SPICE$^2$ – A Spatial Parallel Architecture for Accelerating the SPICE Circuit Simulator

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Abstract

Spatial processing of sparse, irregular floating-point computation using a single FPGA enables up to an order of magnitude speedup (mean $2.8 \times$ speedup) over a conventional microprocessor for the SPICE circuit simulator. We deliver this speedup using a hybrid parallel architecture that spatially implements the heterogeneous forms of parallelism available in SPICE. We decompose SPICE into its three constituent phases: Model-Evaluation, Sparse Matrix-Solve, and Iteration Control and parallelize each phase independently. We exploit data-parallel device evaluations in the Model-Evaluation phase, sparse dataflow parallelism in the Sparse Matrix-Solve phase and compose the complete design in streaming fashion. We name our parallel architecture SPICE$^2$: Spatial Processors Interconnected for Concurrent Execution for accelerating the SPICE circuit simulator. We program the parallel architecture with a high-level, domain-specific framework that identifies, exposes and exploits parallelism available in the SPICE circuit simulator. Our design is optimized with an auto-tuner that can scale the design to use larger FPGA capacities without expert intervention and can even target other parallel architectures with the assistance of automated code-generation. This FPGA architecture is able to outperform conventional processors due to a combination of factors including high utilization of statically-scheduled resources, low-overhead dataflow scheduling of fine-grained tasks, and overlapped processing of the control algorithms.

We demonstrate that we can independently accelerate Model-Evaluation by a mean factor of $6.5 \times (1.4\text{--}23 \times)$ across a range of non-linear device models and Matrix-Solve by $2.4 \times (0.6\text{--}13 \times)$ across various benchmark matrices while delivering a mean combined speedup of $2.8 \times (0.2\text{--}11 \times)$ for the two together when comparing a Xilinx
Virtex-6 LX760 (40nm) with an Intel Core i7 965 (45nm). With our high-level framework, we can also accelerate Single-Precision Model-Evaluation on NVIDIA GPUs, ATI GPUs, IBM Cell, and Sun Niagara 2 architectures.

We expect approaches based on exploiting spatial parallelism to become important as frequency scaling slows down and modern processing architectures turn to parallelism (e.g. multi-core, GPUs) due to constraints of power consumption. Our thesis shows how to express, exploit and optimize spatial parallelism for an important class of problems that are challenging to parallelize.
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Chapter 1

Introduction

This thesis shows how to expose, exploit and implement the parallelism available in the SPICE simulator [1] to deliver up to an order of magnitude speedup (mean 2.8× speedup) on a single FPGA chip. SPICE (Simulation Program with Integrated Circuit Emphasis) is an analog circuit simulator that can take days or weeks of runtime on real-world problems. It models the analog behavior of semiconductor circuits using a compute-intensive non-linear differential equation solver. SPICE is notoriously difficult to parallelize due to its irregular, unpredictable compute structure, and a sloppy sequential description. It has been observed that less than 7% of the floating-point operations in SPICE are automatically vectorizable [2]. SPICE is part of the SPEC92 [3] benchmark collection which is a set of challenge problems for microprocessors. Over the past couple of decades, we have relied on innovations in computer architecture (clock frequency scaling, out-of-order execution, complex branch predictors) to speedup applications like SPICE. It was possible to improve performance of existing application binaries by retaining the exact same ISA (Instruction Set Architecture) abstraction at the expense of area and power consumption. However, these traditional computer organizations have now run up against a power wall [4] that prevents further improvements using this approach. More recently, we have migrated microprocessor designs towards “multi-core” organizations [5, 6] which are an admission that further improvements in performance must come from parallelism that will be explicitly exposed to the hardware. Reconfigurable, spatial architectures like FPGAs provide an alternate platform for accelerating applications such as SPICE.
at lower clock frequencies and lower power. Unlike multi-core organizations which can exploit a fixed amount of instruction-level parallelism, thread-level parallelism and data-level parallelism, FPGAs can be configured and customized to exploit parallelism at multiple granularities as required by the application. FPGAs offer higher compute density [7] by implementing computation using spatial parallelism which distributes processing in space rather than time. In this thesis, we show how to translate the high compute density available on a single-FPGA to accelerate SPICE by $2.8 \times (11 \times \text{max.})$ using a high-level, domain-specific programming framework. The key questions addressed in this thesis as follows:

1. Can SPICE be parallelized? What is the potential for accelerating SPICE?

2. How do we express the irregular parallel structure of SPICE?

3. How do we use FPGAs to exploit the parallelism available in SPICE?

4. Will FPGAs outperform conventional multi-core architectures for parallel SPICE?

We intend to use SPICE as a design-driver to address broader questions. How do we capture, exploit and manage heterogeneous forms of parallelism in a single application? How do we compose the parallelism in SPICE to build the integrated solution? Is it sufficient to constrain this parallelism to a general-purpose architecture (e.g. Intel multi-core)? Will the performance of this application scale easily with larger processing capacities?

### 1.1 Contributions

We develop a novel framework to capture and reason about complex applications like SPICE with heterogeneous forms of parallelism. We use parallel patterns for
expressing and unifying this parallelism effectively. We organize our parallelism on reconfigurable architecture to spatially implement and customize the compute organization to each style of parallelism. We configure this hardware with a compilation flow supported by **auto-tuning** to implement the parallelism in a scalable manner on available resources. We use SPICE as a design driver to experiment with these ideas and quantitatively demonstrate the benefits of this design philosophy.

As shown in the matrix in Table 1.1, we first decompose SPICE into its three constituent phases: (1) Model Evaluation, (2) Sparse Matrix-Solve and (3) Iteration Control. We describe these phases in Chapter 2.

We identify opportunities for parallel operation in the different phases of SPICE and provide design strategies for hardware implementation using suitable parallel patterns. **Parallel patterns** are a paradigm of structured concurrent programming that help us express, capture and organize parallelism. Paradigms and patterns were identified in the classic paper by Floyd [8]. They are invaluable tools and techniques for describing computer programs in a methodical “top-down” manner. When describing parallel computation, we need similar patterns for organizing our parallel program. We can always expose available parallelism in the most general Communicating Sequential Processes (CSP [9]) model of computation. However, this provides no guidance for managing communication and synchronization between the parallel processes. It also does not necessarily provide any insight into choosing the right hardware implementation for the parallelism. In contrast, **parallel patterns** are reusable strategies for describing and composing parallelism that serve two key purposes. First, these patterns let the programmer construct the parallel program from well-known, primitive, building blocks that can compose with each other. Second, the patterns also provide structure for constructing parallel hardware that best implements the computation. This is more restricted than the general CSP model but simplifies the task of writing the parallel program and developing parallel hardware using well-developed solutions to avoid common pitfalls.

We analyze the structure of SPICE in the form of patterns and illustrate how they guide the selection of parallel FPGA organization. The Model-Evaluation phase
of SPICE can be effectively parallelized using the \textit{data-parallel} pattern. We implement this data-parallelism in a software-pipelined fashion on the FPGA hardware. We further exploit the static nature of this data-parallel computation to schedule the computation on a custom, time-multiplexed, VLIW architecture. We extract the irregular, \textit{sparse dataflow} parallelism available in the Matrix-Solve phase of SPICE that is different from the regular, \textit{data-parallel} pattern that characterizes Model-Evaluation phase. We use the KLU matrix-solver package to expose the static dataflow parallelism available in this phase and develop a distributed, token-dataflow architecture for exploiting that parallelism. This static dataflow capture allows us to distribute the graph spatially across parallel compute units and process the graph in a fine-grained fashion. Finally, we express the Iteration-Control phase of SPICE using a \textit{streaming} pattern in the high-level SCORE framework to enable overlapped evaluation and efficiently implement the SPICE analysis state machines.

We now describe some key ideas that help us develop our parallel FPGA implementation:

- **Domain-Specific Framework**: The parallelism in SPICE is a heterogeneous composition of multiple parallel domains. To properly capture and exploit parallelism in SPICE, we develop domain-specific flows which include high-level languages and compilers customized for each domain. We develop a Verilog-AMS compiler to describe the non-linear differential equations for Model-Evaluation. We use the SCORE \cite{10} framework (TDF language and associated compiler and runtime) to express the Iteration-Control phase of SPICE and compose the complete design. This high-level problem capture makes the spatial, parallel structure of the SPICE program evident. We can then use the compiler and develop auto-tuners to exploit this spatial structure on FPGAs.

- **Specialization**: The SPICE simulator requires three inputs: (1) Circuit (2) Device type and parameters (3) Convergence/Termination control options. We exploit the static and early-bound nature of certain inputs to generate optimized hardware implementation and quantify their benefits. The Model Evaluation
computation is dependent on device type and the device parameters. We can
pre-compile, optimize and schedule the static dataflow computation for the few
device types used by SPICE and simply select the appropriate type prior to
execution. We get an additional improvement in performance and reduction
in area by specializing the device evaluation to specific device parameters that
are common to specific semiconductor manufacturing processes. We quantify
the benefits of specialization in Table 3.4 in Chapter 3. The Sparse Matrix
Solve computation operates on a dataflow graph that can be extracted at the
start of the simulation. This early-bound structure of the computation allows
us to distribute and place operations on our parallel architecture once and then
reuse the placement throughout the simulation. We show a $1.35 \times$ improvement
in sequential performance of Matrix-Solve phase through specialization in Ta-
ble 4.5 from Chapter 4. Finally, the simulator convergence and terminal control
parameters are loaded into registers at the start of the simulation and are used
by the hardware to manage the SPICE iterations.

- **Spatial Architectures**: The different parallel patterns in SPICE can be
efficiently implemented on customized spatial architectures on the FPGA. Hence,
we identify and design the spatial FPGA architecture to match the pattern of
parallelism in the SPICE phases. Our statically-scheduled VLIW architecture
delivers high utilization of floating-point resources (40–60%) and is supported
by an efficient time-multiplexed network that balances the compute and com-
 municate operations. We use a Token-Dataflow architecture to spatially dis-
tribute processing of the sparse matrix factorization graph across the paral-
lel architecture. The sparse graph representation also eliminates the need for
address-calculation and sequential memory access dependencies that constrain
the sequential implementation. Finally, we develop a streaming controller that
enables overlapped processing of the Iteration Control computation with the
rest of SPICE.

- **Scalability**: Typical FPGA designs are rigidly tied to a particular FPGA fam-
ily and require a complete redesign to scale to larger FPGA sizes. We are interested in developing scalable FPGA systems that can be automatically adapted to use extra resources when they become available. For speedup calculations in this thesis, our parallel design is engineered to fit into the resources available on a single Virtex-6 LX760 FPGA. We develop an auto-tuner that enables easy scalability of our design to larger FPGA capacities. Our auto-tuner explores area-memory-time space to pick the implementation that fits available resources while delivering the highest performance. The Model-Evaluation and Iteration-Control phases of SPICE can be accelerated almost linearly with additional logic and memory resources. With a sufficiently large FPGA we can spatially implement the complete dataflow graph of the Model-Evaluation and Iteration-Control computation as a physical circuit without using a time-shared VLIW architecture. Furthermore, the auto-tuner can automatically choose the proper configuration for this larger FPGA. This is relevant in light of the recently announced Xilinx Virtex-7 FPGA family [11] that has 2–3× the capacity of the FPGA we use in this thesis. However, the Sparse Matrix-Solve phase will only enjoy limited additional parallelism with extra resources. Additional research into decomposed matrix solvers is necessary to properly scale performance on a larger system.

The quantitative contributions of this thesis include:

1. **Complete simulator**: We accelerate the complete double-precision implementation of the SPICE simulator by 0.2-11.1× when comparing the Xilinx Virtex-6 LX760 (40nm technology node) with an Intel Core i7 965 processor (45nm technology node) with no compromise in simulation quality.

2. **Model-Evaluation Phase**: We implement the Model-Evaluation phase by compiling a high-level Verilog-AMS description to a statically-scheduled custom VLIW architecture. We demonstrate a speedup of 1.4–23.1× across a range of non-linear device models when comparing Double-Precision implementations on a Virtex-6 LX760 compared to an Intel Core i7 965. We also deliver speedups
of 4.5–123.5× for a Virtex-5 LX330, 10.1–63.9× for an NVIDIA 9600GT GPU, 0.4–6× for an ATI FireGL 5700 GPU, 3.8–16.2× for an IBM Cell and 0.4–1.4× for a Sun Niagara 2 architectures when comparing Single-Precision evaluation to an Intel Xeon 5160 across these architectures at 55nm–65nm technology. We also show speedups of 4.5–111.6× for a Virtex-6 LX760, 13.1–133.1× for an NVIDIA GTX285 GPU and 2.8–1200× for an ATI Firestream 9270 GPU when comparing Single-Precision evaluation to an Intel Core i7 965 on architectures at 40–55nm technology.

3. Sparse Matrix-Solve Phase: We implement the sparse dataflow graph representation of the Sparse Matrix-Solve computation using a dynamically-routed Token Dataflow architecture. We show how to improve the performance of irregular, sparse matrix factorization by 0.6–13.4× when comparing a 25-PE parallel implementation on a Xilinx Virtex-6 LX760 FPGA with a 1-core implementation on an Intel Core i7 965 for double-precision floating-point evaluation.

4. Iteration-Control Phase: Finally, we compose the complete simulator along with the Iteration-Control phase using a hybrid streaming architecture that combines static scheduling with limited dynamic selection. We deliver 2.6× (max 8.1×) reduction in runtime for the SPICE Iteration-Control algorithms when comparing a Xilinx Virtex-6 LX760 with an Intel Core i7 965.

1.2 SPICE

We now present an overview of the SPICE simulator and identify opportunities for parallel operation. At the heart of the SPICE simulator is a non-linear differential equation solver. The solver is a compute-intensive iterative algorithm that requires an evaluation of non-linear physical device equations and sparse matrix factorizations in each iteration. When the number of devices in the circuit grows, the simulation time will increase correspondingly. As manufacturing process technologies keep improving we are able to build smaller semiconductor devices and consequently we can pack
larger circuits into a single chip. The ITRS roadmap for semiconductors [12, 13] suggests that SPICE simulation runtimes for modeling non-digital effects [12] and large power-ground networks [13] are a challenge. This will increase the size of the circuits ($N$) we want to simulate. SPICE simulation time scales as $O(N^{1.2})$ which outpaces the increase in computing capacity of conventional architectures $O(N^{0.96})$ as shown in Figure 1.1 (additional discussion in Section 2.2). Our parallel, single-FPGA design scales more favorably with increasing circuit size as $O(N^{0.7})$. We show the peak capacities of the processor and FPGA used in the comparison in Table 1.2. A parallel SPICE solver will decrease verification time and reduce the time-to-market for integrated circuits. However, as identified in [2] parallelizing SPICE is hard. Over the past three decades, parallel SPICE efforts have either delivered limited speedups or sacrificed simulation quality to improve performance. Furthermore, the techniques and solutions are specific to a particular system and not easily portable to newer, better architectures. The SPICE simulator is a complex mix of multiple compute patterns that demands customized treatment for proper parallelization. After a careful performance analysis, it is clear that the Model Evaluation and Sparse Matrix-Solve phases of SPICE are the key contributors to total SPICE runtime. We note that the Model Evaluation phase of SPICE is in fact embarrassingly parallel by itself and should be relatively easy to parallelize. We describe our parallel solution in Chapter 3. In contrast, the Sparse Matrix-Solve phase is more challenging. This is one of the key reasons why earlier studies were unable to parallelize the complete simulator. The underlying compute structure of Sparse Matrix Solve is irregular, unpredictable and consequently performs poorly on conventional architectures. Recent advances in numerical algorithms provide an opportunity to reduce these limitations. We use the KLU solver [14, 15] to generate an irregular, dataflow factorization graph that exposes the available parallelism in the computation. We explain our approach for parallelizing the challenge Sparse Matrix-Solve phase in Chapter 4. Finally, the Iteration-Control phase of SPICE is responsible for co-ordinating the simulation steps. While it is a tiny fraction of sequential runtime, it may become a non-trivial fraction of the accelerated SPICE solver. We show how to efficiently implement the Iteration-
Table 1.2: Peak Floating-Point Throughputs (Double-Precision)

<table>
<thead>
<tr>
<th>Family</th>
<th>Chip</th>
<th>Tech. (nm)</th>
<th>Clock (GHz)</th>
<th>Peak GFLOPS (Double)</th>
<th>Power (Watts)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Intel Core i7</td>
<td>965</td>
<td>45</td>
<td>3.2</td>
<td>25</td>
<td>130</td>
</tr>
<tr>
<td>Xilinx Virtex-6</td>
<td>LX760</td>
<td>40</td>
<td>0.2</td>
<td>26</td>
<td>20–30</td>
</tr>
</tbody>
</table>

Control phase of SPICE in Chapter 5. Finally, in Chapter 6 we discuss how to integrate the complete SPICE solver on a single FPGA.

### 1.3 FPGAs

We briefly review the FPGA architecture and highlight some key characteristics of an FPGA that make it well-suited to accelerate SPICE. A Field-Programmable Gate Array is a massively-parallel architecture that implements computation using hundreds of thousands of tiny programmable computing elements called LUTs (lookup tables) connected to each other using a programmable communication fabric. Typically these LUTs are clustered into SLICEs (2–4 LUTs per SLICE depending on FPGA architecture). Moore’s Law delivers progressively larger FPGA capacities with increasing numbers of SLICEs per chip. Modern FPGAs also include hundreds of embedded memories and integer multipliers distributed across the fabric for concurrent operation. We show a small region inside an island-style FPGA in Figure 1.2 (LUTs...
Figure 1.2: A Small Region of an FPGA
correspond to islands in a sea of interconnect). An FPGA allows us to configure the computation in space rather than time and evaluate multiple operations concurrently in a fine-grained fashion. In Figure 1.3, we show a simple calculation and its conceptual implementation on a CPU and an FPGA. For a CPU implementation, we process the instructions stored in an instruction memory temporally on an ALU while storing the intermediate results (i.e. variables) in a data memory. On an FPGA, we can implement the operations as spatial circuits while implementing the dependencies between the operations physically using pipelined wires instead of variables stored in memory. Additionally, a pipelined FPGA circuit implementation of certain operations in the computation (e.g. divide, sqrt) requires multiple cycles on the CPU while we can configure a custom, pipelined circuit for those operations on the FPGA for higher throughput.

While FPGAs have been traditionally successful at accelerating highly-parallel, communication-rich algorithms and application kernels, they are a challenging target for mapping a complete application with diverse computational patterns such as the SPICE circuit simulator. Furthermore, programming an FPGA with existing techniques for high performance requires months of effort and low-level tuning. Additionally, these designs do not automatically scale for larger, newer FPGA capacities that become available with technology scaling and require re-compilation and re-optimization by an expert.

1.4 Implementing SPICE on an FPGA

We now briefly describe our hardware architecture and mapping flow for implementing SPICE on a single FPGA. We use a high-level framework to express, extract and compose the parallelism in the different phases of SPICE. We develop a hybrid FPGA architecture that combines custom architectures tailored to the parallelism in each phase of SPICE. In Figure 1.4, we show a high-level block diagram of the composite FPGA architecture with the three regions tailored for each phase of SPICE. We show a simple picture of our mapping flow in Figure 1.5. While the logic configuration
of the FPGA is the same for all SPICE circuits, we need to program the FPGA memories in an instance-specific manner for each circuit we simulate. We show how we assemble the FPGA hardware and provide additional details of our mapping flow in Chapter 6.

A completely spatial implementation of the SPICE circuit simulator is too large to fit on a single FPGA at 40nm technology today. On larger FPGAs, in the near future, it will become possible to fit the fully-spatial Model Evaluation graphs entirely on a single FPGA. For these graphs, the cost of a fully-spatial implementation gets amortized across several device evaluations. However, for the Sparse Matrix-Solve phase, the limited amount of parallelism and low reuse does not motivate a fully-spatial design even if we have large FPGAs. Furthermore, configuring the FPGA for every circuit using FPGA CAD tools (synthesize logic, place LUTs, and route wires) can itself take hours or days of runtime defeating the purpose of a parallel FPGA implementation for SPICE. We develop the custom FPGA architecture using virtual organizations capable of accommodating the entire SPICE simulator operation on a single FPGA. Furthermore, we simplify our circuit-specific compilation step by requiring only a memory generation step to program the virtual architecture (static
Figure 1.4: Block-Diagram of the SPICE FPGA Solver

schedule or graph structure).
Figure 1.5: FPGA Mapping Flow for SPICE
Chapter 2

Background

In this chapter, we review the SPICE simulation algorithm and discuss the inner-workings of the three constituent phases of SPICE. We review existing literature and categorize previous attempts at parallelizing SPICE. We argue that these previous approaches were unable to fully capitalize on the parallelism available within the SPICE simulator due to limitations of hardware organizations and inefficiencies in the software algorithms. In the following chapters, we will describe our FPGA-based "spatial" approach that provides a communication-centric architecture for exploiting the parallelism in SPICE.

2.1 The SPICE Simulator

SPICE simulates the dynamic analog behavior of a circuit described by its constituent non-linear differential equations. It was developed at the EECS Department of the University of California, Berkeley by Donald Pederson, Larry Nagel [1], Richard Newton, and many other contributors to provide a fast and robust circuit simulation program capable of verifying integrated circuits. It was publicly announced thirty-seven years ago in 1973 and many versions of the package have since been released. spice2g6 was part of the SPEC89 and SPEC92 benchmark sets of challenge problems for microprocessors. Many commercial versions of analog circuit simulators inspired by SPICE now exist as part of a large IC design and verification industry. For this study, we use the open-source spice3f5 package released from Berkeley in 1995.
SPICE circuit equations model the linear (e.g., resistors, capacitors, inductors) and non-linear (e.g., diodes, transistors) behavior of devices and the conservation constraints (i.e., Kirchoff’s current laws—KCL) at the different nodes and branches of the circuit. SPICE solves the non-linear circuit equations by alternately computing small-signal linear operating-point approximations for the non-linear elements and solving the resulting system of linear equations until it reaches a fixed point. The linearized system of equations is represented as a solution of $A\vec{x} = \vec{b}$, where $A$ is the matrix of circuit conductances, $\vec{b}$ is the vector of known currents and voltage quantities and $\vec{x}$ is the vector of unknown voltages and branch currents.

Spice3f5 uses the Modified Nodal Analysis (MNA) technique to assemble circuit equations into matrix $A$. The MNA approach is an improvement over conventional nodal analysis by allowing proper handling of voltage sources and controlled current sources. It requires the application of Kirchoff’s Current Law at all the nodes in the circuit with voltage at each node being an unknown. It then introduces un-
knowns for currents through branches to allow voltage sources and controlled current sources to be represented.

The simulator calculates entries in $A$ and $\vec{b}$ from the device model equations that describe device transconductance (e.g., Ohm’s law for resistors, transistor I-V characteristics) in the Model-Evaluation phase. It then solves for $\vec{x}$ using a sparse-direct linear matrix solver in the Matrix-Solve phase. We show the steps in the SPICE algorithm in Figure 2.1. The inner loop iteration supports the operating-point calculation for the non-linear circuit elements, while the outer loop models the dynamics of time-varying devices such as capacitors. The simulator loop management controls are handled by the Iteration Control phase. We illustrate the operation of the simulator using an example shown in Figure 2.2. We first write down the non-linear differential equations that capture circuit behavior in Equation 2.1–Equation 2.6. The non-linear (diode) and time-varying (capacitor) devices are represented using linearized forms in Equation 2.2 and Equation 2.3 respectively. We then reassemble these equations into $A\vec{x} = \vec{b}$ form shown in Figure 2.4.

2.1.1 Model-Evaluation

In the Model-Evaluation phase, the simulator computes conductances and currents through different elements of the circuit and updates corresponding entries in the matrix with those values. We expand the equations for the example circuit in Fig-
\[ I(R1) = (V_1 - V_2) \cdot \frac{1}{R_1} \quad (2.1) \]
\[ I(D1) = G_{eq}(D1) \cdot V_2 + I_{eq}(D1) \quad (2.2) \]
\[ I(C1) = G_{eq}(C1) \cdot V_2 + I_{eq}(C1) \quad (2.3) \]
\[ I(V1) = I(R1) \quad (2.4) \]
\[ I(R1) = I(C1) + I(D1) \quad (2.5) \]
\[ V_1 = V_1 \quad (2.6) \]

Figure 2.3: Circuit Equations from Conservation Laws (KCL)

\[
\begin{bmatrix}
  G_1 & -G_1 & 1 \\
  -G_1 & G_1 + G_{eq}(C1) + G_{eq}(D1) & 0 \\
  1 & 0 & 0
\end{bmatrix}
\times
\begin{bmatrix}
  V_1 \\
  V_2 \\
  I(V1)
\end{bmatrix} =
\begin{bmatrix}
  0 \\
  I_{eq}(C1) + I_{eq}(D1) \\
  V_1
\end{bmatrix}
\]

Figure 2.4: Matrix Representation of Circuit Equations in SPICE

For the linear elements (e.g. resistors) this needs to be done only once at the start of the simulation (shown in blue in Figures 2.5 and 2.6). For non-linear elements, the simulator must search for an operating-point using Newton-Raphson iterations which requires repeated evaluation of the model equations multiple times per timestep (inner loop labeled Newton-Raphson Iterations in Figure 2.1 and shown in red in Figure 2.5 and 2.6). For time-varying components, the simulator must recalculate their contributions at each timestep based on voltages at several previous timesteps. This also requires repeated re-evaluations of the device-model (outer loop labeled Transient Iterations in Figure 2.1 and shown in green in Figure 2.5 and 2.6).

\[ I(R1) = (V_1 - V_2) \cdot G_1 \quad (2.7) \]
\[ I(C1) = \left( \frac{2 \cdot C_1}{\delta t} \right) \cdot V_2 - \left( \frac{2 \cdot C_1}{\delta t} \right) \cdot V_{old} + I_{old}(C1) \quad (2.8) \]
\[ I(D1) = \left( \frac{I_s}{v_j} \cdot e^{V_2/v_j} \right) \cdot V_2 + I_s \cdot (e^{V_2/v_j} - 1) \quad (2.9) \]

Figure 2.5: Circuit Equations with contributions from devices
\[
\begin{bmatrix}
G1 & -G1 & 1 \\
-G1 & G1 & 0 \\
+ & (I_s/v_j) \cdot e^{V_2/v_j} & 1
\end{bmatrix}
\cdot
\begin{bmatrix}
V_1 \\
V_2 \\
I(V1)
\end{bmatrix}
=
\begin{bmatrix}
0 \\
-(2 \cdot C1/\delta t) \times V_{old2} + I_{old}(C1) \\
+I_s \cdot (e^{V_2/v_j} - 1) \\
V_1
\end{bmatrix}
\]

Figure 2.6: Matrix contributions from the different devices

\begin{align}
A \cdot \vec{x} &= \vec{b} \tag{2.10} \\
L \cdot U \cdot \vec{x} &= \vec{b} \tag{2.11} \\
L \cdot \vec{y} &= \vec{b} \tag{2.12} \\
U \cdot \vec{x} &= \vec{y} \tag{2.13}
\end{align}

\begin{align}
\begin{bmatrix}
a_{11} & a_{12} & a_{13} \\
a_{21} & a_{22} & a_{23} \\
a_{31} & a_{32} & a_{33}
\end{bmatrix} \cdot
\begin{bmatrix}
x_1 \\
x_2 \\
x_3
\end{bmatrix}
&=
\begin{bmatrix}
b_1 \\
b_2 \\
b_3
\end{bmatrix} \tag{2.14}
\end{align}

\begin{align}
\begin{bmatrix}
1 & 0 & 0 \\
l_{21} & 1 & 0 \\
l_{31} & l_{32} & 1
\end{bmatrix}
\cdot
\begin{bmatrix}
u_{11} & u_{12} & u_{13} \\
0 & u_{22} & u_{23} \\
0 & 0 & u_{33}
\end{bmatrix} \cdot
\begin{bmatrix}
x_1 \\
x_2 \\
x_3
\end{bmatrix}
&=
\begin{bmatrix}
b_1 \\
b_2 \\
b_3
\end{bmatrix} \tag{2.15}
\end{align}

\begin{align}
\begin{bmatrix}
1 & 0 & 0 \\
l_{21} & 1 & 0 \\
l_{31} & l_{32} & 1
\end{bmatrix} \cdot
\begin{bmatrix}
y_1 \\
y_2 \\
y_3
\end{bmatrix}
&=
\begin{bmatrix}
b_1 \\
b_2 \\
b_3
\end{bmatrix} \tag{2.16}
\end{align}

\begin{align}
\begin{bmatrix}
u_{11} & u_{12} & u_{13} \\
0 & u_{22} & u_{23} \\
0 & 0 & u_{33}
\end{bmatrix} \cdot
\begin{bmatrix}
x_1 \\
x_2 \\
x_3
\end{bmatrix}
&=
\begin{bmatrix}
y_1 \\
y_2 \\
y_3
\end{bmatrix} \tag{2.17}
\end{align}

Figure 2.7: Matrix Solve Stages

\subsection{2.1.2 Matrix Solve}

The simulator \texttt{spice3f5} uses the Modified Nodal Analysis (MNA) technique \cite{16} to assemble circuit equations into matrix $A$. Since circuit elements ($N$) tend to be connected to only a few other elements, there are a constant number ($O(1)$) of entries per row of the matrix. Thus, the MNA circuit matrix with $O(N^2)$ entries is highly sparse with $O(N)$ nonzero entries ($\approx 99\%$ of the matrix entries are 0). The matrix structure is mostly symmetric with the asymmetry being added by the presence of independent sources (e.g. input voltage source) and inductors. The underlying non-zero structure of the matrix is defined by the topology of the circuit shown in Figure \ref{fig:circuit} and
consequently remains unchanged throughout the duration of the simulation. In each iteration of the loop shown in Figure 2.1, only the numerical values of the non-zeroes are updated in the Model-Evaluation phase of SPICE with contributions from the non-linear elements. To find the values of unknown node voltages and branch currents \( \vec{x} \), we must solve the system of linear equations \( A\vec{x} = \vec{b} \) as shown in Equation 2.14.

