Aquarius Project

Final Technical Report

Research in the System Architecture of Accelerators for the High Performance Execution of Logic Programs

Alvin M. Despain
Principal Investigator

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University of Southern California
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1. Introduction

This is the final report on research in the system architecture of accelerators for the high performance execution of logic programs. It was conducted by the Electrical Engineering - Systems Department of the University of Southern California, under award number 25879 as subcontractor to the University of California, Berkeley. The research was sponsored by the Defense Advanced Research Projects Agency under contract number N00014-88-K-0579.

The scope of this work included:

- Design of an abstract machine for the execution of Prolog, the Berkeley Abstract Machine (BAM).
- Design, simulation, and implementation of a high-performance VLSI Prolog accelerator chip, the VLSI-BAM.
- A simulator for the Aquarius-II multiprocessor.
- Release of version 1.0 of the Berkeley Extended Prolog (BXP) compiler.
- Design, implementation, evaluation, and release of the Advanced Silicon-Compiler in Prolog (ASP) System.

All of the above work was completed, as reported in the following section of this report.

It was originally proposed that this work would include the design and performance evaluation of the Aquarius-II and Aquarius-III multiprocessors, under options A-II and A-III. As these options were not funded, the research was not performed.
2. Accomplishments

2.1 Aquarius Prolog Compiler

Our work on compilation of Prolog revealed that the language can be implemented an order of magnitude more efficiently than the best existing systems, with the result that its speed approaches that of imperative languages such as C for a significant class of programs. The approach used was to encode each occurrence of a general feature of Prolog as simply as possible. The design of this system, Aquarius Prolog, is based upon four principles:

- Reduce instruction granularity. Use an execution model, the Berkeley Abstract Machine (see below), that retains the good features of the Warren Abstract Machine (WAM).
- Exploit determinism. Compile deterministic programs with efficient conditional branches. Most predicates written by human programmers are deterministic, yet previous systems often compile them in an inefficient manner by simulating conditional branching with backtracking.
- Specialize unification. Compile unification to the simplest possible code. Unification is a general pattern-matching operation that can do many things in the implementation: pass parameters, assign values to variables, allocate memory, and do conditional branching.
- Dataflow analysis. Derive type information by global dataflow analysis to support the above ideas.

The resulting Aquarius Prolog system (Appendix 1) is about five times faster than the high-performance commercial Quintus Prolog compiler. Because of limitation of the dataflow analysis system, Aquarius is not yet competitive with the C language for all programs. This can be addressed in future work.

2.2 Berkeley Abstract Machine (BAM)

The design of the Berkeley Abstract Machine (BAM) was based upon the Programmed Logic Machine (PLM), which was a straightforward microcoded implementation of the Warren Abstract Machine, the most widely-used model for the execution of Prolog. Studies of the PLM found that performance was limited by bus bandwidth. It also proved difficult to perform compiler optimizations on PLM code because of the complexity of the operations. These problems were addressed in the BAM design.

The BAM began with a general-purpose RISC architecture and added a minimal set of extensions to support high-performance Prolog execution. Exploiting these features required simultaneous development of the architecture and an optimizing compiler. While most Prolog-specific operations can be done in software, a crucial set of features that must be supported by the hardware in order to achieve the highest performance:

- Tagging of data, with tags kept in the upper four bits of a 32-bit word.
- Segmented virtual addressing.
- Separate instruction and data buses, with the data bus being double-width.
- Special instructions which can also be used in implementing other languages.
- Instructions to test and manipulate tags.
• Unification support.

The results of this study showed that the special architectural features added 10.6% to the active area of the BAM chip, while increasing performance by 70%. This study is presented in detail in Appendix 2, "Fast Prolog With an Extended General Purpose Architecture."

2.3 Advanced Silicon-Compiler in Prolog (ASP)

The Advanced Silicon-Compiler in Prolog (ASP) is a full-range hardware synthesis system. The goal of ASP is to synthesize a single-chip VLSI processor from a high-level specification of the ISA. The approach is to study a specialized vertical slice of the design space. The design of the system proceeds hierarchically. At each level, many choices are considered for each component, making it convenient to consider the process as a conversion of a conceptual AND-OR tree into an AND tree, with design decisions being the choice of a particular OR branch.

Conceptually, each level of abstraction is composed of a simulator module, a compiler module, a design program (engine) module, and a knowledge base. Each level accepts a specification in a formal specialized language and produces a more detailed and concrete specification in a different specialized language. To determine which design choices should be made, a benchmark program is provided to each level to that the developing architecture can be simulated and measured relative to the design choice.

ASP is a design automation (DA), as opposed to a computer-aided design (CAD) system. In it, the silicon compilation problem is divided into three major problem domains, behavioral, logic, and circuit. The geometric domain is concerned with the lowest level of design, the efficient layout on silicon of a particular logic design. The logic domain produces that logic design, given a behavioral (or register transfer level -- RTL) design. At the highest level, the behavioral domain generates a behavioral description of a particular ISA.

A summary of ASP is presented in Appendix 3, "A CAD Design Environment Based Upon Prolog."

2.4 Aquarius-II Simulator

As a first step toward a Prolog multiprocessor, we developed the NuSim simulator to serve as a testbed for new ideas. Based upon the VLSI-PLM, NuSim provides a framework that permits simulation at many levels, from the instruction set to the memory architecture (including caches and coherency protocols). The simulator's flexibility allows extensive instrumentation and continual updates and changes.

NuSim is an event-driven simulator, with the events being memory accesses ordered by time. This technique simulates a multiprocessor using a uniprocessor. The simulator consists of 16,000 lines of C code and two small machine-dependent routines to save and restore the coroutine stacks. It is fairly portable, currently running under 4.3 BSD Unix on the VAX 785 and the Sun 3, and under System V Unix on an Intel 396-based personal computer.

In Appendix 4, "The Validation of a Multiprocessor Simulator," we report on validating NuSim with respect to the VPSim uniprocessor simulator.
3. Summary

Under this subcontract, the University of Southern California has performed research in accelerators for the high-performance execution of Prolog programs, including compilation techniques, accelerator architecture, multiprocessor design, and application to design automation.

In particular, this project included the design and implementation for a microprocessor for the high-performance execution of Prolog, implementation of a simulator for the Aquarius-II multiprocessor, release of the Aquarius Prolog Compiler, and design, evaluation, and release of the ASP System.
4. References

The following references report the work accomplished under this contract and are attached as appendices:


Appendix 1

"Can Logic Programming Execute as Fast as Imperative Programming?"

Peter L. Van Roy

Ph.D. Dissertation, University of California, Berkeley, November 1990
Can Logic Programming Execute as Fast as Imperative Programming?

By

Peter Lodewijk Van Roy
Graduate (Vrije Universiteit Brussel, Belgium) 1983
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Can Logic Programming Execute as Fast as Imperative Programming?

Peter Lodewijk Van Roy

ABSTRACT

The purpose of this dissertation is to provide constructive proof that the logic programming language Prolog can be implemented an order of magnitude more efficiently than the best previous systems, so that its speed approaches imperative languages such as C, for a significant class of problems. The driving force in the design is to encode each occurrence of a general feature of Prolog as simply as possible. The resulting system, Aquarius Prolog, is about five times faster than Quintus Prolog, a high performance commercial system, on a set of representative programs. The design is based on the following ideas:

(1) Reduce instruction granularity. Use an execution model, the Berkeley Abstract Machine (BAM), that retains the good features of the Warren Abstract Machine (WAM), a standard execution model for Prolog, but is more easily optimized and closer to a real machine.

(2) Exploit determinism. Compile deterministic programs with efficient conditional branches. Most predicates written by human programmers are deterministic, yet previous systems often compile them in an inefficient manner by simulating conditional branching with backtracking.

(3) Specialize unification. Compile unification to the simplest possible code. Unification is a general pattern-matching operation that can do many things in the implementation: pass parameters, assign values to variables, allocate memory, and do conditional branching.

(4) Dataflow analysis. Derive type information by global dataflow analysis to support these ideas.

Because of limitations of the dataflow analysis, the system is not yet competitive with the C language for all programs. I outline the work that is needed to close the remaining gap.

Alvin M. Despain (Committee Chairman)
Acknowledgments

This project has been an enriching experience in many ways. It was a privilege to be part of a team consisting of so many talented people, and I learned much from them. It was by trial and error that I learned how to manage the design of a large program that does not all fit into my head at once. Interaction with my colleagues encouraged the development of the formal specifications of BAM syntax and semantics, which greatly eased interfacing the compiler with the rest of the system. The use of the compiler by several colleagues, in particular the development of the run-time system in Prolog by Ralph Haygood, improved its robustness.

I wish to thank all those who have contributed in some way to this work. Al Despain is a wonderful advisor and a source of inspiration to all his students. Paul Hilfinger's fine-tooth comb was invaluable. Bruce Holmer's unfailing sharpness of thought was a strong support. I also would like to thank many friends, especially Ariel, Bernt, Francis, Hervé, Josh, Mireille, Sue, and Dr. D. von Tischüegel. Veel dank ook aan mijn familie, et gros bisous pour Brigitte.

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Chapter 1

Introduction

"You’re given the form, but you have to write the sonnet yourself. What you say is completely up to you."
-Madeleine L’Engle, A Wrinkle In Time

1. Thesis statement

The purpose of this dissertation is to provide constructive proof that the logic programming language Prolog can be implemented an order of magnitude more efficiently than the best previous systems, so that its speed approaches imperative languages such as C for a significant class of problems.

The motivation for logic programming is to let programmers describe what they want separately from how to get it. It is based on the insight that any algorithm consists of two parts: a logical specification (the logic) and a description of how to execute this specification (the control). This is summarized by Kowalski’s well-known equation Algorithm = Logic + Control [40]. Logic programs are statements describing properties of the desired result, with the control supplied by the underlying system. The hope is that much of the control can be automatically provided by the system, and that what remains is cleanly separated from the logic. The descriptive power of this approach is high and it lends itself well to analysis. This is a step up from programming in imperative languages (like C or Pascal) because the system takes care of low-level details of how to execute the statements.

Many logic languages have been proposed. Of these the most popular is Prolog, which was originally created to solve problems in natural language understanding. It has successful commercial implementations and an active user community. Programming it is well understood and a consensus has developed regarding good programming style. The semantics of Prolog strike a balance between efficient implementation and logical completeness [42,82]. It attempts to make programming in a subset of first-order logic practical. It is a naive theorem prover but a useful programming language because of its mathematical foundation, its simplicity, and its efficient implementation of the powerful concepts of unification (pattern matching) and search (backtracking).
Prolog is being applied in such diverse areas as expert systems, natural language understanding, theorem proving [57], deductive databases, CAD tool design, and compiler writing [22]. Examples of successful applications are AUNT, a universal netlist translator [59], Chat-80, a natural language query system [81], and diverse in-house expert systems and CAD tools. Grammars based on unification have become popular in natural language analysis [55, 56]. Important work in the area of languages with implicit parallelism is based on variants of Prolog. Our research group has used Prolog successfully in the development of tools for architecture analysis [12, 16, 35], in compilation [19, 73, 76], and in silicon compilation [11].

Prolog was developed in the early 70's by Colmerauer and his associates [38]. This early system was an interpreter. David Warren's work in the late 70's resulted in the first Prolog compiler [80]. The syntax and semantics of this compiler have become the de facto standard in the logic programming community, commonly known as the Edinburgh standard. Warren's later work on Prolog implementation culminated in the development of the Warren Abstract Machine (WAM) in 1983 [82], an execution model that has become a standard for Prolog implementation.

However, these implementations are an order of magnitude slower than imperative languages. As a result, the practical application of logic programming has reached a crossroads. On the one hand, it could degenerate into an interesting academic subculture, with little use in the real world. Or it could flourish as a practical tool. The choice between these two directions depends crucially on improving the execution efficiency. Theoretical and experimental work suggests that this is feasible—that it is possible for an implementation of Prolog to use the powerful features of logic programming only where they are needed. Therefore I propose the following thesis:

A program written in Prolog can execute as efficiently as its implementation in an imperative language. This relies on the development of four principles:

1. An instruction set suitable for optimization.
2. Techniques to exploit the determinism in programs.
3. Techniques to specialize unification.
4. A global dataflow analysis.
2. The Aquarius compiler

I have tested this thesis by constructing a new optimizing Prolog compiler, the Aquarius compiler.

The design goals of the compiler are (in decreasing order of importance):

1. High performance. Compiled code should execute as fast as possible.

2. Portability. The compiler's output instruction set should be easily retargetable to any sequential architecture.

3. Good programming style. The compiler should be written in Prolog in a modular and declarative style. There are few large Prolog programs that have been written in a declarative style. The compiler will be an addition to that set.

I justify the four principles given in the thesis statement in the light of the compiler design:

1. Reduce instruction granularity. To generate efficient code it is necessary to use an execution model and instruction set that allows extensive optimization. I have designed the Berkeley Abstract Machine (BAM) which retains the good features of the Warren Abstract Machine (WAM) [82], namely the data structures and execution model, but has a instruction set closer to a sequential machine architecture. This makes it easy to optimize BAM code as well as port it to a sequential architecture.

2. Exploit determinism. The majority of predicates written by human programmers are intended to be executed in a deterministic fashion, that is, to give only one solution. These predicates are in effect case statements, yet systems too often compile them inefficiently by using backtracking to simulate conditional branching. It is important to replace backtracking by conditional branching.

3. Specialize unification. Unification is the foundation of Prolog. It is a general pattern-matching operation that can match objects of any size. Its logical semantics correspond to many possible actions in an implementation, including passing parameters, assigning values to variables, allocating memory, and conditional branching. Often only one of these actions is needed, and it is important to simplify the general mechanism. For example, one of the most common actions is assigning a value to a variable, which can often be simplified to a single load or store.
Dataflow analysis. A global dataflow analysis supports techniques to exploit determinism and specialize unification by deriving information about the program at compile-time. The BAM instruction set is designed to express the optimizations possible by these techniques.

Simultaneously with the compiler, our research group has developed a new architecture, the VLSI-BAM, and its implementation. The first of several target machines for the compiler is the VLSI-BAM. The interaction between the architecture and compiler design has significantly improved both. This dissertation describes only the Aquarius compiler. A description of the VLSI-BAM and a cost/benefit analysis of its features is given elsewhere [34, 35].

3. Structure of the dissertation

The structure of the dissertation mirrors the structure of the compiler. Figure 1.1 gives an overview of this structure. Chapter 2 summarizes the Prolog language and previous techniques for its high performance execution. Chapters 3 through 6 describe and justify the design of the compiler in depth. Chapter 3 discusses its two internal languages: kernel Prolog, which is close to the source program, and the BAM, which is close to machine code. Chapter 4 gives the optimizing transformations of kernel Prolog. Chapter 5 gives the compilation of kernel Prolog into BAM. Chapter 6 gives the optimizing transformations of BAM code. Chapter 7 does a numerical evaluation of the compiler. It measures its performance on several machines, does an analysis of the effectiveness of its optimizations, and briefly compares its performance with the C language. Finally, chapter 8 gives concluding remarks and suggestions for further work.

The appendices give details about various aspects of the compiler. Appendix A is a user manual for the compiler. Appendices B and C give a formal definition of BAM syntax and semantics. Appendix D is an English description of BAM semantics. Appendix E describes the extended DCG notation, a tool that is used throughout the compiler's implementation. Appendix F lists the source code of the C and Prolog benchmarks. Appendix G lists the source code of the compiler.
Figure 1.1 - Structure of the compiler and the dissertation
4. Contributions

4.1. Demonstration of high performance Prolog execution

A demonstration that the combination of a new abstract machine (the BAM), new compilation techniques, and a global dataflow analysis gives an average speedup of five times over Quintus Prolog [58], a high performance commercial system based on the WAM. This speedup is measured with a set of medium-sized, realistic Prolog programs. For small programs the dataflow analysis does better, resulting in an average speedup of closer to seven times. For programs that use built-in predicates in a realistic manner, the average speedup is about four times, since built-in predicates are a fixed cost. The programs for which dataflow analysis provides sufficient information are competitive in speed with a good C compiler.

On the VLSI-BAM processor, programs compiled with the Aquarius compiler execute in 1/3 the cycles of the PLM [28], a special-purpose architecture implementing the WAM in microcode. Static code size is three times the PLM, which has byte-coded instructions. The WAM was implemented on SPUR, a RISC-like architecture with extensions for Lisp [8], by macro-expansion. Programs compiled with Aquarius execute in 1/7 the cycles of this implementation with 1/4 the code size [34].

4.2. Test of the thesis statement

A test of the thesis that Prolog can execute as efficiently as an imperative language. The results of this test are only partially successful. Performance has been significantly increased over previous Prolog implementations; however the system is competitive with imperative languages only for problems for which dataflow analysis is able to provide sufficient information. This is due to the following factors:

- I have imposed restrictions on the dataflow analysis to make it practical. As programs become larger, these restrictions limit the quality of the results.

- The fragility of Prolog: minor changes in program text often greatly alter the efficiency with which the program executes. This is due to the under-specification of many Prolog programs, i.e. their logical meaning rules out computations but the compiler cannot deduce all cases where this happens.
For example, often a program is deterministic (does not do backtracking) even though the compiler cannot figure it out. This can result in an enormous difference in performance: often the addition of a single cut operation or type declaration reduces the time and space needed by orders of magnitude.

- The creation and modification of large data objects. The compilation of single assignment semantics into destructive assignment (instead of copying) in the implementation, also known as the *copy avoidance problem*, is a special case of the general problem of efficiently representing time in logic. A quick solution is to use nonlogical built-in predicates such as `setarg/3` [63]. A better solution based on dataflow analysis has not yet been implemented.

- Prolog's apparent need for architectural support. A general-purpose architecture favors the implementation of an imperative language. To do a fair comparison between Prolog and an imperative language, one must take the architecture into account. For the VLSI-BAM processor, our research group has analyzed the costs and benefits of one carefully chosen set of architectural extensions.

  With a 5% increase in chip area there is a 50% increase in Prolog performance.

4.3. Development of a new abstract machine

The development of a new abstract machine for Prolog implementation, the Berkeley Abstract Machine (BAM). This abstract machine allows more optimization and gives a better match to general-purpose architectures. Its execution flow and data structures are similar to the WAM but it contains an instruction set that is much closer to the architecture of a real machine. It has been designed to allow extensive low-level optimization as well as compact encoding of operations that are common in Prolog. The BAM includes simple instructions (register-transfer operations for a tagged architecture), complex instructions (frequently needed complex operations), and embedded information (allows better translation to the assembly language of the target machine). BAM code is designed to be easily ported to general-purpose architectures. It has been ported to several platforms including the VLSI-BAM, the SPARC, the MIPS, and the MC68020.
4.4. Development of the Aquarius compiler

The development of the Aquarius compiler, a compiler for Prolog into BAM. The compiler is sufficiently robust that it is used routinely for large programs. The compiler has the following distinguishing features:

- It is written in a modular and declarative style. Global information is only used to hold information about compiler options and type declarations.
- It represents types as logical formulas and uses a simple form of deduction to propagate information and improve the generated code. This extends the usefulness of dataflow analysis, which derives information about predicates, by propagating this information inside of predicates.
- It is designed to exploit as much as possible the type information given in the input and extended by the dataflow analyzer.
- It incorporates general techniques to generate efficient deterministic code and to encode each occurrence of unification in the simplest possible form.
- It supports a class of simplified unbound variables, called uninitialized variables, which are cheaper to create and bind than standard variables.

The compiler development proceeded in parallel with the development of a new Prolog system, Aquarius Prolog [31]. For portability reasons the system is written completely in Prolog and BAM code. The Prolog component is carefully coded to make the most of the optimizations offered by the compiler.

4.5. Development of a global dataflow analyzer

The development of a global dataflow analyzer as an integral part of the compiler. The analyzer has the following properties:

- It uses abstract interpretation on a lattice. Abstract interpretation is a general technique that proceeds by mapping the values of variables in the program to a (possibly finite) set of descriptions. Execution of the program over the descriptions completes in finite time and gives information about the execution of the original program.
• It derives a small set of types that lets the compiler simplify common Prolog operations such as variable binding and unification. These types are uninitialized variables, ground terms, nonvariable terms, and recursively dereferenced terms. On a representative set of Prolog programs, the analyzer finds nontrivial types for 56% of predicate arguments: on average 23% are uninitialized (of which one third are passed in registers), 21% are ground, 10% are nonvariables, and 17% are recursively dereferenced. The sum of these numbers is greater than 56% because arguments can have multiple types.

• It provides a significant improvement in performance, reduction in static code size, and reduction in the Prolog-specific operations of trailing and dereferencing. On a representative set of Prolog programs, analysis reduces execution time by 18% and code size by 43%. Dereferencing is reduced from 11% to 9% of execution time and trailing is reduced from 2.3% to 1.3% of execution time.

• It is limited in several ways to make it practical. Its type domain is small, so it is not able to derive many useful types. It has no explicit representation for aliasing, which occurs when two terms have variables in common. This simplifies implementation of the analysis, but sacrifices potentially useful information.

4.6. Development of a tool for applicative programming

The development of a language extension to Prolog to simplify the implementation of large applicative programs (Appendix E). The extension generalizes Prolog's Definite Clause Grammar (DCG) notation to allow programming with multiple named accumulators. A preprocessor has been written and used extensively in the implementation of the compiler.
Chapter 2
Prolog and Its High Performance Execution

This chapter gives an overview of the features of the Prolog language and an idea of what it means to program in logic. It summarizes previous work in its compilation and the possibilities of improving its execution efficiency. It concludes by giving an overview of related work in the area of high performance Prolog implementation.

1. The Prolog language

This section gives a brief introduction to the language. It gives an example Prolog program, and goes on to summarize the data objects and control flow. The syntax of Prolog is defined in Figure 2.2 and the semantics are defined in Figure 2.3 (section 2.1). Sterling and Shapiro give a more detailed account of both [62], as do Pereira and Shieber [56].

A Prolog program is a set of clauses (logical sentences) written in a subset of first-order logic called Horn clause logic, which means that they can be interpreted as if-statements. A predicate is a set of clauses that defines a relation, i.e. all the clauses have the same name and arity (number of arguments). Predicates are often referred to by the pair name/arity. For example, the predicate in_tree/2 defines membership in a binary tree:

\[
\begin{align*}
\text{in_tree}(X, \text{tree}(X, _, _)) &::= X = V, \text{in_tree}(X, \text{Left}) \\
\text{in_tree}(X, \text{tree}(V, \text{Left}, \text{Right})) &::= X < V, \text{in_tree}(X, \text{Left}) \\
\text{in_tree}(X, \text{tree}(V, \text{Left}, \text{Right})) &::= X > V, \text{in_tree}(X, \text{Right})
\end{align*}
\]

(Here "::=" means if, the comma "," means and, variables begin with a capital letter, \text{tree}(V, L, R) is a compound object with three fields, and the underscore "_" is an anonymous variable whose value is ignored.) In English, the definition of \text{in_tree}/2 can be interpreted as: "\text{X is in a tree if it is equal to the node value (first clause), or if it is less than the node value and it is in the left subtree (second clause), or if it is greater than the node value and it is in the right subtree (third clause).}"

The definition of \text{in_tree}/2 is directly executable by Prolog. Depending on which arguments are inputs and which are outputs, Prolog's execution mechanism will execute the definition in different ways. The definition can be used to verify that \text{X is in a given tree, or to insert or look up \text{X in a tree.}
The execution of Prolog proceeds as a simple theorem prover. Given a query and a set of clauses, Prolog attempts to construct values for the variables in the query that make the query true. Execution proceeds depth-first, i.e. clauses in the program are tried in the order they are listed and the predicates inside each clause (called goals) are invoked from left to right. This strict order imposed on the execution makes Prolog rather weak as a theorem prover, but useful as a programming language, especially since it can be implemented very efficiently, much more so than a more general theorem prover.

