a significant amount of overhead and one of our benchmark programs was able to reach the peak performance obtainable on the Cell Broadband Engine Architecture.

Though it is widely believed that automatic parallelisation techniques for imperative programs are infeasible, we showed that automating the parallelisation of sequential matrix language programs is achievable for modern accelerator architectures. Speedups of several magnitudes are possible when various kinds of parallelism, including instruction level, data, pipeline and task parallelism, are exploited. We showed that the main contributors to these speedups are instruction level and data parallelism that are obtained through novel lazy evaluation and lowering techniques.

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A Integer Linear Programming Model

\[
\begin{align*}
\text{min} & \quad z \\
\text{s.t.} & \quad \sum_{i=1}^{n} x_{1i} \leq p \\
& \quad x_{i1} = 0 \quad i \in I \\
& \quad \sum_{j=1}^{n} x_{ij} \leq 1 \quad i \in I \setminus \{1\} \\
& \quad \sum_{j=1}^{n} x_{ji} = 1 \quad i \in I \setminus \{1\} \\
& \quad x_{ii} = 0 \quad i \in I \setminus \{1\} \\
& \quad \sum_{i=1}^{n} (y_{ij} + t_{df}(i)x_{ij}) \leq t_j \quad j \in I \\
& \quad \sum_{i=1}^{n} (y_{ij} + t_{df-ex}(i)x_{ij}) \leq t_j + t_{df}(j) \quad j \in I \\
& \quad \sum_{i=1}^{n} (y_{ij} + t_{df-ex-wb}(i)x_{ij}) \leq t_j + t_{df-ex}(j) \quad j \in I \\
& \quad y_{ij} \leq U x_{ij} \quad i, j \in I \\
& \quad t_i + U x_{ij} - U \leq y_{ij} \leq t_i \quad i, j \in I \\
& \quad t_i + t_{df-ex-wb}(i) \leq z \quad i \in I \\
& \quad t_i \in \mathbb{R}^+ \quad i \in I \\
& \quad x_{ij} \in \{0, 1\} \quad i, j \in I \\
& \quad y_{ij} \in \mathbb{R}^+ \quad i, j \in I \\
& \quad z \in \mathbb{R}^+ 
\end{align*}
\]
B AMPL Script

******************************************************************************
## Input Sets and Parameters
##
## set of matrix instructions
set I;
## set of task precedences
set E within I cross I;
## time parameters for instructions
## time duration for data fetch stage
param t_df{I};
## time duration for execute
param t_ex{I};
## time duration for execute
param t_wb{I};
## number of processors
param num_processors;
******************************************************************************
## Set and Parameters for Model
##
## synthetic start task
param start_symbolic;
## set of matrix instruction with start task
set IS := I union {start};
## set adjacency elements that are not
set A := {(i,j) in (IS cross IS): i <> j};
## upper bound for variables t
param U := sum{i in I} (t_df[i] + t_ex[i] + t_wb[i]);
## set of predecessors
set prede (u in I) := {(v,w) in E: w = u};
******************************************************************************
## Variables
##
## adjacency matrix of stream graph
var x{A}, binary;
## linearization time y[i,j] = x[i,j] * t[i,j]
var y{A}, >=0;
## start time of instruction
var t{I}, >=0;
## makespan
var z, >=0;
******************************************************************************
## Objective
##
## minimize makespan
minimize objective: z;
******************************************************************************
## Constraint
##
## makespan is greater than or equal to the completion of all instructions
subject to makespan {i in I}: t[i] + t_df[i] + t_ex[i] + t_wb[i] <= z;
## instruction precedence constraint
subject to precedence {(i,j) in E}: t[i] + t_df[i] + t_ex[i] + t_wb[i] <= t[j];
## stream succession constraint for each stage
subject to stream_df {j in I}: sum{i in I: i <> j} (y[i,j] + t_df[i] * x[i,j]) <= t[j];
subject to stream_ex {j in I}: sum{i in I: i <> j} (y[i,j] + (t_df[i] + t_wb[i]) * x[i,j]) <= t[j] + t_df[j];
subject to stream_wb {j in I}: sum{i in I: i <> j} (y[i,j] + (t_df[i] + t_ex[i] + t_wb[i]) * x[i,j]) <= t[j] + t_df[j] + t_ex[j];
## stream graph processor constraint
subject to processors: sum{i in I} x[start,i] <= num_processors;
## stream graph successor constraint
subject to successors {i in I}: sum{j in I: i <> j} x[i,j] <= 1;
## stream graph predecessor constraint
subject to predecessors {j in I}: sum{i in IS: i <> j} x[i,j] = 1;

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## Linearization Constraints

- Linearization of quadratic term $y_{i,j} = x_{i,j} \cdot t_{i}$ for all $(i,j)$ in $A$.
- Subject to linearize 1 \{(i,j) in (I cross I): i \not= j\}:
  - $y_{i,j} \leq U \cdot x_{i,j}$.
- Subject to linearize 2 \{(i,j) in (I cross I): i \not= j\}:
  - $t_{i} - U + U \cdot x_{i,j} \leq y_{i,j}$.
- Subject to linearize 3 \{(i,j) in (I cross I): i \not= j\}:
  - $y_{i,j} \leq t_{i}$.
C List of Implemented Octave Functions

The following Octave operators and functions were implemented to work on our custom \texttt{p\_matrix} data type. These operations utilise our framework to execute in parallel.

- \texttt{<p\_matrix> + <p\_matrix>}
- \texttt{<p\_matrix> + <scalar>}
- \texttt{<p\_matrix> - <p\_matrix>}
- \texttt{<p\_matrix> - <scalar>}
- \texttt{<p\_matrix> * <p\_matrix>}
- \texttt{<p\_matrix> * <scalar>}
- \texttt{<p\_matrix> .* <p\_matrix>}
- \texttt{<p\_matrix> ./ <p\_matrix>}
- \texttt{<p\_matrix> .\^ <p\_matrix>}
- \texttt{abs(<p\_matrix>)}
- \texttt{mod(<p\_matrix>, <scalar>)}
- \texttt{sin(<p\_matrix>)}
- \texttt{cos(<p\_matrix>)}
- \texttt{sign(<p\_matrix>)}
- \texttt{round(<p\_matrix>)}
- \texttt{<p\_matrix> == <p\_matrix>}
- \texttt{<p\_matrix> != <p\_matrix>}