The sparse, direct matrix solver used in \textit{spice3f5} first reorders the matrix \( A \) to minimize fill-in using a technique called Markowitz reordering \cite{17}. This tries to reduce the number of additional non-zeroes (fill-in) generated during LU factorization. It then factorizes the matrix by dynamically determining pivot positions for numerical stability (potentially adding new non-zeros) to generate the lower-triangular component \( L \) and upper-triangular component \( U \) such that \( A = LU \) as shown in Equation 2.15. Finally, it calculates \( \vec{x} \) using Front-Solve \( L\vec{y} = \vec{b} \) (see Equation 2.16) and Back-Solve \( U\vec{x} = \vec{y} \) operations (see Equation 2.17).

2.1.3 Iteration Control

The SPICE iteration controller is responsible for two kinds of iterative loops shown in Figure 2.1: (1) \textit{inner loop}: linearization iterations for non-linear devices and (2) \textit{outer loop}: adaptive time-stepping for time-varying devices. The Newton-Raphson algorithm is responsible for computing the linear operating-point for the non-linear devices like diodes and transistors. Additionally, an adaptive time-stepping algorithm based on truncation error calculation (Trapezoidal approximation, Gear approximation) is used for handling the time-varying devices like capacitors and inductors. The controller also implements the loops in a data-dependent manner using customized convergence conditions and local truncation error estimations.

**Convergence Condition**: The simulator declares convergence when two consecutive iterations generate solution vectors and non-linear approximations that are within a prescribed tolerance respectively. We show the condition used by SPICE to determine if an iteration has converged in Equation 2.18 and Equation 2.19. Here, \( \vec{V}_i \) or \( \vec{I}_i \) represent the voltage or current unknowns in the \( i \)-th iteration of the Newton-Raphson loop. The convergence conditions compare the current solution vector in
iteration \((i)\) with the previous iteration \((i - 1)\). SPICE also performs a similar convergence check on the non-linear function described in the Model-Evaluation. The closeness between the values in consecutive iterations is parametrized in terms of user-specified tolerance values: \(\text{reltol} \) (relative tolerance), \(\text{abstol} \) (absolute tolerance), and \(\text{vntol} \) (voltage tolerance).

\[
|\vec{V}_i - \vec{V}_{i-1}| \leq \text{reltol} \cdot \max(|\vec{V}_i|, |\vec{V}_{i-1}|) + \text{vntol} \\
|\vec{I}_i - \vec{I}_{i-1}| \leq \text{reltol} \cdot \max(|\vec{I}_i|, |\vec{I}_{i-1}|) + \text{abstol}
\]  \hspace{1cm} (2.18) \hspace{1cm} (2.19)

Typical values for these tolerances parameters are: \(\text{reltol}=1e^{-3}\) (accuracy of 1 part in 1000), \(\text{abstol}=1e^{-12}\) (accuracy of 1 picoampere) and \(\text{vntol}=1e^{-6}\) (accuracy of 1 \(\mu\)volt). This means the simulator will declare convergence when the changes in voltage and current quantities get smaller than the convergence tolerances.

**Local Truncation Error (LTE):** Local Truncation Error is a local estimate of accuracy of the Trapezoidal approximation used for integration. The truncation-error-based time-stepping algorithm in \texttt{spice3f5} computes the next stepsize \(\delta_{n+1}\) as a function of the LTE \((\epsilon)\) of the current iteration and a Trapezoidal divided-difference approximation \((DD_3)\) of the charges \((x)\) at a few previous iterations. The equation for stepsize is shown in Equation \text{2.20}. For a target LTE, the Iteration Controller can match the stepsize to the rate of change of circuit quantities. If the circuit quantities are changing too rapidly, it can slow down the simulation by generating finer timesteps. This allows the simulator to properly resolve the rapidly changing circuit quantities. In contrast, if the circuit is quiescent (e.g. digital circuits between clock edges), the simulator can take larger timesteps for a faster simulation. When the change in the circuit quantities is small, a detailed simulation at fine timesteps will be a waste of work. Instead, the simulator can advance the simulation with larger stepsizes. A tolerance parameter \(\text{trtol}\) provides the user additional control over tuning the timestep sizes. This parameter (default \(\text{trtol}=7\)) can be increased if necessary to avoid excessively fine timesteps. The stepsize \(\delta_{n+1}\) is added to the current timestep to advance the simulation as shown in Equation \text{2.21}.
2.2 SPICE Performance Analysis

In this section, we discuss important performance trends and identify performance bottlenecks and characteristics that motivate our parallel approach. We use spice3f5 running on an Intel Core i7 965 for these experiments.

2.2.1 Total Runtime

We first measure total runtime of spice3f5 across a range of benchmark circuits on an Intel Core i7 965. We use a lightweight timing measurement scheme using hardware-performance counters (PAPI [18]) that does not impact the actual runtime of the program. We tabulate the size of the circuits used for this measurement along with other circuit parameters in Table 4.2. We graph the runtimes as a function of circuit size in Figure 2.8. We observe that runtime scales as $O(N^{1.2})$ as we increase...
What contributes to this runtime? To understand this, we break down the contribution to total runtime from the different phases of SPICE in Figure 2.9. We observe that, for most cases, total runtime is dominated by the Model-Evaluation Phase. This is because the circuit is mostly composed of non-linear transistor elements. In some cases, the Sparse Matrix Solve phase can be a significant fraction of total runtime. This is true for circuits with large parasitic components (e.g. capacitors, resistors) where the non-linear devices are a small portion of total circuit size. Finally, the control algorithm that sequences and manages SPICE iterations ends up taking a small fraction of total runtime. This suggests that in our parallel solution we must deliver high speedups for the intensive Model-Evaluation and Sparse Matrix Solve phases of SPICE while ensuring that we do not ignore the control algorithms. If we do not parallelize the Iteration Control phase, it may create a sequential bottleneck due to Amdahl’s Law.

2.2.2 Runtime Scaling Trends

Why does SPICE runtime scale super-linearly when we increase circuit size? The Model-Evaluation component of total runtime increases linearly with the number of non-linear devices being simulated (approximately $O(N^{1.1})$) rather than the expected
$O(N)$ due to limited cache capacity of the processor; also see Figure 2.13(a) as shown in Figure 2.10(a). The Matrix-Solve component of runtime empirically grows as $O(N^{1.2})$ as shown in Figure 2.10(b) (also see Figure 2.13(b) for floating-point operation trends). Overall, we observe from Figure 2.8 that runtime grows as $O(N^{1.2})$ where $N$ is the size of the circuit.

What other factors impact overall runtime? In Figure 2.11(a), we observe that floating-point instructions constitute only $\approx 20\%-40\%$ of total instruction count (smaller fraction for larger circuits). The remaining instructions are integer and memory access operations that support the floating-point work. These instructions are purely overhead and can be implemented far more efficiently using spatial FPGA hardware. They also consume instruction cache capacity and issue bandwidth, limiting the achievable floating-point peak. Furthermore, we observe that increasing circuit sizes results in higher L2 cache misses. We see L2 cache miss rates as high as 30\%-70\% L2 for our benchmarks in Figure 2.11(b). As we increase circuit size, the sparse matrix and circuit vectors spill out of the fixed cache capacity leading to a higher cache miss rate. For the benchmarks we use, the sparse matrix storage exceeds the 256KB per-core L2 cache capacity of an Intel Core i7 965 for all except a few small benchmarks. We also show the overflow factor (Memory required/Cache size) for our benchmarks in Figure 2.11(b). A parallel architecture for accelerating SPICE has the
opportunity to properly distribute data across multiple, onchip, embedded memories and manage communication operations (i.e. moving data between memories) to deliver good performance. Our FPGA architecture exploits spatial parallelism, even in the non-floating-point portion of the computation, to deliver high speedup.

2.2.3 CMOS Scaling Trends

As we continue to reap the benefits of Moore’s Law [19], we need to simulate increasingly larger circuits using SPICE. As we saw in the previous Section 2.2.2 sequential SPICE runtimes scales as $O(N^{1.2})$ where $N=$size of the circuit. This means sequential SPICE runtimes will get increasingly slower as we scale to denser circuits. Furthermore, as we shrink device feature sizes to finer geometries, we must model increasingly detailed physical effects in the analog SPICE simulations. This will increase the amount of time spent performing Model-Evaluation [20] as shown in Figure 2.12(a).

For example, the mos1 MOSFET model implements the Shichman-Hodges equations which are adequate for discrete transistor designs. The semi-empirical mos3 model was originally developed for integrated CMOS short-channel transistors at 1-2µm or larger. The new bsim3v3 and bsim4 models are more accurate and commonly used for today’s technology. It may become necessary to use different models for RF simulations (psp model) or bipolar transistors (mextram).
At smaller device sizes, our circuits will suffer the impact of tighter coupling and interference between the circuit elements. This will add additional modeling requirement of parasitic elements (e.g., capacitors, resistors) which increase the size of the matrix. This will, in turn, increase the time spent in the Sparse Matrix-Solve phase. The increase in SPICE runtime per iteration due to inclusion of parasitic effects is shown in Figure 2.12(b).

2.2.4 Parallel Potential

We now try to identify the extent of parallelism available in the two computationally-intensive phases of SPICE. In Figure 2.13(a), we plot the total number of floating-point operations as well as the latency of the Model-Evaluation computation assuming ideal parallel hardware as a function of the number of non-linear elements in the circuit. The amount of work grows linearly ($O(N)$) with the number of non-linear devices. However, the latency of the evaluation remains unchanged at the latency of a single device evaluation. Each non-linear device can be independently evaluated with local terminal voltages as inputs and local currents and conductances as outputs. Thus, the amount of parallelism is proportional to the number of non-linear devices in the circuit. This phase is embarrassingly parallel, and we show how to exploit this parallelism in Chapter 3.
(a) Total Work and Critical Latency Trends for the Model-Evaluation Phase

(b) Total Work and Critical Latency Trends for the Matrix-Solve Phase

Figure 2.13: Identifying Parallel Potential: Work vs. Latency
In Figure 2.13(b) we plot the total number of floating-point operations in the Sparse Matrix-Solve phase of SPICE as well as the critical path latency as a function of circuit size. Here, we observe that the amount of work in the sparse factorization increases as $O(N^{1.4})$ (compared to $O(N^3)$ for dense factorization). This growth rate is an empirical observation that is valid for circuit matrices in our benchmark set (see [21]). This explains the $O(N^{1.2})$ growth in total runtime we previously observed in Figure 2.8. The critical path latency of the factorization only grows as $O(N^{0.7})$. This suggests that a well-designed parallel architecture can extract parallelism from this phase since the critical path is not growing as fast as $N$. An architecture with low-latency communication and lightweight fine-grained spatial processing of operations should allow us to achieve faster operation.
<table>
<thead>
<tr>
<th>Year</th>
<th>Ref.</th>
<th>Name</th>
<th>Key Idea</th>
<th>Parallel Hardware</th>
<th>Sequential Baseline</th>
<th>PEs</th>
<th>Speedup</th>
<th>SPICE phase</th>
<th>Expression</th>
<th>Accuracy Tradeoff</th>
</tr>
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<tbody>
<tr>
<td>1979</td>
<td>[22]</td>
<td>-</td>
<td>Static Dataflow</td>
<td>-</td>
<td>-</td>
<td>100</td>
<td>72.8</td>
<td>Matrix Solve</td>
<td>Dataflow graph</td>
<td>None</td>
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<tr>
<td>1985</td>
<td>[23]</td>
<td>-</td>
<td>Vector task graph</td>
<td>S810 vector</td>
<td>S810 scalar</td>
<td>4</td>
<td>8.9</td>
<td>Matrix Solve</td>
<td>Vectorized graph</td>
<td>None</td>
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<tr>
<td>1986</td>
<td>[24]</td>
<td>Cayenne</td>
<td>Runtime Scheduling</td>
<td>VAX 8800</td>
<td>scalar (spice2)</td>
<td>2</td>
<td>1.7</td>
<td>All</td>
<td>Master-Slave Fortran decomp.</td>
<td>None</td>
</tr>
<tr>
<td>1987</td>
<td>[25]</td>
<td>SAP</td>
<td>Customized Datapath</td>
<td>Custom SIMD</td>
<td>VAX 8650 (spice2)</td>
<td>2</td>
<td>2</td>
<td>All</td>
<td>Compiled-Code</td>
<td>None</td>
</tr>
<tr>
<td>1988</td>
<td>[26]</td>
<td>-</td>
<td>Vector task graph</td>
<td>Alliant FX/8</td>
<td>1 PE (ADVICE)</td>
<td>6</td>
<td>7.3</td>
<td>All</td>
<td>Shared-Memory</td>
<td>None</td>
</tr>
<tr>
<td>1990</td>
<td>[27]</td>
<td>Paraspice</td>
<td>Vector task graph</td>
<td>Alliant FX/80</td>
<td>1 PE (spice2)</td>
<td>8</td>
<td>4.5</td>
<td>All</td>
<td>Shared-Memory, Parallel tasks</td>
<td>None</td>
</tr>
<tr>
<td>1992</td>
<td>[28]</td>
<td>AWSIM-3</td>
<td>Compiled-Code VLIW</td>
<td>Custom prototype</td>
<td>Sun-3/60</td>
<td>-</td>
<td>560</td>
<td>All</td>
<td>Compiled-Code</td>
<td>Table-lookup</td>
</tr>
<tr>
<td>1992</td>
<td>[29]</td>
<td>PSPLAX</td>
<td>Waveform relaxation</td>
<td>Alliant FX/80</td>
<td>1 PE (spice2)</td>
<td>8</td>
<td>6.1</td>
<td>Relax</td>
<td>-</td>
<td>Not spice3</td>
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<tr>
<td>1993</td>
<td>[31]</td>
<td>PACE</td>
<td>Dataflow scheduling</td>
<td>4-chip i860</td>
<td>SPARC2</td>
<td>4</td>
<td>3.7</td>
<td>All</td>
<td>Offline scheduling</td>
<td>None</td>
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<tr>
<td>1995</td>
<td>[32]</td>
<td>Sparta</td>
<td>Dataflow scheduling</td>
<td>Fujitsu AP-1000</td>
<td>1 PE</td>
<td>64</td>
<td>16.5</td>
<td>All</td>
<td>Message-Passing</td>
<td>None</td>
</tr>
<tr>
<td>1995</td>
<td>[33]</td>
<td>OSCAR</td>
<td>Task scheduling</td>
<td>Custom</td>
<td>1 PE</td>
<td>8</td>
<td>4.3</td>
<td>All</td>
<td>Code-generation</td>
<td>Single-Precision</td>
</tr>
<tr>
<td>1999</td>
<td>[34]</td>
<td>Transputer-SPICE</td>
<td>Block Decomposition</td>
<td>Ultra XL Transputer</td>
<td>-</td>
<td>7</td>
<td>-</td>
<td>Matrix Solve</td>
<td>“Parallel C”</td>
<td>None</td>
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</table>

Table 2.1: Taxonomy of Parallel SPICE Approaches in the past
<table>
<thead>
<tr>
<th>Year</th>
<th>Ref.</th>
<th>Name</th>
<th>Key Idea</th>
<th>Parallel Hardware</th>
<th>Sequential Baseline</th>
<th>PEs</th>
<th>Speedup</th>
<th>SPICE phase</th>
<th>Expression</th>
<th>Tradeoff</th>
</tr>
</thead>
<tbody>
<tr>
<td>2000</td>
<td>[35]</td>
<td>Xyce</td>
<td>Iterative Matrix Solve</td>
<td>SGI Origin 2000</td>
<td>1 PE</td>
<td>40</td>
<td>24</td>
<td>All</td>
<td>Message-Passing (MPI)</td>
<td>None</td>
</tr>
<tr>
<td>2002</td>
<td>[36]</td>
<td>SMP-SPICE</td>
<td>Multi-threading</td>
<td>Hitachi 9000 N4000</td>
<td>1 thread</td>
<td>spice3</td>
<td>1</td>
<td>7.4</td>
<td>All</td>
<td>C with PThreads</td>
</tr>
<tr>
<td>2006</td>
<td>[37]</td>
<td>SILCA</td>
<td>Approx.</td>
<td>-</td>
<td>1 thread</td>
<td>spice3</td>
<td>1</td>
<td>7.4</td>
<td>All</td>
<td>Sequential code</td>
</tr>
<tr>
<td>2007</td>
<td>[38]</td>
<td>OpenMP-SPICE</td>
<td>Data-Parallel</td>
<td>UltraSPARC 3</td>
<td>1 thread</td>
<td>spice3</td>
<td>4</td>
<td>1.3</td>
<td>Model Eval.</td>
<td>spice3f5 with OpenMP</td>
</tr>
<tr>
<td>2009</td>
<td>[40]</td>
<td>ACCIT-NSR</td>
<td>Data-parallel</td>
<td>ATI FireStream 9170</td>
<td>4-core AMD Phenom</td>
<td>320</td>
<td>50</td>
<td>Model Eval.</td>
<td>Data-parallel Brook [41]</td>
<td>None</td>
</tr>
<tr>
<td>2009</td>
<td>[45]</td>
<td>DD</td>
<td>Domain-Decomposition</td>
<td>FWGrid [46]</td>
<td>1 thread</td>
<td>spice3f5</td>
<td>32</td>
<td>119</td>
<td>Matrix Solve</td>
<td>C with PETSc package</td>
</tr>
</tbody>
</table>

**FPGA-based Approaches**

<table>
<thead>
<tr>
<th>Year</th>
<th>Ref.</th>
<th>Name</th>
<th>Key Idea</th>
<th>Parallel Hardware</th>
<th>Sequential Baseline</th>
<th>PEs</th>
<th>Speedup</th>
<th>SPICE phase</th>
<th>Expression</th>
<th>Tradeoff</th>
</tr>
</thead>
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<tr>
<td>1995</td>
<td>[48]</td>
<td>TINA</td>
<td>based on AWSIM-3</td>
<td>23 Xilinx XC4005 chips</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>All</td>
<td>Microasm Assembly</td>
<td>Table-lookup</td>
</tr>
<tr>
<td>1997</td>
<td>[49]</td>
<td>-</td>
<td>partial-eval</td>
<td>Altera Flex10K</td>
<td>SPARC station</td>
<td>3</td>
<td>27.7</td>
<td>Model Eval.</td>
<td>PECompiler C subset</td>
<td>Fixed-point</td>
</tr>
<tr>
<td>2003</td>
<td>[50]</td>
<td>SPO</td>
<td>analytical transform</td>
<td>Xilinx Spartan 3</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>All</td>
<td>Simulink graphs fixed-point</td>
<td>None</td>
</tr>
</tbody>
</table>

Table 2.2: Taxonomy of Recent Parallel SPICE Approaches and FPGA-based Systems
2.3 Historical Review

We now review the various studies and research projects that have attempted to parallelize SPICE in the past three decades or so. These studies attempt to accelerate the computation using a combination of parallel hardware architectures and/or numerical software algorithms that are more amenable to parallel evaluation. We tabulate and classify these approaches in Table 2.1 and Table 2.2. We can refine the classification of parallel SPICE approaches by considering underlying trends and characteristics of the different systems as follows:

1. **Compute Organization**: We see parallel SPICE solvers using a range of different compute organizations including conventional multi-processing, multi-core, VLIW, SIMD and Vector.

2. **Precision**: Under certain conditions, SPICE simulations can efficiently model circuits at lower precisions.

3. **Compiled Code**: In many cases, it is possible to generate efficient instance-specific simulations by specializing the simulator for a particular circuit.

4. **Scheduling**: Parallel computation exposed in SPICE must be distributed across parallel hardware elements for faster operation. Many designs develop novel, custom scheduling solutions that are applicable under certain structural considerations of the circuit or matrix.

5. **Numerical Algorithms**: Different classes of circuits perform better with a suitable choice of a matrix factorization algorithm. Our FPGA design may benefit from new ideas for factoring the circuit matrix.

6. **SPICE Algorithms**: Conventional SPICE simulations often perform needless computation across multiple iterations that is not strictly necessary for an acceptable result. In many cases, it is possible to rapidly advance simulation time by exploiting parallelism (e.g. speculation, concurrent alternatives). Our paral-
lel FPGA system can enjoy the benefits of exposing parallelism in the iteration management algorithms.

We now roughly organize the evolution of parallel SPICE systems into five ages and identify the appropriate category for the systems considered:

**Age of Starvation (1980-1990):** In the early days of VLSI scaling, the amount of computing resources available was limited. This motivated designs of custom accelerator systems for SPICE that were assembled from discrete components (e.g. [25, 51, 26]). Numerical algorithms for SPICE were still being developed and not directly suitable for parallel evaluation. The systems scavenged parallelism in SPICE using compiled-code approaches and customized hardware. In a compiled-code approach, the framework produces code for an instance of the circuit and compiles this generated code for the target architecture. This exposes instance-specific parallelism in SPICE to the compiler in two primary ways: (1) it enables static optimization of recurring, redundant work in the Model Evaluation phase and (2) it disambiguates memory references for the sparse matrix access. The resulting compiled-code program is capable of only simulating the particular circuit instance. One of the earliest papers [22] on parallel SPICE sketches a design for the sparse matrix-solve phase by extracting the static triangulation graph for the matrix factorization but ignores communication costs. Other studies have also considered extracting the static dataflow graph for matrix factorization using the MVA (Maximal Vectorization Algorithm) approach [23] (Compiled Code). The resulting graph is vectorized onto a Hitachi S810 supercomputer to get a 8.9× speedup with 4 vector units and a modest increase in storage of intermediate results. Vectorization and simpler address calculation are key reasons for this high speedup. Cayenne [24] maps the SPICE computation to a 2-processor VAX system (Compute Organization) running a multiprocessing operating-system VMS but achieves a limited speedup of 1.7×. A custom design for a SPICE accelerator with 2 PEs [25] running a Compiled Code SPICE simulation shows a speedup of 2× over a VAX 8650 operating at ≈2× the frequency. It emphasizes parallelizing addressing and memory lookup operations within the Processing Element (PE) in a manner similar to our customized FPGA designs. Another Com-
piled Code simulator outlined in [26] delivers a speedup of $7.3 \times$ using 6 processors. This approach exploits multiple levels of granularity in the sparse matrix solve and optimizes locking operations in model-evaluation for greater parallelism. However, performance is constrained by the high operand storage costs of applying a compiled-code methodology to the complete simulator on processing architecture with limited memory resources.

**Age of Growth (1991-1995):** As silicon capacity increased, we saw improved parallel SPICE solvers based on vector or distributed-memory architectures (e.g. [27, 31, 33]). These systems were more general-purpose than the custom accelerators. However, they required careful parallel programming, performance tuning and novel scheduling algorithms to manage parallelism. Awsim-3 [51, 28] again uses a Compiled Code approach and a special-purpose system (Compute Organization) with table-lookup (Precision) Model-Evaluation to provide a speedup of $560 \times$ over a Sun 3/60. The Sun 3/60 implements floating-point operations in software which takes tens of cycles/operation. This means that a bulk of the speedup comes from hardware floating-point units in Awsim-3. Another scheme presented in [27] schedules the complete SPICE computation as a homogeneous graph of tasks (Scheduling). It delivers a speedup of $4.5 \times$ on 8 processors. Parallel waveform-relaxation (SPICE Algorithms) is considered in [29] and [30] and demonstrates good speedups using the different simulation algorithm. A novel row-scheduling algorithm (Scheduling) for processing the sparse matrix solve operations is presented in [31] to obtain modest speedups of $3.7 \times$ on 4 processors. A preconditioned Jacobi technique (Numerical Algorithms) for matrix solve is discussed in [32] but achieves speedups of $16.5 \times$ using 64 processors. This is due to the cost of calculating and applying the preconditioner. In [33], the task scheduling algorithm extracts fine-grained tasks (Scheduling) from the complete SPICE computation to achieve an unimpressive speedup of $4.3 \times$ using 8 processors. Transputers have been used for accelerating block-decomposition factorization (Numerical Algorithms) for the sparse matrix solve phase of SPICE in [34], but no speedups have been reported.
Age of the “Killer Micros” (1996-2005): Moore’s law of VLSI scaling had facilitated the ascendance of general-purpose ISA (Instruction Set Architecture) unprocessors. These “killer micros” \cite{52,53} rode the scaling curve across technology generations and delivered higher compute performance while retaining the ISA programming abstraction. Processors now included integrated high-performance floating-point units which enabled them to deliver SPICE performance that was competitive with custom accelerators. As a result, this age is characterized by a lack of many significant breakthroughs in parallel SPICE hardware designs. A notable exception is the mapping of parallel SPICE to an SGI Origin 2000 supercomputer with 40 nodes (MIPS R10K processors) in \cite{35}. The supercomputer (Compute Organization) was able to speedup SPICE for certain specialized benchmarks by $24\times$ using a message-passing description of SPICE.

Age of Plenty (2006-2010): In this age, the uniprocessor era was starting to run out of steam. The cost of retaining the ISA abstraction while delivering frequency improvements was increasing power consumption to unsustainable levels. This meant that it was no longer possible to simply scale frequency or superscalar Instruction-Level Parallelism to deliver higher performance. The microprocessor vendors turned to putting multiple parallel “cores” on a chip and transferred the responsibility of performance improvement to the programmer. Thus, it became important to explicitly expose parallelism to get high performance. A few studies accelerated SPICE by a modest amount on such systems using multi-threading (e.g. \cite{36,38}). This age also saw the rise of the Graphics Processing Units (GPUs) for general-purpose computing (GP-GPUs) which densely packed hundreds of single-precision floating-points on a single chip. A few studies (e.g. \cite{40,42}) have shown great speedups when accelerating the Model-Evaluation phase of SPICE using GPUs (Compute Organization). In \cite{36}, a multi-threaded version of SPICE is developed using coarse-grained PThreads. It achieves a speedup of $5\times$ using 8 SMP (Symmetric Multi-Processors) on a small benchmark set. SILCA \cite{37} delivers good speedup for circuits with parasitic couplings (resistors and capacitors) using a combination of low-rank matrix updates, piecewise-linear approximations and other accuracy tradeoffs (Precision, Numer-
ical Algorithms). It is possible to achieve limited speedups of $1.3\times$ for SPICE shown in [38] using OpenMP pragma annotations to the Model-Evaluation portion of existing SPICE source-code. GPUs can be used to speedup the data-parallel Model-Evaluation phase of SPICE by $50\times$ ([40]) or $32\times$ ([42]) but can accelerate the complete SPICE simulator in tandem with the CPU by $3\times$ for the GPU-CPU system. A speedup of $2.4\times$ can be achieved using speculative execution of multiple timesteps as demonstrated in [39] (SPICE Algorithms). This approach is orthogonal to the technique discussed in this thesis and can be used to extend our speedups further with additional hardware. In [45], the authors show a speedup of $119\times$ using 32 processors with a domain-decomposition approach (Numerical Algorithms) for accelerating Sparse Matrix-Solve phase. This technique breaks up a large circuit matrix into smaller overlapping submatrices and factorizes them independently. This idea, too, is orthogonal to our approach where we can use our FPGA-based solver to accelerate the individual submatrices.

**Age of Efficiency (2010-beyond)** As Moore’s Law starts to hit physical limits of energy and reliability, we must seek novel architectures for organizing computation. The ISA abstraction is power-hungry and wasteful of silicon resources. Multi-core architectures will eventually run into power-density limits at small feature sizes [54]. The simple SIMD model in GP-GPUs is unsuitable for sparse, irregular computation (e.g. Sparse Matrix-Solve). Instead, we must consider energy-efficient, high-density reconfigurable architectures (e.g. FPGAs) for implementing computation. We need to create customized architectures that match the parallel patterns in SPICE to get high performance. Communication bandwidth and latency become first-class concerns in the compilation flow. This thesis shows how to express, exploit and implement the parallelism available in SPICE using custom FPGA architectures.

### 2.3.1 FPGA-based SPICE solvers

FPGAs have been used extensively in custom-computing machines for accelerating a wide variety of compute-intensive applications. However, they have enjoyed limited use for accelerating SPICE due to limited FPGA resources and lack of tools and
methodology for attacking a problem of this magnitude. In this context, FPGAs were first used in a SPICE accelerator as glue logic \cite{31} to implement the VME interface (Versa Module Europa, IEEE 1014-1987 bus standard) and store communication schedules to support a 4-chip Intel i860 system with a Compiled Code approach. The schedules are implemented using an FPGA-based sequencer. The design in \cite{48} used FPGAs to support the VLIW datapath for accelerating SPICE. Due to limited FPGA capacity, the FPGA was connected to discrete Weitek floating-point units and SRAM chips (Compute Organization). A Compiled Code, partial evaluation approach for timing simulation (lower precision than SPICE) using FPGAs was demonstrated in \cite{49} where the processing architecture was customized for a particular SPICE circuit. The compiler generates optimized fixed-point function units (Precision) and schedules memory access for the simulation. Fixed-point computation may be unsuitable for certain simulation scenarios with high-dynamic range of physical quantities (e.g. leakage simulations with picoampere resolutions). Additionally, this approach demands that the physical FPGA be configured for every circuit being simulated. FPGA mapping algorithms are time-consuming and may themselves take hours to days of runtime. Another recent study \cite{50} explores the use of an FPGA-based, SPICE-like simulator for speeding up transient simulations of SPICE circuits expressed as digital signal processing objects (SPICE Algorithms). This approach converts an analytical analog representation of circuit equations into a graph of variable-rate, discretized, streaming operators. The accuracy of the solution depends on the discretization algorithm (Precision) and may require extremely fine timesteps for correct convergence. Moreover, the current implementation operates on 8-bit/16-bit data-types which are insufficient for detailed leakage simulations.