1.1. Data

The data objects and their manipulation are modeled after first order logic.

1.1.1. The logical variable

A variable represents any data object. Initially the value of the variable is unknown, but it may become known by instantiation. A variable may be instantiated only once, i.e. it is single-assignment. Variables may be bound to other variables. When a variable is instantiated to a value, this value is seen by all the variables bound to it. Variables may be passed as predicate arguments or as arguments of compound data objects. The latter case is the basis of a powerful programming technique based on partial data structures which are filled in by different predicates.

1.1.2. Dynamic typing

Compound data types are first class objects, i.e. new types can be created at run-time and variables can hold values of any type. Common types are atoms (unique constants, e.g. foo, abed), integers, lists (denoted with square brackets, e.g. [Head|Tail], [a,b,c,d]), and structures (e.g. tree(X,L,R), quad(X,C,B,F)). Structures are similar to C structs or Pascal records—they have a name (called the functor) and a fixed number of arguments (called the arity). Atoms, integers, and lists are used also in Lisp.

† Not to be confused with variables of type LOGICAL in Fortran.
1.1.3. Unification

Unification is a pattern-matching operation that finds the most general common instance of two data objects. A formal definition of unification is given by Lloyd [42]. Unification is able to match compound data objects of any size in a single primitive operation. Binding of variables is done by unification. As a part of matching, the variables in the terms are instantiated to make them equal. For example, unifying \( s(X, Y, a) \) and \( s(Z, b, Z) \) (Figure 2.1) matches \( X \) with \( Z \), \( Y \) with \( b \), and \( a \) with \( Z \). The unified term is \( s(a, b, a) \). \( Y \) is equal to \( b \), and both \( X \) and \( Z \) are equal to \( a \).

1.2. Control

During execution, Prolog attempts to satisfy the clauses in the order they are listed in the program. When a predicate with more than one clause is invoked, the system remembers this in a choice point. If the system cannot make a clause true (i.e. execution fails) then it backtracks to the most recent choice point (i.e. it undoes any work done trying to satisfy that clause) and tries the next clause. Any bindings made during the attempted execution of the clause are undone. Executing the next clause may give variables different values. In a given execution path a variable may have only one value, but in different execution paths a variable may have different values. Prolog is a single-assignment language: if unification attempts
to give a variable a different value then failure causes backtracking to occur. For example, trying to unify
s(a, b) and s(X, X) will fail because the constants a and b are not equal.

There are four features that are used to manage the control flow. These are the "cut" operation (denoted by "!" in programs), the disjunction, the if-then-else construct, and negation-as-failure.

1.2.1. The cut operation

The cut operation is used to manage backtracking. A cut in the body of an clause effectively says:
"This clause is the correct choice. Do not try any of the following clauses in this predicate when backtracking." Executing a cut has the same effect in forward execution as executing true, i.e. it has no effect. But it alters the backtracking behavior. For example:

\[
p(A) :- q(A), !, r(A).
p(A) :- s(A).
\]

During execution of \( p(A) \), if \( q(A) \) succeeds then the cut is executed, which removes the choice points created in \( q(A) \) as well as the choice point created when \( p(A) \) was invoked. As a result, if \( r(A) \) fails then the whole predicate \( p(A) \) fails. If the cut were not there, then if \( r(A) \) fails execution backtracks first to \( q(A) \), and if that fails, then it backtracks further to the second clause of \( p(A) \), and only when \( s(A) \) in the second clause fails does the whole predicate \( p(A) \) fail.

1.2.2. The disjunction

A disjunction is a concise way to denote a choice between several alternatives. It is less verbose than defining a new predicate that has each alternative as a separate clause. For example:

\[
q(A) :- ( A=a ; A=b ; A=c ).
\]

This predicate returns the three solutions a, b, and c on backtracking. It is equivalent to:

\[
q(a),
q(b),
q(c).
\]
1.2.3. If-then-else

The if-then-else construct is used to denote a selection between two alternatives in a clause when it is known that if one alternative is chosen then the other will not be needed. For example, the predicate \( p(A) \) above can be written as follows with an if-then-else:

\[
p(A) :- ( \ q(A) \rightarrow r(A) ; s(A) ).
\]

This has identical semantics as the first definition. The arrow \( \rightarrow \) in an if-then-else acts as a cut that removes choice points back to the point where the if-then-else starts.

1.2.4. Negation-as-failure

Negation in Prolog is implemented by negation-as-failure, denoted by \( \neg \) (Goal). This is not a true negation in the logical sense so the symbol \( \neg \) is chosen instead of \( \text{not} \). A negated goal succeeds if the goal itself fails, and fails if the goal succeeds. For example:

\[
x(A) :- \neg t(A).
\]

The predicate \( x(A) \) will succeed only if \( t(A) \) fails. This has identical semantics as:

\[
x(A) :- t(A), !, \text{fail}.
\]

In other words, if \( t(A) \) succeeds then the \( \text{fail} \) causes failure, and the cut ensures that the second clause is not tried. If \( t(A) \) fails then the second clause is tried because the cut is not executed. Note that negation-as-failure never binds any of the variables in the goal that is negated. This is different from a purely logical negation, which must return all results that are not equal to the ones that satisfy the goal. Negation-as-failure is sound (i.e. it gives logically correct results) if the goal being negated has no unbound variables in it.

1.3. Syntax

Figure 2.2 gives a Prolog definition of the syntax of a clause. The definition does not present the names of the primitive goals that are part of the system (e.g. arithmetic or symbol table manipulation). These primitive goals are called "built-in predicates." They are defined in the Aquarius Prolog user
The figure defines the syntax after a clause has already been read and converted to Prolog's internal form. It assumes that lexical analysis and parsing have already been done. Features of Prolog that depend on the exact form of the input (i.e., operators and the exact format of atoms and variables) are not defined here.

To understand this definition it is necessary to understand the four built-in predicates that it uses. The predicates `functor(T, F, A)` and `arg(I, T, X)` are used to examine compound terms. The predicates `var(T)` and `nonvar(T)` are opposites of each other. Their meaning is straightforward: they check whether a term T is unbound or bound to a nonvariable term. For example, `var(_)` succeeds whereas `var(foo(_))` does not.
2. The principles of high performance Prolog execution

The first implementation of Prolog was developed by Colmerauer and his associates in France as a by-product of research into natural language understanding. This implementation was an interpreter. The first Prolog compiler was developed by David Warren in 1977. Somewhat later Warren developed an execution model for compiled Prolog, the Warren Abstract Machine (WAM) [82]. This was a major improvement over previous models, and it has become the de facto standard implementation technique. The WAM defines a high-level instruction set that corresponds closely to Prolog.

This section gives an overview of the operational semantics of Prolog, the principles of the WAM, a summary of its instruction set, and how to compile Prolog into it. For more detailed information, please consult Maier & Warren [43] or Ait-Kaci [1]. The execution model of the Aquarius compiler, the BAM (Chapter 3), uses data structures similar to those of the WAM and has a similar control flow, although its instruction set is different.

2.1. Operational semantics of Prolog

This section summarizes the operational semantics of Prolog. It gives a precise statement of how Prolog executes without going into details of a particular implementation. This is useful to separate the execution of Prolog from the many optimizations that are done in the WAM and BAM execution models. This section may be skipped on first reading.

Figure 2.3 defines the semantics of Prolog as a simple resolution-based theorem prover. For clarity, the definition has been limited in the following ways: It does not assume any particular representation of terms. It does not show the implementation of cut, disjunctions, if-then-else, negation-as-failure, or built-in predicates. It assumes that variables are renamed when necessary to avoid conflicts. It assumes that failed unifications do not bind any variables. It assumes also that the variable bindings formed in successful unifications are accumulated until the end of the computation, so that the final bindings give the computed answer.

Terminology: A goal $G$ is a predicate call, which is similar to a procedure call. A resolvent $R$ is a list of goals $\{ G_1, G_2, \ldots, G_r \}$. The query $Q$ is the goal that starts the execution. The program is a list of


```plaintext
function prolog_execute(Q : goal) : boolean;

var
  B : stack of pair (list of goal, integer); /* the backtrack stack */
  R : list of goal; /* the resolvent */
  i : integer; /* index into program clauses */
begin
  R := [ Q ];
  B := empty;
push (R,1) on B;
while true do begin
  /* Control step: find next clause. */
  if empty(B) then return false else pop B into (R,i);
  if (R = [ ]) then return true;
  if (i+1 ≤ n) then push (R,i+1) on B;

  /* Resolution step: try to unify with the clause. */
  /* At this point, R = [ G₁,...,Gᵢ ] and Aᵢ = (Hᵢ :- A₁,...,Aᵢᵢ) */
  /* Unify the first goal in R with clause Aᵢ. */
  unify G₁ and Aᵢ;
  if successful unification then begin
    /* In R, replace G₁ by the body of Aᵢ. */
    /* If Aᵢ does not have a body, then R is shortened by one goal */
    R := [ A₁,...,Aᵢ₋₁,Gᵢ₊₁,...,Gᵦ ];
push (R,1) on B /* proceed to next goal */
  end
end;
end;
```

Figure 2.3 - Operational definition of Prolog execution

clauses [ A₁ , A₂ , ... , Aᵦ ]. The number of clauses in the program is denoted by n. Each clause Aᵢ has a head Hᵢ, and an optional body given as a list of goals [ Aᵢ₁ , Aᵢ₂ , ... , Aᵦᵢ ].

Execution starts by setting the initial resolvent R to contain the query goal Q. In a resolution-based theorem prover, the resolvent is transformed in successive steps until (1) it becomes empty, in which case execution succeeds, (2) all the clause choices are exhausted, in which case execution fails, or (3) the program goes into an infinite loop. In a single transformation step, a goal G is taken from the current resolvent R and unified with a clause in the program. The next resolvent is obtained by replacing G by the body of the clause.

This process is nondeterministic, and much work has been done in the area of automatic theorem proving to reduce the size of its search space [7]. To get efficiency, the approach of Prolog is to restrict the
process in two ways: by always taking the first goal from \( R \) and by trying clauses in the order they are listed in the program (Figure 2.3). If no successful match is found, then the program backtracks—a previous resolvent is popped off the backtrack stack and execution continues. Therefore the execution flow of Prolog is identical to that of a procedural language, with the added ability to backtrack to earlier execution states.

The function \( \text{prolog\_execute}(Q) \) returns a boolean that indicates whether execution was successful or not (Figure 2.3). If execution was successful, then there is a set of bindings for the variables in \( Q \) that gives the result of the computation. As a definition, \( \text{prolog\_execute}(Q) \) faithfully mirrors the execution of Prolog. As an implementation, however, it is incredibly inefficient. For each clause that is tried, it pushes and pops the complete resolvent (which can be very large) on the backtrack stack. The backtrack stack grows with each successful resolution step. A practical implementation avoids much of this overhead.

- The next section describes the WAM, an execution model that is much more efficient. In the WAM, the resolvents are stored in a compact form on several stacks. Only the differences between successive resolvents are stored, so that memory usage is much less. The stack discipline is used to make backtracking efficient. The WAM also defines a representation for data items that allows an efficient implementation of unification.

2.2. Principles of the WAM

The WAM defines a mapping between the terminology of logic and of a sequential machine (Figure 2.4). Predicates correspond to procedures. Procedures are always written as one large case statement. Clauses correspond to the arms of this case statement. The scope of variable names is a single clause. (Global variables exist; however their use is inefficient and is discouraged.) Goals in a clause correspond to calls. Unification corresponds to parameter passing and assignment. Tail recursion corresponds to iteration. Features that do not map directly are the single-assignment nature and altering backtracking behavior with the cut operation.

The WAM is based on four ideas: use tagged pointers to represent dynamically typed data, optimize backtracking (exploit determinism by doing a conditional branch on the first argument), specialize
Figure 2.4 – Mapping between Prolog and an imperative language (according to WAM)

unification (instead of compiling a general unification algorithm, compile instructions that unify with a known term), and map the execution of Prolog to a real machine. The WAM defines a high-level instruction set to represent these operations.

2.2.1. Implementation of dynamic typing with tags

Data is represented by objects that fit in a register and consist of two parts: the tag field (which gives the type) and the value field (Figure 2.5). The value field is used for different purposes in different types: it gives the value of integers, the address of variables and compound terms (lists and structures), and it ensures that each atom has a unique value different from all other atoms. Unbound variables are implemented as self-referential pointers (that is, they point to themselves) or as pointers to other unbound variables. The semantics of unification allow variables to be unified together, so that they have identical values from then on. In the implementation, such variables can point to other variables. Therefore retrieving the value of a variable requires following this pointer chain to its end, an operation called dereferencing.
2.2.2. Exploit determinism

It is often possible to reduce the number of clauses of a predicate that must be tried. The WAM has instructions that hash on the value of the first argument and do a four-way branch on the tag of the first argument. These instructions avoid the execution of clauses that could not possibly unify with the goal. The four-way branch distinguishes between the four data types—variables, constants (atoms and integers), lists (cons cells), and structures. The hashing instructions hash into tables of constants and tables of structures. For example:

```
week(mondazy).
week(tuesday).
week(wednesday).
week(thursday).
week(friday).
week(saturday).
week(sunday).
```
This is a set of seven clauses with constant arguments. If the argument \( X \) of the call \( \text{week}(X) \) is a constant, then at most one clause can unify successfully with it. Hashing is used to pick that clause. If \( X \) is an unbound variable then no such optimization is possible and all clauses are tried in order.

2.2.3. Specialize unification

Most uses of unification are special cases of the general unification algorithm and can be compiled in a simpler way using information known at compile-time. For example, consider the following clause which is part of a queue-handling package:

\[
\text{\texttt{queue(X,Q) is true if Q is a queue containing the single element X.}}
\]

\[
\text{\texttt{\texttt{\texttt{\texttt{queue(X,q(s(0),(X|C),C)).}}}}}
\]

A queue is represented here as a compound term. The complexity of this term is typical of real programs. In the WAM, a unification in the source code is compiled into a sequence of high-level instructions. The compiled code executes as if the original clause had been defined as follows, with the nested term \( q/3 \) completely unraveled:

\[
\text{\texttt{queue(X,Q) :- Q=q(A,B,C), A=s(0), B=[X|C].}}
\]

(The notation \( P=Q \) means to unify the two terms \( P \) and \( Q \).) The compiled code is:

```
procedure queue/2
  get_structure q/3,r(1)  \% Q=q( <- Start unification of q/3
  unify_variable x(2)    \% A,
  unify_variable x(3)    \% B,
  unify_variable x(4)    \% C
  get_structure s/1,r(2)  \% A=s( <- Start unification of s/1
  unify_constant 0        \% 0)
  get_list x(3)          \% B=  <- Start unification of list
  unify_value x(0)       \% [X
  unify_value x(4)       \% [C
  proceed                \% <- Return to caller
```

(\( r(0) \) and \( r(1) \) are registers holding the arguments \( X \) and \( Q \), and \( r(2), r(3), ... \) are temporary registers.) Unification of the nested structure is expanded into a sequence of operations that do special cases of the general algorithm. These operations are encapsulated in the \texttt{get} and \texttt{unify} instructions.

Unification has two modes of operation: it can take apart an existing structure or it can create a new one.
In the WAM, the decision which mode to use is made at run-time in the `get` instructions by checking the type of the object being unified. A mode flag is set which affects the actions of the following `unify` instructions (up to the next `get`). A more detailed overview of the WAM instruction set is given in section 2.3 below.

2.2.4. Map execution to a real machine

The control flow of Prolog is mapped to multiple stacks. The stack representation holds the resolvents in a form that makes each resolution step as efficient as a procedure call in an imperative language. The stack-based structure allows fast recovery of memory on backtracking. As a result, some applications do not need a garbage collector.

A further optimization maps Prolog variables to registers. The variables in a clause are partitioned into three classes (temporary, permanent, and void) depending on their lifetimes. Void variables have no lifetime and need no storage. Temporary variables do not need to survive across procedure calls, so they can be stored in machine registers. Permanent variables are stored in environments (i.e. stack frames) local to a clause.

2.3. Description of the WAM

The previous section gave an overview of the ideas in the WAM, with a simple example of generated code. This section completes that description by presenting the data storage, execution state, and instruction set of the WAM in full. It also gives a larger example of generated code and a scheme to compile Prolog into WAM.

2.3.1. Memory areas

Memory of the WAM is divided into six logical areas (Figure 2.6): three stacks for the data objects, one stack to support unification, one stack to support the interaction of unification and backtracking, and one area as code space.

(1) The global stack. This stack is also known as the heap, although it follows a stack discipline. This stack holds terms (lists and structures, the compound data of Prolog).
Three kinds of data objects on stacks

<table>
<thead>
<tr>
<th>r(e)</th>
<th>r(b)</th>
<th>r(h)</th>
</tr>
</thead>
<tbody>
<tr>
<td>r(a)</td>
<td>r(tr)</td>
<td>r(pc)</td>
</tr>
<tr>
<td>r(b)</td>
<td>r(b)</td>
<td>r(cp)</td>
</tr>
<tr>
<td>r(h)</td>
<td>r(s)</td>
<td>r(s)</td>
</tr>
</tbody>
</table>

Support for unification and backtracking

<table>
<thead>
<tr>
<th>r(tr)</th>
</tr>
</thead>
</table>

Figure 2.6 – Data structures of WAM and BAM

(2) The environment stack. This stack holds environments (i.e. local frames) which contain variables local to a clause. Because of backtracking (control may return to a clause whose environment is deep inside the stack), this area does not follow a strict stack discipline, however, convention has kept this naming. (The other stacks in the WAM do follow a stack discipline.)

(3) The choice point stack. Also known as the backtrack stack, this stack holds choice points, data objects similar to closures that encapsulate the execution state for backtracking.

(4) The trail. The trail stack is used to save locations of bound variables that have to be unbound on backtracking. Saving variables is called trailing, and restoring them to unbound is called detrailimg.
Not all variables that are bound have to be trailed. A variable must only be trailed if it continues to exist on backtracking, i.e. if its location on the heap or the environment is older than the most recent choice point. This is called the trail condition.

(5) The push-down stack. This stack is used as a scratch-pad during the unification of nested compound terms.

(6) The code space. This area holds the compiled code of a program.

It is possible to vary the organization of the memory areas somewhat without changing anything substantial about the execution. For example, some Prolog systems (including the Aquarius system) combine the environment and choice point stacks into a single memory area. This area is often called the local stack. Since the push-down stack is only used during general unification, it can be kept on the top of the heap.

2.3.2. Execution state

The internal state of the WAM and the BAM is given in Table 2.1. The differences between WAM and BAM are indicated in the table: The BAM adds the register \( r(\text{tmp}_cp) \) for efficient interfacing of Prolog predicates with assembly language. The WAM adds the register \( r(s) \) and the mode flag \( \text{mode} \) for use by the unification instructions. The registers \( p(1) \) are not machine registers, but locations in the current environment, pointed to by \( r(e) \).

<table>
<thead>
<tr>
<th>Register</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>( r(e) )</td>
<td>Current environment on the environment stack.</td>
</tr>
<tr>
<td>( r(a) )</td>
<td>Top of the environment stack (WAM only).</td>
</tr>
<tr>
<td>( r(b) )</td>
<td>Top-most choice point on the choice point stack.</td>
</tr>
<tr>
<td>( r(h) )</td>
<td>Top of the heap.</td>
</tr>
<tr>
<td>( r(hb) )</td>
<td>Top of heap when top-most choice point was created.</td>
</tr>
<tr>
<td>( r(tr) )</td>
<td>Top of the trail stack.</td>
</tr>
<tr>
<td>( r(pc) )</td>
<td>Program counter.</td>
</tr>
<tr>
<td>( r(cp) )</td>
<td>Continuation pointer (return address).</td>
</tr>
<tr>
<td>( r(\text{tmp}_cp) )</td>
<td>Continuation pointer to interface with assembly (BAM only).</td>
</tr>
<tr>
<td>( r(s) )</td>
<td>Structure pointer (WAM only).</td>
</tr>
<tr>
<td>( \text{mode} )</td>
<td>Unification mode flag (value is \texttt{read} or \texttt{write}, WAM only).</td>
</tr>
<tr>
<td>( r(0), \ldots )</td>
<td>Registers for argument passing and temporary storage.</td>
</tr>
<tr>
<td>( p(0), \ldots )</td>
<td>Locations in the current environment (permanent variables).</td>
</tr>
</tbody>
</table>
2.3.3. The instruction set

Table 2.2 contains the WAM instruction set, with a brief description of what each instruction does. The `get(...)` and `unify(...)` instructions echo the `put` instructions, so their listing is abbreviated. `v(N)` is shorthand notation for `r(N)` or `p(N)`. “Globalizing” a variable (see the `put_unsafe_value` instruction) moves an unbound variable from the environment to the heap to avoid dangling pointers.

<table>
<thead>
<tr>
<th>Instruction</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>put_variable v(N), r(I)</code></td>
<td>Create a new variable, put in <code>v(N)</code> and <code>r(I)</code>.</td>
</tr>
<tr>
<td><code>put_value v(N), r(I)</code></td>
<td>Move <code>v(N)</code> to <code>r(I)</code>.</td>
</tr>
<tr>
<td><code>put_unsafe_value v(N), r(I)</code></td>
<td>Move <code>v(N)</code> to <code>r(I)</code> (and globalize).</td>
</tr>
<tr>
<td><code>put_constant C, r(I)</code></td>
<td>Move immediate value <code>C</code> to <code>r(I)</code>.</td>
</tr>
<tr>
<td><code>put_nil r(I)</code></td>
<td>Move nil to <code>r(I)</code>.</td>
</tr>
<tr>
<td><code>put_structure F, r(I)</code></td>
<td>Create functor <code>F</code>, put in <code>r(I)</code>.</td>
</tr>
<tr>
<td><code>put_list r(I)</code></td>
<td>Create a list pointer, put in <code>r(I)</code>.</td>
</tr>
<tr>
<td><code>get(...)</code>, <code>r(I)</code></td>
<td>Unify <code>(...)</code> with <code>r(I)</code>.</td>
</tr>
<tr>
<td><code>unify(...)</code></td>
<td>Unify <code>(...)</code> with structure argument.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Instruction</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>call Label, N</code></td>
<td>Call a predicate.</td>
</tr>
<tr>
<td><code>execute Label</code></td>
<td>Jump to a predicate.</td>
</tr>
<tr>
<td><code>proceed</code></td>
<td>Return from a predicate.</td>
</tr>
<tr>
<td><code>allocate</code></td>
<td>Create local stack frame.</td>
</tr>
<tr>
<td><code>deallocate</code></td>
<td>Remove local stack frame.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Instruction</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>switch_on_term V,C,L,S</code></td>
<td>Four-way branch on <code>r(0)</code>’s tag.</td>
</tr>
<tr>
<td><code>switch_on_constant N, Tbl</code></td>
<td>Hash table lookup of an atomic term in <code>r(0)</code>.</td>
</tr>
<tr>
<td><code>switch_on_structure N, Tbl</code></td>
<td>Hash table lookup of a functor in <code>r(0)</code>.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Instruction</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>try_me_else Label</code></td>
<td>Create a choice point.</td>
</tr>
<tr>
<td><code>retry_me_else Label</code></td>
<td>Change retry address.</td>
</tr>
<tr>
<td><code>trust_me_else fail</code></td>
<td>Remove top-most choice point.</td>
</tr>
<tr>
<td></td>
<td></td>
</tr>
</tbody>
</table>

2.3.4. An example of WAM code

Figure 2.7 gives the Prolog definition and the WAM instructions for the predicate `append/3`. The mapping between Prolog and WAM instructions is straightforward: the `switch` instruction branches to the right clause depending on the type of the first argument, the choice point (`try`) instructions link the clauses together, the `get` instructions unify with the head arguments, and the `unify` instructions unify

...
with the arguments of structures.