2.3.2 Our FPGA Architecture

We now highlight some key features of our architecture that are inspired by the previous studies or superior to the older approaches as appropriate. Our FPGA-based approach accelerates the SPICE computation while retaining the accuracy of spice3f5. Any optimization that sacrifices accuracy by simplifying the computation (e.g. table-
lookups) or reducing the amount of computation (e.g. low-rank updates) is orthogonal to our approach under identical convergence and truncation requirements. We use a **Compiled Code** approach similar to several previous studies for our design. We capture parallelism available in the SPICE simulator using a high-level, domain-specific framework that supports a compiled-code methodology for optimizing the simulation. We then implement this parallelism on available FPGA resources by choosing how to spread this parallelism spatially across the FPGA fabric (**Compute Organization**) while distributing memory accesses over embedded, distributed memories. We are also able to exploit additional parallelism by building custom architectures that are tailored to unique forms of parallelism specific to each SPICE phase. Previous studies did not change their compute organization to adapt to the parallelism for each of the three phases of SPICE. Our approach considers communication as a first-class concern and designs high-bandwidth, low-latency networks that use either a time-multiplexed approach for Model-Evaluation or a dynamically-routed packet-switched design for the Sparse Matrix-Solve. Previous studies either used expensive crossbars or slow packet-switched networks to support communication traffic. We distribute the computation for locality and tune network bandwidth when required to balance compute and communication times.
Chapter 3

Model-Evaluation

In Chapter 2 (Section 2.1), we introduced Model Evaluation as an easily parallelizable phase of the SPICE circuit simulator. It is characterized by abundant data parallelism where each device can be evaluated concurrently in a data-independent manner. In this chapter, we first identify additional computational characteristics of the Model-Evaluation phase that are important for an efficient parallel mapping. We show how to compile the non-linear differential equations describing SPICE device models using a high-level, domain-specific framework based on Verilog-AMS. We then sketch a hypothetical fully spatial design that distributes the complete Model-Evaluation computation in space as a configured circuit to achieve the highest throughput. We develop spatial organizations that can be realized on a single FPGA using statically-scheduled time-multiplexing of FPGA resources. This allows us to use less area than the fully spatial design while still achieving high performance. Our automated compilation and tuning approach can scale the implementation to larger system sizes when they become available. We quantify the impact of loop-unrolling and software pipelining (GraphStep scheduling) optimizations on FPGA costs and performance. We also show how to target other parallel organizations (e.g. multi-core, GPUs, Cell, Niagara) using an automated code-generation and auto-tuning approach. The high-level framework and automated mapping flow allows us to describe the computation at a high-level of abstraction while generating optimized parallel implementations for different parallel architectures. We show speedups up to $23 \times (6.5 \times \text{mean})$ for Double-Precision Model-Evaluation computation when comparing a Virtex-6 LX760 to an
Figure 3.1: Flowchart of a SPICE Simulator with emphasis on Model-Evaluation

Intel Core i7 965 (40–45nm technology). We also demonstrate Single-Precision implementations that deliver speedups of up to $111 \times (31 \times \text{mean})$ for a Virtex-6 LX760, $133 \times (58 \times \text{mean})$ for an NVIDIA GTX285 GPU and $1200 \times (71 \times \text{mean})$ for an ATI Firestream 9270 compared to an Intel Core i7 965.

### 3.1 Structure of Model-Evaluation

The Model-Evaluation phase of SPICE calculates the currents and conductances of all the devices in the circuit. The computed currents and conductances are used to update the matrix $A$ and the vector $\vec{b}$ (in $A\vec{x} = \vec{b}$). At the start, the simulator processes all the devices in the circuit. At subsequent timesteps, *only* the non-linear and time-varying elements change and must be recalculated during the Model-Evaluation phase. The resistors and uncontrolled sources have fixed conductances and do not need to update the matrix in every iteration. A device updates the matrix according to its *stamp* that specifies which entries it defines in the matrix. For an $N$-terminal
device, we need to update at most $N^2$ entries in the matrix. Thus, each device in the
circuit updates a constant number of entries in the matrix corresponding to its node
terminals. We show a 2-terminal diode and 3-terminal transistor stamp in Figure 3.2. 
In this case the two devices share a terminal (D2 for diode and T1 for transistor).
The resulting stamps update the shared matrix entry corresponding to the device 
terminals. We show device equations for a diode in Table 3.4.

For non-linear elements like diodes and transistors, the simulator must search for 
an operating-point using the Newton-Raphson iterations shown in the outer loop of 
Figure 3.1. The conductance of the non-linear and time-varying elements is a function 
of the terminal voltages $\vec{x}$. Since the terminal voltages $\vec{x}$ are computed by the $A\vec{x} = \vec{b}$ 
solve, we need to use Newton-Raphson algorithm to iteratively compute the consistent 
solution vector $\vec{x}$. This requires repeated evaluation of the non-linear model equa-
tions multiple times per timestep. For time-varying components like capacitors and 
inductors, the simulator must recalculate their contributions at each timestep based 
on voltages and charges at several previous timesteps (e.g. Trapezoidal integration). 
This also requires a re-evaluation of the device-model in each timestep.

Each model-evaluation computation only requires local voltages on the device 
terminals as input and is completely data-independent from other devices. This 
embarrassing data parallelism has been exploited extensively in previous studies (see 
Section 3.2). In this chapter, we develop an efficient FPGA implementation that 
exploits this data parallelism and static nature of the Model-Evaluation phase.

For circuits dominated by non-linear transistor devices, the simulator can spend 
almost half its time evaluating the device model.\footnote{1see “no parasitics” case in Table 3.1 we generated datapoints in this table by running spice3f5 
on an Intel Core i7 using Simucad memory benchmarks [55]} For circuits dominated by linear 
parasitics (e.g. parasitic capacitances), simulation time may be dominated by the 
Matrix-Solve phase. Since we are ultimately interested in accelerating both Model-
Evaluation phase and Matrix-Solve phase, it is important to understand how far we 
can improve Model-Evaluation runtimes even in these cases where it is currently not 
the dominant percentage of runtime. Furthermore, as transistor devices shrink in

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\footnote{1see “no parasitics” case in Table 3.1 we generated datapoints in this table by running spice3f5 on an Intel Core i7 using Simucad memory benchmarks [55]}
feature-size, the complexity of the device models required to simulate them correctly grows over time (see [56]). Newer device models often have complexity 4–5× that of the classic bsim3 model [57] (e.g. compare psp [58] and bsim3 models in Table 3.3). This motivates the need to accelerate the Model-Evaluation phase to avoid paying a large modeling cost for future sub-micron netlists. For example, when Model-Evaluation time increases by 5×, the no parasitic ram8k will spend 80% of its time in Model-Evaluation phase, and the parasitic case will spend 36%.

3.1.1 Parallelism Potential

We now enumerate the potential of parallelizing the Model-Evaluation phase.

Data Parallelism: Each individual model evaluation (e.g. for each transistor) within a timestep is completely independent. With ideal hardware (no resource constraints or data distribution latencies), we can process each device in parallel as previously shown in Figure 2.13(a) of Section 2.2.4. Thus, evaluation time will be
<table>
<thead>
<tr>
<th>Benchmark Circuits (bsim3)</th>
<th>Model Evaluation (seconds)</th>
<th>Matrix Solve (seconds)</th>
<th>Model Evaluation (Percent (%))</th>
</tr>
</thead>
<tbody>
<tr>
<td>no parasitics</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>ram2k</td>
<td>55</td>
<td>10</td>
<td>84</td>
</tr>
<tr>
<td>ram8k</td>
<td>237</td>
<td>87</td>
<td>73</td>
</tr>
<tr>
<td>ram64k</td>
<td>2005</td>
<td>1082</td>
<td>64</td>
</tr>
<tr>
<td>with parasitics</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>ram2k</td>
<td>69</td>
<td>149</td>
<td>31</td>
</tr>
<tr>
<td>ram8k</td>
<td>300</td>
<td>2395</td>
<td>11</td>
</tr>
<tr>
<td>ram64k</td>
<td>2597</td>
<td>99487</td>
<td>3</td>
</tr>
</tbody>
</table>

Table 3.1: spice3f5 Runtime Distribution (Core i7 965)

independent of size of the circuit.

**Pipeline Parallelism:** Model evaluation operations can be represented as an acyclic feed-forward dataflow graph (DAG) with nodes representing operations and edges representing dependencies between the operations. This allows us to balance the delays on the different paths in the graph through suitable pipelining. Aggressive pipelining allows us to start multiple device evaluations before the first one is finished thereby improving performance.

**Instruction-Level Parallelism:** The dataflow graphs representing the device equations have Instruction-Level Parallelism (ILP) that can be exploited by concurrent floating-point units on the parallel hardware. We can further increase this ILP by performing loop-unrolling and software-pipelining optimizations on the dataflow graphs.

### 3.1.2 Specialization Potential

We summarize some observations that enable specialization of the Model-Evaluation phase for our parallel design.

- **Fixed-Sized Workload:** The Model-Evaluation phase processes all devices in the circuit in each timestep. While certain simulators can bypass quiescent devices for faster operation, in our design we do not consider this optimization and only handle a fixed-size workload for easier parallelization. Our statically-scheduled
design cannot handle dynamic changes in the workload that may arise from bypassing quiescent devices. Parallelizing dynamically-varying workloads requires a dynamically-scheduled hardware design which may deliver worse performance [59].

- **Early Bound Graph**: The Model-Evaluation compute graphs are known entirely in advance as part of the input netlist. The circuit structure being simulated stays constant throughout the simulation. The conductance and current values of the circuit are the only quantities that change during the simulation.

- **Limited Diversity of Graphs**: Within a simulation, there may be very few unique device models active. (*e.g.* typically all transistors in a circuit will use same `bsim3` model). This allows us to reuse the same parallel architecture and schedule it across multiple device evaluations.

- **Parametrized Reuse of Graphs**: Individual device instances are customized using parameters. Typically the CMOS process determines most of these parameters leaving a handful of parameters which vary from device to device (*e.g.* Width, Length of a transistor). All the constant parameters specified in the process technology can be constant-folded (computation can be simplified using the knowledge of constant values) for faster operation (See Table 3.3).

### 3.2 Related Work

We now review some previous studies that attempted to parallelize the Model-Evaluation phase of SPICE and adopt the category classification introduced in Section 2.3. We tabulate and highlight key features of these studies in Table 3.2.

An early study [25] parallelizes the Model-Evaluation computation using customized vector operations on a custom co-processor board (Compute Organization) but provides a limited speedup of 2×. The computation in the Model-Evaluation phase also contains many non-vectorizable operations that will limit achievable speedups. Our own studies with vectorization on the Intel multi-core architecture (SSE3) show no benefit for Double-Precision evaluation and only modest benefits for Single-Precision (see Section 3.8.1). Thus, vectorization is not the most suitable way
<table>
<thead>
<tr>
<th>Year</th>
<th>Ref</th>
<th>Parallel Hardware</th>
<th>Style</th>
<th>Sequential Hardware</th>
<th>PEs</th>
<th>Speedup</th>
<th>Notes</th>
</tr>
</thead>
<tbody>
<tr>
<td>1992</td>
<td>[28]</td>
<td>Custom Hardware (AWSIM-3)</td>
<td>VLIW</td>
<td>Sun-3/60</td>
<td>-</td>
<td>2500</td>
<td>Table-lookup approximation</td>
</tr>
<tr>
<td>1997</td>
<td>[49]</td>
<td>Altera Flex10K FPGA</td>
<td>VLIW</td>
<td>SPARC station</td>
<td>-</td>
<td>27.7(^2)</td>
<td>Fixed-Point Timing simulation</td>
</tr>
<tr>
<td>1995</td>
<td>[48]</td>
<td>Marc-1 board (TINA)</td>
<td>VLIW</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>No speedups reported</td>
</tr>
<tr>
<td>2002</td>
<td>[36]</td>
<td>Hitachi 9000 N4000</td>
<td>SMP</td>
<td>1 thread</td>
<td>8</td>
<td>5.7</td>
<td>-</td>
</tr>
<tr>
<td>2007</td>
<td>[38]</td>
<td>UltraSPARC 3</td>
<td>SMP</td>
<td>1 thread</td>
<td>4</td>
<td>1.35</td>
<td>-</td>
</tr>
<tr>
<td>2009</td>
<td>[40]</td>
<td>ATI FireStream 9170</td>
<td>GPU</td>
<td>4-core AMD Phenom</td>
<td>320</td>
<td>50</td>
<td>-</td>
</tr>
<tr>
<td>2009</td>
<td>[42]</td>
<td>NVIDIA 8800 GTS</td>
<td>GPU</td>
<td>4-core Intel Core2</td>
<td>128</td>
<td>32</td>
<td>Single-Precision</td>
</tr>
</tbody>
</table>

Table 3.2: Parallel Model-Evaluation Studies

To exploit the parallelism in Model-Evaluation.

Awsim-3 [28] is a custom hardware accelerator (Compute Organization) using table-lookup approximation (Precision) and Compiled Code technique to deliver a speedup of 2500\(\times\) over a Sun 3/60. The Sun 3/60 implements floating-point operations in software which takes 10s of cycles per operation. This slows down the floating-point component of the processing which explains a part of the high speedups. Table-lookup approximations are computationally simpler to implement since they only require memory operations and a few floating-point operations per lookup. They are also easier to parallelize by distributing memory lookups across parallel memory blocks. However, they sacrifice SPICE accuracy for this faster operation. Our single-FPGA implementation implements the full analytic equations on a VLIW scheduled architecture to deliver high speedups without compromising simulation accuracy.

\(^2\)27.7\(\times\) reduction in instruction count does not necessarily translate to speedup.
TINA [48] (inspired from Awsim-3) is the first FPGA-based implementation of SPICE Model-Evaluation. It used the Marc-1 reconfigurable board with 23 XC4005 FPGAs coupled to discrete Weitek FPUs to implement Model-Evaluation using a table-lookup approach. Unfortunately, speedup figures were never unpublished. Another FPGA-based approach in [49], a fixed-point implementation (Precision) of Model-Evaluation computation is implemented using partial evaluation of the Compiled Code timing simulation to deliver a $27.7 \times$ reduction in instructions when comparing an Altera EPK10K50 with a SPARCstation.

We now review some recent literature that parallelizes SPICE Model-Evaluation. Manual multi-threading (PThreads) [36] and automated data-parallelization (OpenMP pragmas) [38] provide limited benefits of $5.7 \times$ and $1.35 \times$ respectively over Symmetric Multi-processing systems. More recent work has focused on parallelizing Model-Evaluation on GPUs (Compute Organization). The use of GPUs for accelerating bsim3 models was first explored in [40] (double-precision) and subsequently in [42] (single-precision). As shown in [40], it is possible to achieve speedups of $10–50 \times$ over a quad-core AMD CPU when using an AMD Firestream 9170 GPU (512 processors). Similarly, we can see in [42] that it is possible to obtain speedups of $32–40 \times$ over a quad-core Intel CPU when using an NVIDIA 8800 GTS GPU (128 processors). Our FPGA implementation uses a different parallelization approach (VLIW) and high on-chip communication and memory bandwidth to deliver speedups in the same ballpark for Model-Evaluation using a single FPGA with a slight compromise in accuracy for Single-Precision implementation (Precision). Additionally, our approach shows a speedup of $18 \times$ for the bsim3 model over a dual-core Intel Xeon processor when using an NVIDIA 9600 GT GPU with 64 processors and a speedup of $8 \times$ when using a Virtex-5 LX330 FPGA. We present additional details in Section 3.7. The GPU implementations in [40, 42] must copy matrix and vector data back-and-forth from the host CPU as the SIMD model is unsuitable for accelerating the Sparse Matrix-Solve phase of SPICE. We later show in Chapter 4 how to accelerate the Sparse Matrix-Solve phase on the same FPGA itself to eliminate this data transfer overhead.
Table 3.3: Verilog-AMS Compiler Optimized Instruction Counts

<table>
<thead>
<tr>
<th>Model</th>
<th>Instruction Distribution (Unoptimized on left and Optimized right)</th>
<th>Add</th>
<th>Multiply</th>
<th>Divide</th>
<th>Sqrt.</th>
<th>Exp.</th>
<th>Log.</th>
<th>Rest</th>
</tr>
</thead>
<tbody>
<tr>
<td>bjt</td>
<td></td>
<td>39</td>
<td>22</td>
<td>65</td>
<td>30</td>
<td>17</td>
<td>3</td>
<td>0</td>
</tr>
<tr>
<td>diode</td>
<td></td>
<td>7</td>
<td>7</td>
<td>7</td>
<td>5</td>
<td>4</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>jfet</td>
<td></td>
<td>31</td>
<td>13</td>
<td>73</td>
<td>31</td>
<td>14</td>
<td>2</td>
<td>0</td>
</tr>
<tr>
<td>mos1</td>
<td></td>
<td>86</td>
<td>24</td>
<td>102</td>
<td>36</td>
<td>36</td>
<td>7</td>
<td>5</td>
</tr>
<tr>
<td>vbic</td>
<td></td>
<td>283</td>
<td>36</td>
<td>347</td>
<td>43</td>
<td>123</td>
<td>18</td>
<td>17</td>
</tr>
<tr>
<td>mos3</td>
<td></td>
<td>89</td>
<td>46</td>
<td>177</td>
<td>82</td>
<td>52</td>
<td>20</td>
<td>11</td>
</tr>
<tr>
<td>hbt</td>
<td></td>
<td>3202</td>
<td>112</td>
<td>4245</td>
<td>57</td>
<td>1101</td>
<td>51</td>
<td>1</td>
</tr>
<tr>
<td>bsim4</td>
<td></td>
<td>770</td>
<td>222</td>
<td>1479</td>
<td>286</td>
<td>482</td>
<td>85</td>
<td>48</td>
</tr>
<tr>
<td>bsim3</td>
<td></td>
<td>585</td>
<td>281</td>
<td>995</td>
<td>629</td>
<td>188</td>
<td>120</td>
<td>23</td>
</tr>
<tr>
<td>mextram</td>
<td></td>
<td>1303</td>
<td>675</td>
<td>3041</td>
<td>1626</td>
<td>758</td>
<td>397</td>
<td>23</td>
</tr>
<tr>
<td>psp</td>
<td></td>
<td>5363</td>
<td>1345</td>
<td>7136</td>
<td>2319</td>
<td>1378</td>
<td>247</td>
<td>350</td>
</tr>
</tbody>
</table>

Column Rest includes MUX, BOOL and INT operations

3.3 Verilog-AMS Compiler

Modern SPICE simulators accept a wide variety of device models that cater to different designer requirements of accuracy and performance. These device models are released as simulator independent Verilog-AMS descriptions [60]. We use open-source Verilog-AMS descriptions of a variety of devices available from Simucad [55]. We developed a Verilog-AMS compiler that supports a subset of the Verilog-AMS language for device models similar to [61]. We compile the device model equations into a flexible intermediate representation that allows us to perform analysis, optimization and code-generation for different architectures easily. The compiler currently performs simple dead-code elimination, mux-conversion, constant-folding, identity simplification and common-subexpression elimination optimizations. Our compiler generates a generic feed-forward dataflow graph of the computation that is processed by architecture-specific backend tools. The unoptimized and optimized instruction counts and operation distribution for different device models is shown in Table 3.3. This flow allows us to automate the compilation of the future device models as well as recompilation of existing models due to updates without requiring manual inter-
\[
I = i_s \cdot (e^{v_j} - 1) \\
G = \frac{di}{dV} = i_s \cdot \frac{1}{v_j} \cdot e^{v_j}
\]

\(v_j = \) junction voltage, \(i_s = \) saturation current

Table 3.4: Diode Dataflow Graph: Equations (top), Unoptimized (left), Optimized (right)

In Table 3.4 we show the device equations for a diode and the corresponding feedforward dataflow graph for those equations. Our Verilog-AMS parser generates the unoptimized graph from the equations which is then optimized into a smaller graph through constant-folding. Our compiler is able to aggressively eliminate boolean and logical operations when using the knowledge of constant input parameters. For larger graphs e.g. bsim4, the compiler optimizations reduce the number of nodes in the graph by \(\approx 7 \times\) (when considering only floating-point operations it is \(4 \times\)).

3.4 Fully-Spatial Implementation

A spatial circuit implementation of computation is a straightforward embodiment of a dataflow graph on an FPGA. Such a circuit contains physical operators for every instruction in the dataflow graph and uses physical wires to implement dependencies between the instructions. These operators can evaluate in parallel and communicate
<table>
<thead>
<tr>
<th>Device Models</th>
<th>Total Speedup</th>
<th>FPGAs Required</th>
<th>Speedup per FPGA</th>
</tr>
</thead>
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<tr>
<td></td>
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<td>No IO Limits</td>
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<tr>
<td>psp</td>
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<td>38</td>
<td>155</td>
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</table>

Table 3.5: Estimated Speedup (vs. Intel Core i7 964) and FPGA costs (Virtex6 LX760) of Multi-FPGA Designs

Figure 3.3: A Spatial Circuit Implementation of Diode Equations
(With pipelined registers along wires with arbitrary delays for illustration purpose)
results directly using the programmable FPGA interconnect. Furthermore, if the computation is data parallel, we can exploit pipeline parallelism by adding a suitable number of registers along the wires to balance dataflow. This will then permit us to start a new evaluation of the dataflow graph in each cycle and deliver results of the computation after the pipeline latency of the graph. This pipelined, spatial circuit implementation of data-parallel computation will deliver the highest possible performance for our Model-Evaluation computation. In contrast, a conventional von-Neumann architecture (e.g. Intel CPUs) will implement this computation by fetching a binary representation of computation stored in memory. The binary implicitly encodes the dataflow structure using a sequence of instructions that communicate results using registers (i.e. memory). The dataflow parallelism hidden in this implicit encoding must be rediscovered by the von-Neumann architecture in hardware often limiting the amount of parallelism that can be exploited from the dataflow graph.

We now consider three different models for estimating the benefits and costs of the multi-FPGA spatial design. We summarize the total speedup, total FPGAs and speedup per FPGA for all devices in Table 3.5.

**Ideal Mapping** We can imagine implementing the data-parallel operations in Model-Evaluation as a pipelined dataflow circuit on the FPGA as shown in Figure 3.3. If cost is not a concern, this approach provides up to two orders of magnitude speedup over an implementation using Intel Core i7 965 microprocessor when using a Xilinx Virtex-6 LX760 FPGA (see Table 3.5). We compute a lower-bound on the number of FPGAs required to implement the dataflow graph based on total operator area (ignoring FPGA external IO limitations and pipelining area costs). This model provides a lower-bound on cost and an upper-bound on the speedup possible with the spatial approach. For the designs that fit in a single FPGA, this model only needs to be refined with pipelining costs and can avoid the complexities of the multi-FPGA distribution. A single-FPGA, fully-spatial implementation of all devices will be eventually possible with the increasing FPGA densities made possible by Moore’s Law. From Table 3.5 the bsim3 model currently requires only 3 Virtex-6 LX 760 FPGAs to fit. This means an FPGA that is $4 \times$ denser will fit the complete device evaluation
graph. This FPGA will become possible two technology nodes into the future at 22nm (Virtex-6 is manufactured at the 40nm technology node).

**Virtual Wires:** When we distribute the dataflow graph across multiple FPGAs, we must consider the cost of accommodating the graph edges (i.e. interconnect requirements) over external FPGA pins. If the number of edges is more than available IO, we must share a physical IO pin across multiple edges of the graph. The idea of time-multiplexing physical chip IO was introduced in the context of prototyping ASIC designs using FPGAs in [62]. We estimate the speedup possible and the number of FPGAs required by time-multiplexing the communication between the FPGAs over external FPGA IO using Virtual Wires. In this case, we compute the sequentialization introduced by the time-multiplexing of communication and reduce ideal speedup by that factor. We use VPR (Versatile Place and Route) [63] to route inter-FPGA communication for a 2D bidirectional mesh organization of the multi-FPGA system shown in Figure 3.4. VPR is originally designed to place and route logic netlists on a single-chip FPGA fabric. For our experiments, we develop a multi-chip implementation model for floating-point netlists and reuse the VPR algorithms for our context. The IO serialization lowerbound is $\lceil \text{channelwidth} / 18 \rceil$ since we estimate that we can fit 18 double-precision channels (graph edges) operating at 250MHz on the external FPGA pins of a Virtex-6 LX760 FPGA. Each of the 1200 external IO pins is capable of 960 MHz operation ($1200 \text{ pins} \cdot 960 \text{ MHz} = 18 \text{ channels} \cdot 64 \text{ bits} \cdot 250 \text{ MHz} \cdot 4 \text{ directions}$).

**Multi-FPGA Fully-Spatial Mesh:** We are ultimately interested in estimating the number of FPGAs required to implement the Model-Evaluation phase of SPICE in a fully-spatial manner. We want to avoid the serialization bottleneck of Virtual Wires while providing a tighter estimate than the lower-bound provided by the Ideal Mapping. When we distribute the graph across a larger number of FPGAs, the amount of IO that must be routed over each FPGA IO will decrease. In this case, we increase system size until the communication requirements can be satisfied with the available FPGA IO capacity with no serialization or time-multiplexing need. We show such a spatial distribution in Figure 3.4. As we just calculated, a $\text{channelwidth} < 18$ meets the bandwidth constraints of the largest Xilinx FPGA. Again, we use VPR
to route inter-FPGA communication for the multi-FPGA model shown in Figure 3.4 with Switch↔Logic communication limits (shown as IO in Figure 3.4) and estimate the minimum number of FPGAs required for a channelwidth-limited mapping.

Performance and design cost are dictated purely by our ability (or inability) to move data across external pins. Hence, we are motivated to consider affordable single-FPGA designs for the problem and avoid external IO entirely.

3.5 VLIW FPGA Architecture

We are interested in designing a single-chip FPGA system capable of accelerating the complete SPICE simulator. In the previous section, we estimated the maximum speedups possible for the Model-Evaluation phase of SPICE without any cost constraints. In this section, we show how to design a cost-effective, single-FPGA solution to deliver good speedups for the Model-Evaluation computation.

In Table 3.5 we observe that some device models require hundreds of FPGAs to
realize the Model-Evaluation computation for that device. It is evident that if we are to implement this computation on a single-FPGA we will have to virtualize (time share) the computation over limited FPGA capacity. Conventional von-Neumann processors (e.g. Intel CPUs) provide a virtualized instruction-set architecture that permits reuse of limited ALU capacity (e.g. a few dedicated floating-point, integer units) across large programs that are stored compactly in a memory. To exploit instruction-level parallelism (ILP) from the computation, the processors attempt to rediscover the dataflow structure of the compute graph using expensive dynamic hardware techniques (e.g. Tomasulu [64]). Furthermore, they attempt to match an application memory-access pattern by speculatively loading and unloading data items into the caches. In Table 3.3 we note that the computation involves certain elementary floating-point operations that are not directly supported in the form of a dedicated unit in the processor’s ALU of an Intel Core i7 965. This means that the elementary functions require several cycles to evaluate on the processor. However, for SPICE Model-Evaluation graphs, we precisely know the dataflow structure of the device equations and also have information about the data-access patterns before execution. A programmable substrate like an FPGA can even be configured to implement the elementary functions directly in hardware. This suggests we may be able to accelerate the computation using a superior architecture that exploits application knowledge and patterns in the computation for an optimized solution. We build an architecture with the proper operator balance and a network which implements the data access patterns in the dataflow graph.

We develop a customized statically-scheduled VLIW (Very Large Instruction Word) architecture [65] that is tailored for the Model-Evaluation computation. The VLIW architecture consists of a heterogeneous collection of floating-point operators coupled to high-bandwidth local memories and interconnected through a time-multiplexed communication network (see Figure 3.7). We choose the floating-point operator balance to match the compute requirements in the dataflow graph. We explicitly distribute data across the local memories to optimize concurrent access as required by the computation. The distribution also exploits locality in the communication to
minimize network bandwidth.

### 3.5.1 Operator Provisioning

Model-evaluation graphs contain a diverse set of floating-point operators such as adds, multiplies, divides, square-roots, exponentials and logarithms as shown in Table 3.3. Not all operators are used equally. We choose an operator mix proportional to the frequency of their use since spatial implementations of floating-point operators can be quite expensive. For example, in Figure 3.5 we provision multiple adders and multipliers and just one each of the remaining operators within a Processing Element (PE). The FPGA consists of multiple tiles of such PEs. Different device models will have different operator distributions and will require suitable operator mixes. We characterize the bisection bandwidth of our network using a tunable Rent parameter $p$ from Rent’s rule $IO = c \cdot N^p$ [66, 67]. Here $IO$=bisection bandwidth, $N$=number of operators, $c$=channels in the network. For our experiments we consider multiple topologies with $0 < p < 1$ with $c = 1$ as shown in Figure 3.6. A network with a $p = 0$ provides bisection bandwidth similar to a bus with small area requirement while a network with a $p = 1$ provides bisection bandwidth of a crossbar with a larger area.
Figure 3.6: Butterfly Fat Tree Interconnect Examples with three Rent parameters (p)

Figure 3.7: VLIW Processing Element for SPICE Model Evaluation
requirement. We match communication requirements of the application by picking the appropriate $p$ (balance bandwidth benefits with area requirement) through an automated design-space exploration (See Section 3.7.4).

### 3.5.2 Static Scheduling

The Model-Evaluation computation in SPICE is known well in advance of the execution of the simulator. The device equations are described in Verilog-AMS and are available as feed-forward dataflow graphs. Since this graph structure is known, we can statically schedule our FPGA architecture to ensure high utilization of limited resources. It is indeed possible to evaluate this graph using a dynamically-scheduled hardware architecture, but that reduces performance by $2–3\times$ [59]. This is because the logic to dynamically mediate access to shared resources at runtime costs area and introduces additional latency into the execution. Our static scheduling approach eliminates this unnecessary overhead by avoiding any runtime decision-making. Furthermore, the VLIW configuration can be efficiently stored in on-chip memories.

The scheduler generates a VLIW configuration [68] for the execution that contains precomputed datapath, memory and switch controls (see Figure 3.7). A VLIW instruction for the PE consists of read/write addresses for the input and output memories along with multiplexer select signals for the datapath. The time-multiplexed
<table>
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</table>

Table 3.6: Naive Schedule Performance

switch also contains configuration instructions that provide routing information to schedule communication between the input and output ports. We show how a simple dataflow graph may be scheduled on a simple VLIW architecture in Figure 3.8. The nodes of the graph are assigned to the appropriate PE while communication between the nodes is handled by the appropriate switches in the network. The static scheduler provides a cycle-by-cycle configuration for the system represented by the occupied boxes in the figure. The unoccupied boxes represent idle PEs or switches. The efficiency of the design can be measured by the amount of idle time. Assuming this naïve scheduling technique, we can compare the performance of the FPGA with an Intel Core i7 CPU in Table 3.6. We can see that this simple approach can be worse than the CPU implementation by as much as $50\times$. The long latency of the floating-point operators and time-multiplexed routing fabric combined with low throughput evaluation of the devices results in poor FPGA performance. We will now investigate techniques for improving utilization of the FPGA hardware for high throughput evaluation of a large number of devices.

### 3.5.3 Scheduling Techniques

We now consider strategies for reducing the amount of idle time in the static schedule. The parallelism profile of Model-Evaluation graphs is shown in Figure 3.9. We note that 20–50% of the graph operations can be issued in the first few steps of the graph while the remaining operations must execute sequentially along the long tail. A naïve
schedule of this graph as shown in Figure 3.8 will leave many idle slots leading to poor utilization of resources. In fact as we saw in Table 3.6, the performance of this unoptimized schedule is worse than a processor implementation. A key challenge for the scheduler is to expose work to use these slots productively. We consider two strategies for efficiently scheduling our VLIW architecture: (1) Loop Unrolling (2) Software Pipelining with GraphStep Scheduling.

3.5.3.1 Conventional Loop Unrolling

When scheduling single loop iterations on fully-pipelined hardware, the total number of active pipeline stages doing useful work may be limited. For Model Evaluation, each iteration evaluates an individual non-linear device in the circuit. We can create additional work for the scheduler to fill these empty pipeline slots by unrolling multiple iterations of the loop. Loop Unrolling on Model-Evaluation graphs is possible with no increase in the logical critical path of the logical graph since iterations are independent of each other. This allows the scheduler to get better utilization of provisioned hardware resources. The per-iteration efficiency gains more than compensates for the slight increase in scheduling latency of the mapped graph (due to congestion and use of greedy scheduling algorithms). For example, in Figure 3.10, the latency to get the output of a single iteration is 8 cycles. After unrolling 3 iterations, the total

Figure 3.9: Dataflow Graph Profile for Model-Evaluation
latency increases to 12 cycles, but the average per-iteration latency drops to 4 cycles. However, these efficiency gains come at the expense of increased memory cost for storing intermediate state and instruction context. We now need to store 12 VLIW configuration instructions instead of just 6 for the single-iteration case. Also, the intermediate state requirements increase from 3 registers to 9 registers. For 100 total iterations of example graph, this unrolled design will require $100 \times 4 = 400$ cycles. We empirically determine the extent of the unroll using an auto-tuning approach that provides the most judicious use of resources (Section 3.6).

3.5.3.2 Software Pipelining with GraphStep Scheduling

Software Pipelining [69, 70] improves the per-iteration performance by initiating execution of successive loop iterations at a rate faster than their individual execution latencies (which in our case is the resource-constrained initiation interval) without requiring any unrolling. It overlaps execution of different portions of the loop in a single repetitive macro-cycle. The benefit of a software-pipelined schedule is that
a single schedule is valid for all iterations thereby saving instruction storage costs.
It does increase the amount of intermediate state for instructions communicating
across macro-cycle boundaries as well as latency of evaluation. For example, in Figure 3.11 we need only 2 cycles to schedule all the instructions and communication
between instructions in a macro-cycle (throughput is 1 result every 2 cycles) while
the result of the first iteration is available after 5 macro-cycles (latency is 10 cycles).
For 100 iterations of the example graph, our software pipelined design will require
\((100+4) \times 2 = 208\) cycles which is a speedup of almost 2× over loop unrolling example
(note that the +4 accounts for the initial macro-cycles required to fill the scheduled
pipeline). We use the GraphStep scheduling algorithm for implementing the software
pipelining optimization. It is inspired by the GraphStep system architecture [71]
developed for efficient parallel processing of sparse graphs.

In Figure 3.12, we show how the performance of Loop-Unrolling (unroll=10) compares with that of Software Pipelining as a function of device count for a VLIW
architecture with 16 operators and $p = 0$. We see that Software Pipelining scheduling outperforms Loop-Unrolling when the number of devices exceeds $\approx 100$ devices. When combining both optimizations together we can improve performance by $2.7 \times$ at $10^4$ devices. We get this additional performance boost through better balanced distribution of operators and fewer idle cycles per operator. We analyze performance in greater detail in Section 3.6.

With loop-unrolling we can generate sufficient work to better overlap communication latency and fill the pipelines of long-latency floating-point operators like divide (57 cycles).

For our scheduling problem:

- We must typically evaluate a large number of devices compared to depth of the graph (number of instructions along the critical path). This allows us to pipeline the graph deeply for high throughput while ensuring high utilization.

- We have access to several on-chip distributed FPGA memories to store intermediate state. This capacity allows us to store the increased intermediate state generated from loop-unrolling and software pipelining. We show the memory requirements for the different cases in Figure 3.13. As expected, we observe a proportional increase in memory requirements when we increase the unroll factor. We also observe a $1.9 \times$ increase ($1.3-4 \times$) in storage costs for accommodating the retiming registers for GraphStep scheduling. When implementing this on a Xilinx FPGA, we are limited by the amount of onchip memory capacity available in relation to logic resources on
the FPGA. The increase in memory requirement does not affect our performance as long as the memory-logic balance is preserved. Our design-space exploration 3.7.4 will automatically pick the best configuration to maximize performance.

- We have to generate a schedule for the switching network in addition to compute scheduling. Without software pipelining, we have to schedule the PE and the switching network in an interdependent fashion. Software Pipelining with GraphStep scheduling allows separate scheduling of compute and switching network allowing overlapped execution.

- We are able to provision interconnect resources as necessary to meet the demands of the application. This is important to avoid communication bandwidth bottlenecks for large graphs.

For data-independent loop-iterations, conventional modulo scheduling [72] will generate a schedule while obeying intra-iteration dependencies. Based on our observations, we propose a simplified scheduler described here:

- We distribute instructions to different operators with load-balancing and schedule all instructions on an operator without precedence constraints. This is because Software Pipelining allows us to stagger the schedule for a pipelined path across multiple macro-cycles. This gives us scheduling freedom in each macro-cycle to tightly schedule as many pipelined operator slots as possible by ignoring precedence constraints. This improves operator packing density at the expense of latency (additional macro-cycles).
Within a macro-cycle, all instructions are processed on the operators and all instruction dependencies are routed on the communication network in parallel with each other. Typically Software Pipelining is used to distribute instructions (graph nodes) across multiple macro-cycles. In our case, since we must schedule both compute and communicate, we can treat them uniformly for scheduling.

We schedule data movement between operators concurrently but independently from the computation. This allows us to overlap the compute and communicate scheduled cycles thereby improving performance by $2 \times$ in the ideal case. However, this adds an extra macro-cycle of latency before the dependent instructions can see their inputs.

We levelize the instructions in the compute graph based on an ASAP ordering of the instructions within an iteration. We retime the inputs to each instruction based on these levels and ensure they receive inputs from the correct iteration. These are called rotating registers in the schedule [69].

These simplifications allow us to densely pack both the floating-point operators and the communication network between these operators with high efficiency.

### 3.6 Methodology and Performance Analysis

In this section, we discuss our experimental methodology and explain our results for double-precision implementation of Model-Evaluation.

#### 3.6.1 Toolflow

We setup a shared toolflow for the two FPGA static scheduling strategies as shown in Figure 3.14. We start by first deciding system size (i.e. number of floating-point operators) and partitioning the nodes based on locality using a high-quality partitioner MLPart [73]. At this point we do not partition the nodes according to operation type. Next, we provision the number of hardware operators of each type according to instruction frequency and allocate them to partitions using need-proportional distribution [74, 75]. Each partition can process operations of a single operator type. We
then reassign nodes paired with invalid operators to the nearest valid operator that is least occupied. The range of FPGA configurations possible is large and we use an auto-tuner to automatically select the best configuration.

- For loop unrolling, we provide an unrolled graph to the partitioner/placer. Once instructions have been placed on proper operators, we then use a greedy list scheduler to assign those instructions to schedule slots on the operator. We use a priority function that prefers nodes along the circuit critical path. We schedule communication between the nodes using a greedy time-multiplexed router that uses A* routing. We developed this scheduler and router as part of the Graph Machine project [71, 59, 76].

- For the GraphStep scheduler, we separately schedule computation and communication. The compute scheduler simply assigns all instructions to consecutive scheduling slots on the fully-pipelined hardware operator. The communication scheduler routes every edge with A* routing without any precedence constraints.

### 3.6.2 FPGA Hardware

We use spatial implementations of individual floating-point *add*, *multiply*, *divide* and *square-root* operators from the Xilinx Floating-Point library in CoreGen [77, 78]. For the *exp* and *log* operators we use FPLibrary from the Arénaire [79] group. Neither of these implementations support denormalized (subnormal) numbers. We use the Xilinx Virtex-5 LX330T and Virtex-6 LX760 for our experiments. We limit our implementations to fit on a single-chip and use only on-chip memory resources for storing intermediate results. The time-multiplexed switches are a collection of multiplexers whose select bits are generated by a configuration context memory on...
<table>
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Table 3.7: Model-Evaluation FPGA Cost Model (Virtex-6 LX760)

each cycle. We pipeline the wires between the switches and between the floating-point operator and the coupled-memories for high-performance. You can find additional details of our time-multiplexed switches in [59]. We synthesize and implement a sample double-precision, 8-operator design for the bsim3 model on a Xilinx Virtex-5 device using Synplify Pro 9.6.1 and Xilinx ISE 10.1. We provide placement and timing constraints to the backend tools and attain a frequency of 200 MHz (See Table 3.7). Aggressive pipelining of \( \exp \) and \( \log \) operators should enable higher rates.

### 3.6.3 Optimized Processor Baseline

We compile Verilog-AMS models into loop-unrolled, multi-threaded C-code for our sequential baseline comparison. We measure sequential performance on two machines: (1) a dual-core 65nm, 3 GHz Intel Xeon 5160 processor with a 4MB shared L2 cache and 16GB main memory running 64-bit Debian Linux and (2) a quad-core 45nm, hyper-threaded 2.67 GHz Intel Core i7 965 with an 8MB shared L3 cache and 12GB memory. We use gcc-4.4.3 (-O3) with either the GNU libm math library or Intel MKL library to compile device models. We use PAPI 4.0.0 [18] performance counters to measure runtimes and report runtime averaged across multiple device evaluations.

We show peak double-precision floating-point capacities of these devices in Table 3.8. From the table, we note that both the CPU and FPGA architectures have
comparable peak floating-point peaks despite the FPGA having more than an order of magnitude lower clock frequency than the CPU (200 MHz vs 3 GHz). We now show how the FPGA architecture makes better use of its peak than the CPU to deliver higher speedups.

### 3.6.4 Overall Speedups

In Table 3.9, we tabulate minimum, maximum and mean speedups across our benchmark set for two comparisons (1) Xilinx Virtex-5 vs. Intel Xeon 5160 (65nm technology) and (2) Xilinx Virtex-6 vs. Intel Core i7 965 (45 nm technology). We achieve modest mean speedups of $2 \times$ when using the Loop-Unrolling optimization and in the worst case a slowdown of $0.3 \times$. The mean speedup increases to $5 \times$ if we use the GraphStep scheduling optimization while the worst case a slowdown of $0.9 \times$. If we combine both together, we are able to achieve a higher mean speedup of $6.5 \times$ with
a worst case slowdown of 0.9×. For a given optimization, the FPGA speedups are almost identical at both technology nodes suggesting a sustained performance benefit for the FPGA with technology scaling. This is expected due to our spatial formulation where the performance smoothly scales with increasing capacity for data-parallel computation.

### 3.6.5 Speedups for Different Verilog-AMS Device Models

We plot the speedup achieved by our optimized FPGA designs compared to optimized sequential implementations in Figure 3.15. We also show the percentages of floating-point peak achieved by our design across the different non-linear models and scheduling strategies. In the best case, we observe FPGA speedups as high as 23× while FPGA floating-point utilization as high as 60%. In contrast, the optimized CPU implementations never achieve more than 30% floating-point utilization for any non-linear model. Both loop-unrolling and GraphStep scheduling contribute towards this performance benefit. We show the key reasons behind this benefit in Figure 3.16. As we see in Figure 3.16(a), Loop-Unrolling delivers performance improvement as we increase the extent of unroll. The extra work introduced at large unroll factors compensates for the large latency of the dataflow graph for the cases characterized by high latency. At large unroll factors, we see a saturation in performance as the extra work exposed matches the latency of the graph. In this case, we merely increase memory requirements without significant performance improvements. In Figure 3.16(b), we separately show how GraphStep scheduling allows us to overlap compute and communication phases of the schedule. When these times match, we can achieve performance close to the computational peak as neither phase dominates total system performance (i.e., no communication bottleneck). Why do they work better together? When distributing the graph nodes across the VLIW architecture, we may end up with poor utilization of certain operator resources. This happens when we have insufficient floating-point work compared to the scheduled latency of dataflow graph. When compute and communicate phases are mostly balanced, long-latency floating-point operators like `divide` and `sqrt` (57 cycles) may not have sufficient pipeline
Figure 3.15: Impact of Static Scheduling Optimizations (Double-Precision)
slots performing useful work. For example, the bsim3 device evaluation with 128 PEs and a $p = 0.8$ (marginally compute-dominated), requires 74 compute cycles to schedule the floating-point operators (57 cycles of operator latency accounts for most of this time as average nodes/PE in this case is $\approx 9$ nodes). With an unroll factor of 2, we can increase total scheduled cycles to 84 compute cycles to deliver a better per-device throughput of $84/2=42$ cycles. The increased work is easily overlapped with the long-latency floating-point operations to deliver higher utilization of the long latency pipeline. In this case, we can further unroll the graph until the long-latency pipelines are saturated and the bottleneck shifts elsewhere. Thus, GraphStep scheduling with Loop-Unrolling together can deliver higher utilization of the custom VLIW architecture.

Finally in Figure 3.16(c) we see that the CPU instructions are dominated by load/store and data movement instructions. This means that a small fraction of instructions are performing useful floating-point work. In contrast, our VLIW FPGA implementation spatially distributes memory operations, floating-point calculation and communication operations. This means that all these operations are not simultaneously competing for instruction memory bandwidth. We observe a maximum IPC (Instructions per Cycle) of only 0.8 per core (peak of 4 instructions/cycles per core) resulting in the low floating-point utilization we saw in Figure 3.15.

3.6.6 Effect of Device Complexity on Speedup

Across all graphs in Figure 3.15, we observe that the smaller device models (e.g. diode, bjt, and others) achieved much higher speedups that the larger device models (e.g. bsim3, psp). Why is this so? Smaller devices have lower communication requirements in the dataflow graph and consequently fit in smaller memories and smaller networks. Let us now analyze this in greater detail.

In Figure 3.15(a) we show the speedups of the Loop Unrolling FPGA implementation and observe speedups of 0.3-8.3×. For small devices, we are able to deliver high speedups by unrolling the device evaluation graphs by 10-30 unrolls depending on the type of graph. The hbt benchmark delivers the highest speedup as it has
Figure 3.16: Understanding Performance Trends

(a) Loop Unrolling Improvements (8 operators)

(b) GraphStep Scheduling Improvements

(c) Intel Core i7 Analysis
sufficient parallel work and contains 6% exponential and logarithm operations (average 2.8%, see Table 3.3) that are well-suited for parallel FPGA operation. For large non-linear devices like mextram, bsim3, psp we suffer a slowdown. This disparity can be explained by our inability to unroll the large device models due to limited onchip FPGA memory capacity. We can see in Figure 3.17(a) how the peak unroll factor reduces from the high of 15 to the lows of 2–3 as we increase device complexity. Furthermore, the average work per pipeline stage in the dataflow graph representing the device evaluation computation is high for the large devices (also see Figure 3.9). This means that loop unrolling will be less effective at exposing parallelism in this case. In contrast, the smaller device graphs are tall and skinny with limited amount of work per stage. Such graphs can benefit from loop unrolling by balancing the large graph depth with additional work from unrolling.

In Figure 3.15(c) we show the speedups of the GraphStep FPGA implementation and observe speedups of 0.9–16.1×. As expected, this is higher than the speedups achieved from Loop Unrolling. GraphStep implementation allows overlapped scheduling of compute (graph nodes) and communicate (graph edge) operations. However, speedups for the large devices still remain low. In Figure 3.17(b), we show why speedups start to reduce for the large devices. As explained earlier, the peak unroll factor is low for large devices due to limited onchip memory capacity. For GraphStep
scheduling, our peak unroll factors reduce even more since we must accommodate additional rotating-register state in the onchip memories. In Figure 3.17(b), we see that the peak unroll factor drops to the low of 1–2 for the large device models. At the same time, the larger dataflow graph size requires devoting a larger fraction (as much as 60% for the largest device models) of the FPGA area to the BFT network for balanced overall performance. The Loop-Unrolling optimization needs to only devote 40% of its area to the network in the worst case. This is particularly important for the GraphStep scheduling optimization since communication and compute are overlapped. In most cases, we willingly perform this tradeoff to achieve higher floating-point utilization. For the large devices, network area dominates compute area which substantially reduces the peak floating-point capacity (around 2.5× when comparing psp with diode). Thus, when we achieve high floating-point utilization with GraphStep scheduling, we end up with fewer floating-point units on the FPGA for the large devices.

3.6.7 Understanding the Effect of Scheduling Strategies

In Figure 3.18, we separate out the effect of the scheduling strategies on performance of bsim4 evaluation as a function of PE size. We note that without either scheduling optimization the performance of the FPGA actually gets worse as we distribute processing over multiple PEs. This is because, the dataflow graph gets scattered across the system with multiple cycles of network latency and insufficient parallelism to balance the latency (see Figure 3.9). When we unroll the computation by a factor of 8, we generate 8× as much parallelism without changing overall latency of the data-parallel computation. This by itself improves performance by ≈5.6× but does not scale as we increase PE count. Now, if we perform GraphStep scheduling instead, we see a higher overall benefit of ≈100× over the unoptimized mapping at 128 PEs. But performance scales by only 3× from 8 PEs to 128 PEs (an ≈16× increase in area). If we perform loop unroll and GraphStep scheduling together, we get better scaling behavior and observe an improvement of 11× as we scale from 8 PEs to 128 PEs. This explains the higher composite speedups shown in Figure 3.15(e).
3.6.8 Effect of PE Architecture on Performance

We now discuss the impact of key architecture features that contribute to our speedup. The internal design elements in our PE (i.e. parallel operators), distributed memories and spatial floating-point operators, the design of the network and software optimization are all crucial for delivering high speedup. In Table 3.10 and corresponding Figure 3.19 we illustrate the impact of these key features on performance of the bsim4 device model. The column **Ratio to Baseline** indicates the loss in performance by eliminating a design elements indicated by the other columns in **Architecture Features**. In the **Single Datapath** case, the processing architecture consists of 8 floating-point operators (2 adds, 2 multiplies, 1 divide, 1 square-root, 1 exponential, 1 logarithm) fused into a single datapath. For this configuration, we can access only one operator at time. We observe a performance loss of 4× when sequentializing operations onto a unified datapath pipeline. To put it another way, we get a 4× benefit from having simultaneous access to the 8 floating-point operators in 8 separate PEs. In the **Single Memory** case, the internal PE memories are fused into one for the scheduler. All arriving messages, outgoing messages and datapath operations
must share a single dual-ported memory. This results in a loss in performance of $2.2\times$. Thus, streamlined message processing (simultaneously performing 2 network receives and a network send) with overlapped compute operations using distributed local memories within the PE delivers an additional $2.2\times$ performance improvement (compare Single Memory with Single Datapath). Next, in the Non-Spatial Operators case, we replace fully-pipelined implementations of elementary floating-point functions like divide, sqrt, exponential and logarithm with non-pipelined low-throughput versions while retaining streamlined message processing. We observe that pipelined spatial implementations of elementary floating-point operations contribute a significant $6\times$ performance improvement (compare Non-Spatial Operators with Single Datapath). Finally, spatial floating-point operators combined with streamlined message-processing together deliver a $7\times$ improvement over the non-spatial, non-streamlined version (compare Non-Spatial and Single Memory with Single Datapath). This means we get a modest $1.1\times$ improvement by adding streamlined message-processing to the non-spatial implementation (compare Non-Spatial and Single Memory with Non-Spatial Operators). Streamlined message-processing becomes important only when the compute bottleneck introduced by non-spatial processing is eliminated. When we compare the effect of having all architecture features we observe that the we can
Table 3.10: Impact of Architecture Features of the PE on Performance

<table>
<thead>
<tr>
<th>Experiment</th>
<th>PEs</th>
<th>Ops. per PE</th>
<th>Spatial Op.</th>
<th>Mem. per PE</th>
<th>Sched. Cycles</th>
<th>Baseline</th>
<th>Single Datapath</th>
</tr>
</thead>
<tbody>
<tr>
<td>Baseline</td>
<td>8</td>
<td>1</td>
<td>Yes</td>
<td>3</td>
<td>294</td>
<td>1</td>
<td>0.25</td>
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<tr>
<td>Single Datapath</td>
<td>1</td>
<td>8</td>
<td>Yes</td>
<td>3</td>
<td>1201</td>
<td>4</td>
<td>1</td>
</tr>
<tr>
<td>Single Memory</td>
<td>1</td>
<td>8</td>
<td>Yes</td>
<td>1</td>
<td>2760</td>
<td>9</td>
<td>2.2</td>
</tr>
<tr>
<td>Non-Spatial Operators</td>
<td>1</td>
<td>8</td>
<td>No</td>
<td>3</td>
<td>7148</td>
<td>24</td>
<td>6</td>
</tr>
<tr>
<td>Non-Spatial and Single Mem.</td>
<td>1</td>
<td>8</td>
<td>No</td>
<td>1</td>
<td>8244</td>
<td>28</td>
<td>7</td>
</tr>
</tbody>
</table>

deliver performance as high as 28× compared to the non-spatial, non-streamlined, unoptimized implementation.

3.6.9 Auto-Tuning System Parameters

Our auto-tuning framework selects implementation configurations that are feasible on the target FPGA while maximizing performance. As we saw in Figure 3.16(a), increasing unroll factor can improve performance. However, this also increases the amount of on-chip memory resources required. In Figure 3.20(a), we show both performance improvements and memory resource requirement for the bsim4 device model as a function of unroll factor. For Figure 3.16(a), we consider a 7-tile design where each tile contains 8 PEs (operators). We observe that at an unroll factor of \( \approx 12 \) we run out of on-chip capacity of the Xilinx Virtex-6 LX760. Moreover, performance starts to saturate at higher unroll factors as explained previously. Our auto-tuner limits the unroll factor to fit on-chip capacity.

In Figure 3.20(b) we measure the impact of scaling PE count per tile for a 7-tile design and network richness (Rent parameter) within each tile on performance and area cost (FPGA Slices). We observe that we can accommodate PE counts below 16 before we run out of on-chip logic capacity of the Xilinx Virtex-6 LX760. For \( p=0 \) (bandwidth of a ring), we see no performance improvements while total area cost increases linearly with PE count. This is an example of area-time tradeoff that simply wastes area. At the other end, \( p=1.0 \), we see performance improvements as we
increase system size. In this case, the richer network adds as much as $2 \times$ additional area compared to the $p=0$ system. However, this additional network area allows us to deliver performance close to the ideal scaling curve.

The static scheduling strategies generate FPGA configurations with a range of area-time and memory-time tradeoffs. While Loop Unrolling generates work for the scheduler by filling-in idle slots, it increases the memory requirement for storing the intermediate results along the edges of the unrolled dataflow graph. Similarly, Software Pipelining generates a dense schedule by tight packing of compute and communication operations but it increases network bandwidth and also increases memory requirement for storing the rotating registers. We show the different area-time GraphStep configurations possible for the `bsim4` device model in Figure 3.21(a) and memory-time Loop-Unrolling configuration in Figure 3.21(b). Each point in these graphs corresponds to an arrangement of floating-point operators, network richness and unroll factor. Our auto-tuner picks the best configuration through an exhaustive design-space exploration of this space. This time-consuming exploration is acceptable as we know the dataflow graphs for the devices when setting up the simulator much in advance of the actual simulation run. Our framework also generates some configurations that will be feasible only on future FPGAs and allows us to make robust

Figure 3.20: Understanding Performance Trends
design decisions. Thus, we can easily scale our design to fit newer, larger FPGAs in the future without requiring a redesign from scratch.

### 3.7 Parallel Architecture Backends

We have seen how SPICE Model Evaluation is characterized by abundant data parallelism. Modern parallel architectures such as GPUs and multi-core processors can exploit this data parallelism with specifically-tailored parallel descriptions. How can we target these different parallel organizations to implement the Model-Evaluation phase? How does the FPGA implementation compare against these architectures? In this section we show how to explore multiple parallel implementations on these architectures through a combination of automated code-generation and auto-tuning operations.

In Table 3.11, we compare the peak floating-point capacities of the different architectures used in this study. We observe that the raw floating-point peaks vary as much as two orders of magnitude across the architectures. However, floating-point peak throughputs do not always characterize achieved application performance. Each architecture supports a different compute organization (SIMD, Vector, VLIW) for delivering this throughput to the application. This suggests that SPICE Model
<table>
<thead>
<tr>
<th>Family</th>
<th>Chip</th>
<th>Tech. (nm)</th>
<th>Clock (GHz)</th>
<th>Peak GFLOPS</th>
<th>Power (Watts)</th>
<th>GFLOPs per Watt</th>
</tr>
</thead>
<tbody>
<tr>
<td>Intel Xeon</td>
<td>5160</td>
<td>65</td>
<td>3</td>
<td>12</td>
<td>24</td>
<td>80</td>
</tr>
<tr>
<td>Xilinx Virtex-5</td>
<td>LX330T</td>
<td>65</td>
<td>0.2</td>
<td>11</td>
<td>33</td>
<td>20–30</td>
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<tr>
<td>IBM Cell</td>
<td>PS3</td>
<td>65</td>
<td>3.2</td>
<td>10</td>
<td>204</td>
<td>135</td>
</tr>
<tr>
<td>Sun Niagara</td>
<td>Ultraspac</td>
<td>65</td>
<td>1.2</td>
<td>8</td>
<td>9.6</td>
<td>95–123</td>
</tr>
<tr>
<td></td>
<td>T2</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>0.1</td>
</tr>
<tr>
<td>NVIDIA GPU</td>
<td>9600GT</td>
<td>65</td>
<td>1.6</td>
<td>-</td>
<td>312</td>
<td>59–96</td>
</tr>
<tr>
<td>AMD FireGL GPU</td>
<td>5700</td>
<td>55</td>
<td>0.6</td>
<td>120</td>
<td>144</td>
<td>-</td>
</tr>
<tr>
<td>Intel Core i7</td>
<td>965</td>
<td>45</td>
<td>3.2</td>
<td>25</td>
<td>51</td>
<td>130</td>
</tr>
<tr>
<td>Xilinx Virtex-6</td>
<td>LX760</td>
<td>40</td>
<td>0.2</td>
<td>26</td>
<td>75</td>
<td>20–30</td>
</tr>
<tr>
<td>NVIDIA GPU</td>
<td>GTX285</td>
<td>55</td>
<td>1.4</td>
<td>132</td>
<td>1062</td>
<td>204</td>
</tr>
<tr>
<td>AMD Firestream GPU</td>
<td>9270</td>
<td>55</td>
<td>0.75</td>
<td>240</td>
<td>1200</td>
<td>160–220</td>
</tr>
</tbody>
</table>

Table 3.11: Peak Floating-Point Throughput

Evaluation may achieve better overall performance even on architectures with poor floating-point peaks due to higher utilization of available resources. In this experiment, we focus on comparing the performance of the Intel Xeon 5160, NVIDIA GPU 9600 GT, IBM Cell (1st generation), Sun Niagara 2 with the Xilinx Virtex 5 FPGAs (65nm technology) and the Intel Core i7 965, NVIDIA GTX 285, ATI Firestream 9270 along with the Xilinx Virtex 6 FPGAs (55nm technology or smaller). We customize the code-generators and performance tuners to match the programming models for the different architectures and perform this experiment with a completely automated flow.

### 3.7.1 Parallel Architecture Potential

We now attempt to understand the compute organizations of the different parallel architectures. Each architecture we consider contains multiple floating-point units which are essential for a parallel implementation of Model Evaluation. They differ in their operating frequencies, pipeline depths and data access mechanisms. Our parallel implementation must aspire to maximize performance by efficiently partitioning device clusters across parallel hardware and keeping resource utilization properly
Table 3.12: A comparison of data-parallel constructs across three architectures
balanced.

The **Intel Core i7 965** processor provides four parallel cores with hyper-threading capability on a single-chip and has 128-bit vector floating-point SSE pipelines per core. Our parallel implementation partitions the devices across the cores using multiple parallel threads. Within a core, we can pack computation into SSE instructions to keep the vector units busy. We can strategically use loop-unrolling to generate additional work for higher utilization.

The **IBM Cell** processor contains a single Power Processing Element (PPE) and eight Synergistic Processing Elements (SPE) with 128-bit vector floating-point pipelines. However, the Cell processor in the Sony PlayStation 3, which we use in this paper, allocates one SPE for running a proprietary hypervisor and keeps another SPE as spare for improving yield. We statically partition devices across the 6 SPEs. Furthermore, we only consider single-precision implementations due to higher peak throughput and bugs in the Cell vector library. Since there is no shared memory, we must explicitly load inputs and unload outputs of the Model Evaluation computation. We pack device computation into vector operations provided by the Cell SPEs for higher utilization.

The **NVIDIA GPUs** used in our experiments are CUDA-capable (Compute Unified Device Architecture) graphics processors that support general-purpose computation expressed in C/C++ with the CUDA API. The 9600 GT GPU contains 64 stream processors organized into 8 multiprocessors each capable of running multiple threads managed by the GPU thread scheduler. The GTX 285 is a larger GPU that contains 240 stream processors organized into 30 multiprocessors. The high-level CUDA implementation runs the kernel in a data-parallel SIMD (single-instruction multiple-data) fashion across the parallel GPU resources. The devices get partitioned across the different processors depending on a specified distribution pattern.

The **ATI GPUs** used in our experiments support the ATI Stream Computing platform that allows computation described in the Brook language to run in parallel on the GPU hardware. The FireGL 5700 GPU is a low-power GPU for mobility applications that contains 120 shader cores while the Firestream 9270 GPU
is a higher-performance GPU with 800 stream cores. We express our data-parallel computation as stream kernels in the Brook language to distribute processing across these parallel cores.

The **Xilinx FPGA** contains hundreds of thousands of logic cells, hundreds of on-chip memories and embedded fixed-point multipliers that can be configured to implement irregular floating-point datapaths. We statically distribute the devices across multiple *custom VLIW tiles* (see Section 3.5). Each tile is statically-scheduled using a combination of loop-unrolling and software-pipelining to design a well-balanced system.