The same instruction sequence is used to take apart an existing structure (read mode) or to build a new structure (write mode). The decision which mode to use is made in the get instructions, which set a mode flag. For example, if get_list r(0) sees an unbound variable argument, it sets the flag to write mode. If it sees a list argument, it sets the flag to read mode. If it sees any other type, it fails, i.e. it backtracks by restoring state from the most recent choice point.

Choice point handling is done by the try instructions. The try_me_else L instruction creates a choice point, i.e. it saves all the machine registers on a stack in memory. It is compiled before the first clause in a predicate. It continues execution with the next instruction and backtracks to label L. (The try L instruction is identical to try_me_else, except that it continues execution at L and backtracks to the next instruction.) The retry_me_else L instruction modifies a choice point that already exists by changing the address that it jumps to on backtracking. It is compiled before all clauses after the first but not including the last. The trust_me_else fail instruction removes the top-most choice point from the stack. It is compiled before the last clause in a predicate.

2.3.5. Compiling into WAM

Compiling Prolog into WAM is straightforward because there is almost a one-to-one mapping between items in the Prolog source code and WAM instructions. Figure 2.8 gives a scheme for compiling Prolog to WAM. This compilation scheme generates suboptimal code. One can optimize it by generating switch instructions to avoid choice point creation in some cases [73].

The clauses of predicate p/3 are compiled into blocks of code that are linked together with try instructions to manage choice points. Each block consists of a sequence of get instructions to do the unification of the head arguments, followed by a sequence of put instructions to set up the arguments for each goal in the body, and a call instruction to execute the goal. The block is surrounded by allocate and deallocate instructions to create an environment for permanent variables.

The last call optimization, or LCO (also called tail recursion optimization, although it is applicable to all predicates, not just recursive ones) converts a call instruction followed by a return into a jump, i.e. it
append([], L, L).
append([X|L1], L2, [X|L3]) :- append(L1, L2, L3).

Prolog definition of append/3

append/3:
  switch_on_term V1, C1, C2, fail ; Go to V1 if r(0) is a variable.
                    ; Go to C1 if r(0) is a constant.
                    ; Go to C2 if r(0) is a list.
                    ; Fail if r(0) is a structure.
  V1: try_me_else V2 ; Create a choice point.
    Cl: get-nil r(0) ; Unify r(0) with nil.
        get_value r(1), r(2) ; Unify r(1) and r(2).
        proceed ; Return to caller.
  V2: try_me_else fail ; Remove choice point.
    C2: get_list r(0) ; Start unification of r(0) with a list.
        unify_variable r(3) ; Load head of list into r(3).
        unify_variable r(0) ; Load tail of list into r(0).
        get_list r(2) ; Start unification of r(2) with a list.
        unify_value r(3) ; Unify head of list with r(3).
        unify_variable r(2) ; Load tail of list into r(2).
        execute append/3 ; Jump to append/3 (last call optimization).

WAM code for append/3

Figure 2.7 – Compiling append/3 into WAM code

reduces memory usage on the environment stack. For recursive predicates, the LCO converts recursion into iteration, since the jump is to the first instruction of the predicate. The WAM implements a generalization of last call optimization called environment trimming that allows the environment to become smaller after each call.

3. Going beyond the WAM

Prolog implementations have made great progress in execution efficiency with the development of the WAM [82]. However, these systems are still an order of magnitude slower than implementations of popular imperative languages such as C. To improve the execution speed it is necessary to go beyond the WAM. This section discusses the limits of the WAM and how the four principles of the Aquarius compiler build on the WAM to achieve higher performance.
3.1. Reduce instruction granularity

The WAM is an elegant mapping of Prolog to a sequential machine. Its instructions encapsulate parts of the general unification algorithm. However, these parts are quite large, so that many optimizations are not possible. For example, consider the predicate:

\[ p(\text{bar}). \]

This is compiled as:

```
L1: try_me_else L2
    code for clause 1

L2: retry_me_else L3
    code for clause 2

Ln: trust_me_else fail
    code of last clause
```

Figure 2.8 – Compiling Prolog into WAM

```
p(E, F, G) :- k(X, F, F), m(S, T), ...
p(A, B, C) :- q(A, Z, H), r(W, T, B), ..., z(A, X).
```

Original Prolog predicate

```
Compiled WAM code
```

Allocate Create environment.
Allocate (get arguments) Unify with caller arguments.
Allocate (put arguments) Load arguments and call.
call q/3
Allocate (put arguments) Load arguments and call.
call r/3
allocate Remove environment.
deallocate Last call is a jump.
exectute z/2

A single compiled clause

```
```

L1: try_me_else L2
    code for clause 1

L2: retry_me_else L3
    code for clause 2

Ln: trust_me_else fail
    code of last clause

Figure 2.8 – Compiling Prolog into WAM

3.1. Reduce instruction granularity

The WAM is an elegant mapping of Prolog to a sequential machine. Its instructions encapsulate parts of the general unification algorithm. However, these parts are quite large, so that many optimizations are not possible. For example, consider the predicate:

\[ p(\text{bar}). \]

This is compiled as:
The `get_constant` instruction encapsulates a series of operations: dereference `r(0)` (follow the pointer chain to its end), test its type, and do either read mode unification (check that the value of `r(0)` is `bar`) or write mode unification (trail `r(0)` and store `bar` in its cell). All this generality is often unnecessary. For example, if the predicate `p(X)` is always called with a dereferenced atom, then unification reduces to a simple check that the value is correct. The other operations are superfluous.

The Aquarius compiler's execution model, the BAM, is designed to retain the good features of the WAM while allowing optimizations such as this one. It retains data structures and an execution flow similar to the WAM, but it has an instruction set of finer granularity (Chapter 3). The compiler does not use the WAM during compilation, but directly compiles to the BAM. It is of fine enough grain to allow extensive optimization, but it also encodes compactly the operations common in Prolog. For example, it includes an explicit dereferencing instruction, which makes it possible to reduce the amount of dereferencing significantly by only doing it when it is necessary and not in every instruction.

3.2. Exploit determinism

The majority of predicates written by human programmers are intended to give only one solution, i.e. they are deterministic. However, too often they are compiled in an inefficient manner using shallow backtracking (backtracking within a predicate to choose the correct clause), when they are really just case statements. This is inefficient since backtracking requires saving the machine state and restoring it repeatedly.

3.2.1. Measurement of determinism

Measurements of Prolog applications support these assertions:

1. *Tick* shows that choice point references constitute about half (45-60%) of all data references [69].

2. Touati and Despain show that at least 40% of all choice point and fail operations can be removed through optimization [70].

The latter result is especially interesting because it attempts to quantify how often shallow backtracking is
optimizable. It considers a choice point to be avoidable if between the access of a choice point and its removal by a cut there are no calls to non-built-in predicates, no returns, and only binding of variables that do not have to be restored on backtracking. Avoidable choice points do not have to be created because they are removed immediately. For a set of medium-sized programs, on average the following percentages of choice point creations are avoidable: 57% of the ones removed by cut, 43% of the ones removed by trust, and 48% of the ones restored by fail. The variance of these numbers is large, but the potential for optimization when these situations do occur is significant. The Aquarius compiler is able to take advantage of these optimizations and more, e.g. due to the factoring transformation (Chapter 4) it is able to compile the partition/4 predicate in Warren's quicksort benchmark [30] into deterministic code. The optimizations are synergistic, that is, doing them makes other improvements possible:

1. Less stack space is needed on the environment/choice point stack. Choice points and environments are both stored on this stack, which means that often a clause's environment is hidden underneath a more recently created choice point. When this happens the last call optimization is not able to recover space. If fewer choice points are created, then last call optimization is effective more often.

2. There are fewer memory references to the heap because binding a variable is postponed until a clause is chosen.

3. There is less trailing because it is only needed for bindings that cross a choice point.

4. Garbage collection is more efficient, since the creation of fewer choice points means that there are fewer starting points for marking.

3.2.2. Ramifications of exploiting determinism

The goal of compiling deterministic predicates into efficient conditional branches affects a large part of the compiler. Many of the transformations done in the compiler are intended to increase the amount of determinism that is easily accessible. This includes formula manipulation, factoring, head unraveling, the determinism transformation (all in Chapter 4), the determinism compiler (Chapter 5), and the determinism optimization (Chapter 6).
Through these transformations the compiler creates a decision graph to index the arguments of a predicate. Type information derived by dataflow analysis is exploited to simplify the graph. The graph is created in an architecture-independent way through the concept of the test set (Chapter 4). Intuitively, a test set is a set of Prolog predicates that are mutually disjoint (only one can succeed at any given time) and that correspond to a multi-way branch in the architecture.

3.3. Specialize unification

The WAM unification instructions (get and unify) are complex. They operate in two modes (read mode and write mode) depending on the type of the object being unified, they dereference their arguments, and they trail variable bindings. It is better to compile unification directly into simpler instructions.

In the Aquarius compiler, unification is compiled into the simplest possible BAM code taking the type information into account (Chapter 5). Often it is possible to reduce a unification to a single load or store. The use of uninitialized variables (see below) to simplify variable binding greatly improves the generated code.

![Diagram of three categories of unbound variables](image)

Figure 2.9 – Three categories of unbound variables
3.3.1. Simplifying variable binding

A major source of inefficiency in WAM implementations is that logical variables are often created as
unbound (i.e. as self-referential pointers) and then unified soon afterwards. Creating and unifying does
much unnecessary work; it would be faster just to reserve a memory location and then write to it. The
Aquarius compiler defines such a representation, called uninitialized variables. Conceptually, uninitialized
variables are defined at two levels:

(1) At the logical level, an uninitialized variable is an unbound variable that is not aliased, i.e. there are
no other variables bound to it. The dataflow analyzer (Chapter 4) uses this definition to derive unin-
itialized variable types.

(2) At the implementation level, an uninitialized variable is a location that is allocated to contain an
unbound variable, but the location is not given a value. The kernel Prolog compiler (Chapters 4, 5,
and 6) uses this definition to compile uninitialized variables efficiently.

The location containing an uninitialized variable can either be a register or a memory word, resulting in
two kinds of uninitialized variables, namely uninitialized register and uninitialized memory variables. The
first are registers whose contents are ignored. The second are pointers to memory locations whose contents
are ignored. Standard unbound variables are called initialized variables; they are pointers to locations
pointing to themselves. Figure 2.9 illustrates the three categories of unbound variables.

<table>
<thead>
<tr>
<th>Type of variable</th>
<th>Cost (VLSI-BAM cycles)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>For Unification</td>
</tr>
<tr>
<td></td>
<td>Creation</td>
</tr>
<tr>
<td>Uninitialized Register</td>
<td>0</td>
</tr>
<tr>
<td>Uninitialized Memory</td>
<td>1</td>
</tr>
<tr>
<td>Initialized Variable</td>
<td>2</td>
</tr>
</tbody>
</table>

The dataflow analyzer derives both uninitialized register and uninitialized memory types. It is often
able to determine that an argument is uninitialized; for a representative set of programs it finds that 23% of
all predicate arguments are uninitialized. Of these, two thirds have uninitialized memory type and one
third have uninitialized register type.
Table 2.3 gives the minimum run-time costs on the VLSI-BAM processor for the three categories of unbound variables. Costs are given for unification support (creation and binding) and for backtracking support (trailing and detrailing). Binding an initialized variable is expensive because the variable must be dereferenced before the new value can be stored in the memory cell. Binding an uninitialized memory variable reduces to a single memory store operation. Binding an uninitialized register variable is free if it is created in the register that needs it. The cost of detrailing (restoring a variable to an unbound state on backtracking) is zero for uninitialized variables. For initialized variables it depends strongly on the effectiveness of the compiler in generating deterministic code. It is 0 cycles if the variable does not have to be unbound on backtracking, and 4 cycles otherwise.

3.4. Dataflow analysis

The Aquarius compiler implements a dataflow analyzer that is based on abstract interpretation. It translates the program to one in which predicate arguments range over a finite set of values. Each of the values corresponds to an infinite set of values (i.e. a type) in the original program. The analyzer derives a small set of types—uninitialized, ground (the argument contains no unbound variables), nonvariable (the argument is not an unbound variable) and recursively dereferenced (the argument is dereferenced, i.e. it is accessible without pointer chasing, and if it is compound, then all its arguments are recursively dereferenced). These types have been chosen carefully to be useful during compilation.

Dataflow analysis by itself is not enough. The rest of the system must be able to use the information derived by the analysis. The techniques to exploit determinism and specialize unification in the Aquarius compiler have been developed in tandem with the analyzer for this purpose. In addition, the fine instruction granularity of the BAM is designed to support these optimizations.

4. Related work

First a survey is given of work that is related to the four principles of the Aquarius compiler. Then an overview is given of Prolog implementations that are interesting in some way.
4.1. Reduce instruction granularity

Tamura et al [39, 65] have done fundamental work at IBM Japan in reducing the grain size of compiled operations for Prolog. Their compilation is done in three steps. The first step is to compile Prolog into WAM. In the second step the intermediate code is translated into a directed graph. Each WAM instruction becomes a subgraph containing simple operations such as case selection on tags, jumps, assignments, and dereferencing. The graph is optimized through rewrite rules. Case selections based on a tag value, never-selected cases, redundant tests, case statements with only one branch, and unreachable instructions are eliminated. Known values are propagated. These rewrites are applied several times and the resulting graph is then translated back into intermediate code. In the third step the intermediate code is translated into a PL.8 program which is sent to a high-quality PL.8 optimizing compiler [3]. Performance results are given for a few small programs and are quite good. There are several problems in their approach. They still use the WAM as an intermediate language, and compiling is prohibitively slow because their system is experimental. Without compile-time hints their performance drops significantly.

4.2. Exploit determinism

Significant improvements over the WAM are possible to avoid choice point creation in deterministic predicates. The WAM indexes on only the first argument and saves all registers in choice points. Turk [72] describes several optimizations that reduce the time necessary to restore machine state when backtracking. In [74], I describe a compilation scheme that attempts to take advantage of the fact that most Prolog predicates are deterministic. Choice point creation and moves to and from choice points are minimized. Clauses are compiled with multiple entry points and predicates are compiled as decision trees. The techniques used in the Aquarius system are inspired by this work. Carlsson [15] measures the performance improvement of a scheme for creating choice points in two parts, saving only a small part of the machine state first, and postponing saving the remainder until later in the clause when it can be determined that the head unification and any simple tests have succeeded. Implemented in the SICStus Prolog system, this reduces execution time by 7-15% on four large programs.

Recently there have appeared several commercial Prolog-like languages (Trilogy and Turbo Prolog)
that generate efficient code for programs annotated with type and determinism declarations. In this regard Trilogy [79] is noteworthy because it gives a logical semantics to programs written in a Pascal-like notation. Typed predicates that are annotated as being deterministic are compiled into efficient native code. The achievement of Trilogy is reassuring; since many predicates in standard Prolog are intended to be executed in a deterministic way, with some analysis it should be possible to obtain the same efficiency for standard Prolog.

Several systems have generalized the first argument indexing of the WAM. BIM_Prolog [4] can index on any argument when given appropriate declarations. SEPIA [29] incorporates heuristics to decide which predicate arguments are important for deterministic selection. It uses the first “indexable” argument of a predicate. If there are several possibilities it first uses the argument where it is more likely that fewer clauses will be selected.

Several papers describe fast implementations of the cut operation. Bowen et al [9] implement cut by adding a register that holds the address of the most recent choice point before entering the predicate. This register is updated by each call and execute instruction. Cut is implemented by moving this register to the WAM’s choice point register r(b). Mariën and Demoen [46] implement cut in a similar fashion. These schemes suffer from having to do an additional register move for each procedure call, unless a different call instruction is used for predicates with and without cut. The scheme implemented in the Aquarius compiler does not slow down procedure calls and does not need an additional register.

4.3. Specialize unification

Significant improvements over the WAM are possible for unification. Turk [72] describes several optimizations related to compilation of unification, to reduce the overhead of explicitly maintaining a read/write mode bit and remove some superfluous dereferencing and tag checking. Mariën [44] describes a method to compile write mode unification that uses a minimal number of memory operations and avoids all superfluous dereferencing and tag checking. In [75], I build on this work by introducing a simplified notation and extending it for read mode unification, but my scheme suffers from a large code size expansion. The Aquarius system modifies this technique to limit the code size expansion at a slight execution time.
cost. Meier [48] has developed a technique that generalizes Mariën’s idea for both read and write mode and achieves a linear code size, also with a slight execution time cost. This technique is implemented in the SEPIA system [29].

Beer [5] has suggested the use of a simplified representation of Prolog variables for which binding is much faster. He introduces several new tags for this representation, which he calls uninitialized variables, and keeps track of them at run-time. He shows that both dereferencing and trailing are reduced significantly. This idea was a strong influence on the Aquarius compiler. At the Prolog level, logical semantics are preserved, but at the code level there is now a coherent integrated use of destructive assignment for values that fit in a register. My scheme is different from Beer’s—it uses the same tag for both uninitialized and standard Prolog variables. The analyzer finds uninitialized variables at compile-time and the compiler determines when it is safe to use destructive assignment to bind them.

4.4. Dataflow analysis

R. Warren et al [84] have done the most comprehensive work measuring the practicality of global dataflow analysis in logic programming. Their paper describes two dataflow analyzers: (1) MA3, the MCC And-parallel Analyzer and Annotator, and (2) Ms, an experimental analysis scheme developed for SB-Prolog. MA3 derives aliasing and ground types and keeps track of the structure of compound terms, while Ms derives ground and nonvariable types. The paper concludes that both dataflow analyzers are effective in deriving types and do not increase compilation time by too much. My dataflow analyzer differs from both MA3 and Ms in three ways. First, the analyzer works over a different domain. Second, it avoids problems with aliased variables by deriving only limited type information for them. Third, it is integrated into a compiler which has been developed to take full advantage of the types it derives.

For correctness, it is imperative to consider the effects of variable aliasing on dataflow analysis. Aliasing occurs when two variables are bound to terms that have variables in common. Finding accurate aliasing information is an important topic in current research [18, 36]. However, aliasing complicates the implementation of dataflow analysis. My analyzer considers only unaliased variables as candidates for unbound variable types. Measurements of the analyzer show that unaliased variables occur often enough
to make the analysis worthwhile. This conservative treatment of aliasing simplifies the implementation,
since it is not necessary to explicitly represent and propagate aliasing information. Of course, it also
reduces the effectiveness of the analysis. Thus aliasing needs to be studied further.

Mariën et al [45] have performed an interesting experiment in which several small Prolog predicates
(recursive list operations) were hand-compiled with several levels of optimization based on information
derivable from a dataflow analysis. The analysis was done by hand at four levels: The first level derives
unbound variable and ground modes. The second level also derives recursively defined types. The third
level also derives lengths of dereference chains (pointer chains that must be followed at run-time). The
fourth level also derives liveness information for compound data structures and is used to determine when
they are last used so that their memory may be recovered (compile-time garbage collection). Execution
time measurements show that each analysis level improves speed over the previous level. This experiment
shows that a simple analysis can achieve good results on small programs.

4.5. Other implementations

This section gives an overview of interesting Prolog implementations that are related to this disserta-
tion in some way. Most existing implementations of Prolog, both on general-purpose and special-purpose
machines, are based on the Warren Abstract Machine (WAM) or are derived from it. The general-purpose
and special-purpose approaches are presented separately. The first subsection describes some important
software implementations and their ideas. The second subsection summarizes some important architec-
tures and their innovations.

4.5.1. Implementing Prolog on general-purpose machines

As far as I know, the earliest WAM compiler was my PLM compiler, completed and published in
August 1984 [73].† The compiler was interesting as it was itself written in Prolog, unlike many later Prolog
compilers. The first commercial implementation of the WAM was Quintus Prolog, announced in
November 1984.

† The PLM compiler is still available from us, but is now obsolete and not recommended for current research work. Our
research group expects to release soon a complete Prolog system based on the Aquarius compiler.
Among the highest performance commercial implementations available today are IBM Prolog, Quintus Prolog [58], BIM_Prolog [4], and ALS Prolog [2]. There are three significant implementations of Prolog available today that were developed at research institutions: SICStus Prolog [63], SEPIA [29], and SB-Prolog [83]. All of these systems are based on extensions of the WAM (except possibly IBM Prolog, of which I have little information) and compile to WAM-like instructions which are either emulated on the target machine or macro-expanded to native code. Some of these systems (e.g. SB-Prolog and IBM Prolog) are able to compile special cases of deterministic programs into efficient code.

4.5.1.1. Taylor's system

Independently of this research, Andrew Taylor is implementing a high performance Prolog compiler for the MIPS processor [67]. The compiler includes a dataflow analyzer that explicitly represents type, aliasing, dereference chain lengths, and trailing information [66]. His preliminary results indicate that it is of comparable performance to the compiler presented in this dissertation. Running a set of small benchmark programs on the MIPS R2030 processor, the system is 24 times faster than compiled SICStus Prolog version 0.6 and the code size is similar to that of the KCM.

4.5.1.2. IBM Prolog

IBM Prolog accepts mode declarations, implements more general indexing than the WAM, does a limited global analysis (however, it does not derive any types), and generates high performance native code. It is able to compile some kinds of deterministic programs with conditional branches.

4.5.1.3. SICStus Prolog

SICStus Prolog was developed at the Swedish Institute of Computer Science in Stockholm. A backend module was written for it by Mats Carlsson which generates native code avoiding the superfluous memory references of a naive WAM translation [14,44]. It is comparable in performance to Quintus Prolog when no built-in predicates are used.
4.5.1.4. SB-Prolog

SB-Prolog was developed at SUNY in Stony Brook. It recognizes a special case of the general techniques for extracting determinism discussed in this dissertation: it recognizes when arithmetic tests that are each other’s opposites appear, and compiles a conditional branch. It also incorporates a simple partial evaluator which is used for macro expansion and a simple dataflow analysis scheme has recently been developed for it [84].

4.5.2. Implementing Prolog on special-purpose machines

In the past, because the WAM was regarded as the best way to implement Prolog, the performance gap between special-purpose architectures and general-purpose architectures was large. Much of the effort in high performance Prolog implementation was put into architecture design, and in particular in hardware support for the WAM instructions. This dissertation shows that a better understanding of Prolog execution narrows the performance gap. The implications of this development for the future of special-purpose architectures are discussed in the VLSI-BAM paper [34] and summarized in this section.

4.5.2.1. PLM

The first special-purpose Prolog architecture that was built is the PLM (Programmed Logic Machine), due to Dobry et al [26-28]. Its design was inspired by a proposal of Tick & Warren [68]. The PLM implements the WAM in microcode with a 100 ns clock cycle. It was built on wire-wrap boards and ran a few small programs in 1985. Spin-offs of this project included the VLSI-PLM single-chip implementation [60] and the Xenologic X-1, a commercial coprocessor for Sun workstations.