In Figure 3.22, we show our mapping flow for our experiments with parallel architectures. We share the Verilog-AMS compiler with the FPGA backend. The rest of the flow is customized for the software environments specific to each parallel architecture. We tabulate the different compilers, tools, libraries and timing-functions used in our experiments in Table 3.13. We measure runtime of our generated code averaged across a large number of device evaluations to minimize the effect of startup costs, OS overheads and measurement noise. The two key components of this flow are (1) the code-generator and (2) the auto-tuner. We explain these in detail in Section 3.7.2 and Section 3.7.4 respectively.

### 3.7.2 Code Generation

We target multiple architecture backends in our study to enable a fair comparison of the different compute organizations. The software development framework for each backend supports unique parallel constructs to expose parallelism within the application. Instead of manually rewriting code using these constructs for each ar-
Architecture, we use automated code generation to simplify the mapping process. Our code-generator backend accepts an intermediate representation of computation generated by the Verilog-AMS compiler. This allows us to target a single Verilog-AMS description of computation across all the architectures considered in this study. For data-parallel computation, the constructs extract the number of data-parallel units of work available and the computation for each of these units. The compilers or runtimes distribute the data-parallel units of work across the parallel computing elements. Each computing element then evaluates the computation for the subset of work it is assigned.

Each architecture provides a different data-parallel constructs to expose parallelism to the compiler as shown in Table 3.12. We generate code with simple OpenMP pragma `omp parallel for` shown in Table 3.12(a) to distribute Model-Evaluation across 8 threads on the Intel Core i7 processor. We express each individual device evaluation as a scalar kernel and let the GPU thread scheduler distribute these threads across the GPU using the CUDA API constructs shown in Table 3.12(b). We distribute processing across the six user-programmable PS3 Cell SPUs by using PThreads [84] shown in Table 3.12(c) to create and manage parallel threads. We generate custom VLIW instructions for our FPGA architecture described in [85] and

<table>
<thead>
<tr>
<th>Arch.</th>
<th>Compiler</th>
<th>Libraries</th>
<th>Timing</th>
</tr>
</thead>
<tbody>
<tr>
<td>Intel CPUs</td>
<td>gcc-4.4.3 (-O3)</td>
<td>OpenMP 3.0 [80], GNU libm, Intel MKL 10.1</td>
<td>PAPI 4.0.0 [18], PAPI_flops()</td>
</tr>
<tr>
<td>Nvidia GPUs</td>
<td>nvcc, CUDA SDK 2.3</td>
<td>CUDA libraries</td>
<td>cudaEventRecord()</td>
</tr>
<tr>
<td>ATI GPUs</td>
<td>brc, g++-4.1.2, ATI Stream CAL 1.4beta [81]</td>
<td>ATI Brook libraries</td>
<td>gettimeofday()</td>
</tr>
<tr>
<td>IBM Cell</td>
<td>spu-gcc, ppu-gcc, Cell SDK 3.1 [82]</td>
<td>Simdmath, MASS</td>
<td>gettimeofday()</td>
</tr>
<tr>
<td>Sun Niagara2</td>
<td>cc, Sun Studio 12.1 [83]</td>
<td>OpenMP [80], libm</td>
<td>PAPI 3.7.0 [18], PAP_flops()</td>
</tr>
<tr>
<td>Xilinx FPGA</td>
<td>Synplify Pro 9.6.1, Xilinx ISE 10.1</td>
<td>Xilinx Coregen [77], Arénaire [79]</td>
<td>-</td>
</tr>
</tbody>
</table>

Table 3.13: Software Environments
3.7.3 Optimizations

In addition to data-parallelism, we can exploit other characteristics of SPICE Model-Evaluation graphs to get better performance. For example, we notice that SPICE model-evaluation graphs are characterized by long critical paths with little work off this path which may significantly underutilize processor capacity (see Figure 3.9). To increase utilization and improve performance, we can perform loop unrolling or vectorize the device loop so that multiple devices are scheduled together. For example, the \texttt{bsim3} device model implemented in Single-Precision on 8-operators ($p = 0.8$) with GraphStep scheduling requires 481 cycles per evaluation with no unrolling, which reduces to 477 cycles per evaluation when unrolled twice.

3.7.4 Auto Tuning

To enable a fair comparison of performance across the different architectures, we must be able to generate optimized code for each architecture. We can achieve the best performance on each parallel architecture by generating custom code for a target architecture and then tuning the performance for each target in an architecture-specific manner. The data-parallel computation distributed across the parallel compute elements must be tuned to achieve best performance. The tuning process identifies and assigns values to optimization parameters that influence performance of the mapped application (e.g. loop unroll factor, vector length). Typically, tuning is a manual programmer-intensive process requiring trial-and-error or iterative approach to choose the best value for an optimization parameter. This can be a non-trivial amount of time if the number of parameters and their interactions are large. Furthermore the results of the tuning process are tied to the particular combination of compiler, runtime, operating-system and system configuration and not trivially portable to a new system.

For our experiments, we choose an automated approach that empirically tunes
the mapping for each architecture. The approach is similar to the auto tuner used in the ATLAS framework [86] for optimizing dense linear algebra kernels. Our auto tuner exhaustively explores several implementation parameters for the different architectures as shown in Table 3.14. The set of possible configurations we explore makes it infeasible for a manual tuning process and automated approach is required. Such an exhaustive approach is possible in our case since the Model Evaluation graphs are completely known in advance. Furthermore, the entire space of implementations can be explored automatically in a few hours. In Figure 3.23 we show the dynamic performance range that our auto-tuner explores while determining the best parallel implementation. As we can see in some cases the dynamic range can be as high as 100×. We envision an installation phase during initial setup on a parallel system, where we spend time generating an optimized configuration. This offline install-time tuning will pay off during an actual SPICE simulation by delivering the best possible performance on that system.

Figure 3.23: Auto-tuning Performance-Ranges for the Parallel Architectures at 65nm (9493 configurations)
Table 3.14: Auto-Tuning Parameters

<table>
<thead>
<tr>
<th>Architecture</th>
<th>Parameter</th>
<th>Range</th>
<th>Increment</th>
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<tbody>
<tr>
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<td>+1</td>
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<tr>
<td></td>
<td>MKL Vector</td>
<td>true/false</td>
<td></td>
</tr>
<tr>
<td>NVIDIA GPU</td>
<td>Loop-Unroll Factor</td>
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<td>+1</td>
</tr>
<tr>
<td></td>
<td>Threads per block</td>
<td>8–512</td>
<td>×2</td>
</tr>
<tr>
<td></td>
<td>Registers/Thread</td>
<td>16–128</td>
<td>×2</td>
</tr>
<tr>
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<td>Loop-Unroll Factor</td>
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<td>+1</td>
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</tr>
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<td></td>
<td>MASS Vector</td>
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<tr>
<td>Sun Niagara2</td>
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<td>+1</td>
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<td>Operators per PE</td>
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<td>×2</td>
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<td></td>
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</tbody>
</table>

Let us now consider a few examples of how this tuning works. Our GPU implementation organizes device evaluations into threads (mapped to an ALU: a floating-point unit) which must be grouped into blocks (mapped to multiprocessor: collection of ALUs) and grids (mapped to GPU: collection of multi-processors) for a CUDA implementation. Our auto-tuner picks the number of threads in each block (grid configuration) to maximize GPU usage and deliver best performance. Similarly, our FPGA architecture includes several different parameters that can be chosen to make best use of available resources for a given problem.

3.8 Results: Single-Precision Parallel Architectures

In this section, we discuss our results of mapping SPICE Model-Evaluation code to multiple parallel architectures. For these experiments we use runtimes for our models averaged across multiple device evaluations to eliminate measurement noise and other artifacts.

In Table 3.15 we show the minimum, maximum and mean speedups across different non-linear devices for the different parallel architectures when compared to the optimized Intel microprocessor implementations. We also show mean percent
floating-point peak achieved by the different architectures. We observe that at 65nm, the Virtex-5 LX330T achieves the highest mean speedup of $35\times$ across the different architectures at that technology. We also see mean peak floating-point usage of around 40%. The rest of the architectures provide modest speedups with the exception of the NVIDIA 9600GT GPU and only achieve a mean peak floating-point usage of a few percent. Thus, the architecture with the highest peak does not necessarily deliver the best performance. This changes for the 45nm architecture experiments. While the FPGA still achieved good speedup of around $30\times$ across the benchmark set, we observe a noticeable increase in the performance of the two GPUs. The NVIDIA GTX285 delivers a mean speedup of $58\times$ while the ATI Firestream 9270 delivers a mean speedup of $71\times$ at sufficiently large number of device evaluations. We attribute this increase to the substantially superior floating-point peak throughput. The data-parallelism in the Model-Evaluation computation matches favorably with the SIMD and Streaming compute models of these GPU architectures. Despite this high speedup the FPGA implementation still manages to achieve a high floating-point utilization of 43% while the GPUs only manage 6% usage. In this case, the performance difference between the FPGA and the GPU is much smaller than what the peaks might suggest. We explain some of the reasons behind this difference in Section 3.8.2. In Figure 3.24 we plot the speedups and percent floating-point peak achieved by each non-linear device model across the different architectures.

### 3.8.1 Performance Analysis

In Figure 3.25 we plot achieved floating-point throughputs against the peak floating-point throughputs of the different parallel architectures. As expected, we are unable to fully exploit the peak floating-point throughput for any architecture due to bottlenecks in other components of the architecture (all achieved data-points are below Actual=Peak line in the figure). We observe that at both 65nm and 45nm, the FPGAs are the closest to achieving their peak potential. At 65nm, not only does the FPGA achieve a larger fraction of the peak, it also achieves the highest overall floating-point throughput. At 45nm, the GPUs do achieve a higher actual floating-point throughput
Figure 3.24: Total Speedups and Percent Peak Floating-Point Throughputs (Single-Precision)
<table>
<thead>
<tr>
<th>Technology</th>
<th>Speedup</th>
<th>% Peak</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Min</td>
<td>Max</td>
</tr>
<tr>
<td>65nm</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Intel Xeon 5160</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>Virtex-5 LX330T</td>
<td>4.5</td>
<td>123.5</td>
</tr>
<tr>
<td>NVIDIA 9600GT GPU</td>
<td>10.1</td>
<td>63.9</td>
</tr>
<tr>
<td>ATI FireGL 5700 GPU</td>
<td>0.4</td>
<td>6.0</td>
</tr>
<tr>
<td>IBM Cell PS3</td>
<td>3.8</td>
<td>16.2</td>
</tr>
<tr>
<td>Sun Niagara2</td>
<td>0.4</td>
<td>1.4</td>
</tr>
<tr>
<td>45nm</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Intel Core i7 965</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>Virtex-6 LX760T</td>
<td>4.5</td>
<td>111.6</td>
</tr>
<tr>
<td>NVIDIA GTX285 GPU</td>
<td>13.1</td>
<td>133.1</td>
</tr>
<tr>
<td>ATI Firestream 9270</td>
<td>2.8</td>
<td>1200</td>
</tr>
</tbody>
</table>

Table 3.15: Speedups for Single-Precision Model Evaluation across Parallel Architectures (Compared to optimized Intel implementations at each technology node)

resulting in better performance than the FPGAs. However, this comes at the cost of almost $100\times$ higher peak floating-point throughput.

In Figure 3.26(a), we show how performance scales as a function of number of threads of the Sun Niagara2. While the overall performance of this processor is poor, we believe this scaling behavior is indicative of the kind of tradeoffs that will be necessary to efficiently program multi-core systems with 100s of cores in the future. For small devices like diodes performance saturates at 16 threads while it continues to scale up to 64 threads for larger devices like mextram and psp.

In Figure 3.26(b), we show the impact of using the Intel MKL library [87] for accelerated processing of vectorizable floating-point functions (e.g. divide, sqrt, exp, log). We observe a limited performance improvement of at most $\approx3\times$ for using the vector library only for single-precision evaluation. The limited speedups possible are due to the inability to implement all the operations in the Model-Evaluation computation in vectorized form. For the large devices, only 40–50% of the operations are vectorizable. We see no practical improvements for double-precision evaluation using the MKL library. The peak floating-point throughput for single-precision evaluation on Intel CPUs is twice as high as double-precision evaluation (see Table 3.11).

In Figure 3.26(c), we show how performance scales as a function of number of
Figure 3.25: Comparing Actual and Peak Floating-Point Throughputs

(a) Architectures 65nm (Actual vs. Peak)
(b) Architectures 65nm (Actual vs. Peak)

Figure 3.26: Additional Analysis

(a) Thread Scaling for Sun Niagara2
(b) Intel MKL Library
(c) Performance Scaling with Device Count
(d) Accounting for Memory Transfer time
devices being evaluated for the \texttt{bsim4} device model. As expected, at low device counts, the performance per device is poor. At these low device counts, the cost of distributing evaluation across parallel resources does not justify the performance improvement from parallelism. As we increase the number of devices, the cost is amortized by the increased amount of parallel work. Thus at large device counts, we see markedly improved performance. The ATI Firestream runtime has a high overhead for launching kernels and transferring data to the GPU. We are able to realize the architecture’s potential only at much higher device counts than the other architectures. We also observe a performance saturation for the Intel Core i7 at 256 devices. The performance improvement for the NVIDIA GTX285 GPU saturate at much higher 4096 devices while the Xilinx Virtex-6 FPGA saturates at 16384 devices.

In Figure 3.26(d), we include the cost of transferring data when computing total speedup. For the FPGA, we estimate loading time for the device parameters from the offchip interface (capable of transferring 32 bytes per cycle using the BEE3 memory controller \cite{88}). For the two GPU architectures, we include the cost of copying the input arrays from the host CPU to the device and the output arrays from the device to the host CPU. We note that the ATI Firestream 9270 performance drops dramatically when including these memory transfer times. The performance of the NVIDIA GTX285 also drops significantly and allows the FPGA to close the performance gap for large device sizes.

### 3.8.2 Explaining Performance

In Table 3.16, we show assembly instruction analysis for a few candidate architectures. We separate floating-point instructions (column ‘Float’ or ‘ALU’) from memory load-store instructions (column ‘Ld/Str’ or ‘Fetch’) from the binary assembly codes generated by \texttt{gcc} for the Intel processors to estimate a weak lower-bound on floating-point utilization. For the NVIDIA GPU, we use the CUDA occupancy calculator \cite{89} to predict the utilization of our floating-point resources. For the AMD/ATI GPUs, we use the Stream KernelAnalyzer \cite{90} to calculate the distribution of instructions in the GPU assembly.
Table 3.16: Assembly Code Analysis for SPICE Model-Evaluation

<table>
<thead>
<tr>
<th>Model</th>
<th>Intel Xeon</th>
<th>IBM Cell</th>
<th>NVIDIA GPU</th>
<th>ATI GPU</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Float. (%)</td>
<td>Ld/Str (%)</td>
<td>Float. (%)</td>
<td>Ld/Str (%)</td>
</tr>
<tr>
<td>bjt</td>
<td>10</td>
<td>23</td>
<td>13</td>
<td>20</td>
</tr>
<tr>
<td>diode</td>
<td>4</td>
<td>14</td>
<td>9</td>
<td>26</td>
</tr>
<tr>
<td>jfet</td>
<td>9</td>
<td>20</td>
<td>14</td>
<td>24</td>
</tr>
<tr>
<td>mos1</td>
<td>8</td>
<td>29</td>
<td>11</td>
<td>23</td>
</tr>
<tr>
<td>vbic</td>
<td>17</td>
<td>22</td>
<td>17</td>
<td>19</td>
</tr>
<tr>
<td>mos3</td>
<td>20</td>
<td>44</td>
<td>19</td>
<td>15</td>
</tr>
<tr>
<td>hbt</td>
<td>11</td>
<td>49</td>
<td>14</td>
<td>17</td>
</tr>
<tr>
<td>bsim4</td>
<td>28</td>
<td>47</td>
<td>25</td>
<td>16</td>
</tr>
<tr>
<td>bsim3</td>
<td>36</td>
<td>50</td>
<td>37</td>
<td>17</td>
</tr>
<tr>
<td>mextram</td>
<td>26</td>
<td>59</td>
<td>29</td>
<td>25</td>
</tr>
<tr>
<td>psp</td>
<td>41</td>
<td>52</td>
<td>43</td>
<td>24</td>
</tr>
<tr>
<td>Mean</td>
<td>16</td>
<td>34</td>
<td>18</td>
<td>20</td>
</tr>
</tbody>
</table>

Ld/Str=Load Store, Float.=Floating-Point

We measured mean floating-point fractions of total instruction to be around 15% irrespective of architecture. For small models like diode and bjt, this fraction may even be in the below 10%. For larger models like bsim3 and psp, the fraction rises to 25–40% of total instructions. Modern processors contain a fixed combination of Load/Store and ALU capacity. This imbalanced fraction suggests that we may never be able to fully achieve the peak utilization of floating-point units in these processing architectures. On the FPGA, we are able to spatially distribute processing of the Load/Store operations by creating address generators and datapaths for the computation. We customize the distribution of floating-point resources to match the required application balance and to avoid hotspots. We overlap compute and communicate operations to improve performance. We also ensure high resource utilization by software-pipelining and loop-unrolling.

3.9 Conclusions

In this chapter, we show how to implement the data-parallel Model-Evaluation phase of SPICE on an FPGA and other parallel architectures for a range of non-linear SPICE
device models. We demonstrate speedups of 1.4–23.1× when comparing a Xilinx Virtex-6 LX760 FPGA with an Intel Core i7 965 for double-precision floating-point evaluation. Our model-specific, statically-scheduled VLIW FPGA architecture is able to deliver these speedups due to need-proportional provisioning of floating-point operators, software-pipelining and loop-unrolling optimizations, overlapped scheduling of compute and communicate phases resulting in higher utilization of floating-point resources (40%–60%). Our mapping framework can also target other parallel architectures apart from FPGAs since we capture the Model-Evaluation computation in a high-level framework based on Verilog-AMS. We use code-generators and auto-tuners to generate optimized implementations across these different architectures. We deliver speedups of 4.5–123.5× for a Virtex-5 LX330, 10.1–63.9× for an NVIDIA 9600GT GPU, 0.4–6× for an ATI FireGL 5700 GPU, 3.8–16.2× for an IBM Cell and 0.4–1.4× for a Sun Niagara 2 architectures when comparing Single-Precision evaluation across these architectures at 55nm–65nm technology. We also show speedups of 4.5–111.6× for a Virtex-6 LX760, 13.1–133.1× for an NVIDIA GTX285 GPU and 2.8–1200× for an ATI Firestream 9270 GPU when comparing Single-Precision evaluation on modern architectures at 40–55nm technology. We observe that at 55nm–65nm FPGAs deliver the highest performance for Model-Evaluation and are within 2.5× of the GPUs at 40–55nm comparisons. We expect to improve FPGA speedups in the future using higher quality distribution of floating-point computation, use of offchip memory for storing intermediate state, superior floating-point operators [91, 92] and better static scheduling algorithms.
Chapter 4

Sparse Matrix Solve

In Chapter 2 (Section 2.1), we identified the Matrix-Solve phase of the SPICE circuit simulator as the most challenging phase for parallelization. The computation is characterized by sparse, irregular operations that are too fine-grained to be effectively exploited on conventional architectures (e.g. multi-cores). In this chapter, we show how to parallelize Sparse Matrix Solve using a combination of the KLU algorithm (better software) and an efficient dataflow FPGA architecture (better hardware). We start by introducing the KLU algorithm that extracts the exact compute graph of matrix factorization and then describing a dataflow architecture for efficiently mapping the compute graph to an FPGA. We identify and quantify different approaches for distributing the dataflow graph which is a subject of continuing research.

4.1 Structure of Sparse Matrix Solve

The open-source spice3f5 package assembles the matrix and the right-hand side (RHS) vectors in $A\vec{x} = \vec{b}$ using the Modified Nodal Analysis approach. Since circuit elements ($N$) tend to be connected to only a few other elements, there are a constant number ($O(1)$) of entries per row of the matrix. Thus, the MNA circuit matrix with $O(N^2)$ entries is highly sparse with $O(N)$ nonzero entries ($\approx 99\%$ of the matrix entries are 0 as shown in Table 4.2). spice3f5 uses the Sparse 1.3 [17] solver optimized for efficient sequential processing of sparse matrices to compute $\vec{x}$. This solver uses dynamic pivoting for numerical accuracy which changes the non-zero pattern in every
step of the solve. A change in the non-zero pattern changes the sequence of operations for the Matrix-Solve step. This makes it difficult to distribute the computation in parallel as the compute graph evolves and changes in each iteration. Furthermore, the fine granularity of the underlying matrix solve computation is another factor that prevents efficient parallel operation. Advances in numerical techniques during the past decade have delivered newer, faster solvers. A few studies have successfully used the highly-parallelizable preconditioned Krylov techniques for Sparse Matrix Solve like BiConjugate-Gradient and GMRES for certain classes of circuits. In our experiments, these techniques delivered mixed results and need additional research for proper consideration (see Figure 4.12 in Section 4.5.1). Instead we consider the newer KLU solver that is a sparse, direct method that can work across a broad range of circuits. While a direct method is not as parallelizable or scalable as a Krylov technique, we choose this approach as it is robust across a broader range of circuits.

KLU performs a one-time partial pivoting at the start of the simulation and delivers a static compute graph that can be efficiently distributed and evaluated in parallel.
The KLU solver uses matrix preordering algorithms (Block Triangular Factorization-BTF [94] and Column Approximate Minimum Degree-COLAMD [95] vs. Markowitz reordering used by Sparse 1.3) that are better at minimizing the fillin during the factorization phase. It employs the left-looking Gilbert-Peierls [96] algorithm to compute the LU factors of the matrix for each SPICE iteration. The solver attempts to reduce the factorization runtimes for subsequent iterations (refactorization). It uses a partial pivoting technique to generate a fixed non-zero structure in the LU factors at the start of the simulation (during the first factorization). This static pattern is reused across all SPICE iterations and enables us to generate a specialized static dataflow architecture that processes the graph in parallel. It even enables faster sequential evaluation by specializing the data-structures that hold the LU factor locations and non-zero entries. The preordering and symbolic analysis step (labeled as Step 1 in Figure 4.1) computes non-zero positions of the factors at the start while the refactoring and solve steps (labeled as Step 2) solve the system of equations in each iteration. Our FPGA solution parallelizes the refactorization and solve phases of the KLU solver.

### 4.1.1 Structure of the KLU Algorithm

At the start of the SPICE simulation (in Step 1 of Figure 4.1), the KLU solver performs symbolic analysis and reordering in software to determine the exact non-zero structure of the L and U factors. We show an example matrix and the locations...
(Gilbert-Peierls)

```matlab
L=I; % I=identity matrix
for k = 1:N
   b = A(:,k); % kth column of A
   x = L \ b; % \ is Lx=b solve
   U(1:k) = x(1:k);
   L(k+1:N) = x(k+1:N) / U(k,k);
end;
```

Listing 4.1: Gilbert-Peierls Algorithm (A=LU)

(Sparse L-Solve)

```matlab
x=b;
% symbolic analysis predicts non-zeros
for i = 1:k-1 where x(i)!=0
    for j = i+1:N where L(j,i)!=0
        x(j) = x(j) - L(j,i)*x(i);
    end;
end;
% returns x as result
```

Listing 4.2: Sparse L-Solve (Lx=b, x= unknown )
%---------------------------------------------------------------
% input: lower triangular matrix L, vector b
% output: intermediate vector y
%---------------------------------------------------------------
y = b;
for i = 1:N-1
    y(i+1:N) = y(i+1:N) - L(:,i)*y(i);
end;

Listing 4.3: Front Solve (Ly=b)

%---------------------------------------------------------------
% input: upper triangular matrix U, vector b
% output: final solution vector x
%---------------------------------------------------------------
x = y;
for i = N:1
    x(i-1:1) = x(i-1:1) - U(:,i)*x(i)/U(i,i);
end;

Listing 4.4: Back Solve (Ux=y)
A_1 \cdot \bar{x}_1 = \bar{b}_1 \quad (4.1)

\begin{bmatrix}
1 & -1 & 1 \\
-1 & 4923.1 & 0 \\
1 & 0 & 0
\end{bmatrix} \cdot \begin{bmatrix} x_1 \\ x_2 \\ x_4 \end{bmatrix} = \begin{bmatrix} 0 \\ 4336 \\ 3 \end{bmatrix} \quad (4.2)

L_1 \cdot U_1 \cdot \bar{x}_1 = \bar{b}_1 \quad (4.3)

\begin{bmatrix}
1 & -1 & 1 \\
-0.2E^{-3} & 1 & - \\
1 & 1.0002 & 1
\end{bmatrix} \cdot \begin{bmatrix}
1 & -1 & 1 \\
- & - & 0.2E^{-3} \\
- & - & -1.0002
\end{bmatrix} \cdot \begin{bmatrix} x_1 \\ x_2 \\ x_3 \end{bmatrix} = \begin{bmatrix} 0 \\ 0.880 \\ 3 \end{bmatrix} \quad (4.4)

A_2 \cdot \bar{x}_2 = \bar{b}_2 \quad (4.5)

\begin{bmatrix}
1 & -1 & 1 \\
-1 & 6736.1 & 0 \\
1 & 0 & 0
\end{bmatrix} \cdot \begin{bmatrix} x_1 \\ x_2 \\ x_4 \end{bmatrix} = \begin{bmatrix} 0 \\ 5936 \\ 3 \end{bmatrix} \quad (4.6)

L_2 \cdot U_2 \cdot \bar{x}_2 = \bar{b}_2 \quad (4.7)

\begin{bmatrix}
1 & -1 & 1 \\
-0.1E^{-3} & 1 & - \\
1 & 1.0001 & 1
\end{bmatrix} \cdot \begin{bmatrix}
1 & -1 & 1 \\
- & - & 0.1E^{-3} \\
- & - & -1.0001
\end{bmatrix} \cdot \begin{bmatrix} x_1 \\ x_2 \\ x_3 \end{bmatrix} = \begin{bmatrix} 0 \\ 0.881 \\ 3 \end{bmatrix} \quad (4.8)

Figure 4.3: A Matrix Solve Example using KLU (Circuit from Figure 2.2)

of the LU factors obtained from KLU symbolic analysis in Figure 4.2. The original non-zero pattern is shown in the matrix on the left with the shaded boxes being the non-zero locations. The non-zero pattern of the L and U factors with fillin is shown on the right. The darker shaded boxes are the lower triangular non-zero entries while the lighter shaded boxes are the upper triangular non-zero entries. The box with the cross is a fillin generated during the pre-processing phase. This pre-processing phase is a tiny fraction of total time and needs to be run just once at the start (see column ‘Analysis’ in Table 4.5). In our parallel approach, we start with knowledge of the non-zero pattern of the reordered circuit matrix. We are parallelizing Step 2 of the KLU Matrix-Solver. This step consists of a Factorization, Front-Solve and Back-Solve phases. We show an example matrix and the key steps involved in Figure 4.3. Here we show the original matrix A_1 in a SPICE iteration and its non-zero pattern in Equation 4.2. The first LU factorization in Equation 4.4 processes this matrix to generate the L_1 and U_1 factors. In another SPICE iteration shown in Equation 4.6
we get a new matrix $A_2$ with only the non-zero values changed. The factorization of $A_2$ performs the same sequence of factorization operations on the changed non-zero values to get new factors $L_2$ and $U_2$ as shown in Equation 4.8. In this small example, the sparsity in the matrices is not typical of large SPICE circuits. In Figure 4.4, we show the non-zero pattern of a more typical large SPICE circuit $s1196$. Each black pixel in the figure represents a non-zero value. We show the non-zero pattern of the original matrix on the left and the reordered matrix on the right. The high density of non-zeros along the diagonal is evidence of locality in the matrix. The reordering enhances this clustering effect along the diagonal to maximize locality. This matches the natural real-world circuit structure where elements are connected to only a few other elements (limited fanin and fanout) which are in its spatial neighborhood. The concentration of non-zeros in the final few columns of the reordered matrix are due to high fanout nets in the circuit such as power nets and clock trees.

In Listing 4.1, we illustrate the key steps of the factorization algorithm. It is the Gilbert-Peierls left-looking algorithm that factors the matrix column-by-column from left to right (shown in the figure accompanying Listing 4.1 by the sliding column
For each column \( k \), we must perform a sparse lower-triangular matrix solve shown in Listing 4.2. The algorithm exploits knowledge of non-zero positions of the factors when performing this sparse lower-triangular solve (the \( x(i) \neq 0 \) checks in Listing 4.2). This feature of the algorithm reduces runtime by only processing non-zeros and is made possible by the early symbolic analysis phase. It stores the result of this lower-triangular solve step in \( x \) (Line 8 of Listing 4.1). The \( k \)th column of the \( L \) and \( U \) factors is computed from \( x \) after a normalization step on the elements of \( L_k \). Once all columns have been processed, \( L \) and \( U \) factors for that iteration are ready. The sparse Front-Solve and Back-Solve steps are shown in Listing 4.3 and Listing 4.4 respectively. The compute structure of these steps is similar to the L-Solve step in Listing 4.2. These solve steps use the \( L \) and \( U \) matrices obtained from the factorization phase and generate the final result vector \( x \).

### 4.1.2 Parallelism Potential

From the pseudo-code in Listing 4.1—Listing 4.4 it may appear that the matrix solve computation is inherently sequential. However, if we unroll those loops we can expose the underlying dataflow parallelism available in the sparse operations. In Figure 4.5, we compare the apparent sequential ordering (first column) of operations with the parallel grouping of operations (second column) in the processing. These operations are obtained by stepping through the computation on Line 8–10 of Listing 4.1. We can see each evaluation of Line 10 (divide) in Listing 4.1 and Line 9 (multiply-subtract) in Listing 4.2. We count a total of 6 steps in the sequential execution while we can reduce this to 4 steps in the parallel evaluation. In the parallel evaluation, we schedule operations purely on the basis of their dataflow dependencies rather than the artificial ordering imposed by sequential description. We note that the divides (in Step 1 of Parallel Column) can be issued in parallel since they have no dependencies between each other. Similarly the multiply-subtract operations (in Step 2 and Step 3 of Parallel Column) can also partially overlap. This fine-grained dataflow parallelism allows concurrent evaluation of operations in the computation. We observe there are two forms of parallel structure in the dataflow graph that we can exploit in our
parallel design:

1. Parallel Column Evaluation: Columns 1, 2 and 3 in Figure 4.5 can all be processed in parallel. This is possible when the row corresponding to the diagonal pivot element contains zeros for all earlier columns. This means that the non-zero pivot value will be free of dependencies on earlier columns. These create parallel dataflow paths rooted at the dependency-free pivot elements that can be evaluated concurrently.