Several papers have compared the number of cycles needed by the PLM to that of general-purpose architectures. These ratios are valid measurements of the effect of the PLM’s architectural support for WAM implementation. Mulder & Tick [51] and Patt & Chen [54] have compared the performance of the PLM [28], a microcoded implementation of the WAM, to a macro-expanded WAM on the MC68020 processor. They find that the MC68020 needs 3 to 4 times the number of cycles as the PLM to execute the WAM. Patt and Chen find that static code size on the MC68020 is about 20 times the PLM.
4.5.2.2. SPUR

Borriello et al [8] have implemented a macro-expanded WAM on the SPUR processor (Symbolic Processing Using RISCs). They find that the SPUR takes about 2.0 times the number of cycles as the PLM and that static code size is about 14 times the PLM. These numbers include local optimizations implemented by Chen and Nguyen [20] that improve the original numbers by about 10%.

4.5.2.3. PSI-II and PIM/p

In the context of the FGCS (Fifth Generation Computer System) project, researchers of ICOT (the Japanese Institute for New Generation Computer Technology) have designed and built several sequential and parallel architectures for logic programming [64, 71]. One of the more interesting sequential machines is the PSI-II (Personal Sequential Inference machine II) [52] a microcoded implementation of the WAM which executes at speeds similar to the PLM. The processing elements of the PIM/p (Parallel Inference Machine) architecture are currently the highest performance sequential logic machines at ICOT. They execute at two to three times the speed of the PLM.

4.5.2.4. KCM

Benker et al [6] describe a special-purpose Prolog machine, the KCM (Knowledge Crunching Machine), which is based on an extended WAM. Its instruction set consists of two parts: a general-purpose instruction set, and a microcoded Prolog-specific instruction set. It has a cycle time of 80 ns and executes in about 1/3 the number of cycles of the PLM. Its code size is about three times greater. The KCM project was done together with the development of a Prolog system and environment called SEPIA (see previous section). About 60 KCM machines were constructed and delivered to the ECRC member companies.

4.5.2.5. VLSI-BAM

Holmer et al [34] describe a single-chip microprocessor with extensions for Prolog, the VLSI-BAM (VLSI Berkeley Abstract Machine). It is a pipelined load-store processor with a cycle time of 33 ns. It takes about 1/3 the number of cycles to run programs as the PLM and its code size is about three times...
greater, results similar to the KCM. However, they are achieved largely through the effort of the compiler.

The goal of the BAM project is to find the minimal extensions to a general-purpose architecture to support a high performance Prolog implementation. The rationale for the VLSI-BAM architecture is that existing general-purpose architectures are designed to execute imperative languages like C and do not have adequate support for Prolog. The compiler described in this dissertation was developed simultaneously with the architecture, and interaction between the two designs has significantly improved both.

The BAM project has determined that a small amount of architectural support (5% increase in chip area) gives a large performance boost (50% performance increase) for programs that use Prolog-specific features. The support does not interfere with the general-purpose architecture, so it is possible for future general-purpose machines to incorporate this support for high performance symbolic computing. The support is designed specifically to support the logical variable, dynamic typing, unification, and backtracking. A language that uses any of these features can benefit from it.
Chapter 3
The Two Representation Languages

1. Introduction

This chapter defines the two languages used by the compiler to represent programs: kernel Prolog, a simplified form of Prolog, and the Berkeley Abstract Machine (BAM), a low-level instruction set and execution model that is close to a standard sequential processor. Kernel Prolog is an internal language that is not accessible to the user. BAM is the output language of the compiler.

2. Kernel Prolog

The first representation language in the compiler is kernel Prolog, a simplified, canonical form of Prolog. The syntax of kernel Prolog is given in Figure 3.1. This should be compared with the definition of full Prolog syntax given in Chapter 2. The control flow of kernel Prolog is simpler, a set of internal primitives is defined that are only used inside the compiler, and a case statement is defined. Kernel Prolog does not have nested disjunctions, if-then-else, cut, negation, or arithmetic expressions. Each predicate is represented as a single term \((H : -D)\) containing a head \(H\) with distinct variable arguments and a body \(D\) that is a single disjunction (an OR choice). Each alternative of the disjunction is a conjunction, i.e. an AND sequence of goals. Unifications in the head of the original predicate are represented as explicit unifications in the arms of the disjunction. Disjunctions, negations, and if-then-else forms in the original predicate are converted into dummy predicates. Cut and arithmetic expressions are converted into simpler internal built-in predicates.

For example, the predicate:

\[
\begin{align*}
a(b) & \\
a(X) & : - (0 \ mod \ 2 \ \rightarrow e(X) \ ; \ f(X))
\end{align*}
\]

is represented as follows in kernel Prolog:

\[
\begin{align*}
a(b) & \\
a(X) & : - (0 \ mod \ 2 \ \rightarrow e(X) \ ; \ f(X))
\end{align*}
\]
predicate((H:-D)) :- head(H), disjunction(D).

head(H) :- goal_term(H).

disjunction(fail).

disjunction((C;D)) :- conjunction(C), disjunction(D).

conjunction(true).

conjunction((G,C)) :- goal(G), conjunction(C).

goal(G) :- case_goal(G).

goal(G) :- internal_goal(G).

goal(G) :- external_goal(G).

case_goal('Scase'(Name,Ident,CB)) :- test_set(Name, Ident), case_body(CB).

case_body('Selse'(D)) :- disjunction(D).

case_body('Selse'(T,D):CB)) :- test(T), disjunction(D), case_body(CB).

external_goal(G) :- goal_term(G), \+case_goal(G), \+internal_goal(G).

term(T) :- var(T).

term(T) :- goal_term(T).

goal_term(T) :- nonvar(T), functor(T, _, A), term_args(I, A, T).

term_args(I, A, _) :- I>A.

term_args(I, A, T) :- I=<A, arg(I, T, X), term(X), I is I+1, term_args(I, A, T).

\ % Predicates defined in 'tables:

internal_goal(G) :- (Defined in Table 3.1).

test_set(Name, Ident) :- (Defined in Table 4.1).

test(T) :- (Defined in Table 4.1).

\ % Built-in predicates needed in the definition:

functor(T, F, A) :- (Term T has functor F and arity A).

arg(I, T, X) :- (Argument I of compound term T is X).

var(T) :- (Argument T is an unbound variable).

nonvar(T) :- (Argument T is a nonvariable).

Figure 3.1 – Syntax of kernel Prolog
All predicates that start with the character ' $' are created internally. Cut is implemented with the two built-ins '$cut_load'(X) and ' $cut'(X). The arithmetic expression 0 is X mod 2 is replaced by a call to an explicit arithmetic built-in ' $mod'(X, 2, 0). The if-then-else is replaced by a call to the dummy predicate ' $d'(X). All dummy predicates are given unique names.

Kernel Prolog has many advantages over standard Prolog. The scope of variables is not limited to a single clause, but is extended over the whole predicate. Many optimizations are easier to do—for example, dataflow analysis and determinism extraction. Compilation to BAM code and register allocation are simplified.

The following two sections describe the internal predicates of kernel Prolog and how standard Prolog is converted to kernel Prolog.

2.1. Internal predicates of kernel Prolog

The kernel Prolog form of a program contains predicates that are not part of standard Prolog and that are invisible to the user. The internal predicates always begin with the character ' $'. They are of three kinds:

(1) Internal built-in predicates (Table 3.1). These are classified into three categories depending on their use: (1) implementation of cut, (2) type checking, and (3) arithmetic. They are expanded into BAM instructions before being output, so the user never sees them.

(2) A case statement. This control structure is designed to express deterministic selection in Prolog.

Chapter 4 describes how the case statement is created. It is translated directly into conditional
Table 3.1 - Internal built-ins of kernel Prolog

<table>
<thead>
<tr>
<th>Built-in</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>'Scut_load' (X)</td>
<td>Load the choice point register r (b) into X.</td>
</tr>
<tr>
<td>'Scut' (X)</td>
<td>Make the choice point pointed to by X the new top of the choice point stack.</td>
</tr>
<tr>
<td>'Sname arity' (X, Na, Ar)</td>
<td>Test that X has functor Na and arity Ar. This only does a check; it never binds X.</td>
</tr>
<tr>
<td>'Stest' (X, T)</td>
<td>General type-checking predicate that tests whether the type of X is in the set T, where T ⊆ {unbound variable, nil, non-nil atom, negative integer, nonnegative integer, float, cons, structure}.</td>
</tr>
<tr>
<td>'Sequal' (X, Y)</td>
<td>Test that X and Y are identical simple terms.</td>
</tr>
<tr>
<td>'Sadd' (S1, S2, D)</td>
<td>Integer addition D ← S1+S2.</td>
</tr>
<tr>
<td>'Ssub' (S1, S2, D)</td>
<td>Integer subtraction D ← S1–S2.</td>
</tr>
<tr>
<td>'Smul' (S1, S2, D)</td>
<td>Integer multiplication D ← S1*S2.</td>
</tr>
<tr>
<td>'Sdiv' (S1, S2, D)</td>
<td>Integer division D ← S1/S2.</td>
</tr>
<tr>
<td>'Smod' (S1, S2, D)</td>
<td>Integer remainder D ← S1 mod S2.</td>
</tr>
<tr>
<td>'Sand' (S1, S2, D)</td>
<td>Bitwise integer &quot;and&quot; D ← S1 &amp; S2.</td>
</tr>
<tr>
<td>'Sor' (S1, S2, D)</td>
<td>Bitwise integer &quot;or&quot; D ← S1 v S2.</td>
</tr>
<tr>
<td>'Sxor' (S1, S2, D)</td>
<td>Bitwise integer exclusive-or D ← S1 ⊕ S2.</td>
</tr>
<tr>
<td>'Ssll' (S1, S2, D)</td>
<td>Logical left shift D ← S1&lt;&lt;S2.</td>
</tr>
<tr>
<td>'Ssra' (S1, S2, D)</td>
<td>Arithmetic right shift D ← S1&gt;&gt;S2.</td>
</tr>
<tr>
<td>'Snot' (S, D)</td>
<td>Bitwise integer negation D ← not S.</td>
</tr>
</tbody>
</table>

branches in the BAM code and has the following syntax:

```
'Scase' (Name, Ident, CaseBody)
```

where:

```
CaseBody = ('Stest' (Test, Code)
             ; ... ;
             ; 'Selse' (Code)
         ) .
```

CaseBody is a disjunction of 'Stest' goals, terminated with an 'Selse' goal. Code is any valid kernel Prolog disjunction. Name and Ident identify the test set, and Test is a Prolog predicate (Table 4.11). Test is the test that is valid along the branch. For example, for the hashing function it will be the goal X=a where a is the atom or structure used in that direction.

(3) "Dummy" predicates. Kernel Prolog does not allow control structures (i.e. disjunctions, if-then-else, and negation) in clauses, but only calls. The control structures are transformed into calls to dummy predicates, which are predicates that exist only inside the original predicate. Dummy predicates are created with unique names that are derived from the predicate they are contained in.
2.2. Converting standard Prolog to kernel Prolog

The first stage of compilation is a sequence of five source transformations that converts raw input clauses into kernel Prolog. An input predicate in standard Prolog is transformed into a tree that contains a kernel Prolog form of the original predicate and a set of dummy predicates in kernel form created during the transformation. Care is taken to put the predicate in a form that maximizes opportunities for determinism extraction. The five transformations are:

(1) **Standard form transformation.** Convert the raw Prolog input to a convenient standard notation. This does several housekeeping tasks: it properly terminates conjunctions (with `true`) and disjunctions (with `false`), and it converts negation-as-failure into if-then-else.

(2) **Head unraveling.** Rewrite the head of each clause as a new head and a list of unification goals such that all the arguments of the new head are distinct variables and the head unifications are unification goals.

(3) **Arithmetic transformation.** Compile arithmetic expressions to internal arithmetic built-ins.

(4) **Cut transformation.** Implement cut by converting all uses of cut and if-then-else to internal cut built-ins.

(5) **Flattening.** At this point all complex control has been converted to disjunctions. Convert nested disjunctions to dummy predicates.

### 2.2.1. Standard form transformation

The standard form of a clause is intended to simplify its syntax so that traversing it is as simple as possible. The standard form satisfies the rules in Table 3.2. These rules are ignored in the presentation of most of the examples in this dissertation because they make the examples less readable (although they are always satisfied in the compiler).

### 2.2.2. Head unraveling

Unraveling the head of a clause consists of rewriting it as a new head and putting a series of unification goals in the clause's body so that all the head's arguments are distinct variables and all the head
Table 3.2 – Standard form of a clause

<table>
<thead>
<tr>
<th>Rule</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Conjunctions and disjunctions are right associative.</td>
</tr>
<tr>
<td>2</td>
<td>Conjunctions have no internal true and are terminated by true.</td>
</tr>
<tr>
<td>3</td>
<td>Disjunctions have no internal fail and are terminated by fail.</td>
</tr>
<tr>
<td>4</td>
<td>Single goals inside disjunctions are considered as conjunctions (and therefore rule 2 applies).</td>
</tr>
<tr>
<td>5</td>
<td>There is no negation (it is converted to if-then-else).</td>
</tr>
<tr>
<td>6</td>
<td>Arguments of if-then-else are considered as conjunctions (and therefore rule 2 applies).</td>
</tr>
<tr>
<td>7</td>
<td>((A \rightarrow B)) as a goal in a conjunction is converted to ((A \rightarrow B; fail)).</td>
</tr>
<tr>
<td>8</td>
<td>The first argument of all unify goals is a variable.</td>
</tr>
</tbody>
</table>

The first argument of all unify goals is a variable.

If this is not done correctly then much opportunity for later optimization is lost. From the predicate's type formula, the compiler knows which head arguments are nonvariable and which head arguments are unbound. Unification goals are created that satisfy two constraints:

1. Maximize the number of nonvariable arguments that are unified together. Put these unifications first in the unraveled clause.

2. Minimize the number of unification goals that contain unbound variables. Put these unifications last in the unraveled clause.

For example, consider the clause:

\[ :-mode((a(A,B,C):-nonvar(A),nonvar(B),var(C))). \]

\[ a(A,A,A) :- atomic(A), \ldots \]

The type declaration says that the first two arguments are nonvariables and the third argument is an unbound variable. The argument \(A\) appears three times in the head. Therefore there are three ways to unravel this clause: \((a(X,Y,Z):-X=Y,X=Z),\) \((a(X,Y,Z):-Y=X,Y=Z),\) and \((a(X,Y,Z):-Z=X,Z=Y).\) Considering the mode declaration, the head is transformed into the first of the three unraveled versions:

\[ a(A,B,C) :- A=B, A=C, atomic(A), \ldots \]

The first unification \(A=B\) is of two nonvariables. The second unification \(A=C\) is of a nonvariable and an unbound variable. This satisfies both constraints.
expression((X is Expr), Code) :- expr(Expr, X, Code, []).
expr(V, V) --> {var(V)}, !.
expr(A, A) --> {integer(A)}, !.
expr(A+B, C) --> expr(A, Ta), expr(B, Tb), ['$add' (Ta, Tb, C)].
expr(A-B, C) --> expr(A, Ta), expr(B, Tb), ['$sub' (Ta, Tb, C)].
expr(A*B, C) --> expr(A, Ta), expr(B, Tb), ['$mul' (Ta, Tb, C)].
expr(A/B, C) --> expr(A, Ta), expr(B, Tb), ['$div' (Ta, Tb, C)].

Figure 3.2 – Compiling an arithmetic expression

2.2.3. Arithmetic transformation

The is42 predicate is translated into internal three-argument arithmetic built-ins (Table 3.1). Figure 3.2 gives a simplified but fully functional version of the algorithm used to compile expressions. It handles arbitrary expressions containing the four basic arithmetic operations. For example, the call:

expression(X is 23*(Y+2), Code)

gives the code:

Code = ['$add' (Y, Z, T), '$mul' (23, T, X)]

The full algorithm handles all the arithmetic primitives of Table 3.1 and does partial constant folding.

2.2.4. Cut transformation

The cut operation modifies control flow by removing all choice points created since entering the predicate containing the cut, including the choice point of the predicate itself. Cut is implemented by means of a source transformation. It requires no support from the architecture except the ability to access and modify the register r(b), which points to the most recent choice point.

The cut transformation is given in Figure 3.3. A call to the built-in "$cut_load"(X) is put at the entry of a predicate containing a cut. This built-in moves the r(b) register to X, which marks the top of the choice point stack on entry to the predicate. The argument X is passed to the predicate’s body. Each occurrence of cut in the body is replaced by a call to the built-in "$cut"(X). This built-in loads r(b)


procedure cut_transformation;  
var  $P'$: list of clause;  
begin  
  for each predicate $P$ in the program do begin  
    if $P$ contains a cut then begin  
      /* At this point $P = \{ C_1, \ldots, C_n \}$ (list of clauses) and $C_i = (H_i, :- B_i)$ */  
      Add the argument $X$ to all $H_i$ in $P$;  
      Replace each occurrence of '!' in $P$ by 'Scut' ($X$);  
      $P' := P$;  
      Add the predicate $P'$ to the program;  
      $H := \text{new head with same functor and arity as all } H_i$;  
      $H' := (H \text{ with the additional argument } X)$;  
      $P := [(H := 'Scut\_load' (X), H')]$  
    end  
  end  
end;  

Figure 3.3 – The cut transformation

from $X$, which restores the original top of the choice point stack. For example, consider the predicate:

$$p := q, !, r.$$  
$$p := s.$$  

This is transformed into:

$$p := 'Scut\_load'(X), p' (X).$$  
$$p' (X) := q, 'Scut' (X), r.$$  
$$p' (X) := s.$$  

Compilation then continues in the usual manner. This method is simple and efficient. Variations of it have been implemented in other Prolog systems [4, 13, 45]. This method differs from these variations in that the compiler does not always store the value of $r(b)$ on the environment stack, but puts it in a predicate argument $X$. It is stored in an environment only if the clause is compiled with an environment.

2.2.5. Flattening

At this point, all the complex control in a predicate (disjunctions, if-then-else, and negation-as-failure) has been translated to disjunctions. Flattening replaces the disjunctions by calls to dummy predicates. For example, the definition:
\[ a(X, Y) := (b1(X, A) ; b2(X, B), t(B) ), d(Y, A). \]

is transformed into:

\[ a(X, Y) := 'Sflatten_a/2_1'(X, A), d(Y, A). \]

\[ 'Sflatten_a/2_1'(X, A) := b1(X, A). \]

\[ 'Sflatten_a/2_1'(X, A) := b2(X, B), t(B). \]

Compilation then continues in the usual manner and the dummy predicate \( 'Sflatten_a/2_1'(X, A) \)
is compiled as in-line code. The dummy predicate is created with a unique name derived from the name of
the original predicate. The argument list of the dummy predicate is the intersection of the set of variables
used inside the disjunction and the set of variables used outside it. In this example the argument list is the
intersection of \( \{ X, Y, A \} \) and \( \{ X, A, B \} \), which is \( \{ X, A \} \).
3. The Berkeley Abstract Machine (BAM)

The foundation of the efficiency of the compiler is its execution model, the BAM. The BAM has been designed to support all compiler optimizations and to make the system easily retargetable to the VLSI-BAM and general-purpose machines. The design evolved by interaction with the development of the compiler, the architecture design of the VLSI-BAM processor, and the requirement of portability to other architectures. The BAM was developed in tandem with the VLSI-BAM processor, but the two instruction sets are quite different. The VLSI-BAM is constrained by its hardware implementation; the BAM evolved by looking at the requirements of Prolog and is designed to allow a great deal of low-level optimization.

The Aquarius compiler uses a simple output language and not an existing high-level language such as C or an existing low-level language such as an assembly for a particular machine. There are several reasons for this:

(1) Choosing an existing language requires choosing representations for tags and data structures, and writing frequently used Prolog-specific operations as subroutines. This is undesirable for two reasons: First, the VLSI-BAM is one of the target machines and its architecture has a more abstract representation for tags and Prolog-specific operations than general-purpose processors. Second, these representations are not necessarily the best for all machines.

(2) Choosing an existing high-level language is unsatisfactory for the VLSI-BAM processor since the only compiler for it is currently the Aquarius compiler.

(3) An unpredictable factor is introduced when doing performance evaluations. The performance on different machines varies depending on the sophistication of the implementation of the existing language. It is not always easy to determine the performance of the existing language from inspection of its source code.

The syntax and semantics of the BAM is presented at several levels of detail, from a discussion of its features in English down to a detailed formal specification of its semantics in Prolog. The body of the dissertation defines the data types of the BAM, gives an overview of its instruction set, and justifies the choice of instructions. Appendices B and C give formal specifications of BAM syntax and semantics, and
Appendix D gives a concise but complete English description of BAM semantics.

This section has four parts. The first part presents the data types of the BAM. The second part summarizes the BAM instruction set. The instruction set consists of four parts: simple instructions (tagged load-store architecture), complex instructions (Prolog-specific operations), pragmas (embedded information to allow better translation to a real machine), and user instructions (intended to allow the complete run-time system to be written in BAM). The third part justifies the complex instructions. The fourth part justifies the instructions needed to implement unification by showing how they are constructed from a unification algorithm given a few simple assumptions about the architecture.

3.1. Data types in the BAM

The data types of the BAM are classified into two groups: the types used during execution and the types used to represent instructions (Table 3.3). The BAM has four data types that are used during execution: words, natural numbers, symbolic labels, and mappings. These are denoted as the set of all words $W$, the set of natural numbers $N$, the set of mappings $M$, and the set of symbolic labels $L$. A word is a pair $T \cdot N$ where $T$ is the tag and $N$ is the value. A natural number is a nonnegative integer. A mapping (not shown in Table 3.3) is a correspondence between a set of objects and their values (which are often words). A symbolic label marks a position in the program.

Several definitions in Table 3.3 require some clarification. Sets are denoted by bold capital letters, variables by capital letters, and constants by lower case letters. Addressing modes are defined recursively, with a base case consisting of registers and atomic terms, and a recursive case consisting of three parts: tag insertion ($T \cdot X$), indirection ($X$), and offset ($X+N$). The BAM uses only a subset of the infinite set of addressing modes defined here. Of all the internal registers of the BAM, only the argument registers $r(I)$, the heap pointer $r(h)$, and the backtrack pointer $r(b)$ are visible in the instruction set. Appendix B gives a precise definition of instruction syntax including the addressing modes that are actually used. The meaning of the instructions is defined informally in section 3.2 and formally in Appendix C.