2. Fine-Grained Dataflow Parallelism: Multiply operations for updating A(3,4) could proceed before all three earlier columns are evaluated. The non-zero pattern in a column is defined by the sparsity of a circuit node i.e. a circuit node will connect to only few circuit elements. This generates sparse dataflow paths with dependent operations with internal parallelism within the path.

The parallelism comes from the non-zero pattern of circuit matrices. The sparsity and natural clustering of circuit elements generates independent subtrees with little or no communication with each other. We represent the complete dataflow compute graph for this example in Figure 4.6. We also mark the sequential chain as well as parallel wavefronts on the dataflow graph for this example matrix. We note that all operations in the first wavefront (e.g. divides) can be issued in parallel.

Next, we show the execution profile of an example benchmark psadmit2 in Figure 4.7(a) to illustrate potential for parallel operation. We observe that we can issue as many as 6% of the operations in the first few steps of the graph while on average we can issue as many as 75 operations/step. Each step in this computation is a floating-point operation which requires 10s of cycles on an FPGA (see Table 4.4). Thus, the effective number of operations/cycle is \( \approx 7.5 \). This compares somewhat favorably with architecture sizes of 9-25 PEs that can fit on an Virtex-5 LX330T and Virtex-6 LX760 FPGA respectively. However, the critical chain of dependencies in the evaluation can be long and may limit achievable performance (long tail of Figure 4.7(a)). We must take care to avoid a bad distribution of operations as it may spread the critical path across the machine introducing unnecessary high-latency communication (see Figure 4.18(c)). Assuming unbounded hardware and no commu-
Figure 4.5: Steps of Matrix-Solve Evaluation: Sequential (left) and Parallel (right)
Figure 4.6: Execution Flow for Sparse Matrix-Solve Dataflow Graph
Figure 4.7: Profiles of Dataflow graphs

We can estimate peak speedups possible for our benchmark matrices as shown in Figure 4.8(a). We observe that we can accelerate the matrix solve benchmarks by $1.45 - 189 \times$ (mean $10.5 \times$) using this ideal parallel hardware. We also show the predicted utilization of floating-point capacity in Figure 4.8(b). This shows we can expect mean speedups of $10 \times$ and mean utilization of $12\%$ when mapping the Sparse Matrix-Solve computation to ideal hardware. However, communication latency, poor distribution of processing and queueing delays in the processing element will limit realizable speedups and floating-point utilization. Additionally, certain rows and columns in the matrix may be substantially dense (due to high-fanout nets like power lines, clock, etc) that may create bottlenecks in the compute graph (high-fanin and high-fanout nodes). In future work, we plan to attack this bottleneck through associative reduce decomposition.

4.2 Dataflow FPGA Architecture

Our goal is to accelerate the Sparse Matrix Solve computation. This computation can be represented as a sparse dataflow graph once we have performed symbolic analysis to lock down non-zero locations. The nodes represent floating-point operations and edges represent dependencies between the operations as previously shown in Figure 4.6(b). The sequential microprocessor implementation of this computation
Figure 4.8: Predicted Performance of Parallel Sparse Matrix-Solve

under-utilizes the floating-point capacity of the processor due to non-floating-point operations like address calculation and memory fetches (see Figure 4.18(a)). By extracting the dataflow graph and mapping it to a custom parallel architecture, we reduce these overheads. We limit the sparse matrix access to local onchip FPGA memories and implicitly encode instructions in the graph nodes. We expose the inherent dataflow parallelism available in the computation and route dataflow dependencies over the packet-switched network.

The Sparse Matrix Solve graph must be evaluated once per SPICE iteration. Unlike the Model-Evaluation graphs that must be repeatedly evaluated several times per SPICE iteration in order to process a larger number of identical devices in the circuit, the sparse matrix graph needs to be processed only once in each iteration. Hence, we must design an architecture that is optimized for low latency evaluation of the dataflow graph rather than for high throughput.

The KLU solver provides a static dataflow graph of the matrix factorization. Ideally, we should be able to perform an optimized static schedule of this graph on our
Figure 4.9: FPGA Dataflow Architecture for SPICE Sparse-Matrix Solve

Figure 4.10: PE Organization for SPICE Sparse-Matrix Solve
parallel architecture *i.e.* the scheduler specifies the timing and location of each operation across the parallel architecture. However, our scheduling infrastructure is too slow (≈ days of runtime for the largest benchmarks) to handle million-node matrix factorization graphs. Hence, we use a lightweight, dynamically-scheduled design that **self-times** the operations in the dataflow graph. A node in the dataflow graph is ready for processing when it receives all its inputs. This is the *dataflow firing rule*. This technique eliminates the need for static scheduling at the expense of some additional area and slightly lower performance. We need to devote additional area to implement the triggering logic and to support a dynamically-routed packet-switched network. We sacrifice performance compared to the statically-scheduled design because we must now make local scheduling and routing decisions with incomplete information about global system conditions at runtime which may result in longer latency operation.

Our parallel FPGA architecture contains dataflow-scheduled parallel floating-point operators coupled to local memories and interconnected by a packet-switched network (Figure 4.9). This organization is inspired by the Tagged-Token Dataflow style [97]. This architecture processes dataflow graphs by explicitly passing tokens between dataflow graph nodes (over the network) and making independent, local firing decisions to process computation at each node. Each PE processes one dataflow graph node at a time but manages multiple nodes in the dataflow graph (virtualization) to handle dataflow graphs much larger than the physical PE count. When the dataflow-firing rule is satisfied, the floating-point computation at the node is processed by the PE datapath. The results are then routed to the destination nodes as specified in the dataflow graph over a packet-switched network using 1-flit packets [59]. Each packet contains the destination address and the floating-point result.

An FPGA implementation of this computation enables concurrent evaluation of high-throughput floating-point operations, control-oriented dataflow conditions as well as pipelined, low-latency on-chip message routing using the same substrate. The PE shown in Figure 4.9 supports double-precision floating-point add, multiply and divide and is capable of issuing one floating-point operation per cycle. The network
interfaces are streamlined to handle one message per cycle (non-blocking input). We explicitly store the Matrix-Solve graph structure (shown in Figure 4.6) in local FPGA on-chip memories. The Dataflow Logic in the PE keeps track of ready nodes and issues floating-point operations when the nodes have received all inputs (dataflow firing rule). The Send Logic in the PE inspects network busy state before injecting messages for nodes that have already been processed. We map the Matrix-Solve graphs to this architecture by assigning multiple nodes to PEs so as to maximize locality and minimize network traffic. We route packets between the PEs in packet-switched manner over a Bidirectional Mesh network. For large graphs, we may not be able to fit the graph structure entirely on-chip. We can fit the graphs by partitioning them and then loading the partitions one after another. This is possible since the graph is completely feed forward (DAGs) and we can identify the order of loads. We estimate such loading times over a DDR2-500 memory interface.
4.3 Related Work

We summarize several recent parallel Sparse Matrix solvers in Table 4.1 and identify the categories for understanding each design as introduced in Section 2.3.

4.3.1 Parallel Sparse Matrix Solve for SPICE

First, we consider software approaches for parallelizing sparse matrix solvers that are specific to SPICE circuit simulation. These solvers exploit structural properties of the SPICE simulation to accelerate the sparse matrix computation. In [98], a hybrid direct-iterative solver (Numerical Algorithms) is used to parallelize SPICE Matrix-Solve phase but requires modifications to the matrix structure (dense row/column removals) and is able to deliver only 2–3× speedup using 4 SGI R10000 CPUs.

The preconditioned sparse iterative technique GMRES (Numerical Algorithms) used in [99] is shown to provide 3–18× speedup primarily for circuits dominated by linear parasitics. It is able to deliver these speedups by reformulating the SPICE integration and linearization algorithms (SPICE Algorithms). Our technique is applicable to a broader class of circuits and requires no modification to SPICE algorithms but provides lower speedups. SILCA [37] demonstrates good speedup for circuits with parasitic couplings by pruning unnecessary work and trading off accuracy of the simulation (Precision). Our approach does not sacrifice accuracy and shows speedups across a large benchmark set. These work-saving techniques can be adapted for our design to achieve similar benefits.

In [45], a coarse-grained domain-decomposition technique (Numerical Algorithms) is used to achieve 31×-870× (119× geometric mean) speedup for full-chip transient analysis with 32 processors at SPICE accuracy. The key idea is to decompose the large circuit matrix into smaller submatrices (domains) using Additive Schwarz decomposition [15] and then solve the smaller matrices (domains) using a KLU solver. Our technique accelerates the KLU solve step and can be used in conjunction with domain-decomposition to accelerate each domain evaluation.
4.3.2 Parallel FPGA-based Sparse Matrix Solvers

Next, we review FPGA-based approaches for parallelizing sparse matrix solve computation. FPGA-based accelerators for sparse direct methods have been considered in [100, 101] and [104, 102] in the context of Power-system simulations. In [100], a parallel architecture based on embedded processors (Altera NIOS) performs single-precision LU Factorization (Precision) of symmetric Power-system matrices using coarse-grained block-diagonal decomposition to deliver a speedup of $4 \times$ using 6 NIOS RISC processor cores (Compute Organization) compared to a uni-processor implementation on a single NIOS core when considering small matrices of size up to 102x102. This comparison does not include the highly-optimized sequential implementation on an Intel CPU which is likely to be an order of magnitude faster. The embedded processor requires several cycles to perform network send and receive operations and cannot spatially implement the non-floating-point operations. Furthermore it must serialize compute and communicate instructions due to a single instruction issue pipeline. Our processing element design supports double-precision operations and is streamlined to process an incoming message, an outgoing message and a floating-point operation per cycle for higher throughput. A different custom architecture (Compute Organization) implements the block-diagonal decomposition algorithm (Numerical Algorithm) in [101] but compares performance of a Xilinx XC2VP125 against a TMS320C6711 DSP to demonstrate $10 \times$ speedups for small matrices up to size 32x32. The custom architecture does not use embedded processors like the NIOS but is still organized as a RISC architecture preventing spatial implementation of non-floating-point overhead operations.

In [102], a right-looking LU factorization technique using Block-Diagonal decomposition (Numerical Algorithms) delivers 5–6× speedup for the single-precision LU Factorization (Precision) using 2 parallel processing units when compared to an Intel Pentium 4 2.6 GHz for symmetric Power-system matrices. Our approach parallelizes all three phases (LU Factorization, Front-Solve and Back-Solve) on FPGAs in double-precision arithmetic across 9–25 PEs with faster FPGA hardware (125 MHz vs.
200 MHz) and compares performance to the newer Intel Core i7 965. In future work, we expect to enhance our current FPGA implementation with domain-decomposition techniques for further accelerating the sparse matrix factorization of asymmetric circuit matrices.

4.3.3 General-Purpose Parallel Sparse Matrix Solvers

Finally, we briefly discuss techniques used in general-purpose parallel software solvers for sparse matrices. We report these with the intent of comparing our approach with non-SPICE-specific parallel sparse solvers and identifying key differences. The parallel sparse unsymmetric matrix solver HSL_MP48 \cite{103, 105} uses a coarse-grained strategy to decompose the matrix into diagonal blocks (Numerical Algorithms) and factors the blocks in parallel using MPI. They report a speedup of 1–9× over 16 processors on a Cray T3E in \cite{103}. While MPI is not directly applicable to tightly-integrated architectures due to higher latency, we will consider the underlying algorithm in the future to obtain higher speedups. Task queues for scheduling dense matrix computation (Scheduling) across parallel multi-core architectures are considered in \cite{106}. This solver organizes processing into DAGs (directed acyclic graphs) of block operations and uses the task queue to dynamically schedule dependencies between the block operations to maximize performance. Our FPGA approach performs sparse LU factorization of matrices whose structure is known \textit{a priori} and can determine dependencies between operations on individual matrix entries instead of blocks for finer-grained parallelism. Scheduling operations at a block-level is unsuitable for sparse matrices due to the unnecessary sequentialization of possibly concurrent operations within a block. Our approach exposes all available parallelism in the underlying sparse matrix dataflow graph without constraining it into blocks. Furthermore, the communication network between the parallel elements on the FPGA provides a faster mechanism for moving fine-grained data items across the parallel architecture than relying on cache-coherency protocols on conventional multi-core processors.
4.4 Experimental Methodology

We now explain the experimental framework used for the Matrix-Solve design. We show the entire flow in Figure 4.11. In our experimental flow, we use spice3f5 simulator with the Sparse 1.3 [17] package for a functional baseline. We integrate the KLU solver with spice3f5 to replace Sparse 1.3 to measure optimized sequential performance. For our parallel FPGA design, we measure performance of the dataflow graph extracted from the KLU [15] solver using a cycle-accurate simulator that models the dataflow architecture. We compare the sequential and parallel implementation across a range of benchmark matrices generated from spice3f5, circuit-simulation matrices from the University of Florida Sparse-Matrix Collection [111] as well as Power-system matrices from the Harwell-Boeing Matrix-Market Suite [112]. For matrices generated from spicef5, we use circuit benchmarks provided by Simucad [107], Igor Markov [108], Paul Teehan [109] and Sani Nassif [110]. Our benchmark set captures matrices from a broad range of problems that have widely differing structure. We tabulate key characteristics of these benchmarks in Table 4.2 and Table 4.3.
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Table 4.2: Circuit Simulation Benchmark Matrices
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### Table 4.3: UFL Matrix Circuit-Simulation Benchmarks

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<th>BRAM (blocks)</th>
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### Table 4.4: Area and Latency model for Sparse Matrix-Solve Hardware (Virtex-6 LX760)

Our parallel FPGA architecture handles dataflow graphs of the sizes shown in column ‘Total Ops.’ of Table 4.2 and Table 4.3. We generate the dataflow graphs for LU factorization as well as Front/Back solve steps from the initial symbolic analysis and reordering phase of the KLU solver. Once we have the dataflow graphs, we assign nodes to PEs of our parallel architecture. We consider two strategies for placing nodes on the PEs: random placement and placement for locality using MLPart with fanout decomposition. We will consider additional strategies for distributing the graph nodes across our parallel dataflow architecture for higher performance as future work.
4.4.1 FPGA Implementation

We consider a single-chip design of our dataflow architecture on a Xilinx Virtex-5 SX95T and a Xilinx Virtex-6 LX 760 FPGAs. We allow the dataflow design to use the complete FPGA area for these experiments, but when composing the complete SPICE design, we partition the FPGA accordingly (see Chapter 6). In some cases, the dataflow graph will not fit entirely in the FPGA onchip memories. As identified earlier, we can statically compute a loading order and stream the dataflow graph from an offchip DRAM memory.

For our PE design, we use spatial implementations of individual floating-point add, multiply and divide operators from the Xilinx Floating-Point library in CoreGen [77]. These implementations do not support denormalized (subnormal) numbers. We compose the two-dimensional, packet-switched network organized as an 84-bit Bidirectional Mesh (64-bit data, 20-bit address). Each switch in the network is assembled using simple split and merge blocks as described in [59]. The switches implement Dimension-Ordered Routing (DOR [113]). We pipeline the wires between the switches for high performance. Every Manhattan hop in the mesh takes 9 cycles including switch and wire pipeline delays. We estimate memory load time for large matrices using streaming loads over the external DDR2-500 MHz memory interface using lowerbound bandwidth calculations. We do not require a detailed cycle-accurate simulation for the memory controller since it is a simple, sequential, streaming access pattern that we know and precompute. We show the area and latency model in Table 4.4.

We synthesize and implement a sample double-precision 4-PE design on a Xilinx Virtex-5 device using Xilinx ISE 12.1. We provide placement and timing constraints to the backend tools and attain a frequency of 250 MHz. We can fit a system of 9 PEs on a Virtex-5 SX240T (77% logic occupancy) while systems with 25 PEs are easily possible on a Virtex-6 LX760 (estimated 65% logic occupancy). We note that our architecture is frequency limited primarily by the Xilinx floating-point divide operator and the DDR2 controller bandwidth. We can improve our clock frequency
further by using better division algorithms \((e.g.)\) SRT-8 and better memory interfaces \((e.g.)\) DDR3 for additional external memory bandwidth.

4.5 Results

We now discuss experimental results for parallelizing the Sparse Matrix Solve phase of SPICE. We first show results for some initial exploratory studies that attempted to use Iterative algorithms but were not robust enough for circuit simulation. Next, we look at the performance of the sequential SPICE solver with the KLU solver integrated into the package. Finally, we present detailed analysis of the performance of our FPGA design and identify the underlying factors to explain our results.

4.5.1 Exploring Opportunity for Iterative Solvers

It is well-known that \textit{iterative algorithms} for Sparse Matrix Solve based on Krylov-subspace techniques can be parallelized more effectively \[93]. In contrast with \textit{direct solvers} that perform an LU decomposition, the iterative algorithms use repeat evaluations of Sparse Matrix-Vector multiplications. These Matrix-Vector multiplications can be parallelized very effectively on FPGAs \[114]\, GPUs \[115]\ and even Multi-Cores \[87]. Sparse direct solvers are not as easy to parallelize and require additional effort. However, iterative techniques are not always sufficiently robust across different
application scenarios. A lack of robustness may result in SPICE simulations that do not converge at all or converge too slowly. It is possible to improve the robustness of these iterative solvers using preconditioning. We experimented with a range of preconditioned iterative algorithms and sparse direct solvers to evaluate their effectiveness for solving circuit simulation matrices. We show the residues \((\vec{b} - A\vec{x})\) from our experiments in Figure 4.12. The smaller the residue the better the result. We observe that the KLU solver and the preconditioned-BiCG solvers deliver the most robust results across our benchmark set. The preconditioned-GMRES algorithm is consistently generating poor solutions with high residues in a majority of the cases. While preconditioned BiCG iterative algorithm does deliver reasonable results, the cost of calculating the incomplete LU preconditioner (using a sparse, direct method) is expected to be comparable to the cost of complete LU factorization using a sparse direct method. This initial exploration suggested that we will need to parallelize the sparse direct solver in any case. For the purpose of this thesis, we focus on parallelizing the KLU sparse direct solver. In future work, we will fully explore the opportunities for using a parallel preconditioned iterative solver for SPICE.

### 4.5.2 Sequential Baseline: KLU with spice3f5

For our experiments, we first compare the sequential performance of the KLU solver with the Sparse 1.3 solver (default in spice3f5). We replace the Sparse 1.3 solver with the newer, improved KLU solver for all transient iterations. Note that both techniques use the exact same convergence conditions defined in spice3f5 without any loss in accuracy nor an increase in Newton-Raphson iterations. For simplicity, we currently retain Sparse 1.3 to produce the DC operating point at the beginning of the simulation. We quantify the performance benefits of using the newer solver by measuring the runtimes of Matrix-Solve phase of spice3f5 with PAPI 4.0.0 [18] performance counters. We measure runtimes of these sequential solvers when using a single core of the Intel Core i7 965 processor. We compute speedup for the KLU implementation using the formula in Figure 4.13.

We show the speedup for using the KLU solver over the Sparse 1.3 solver in
\[
Software_{\text{Speedup}} = \frac{T_{\text{sparse1.3}}(\text{refactor}) + T_{\text{sparse1.3}}(\text{front solve}) + T_{\text{sparse1.3}}(\text{back solve})}{T_{\text{klu}}(\text{refactor}) + T_{\text{klu}}(\text{front solve}) + T_{\text{klu}}(\text{back solve})}
\]

- \(T_{\text{sparse1.3}}(\text{refactor})\) = Sparse 1.3 Solver LU Refactorization Time on CPU
- \(T_{\text{sparse1.3}}(\text{front solve})\) = Sparse 1.3 Solver Front-Solve Time on CPU
- \(T_{\text{sparse1.3}}(\text{back solve})\) = Sparse 1.3 Solver Back-Solve Time on CPU
- \(T_{\text{klu}}(\text{refactor})\) = KLU Solver LU Refactorization Time on CPU
- \(T_{\text{klu}}(\text{front solve})\) = KLU Solver Front-Solve Time on CPU
- \(T_{\text{klu}}(\text{back solve})\) = KLU Solver Back-Solve Time on CPU

\[
FPGA_{\text{Speedup}} = \frac{T_{\text{klu}}(\text{refactor}) + T_{\text{klu}}(\text{front solve}) + T_{\text{klu}}(\text{back solve})}{T_{\text{fpga}}(\text{refactor}) + T_{\text{fpga}}(\text{front solve}) + T_{\text{fpga}}(\text{back solve})}
\]

- \(T_{\text{fpga}}(\text{refactor})\) = KLU Solver LU Refactorization Time on FPGA
- \(T_{\text{fpga}}(\text{front solve})\) = KLU Solver Front-Solve Time on FPGA
- \(T_{\text{fpga}}(\text{back solve})\) = KLU Solver Back-Solve Time on FPGA

Figure 4.13: Speedup Calculation Equation for Sparse Matrix-Solve Implementations

---

**Figure 4.14: Sequential Sparse Matrix-Solve Speedup for KLU over Sparse 1.3 (Intel Core i7 965)**
We show additional details in Table 4.5 where we include runtimes for the different components of Sparse Matrix-Solve phase when using the Sparse 1.3 solver and the KLU solver. We see that KLU improves the per-iteration matrix solve time by as much as 4.6× for the largest ram2k benchmark while delivering a geometric mean improvement of 1.35× on the whole benchmark set. We also observe that, for some matrices, KLU delivers similar performance as Sparse 1.3 (e.g. 10stages, s298). For very small matrices ≤ 200 rows (e.g. mux8, s27, ringosc), the KLU solver actually slows down the Matrix-Solve phase by ≈50%. The symbolic analysis time (sub-column Analyze in column KLU) is ≈5× the runtime of a single KLU Matrix-Solve iteration. This means that even if the analysis phase remains sequentialized and we speedup iterations by 10× (see Figure 4.16), analysis accounts for only 0.5% of total runtime after 10K iterations (3.3K timesteps for an average of 3 iterations/timestep seen in our benchmark set). We show this calculation in Equation 4.15. We use this faster KLU solver as the sequential baseline for computing speedups of our FPGA architecture.

4.5.3 Parallel FPGA Speedups

For our Dataflow FPGA implementation we compute speedup using the formula previously described in Figure 4.13. In Figure 4.16, we show the speedup of our FPGA architecture using the best placed design that fits in the FPGA over the sequential software version running on an Intel Core i7 965. We obtain speedups between 0.6–7.1× (geomean 2.4×) for Virtex-6 LX760. We observe a slight slowdown for very small matrices mux8 and ringosc. We deliver a speedup of 1.5–5.8× (geomean 3.4×) for Power-system simulation matrices accelerated using FPGAs in [101]. Our solution is 2× faster than the one presented in [101] (estimated from MFLOPs since straight
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Geometric Mean 1.35 5

Table 4.5: Runtime Per Iteration of KLU and Sparse 1.3
Figure 4.16: FPGA Speedups for Sparse Matrix-Solve for Circuits and UFL Matrices (Virtex-6 LX760 vs. Intel Core i7 965)

Figure 4.17: Achieved Floating-Point Peaks
runtime figures were not reported) due to independent processing of routing and floating-point operations. For these speedup calculations we include memory loading time for the large dataflow graph from external DRAM. We also compute speedups in the case where we can overlap memory loading with computation. This is possible since the graph is entirely feed-forward and we can compute a load order in advance. Under this optimistic assumption we can achieve speedups between $0.64 - 13.49 \times$ (geomean $3.8 \times$). We also wish to explore the opportunity for placing large dataflow graphs using greedy packers to avoid the bad random distributions as part of our future work.

In Figure 4.17, we show the percent of peak floating-point throughput of our 25-PE Virtex-6 LX760 FPGA design as well as 1 core of the Intel Core i7 965 processor. A multi-core implementation for the KLU solver currently does not exist. Our experiments with exposing the dataflow structure directly to the multi-core architecture resulted in poor performance. Hence, for this study, we limit our comparisons to the 1-core implementation. The peak double-precision floating-point throughput of our 25-PE design is 12.5 GFLOPs while that of 1 core of the Intel Core i7 965 is 6.25 GFLOPs. We see that the FPGA is able to achieve a geometric mean 4.5% of the floating-point peak of the FPGA while the CPU can only achieve 2.2% of its peak. This translates into 33–1104 MFLOPS (geomean of 520 MFLOPS) on the 25-PE while the processor is only able to achieve rates of 21–502 MFLOPS (geomean of 137 MFLOPS). The FPGA mean speedups and floating-point utilization are within $2–3 \times$ of their predicted values shown earlier in Figure 4.8. This indicates we need to explore different parallelization strategies such as domain decomposition on top of the sparse dataflow parallelism for further accelerating the Sparse Matrix-Solve phase of SPICE.

### 4.5.4 Limits to Parallel Performance

We now attempt to provide an understanding of sequential and parallel performance and identify opportunities for improvement.

We measure the dynamic instruction distribution in the Intel Core i7 implementation of computation in Figure 4.18(a). We observe that only of 1.3% (geomean) of
total instructions are actually floating-point instructions. This partly explains why we observe poor actual floating-point performance in Figure 4.17. The non-floating-point instructions are devoted to address-calculation and memory fetch operations on the sparse matrix data-structures. We can see in Line 9 from Listing 4.2 that we must perform 3 reads and 1 write for each execution of that statement which may require several cycles if not in the cache. Furthermore, we must lookup the non-zero location corresponding to the \( j, i \)th entry in the matrix \( L \) through an address table which can, in turn, take multiple cycles.

Our FPGA implementation achieves a higher utilization and delivers speedup by spatially distributing these address-calculation overheads and by directly routing messages between graph nodes where needed. Next, we measure runtimes of the different phases of the FPGA implementation in Figure 4.18(b) to identify bottlenecks. This shows that memory loading times can account for as much as 38% of total runtime while the LU-factorization phase can account for as much as 30%. Memory load times (limited by DRAM bandwidth) can be reduced by around 3–4× using a higher-bandwidth DRAM interface (saturating all FPGA IO and using DDR3-800) while the LU-factorization runtime can be improved in a limited fashion with better placement of graph nodes.

In Figure 4.18(c), we attempt to bound the additional improvement that may be possible in runtime for the dominant phase: LU factorization. For these estimates we ignore communication bottlenecks. We first compute the latency of the critical path of floating-point instructions in the compute graph. This critical path latency \( (T_{\text{critical}}) \) is the ideal latency that can be achieved assuming infinite PEs, unlimited DRAM bandwidth and no routing delays. This provides an upper bound on achievable speedup \( (\text{Speedup}_{\text{latency}} = T_{\text{sequential}}/T_{\text{critical}}) \). When we account for serialization in each PE arising from reuse of PE resources, we can compute another upper-bound on achievable performance. The floating-point operators in each PE must be shared by multiple dataflow graph nodes. Next, we consider IO serialization latency \( (T_{\text{serialization}}) \) which measures the number of cycles required to send and receive messages into the network. We compute another idealized (approximate) estimate of
Figure 4.18: Understanding FPGA Sparse Matrix-Solve Performance
these serialization overheads at 25 PEs to get a tighter bound on achievable speedup ($Speedup_{serialization} = \frac{T_{sequential}}{T_{serialization}}$). The estimated speedup is the lower of the two idealized estimates ($Speedup = \text{MIN}(Speedup_{latency}, Speedup_{serialization})$).

For most of our benchmarks we are within a 2× of the ideal latency scenario. This suggests that a limited amount of additional speedup is achievable for such benchmarks using better distribution and placement of sparse dataflow parallelism. For larger benchmarks such as $s1196$, $mempplus$ higher additional speedups are possible. For netlists with high potential speedups, we are currently unable to contain the critical paths effectively into few PEs. We expect to extract higher performance for large sparse dataflow graphs through novel placement and clustering strategies as part of our future work.

### 4.5.5 Impact of Scaling

In Figure 4.19, we show the impact of scaling system size (PE count) on performance. We observe a wide range of scaling trends ranging from 14× speedup at 36 PEs (requires 2 Virtex-6 LX760 FPGAs) to no speedups at all in some cases (in fact some slowdown). Large netlists like $40\text{stages}$, $50\text{stages}$ which are relevant and challenging for parallelization show the best scaling. These netlists have low fanout
8–10 and modest fanin within a column ≈400-9000. Small netlists like \texttt{mux8}, \texttt{ringosc} and \texttt{bcspwr01} show poor scaling due to the small dataflow graph sizes ≈600–2000. We see mean speedup scaling saturates at around 4× at 16 PEs with modest improvement to 5× at 36 PEs. This modest speedup scaling is primarily due to the long tail of the dataflow graph. We saw in Figure 4.7 that bulk of the nodes in the graph have a depth of 1–2 in the graph. We also saw in Figure 4.18(c) that for most matrices we are only ≈2× away from ideal critical-path latency in the dataflow graph. This suggests we must investigate alternate strategies for exploiting parallelism. We may be able to improve the latency of the packet-switched network (currently 9 cycles per Manhattan hop) with an alternate design. We discuss a few of our ideas for future work in Chapter 8.

In Figure 4.20 we estimate the impact of two architecture changes on final speedups. We first reduce the latency of the floating-point operations to 1 cycle per operation to minimize the impact of operator latency on total performance. Then, we increase offchip IO bandwidth available on the FPGA by 10× to reduce memory load times. We observe that reducing operator latency improves performance marginally from 2.4× to 2.9×. This suggests that a significant portion of the total cycles is due to packet-switched interconnect. In future work, we must also consider lower latency network designs to improve performance. We also observe a higher improvement in performance from 2.4× to 4.7× when we increase offchip IO bandwidth.
4.6 Conclusions

In this chapter, we show how to implement the irregular Sparse Matrix-Solve phase of SPICE on an FPGA for a range of matrix and circuit benchmarks. We demonstrate speedups of 0.6–13.4× when comparing a 25-PE parallel implementation on a Xilinx Virtex-6 LX760 FPGA with a 1-core implementation on an Intel Core i7 965 for double-precision floating-point evaluation. Our token dataflow FPGA architecture permits lightweight distributed processing of the sparse, irregular factorization graph to deliver better performance. At present, our speedups for the Sparse Matrix-Solve phase are limited primarily due to limited parallelism in the sparse dataflow graph. Additionally, we are also affected by a combination of factors including memory load times, long latency floating-point operations and poor distribution of critical path across the parallel elements.

In future work, we will investigate different numerical algorithms (e.g. iterative, domain-decomposition) to explore a different strategy to exploit higher parallelism than that available in the sparse dataflow graph. In the short term, we expect to improve FPGA speedups in the future using better placement algorithms that optimize latency of evaluation, overlapped streaming fetch from offchip-memory, and lower latency floating-point operators [91, 92]. We will also develop vector implementations of the levelized dataflow graphs to compare performance against GPUs and Intel multi-core CPUs.
Chapter 5

Iteration Control

In Chapter 3 and 4, we discussed the two computationally-intensive phases of the SPICE simulator. In this chapter, we explain how to implement the sequential, control-intensive SPICE state-machines. We also discuss a streaming approach that will permit a high-level expression of the Iteration Control computation using the SCORE framework. We show that ignoring this phase for parallelization results in mean speedups of $2.4 \times$ for the composed SPICE simulator. We caution against mapping this sequential Iteration Control computation to a lightweight embedded microcontroller (e.g. Microblaze) as it creates a performance bottleneck and decreases overall speedups to $1.9 \times$. We discuss our FPGA organization that uses a combination of static and dynamic scheduling to deliver balanced speedups of $2.6 \times$ for the integrated design.

5.1 Structure in Iteration Control Phase

As discussed in Chapter 2, SPICE is an iterative algorithm that solves non-linear differential equations. SPICE solves these equations using an iterative approach that first linearizes the non-linear circuit elements and then performs a numerical integration involving time-varying quantities. The space of algorithms for linearization and numerical integration is vast, and it covers conflicting requirements of convergence speed, accuracy and stability while demanding different amounts of computation and memory storage costs. The choice of a suitable algorithm applicable to circuit sim-
ulation is the subject of continued research and is beyond the scope of this thesis. Using the framework and methodology described in this chapter, we can support newer algorithms for managing the SPICE simulation. For the purpose of this thesis and as a proof-of-concept, we pick the algorithms used in the \texttt{spice3f5} package: the Newton-Raphson algorithm for handling non-linear elements and the Trapezoidal approximation for numerical integration.

The \texttt{spice3f5} iteration controller manages two kinds of iterative loops: (1) a loop for linearizing the non-linear elements of the circuit, and (2) another loop for advancing the timestep of the simulation. We show these loops in Figure\texttt{5.1}. The convergence conditions for the Newton-Raphson algorithm are implemented in the block (a) of the figure. SPICE employs the Newton-Raphson algorithm for computing the linear operating points of non-linear devices like diodes and transistors. The equation for next timestep calculation is implemented in block (b) of the figure. SPICE uses an adaptive timestep-control algorithm that adjusts the timestep of the simulation based on an estimate of local truncation error as previously introduced in Section \texttt{2.1.3}.
block (C), SPICE implements the dynamic breakpoint processing logic for handling source transition timesteps in the voltage and current sources. Finally, in block (D), the analysis state machines implement the loop control algorithms for performing DC and transient analysis. The Iteration Control computation constitutes a small (7%) fraction of total SPICE runtime. We reported these measurements earlier in Figure 2.9 shown in Section 2.2.

5.2 Performance Analysis

We now show the benefit of carefully parallelizing the Iteration Control phase of SPICE. In Figure 5.2 we show the runtimes of the different phases of SPICE with and without parallelization for the s641 and r4k SPICE netlists. In the “Sequential” column we observe that sequential runtime is dominated by the Model-Evaluation computation and the Iteration Control is a small 7–8% fraction of total runtime. When we parallelize the Model-Evaluation and the Sparse Matrix-Solve phases on a Virtex-6 LX760, we notice that the Iteration Control is now a significant 24–50% portion of total runtime (see Column “Ignore IterCtrl”). This is especially true for the r4k SPICE netlist where it is 50% of total runtime. This suggests, that we must parallelize this phase to achieve high overall speedup. This is very relevant as we scale to larger FPGA sizes in the future. In that case, since we have more area at our disposal, we can further parallelize the Model-Evaluation and Sparse Matrix-Solve phases of SPICE. The sequential Iteration Control phase will be an even greater fraction of parallel runtime. In column “Spatial IterCtrl” we show the effect of spatially parallelizing the Iteration Control phase on total performance. Like other FPGA designs [116], we may choose to implement the Iteration Control portion on an embedded processor e.g. Microblaze. However, in Column “Microblaze IterCtrl” for the s641 netlist, we observe that this is a poor implementation choice and results in lost speedup. We discuss performance comparison in greater detail in Section 5.6. These two distributions suggest that we must avoid a sequential Microblaze implementation of the Iteration Control phase while parallelizing this phase properly.
We present another way to look at the performance of Iteration Control in Figure 5.3. We compare the contribution of the Iteration Control phase of SPICE before (Figure 5.3(a)) and after (Figure 5.3(b)) parallelizing the other two SPICE phases. We observe that once the rest of SPICE is parallelized, the contribution of the Iteration Control phase increases from 7% to 19% of total parallel runtime. Hence it is important to seek ways to extract parallelism out of this phase of SPICE to deliver overall speedups.

What will happen to overall speedups if we do not parallelize Iteration Control? In Figure 5.4, we see that a speculative $25 \times$ speedup for the Model-Evaluation and
Sparse Matrix-Solve phase of SPICE results in an overall speedup of only $\approx 9\times$. This is due to Amdahl’s Law (shown in Equation 5.1–5.2) which states that the maximum speedup possible in this case is $9.3\times$ assuming 7% of total runtime is spent in the Iteration Controller. We later show actual speedups achieved by our Virtex-6 LX760 implementation in Section 5.6.

\[
\text{Speedup} = \frac{1}{(1 - P_{frac}) + (P_{frac}/25)} \quad (5.1)
\]

\[
P_{frac} = \frac{T_{seq}(modeeval) + T_{seq}(matsolve)}{T_{seq}(modeeval) + T_{seq}(matsolve) + T_{seq}(iterctrl)} \quad (5.2)
\]

- $P_{frac}$ = Parallelized fraction of SPICE
- $T_{seq}(modeeval)$ = Sequential Model-Evaluation time
- $T_{seq}(matsolve)$ = Sequential Matrix-Solve time
- $T_{seq}(iterctrl)$ = Sequential Iteration-Control time
When architecting the complete FPGA-based system, we must implement this computation in a manner that provides best overall speedup for the complete SPICE simulator. We observed in Figure 5.3(b) that this phase is a non-dominant runtime fraction of the complete simulation. We must take care to limit the amount of area used by this component and devote FPGA resources to the compute-intensive Model-Evaluation and Sparse Matrix Solve phases of SPICE. This will allow us to achieve the highest composite performance. At the same time, we must be careful not to excessively sequentialize the SPICE control algorithms.

5.3 Components of Iteration Control Phase

We now describe the key components of the Iteration Control phase of spice3f5. We show the precise convergence conditions and the local truncation error estimate equations used in spice3f5.

Convergence Conditions The simulator declares convergence when two consecutive Newton-Raphson iterations generate solution vectors and non-linear approximations that are within a prescribed tolerance respectively. In Equation 5.3 and Equation 5.4, \(V_i\) or \(I_i\) represent the voltage or current unknowns in the \(i\)-th iteration of the Newton-Raphson loop. The convergence conditions compare quantities in iteration \(i\) with the previous iteration \((i - 1)\). We show the convergence component in block \(\Box\) in Figure 5.1.

\[
|\vec{V}_i - \vec{V}_{i-1}| \leq \text{reltol} \cdot \max (|\vec{V}_i|, |\vec{V}_{i-1}|) + \text{vntol} \tag{5.3}
\]

\[
|\vec{I}_i - \vec{I}_{i-1}| \leq \text{reltol} \cdot \max (|\vec{I}_i|, |\vec{I}_{i-1}|) + \text{abstol} \tag{5.4}
\]

The simulator also compares the non-linear function \(f_{i-1}\) in the Model-Evaluation of the previous iteration with the linear approximation \(\hat{f}_i\) of the current iteration. This condition, represented in Equation 5.5, checks the consistency of the non-linear
quantities after the Matrix-Solve step.

\[ |\hat{f}_i - f_{i-1}| \leq \text{reltol} \cdot \max(|\hat{f}_i|, |f_{i-1}|) + \text{abstol} \quad (5.5) \]

The closeness between the values is parameterized in terms of user-specified tolerance values: \text{reltol} (relative tolerance), \text{abstol} (absolute tolerance), and \text{vntol} (voltage tolerance). Typical values for these tolerance parameters are: \text{reltol} = 1e^{-3} (accuracy of 1 part in 1000), \text{abstol} = 1e^{-12} (accuracy of 1 picoampere) and \text{vntol} = 1e^{-6} (accuracy of 1 \mu\text{volt}). This means the simulator will declare convergence when the changes in voltage and current quantities get smaller than the convergence tolerances.

**Local Truncation Error (LTE):** The truncation-error-based time-stepping algorithm in \texttt{spice3f5} advances the simulation timestep based on the rate of change of the circuit quantities (e.g. charge, flux). LTE is a local estimate of accuracy of the circuit quantities. If these circuit quantities are changing too rapidly, a smaller timestep limits the amount of error. Conversely, if the circuit is mostly quiescent (typical of digital circuit simulation between clock edges), a larger timestep can be permitted without increasing error. The LTE calculation is shown in block \(\oplus\) in Figure 5.1.

The algorithm computes the stepsize \(\delta_{n+1}\) so as to achieve a target LTE for the observed divided-difference approximation \(DD_3(x)\) of the varying circuit quantity. This stepsize then advances the simulation timestep \(t_{n+1}\) as shown in Equation 5.6 and Equation 5.7. A tolerance parameter \text{trtol} provides the user additional control over tuning the stepsize. A simple state machine combines the results of convergence detection and stepsize calculation to decide how to advance the simulation.

\[ t_{n+1} = t_n + \delta_{n+1} \quad (5.6) \]

\[ \delta_{n+1} = \sqrt{\frac{\text{trtol} \cdot \epsilon}{\max \left( \frac{\text{DD}_3(x)}{12}, \text{abstol} \right)}} \quad (5.7) \]

In Equation 5.10 we show how to compute the Local Truncation Error (LTE,
\( \epsilon \), as a function of currents \( I \) and charges of \( Q \) capacitors (or fluxes and voltages of inductors). It is the maximum of two errors: current error \( (\epsilon_I) \) and charge error \( (\epsilon_Q) \).

\[
\begin{align*}
\epsilon_I &= \text{reltol} \cdot \max (|I_i|, |I_{i-1}|) + \text{abstol} \\
\epsilon_Q &= \text{reltol} \cdot \max (|Q_i|, |Q_{i-1}|, \text{chgtol}) / \delta_n \\
\epsilon &= \max (\epsilon_I, \epsilon_Q)
\end{align*}
\]

The divided difference approximation is recursively defined in Equation (5.11). For our implementation, we consider second-order Trapezoidal approximations only \((1 \leq k \leq 2)\).

\[
DD_k = \frac{DD_{k-1}(t_{n+1}) - DD_{k-1}(t_{n+1})}{\sum_{i=1}^{k} \delta_{n+1-k}}
\]

**Breakpoints:** The calculation of the timestep based on divided differences in Equation (5.7) assumes that the physical circuit quantities being approximated are continuously differentiable. However, when the source elements suddenly change value (e.g. Piece-Wise Linear sources), they introduce a discontinuity. SPICE stores these timepoints as breakpoints and forces a circuit evaluation at the breakpoint using a first-order backward-Euler integration. This ensures timepoints beyond the discontinuity are avoided. We represent this as the block labeled \( \circ \) in Figure 5.1.

**Analysis State Machines** The loop control logic is managed by the SPICE analysis state machines. These state machines are responsible for organizing the simulation steps, handling error conditions, determining convergence and announcing termination. We separate these state machines into two parts: (1) a high-level controller `spicestmc` that manages the DC and transient analysis along with the timestepping algorithm (the outer-loop in Figure 5.1), and (2) the iteration controller `nistmc` that invokes Model-Evaluation and Sparse Matrix-Solve phases at the right time (the inner loop in Figure 5.1). These are represented as \( \dagger \) in Figure 5.1.
We now show a runtime breakdown of these different components of the SPICE simulation in Figure 5.5. We observe that runtime is dominated by the calculation of convergence conditions and local truncation error. These two operators together account for 99% of total runtime. The breakpoint logic and SPICE control algorithms take up less than 1% of total runtime.

Let us now attempt to understand and identify the sources of parallelism in the Iteration Controller. We note that the Iteration Control state machine evaluation is sequential and data-dependent in nature. The exact path through the state machines depends on the numerical properties of the circuit equations and matrix solutions. However, we can indeed statically schedule and extract limited instruction-level parallelism from the individual state actions of the Iteration-Control state machines. Furthermore, the LTE calculations and convergence detection conditions are data-parallel operations over the voltage, current and charge vectors of the circuit. We can statically schedule these operations since the equations must be evaluated for all elements on the circuit vectors. These observations motivate the use of a streaming abstraction for expressing the Iteration Control computation. In this chapter, we discuss our approach for efficiently implementing the SPICE Iteration Control
Table 5.1: SCORE Compiler Optimized Instruction Counts for Iteration Control

<table>
<thead>
<tr>
<th>Operator</th>
<th>Add</th>
<th>Mult.</th>
<th>Divide</th>
<th>Sqrt.</th>
<th>If-Mux</th>
<th>Cmp.</th>
<th>Bool</th>
<th>Rest</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>converge</td>
<td>7</td>
<td>1</td>
<td>0</td>
<td>0</td>
<td>6</td>
<td>5</td>
<td>1</td>
<td>0</td>
<td>20</td>
</tr>
<tr>
<td>LTE</td>
<td>16</td>
<td>8</td>
<td>9</td>
<td>1</td>
<td>21</td>
<td>20</td>
<td>0</td>
<td>0</td>
<td>75</td>
</tr>
<tr>
<td>breakpoint</td>
<td>95</td>
<td>2</td>
<td>1</td>
<td>0</td>
<td>110</td>
<td>76</td>
<td>35</td>
<td>11</td>
<td>330</td>
</tr>
<tr>
<td>nistmc</td>
<td>2</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>8</td>
<td>7</td>
<td>5</td>
<td>2</td>
<td>24</td>
</tr>
<tr>
<td>spicestmc</td>
<td>29</td>
<td>15</td>
<td>6</td>
<td>0</td>
<td>79</td>
<td>42</td>
<td>24</td>
<td>17</td>
<td>212</td>
</tr>
<tr>
<td>Total</td>
<td>149</td>
<td>26</td>
<td>16</td>
<td>1</td>
<td>224</td>
<td>150</td>
<td>65</td>
<td>32</td>
<td>513</td>
</tr>
</tbody>
</table>

Column Rest includes floor, ceiling, and other special functions.

5.4 Iteration Control Implementation Framework

We express the SPICE Iteration Control algorithms in a stream-based framework called SCORE [10] (Stream Computation Organized for Reconfigurable Execution). The SCORE programming model allows us to capture the SPICE iteration control algorithm at a high-level of abstraction and permits exploration of different implementation configurations for the parallel SPICE solver. The streaming abstraction naturally matches the processing structure of the control algorithms and the overall composition of the solver. However, the SCORE compute model was originally designed for rapidly-reconfigurable, time-multiplexed FPGAs. Modern FPGAs offer poor dynamic reconfiguration support and are unsuitable for the coarse-grained, dynamically-reconfigurable implementation of SCORE. Consequently, we develop a new implementation model for SCORE based on resource-sharing and static scheduling. We adapt the backend flow from our Model-Evaluation infrastructure described in Chapter 3 to support dataflow graphs generated from the SCORE description of the Iteration Control computation.

SCORE allows description of streaming applications using dynamic dataflow. A SCORE program consists of a graph of operators (compute) and segments (memory) linked to each other via streams (interconnect). Computation within an operator is described as a finite-state machine (FSM). The operations within a state can be
described as a dataflow graph, while the state machine transitions are captured using a state transition graph. This suits the control-intensive nature of the SPICE iteration control algorithm.

We show the high-level SCORE representation of the SPICE Iteration Controller in Figure 5.6. We describe the control algorithms as SCORE operators and state-machines interconnected by streams. The stream connection allows pipelined, parallel evaluation of the different operators when possible. The white nodes in Figure 5.6 represent the state-machine and breakpoint logic. For calculating convergence and local truncation error, we stream voltages, currents and charges through the operation graph for the respective equations. The gray nodes are the data-parallel stateless nodes that calculate Local Truncation Error (LTE) and compute convergence as a function of voltage $\vec{x}$, current $\vec{b}$ and charge $\vec{Q}$ vectors. We represent the Model-Evaluation and Sparse Matrix-Solve phases of SPICE as black boxes. Internally these are implemented differently using FPGA organizations described earlier in Chapter 3 and Chapter 4.

In Table 5.1 we show the number of floating-point instructions and their types in the different SCORE operators. These statistics are obtained from the optimized operation graphs generated by tdfc, the SCORE compiler. As expected, we observe that the If-Mux, Comparison and Boolean instructions constitute the bulk of
<table>
<thead>
<tr>
<th>Operator</th>
<th>Total Activations/Iteration</th>
<th>Percent of Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>converge</td>
<td>1088465</td>
<td>64.394</td>
</tr>
<tr>
<td>LTE</td>
<td>601076</td>
<td>35.560</td>
</tr>
<tr>
<td>accept</td>
<td>299</td>
<td>0.017</td>
</tr>
<tr>
<td>breakpoint</td>
<td>48</td>
<td>0.002</td>
</tr>
<tr>
<td>nistmc</td>
<td>152</td>
<td>0.009</td>
</tr>
<tr>
<td>spicestmc</td>
<td>262</td>
<td>0.015</td>
</tr>
</tbody>
</table>

Table 5.2: SCORE Operator Activation Frequency for a simple Resistor-Capacitor-Diode circuit

the control-intensive computation in this phase of SPICE. We also note that we need only one \textit{SQRT} floating-point operation and no other expensive elementary floating-point functions. In Table 5.2, we show the dynamic activation counts for the different SCORE operators in the Iteration Control phase of SPICE. An activation is when a state within that SCORE operator gets fired. We observe that the LTE and Convergence calculation dominate the dynamic activation counts. This explains the runtime distribution shown in Figure 5.5.

5.5 Hybrid FPGA Architecture

We now describe the FPGA architecture for efficiently implementing the Iteration Control phase of SPICE. As identified earlier in Section 5.1, this phase of SPICE is responsible for a small fraction of total runtime. The control algorithms and SPICE state-machines are infrequent operations that are sparsely activated per iteration as shown in Table 5.2 and Figure 5.5. We also note that LTE calculation and convergence evaluation are data-parallel operations that constitute the bulk to total time spent in Iteration Control. A fully-spatial circuit implementation will be wasteful and occupy a large amount of area while staying underutilized. An inexpensive embedded microprocessor (\textit{e.g.} Microblaze) implementation that sequentially processes the computation will be too slow for the parallel design (see Section 5.6.4).

We develop a hybrid FPGA architecture that exploits the (1) streaming parallelism between SCORE operators and (2) dataflow parallelism within each SCORE
state to deliver an area-efficient implementation that achieves modest speedups. Our hybrid design combines statically-scheduled components with dynamic logic specialized for state selection and stream management. The SCORE description allows a straightforward separation of computation into static and dynamic portions. The dataflow within a state is considered for static scheduling while the data-dependent, dynamic state selection is mapped to dynamic logic. We couple SCORE with a back-end that schedules static feedforward dataflow graphs to generate configurations for each state. For certain SCORE operators, like LTE calculation and convergence evaluation, we perform loop unrolling to improve throughput. We show the architecture in Figure 5.7. The design of this architecture is similar to the design of the Model-Evaluation VLIW hardware described in Chapter 3.
5.6 Methodology and Results

We now describe our experimental methodology and explain our results. We show a high-level representation of our flow in Figure 5.8.

5.6.1 Integration with spice3f5

We re-express the SPICE simulator in the high-level SCORE framework. To verify correctness of our implementation, we perform a functional SCORE simulation of the Iteration Control algorithms and compare our results with spice3f5. We generate multi-threaded C++ code from the SCORE compiler to obtain a functionally-correct implementation of the SCORE description of Iteration Control. Then, we integrate the SCORE runtime into spice3f5 to communicate relevant SPICE state to our SCORE implementation using Inter-Process Communication (IPC). We then empirically compare the sequence of visited states in the two processes to determine correct operation. Barring a few cases involving floating-point rounding, we achieved the same operation for our SCORE implementation on a sample benchmark. This means that our SCORE implementation visited the same states, made the same branching and control decisions and generated the same timesteps. We also perform modular
verification of individual SCORE operators by letting the \texttt{spice3f5} process handle the remaining Iteration Control computation.

### 5.6.2 Mapping Flow

We implement our SCORE computation on the FPGA in two ways:

1. Hybrid VLIW FPGA version: Statically-scheduled custom VLIW implementation shown in Figure 5.7.


For both cases, we measure the cycles required to implement each SCORE operator. We also count the number of state activations corresponding to the SCORE operator graph of the Iteration Control computation from a \texttt{spice3f5} run. This allows us to determine the runtime of the two FPGA implementations for various circuits in our circuit benchmark set by multiplying the state activations with the cycle count per state.

\[
\text{time} = \text{clock-period} \cdot \left( \sum_{i \in \text{operator}} \left( \sum_{n \in \text{state}} \text{activations}(i,n) \cdot \text{cycles}(i,n) \right) \right)
\]

For the statically-scheduled implementation, we obtain the cycle counts from the scheduler for each state of every SCORE operator. We implement the data-parallel computation in LTE and Convergence operators using the GraphStep scheduler with Loop-Unrolling (described in Chapter 3). We implement the sequential state-machine logic in \texttt{nistmc} and \texttt{spicestmc} along with the \texttt{breakpoint} operators using a simple Dataflow scheduler (without any unrolling). We combine these two schedules to assemble the spatial implementation of the Iteration Controller. Presently, our statically-scheduled implementation operates at 200 MHz due to the limitations of the Xilinx Coregen double-precision floating-point divider. We use the scheduled cycle counts to compute the total time required for the Iteration Controller.

For the Microblaze implementation, we develop a SCORE runtime customized for the Microblaze micro-controller. We perform automated code-generation of the
Table 5.3: Resource Usage for Iteration Control Implementation (Virtex-6 LX760)

<table>
<thead>
<tr>
<th>Block</th>
<th>Area (Slices)</th>
<th>Memory (BRAMs)</th>
</tr>
</thead>
<tbody>
<tr>
<td>LTE (12 operators)</td>
<td>10917</td>
<td>28</td>
</tr>
<tr>
<td>convergence (12 operators)</td>
<td>10389</td>
<td>9</td>
</tr>
<tr>
<td>breakpoint, nistmc, spicestmc (4 operators)</td>
<td>3644</td>
<td>6</td>
</tr>
<tr>
<td>Microblaze and Peripherals</td>
<td>1504</td>
<td>16</td>
</tr>
<tr>
<td>Spatial Total (LTE + convergence + breakpoint, nistmc, spicestmc)</td>
<td>24950</td>
<td>43</td>
</tr>
<tr>
<td>Microblaze Total (LTE + convergence + Microblaze and Peripherals)</td>
<td>22810</td>
<td>53</td>
</tr>
</tbody>
</table>

SCORE description of the Iteration Control computation. The code-generator produces a flavor of C suitable for use with a light-weight embedded operating system (Xilkernel [117]). We measure the number of Microblaze clock cycles to implement each state of every SCORE operator using a hardware counter. The Xilinx Microblaze controller along with supporting logic is designed to operate at 100 MHz by Xilinx Core Generator [77]. We tabulate the cost model for these two designs in Table 5.3.

Finally, we compare both these implementations with a spice3f5 running on an Intel Core i7 965 running at 2.67 GHz. We measure the runtime of the Iteration Control phase using the PAPI 4.0 [18] performance counters on a 64-bit Linux workstation running Ubuntu Lucid Lynx 10.04.

5.6.3 Hybrid VLIW FPGA Implementation

We compute speedup of our Hybrid VLIW implementation using the formula described in Figure 5.11.

In Figure 5.9, we plot the speedup achieved by our hybrid FPGA architecture over the sequential microprocessor implementation on an Intel Core i7 965 for the Iteration Control phase in spice3f5. We are able to accelerate this phase of SPICE by 1.07–3.3× across the benchmark set (mean of 2.12×). We deliver the higher speedups of around 3.3× for the larger circuit sizes.

In Figure 5.10, we show the effect of parallelizing the Iteration Control phase of SPICE on overall SPICE performance. With 12 operators (3 ADDs, 3 MULTIPLYs,
Figure 5.9: Speedup for the Iteration Control Phase of SPICE
(12-operator Virtex-6 LX760 vs. Intel Core i7 965)

Figure 5.10: Speedup for the Complete SPICE Simulator
(Virtex-6 LX760 vs. Intel Core i7 965)

\[ \text{Speedup} = \frac{T_{\text{seq}}(\text{LTE} + \text{convergence} + \text{breakpoint} + \text{nistmc} + \text{spicestmc})}{T_{\text{spatial}}(\text{LTE} + \text{convergence} + \text{breakpoint} + \text{nistmc} + \text{spicestmc})} \]

- \( T_{\text{seq}}() \) = Sequential Iteration Control time on an Intel Core i7 965
- \( T_{\text{spatial}}() \) = Parallel time of the Hybrid VLIW FPGA Implementation on a Virtex-6 LX 760

Figure 5.11: Hybrid VLIW Implementation Speedup Calculation Equations
We now show how performance scales as we increase the amount of resources provided to the Iteration Control phase. In Figure 5.12(b) and Figure 5.12(a) we show the impact of increasing PE count (area) and Loop Unroll factor (memory) on the cycles required by the static schedule averaged for each evaluation.

### 5.6.4 Microblaze Implementation

As an alternative, we consider a Microblaze implementation of the state-machine and breakpoint-processing logic. In this arrangement, the LTE and Convergence operations for the Hybrid VLIW Implementation

3 DIVIDEs, 3 SQRTs) implemented on a Virtex-6 LX760, we can deliver a mean speedup of $2.6 \times$ across our benchmark set (1.29–8.1×). If we ignore this phase our speedups are limited to $2.4 \times$ (1.26–5.9×). Thus parallelizing this phase improves overall performance by a mean of 8% (max 35%). In Chapter 6, we discuss overlapping the LTE and Convergence operations with the Model-Evaluation and Sparse Matrix-Solve phase as a strategy for improving composite SPICE speedups. Finally, if we could arbitrarily parallelize the Iteration Controller (without overlapping), we can achieve mean speedups of $3 \times$ (max 11×). Thus, with sufficient parallel hardware, we can improve performance of the composite SPICE implementation by a mean of 25% (max 83%) compared to the sequential microprocessor implementation.
\[
\text{Speedup} = \frac{T_{\text{seq}}(\text{LTE} + \text{convergence} + \text{breakpoint} + \text{nistmc} + \text{spicestmc})}{T_{\text{spatial}}(\text{LTE} + \text{convergence}) + T_{\text{microblaze}}(\text{breakpoint} + \text{nistmc} + \text{spicestmc})}
\]

\[T_{\text{seq}}() = \text{Sequential Iteration Control time}\]
\[T_{\text{spatial}}() = \text{Parallel time of the Hybrid VLIW Implementation}\]
\[T_{\text{microblaze}}() = \text{Sequential time of the Microblaze Implementation}\]

Figure 5.13: Microblaze Speedup Calculation Equations

operations continue to be implemented over statically-scheduled hardware. A Microblaze implementation of the data-parallel, floating-point intensive computation in LTE and Convergence blocks will result in extremely poor performance that is substantially worse than what we present here. Hence, we do not consider that implementation for our comparisons. The embedded Microblaze controller runs at 100 MHz and sequentially processes the infrequent state-machine and breakpoint logic using 3734 slices including supporting logic. We compute speedup for this phase using the formula represented in Figure 5.13.

The performance of this lightweight implementation is shown in Figure 5.14. Unfortunately, this implementation actually slows down the computation by as much as 30× for small circuits while delivering speedups of 2.9× for the larger benchmarks. In contrast, we can achieve speedups for all circuit sizes for the statically-scheduled hybrid FPGA architecture as shown in Figure 5.9. The reasons for this slowdown include (1) lower clock frequency of the processor, (2) sequential nature of the processor architecture and (3) poor double-precision floating-point support (10s-100s of cycles for double-precision floating-point functions). We expect a faster embedded controller with superior double-precision support to deliver better performance.

5.6.5 Comparing Application-Level Impact

In Figure 5.15 we summarize the speedups achieved for the complete SPICE simulator under three scenarios: (1) no Iteration-Control parallelization (2) Microblaze implementation of Iteration-Control and (3) Hybrid VLIW Implementation of Iteration-
Figure 5.14: Speedup for the Microblaze Implementation of the Iteration Control Phase of SPICE (increasing circuit size)

Figure 5.15: Comparing Microblaze Speedup with other Implementations
(Virtex-6 LX760 vs. Intel Core i7 965)
Control. It is clear that the spatial implementation provides the highest overall mean speedups of $2.6 \times$. The Microblaze implementation is outperformed by even the Sequential implementation of Iteration Control. We are able to achieve a mere $1.9 \times$ mean speedups ($0.94-7.8 \times$) across our benchmark set if we implement the Iteration Controller on the Microblaze. This approach only delivers this speedup for very large circuits where the Iteration Control phase is a tiny fraction of total runtime. For the circuits in our benchmark set, even the naïve Sequential implementation of Iteration Control achieves a mean speedup of $2.4 \times$.

5.7 Conclusions

In this chapter, we showed how to express and parallelize the Iteration Control phase of SPICE using the streaming SCORE framework. If we do not parallelize the Iteration Controller and implement it on the host CPU, our overall SPICE speedups are limited to $2.4 \times$ (max $6 \times$). If we implement the Iteration Controller on an embedded, sequential Microblaze controller, our mean speedups get reduced to $1.9 \times$ (max $7.8 \times$). A parallel implementation of the Iteration Controller that exploits data-parallelism and instruction-level parallelism in this phase of SPICE allows the complete SPICE simulator to achieve composite speedups of $2.6 \times$ (max $8.1 \times$).
Chapter 6

System Organization

In this chapter, we explain the complete mapping flow and organization of the FPGA-based system for accelerating SPICE. In the previous chapters, we studied the designs of the individual SPICE phases. We now show how to compose parallelism in the different phases together to assemble the complete design on a single FPGA.