A term can be of arbitrary size. A term that fits completely in a register is called simple. All other terms are called compound. A register cannot store all possible terms, but it can contain encoded informa-
Table 3.3 - Types in the BAM

<table>
<thead>
<tr>
<th>Name</th>
<th>Definition</th>
</tr>
</thead>
<tbody>
<tr>
<td>Word</td>
<td>[ W = { T \cdot N</td>
</tr>
<tr>
<td>Symbolic label</td>
<td>[ L = { \text{fail} } \cup { F/N, l(F/N), I</td>
</tr>
<tr>
<td>Natural number</td>
<td>[ N = { \text{atom}(V) \lor \text{natural}(V) } \cup { V \mid \text{integer}(V) } \cup { \text{float}(V) } ]</td>
</tr>
<tr>
<td>Atomic term</td>
<td>[ A = { \text{atom}(V) \lor (V=(F/N) \land \text{atom}(F) \land \text{natural}(N)) } \cup { V \mid \text{integer}(V) } \cup { \text{float}(V) } ]</td>
</tr>
</tbody>
</table>

Types used to represent instructions

<table>
<thead>
<tr>
<th>Name</th>
<th>Definition</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tag</td>
<td>[ T = { \text{tvar}, \text{tlst}, \text{tstr}, \text{tatom}, \text{tint}, \text{tneg}, \text{tflt} } = T_p \cup T_a ]</td>
</tr>
<tr>
<td>Pointer tag</td>
<td>[ T_p = { \text{tvar}, \text{tlst}, \text{tstr} } ]</td>
</tr>
<tr>
<td>Atomic tag</td>
<td>[ T_a = { \text{tatom}, \text{tint}, \text{tneg}, \text{tflt} } ]</td>
</tr>
<tr>
<td>Condition</td>
<td>[ C = { \text{eq}, \text{ne}, \text{lt}, \text{les}, \text{gt}, \text{ges} } ]</td>
</tr>
<tr>
<td>Equality condition</td>
<td>[ C_e = { \text{eq}, \text{ne} } ]</td>
</tr>
<tr>
<td>Arithmetic operation</td>
<td>[ E = { \text{add}, \text{sub}, \text{mul}, \text{div}, \text{mod}, \text{and}, \text{xor}, \text{sll}, \text{sra} } ]</td>
</tr>
<tr>
<td>State register</td>
<td>[ R_s = { r(\text{h}), r(\text{b}), r(\text{e}), r(\text{hb}), r(\text{pc}), r(\text{cp}), r(\text{tmp}_\text{cp}), r(\text{tr}) } ]</td>
</tr>
<tr>
<td>Argument register</td>
<td>[ R_a = { r(\text{I})</td>
</tr>
<tr>
<td>Permanent register</td>
<td>[ R_p = { p(\text{I})</td>
</tr>
<tr>
<td>Addressing mode</td>
<td>[ X = { \text{A} \cup R_s \cup R_p \cup { r(\text{h}), r(\text{b}) } \cup { T \cdot X</td>
</tr>
<tr>
<td>Instruction</td>
<td>[ \text{I} ] (The set of BAM instructions is defined in section 3.2 and Appendix B)</td>
</tr>
</tbody>
</table>

The encoding of information in tags is designed to simplify common operations. It is similar to the encoding used in the WAM (Figure 2.5). Atoms are represented as immediate values with a \text{tatom} tag. Integers are represented as themselves, and are considered to have \text{tint}, \text{tneg}, or \text{tneg} tags for the conditional branches that look at tags. Unbound variables are represented as pointers with a \text{tvar} tag that point to themselves or another unbound variable. Structures and lists are represented as pointers with tags \text{tstr} or \text{tlst}. They point to a contiguous block of their arguments on the heap. The main functor and arity of a structure are stored there encoded in a single word. The main functor and arity of a list (cons cell) are not stored since they are known implicitly.

The BAM defines five mappings to represent and access all data structures used during execution (Table 3.4). These mappings are the Register Set, the Heap, the Trail, the Code Space, and the Label Map. An infinite number of argument and permanent registers is assumed to exist. Of all registers, only the heap
Table 3.4 - Run-time data structures of the BAM

<table>
<thead>
<tr>
<th>Name</th>
<th>Definition</th>
</tr>
</thead>
<tbody>
<tr>
<td>Register Set</td>
<td>((R_u \cup R_e \cup R_p) \rightarrow W)</td>
</tr>
<tr>
<td>Heap</td>
<td>(W \rightarrow W)</td>
</tr>
<tr>
<td>Trail</td>
<td>(N \rightarrow W)</td>
</tr>
<tr>
<td>Code Space</td>
<td>(N \rightarrow I)</td>
</tr>
<tr>
<td>Label Map</td>
<td>(L \rightarrow N)</td>
</tr>
</tbody>
</table>

The pointer \(r(h)\) and the backtrack pointer \(r(b)\) are made explicit in the instruction set. The others are implicit in its execution. Environments and choice points are represented as register sets that are stored in registers \(r(e)\) and \(r(b)\), respectively. Prolog terms are stored in registers, on the heap, and on the trail. Compound terms are stored on the heap as sequences of words in the same manner as is done in the WAM (Figure 2.5). For all types except atoms, the value field of a word is a natural number that indexes into the heap, and therefore points to terms on the heap. For atoms, the value field is the symbolic atom itself. The correspondence between tags and Prolog data types is given in Table 3.5.

Table 3.5 - Correspondence of tags with Prolog data types

<table>
<thead>
<tr>
<th>Tag</th>
<th>Data type</th>
</tr>
</thead>
<tbody>
<tr>
<td>tvar</td>
<td>An unbound variable or a general pointer.</td>
</tr>
<tr>
<td>tstr</td>
<td>Pointer to a structure—a compound term with a functor and fixed number of arguments.</td>
</tr>
<tr>
<td>tlist</td>
<td>Pointer to a cons cell—a compound term consisting of two parts, a head and a tail.</td>
</tr>
<tr>
<td>tatm</td>
<td>An atom.</td>
</tr>
<tr>
<td>tpos</td>
<td>A nonnegative integer.</td>
</tr>
<tr>
<td>tneg</td>
<td>A negative integer.</td>
</tr>
<tr>
<td>tint</td>
<td>An integer.</td>
</tr>
<tr>
<td>tflt</td>
<td>A floating point number.</td>
</tr>
</tbody>
</table>

The following descriptions clarify the correspondence between BAM types and Prolog types:

1. The value corresponding to a pointer tag is an index into an array of words. This is normally implemented as an address.

2. The value corresponding to a tatm tag is a symbol that uniquely identifies an atom or the main functor of a structure. It is a Prolog atom or a Prolog structure of the form \(F/N\) where \(F\) is a Prolog atom representing the functor and \(N\) is a nonnegative integer representing the arity. For correctness, the assembler and run-time system must guarantee an exact correspondence between this symbol and the contents of the run-time symbol table, so that the built-ins name/2, functor/3, arg/3, and .../2 all work correctly.
(3) The value corresponding to a tpos or tneg tag is a nonnegative integer that represents the absolute value of the integer represented by the word.

(4) The value corresponding to a tint tag is an integer that represents the value of the integer represented by the word.

(5) The value corresponding to a tflt tag is a floating point number that represents the value of the number represented by the word.

Nothing is assumed about how these types are represented on a real machine. When the BAM is targeted to a real machine then the representation of types on the machine must be defined. The representation of types changes with different target machines, different versions of the system, and even different programs. The Implementation Manual [31] discusses how to port the BAM. Symbolic labels are pointers to code. Since mappings can be of any size, they are pointers to data stacks in memory. The representation of a word depends on the encoding used to represent tags on the machine, the word size of the machine, and on the encoding of Prolog atoms into unique bit patterns. For the VLSI-BAM processor, all four types are mapped into 32 bits and words consist of 4 bit tags and 28 bit values.

<table>
<thead>
<tr>
<th>Argument</th>
<th>Type</th>
</tr>
</thead>
<tbody>
<tr>
<td>X, Y, Z</td>
<td>Addressing modes, elements of X. Most instructions use a subset of all possible addressing modes.</td>
</tr>
<tr>
<td>L, L1, L2, L3</td>
<td>Branch destinations, elements of L.</td>
</tr>
<tr>
<td>N</td>
<td>A natural number, element of N.</td>
</tr>
<tr>
<td>A</td>
<td>A Prolog atom, element of A.</td>
</tr>
<tr>
<td>Tag</td>
<td>A tag value, element of T.</td>
</tr>
<tr>
<td>Eq</td>
<td>An equality condition, element of C_ε.</td>
</tr>
<tr>
<td>Cond</td>
<td>A condition, element of C.</td>
</tr>
<tr>
<td>Op</td>
<td>An arithmetic operation, element of E.</td>
</tr>
<tr>
<td>Reglist</td>
<td>A list of registers used in choice point management.</td>
</tr>
<tr>
<td></td>
<td>RegList ∈ {[α_0, α_1, ..., α_n]</td>
</tr>
</tbody>
</table>

3.2. An overview of the BAM

The BAM uses types and data structures similar to the WAM. It has registers and stacks similar to the WAM and uses a similar execution strategy. However, the instruction set is completely different. The BAM has a load-store instruction set that is extended with tagged addressing modes and a few primitive Prolog-specific instructions. A summary of the addressing modes and instructions is given in Tables 3.6.
through 3.10. All instructions use only a subset of the addressing modes given in Table 3.3. The instruction set includes:

- **Simple instructions** (Table 3.7). These are simple register-transfer level operations for a tagged architecture. They include move, push, conditional branch, and arithmetic. These instructions are used to implement many cases of unification and many built-in predicates.

- **Complex instructions** (Table 3.8). There are five frequently-used operations defined as single instructions: dereferencing (following a pointer chain to its end), trailing (saving a variable's address so it can be restored on backtracking), general unification (when the compiler cannot simplify the general case), choice point handling (saving and restoring state for backtracking), and environment handling (creating and removing local stack frames).

- **Embedded information** (Table 3.9). This allows a better translation to the assembly language of the target machine. This information is expressed in two ways: (1) with pragmas, which resemble instructions but are not executable, and (2) by extending instructions with additional arguments. An example of (1) is the tag pragma, which gives the tag of a load or a store, e.g.:

  ```
  pragma(tag(r(1), tvar)). % Register r(1) contains a tvar tag.
  move([r(1)], r(0)). % Load register r(0) from register r(1).
  ```

  By giving the tag at compile-time, this avoids tag masking on a general-purpose processor and allows the load to be done in a single cycle. An example of (2) is:

  ```
  unify(r(0), r(1), ?, nonvar, fail). % Register r(1) is nonvariable.
  ```

  This gives no information about r(0) but says that r(1) is nonvariable. This allows the unification to be done more efficiently because no check has to be done whether r(1) is unbound.

- **User instructions** (Table 3.10). The BAM language is extended with several instructions, registers, and tags that are never output by the compiler, but are intended for use only by a BAM assembly programmer. This allows the non-Prolog component of the run-time system to be written completely in BAM assembly. These instructions are described in Appendix D.
Table 3.7 - Simple instructions

<table>
<thead>
<tr>
<th>Instruction</th>
<th>Meaning</th>
</tr>
</thead>
<tbody>
<tr>
<td>equal(X,Y,L)</td>
<td>Branch to L if X and Y are not equal.</td>
</tr>
<tr>
<td>move(X,Y)</td>
<td>Move X to Y.</td>
</tr>
<tr>
<td>push(X,Y,N)</td>
<td>Push X on stack with stack pointer Y and post-increment N.</td>
</tr>
<tr>
<td>Op(X,Y,Z)</td>
<td>Perform the arithmetic operation Op on X and Y and store the result in Z. Trap if an operand or the result is not integer.</td>
</tr>
<tr>
<td>adda(X,Y,Z)</td>
<td>Full-word non-trapping add of a word X and an offset Y, giving a word Z.</td>
</tr>
<tr>
<td>pad(N)</td>
<td>Add N to the heap pointer.</td>
</tr>
<tr>
<td>switch(Tag,X,L1,L2,L3)</td>
<td>Three-way branch; branch to L1, L2, L3 depending on whether the tag of X is tvar, Tag, or any other value.</td>
</tr>
<tr>
<td>test(Eq,Tag,X,L)</td>
<td>Branch to L if the tag of X is equal or not equal to Tag.</td>
</tr>
<tr>
<td>hash(T,X,N,L)</td>
<td>Look up X in a hash table of length N located at L. If X is in the table then branch to the label in the table, else fall through. T ∈ {atomic, structure}.</td>
</tr>
<tr>
<td>pair(E,L)</td>
<td>A hash table entry. E is either an atom or a pair functor/arity.</td>
</tr>
<tr>
<td>jump(Cond,X,Y,L)</td>
<td>Jump to L if the arithmetic comparison of X and Y is true. Trap if an operand is not integer.</td>
</tr>
<tr>
<td>jump(L)</td>
<td>Jump unconditionally to L.</td>
</tr>
<tr>
<td>label(L)</td>
<td>L is a branch destination.</td>
</tr>
<tr>
<td>procedure(Name/Arity)</td>
<td>Mark the beginning of a procedure.</td>
</tr>
<tr>
<td>call(Name/Arity)</td>
<td>Call the procedure Name/Arity.</td>
</tr>
<tr>
<td>jump(Name/Arity)</td>
<td>Jump to the procedure Name/Arity.</td>
</tr>
<tr>
<td>return</td>
<td>Return from a procedure call.</td>
</tr>
<tr>
<td>simple_call(Name/Arity)</td>
<td>Non-nestable call used to interface with routines written in BAM assembly.</td>
</tr>
<tr>
<td>simple_return</td>
<td>Non-nestable return used for routines written in BAM assembly.</td>
</tr>
</tbody>
</table>

3.3. Justification of the complex instructions

The execution of Prolog requires five complex operations: dereferencing, trailing, unification, backtracking, and environment management. These operations are represented as single instructions in the BAM. In the WAM, dereferencing, trailing, and unification are done implicitly by many instructions even when they are not needed. Making them explicit allows the compiler to minimize their use as much as possible by doing them only when they are really needed.

The complex instructions could be expanded into sequences of simple instructions; however, this expansion is not done at the BAM level but is delayed to the machine level. There are two reasons for this:

1. Some machines may implement part or all of a complex instruction directly. Expanding it into simple instructions is therefore premature since it would make this harder to detect. For example, the VLSI-BAM processor has support for some complex instructions (e.g. dereferencing, trailing, and unification).
Table 3.8 – Complex instructions

<table>
<thead>
<tr>
<th>Instruction</th>
<th>Meaning</th>
</tr>
</thead>
<tbody>
<tr>
<td>deref(X, Y)</td>
<td>Dereference X and store result in Y.</td>
</tr>
<tr>
<td>trail(X)</td>
<td>Push X on the trail stack if the trail condition is satisfied.</td>
</tr>
<tr>
<td>unify(X, Y, Tx, Ty, L)</td>
<td>General unification of X and Y, branch to L if fail. Trailing is done by this instruction. Extra parameters Tx, Ty ∈ {?, var, nonvar} give information to improve the translation. They are not needed for correctness. Unify X with the atom A and branch to L if fail. No trailing is done by this instruction.</td>
</tr>
<tr>
<td>unify_atomic(X, A, L)</td>
<td></td>
</tr>
<tr>
<td>allocate(N)</td>
<td>Create an environment of size N on the local stack.</td>
</tr>
<tr>
<td>deallocate(N)</td>
<td>Remove the top-most environment from the local stack.</td>
</tr>
<tr>
<td>choice(I/N, RegList, L)</td>
<td>Create a choice point containing the registers listed in RegList and set the retry address to L.</td>
</tr>
<tr>
<td>choice(I/N, RegList, L)</td>
<td>(I&lt;≤N) Restore the argument registers listed in RegList from the current choice point, and modify the retry address to L.</td>
</tr>
<tr>
<td>choice(N/N, RegList, fail)</td>
<td>Restore the argument registers listed in RegList from the current choice point, and pop the current choice point from the choice point stack.</td>
</tr>
<tr>
<td>fail</td>
<td>Restore the machine state (except the argument registers) from the most recent choice point, restore to unbound all variables on the trail that were bound and trailed since the creation of this choice point, and transfer control to the retry address.</td>
</tr>
<tr>
<td>move(r(b), X)</td>
<td>Move the backtrack pointer to X. This must be done at the entry of any predicate containing a cut.</td>
</tr>
<tr>
<td>cut(X)</td>
<td>Make the choice point pointed to by X the new top of the choice point stack.</td>
</tr>
</tbody>
</table>

Table 3.9 – Embedded information (pragmas)

<table>
<thead>
<tr>
<th>Instruction</th>
<th>Meaning</th>
</tr>
</thead>
<tbody>
<tr>
<td>pragma(align(X,N))</td>
<td>The contents of location X are a multiple of N.</td>
</tr>
<tr>
<td>pragma(tag(X,Tag))</td>
<td>The contents of location X have tag Tag.</td>
</tr>
<tr>
<td>pragma(push(term(N)))</td>
<td>A term of size N is about to be created on the heap.</td>
</tr>
<tr>
<td>pragma(push(cons))</td>
<td>A cons cell is about to be created on the heap.</td>
</tr>
<tr>
<td>pragma(push(structure(N)))</td>
<td>A structure of arity N is about to be created on the heap.</td>
</tr>
<tr>
<td>pragma(push(variable))</td>
<td>An unbound variable is about to be created on the heap.</td>
</tr>
<tr>
<td>pragma(hash_length(N))</td>
<td>A hash table of length N is about to be created.</td>
</tr>
</tbody>
</table>

(2) For best performance, optimizations should be done at all levels. The BAM level makes certain optimizations easy, e.g. the determinism optimization in Chapter 6. Keeping the complex operations as single instructions allows them to be optimized directly. For example, if a variable is dereferenced twice then the second dereference can be removed. This is much harder to detect if the dereference instruction is expanded into a loop.

It is best to avoid assumptions about the characteristics of the target machine. In the cases where such assumptions would be useful, the BAM uses pragmas to give the information without compromising the
machine independence. The translator is free to use or ignore this information.

3.4. Justification of the instructions needed for unification

This section constructs the BAM instructions that contain the required instructions and addressing modes to support unification. It turns out that both simple and complex instructions are necessary to support unification. The instructions are constructed starting from an algorithm for unification and a very general intermediate language. The algorithm is decomposed into specialized instructions depending on the form of the data known at compile-time.

The two starting points are (1) an algorithm for unification (a specification of a unification algorithm is given in Appendix C), and (2) a very general instruction set. The method proceeds in a top-down manner by decomposing the unification algorithm into specialized instructions depending on information about the form of the data known at compile-time (Figure 3.4).

This method is inspired by Kursawe [41] and Holmer [32]. Kursawe applies partial evaluation and specialization in a top-down manner starting from a Prolog program and obtains an instruction set resembling the WAM. Holmer describes several techniques for the automatic design of instruction sets, of which decomposition is one. To go beyond the WAM it is necessary to make assumptions about the architecture, a step that Kursawe does not take. The design of the BAM starts with a general instruction set that does make these assumptions.

The choice of what general instruction set to start with is important. It is not useful to start with an instruction set that has too little expressive power, for example one with a limited set of addressing modes.

<table>
<thead>
<tr>
<th>Instruction</th>
<th>Meaning</th>
</tr>
</thead>
<tbody>
<tr>
<td>ord(X,Y)</td>
<td>Extract the value of X and move it to Y.</td>
</tr>
<tr>
<td>val(T,X,Y)</td>
<td>Create the word Y from the tag T and the value X.</td>
</tr>
<tr>
<td>jump_reg(R)</td>
<td>Jump to address stored in register R.</td>
</tr>
<tr>
<td>jump_nt(Cond,X,Y,L)</td>
<td>Jump to L if the full word comparison of X and Y is true. Never trap.</td>
</tr>
<tr>
<td>Op_nt(X,Y,Z)</td>
<td>Perform the full word arithmetic operation Op (except multiply and divide) on X and Y and store the result in Z. Never trap.</td>
</tr>
<tr>
<td>trail_bda(X)</td>
<td>Push address X and the value stored there on the trail stack if the trail condition is satisfied. This is a special trail instruction for backtrackable destructive assignment.</td>
</tr>
</tbody>
</table>
because the required addressing modes are not yet known. Prematurely decomposing complex instructions into simple ones side-steps the results.

The following assumptions are made:

1. The architecture is sequential and of Von Neumann design with multiple registers.
2. The basic data element is a word, which is large enough to contain an address. A register holds one word.
3. The instructions have three parts:
   - An action. Some sample actions are data movement (move, push), conditional branching (equal), and general unification (unify). Other important actions are multi-way branching (switch) and several Prolog-specific operations (deref, trail).
   - A set of arguments. Unification acts on two operands, so typically two arguments are sufficient.
   - A set of destination addresses. Depending on the outcome of the action, control continues at one of the destinations. The size of the set and the meaning of its members depends on the action. The address of the next instruction in the instruction stream is an implicit member of
Arguments are referenced with multiple addressing modes. An infinite set of addressing modes are defined in Table 3.3. The instructions derived in this section will need only finite subset. For clarity, Table 3.11 gives some abbreviations useful for this subset.

<table>
<thead>
<tr>
<th>Notation</th>
<th>Meaning</th>
</tr>
</thead>
<tbody>
<tr>
<td>Disp</td>
<td>a positive heap displacement (bounded by the size of a term).</td>
</tr>
<tr>
<td>Offset</td>
<td>a nonnegative offset into a structure (bounded by the arity).</td>
</tr>
<tr>
<td>Imm</td>
<td>an immediate value; an atom or a numeric constant.</td>
</tr>
<tr>
<td>Var</td>
<td>a variable local to a clause, i.e. r(I) or p(J).</td>
</tr>
<tr>
<td>Arg</td>
<td>denotes Var or [Var+Offset].</td>
</tr>
</tbody>
</table>

Construction of the instruction set proceeds in the following steps. The data representation has already been fixed (section 3.1). The existence of two forms of unification (read mode and write mode) and the need for dereferencing and a three-way branch is shown. The instructions required for read mode and write mode are constructed. Finally, the effects of variable representation (in registers or on the environment) on the instruction set are discussed.

3.4.1. The existence of read mode and write mode

The compilation of the unification \( T_1 = T_2 \), where \( T_1 \) and \( T_2 \) are two arbitrary terms, is reduced to the compilation of \( V = T \) where at compile-time \( V \) is a variable and \( T \) is any term. At run-time there are two values of \( V \) that result in different actions of the unification algorithm:

1. \( V \) is an unbound variable, in which case \( T \) is constructed on the fly and bound to \( V \) (this is called write mode). To satisfy the standard definition of unification, when \( T \) is bound to \( V \) a check needs to be done (the occur check) that \( T \) does not contain \( V \). Following Prolog implementation convention, this check is ignored for efficiency reasons.

2. \( V \) is a nonvariable term, in which case it is checked that the form of \( V \) matches \( T \), and the algorithm is invoked recursively for the term’s arguments (this is called read mode).
3.4.2. The need for dereferencing

Unifying two unbound variables makes one point to the other. Doing this several times leads to pointer chains, with the common value of all the variables in a single location at the end of the chain. To get a variable's value, the pointer chain is followed to its end, an operation known as dereferencing. It can be provided as an addressing mode or as a separate instruction. Making it an instruction avoids repeated dereferencing. Therefore the following instruction is added:

\[ \text{deref(Var1,Var2)} \]

First \( \text{Var1} \) is moved to \( \text{Var2} \). Then the tag of \( \text{Var2} \) is checked. If it is an unbound variable (\( t\text{var} \)) it reads memory and a loop is entered replacing \( \text{Var2} \) by the referenced value while its tag is \( t\text{var} \) and its pointer part is different from \( \text{Var2} \). A two-argument dereference is chosen over a single-argument dereference because it allows a more compact representation of write-once variables (Chapter 5).

It is assumed in what follows that \( V \) and \( T \) are dereferenced when necessary, in particular that both the \text{trail} and \text{unify} instructions are always given dereferenced arguments.

3.4.3. The need for a three-way branch

The code for a unification \( V = T \) consists of three parts: (1) a check whether \( V \) is an unbound variable or a nonvariable for choosing between write mode and read mode unification, (2) the instructions for read mode unification, and (3) the instructions for write mode unification.