6.1 Modified SPICE Solver

We have previously shown how to parallelize and implement the three phases of SPICE: Model-Evaluation in Chapter 3, Sparse Matrix-Solve in Chapter 4 and the Iteration Control in Chapter 5. We suitably modified the simulation flow to allow us to expose the parallelism available in the different SPICE phases for parallel operation. We perform offline analysis and optimization of the non-linear device models and the Iteration Control algorithms to generate static schedules for FPGA implementations. We also change runtime behavior to include a one-time analysis of the sparse matrix structure to extract parallel dataflow graphs to be shared across all factorizations. We show this modified flow in Figure 6.1. In the figure, we partition processing into Offline and Runtime components depending on the binding time of the parallelism information into our FPGA solver. We compile the Iteration Controls state-machines and algorithms from a high-level SCORE description in Step 1a to generate a static configuration. This FPGA configuration is used in Step 1b at Runtime to make convergence and timestepping decisions. We generate optimized dataflow graphs for
Model-Evaluation from the high-level Verilog-AMS description in Step 2a. Based on the type of non-linear device used in the circuit being simulated, we load the appropriate configuration for use in Step 2b. We perform matrix reordering and symbolic analysis of the sparse circuit matrix at Runtime in Step 3a to extract the sparse dataflow graph. This only needs to be done once for the entire simulation run at the beginning of the simulation. This sparse dataflow graph is reused across all factorizations in Step 3b.

### 6.2 FPGA Mapping Flow

We now explain our FPGA mapping flow for generating a configuration for the FPGA for implementing SPICE. We show the complete FPGA mapping flow in Figure 6.2. At a high level, our FPGA flow is organized into paths that are customized for the specific SPICE phase. These are represented with unique characteristic colors in Figure 6.2. In this figure, we also partition the mapping flow based on binding time of...
the operation being performed. We can optimize the Model-Evaluation computation as well as compile the Iteration-Control processing statically offline. However, most of the Sparse Matrix-Solve processing must be handled at runtime on a per-circuit basis.

We statically generate a hardware configuration for the parallel SPICE simulator and build the onchip memory images for each circuit at runtime. This means that we need to run the time-consuming FPGA CAD tools offline to create an FPGA configuration for the simulator. This is possible because we can limit the total number of configurations and simply select the configuration to load based on the kind of circuit being simulated. We list some observations about the simulator that make this possible:

- SPICE circuits in most real-world environments as well as our benchmark set use a single type of non-linear transistor model (e.g. bsim3, bsim4 or mos3). Moreover, the bulk of the model-specific parameters are constant across all non-linear devices.
This implies that we need a single, constant-folded static-schedule for non-linear device processing on our VLIW architecture. This architecture will load the device-specific parameters at execution time as described in Chapter 3.

- The Sparse Matrix-Solve computation for all circuits is converted to a sparse dataflow graph that is processed on our Dataflow architecture. This does require a circuit-specific extraction and distribution step, but the design of the Packet-Switched Dataflow hardware does not need to be customized to the graph structure.

- The Iteration Controller is parameterized in terms of a few constants (e.g. \texttt{abstol, reltol, final\_time}) but the underlying control algorithms do not need to be recompiled per circuit.

Our mapping flow generates a few FPGA circuit configurations for SPICE customized for specific non-linear device models (e.g. \texttt{bsim3, bsim4, mos3}). We pick the appropriate configuration at the start of the simulation, when configuring the FPGA, based on the device type being used in the SPICE circuit. Once we choose a distribu-
tion of area and memory resources across the different SPICE phases, we can generate an FPGA circuit configuration that can be reused across all circuits. This includes the static VLIW configurations for the PEs and switches of the statically-scheduled Model-Evaluation and Iteration Control processing elements (output of the “Static Scheduler” blocks shown in Figure 6.2).

For each new circuit we simulate, we only need to load and program certain memory resources appropriately. In Figure 6.2, this is shown by the paths that lead into the “Memory Builder” block. For the non-linear devices and independent sources, we must store the device-specific constant parameters specified in the circuit netlist in FPGA onchip memory or offchip DRAM memory if necessary. We also need to distribute the sparse dataflow graph across the Matrix-Solve processing elements (shown by the “Placement” block in Figure 6.2) and store the graph in offchip DRAM memory when it does not fit onchip capacity. Finally, we must load a few simulation control parameters (e.g. abstol, reltol, final_time) to help the Iteration Control phase declare convergence and termination of the simulation.

6.3 FPGA System Organization and Partitioning

We represent the high-level FPGA organization of the Parallel SPICE Solver in Figure 6.3. We partition the FPGA into regions that are customized for solving the individual phases of SPICE. Our goal is to minimize the runtime of the parallel FPGA system while fitting the complete design onto a single FPGA chip. We represent the optimization problem in Equation 6.4. Since our set of feasible configurations is very small, we solve this optimization problem through a simple exhaustive search over all possible configurations and pick the configuration that minimizes time. In Figure 6.5 we show Area-Time tradeoffs for the different phases of SPICE. We tabulate the resource distribution between the three phases for the configuration with best mean performance in Table 6.1. Our composite design is projected to use $\approx 90\%$ of the total FPGA resources of the Virtex-6 LX760. While we have shown how to build the individual computing elements for the three phases of SPICE, we have not
Minimize
\[ \max(T_{par}(modeeval) + T_{par}(matsolve), T_{par}(iterctrl)) \]

Subject to
\[
\begin{align*}
\text{Area}(modeeval) + \text{Area}(matsolve) + \text{Area}(iterctrl) &< \text{Area}(\text{FPGA}) \\
\text{Mem}(modeeval) + \text{Mem}(matsolve) + \text{Mem}(iterctrl) &< \text{Mem}(\text{FPGA}) \\
\text{Area}(\text{FPGA}) &= 118560 \text{ slices} \\
\text{Memory}(\text{FPGA}) &= 720 \text{ BRAMs} \\
< A_i, T_i > &: \text{Area-Time mappings}
\end{align*}
\]

Figure 6.4: Simplified Optimization Formulation

<table>
<thead>
<tr>
<th>SPICE Phase</th>
<th>Area</th>
<th>Memory</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Slices</td>
<td>%</td>
</tr>
<tr>
<td>Model-Evaluation (bsim4)</td>
<td>62512</td>
<td>53</td>
</tr>
<tr>
<td>Sparse Matrix-Solve</td>
<td>27090</td>
<td>23</td>
</tr>
<tr>
<td>Iteration Control</td>
<td>17848</td>
<td>15</td>
</tr>
<tr>
<td><strong>Total</strong></td>
<td>107450</td>
<td>91</td>
</tr>
</tbody>
</table>

Table 6.1: FPGA Resource Distribution for complete SPICE Solver (Virtex-6 LX760)

conducted the final engineering effort required to integrate the whole design along with the software infrastructure required to support the integrated design.

6.4 Complete Speedups

We show complete FPGA speedups for the integrated design on a Virtex-6 LX760 FPGA in Figure 6.7. We use the performance model described in Equation 6.6 to calculate speedups. Unlike speedup calculations in Chapter 5 in this composite design, we are able to overlap the timestep and convergence processing components of Iteration Control phase with the Model-Evaluation and Matrix-Solve phases of SPICE. These operations operate on the conductance and current updates from Model-Evaluation along with voltage and current vectors from the Matrix-Solve in a streaming fashion. With these changes, we observe a mean speedup of 2.8× across our benchmark set with a peak speedup of 11× for the largest benchmark. As we
Table 6.2: Area and Power consumption of the two architectures

<table>
<thead>
<tr>
<th>Architecture</th>
<th>Area ($mm^2$)</th>
<th>Power (Watts)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Intel Core i7 965</td>
<td>263</td>
<td>130</td>
</tr>
<tr>
<td>Xilinx Virtex-6 LX760</td>
<td>576</td>
<td>35</td>
</tr>
</tbody>
</table>

need to simulate increasingly larger circuits, we are particularly concerned with the performance of the parallel FPGA solution at large circuit sizes. If we neglect the three small benchmarks s27, mux8 and ringosc, the mean speedups increase slightly to 3×.

### 6.5 Comparing different Figures of Merit

The primary figure of merit for this thesis is **speedup per chip**. However, we can compare the two architectures using other important cost metrics such as area and energy. We tabulate the estimated area of the chip and power consumption of the two architectures in Table 6.2. We observe that the FPGA is able to deliver its speedup using a larger chip ($\approx 2\times$) and lower power ($\approx 3.5\times$). We estimate area assuming that
\[
\text{Speedup} = \frac{T_{\text{seq}(\text{modeleval})} + T_{\text{seq}(\text{matsolve})} + T_{\text{seq}(\text{iterctrl})}}{\max(T_{\text{par}(\text{modeleval})} + T_{\text{par}(\text{matsolve})}, T_{\text{par}(\text{iterctrl})}) + T_{\text{parseq}(\text{iterctrl})}}
\]

- \(T_{\text{seq}(\text{modeleval})}\) = Sequential Model-Evaluation time
- \(T_{\text{seq}(\text{matsolve})}\) = Sequential Matrix-Solve time
- \(T_{\text{seq}(\text{iterctrl})}\) = Sequential Iteration-Control time
- \(T_{\text{par}(\text{modeleval})}\) = Parallel Model-Evaluation time
- \(T_{\text{par}(\text{matsolve})}\) = Parallel Matrix-Solve time
- \(T_{\text{par}(\text{iterctrl})}\) = Iteration-Control time for parallelizable portion
- \(T_{\text{parseq}(\text{iterctrl})}\) = Iteration-Control time for sequential portion

Figure 6.6: Speedup Calculation Equation

Figure 6.7: Complete Speedup (Estimated) for SPICE on Xilinx Virtex-6 LX760 FPGA
\[
\frac{\text{Area}(6\text{LUT}) \cdot \text{Number}(6\text{LUT})}{\text{Area}(4\text{LUT}) \cdot \text{Number}(4\text{LUT})} = 1.7 \quad (\text{from } \text{[118]}) \quad (6.1)
\]
\[
\frac{\text{Num}(4\text{LUT})}{\text{Num}(6\text{LUT})} = 1.4 \quad (\text{from } ABC \text{ synthesis}) \quad (6.2)
\]
\[
\text{Area}(4\text{LUT}) \approx 10^6 \lambda^2 \quad (\text{from } \text{[7]}) \quad (6.3)
\]
\[
\Rightarrow \text{Area}(6\text{LUT}) \approx 1.7 \cdot 1.4 \cdot 10^6 \lambda^2 \quad (6.4)
\]
\[
\Rightarrow \text{Area}(6\text{LUT}) \approx 2.4 \cdot 10^6 \lambda^2 \quad (6.5)
\]

Figure 6.8: Estimating area of a 6-LUT

\[
\text{Virtex6 LX760} = 118560 \text{ SLICES} \quad (6.6)
\]
\[
= 474240 \text{ 6LUTs} \quad (6.7)
\]
\[
= 474240 \cdot 2.4 \cdot 10^6 \lambda^2 \quad (6.8)
\]
\[
= 474240 \cdot 2.4 \cdot 10^6 \cdot (20nm)^2 \quad (6.9)
\]
\[
= 471 \text{mm}^2 \quad (6.10)
\]

Figure 6.9: Estimating area of FPGA die

the largest FPGAs are manufactured near the reticule size (24mm×24mm=576mm²).

We validate the ballpark estimate with a bottom-up area calculation from Equation 6.1 and Figure 6.9. We show the steps of this calculation in Figure 6.8. We then estimate the actual area of the FPGA die in Figure 6.9 (estimated 471mm² close to reticule limit of 576mm²). Using these estimates, we can compare area efficiency (performance per unit area) as shown in Figure 6.10. Thus, when considering speedup per unit area, the FPGA is able to deliver lower speedups up to 4.8× with a geometric mean speedup of only 1.05×.

We estimate power consumption of the FPGA using the Xilinx XPower tool assuming 20% activity on the Flip-Flops, Onchip-Memory ports and external IO ports. We compute energy savings (\(\text{Energy} = \text{Power} \times \text{Time}\), as shown in Figure 6.11) of our FPGA design with the microprocessor. We observe that the FPGA consumes up to 40.9×(geomean 8.9×) lower energy than the microprocessor implementation.
Figure 6.10: Speedup per Unit Area (Estimated) for SPICE on Xilinx Virtex-6 LX760 FPGA

Figure 6.11: Energy Savings (Estimated) for SPICE on Xilinx Virtex-6 LX760 FPGA
6.6 FPGA Capacity Scaling Analysis

Our auto-tuner allows our design to easily scale to larger FPGA sizes. The larger Virtex-7 XC7V2000T FPGA (part of the new Virtex-7 series [11] recently announced by Xilinx) can accommodate a design with 2× the resources required of our current Virtex-6 LX760 design. We estimate speedups achieved by this FPGA in Figure 6.12.

We observe speedups between 0.2–14.1× across our benchmark set (mean speedup of 3×). This represents a modest increase of 6% in mean speedup with a 2× increase in area. Now, if we remove the small benchmarks s27, mux8, and ringosc from the mean speedup calculation, our mean speedup is 4× which represents a more respectable 30% increase in speedup with a 2× increase in area. The small benchmarks have insufficient parallelism to justify a spatial mapping and can actually slow down (≈1–2% in this case) when distributed across a large FPGA.
Chapter 7

Conclusions

The concrete goal of this thesis was to parallelize the SPICE circuit simulator using a single FPGA. In this thesis, we show how to achieve 0.2–11× speedup for SPICE circuits when comparing a Xilinx Virtex-6 LX760 FPGA with an Intel Core i7 965. We now summarize the answers to the key questions from Chapter 1 that we address in this thesis:

1. Can SPICE be parallelized? What is the potential for accelerating SPICE? We observe that there is parallelism in all phases of SPICE. Each phase of SPICE is characterized by a specific parallel pattern i.e. data-parallel computation in SPICE Model-Evaluation phase, sparse dataflow pattern in SPICE Matrix-Solve phase and streaming, sequential controller pattern in the SPICE Iteration-Control phase. Once we identify the parallel pattern, we can pick a suitable high-level framework and corresponding implementation model for exploiting that parallelism. Our performance analysis from Section 2.2 suggests that the Model-Evaluation phase is completely data-parallel (O(N) parallelism) and relatively easy to parallelize. We exploit specialization potential of device types, parameters and constants to generate optimized parallel implementations. The Sparse Matrix-Solve phase has a limited amount of parallelism (O(N^{0.7})) which is challenging to capture and exploit. In this case, we exploit the early-bound nature of the factorization graph to distribute the computation across parallel compute elements. Finally, we exploit the streaming, data-parallel components of the Iteration-Control phase to generate effective parallel implementations.
We also explore opportunities for overlapping the Iteration-Control computation with the other two phases of SPICE.

2. How do we express the irregular, parallel structure of SPICE? As identified earlier, SPICE is a composite mixture of multiple forms of parallel patterns. Instead of attempting to scavenge parallelism from dusty-deck C code we choose to re-express SPICE in these high-level languages. We capture this parallelism directly from the circuit computation graph (Verilog-AMS, KLU solver and SCORE). We compile the high-level Verilog-AMS descriptions of the device models into the static dataflow graph for data-parallel Model Evaluation computation. We extract dataflow parallelism in the Sparse Matrix-Solve phase directly from the SPICE circuit using the KLU solver and distribute this parallelism across a Dataflow FPGA architecture. This form of sparse, irregular parallelism is unsuitable for direct implementation on multi-core and GPU architectures due to high synchronization overheads. Finally, we describe the SPICE Iteration-Control algorithms in SCORE and identify additional opportunities for overlapped, parallel evaluation.

3. How do we use FPGAs to exploit the parallelism available in SPICE? FPGAs can be configured to support any kind of spatial parallelism in the application. Once we capture the parallelism in SPICE, we implement this parallelism on a combination of spatial architectures customized for each phase of SPICE i.e. Custom VLIW architecture for SPICE Model-Evaluation phase, Token Dataflow architecture for SPICE Matrix-Solve phase and a Hybrid VLIW architecture for the SPICE Iteration Control phase of SPICE. We exploit the high memory bandwidth available from hundreds of distributed onchip Block-RAMs on the FPGA and spatially implement the communication operation in the different phases using either a time-multiplexed or a packet-switched network. Our framework allows us to compose these heterogeneous implementations into a single integrated design in a streaming fashion as described in Chapter 6. Apart from FPGAs, we also explore multiple implementation tar-
gets for Model-Evaluation phase including GPUs, IBM Cell, Sun Niagara and Intel Multi-core processors with our automated code-generation and auto-tuning framework.

4. Will FPGAs outperform conventional multi-core architectures for parallel SPICE?

We show how to accelerate SPICE by $0.2\text{–}11 \times$ across a range of benchmark circuits when comparing a Xilinx Virtex-6 LX760 to the state-of-the-art quad-core Intel Core i7 965. We can parallelize large circuits with low fanout very effectively. Small circuits have insufficient parallelism to justify a spatial implementation. Similarly, circuits with high-fanout nets perform poorly and deliver limited speedups. Thus, for well-behaved circuits we can outperform the multi-cores by almost an order of magnitude using a single chip. We expect multi-core architectures to be able to accelerate the Model-Evaluation phase with additional parallel cores (see Figure 3.26(a)). The key performance bottleneck in our current spatial design is the Sparse Matrix-Solve phase. We believe efficient implementation of this sparse, irregular parallelism on multi-processing architectures will be possible only with suitable low-latency, message-passing network support [119]. As we scale to larger FPGA capacities, our auto-tuning framework will enable automated performance scaling of the SPICE design to the newer, larger FPGAs.

Apart from showing how to parallelize SPICE using FPGAs, we are also broadly interested in understanding the design space for mapping any parallel application to suitable parallel organizations. We must describe applications like SPICE using a composition of multiple parallel patterns using suitable high-level languages. We must emphasize ease of composability using frameworks such as SCORE for integrating the complete design and delivering the overall parallel solution. To fully exploit available parallelism we must be able to customize the underlying compute organization to match the pattern of parallelism in the application (e.g. Dataflow architectures for sparse irregular computation, VLIW architectures for static data-parallel computation and Streaming architectures for composing heterogeneous kinds
of parallelism). We expect this thesis to serve as a starting point for further research in this direction.

### 7.1 Contributions

We summarize the key quantitative contributions of this thesis:

1. **Complete simulator**: We accelerate the complete double-precision implementation of the SPICE simulator by $0.2–11\times$ when comparing a Xilinx Virtex-6 LX760 with an Intel Core i7 965 processor for a Double-Precision evaluation. We also deliver an estimated energy saving of up to $40.9\times(8.9\times$ geomean) when comparing the same architectures.

2. **Model-Evaluation Phase**: We demonstrate a speedup of $1.4–23.1\times$ across a range of non-linear device models when comparing Double-Precision implementations on a Virtex-6 LX760 compared to an Intel Core i7 965. We also deliver speedups of $4.5–123.5\times$ for a Virtex-5 LX330, $10.1–63.9\times$ for an NVIDIA 9600GT GPU, $0.4–6\times$ for an ATI FireGL 5700 GPU, $3.8–16.2\times$ for an IBM Cell and $0.4–1.4\times$ for a Sun Niagara 2 architectures when comparing Single-Precision evaluation across these architectures at 55nm–65nm technology. We also show speedups of $4.5–111.6\times$ for a Virtex-6 LX760, $13.1–133.1\times$ for an NVIDIA GTX285 GPU and $2.8–1200\times$ for an ATI Firestream 9270 GPU when comparing single-precision evaluation on architectures at 40–55nm technology.

3. **Sparse Matrix-Solve Phase**: We show how to improve the performance of irregular, sparse matrix factorization by $0.6–13.4\times$ when comparing a 25-PE parallel implementation on a Xilinx Virtex-6 LX760 FPGA with a 1-core implementation on an Intel Core i7 965 for double-precision floating-point evaluation.

4. **Iteration-Control Phase**: Finally, we deliver $1.07–3.3\times$ (mean $2.12\times$) reduction in runtime for the SPICE Iteration-Control algorithms when comparing a Xilinx Virtex-6 LX760 with an Intel Core i7 965.
7.2 Lessons

Finally, we highlight some key lessons from the thesis:

**Sparse Matrix Solve Performance Scaling:** The matrix factorization computation is the key performance bottleneck for parallelization. Our Token Dataflow architecture is only able to deliver a mean speedup of only $2.4 \times$ for our benchmark matrices at 25 PEs. We are unable to significantly improve performance at larger PE counts (see Section 4.5). The scheduled latency of the critical chain of dependencies through the factorization graph limits overall performance. While the logical latency of the factorization graph scales as $O(N^{0.7})$ (see Section 2.2), the FPGA floating-point operator latency (8 cycles for add, 10 cycles for multiply) and comparable network latency (9 cycles including switch and wire delays) result in high latency paths. In future work, we need to identify ways for reducing this physical mapped latency (and possible even the logical latency of the graph).

**Auto-Tuning:** Our parallel framework is capable of exploring multiple parallel architectures and scaling the design to larger FPGAs with no architecture-specific manual customization. For the FPGA implementations, the auto-tuner is provided a resource constraint for selecting the best design. The auto-tuner enables us to automate the process of considering multiple implementation choices and selecting the best one.

**GPUs:** It may appear that GPUs are an ideal target for implementing data-parallel Model-Evaluation computation. However, we observe that FPGAs can outperform GPUs for most cases and are slower than GPUs in a few cases by $2$–$3 \times$ in the best case. GPUs work well for regular operations rich in floating-point multiply-add with high arithmetic intensity (few memory operations per compute operation). Model-Evaluation computation is characterized by a mixture of elementary floating-point functions (e.g. exponential, logarithm), relatively large amount of internal state and high thread divergence due to control-flow statements. Furthermore, GPUs lack a flexible, low-latency, message-passing network for rapidly moving data between the ALUs. Our exploratory mapping studies for implementing vectorized Sparse Matrix-
Solve computation on the GPU yielded very poor results. Finally, GPUs consume up to an order of magnitude more power than the FPGA, resulting in far lower energy efficiency for sparse factorization computation. This makes them unsuitable for implementing sparse, irregular, fine-grained parallelism. This suggests that GPUs are a poor target for accelerating the complete SPICE simulator by themselves. They may work well in tandem with a CPU and/or FPGA as part of the complete system for accelerating SPICE.

**Programming FPGAs for General-Purpose Applications:** In this thesis, we demonstrate how to parallelize SPICE on FPGAs as a stepping stone towards understanding how to broadly use FPGAs for general-purpose computing. Typical real-world applications are a heterogeneous mix of multiple parallel patterns. We must expose the pattern structure using suitable languages or language extensions (e.g. Model-Evaluation computation using Verilog-AMS, streaming composition using SCORE TDF language). These patterns must be efficiently supported in a few high-level programming languages. Modern programming languages support few patterns of parallelism with libraries (e.g. OpenMP) that are added as an after thought. When programming FPGAs today, we must manually explore multiple implementation configurations. This requires expertise and experience for good mapping results. We believe auto-tuners need to play a vital role in hiding this complexity and automating the task of searching a good mapping. Thus, a combination of problem capture using pattern and design optimization auto-tuning support will enable general-purpose application development using FPGAs.
Chapter 8

Future Work

8.1 Parallel SPICE Roadmap

This thesis serves as a starting point for further research in understanding how to implement the SPICE simulator in parallel using FPGAs. Specifically, we will now identify a roadmap for additional research that are highlighted by this thesis. We also identify the appropriate categorization that we introduced in Section 2.3.

8.1.1 Precision of Model-Evaluation Phase

Double-precision floating-point operators consume a large amount of area on FPGAs. Custom floating-point or fixed-point operators (Precision) that operate at just enough precision might provide an opportunity for improving the compute density on FPGAs. For SPICE, it may be possible to apply or modify existing tools [120, 121] to estimate the precision requirements of the Model Evaluation graphs (and possibly Sparse Matrix-Solve graphs). Additional work may be necessary to make some of these tools [121] work with elementary floating-point operations. It may even be possible to implement the Model-Evaluation computation entirely in fixed-point arithmetic. These additional optimizations will help us reduce cost (e.g. area, energy) of the FPGA implementation while delivering a higher performance result.
8.1.2 Scalable Dataflow Scheduling for Matrix-Solve Phase

Sparse matrix solve operations on large matrices can generate large dataflow graphs with millions of nodes and edges. How do we rapidly distribute and schedule these operations on a dataflow architecture? Furthermore, how do we make the performance of a dataflow design scale to even larger system sizes? We believe greedy clustering algorithms [122] (Scheduling) might provide a starting point for further research. We could even investigate the opportunity for parallel clustering algorithms (possibly running on the FPGA itself) that can handle large graph sizes efficiently. Alternately, we can consider strategies for a top-down clustering strategy based on matrix column dependency graph to enable rapid scheduling. These optimizations enable us to close the performance gap with ideal parallel performance inherent in the dataflow structure.

8.1.3 Domain Decomposition for Matrix-Solve Phase

SPICE circuits for entire chips will generate large matrices for the factorization phase. To get high performance for such large matrices, we will need to partition the processing across multiple FPGAs. Offchip IO limits may constrain performance for dataflow communication between these large matrices. It will also be challenging to schedule these large graphs in a reasonable amount of time. Domain-decomposition approaches [45] (Numerical Algorithms) that break up the large matrix into multiple submatrices can offer a solution. The decomposed submatrices can be independently solved in dataflow fashion on individual FPGAs or distributed across multiple FPGAs if available.

8.1.4 Composition Languages for Integration

For SPICE, we composed the complete application using SCORE’s TDF (Task Description Format) language. This may not necessarily be the best language to capture certain control-intensive and sparse dataflow-oriented forms of computation. We need to develop languages that allow composing multiple expression patterns effortlessly.
This may be in the form of *embedded languages* or a completely new custom coordination language.

### 8.1.5 Additional Parallelization Opportunities in SPICE

We currently expose most, but not all, of the parallelism available in the SPICE simulator. We must investigate the following key opportunities for additional improvement in parallel SPICE performance:

1. We can overlap the Model-Evaluation phase with the Sparse Matrix-Solve phase of SPICE. Our streaming high-level capture in SCORE offers the ability to integrate a scheduler (*Scheduling*) that can facilitate this overlap. The scheduler needs to statically compute a suitable ordering of the device evaluation in Model-Evaluation to match the dataflow ordering in the Sparse Matrix-Solve computation.

2. Additionally, we can improve the performance of the Model-Evaluation phase with extra loop-unrolling and the use of offchip memory capacity (*Scheduling*). We need to develop an extension to our VLIW architecture to migrate data offchip when necessary.


4. Apart from these approaches, it may be useful to consider completely different algorithms (iterative matrix-free fixed-point simulation [124] or constant-Jacobian [125]; *SPICE Algorithms*) for SPICE simulations that completely eliminate the need for performing per-iteration matrix factorization.
8.2 Broad Goals

The goals of this thesis are not limited to implementing SPICE or FPGAs. Ultimately, we are interested in understanding how to implement different kinds of computation on multiple computing organizations e.g. multi-cores, GPUs. We now identify a few areas of research that can help us support this long-term vision. These areas are partly inspired from the thesis.

8.2.1 System Partitioning

For this thesis, we consider a solution which uses FPGA logic for the complete SPICE solver. In the future, we want develop high-level frameworks that combine domain-specific languages and auto-tuners to and build system-level backends for heterogeneous systems. We envision parallel processing systems which integrate FPGAs, GPUs and Multi-Core processors to become important for many performance-critical application. We can use SPICE as a design driver to show how to effectively program and use such systems. We sketch some system partitioning strategies in Figure 8.1.

8.2.2 Smart Auto-Tuners

When mapping to multiple parallel architectures, we used automated code-generation and auto-tuning approaches in this thesis to enable a fair comparison. This auto-tuner makes an exhaustive sweep of the parameter space which in general evaluates the cartesian product set of the parameter ranges. This is needless in most cases and limits the utility to offline tuning. We may be able to use intelligent auto-tuners that efficiently prune the search space and may even make it possible to perform online auto-tuning like that in [86] or a mixed offline-online technique as in [126].

8.2.3 Fast Online Placement and Routing

In this thesis, for the Sparse Matrix-Solve computation, we currently generate the dataflow graph and perform placement on the CPU and then transfer the placed
Figure 8.1: Implementation Choices for the Complete SPICE Solver
graph over to the FPGA. This may be acceptable for long-running simulations but can become a bottleneck for short runs. We must be able to generate the dataflow graph on the FPGA as well as rapidly place the dataflow graph for locality. These ideas have been previously explored in [127]. We currently perform dynamically-routed packet-switched dataflow processing of the large matrix factorization dataflow graph. Since we know the graph structure entirely in advance, we should be able to schedule (place and route) the graph statically. Again, we can build on ideas explored earlier in [128].

8.2.4 Fast Simulation

A scalable, fast cycle-accurate software simulator is crucial for demonstrating and understanding performance scaling of novel computer architecture organizations on large real-world problems. This is important before we invest significant time and energy into building prototype hardware. We need to obtain additional insight into performance trends for large matrix workloads to expose potential bottlenecks at large problem sizes. We can exploit multi-core parallelism to build fast simulators for our Dataflow FPGA architecture.

8.2.5 Fast Design-Space Exploration

In this thesis, we show the utility and benefits of using auto-tuning for generated efficient, optimized FPGA implementation of the Model-Evaluation hardware. For our case, we can perform an exhaustive exploration of the design space due the small size of the space and quick compilation of each configuration. For general problems, we must develop smarter exploration algorithms with rapid performance estimation for faster optimization. We can even consider FPGA-based simulators [129] that can quickly provide performance estimates.
8.2.6 Dynamic Reconfiguration and Adaptation

The SPICE algorithm we use for this thesis has a mostly static compute structure. We might be able to exploit delta dataflow [130] and tradeoff accuracy of the simulation for higher performance (less work). For more general matrices (not limited to circuit simulation), we will need to support dynamic runtime pivoting for the LU factorization. This will change the dataflow structure at runtime. We will need to develop lightweight runtimes to support dynamic changes in the application compute structure. This will allow us to extend the generality of this thesis to a broader range of applications and even provide a richer quality-performance tradeoff space for SPICE.
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