The tag field is available directly for the check of (1). The check has three possible results: the tag of \( V \) matches a known tag (read mode), the tag is an unbound variable tag (write mode), or the tag is neither (failure). This implies the following three-way branch:

\[ \text{switch(Tag, Var, VarLbl, TagLbl, FailLbl)} \]

If the tag of \( \text{Var} \) is \( t\text{var} \) (an unbound variable) then jump to \( \text{VarLbl} \). If the tag of \( \text{Var} \) matches \( \text{Tag} \) then jump to \( \text{TagLbl} \). Otherwise jump to \( \text{FailLbl} \). The failure address is explicit instead of implicit to allow the implementation of fast incremental shallow backtracking.
3.4.4. Constructing the read mode instructions

The general case of read mode unification is \( V = T \), where at compile-time \( V \) is a variable or an argument of a compound term, and \( T \) is a term. The first argument of each instruction is the value of \( V \).

Two locations are possible for its value:

- \( \text{Var} \)  
  \( V \) is a variable
- \( \text{Var} + \text{Offset} \)  
  \( V \) is an argument of a compound term

The abbreviation \( \text{Arg} \) is used to denote one of these two addressing modes (Table 3.11). The second argument and the action are determined by the compile-time knowledge of \( T \). The possibilities are:

1. \( T \) is partially or wholly known at compile-time. The possible information known about \( T \) is:
   - \( T \) is an unbound variable that has not yet been initialized, e.g. because it is the first occurrence in the clause. \( V \) is moved directly to \( T \).
   - \( T \) is an unbound variable. \( V \) is stored to \( T \)'s location in memory.
   - \( T \) is atomic. Unification reduces to a check that \( T \) and \( V \) have the same atomic value. If the values do not match the unification fails.
   - \( T \) is compound. Unification reduces to a check that \( V \) has the correct functor and arity, followed by a unification of its arguments with \( T \)'s arguments. If \( V \)'s arguments are loaded into registers then the unification can be compiled recursively. It follows that arbitrarily deep nesting of addressing modes is not necessary if one instruction is added:

   \[
   \text{move}([\text{Var} + \text{Offset}], \text{Var})
   \]

2. Nothing is known about \( T \) at compile-time. The unification of \( V \) and \( T \) requires a general unification.

The following table of primitive instructions summarizes the action and both arguments:
The instructions equal and unify both can fail, so they have a failure address as third argument. The equal instruction compares its arguments and jumps to FailLbl if they are not equal.

General unification (unify) is the most complex instruction. If the unification fails it jumps to FailLbl. This instruction can be implemented using only the other instructions. However, it seems that one additional instruction is useful: a multi-way branch with a different destination for each possible tag value. If there are many possible tags this implies the existence of a jump table in memory, so that the instruction must do a memory reference before it can branch. Instead of using this instruction, another approach is to use a multilevel tree based on the three-way branch. Both approaches are viable since general unification is used rarely in real programs. According to measurements done by Holmer for several large programs [33], general unification takes about 4% of the total execution time of the VLSI-PLM [61]. More than 95% of these calls have arguments that are not compound terms of the same type and therefore do not need the recursive algorithm.

3.4.5. Constructing the write mode instructions

The general case of write mode unification is \( V = T \), where \( V \) is known to be an unbound variable at run-time and \( T \) is a term. Assume that the term \( T \) is created on a stack (called the heap) with a minimal number of move instructions. This assumption forces us to derive the form that a compound term has on the heap. The following are the possible values of words of a compound term:

- \( \text{Var} \): a variable (assumed initialized)
- \( \text{Tag} \cdot \text{Imm} \): a simple subterm of \( T \)
- \( \text{Tag} \cdot (r \cdot (h) \cdot \text{Disp}) \): a pointer to a compound subterm of \( T \)

These are the source addressing modes for the move instructions. A variable \( \text{Var} \) does not have to be
dereferenced when it is stored on the heap because its value is not read. The destination of the move instruction is a location on the heap. This location can be addressed either by a displacement addressing mode offset from the heap pointer \( r(h) \), i.e. \( r(h) - \text{Disp} \), or by an auto-increment addressing mode, i.e. a push instruction. The BAM uses the auto-increment addressing mode, for these reasons:

1. Preliminary studies using exhaustive search [32] show that with the VLSI-BAM microarchitecture the optimal way to create structures in write mode is by means of the idiom "load register, load register, double-word push", i.e. two registers are loaded and then pushed in a single instruction.

2. Instruction encoding is compacter, i.e. a push does not need a displacement field.

3. In the VLSI-BAM architecture the push instruction is given a displacement field anyway. This allows efficient implementation of uninitialized variables. For example, a cons cell whose cdr is uninitialized can be created with a single push that has a displacement of 2.

4. In the VLSI-BAM architecture the use of a push instruction allows a cache optimization: when pushing a dirty line it is not necessary to flush the line first [17]. This optimization was first done in the PSI-II architecture [52].

To summarize, to create a term on the heap it is sufficient to choose from the following set of three instructions (where \( r(h) \) is the stack pointer and \( l \) is the increment):

\[
\begin{array}{l}
\text{push(Var, } r(h), l) \\
\text{push(Tag"Imm, } r(h), l) \\
\text{push(Tag"(r(h)-Disp), } r(h), l)
\end{array}
\]

It is also necessary to bind the term to \( V \). This requires us to consider the form an unbound variable can take. There are two possibilities:

1. \( V \) has not yet been initialized, e.g. because it is the first occurrence in the clause. The term is moved directly to \( V \).

2. \( V \) has been initialized; it points to a location in memory. The term is stored in this location.

These two possibilities result in the following two instructions:
move(A, Var) store directly to a variable
(variable is not initialized)
move(A, [Var]) store to variable's location
(variable is initialized)

The addressing mode of the argument A depends on whether the term is compound or simple, and if it is simple, whether it is an atom or a variable. This results in three possible values for A:

Var a simple term (variable)
Tag·Imm a simple term (nonvariable)
Tag·r(h) a compound term (on the heap)

In addition to the above instructions, it is also necessary to initialize the first occurrence of a variable. One way to do this is:

```
move(tvar·(r(h)-Disp), Var)
push(Var, r(h), 1)
```

With these instructions it is possible to create a term of size n on the heap in n pushes, a great improvement over the WAM, which requires \( n + f - 1 \) stores, \( f - 1 \) dereference operations, and \( f - 1 \) trail checks, where \( f \) is the number of functors in the term. This idea was first proposed by André Mariën [44].

3.4.6. Representation of variables

Assume that the execution model represents variables local to a clause in an environment, or stack frame. There is a dedicated register \( r(e) \), called the environment pointer, that points to the current environment in the environment stack. Variables local to a clause are stored either in registers or in an environment, so the notation Var denotes one of the following two addressing modes:

\[
\begin{align*}
\text{r(I)} & \quad \text{a variable in a register} \\
\text{p(J)} & \quad \text{a variable on the environment stack}
\end{align*}
\]

where \( p(J) \) is implemented as an offset into the environment, i.e. as \( [r(e)+J'] \) for some \( J' \). This implies that double indirection is possible: the addressing mode \( [\text{Var}+\text{Offset}] \) is \( [p(J)+\text{Offset}] \), when \( \text{Var} \) is an environment variable. The double indirection is avoided by including one instruction:

```
move(p(J), r(I))
```
### Table 3.12 - Data movement instructions for unification

<table>
<thead>
<tr>
<th>Read mode</th>
<th>Write mode</th>
</tr>
</thead>
<tbody>
<tr>
<td>move(Arg, Var)</td>
<td>push(Tag-Imm, r(h), 1)</td>
</tr>
<tr>
<td>move(Arg, [Var])</td>
<td>push(Tag-Imm, r(h), 1)</td>
</tr>
<tr>
<td>equal(Arg, Var, F)</td>
<td>push(Tag-ax, r(h) + Displ), r(h), 1</td>
</tr>
<tr>
<td>equal(Arg, Tag-Imm, F)</td>
<td>move(Tag-Imm, [Var])</td>
</tr>
<tr>
<td>unify(Arg, Var, F)</td>
<td>move(Tag-Imm, [Var])</td>
</tr>
</tbody>
</table>

### Table 3.13 - Control flow and other instructions for unification

<p>| | |</p>
<table>
<thead>
<tr>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>switch(Tag, Var, VarLbl, TagLbl, F)</td>
<td>three-way branch</td>
</tr>
<tr>
<td>jump(Lbl)</td>
<td>join read and write mode paths</td>
</tr>
<tr>
<td>deref(Var1, Var2)</td>
<td>dereference a pointer chain</td>
</tr>
</tbody>
</table>

### 3.4.7. Summary of the unification instructions

This section summarizes the BAM instructions necessary to support unification. Tables 3.12 and 3.13 present the instructions. They use only a small finite subset of the addressing modes of Table 3.3.

The following typical instructions illustrate the meaning of the notation:

- `move(tatm-axe, r(3))` Move the atom axe into register r(3).
- `move([r(3)+5], r(4))` Move the word located at address r(3)+5 into r(4).
- `equal(r(2), tatm-cat, F)` If r(2) is equal to the atom cat then fall through, else jump to label F.
- `unify(p(2), p(3), F)` Unify the term located in p(2) with the term located in p(3). Jump to label F if the unification fails.
- `switch(tatm, r(3), V, T, F)` If r(3)'s tag is tvar then jump to label V. If r(3)'s tag is tatm then jump to label T. Otherwise, jump to label F.
Chapter 4

Kernel transformations

I. Introduction

Four optimizing transformations are done on the kernel Prolog representation of programs: formula manipulation, factoring, global dataflow analysis, and determinism extraction. The goal of the transformations is to reduce a single metric: the total execution time of all unifications in the program. This metric is approximated by the number of unifications and by the size of the terms being unified. The chapter first describes the representation of types as logical formulas in the compiler. This is followed by a description of each of the four transformations:

(1) Formula manipulation. The compiler implements a set of primitive transformations to replace Prolog code and types (both are represented as logical formulas) with simpler versions that have identical semantics. The simplicity of a formula is defined as the number of goals in the formula. These transformations are done whenever there is a possibility that the code is too complex, i.e., upon reading in a program and after other transformations such as the determinism transformation (see below).

(2) Factoring. This transformation groups sets of clauses in a predicate together if they have head unifications in common. This reduces the number of head unifications and shallow backtracking steps.

(3) Global dataflow analysis. This stage analyzes the program, annotates it with types, and restructures it. The analyzer uses abstract interpretation to determine the types of predicate arguments.

(4) Determinism transformation. This stage rewrites the program to make its determinism explicit, i.e. it replaces shallow backtracking by conditional branching. Many of the other transformations in this chapter are chosen to make this transformation possible more often. The transformation converts the predicate into a series of nested case statements. Sometimes this is only partially successful; certain branches of the case statements may still retain disjunctions (OR choices) that could not be converted into deterministic code.
To improve readability, the examples in this chapter are given in standard Prolog notation. It is understood that they are represented internally in kernel Prolog.

2. Types as logical formulas

Throughout the compiler, type information about variables is represented with logical formulas. During compilation, any information learned is added to the formula, and deduction based on the formula simplifies the generated code. It is a simple and powerful approach to avoid doing redundant operations at run-time. For example, if a variable is dereferenced once, then it should never be dereferenced again.

Types in the compiler are defined as follows:

Definition T: Given a predicate \( f/\mathbf{X} \) with main functor \( f \) and arity \( \mathbf{n} \), a type of \( f/\mathbf{X} \) is a term 
\[
(f(A_1, A_2, \ldots, A_n) \leftarrow \text{Formula})
\]
where the \( A_1, A_2, \ldots, A_n \) are \( \mathbf{n} \) distinct variables and Formula is a logical formula (i.e. a Prolog term).

For example, the type \( \text{range}(A, B, C) \leftarrow \text{integer}(A), \text{var}(B), \text{integer}(C) \) says that the first and third arguments of \text{range}/3 are integers and the second argument is an unbound variable. The compiler recognizes all Prolog type-checking predicates in the type formula. Appendix A gives a table of the types recognized by the compiler. In addition to these types, several other types are recognized that do not correspond to Prolog predicates. These types introduce distinctions between objects that depend on the implementation and are indistinguishable in the language, for example, the difference between an integer and a dereferenced integer, and the difference between an unbound variable that is not aliased to any other and an unbound variable that may be aliased. The following types are recognized that do not exist as Prolog predicates:

<table>
<thead>
<tr>
<th>Internal Type</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>uninit(X)</td>
<td>( X ) is an uninitialized memory argument.</td>
</tr>
<tr>
<td>uninit_mem(X)</td>
<td>( X ) is an uninitialized memory argument.</td>
</tr>
<tr>
<td>uninit_reg(X)</td>
<td>( X ) is an uninitialized register argument.</td>
</tr>
<tr>
<td>unbound(X)</td>
<td>( X ) is one of the types ( \text{uninit_mem}(X), \text{uninit_reg}(X), \text{var}(X) ).</td>
</tr>
<tr>
<td>deref(X)</td>
<td>( X ) is dereferenced, i.e. it is accessible without following any pointers.</td>
</tr>
<tr>
<td>rderef(X)</td>
<td>( X ) is recursively dereferenced, i.e. it is dereferenced, and if it is compound then all its arguments are recursively dereferenced.</td>
</tr>
</tbody>
</table>
These types should not be given by the programmer since incorrect code or a significant loss of efficiency may result if they are used incorrectly. For example, declaring an argument of a predicate to be of uninitialized register type, i.e. the argument is an output that is passed in a register, may cause a large increase in stack space used by the program if that predicate is the last goal in a clause, because last call optimization is not possible in that instance. The safe approach is to leave the use of these types up to the compiler.

The use of logical formulas to hold information during compilation can be contrasted with the use of a symbol table in a compiler for an imperative language. Representing types as logical formulas has two advantages over a symbol table: (1) They are more flexible during compiler development. The kind of information stored in a symbol table must be known when the compiler is designed. Formulas can contain kinds of information that are not known during the compiler's design. (2) They lend themselves to powerful, symbolic manipulation such as deduction. Improving the deductive abilities leads to better code without having to change any other part of the compiler. The disadvantage of this representation is that its manipulation is slow. Future versions of the compiler could use a representation that is faster in the common cases.

Type formulas are used in the following ways in the compiler:

1. Representing type information known about a set of variables. For example, the formula \( \text{var}(X), \text{atom}(Y) \) means that \( X \) is an unbound variable and \( Y \) is an atom. The user manual (Appendix A) lists the types recognized by the compiler.

2. Using a primitive form of deduction to simplify the generated code. For example, assume the formula is \( \text{list}(X), \text{var}(Y), \text{deref}(Z), \ldots \). To compile a run-time check that \( X \) is a non-variable, the compiler first checks whether this formula implies \( \text{nonvar}(X) \). This is true because \( \text{list}(X) \) implies \( \text{nonvar}(X) \), so no run-time check is necessary.

3. Updating the type formula when new information is learned. After compiling a goal, the formula is updated to represent the new knowledge that is gained. For example, after executing the arithmetic

---

* Of course, both the assembler and the run-time system use standard symbol tables.
expression \( X \) is \( A+B \) it is known that \( X \) is an integer, so the formula is extended with \( \text{integer}(X) \).

In most cases, logical formulas are immutable, e.g. when a variable \( X \) is known to be a list (represented as \( \text{list}(X) \)), that fact remains true forever. This is not true for all types. The types used to denote unbound variables (e.g. \( \text{var}(X) \) and \( \text{uninit}(X) \)) become false as a result of an instantiation. This is also true of the standard order comparisons (e.g. \( X < Y, X > Y \), and so forth) and the types \( \text{deref}(X) \) and \( \text{reref}(X) \). The compiler is careful to take this into account when updating the type formula.

| Table 4.1 – Primitives to manipulate logical formulas and Prolog formulas |
|-----------------|---------------------------------|
| Primitive       | Description                      |
| \( F_1 \) implies \( F_2 \) | Implication: Succeeds if it can determine that there does not exist an assignment to variables in \( F_1 \) and \( F_2 \) that causes both \( F_1 \) and \( \text{not}(F_2) \) to succeed. |
| \( F_2 := \text{simplify}(F_1) \) | \( F_2 \) is a simplification of \( F_1 \). |
| \( F_2 := \text{subsume}(F,F_1) \) | \( F_2 \) is a simplification of \( F_1 \), given that \( F \) is true. |
| \( F_2 := \text{update_formula}(F,F_1) \) | \( F_2 \) is the result of removing information contradicted by \( F \) from \( F_1 \) and adding \( F \) to \( F_1 \). |

3. Formula manipulation

The compiler implements a set of primitive transformations to manipulate formulas. They are summarized in Table 4.1, where \( F, F_1, \) and \( F_2 \) are logical formulas. Each of these primitives has two versions: a pure logical and a Prolog version. The logical version is used to manipulate types (see previous section). It assumes the formula has a purely logical meaning, i.e. that the operational concepts of execution order of goals, number of solutions, and backtracking behavior are not important. The Prolog version is used to manipulate kernel Prolog code. It assumes the formula must keep Prolog’s operational semantics.

Implication is implemented to work well with most combinations of Prolog predicates that are used in type declarations. The following examples all return with success:
Table 4.2 – Examples of simplification

<table>
<thead>
<tr>
<th>Formula</th>
<th>Simplified formula logical</th>
<th>Simplified formula Prolog:</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>(true; true)</td>
<td>true</td>
<td>(true; true)</td>
<td>The Prolog version is unchanged unless the compiler option same_number_solutions is disabled.</td>
</tr>
<tr>
<td>(p,fail)</td>
<td>fail</td>
<td>(p,fail)</td>
<td>The Prolog version is unchanged unless the compiler can deduce that p has no side effects (read / write or assert / retract).</td>
</tr>
<tr>
<td>(!,p ; q)</td>
<td>(p ; q)</td>
<td>(!,p)</td>
<td>Cut is logically identical to true, but it must be retained since it modifies backtracking behavior in the entire clause containing it.</td>
</tr>
</tbody>
</table>

atom(X) implies nonvar(X)
X<Y implies integer(X)
X<5 implies X<10
uninit(X) implies deref(X)
functor(X, _, 0) implies atomic(X)
(X==a; X==b) implies atom(X)

Simplification is done on standard Prolog, on kernel Prolog, and on type formulas. Table 4.2 gives some examples to illustrate the difference between logical and Prolog semantics. A single function simplify(F) handles both logical and Prolog semantics (Figure 4.1). For conciseness, the definition of simplify(F) uses the compound terms (A, B), (A; B), (A->B), and (\+(A)) both as selectors (to choose the branch of the case statement) and constructors (in the calls to simp_step(F)). Tables 4.3 and 4.4 define part of the definition of simp_step(F), the primitive simplification step. The complete definition contains about 50 rules. The functions subsume(F, F1) and update_formula(F, F1) are implemented in a similar way.

```
function simplify(F : formula) : formula;
begin
    case /* decompose the formula */
        F = (A, B) : return simp_step( simplify(A), simplify(B) ) ; /* and */
        F = (A; B) : return simp_step( simplify(A); simplify(B) ) ; /* or */
        F = (A->B) : return simp_step( simplify(A)->simplify(B) ) ; /* implies */
        F = \+(A) : return simp_step( \+(simplify(A)) ) ; /* negation */
    otherwise : return simp_step(F);
end
end;
```

Figure 4.1 – Simplification of a formula
### Table 4.3 – Simplification rules (part of simp_step's definition)

<table>
<thead>
<tr>
<th>Rule</th>
<th>Input formula</th>
<th>Output formula</th>
<th>Condition to apply this rule</th>
</tr>
</thead>
<tbody>
<tr>
<td>(true,A)</td>
<td>A</td>
<td>(none)</td>
<td>no_side_effects(A) ∧ diff_sol ∧ no_bind(A)</td>
</tr>
<tr>
<td>(A,true)</td>
<td>A</td>
<td>(none)</td>
<td>semantics(prolog)</td>
</tr>
<tr>
<td>(true:A)</td>
<td>true</td>
<td>semantics(prolog) ∧ no_side_effects(A) ∧ diff_sol ∧ no_bind(A)</td>
<td></td>
</tr>
<tr>
<td>(true:A)</td>
<td>true</td>
<td>semantics(logical)</td>
<td></td>
</tr>
<tr>
<td>(A,fail)</td>
<td>fail</td>
<td>semantics(prolog) ∧ no_side_effects(A)</td>
<td></td>
</tr>
<tr>
<td>(A,fail)</td>
<td>fail</td>
<td>semantics(logical)</td>
<td></td>
</tr>
<tr>
<td>(fail,A)</td>
<td>fail</td>
<td>(none)</td>
<td></td>
</tr>
<tr>
<td>(fail;A)</td>
<td>A</td>
<td>(none)</td>
<td></td>
</tr>
<tr>
<td>(A-&gt;true:B)</td>
<td>A</td>
<td>semantics(prolog) ∧ succeeds(A) ∧ deterministic(A)</td>
<td></td>
</tr>
<tr>
<td>(A-&gt;true:B)</td>
<td>A</td>
<td>semantics(logical) ∧ succeeds(A)</td>
<td></td>
</tr>
<tr>
<td>(fail;A)</td>
<td>A</td>
<td>semantics(prolog) ∧ fails(A) ∧ no_side_effects(A)</td>
<td></td>
</tr>
<tr>
<td>(fail;A)</td>
<td>A</td>
<td>semantics(logical) ∧ fails(A)</td>
<td></td>
</tr>
</tbody>
</table>

### Table 4.4 – The conditions for applying simplification rules

<table>
<thead>
<tr>
<th>Condition</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>semantics(S)</td>
<td>Simplify according to semantics S where S ∈ {prolog, logical}.</td>
</tr>
<tr>
<td>no_side_effects(A)</td>
<td>Formula A does not have side effects when executed.</td>
</tr>
<tr>
<td>deterministic(A)</td>
<td>Formula A gives only one solution when executed.</td>
</tr>
<tr>
<td>no_bind(A)</td>
<td>Formula A does not bind any variables.</td>
</tr>
<tr>
<td>diff_sol</td>
<td>Relax semantics of Prolog to allow a different number of solutions.</td>
</tr>
<tr>
<td>succeeds(A)</td>
<td>Formula A always succeeds when executed.</td>
</tr>
<tr>
<td>fails(A)</td>
<td>Formula A always fails when executed.</td>
</tr>
</tbody>
</table>

4. Factoring

Factoring is based on the operation of finding the most-specific-generalization, or MSG, of two terms. Factoring collects groups of clauses whose heads can be combined in nontrivial fashion using the MSG operation. The advantage of factoring is that it reduces the number of unifications performed during execution. Figure 4.2 defines the MSG in terms of unification. Given two terms $T_1$ and $T_2$, consider the set $M$ of all terms that unify with both of them. The MSG of $T_1$ and $T_2$ is the unique element $T_m$ of $M$ which unified with any other element $U$ of $M$ gives $T_m$. Intuitively, this says that $T_m$ contains the maximal common information of $T_1$ and $T_2$.

The MSG (also called anti-unification) is the dual operation to unification. Given two terms, unification finds a term that is a more instantiated case of each of the two, i.e. the most general common instance of the two. The MSG is a term of which each of the two is a more instantiated case. For example, consider the two compound terms $s(A,x,C)$ and $s(A,B,y)$. Unifying these two terms results in $s(A,x,y)$. The MSG of the two terms is $s(A,B,C)$. Unification may fail, i.e. the most general unifier
function msg(T₁, T₂ : term) : term;
var
    M : set of term;
    Tₘ, U : term;
begin
    M := { T \mid T unifies with T₁ and T unifies with T₂ };
    Find Tₘ ∈ M such that ∀ U ∈ M : unify(U, Tₘ) = Tₘ;
    return Tₘ
end;

Figure 4.2 – The most specific generalization

is the empty set. Finding the MSG never fails. In the worst case, the generalization of the two terms is an unbound variable, which represents the set of all terms. For example, consider the two atomic terms x and y. Unifying these two results in failure, whereas the MSG is an unbound variable.

Another way of viewing the MSG operation is as an approximation to the union of two sets. Every term corresponds to a set by instantiating the variables in the term to all possible ground values. In general, the union of two of these sets does not correspond to any term. The MSG finds the smallest superset of the union that is represented by a term. A similar property holds of unification: it finds the largest subset of the intersection that is represented by a term.

For all arguments of the predicate, the factoring transformation finds the largest contiguous set of clauses whose MSG is a compound term. This set is used to define a dummy predicate and the definition of the original predicate is modified to call the dummy predicate. The algorithm is given in Figure 4.3. As an example of factoring, consider the predicate:

h([x_1]).
h([y_1]).
h([1]).

The lists in the heads of the first two clauses are combined: the MSG of \{x_1\} and \{y_1\} is \{_1\}.

The result after factoring is:
procedure factoring:
var
M : term;
C_i , C_i' : clause;
\pi, P_\pi : list of clause;
a, i, p, q : integer;
begin
for each predicate \( P \) in the program do begin
  /* At this point \( P = \{ C_1, C_2, ..., C_n \} \) (list of \( n \) clauses) */
  /* and \( C_i = (H_i ; : B_i) \) (Each clause has head \( H_i \) and body \( B_i \)) */
  for \( a := 1 \) to arity(\( P \)) do begin
    Partition \( P \) such that each contiguous group \( \pi = \{ C_p, C_{p+1}, ..., C_q \} \) (1 \( \leq p \leq q \leq n \)
    satisfies exactly one of the two properties:
    1. Either \( p = q \) (\( \pi \) contains only one clause), or
    2. \( \pi \) is the largest group for which \( M = \sigma_i S G \) (argument \( a \) of \( H_i \)) is compound.
    for each contiguous group \( \pi \) do if \( p < q \) then begin
      /* Create the dummy predicate \( P_\pi \) */
      for \( i := p \) to \( q \) do begin
        \( C_i' := C_i \);
        Remove \( M \) from \( H_i' \);
        Add all variables in \( M \) as arguments to \( H_i' \);
      end;
      \( P_\pi := \{ C_p', ..., C_q' \} \);
      /* Create the call to the dummy predicate */
      \( H := \) (new head with same functor and arity as \( P \) and \( M \) in argument \( a \));
      \( H_a := \) (new head with same functor and arity as \( P_a \));
      for \( i := 1 \) to arity(\( P \)) do if \( i \neq a \) then begin
        Make argument \( i \) of \( H \) and \( H_a \) identical
      end;
      Replace \( \pi \) in \( P \) by the single clause \( C_\pi = (H := H_a) \)
    end
  end
end
end;

Figure 4.3 – The factoring transformation

\[ h([A\, B]) := h'(B, A). \]
\[ h([I]). \]
\[ h'(B, x). \]
\[ h'(B, y). \]

Factoring reduces the number of unifications done at run-time in two ways: (1) compound terms are only
created once during predicate execution, instead of being repeated for each clause (e.g. the list \([A, B] \) in
the example), and (2) the arguments of compound terms become predicate arguments, which more often
allows the determinism transformation to convert shallow backtracking into deterministic selection (e.g. the value of the second argument of the predicate \( h' \) determines the correct clause directly without any superfluous unifications). The following heuristic is used:

**Factoring Heuristic:** For each argument in a predicate, factor the largest set of contiguous clauses whose MSG is a compound term. Repeat this operation until no more factoring is possible.

This heuristic needs refinement in some cases to avoid superfluous choice point creation which may slow down execution. The savings of multiple structure creation (how many fewer unifications are done) should be weighed against how much deterministic selection is possible in the dummy predicates.

If the compiler option `same_order_solutions` is enabled (the default) then the operational semantics is that of standard Prolog, i.e. the order of solutions returned on backtracking is identical to that of standard Prolog. Disabling the option relaxes the semantics of standard Prolog by also factoring non-contiguous clauses whose MSG is a compound term. This may change the ordering of solutions on backtracking. This option allows experimentation with variations of standard Prolog semantics.

To illustrate how factoring can reduce the amount of shallow backtracking, consider the following predicate, which is part of a definition of quicksort:

\[
\text{partition}([Y|L], X, [Y|L1], L2) :- Y<X, \text{partition}(L, X, L1, L2).
\]

\[
\text{partition}([Y|L], X, L1, [Y|L2]) :- Y>X, \text{partition}(L, X, L1, L2).
\]

\[
\text{partition}([], _, [], []). 
\]

The first argument of the first two clauses can be factored, resulting in:

\[
\text{partition}([Y|L], X, L1, L2) :- \text{partition'}(L, X, L1, L2, Y).
\]

\[
\text{partition'}([], _, [], []). 
\]

\[
\text{partition'}(L, X, [Y|L1], L2, Y) :- Y<X, \text{partition}(L, X, L1, L2).
\]

\[
\text{partition'}(L, X, L1, [Y|L2], Y) :- Y>X, \text{partition}(L, X, L1, L2).
\]

(In the compound term \([Y|L]\) the rightmost variable \( L \) is kept in the same argument position and the other variable \( Y \) is put at the end of the goal.) The transformation results in only a single unification of \([Y|L]\) instead of two in the original definition. In the dummy predicate the comparisons \( Y<X \) and \( Y>X \) use arguments of the predicate, not arguments of a compound term. This makes it possible to compile `partition/4` with a conditional branch instead of with shallow backtracking.
5. Global dataflow analysis

It is difficult to obtain information about a program by executing it in its original form, since the range of possible behaviors is potentially infinite, and even simple properties of programs may be undecidable. To get around this problem, the idea of abstract interpretation is to transform the program into a simpler form which allows practical analysis. After the analysis the inverse transformation gives information about the original program. The fundamentals of a general method based on this approach and its mathematical underpinning are explained by Kildall [37] and Cousot & Cousot [23]. Marriott and Sondergaard [47] give a lucid explanation of the basic ideas. This method has been studied extensively and developed into a practical tool for Prolog [18, 21, 24, 25, 49, 50, 53, 66, 67, 76, 84].

The four sections that follow summarize the relevant parts of the theory of abstract interpretation, present my application of it to Prolog, describe the analysis algorithm in detail, and discuss the integration of the algorithm into the body of the compiler. In Chapter 7 an evaluation is done of the effectiveness of the algorithm.

5.1. The theory of abstract interpretation

The transformed program should mimic the original faithfully. This is made rigorous by introducing the concept of descriptions of data objects. Let $E$ be the powerset, i.e. the set of all subsets, of a set of data objects, and $D$ be a partially ordered set of descriptions. Then an abstract interpretation is defined by the following conditions:

1. $E_p : E \rightarrow E$, $D_p : D \rightarrow D$
2. $\alpha : E \rightarrow D$, $\gamma : D \rightarrow E$
3. $\alpha$ and $\gamma$ are monotonic.
4. $\forall d \in D : d = \alpha(\gamma(d))$
5. $\forall e \in E : c \leq \gamma(\alpha(e))$
6. $\forall d \in D : E_p(\gamma(d)) \leq \gamma(D_p(d))$
The operator $E_P$ in the first condition describes a single step of the execution of the program $P$ as a state transformation. Symbolic execution of the transformed program is described by the operator $D_P$. Except for the conditions given above, the choice of $E_P$ and $D_P$ is completely free. The choice is guided by several trade-offs, for example: (1) speed versus precision of the analysis, (2) complexity versus confidence in the correctness of the analysis.

As an example of $E_P$ (from Cousot & Cousot [23]), consider a program in an imperative language represented as a graph where each node is a simple statement such as an assignment or a conditional. Let an environment be defined as a correspondence between each variable in the program and a possible value. Then for each edge of the graph a set of possible environments (called a context) is given. Initially they are all unknown. An application of $E_P$ transforms all contexts to their new values reached after one execution step.

For Prolog, a natural choice is to identify $E_P$ with the standard operator $T_P : 2^{B_P} \rightarrow 2^{B_P}$ which describes its procedural semantics. In this case $E$ is $2^{B_P}$, where $B_P$ is the Herbrand universe of the program $P$, i.e. the set of all ground goals† that can be constructed using predicates, functors, and constants of the program. $T_P$ does a single "forward chaining" step to find the conclusions that can be inferred from a given set of ground goals. Formally, $T_P$ maps any $I \subseteq B_P$ into $T_P(I) = \{ A \in B_P : A :- A_1, \cdots, A_n \}$ is a ground instance of a clause in $P$ and $\{ A_1, \cdots, A_n \} \subseteq I$. In other words, an application of $T_P$ transforms a subset of $B_P$ into a new subset containing the new goals inferred from the program's clauses given the old goals. The meaning of a program $P$ is defined as $\text{lfp}(T_P)$ (where $\text{lfp}$ is the least fixpoint operator). This is the set of all ground goals that can be derived from the program clauses. For example, consider the following program:

$$
\text{nat}(0).
\text{nat}(s(X)) :- \text{nat}(X).
$$

which states that $\text{nat}(X)$ is true if $X$ is zero or $X$ is the successor of a natural number. The program's meaning is:

† These are called "atoms" in mathematical logic. To avoid confusion with the atom data type in Prolog, this dissertation uses the Prolog terminology.
\{ \text{nat}(0), \text{nat}(s(0)), \text{nat}(s(s(0))), \text{nat}(s(s(s(0)))), \ldots \}

which represents the set of natural numbers.

The second and third conditions introduce the operators \( \alpha \) (the abstraction function) and \( \gamma \) (the concretization function). The operator \( \alpha : E \rightarrow D \) determines the description corresponding to a particular set of data objects. The operator \( \gamma : D \rightarrow E \) determines the set of data objects corresponding to a particular description.

The fourth and fifth conditions ensure that \( \alpha \) and \( \gamma \) behave correctly with respect to each other. Condition four means that in going from descriptions to data objects and back no information is lost. Condition five means that in going from a data object to a description and back that the resulting set of data objects includes the original data object. The sixth condition is known as the \textit{safeness criterion}. It is necessary to ensure that the symbolic execution (through \( D_P \)) mimics the execution of \( P \) accurately (through \( E_P \)). In other words, the abstract interpretation gives descriptions that include all the data objects that the execution of the original program gives.

To illustrate what the conditions mean consider the abstract domain of signs of real numbers. The data objects are real numbers. Let there be three possible signs for numbers: + (positive), - (negative), and 0 (zero). The set of descriptions \( D \) describes the possible states of a set of reals, so it contains all combinations of the three signs:

\[ D = \{ \{ \}, \{0\}, \{+\}, \{-\}, \{+,-\}, \{-,0\}, \{+,0\}, \{+,-,0\} \} \]

According to the second condition \( \alpha \) maps a set of reals onto its signs, and \( \gamma \) maps a set of signs onto a set of reals. For example:

\[ \alpha([-5]) = (-) \]

\[ \alpha([-3.5]) = (+,-) \]

\[ \gamma([+]) = \{ r \in \mathbb{R}, r > 0 \} \]

The fourth condition says that going from a sign to a set of reals and back will give the same sign. The fifth condition says that going from a set of reals to a sign and back will give a set of reals that includes the
original set. So for example:

\((+) = \alpha(\gamma(\{+\}))\)

since \(\gamma(\{+\})\) = the set of positive reals, whose sign is \(+\), and:

\((5) \subseteq \gamma(\alpha(\{5\}))\)

since \(\alpha(\{5\}) = \{+\}\), and \(\gamma(\{+\})\) is the set of positive reals, which contains 5. In order to explain condition six, consider the equation \(27 \times 37\). Here \(E_P\) is multiplication of reals, and \(D_P\) is the corresponding operation in the abstract domain of signs. The multiplication corresponds to \((+) \times (+)\) in the abstract domain. The result of the abstract multiplication should be \((+)\), since \(27 \times 37 = 999\), which is positive.

Condition six is a formalization of this requirement.

Dataflow analysis is done by transforming the original program over the domain \(E\) described by \(E_P\) to a new version over the domain \(D\) described by \(D_P\). Then \(\gamma(lfp(D_P))\) (lfp = the least fixpoint operator) gives a conservative estimate of the required information. Much work has been done in discovering useful domains \(D\) for particular applications and efficient algorithms for finding fixpoints of \(D_P\) [10, 53].

5.2. A practical application of abstract interpretation to Prolog

The implementation of abstract interpretation presented in this dissertation uses a very different \(E_P\) from the one suggested in the previous section by the formal definition of Prolog's procedural semantics. The choice of \(E_P\) used in the Aquarius compiler closely follows execution on a machine. Consider a program with \(n\) predicates \(P_i\). The data objects are the \(n\)-tuples \((T_1, T_2, \ldots, T_n)\) where each \(T_i\) is a functor of same name and arity as \(P_i\) and the arguments of \(T_i\) are terms constructed using data functors and atomic terms in the program and possibly containing unbound variables. \(E\) is the powerset of these data objects. The descriptions are the \(n\)-tuples \((L_1, L_2, \ldots, L_n)\) where each \(L_i\) is a functor of same name and arity as \(P_i\) and the arguments of \(L_i\) are constrained to be on a given finite lattice. \(D\) is the set of these descriptions. A lattice is a partially ordered set in which every nonempty subset has a least upper bound (denoted as the lub) and a greatest lower bound (denoted as the glb). Each of the elements of the lattice corresponds to a set of possible values in the original program. This lattice is called an argument lattice, since it is used to represent the possible values of a predicate argument. A predicate lattice (such as \(L_+\)) is the Cartesian
product of the lattices of all the predicate’s arguments.

The operator $E_p$ that mirrors execution of the program corresponds to a single resolution step. It is a transformation of a set of data objects and an execution state to another set of data objects and a new execution state, following Prolog’s depth-first execution semantics, that is, its left-to-right execution of goals in a clause, and its top-to-bottom selection of clauses in a predicate. The operator $D_p$ that mirrors execution of the program over the descriptions is similar, except that the arguments are lattice values.

If the conditions of abstract interpretation hold, then the least fixpoint of the symbolic execution over the lattice is a conservative approximation to the global information, in other words the set of values that a variable can have during execution is a subset of what is derived in the analysis.

The three sections that follow describe the lattice used by the analysis algorithm. The first section introduces and defines the lattice elements and the types with which they correspond. The next section gives an example to show how to derive the types. The last section summarizes the properties of the types that are used by the algorithm.

5.2.1. The program lattice

Dataflow analysis for Prolog differs from that of statically typed languages because it does not check types, but it infers them. The most important information that can be deduced about an argument is whether it is used as an input or an output argument of a predicate, i.e. the mode of the argument. After the mode is determined, it is useful to find its type, i.e. the set of values that it can have. The remainder of this chapter refers only to the type of an argument, in the assumption that this implies the mode as well. I have experimented with four lattices of varying complexity in the analyzer, and the lattice that is currently implemented has been chosen to give the most information while keeping analysis fast.

During the analysis the algorithm maintains two lattices for each predicate in the program. These lattices correspond to the entry and exit types of the predicate, i.e. the value of the variable valid upon entering the predicate and upon successful exit from the predicate. The lattice describing the entire program is the Cartesian product of the predicate lattices.
The argument lattice of the entry and exit types in the current analyzer is shown in Figure 4.4. In this lattice, any (the top element) denotes the set of all values, impossible (the bottom element) denotes the empty set (i.e. this predicate is unreachable during execution), uninit denotes the set of uninitialized variables (unbound variables that are not aliased; see Chapter 2), ground denotes the set of values that are ground (i.e. the term contains no unbound variables), nonvar denotes the set of nonvariables, rderef denotes the set of values that are recursively dereferenced (i.e. the term is dereferenced, which means that it is accessible without any pointer chasing, and if it is compound then all its arguments are recursively dereferenced), and ground+rderef denotes the set of values that are both ground and recursively dereferenced.

5.2.2. An example of generating an uninitialized variable type

This section gives a simple example of the generation of uninitialized variable types to give an idea of what abstract interpretation does and to illustrate the argument lattice. Uninitialized variables are generated whenever the analyzer deduces that an unbound variable cannot be aliased to another. For example, consider the following program fragment:
\texttt{pred(...)} := \ldots \texttt{goal(Z)}, \ldots

\texttt{goal(X)} := \texttt{X=s(Y)}, \texttt{goal(Y)}.

If Z is the first occurrence of that variable in the \texttt{pred(...)} clause then it is considered a candidate uninitialized variable. This is possible because it is certainly not aliased to any other variable. In the definition of \texttt{goal(X)}, if X is uninitialized then the argument Y of the structure \texttt{s(Y)} may be considered uninitialized as well. This Y is passed on as an argument to \texttt{goal(Y)}. Therefore both calls of \texttt{goal(X)} are with an uninitialized argument, so it is consistent to give the argument X an uninitialized variable type.

It may happen that elsewhere in the program there is a call of \texttt{goal(X)} where X is not uninitialized (for example it may be a compound term, or it may be aliased). In that case, the assumption that X is uninitialized is invalidated. This may invalidate assumptions about other arguments of other predicates, so it is necessary to propagate this information. For correctness, it is necessary to iterate until the least fixpoint is reached. At that point symbolic execution of the program does not change any of the derived types.

5.2.3. Properties of the lattice elements

The example given above already gives an inkling of the relevant properties of ground, uninitialized, and recursively dereferenced variables that simplify the analysis. Here is a more complete list of these properties:

- The property of being ground, uninitialized, or recursively dereferenced propagates through explicit unifications. The propagation is bidirectional:

  (1) If X is ground, uninitialized, or recursively dereferenced, then after executing an explicit unification with a compound term (e.g. \texttt{X=s(A,B)}), all of its variables (e.g. A and B) are ground, all of the new variables (e.g. A and B) are uninitialized, or all of the new variables are recursively dereferenced.

  (2) In the other direction, if all the variables in the compound term are ground, then X is ground. If all the variables are recursively dereferenced, then X is recursively dereferenced if it was...
previously uninitialized.

- The property of being ground is independent of aliasing. For example, if \( X \) is ground, then it remains ground after executing the unification \( X = Y \). This is not true of recursively dereferenced or uninitialized variables.

- An uninitialized variable is not aliased to any other variable. Lattice calculations for uninitialized variables do not affect each other.

5.3. Implementation of the analysis algorithm

Previous sections have introduced the ideas underlying the algorithm, the program lattice used by the algorithm, an example of how types are derived, and the properties of the lattice elements. This section gives a more complete explanation of the algorithm. The presentation starts with an overview of the data representation. It then describes the algorithm, and finally it gives a detailed example of analysis.

<table>
<thead>
<tr>
<th>Table 4.5 - The components of the variable set VS</th>
</tr>
</thead>
<tbody>
<tr>
<td>Name</td>
</tr>
<tr>
<td>------</td>
</tr>
<tr>
<td>S</td>
</tr>
<tr>
<td>G</td>
</tr>
<tr>
<td>N</td>
</tr>
<tr>
<td>U</td>
</tr>
<tr>
<td>D</td>
</tr>
</tbody>
</table>

5.3.1. Data representation

During analysis the types are represented in two ways:

(1) As lattice elements. For each predicate, there are two structures containing a lattice element in each
argument. These structures represent the entry and exit types of the predicate. For example, the predicate `concat/3` has two structures which could have the values:

```
entry: concat(any, ground, uninit)
exit:  concat(ground, ground, any)
```

This says that upon entering `concat/3` the second argument is ground and the third argument is uninitialized. When the predicate is exited the first two arguments are ground.

(2) As sets of variables. Type information can also be stored as a set for each type that contains the variables of that type.

These two different representations each have their advantages. The lattice representation makes it easy to calculate the lub (least upper bound). The variable set representation makes it easy to symbolically execute a clause, i.e. to propagate and update information about variables' types through the clause. Functions are provided to convert between the two representations (Figure 4.7). For the lattice in Figure 4.4, there are five sets of variables which are updated during the symbolic execution of a predicate. Conceptually they are part of a 5-tuple $VS = (S, G, N, U, D)$ that holds the current type information (Table 4.5).

5.3.2. Evolution of the analyzer

The current analyzer was preceded by three simpler versions. The lattice of the first analyzer represented only entry types and had three elements: impossible, uninit, and any. The second analyzer added the ground type in the entry lattice and an exit lattice of the same structure. The third analyzer added the rderef type to these lattices. The current (fourth) analyzer added the nonvar type. Despite not using a representation for variable aliasing, the third and fourth analyzers are able to derive many nontrivial rderef and nonvar types. The added types are independent, i.e. each version of the analyzer does no better than previous versions on types that previous versions also derive.

The choice of what lattice types to add was done by inspecting the compiled code of programs and by deciding what types were easy to derive in the context of the structure of the existing analyzers. Types were added that are present in many programs. Measurements show that having an exit lattice and doing back propagation (see below) are essential features to derive good ground, rderef, and nonvar
types. A numerical evaluation of the efficiency of the analysis (the percentage of arguments for which
types are derived) and the effect of analysis on execution time and code size is given in Chapter 7.

For the next version of the analyzer the added types \texttt{rlist} (recursive list, i.e. the term is either \texttt{nil}
or a cons cell whose tail is a recursive list), \texttt{integer}, and ((nonvar+deref) or \texttt{uninit}) (the
term is either a dereferenced nonvariable or uninitialized) are contemplated.

```plaintext

\begin{verbatim}

  type varset = (set, set, set, set);  /* 5-tuple */
  var Program : set of predicate;
  \quad L_entry : mapping predicate \rightarrow lattice;
  \quad L_exit : mapping predicate \rightarrow lattice;
  \quad P : predicate;

  procedure analysis:
  \begin{var}
  \quad E : set of predicate;
  \quad VS : varset;
  \begin{begin}
  \quad E := \{ P | \text{arity}(P) = 0 \} \cup \text{(declared entry points)};
  \quad \text{Initialize } L_{entry} \text{ with the types of the declared entry points;}
  \quad \text{Initialize } L_{exit} \text{ to impossible for all predicate arguments;}
  \quad \text{while } E \neq \emptyset \text{ do begin}
  \quad \quad \text{for each predicate } P \in E \text{ do begin}
  \quad \quad \quad VS := \text{lattice_to_varset}(L_{entry}[P], P);
  \quad \quad \quad VS := \text{update_exit}(VS, \text{predicate_analyze}(P, VS), P)
  \quad \quad \quad end;
  \quad \quad E := \{ P | L_{entry}[P] \text{ has changed or } \exists \ G \in P : L_{exit}[G] \text{ has changed} \}
  \quad \text{end;
  \quad end:

Figure 4.5 – The analysis algorithm: top level

\end{verbatim}

\end{verbatim}

5.3.3. The analysis algorithm

The analysis algorithm is presented at three levels of detail. An English-language description is
given of the basic ideas. A detailed pseudocode definition (Figures 4.5 through 4.7) describes the complete
algorithm at a high level of abstraction. Appendix G gives the implementation in the compiler.

The algorithm maintains entry and exit lattice elements for each predicate argument in the program.
Analysis proceeds by traversing the call graph starting from a set of entry points that have known types.
The entry points include all predicates of arity 0 and any entry declarations given by the programmer
(Appendix A). The traversal is repeated until there are no more changes in the lattice values, that is, until a
function predicate_analyze(P : predicate; VS : varset): varset;
var
F : formula;
VS : array [1 .. n] of varset;
G, : goal;
i, j : integer;
begin
/* At this point $P = \{ C_1, ..., C_n \}$ (list of $n$ clauses) */
for each non-active clause $C_i \in P$ do begin /* Symbolic execution of clause $C_i$ */
  /* At this point $C_i = (G_{i1}, ..., G_{in})$ (conjunction of $n$ goals) */
  VS := VS;
  for j := 1 to n do begin /* Symbolic execution of goal $G_{ij}$ */
    if ($G_{ij}$ is a unification) then begin
      VS := symbolic_unify(VS, $G_{ij}$) /* Figure 4.8 */
    end else if $G_{ij} \in Program$ then begin /* $G_{ij}$ is defined in the program */
      $L_{entry}[G_{ij}] := \text{lub}(L_{entry}[G_{ij}], \text{varset_to_lattice}(VS, G_{ij}))$;
      if non-exponentiality constraint then begin
        VS := update_exit(VS, predicate_analyze($G_{ij}, VS_i$), $G_{ij}$)
      end end else begin /* $G_{ij}$ is not defined in the program */
      F := varset_to_type(VS, $G_{ij}$);
      $G_e := \text{entry_specialize}(G_{ij}, F)$;
      VS := update_exit(VS, exit_varset($G_e$, $G_{ij}$)
    end end;
  VS := back_propagate(VS, $C_i$) /* To obtain more precision */
end;
return $\bigcup_{i=1}^{n} VS_i$ /* Merge the exit values of all VS */
end;

Figure 4.6 – The analysis algorithm: analyzing a predicate

fixpoint is reached. With suitable conditions (i.e. all type updating is monotonic and types are propagated correctly) this fixpoint is the least fixpoint and the resulting types give accurate information about the original program. When a goal is encountered during a traversal three things are done: (1) the goal’s entry lattice type is updated using the current value of VS, (2) if the goal’s definition is part of the program then the definition is entered, and (3) upon return, the new value of VS is used to update the goal’s exit lattice type.

A correct value of VS is maintained at all times during the traversal of a goal’s definition.

The definition of the algorithm in Figures 4.5 through 4.7 leaves out some details but is a faithful description of the analysis. The two conditions non-active and non-exponentiality are explained in the next section. The following sections describe what happens in symbolic execution of a predicate (including back propagation) and symbolic execution of a goal.
function update_exit(VS₁, VS₂ : varset; G : goal) : varset;
var VS : varset;
begin
  /* Calculate new VS from old VS₁ and exit VS₂ */
  VS.nonvar := VS₁.nonvar ∪ VS₂.nonvar;
  VS.ground := VS₁.ground ∪ VS₂.ground;
  VS.rdercf := (VS₁.rdercf ∩ VS₁.ground) ∪ VS₂.rdercf;
  VS.sofar := VS₁.sofar ∪ vars(G);
  VS.uninit := VS₁.uninit − vars(G);
  /* Calculate new exit lattice */
  L.exit [G] := lub(L.exit [G], varset_to_lattice(VS, G));
  return VS
end;

function lub(L₁, L₂ : lattice) : lattice;
return (least upper bound of L₁ and L₂);

function varset_to_lattice(VS : varset; G : goal) : lattice;
return (lauice corresponding to VS using variables of G);

function entry_specialize(G : goal; F : formula) : goal;
return (specialized entry point of G when called with type F);

function exit_varset(G : goal) : varset;
return (exit varset stored for the known goal G);

Figure 4.7 – Utility functions needed in the analysis algorithm

5.3.4. Execution time of analysis

This section shows that the average analysis time for programs that contain only linearly recursive predicates (i.e. no clauses contain more than one recursive call) and that have bounded arity is proportional to the size of the program. The analysis time $T_{\text{analysis}}$ is proportional to the time of each iteration $T_{\text{iter}}$ and the number of iterations $N_{\text{iter}}$ needed to reach the least fixpoint:

$$T_{\text{analysis}} = O(T_{\text{iter}} \cdot N_{\text{iter}})$$

For programs that contain only linearly recursive predicates, the time of each iteration is:

$$T_{\text{iter}} = O(S \cdot A)$$
where \( S \) is the total number of goals in the program and \( A \) is the maximum number of times a predicate is traversed. (Programs with non-linearly recursive predicates are discussed below.) This is true because the algorithm traverses each clause at most once in an iteration. It assumes that the symbolic execution of a goal whose definition is not traversed is a constant time operation. A predicate is traversed only if the current entry type is worse than the previous worst entry type. The number of times this situation can occur is bounded by the depth of the entry lattice of the predicate, which is proportional to the maximal arity in the program. Therefore:

\[
S = \sum_{i=1}^{n} \sum_{j=1}^{n} \text{length}(C_{ij})
\]

\[
A = O(\max_{i=1}^{n} \text{arity}(P_i))
\]

where the program contains \( n \) predicates, and each predicate \( P_i \) contains \( n_i \) clauses \( C_{ij} \). The arity of a predicate is denoted by \( \text{arity}(P_i) \) and the number of goals in a clause is denoted by \( \text{length}(C_{ij}) \). The number of iterations is trivially bounded by the depth of the program lattice:

\[
N_{\text{iter}} = O(D_{\text{total}})
\]

where \( D_{\text{total}} \) is given by:

\[
D_{\text{total}} = \sum_{i=1}^{n} 2 \cdot 4 \cdot \text{arity}(P_i)
\]

In this equation, 2 counts the entry and exit lattices, \( \text{arity}(P_i) \) is the number of arguments in the predicate lattice, and 4 is the depth of each argument lattice. This bound on \( N_{\text{iter}} \) is wildly pessimistic. For most real programs \( N_{\text{iter}} \) is bounded by a small constant. All the benchmark programs satisfy \( N_{\text{iter}} \leq 7 \) (Chapter 7). However, there exist pathological predicates \( P_x \) for which \( N_{\text{iter}} = \Theta(\text{arity}(P_x)) \). For example, consider the program:

```
main :- a(9,a,-,-,-,-,-,-,-,-).
a(0,-,-,-,-,-,-,-,-).
a(N,A,C,D,E,F,G,H,I,J) :- N1 is N-1, a(N1,A,C,D,E,F,G,H,I,J).
```

The analyzer requires 10 passes to determine that all arguments of \( a/11 \) are ground and dereferenced upon exit.
To summarize these results, the worst-case and average case total execution times of analysis for programs without non-linearly recursive predicates are:

\[ T_{\text{analysis, worst}} = O(A \cdot S \cdot D_{\text{local}}) \]

\[ T_{\text{analysis, ave}} = O(A \cdot S) \]

If the arity is bounded, then the average execution time of analysis is proportional to the program's size.

For programs that contain non-linearly recursive predicates this result needs to be amended. There is a trade-off between precision and execution time of the analysis. If not enough predicates are traversed then analysis information is lost. If too many predicates are traversed then analysis time becomes too long.

Two constraints are used to prune the traversal of the call graph:

1. The non-active constraint. A clause that is in the process of being traversed is called an active clause. During recursive calls of predicate_analyze, the algorithm maintains a set of the active clauses and will not traverse an active clause twice.

2. The non-exponentiality constraint. Traverse a predicate (i.e. call predicate_analyze) only if one of two conditions hold: (a) The entry type has changed since the last traversal of the predicate, or (b) At least one of the predicate's clauses is active.

Condition (a) is understandable: it is needed to ensure that an updated type is propagated correctly. The rationale for condition (b) is more subtle. If it did not hold, then the exit types derived by the analysis would be significantly worse because the base case of a recursive predicate may not be reached during the traversal. Running the analyzer both with and without this condition shows this to be true for most programs.

The problem with condition (b) is that it leads to an analysis time that is exponential in the number of non-linearly recursive clauses in a predicate. For many programs this is not serious. However, it occurs often enough that it should be solved. One of the benchmark programs, the nand benchmark, has this problem. A better condition is needed to replace condition (b). It must (1) ensure that the base case of all recursive predicates is reached (for good exit types), and (2) not result in time exponential in the number of non-linearly recursive predicates.
5.3.5. Symbolic execution of a predicate

The heart of the dataflow analysis algorithm is the symbolic execution of a predicate (Figure 4.6). Each clause of the predicate is traversed from left to right. During the traversal the type information is kept in the variable set VS. Symbolic execution of the predicate consists of four steps:

1. For each clause of the predicate, translate the lattice entry type of the predicate into the variable set VS, and start traversing the clause.

2. Symbolically execute each goal in the clause and update VS.

3. At the end of each clause, back propagation improves VS by deducing information that only becomes available at the end of the clause. For example, consider the clause:

   a(X) :- X=[Y|L], b(Y, L).

   If both Y and L are in the ground set G of VS at the end of the clause then this is also true of X because of the unification X=[Y|L]. Back propagation is used to improve the exit types for ground, recursive dereference, and nonvariable types. Measurements show that it is a necessary step to get good exit types.

4. At the end of the predicate, combine the variable sets of all clauses by intersecting their corresponding components. Convert the result back to the lattice representation and update the exit type for the predicate.

5.3.6. Symbolic execution of a goal

Symbolic execution of a goal is done in three ways, depending on whether the goal is a unification, the goal is defined in the program, or the goal is not defined in the program.

5.3.6.1. Unification goals

Symbolic execution of unification is defined by the function symbolic_unify(VS, X=T) in Figure 4.8, which converts VS = (S, G, N, U, D) into VS' = (S', G', N', U', D'). These equations use the utility functions of Table 4.6. For each component of VS, any equation in Figure 4.8 with a true condition can be
Table 4.6 - Utility functions of a term $T$

<table>
<thead>
<tr>
<th>Notation</th>
<th>Definition</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\text{vars}(T)$</td>
<td>The set of variables in the term $T$.</td>
</tr>
<tr>
<td>$\text{dups}(T)$</td>
<td>The set of variables that occur at least twice in the term $T$.</td>
</tr>
<tr>
<td>$\text{new}(T) = \text{vars}(T) - S$</td>
<td>The set of all variables in $T$ that have not occurred before.</td>
</tr>
<tr>
<td>$\text{old}(T) = \text{vars}(T) \cap S$</td>
<td>The set of all variables in $T$ that have occurred before.</td>
</tr>
<tr>
<td>$\text{deref}(T) = \text{vars}(T) - (S - U)$</td>
<td>The set of all variables in $T$ that are candidates to be recursively dereferenced. This is the same as $\text{new}(T) \cup (\text{vars}(T) \cap U)$, i.e. $\text{new}(T)$ supplemented with the variables in $T$ that are uninitialized.</td>
</tr>
</tbody>
</table>

\[
S' = S \cup \text{vars}(X=T)
\]

\[
G' = \begin{cases} 
G \cup \text{vars}(T) & \text{if } X \in G \\
G & \text{otherwise}
\end{cases}
\]

\[
N' = \begin{cases} 
N \cup G' \cup \{X\} & \text{if } \text{nonvar}(T) \text{ or } (\text{var}(T) \text{ and } T \in N) \\
N \cup G' & \text{otherwise}
\end{cases}
\]

\[
U' = \begin{cases} 
U \cup \text{new}(T) - \text{old}(T) - \{X\} - \text{dups}(T) & \text{if } (X \in S \text{ or } X \in U) \\
U - \text{vars}(X=T) & \text{otherwise}
\end{cases}
\]

\[
D' = \begin{cases} 
D \cup \text{deref}(T) \cup \{X\} & \text{if } (X \in S \text{ or } X \in U) \text{ and } \text{old}(T) \subset (D \cup U) \\
D \cup \text{deref}(T) & \text{if } (X \in S \text{ or } X \in U) \text{ or } X \in (D \cap G) \\
D \cup \text{deref}(T) - \{X\} & \text{if } \text{dups}(T) = \emptyset \text{ and } X \in D \text{ and } \text{old}(T) \subset U \\
D \cap G & \text{otherwise}
\end{cases}
\]

Figure 4.8 - Symbolic unification $V\Sigma' := \text{symbolic_unify}(V\Sigma, X=T)$

used. In practice, if more than one condition is satisfied, an equation giving more information (i.e. the resulting set is larger) is used first. These equations are listed first. For example, the first equation of $D'$ gives a larger set, so it is preferred over the others. If both $X$ and $T$ are variables, then the algorithm switches $X$ and $T$ is to see if one of the more desirable equations is satisfied before attempting one of the lesser equations.
Table 4.7 - Conditions for the lattice entry type

<table>
<thead>
<tr>
<th>Name</th>
<th>Condition</th>
</tr>
</thead>
<tbody>
<tr>
<td>( C_{\text{ground}} )</td>
<td>( \text{vars}(X) \subseteq G )</td>
</tr>
<tr>
<td>( C_{\text{var}} )</td>
<td>( \text{var}(X) )</td>
</tr>
<tr>
<td>( C_{\text{old}} )</td>
<td>( X \in (\text{dups}(P) \cup S - U) )</td>
</tr>
<tr>
<td>( C_{\text{deref}} )</td>
<td>( (\text{vars}(X) \cap S) \subseteq D )</td>
</tr>
<tr>
<td>( C_{\text{nonvar}} )</td>
<td>( (X \in N) )</td>
</tr>
</tbody>
</table>

Table 4.8 - Calculation of the lattice entry type

<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th>( C_{\text{deref}} )</th>
<th>( C_{\text{nonvar}} )</th>
<th>Latice value</th>
</tr>
</thead>
<tbody>
<tr>
<td>yes</td>
<td>yes</td>
<td>yes</td>
<td>yes</td>
<td>ground+rderef</td>
<td>ground+rderef</td>
</tr>
<tr>
<td>yes</td>
<td>no</td>
<td>yes</td>
<td>no</td>
<td>ground</td>
<td>ground</td>
</tr>
<tr>
<td>no</td>
<td>no</td>
<td>yes</td>
<td>no</td>
<td>nonvar+rderef</td>
<td>nonvar+rderef</td>
</tr>
<tr>
<td>no</td>
<td>yes</td>
<td>no</td>
<td>yes</td>
<td>nonvar</td>
<td>nonvar</td>
</tr>
<tr>
<td>no</td>
<td>yes</td>
<td>yes</td>
<td>yes</td>
<td>nonvar+rderef</td>
<td>nonvar+rderef</td>
</tr>
<tr>
<td>no</td>
<td>yes</td>
<td>yes</td>
<td>no</td>
<td>rderef</td>
<td>rderef</td>
</tr>
<tr>
<td>no</td>
<td>yes</td>
<td>yes</td>
<td>no</td>
<td>nonvar</td>
<td>nonvar</td>
</tr>
<tr>
<td>no</td>
<td>yes</td>
<td>yes</td>
<td>no</td>
<td>any</td>
<td>any</td>
</tr>
</tbody>
</table>

5.3.6.2. Goals defined in the program

Symbolic execution of a goal with a definition is done by symbolically executing the definition. Information is kept about the part of the call graph that has already been traversed, so that analysis will not go into an infinite loop. The function \( \text{varset_to_lattice}(V, P) \) is defined by Tables 4.7 and 4.8. For each argument \( X \) of \( P \), first determine the values of the five conditions in Table 4.7. Then use these conditions to look up the lattice value for the argument in Table 4.8.

5.3.6.3. Goals not defined in the program

Examples of goals that are not defined in the program being analyzed are built-ins and library predicates. Symbolic execution of these goals is done in two parts. First, entry specialization replaces the goal by a faster entry (section 5.4.1). Second, the type declarations that the programmer has given for the entry are used to continue the analysis. If there are none, then worst-case assumptions are made.

5.3.7. An example of analysis

The following program is interesting because it is mutually recursive:
Table 4.9 – Analysis of an example program

<table>
<thead>
<tr>
<th>incl_2(A, B, C)</th>
<th>incl_3(A, B, C, D)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Start</td>
<td></td>
</tr>
<tr>
<td>entry</td>
<td>impossible</td>
</tr>
<tr>
<td>exit</td>
<td>impossible</td>
</tr>
<tr>
<td></td>
<td>impossible</td>
</tr>
<tr>
<td></td>
<td>impossible</td>
</tr>
<tr>
<td>After pass 1</td>
<td></td>
</tr>
<tr>
<td>entry</td>
<td>rderef</td>
</tr>
<tr>
<td></td>
<td>nonvar</td>
</tr>
<tr>
<td></td>
<td>uninit</td>
</tr>
<tr>
<td>exit</td>
<td>rderef</td>
</tr>
<tr>
<td></td>
<td>any</td>
</tr>
<tr>
<td></td>
<td>nonvar</td>
</tr>
<tr>
<td>After pass 2</td>
<td></td>
</tr>
<tr>
<td>entry</td>
<td>rderef</td>
</tr>
<tr>
<td></td>
<td>nonvar</td>
</tr>
<tr>
<td></td>
<td>uninit</td>
</tr>
<tr>
<td></td>
<td>rderef</td>
</tr>
<tr>
<td>exit</td>
<td>rderef</td>
</tr>
<tr>
<td></td>
<td>any</td>
</tr>
<tr>
<td></td>
<td>nonvar</td>
</tr>
<tr>
<td>After pass 3</td>
<td></td>
</tr>
<tr>
<td>entry</td>
<td>rderef</td>
</tr>
<tr>
<td></td>
<td>nonvar</td>
</tr>
<tr>
<td></td>
<td>uninit</td>
</tr>
<tr>
<td></td>
<td>rderef</td>
</tr>
<tr>
<td>exit</td>
<td>rderef</td>
</tr>
<tr>
<td></td>
<td>any</td>
</tr>
<tr>
<td></td>
<td>nonvar</td>
</tr>
</tbody>
</table>

main :- incl_2([A, B], C, D).
incl_2([], C, [C]).
incl_2([A,E], C, D) :- incl_3(C, A, E, D).
incl_3(C, A, E, [A|D]) :- incl_2(E, C, D).

The predicates incl_2/3 and incl_3/4 are extracted from a definition of set inclusion. Three analysis passes are necessary to reach the fixpoint (Table 4.9). The entries that have changed with respect to the previous pass are in italics. The final types are given in Table 4.10. Most of the correct types are determined after the first pass. A single exit type of incl_3/4 is corrected in the second pass. This is necessary because the third argument of incl_3/4 is the same as the first argument of incl_2/3.

Table 4.10 – Final results of analysis

<table>
<thead>
<tr>
<th>incl_2(A, B, C)</th>
<th>incl_3(A, B, C, D)</th>
</tr>
</thead>
<tbody>
<tr>
<td>entry type:</td>
<td>rderef(A), uninit(B), uninit(C)</td>
</tr>
<tr>
<td>exit type:</td>
<td>nonvar(A), rderef(B), nonvar(C)</td>
</tr>
<tr>
<td>entry type:</td>
<td>uninit(A), rderef(B), rderef(C), uninit(D)</td>
</tr>
<tr>
<td>exit type:</td>
<td>rderef(A), nonvar(C), nonvar(D)</td>
</tr>
</tbody>
</table>

5.4. Integrating analysis into the compiler

Deriving type information is only the beginning. The analyzer must be integrated into the compiler to take advantage of the type information. The dataflow analysis module itself does four source transformations (Figure 4.9) before passing the result to the next stage, which does determinism extraction. The
following source transformations are done in the dataflow analysis module:

(1) **Entry specialization.** Determine a fast entry point for each occurrence of a call whose definition is not in the program being analyzed and continue analysis with this entry point.

(2) **Uninitialized register conversion.** Convert uninitialized memory types to uninitialized register types when it results in a speedup. It is done when an argument can be returned in a register without giving up last call optimization.

(3) **Head unraveling.** Unravel the heads of all clauses again in the light of the derived type information.

For example, the head `a(A, A, A)` can be unraveled in three different ways, namely
\[(a(A, B, C) : -A=B, C=B) \] or \[(a(A, B, C) : -A=C, B=C) \] or \[(a(A, B, C) : -B=A, C=A). \] If both \(A\) and \(B\) are nonvariables and \(C\) is unbound, then the first or third possibilities allow the compiler to do argument selection. Unraveling is already done during the conversion to kernel Prolog, but it must be done again after dataflow analysis since the new types may allow it to be done better.

4) Type updating. Supplement the type declarations given by the programmer (if any) by the derived types. All inconsistencies are reported and compilation continues with the corrected types.

The first three of these transformations are discussed in more detail in the following sections.

5.4.1. Entry specialization

During analysis, a fast entry point is determined for each call whose definition is not in the program being analyzed (i.e. each dangling call). For example, the call `sort(A, B)` is replaced by the entry point `\$('sort *2' A, B\')` if \(B\) is uninitialized. Analysis continues with the types of the fast entry point. The program is unchanged until the end of analysis, so the determination of the fast entry point is repeated in each analysis iteration whenever a dangling call is encountered. This mechanism is intended to speed up execution of built-in predicates and library routines, but it is also available to the programmer.

The fast entry point is determined by calculating the type formula corresponding to the variable set \(VS\) with the function `varset_to_type(VS, G)` (Figures 4.6 and 4.7). This type formula is used to traverse the modal entry tree for the goal. The modal entry tree is a data structure that contains a set of entry points and the types that each requires (Appendix A). Entry specialization is also done in the clause compiler, and a detailed example of the use of the modal entry tree is given in Chapter 5 (section 3.4).

5.4.2. Uninitialized register conversion

Often an uninitialized memory type can be converted to an uninitialized register type. The compiler uses four conditions to guide the conversion process. Define a survive goal as one that does not alter any temporary registers (except for arguments with uninitialized register type, which are outputs). A goal that potentially alters temporary registers is a non-survive goal. The compiler maintains a table of survive goals. With these definitions the four conditions for a predicate \(P\) are:
(1) All arguments of $P$ with uninitialized memory type are candidates to be converted.

(2) A candidate argument of $P$ must occur at most once in the body of each clause of $P$. In each clause where it occurs, the argument must be in the last non-survive goal or any survive goal beyond it.

(3) For each clause of $P$, if the last goal $G$ is a non-survive goal, then the candidate argument of $P$ must be in the same argument position in $G$ as in the head of $P$. This is necessary to avoid losing the opportunity for last call optimization (LCO): if the argument positions are different then a move instruction is needed between the last call and the return. If the last goal is a survive goal then the condition is unnecessary because it is not as important to retain LCO: a survive goal can never be mutually recursive with the predicate it is part of.

(4) Often the last goal $G$ has candidate arguments that are not candidate arguments of $P$, so they have to be initialized when returning from $G$. This has two disadvantages: $P$ loses LCO and $P$ must allocate an environment (which may not exist otherwise). The solution to this problem involves a trade-off: is it better to have LCO in $P$ and fewer uninitialized register arguments in $G$, or to have no LCO in $P$ and more uninitialized register arguments in $G$? The compiler recognizes a class of predicates $G$ for which the first is true: Define a fast predicate as one whose definition contains only built-ins and survive goals. If $G$ is fast then reduce the set of $G$'s candidate arguments to include only those that are candidate arguments of $P$.

A transitive closure is done until all four conditions are satisfied. These conditions can be relaxed slightly in several ways. However, even with the existing conditions it is possible to convert about one third of all uninitialized types into uninitialized register types (Chapter 7). The third and fourth conditions are not needed for correctness, but only for execution speed. The third condition ensures that LCO is not lost. The fourth condition speeds up the chat_parser benchmark by 1% and was added after code inspection discovered cases where the use of uninitialized registers slows down execution.

5.4.3. Head unraveling

This transformation repeats the head unraveling transformation (Chapter 3) with the information gained from dataflow analysis. This increases the opportunities for determinism extraction. For example,
before analysis the clause:

\[
a(X, X, X).
\]

is transformed to the following kernel Prolog by making the head unifications explicit (i.e. "unraveling" the head unifications):

\[
a(X, Y, Z) :- X = Y, \ X = Z.
\]

If analysis derives that \(X\) is unbound and both \(Y\) and \(Z\) are nonvariable, then the above expansion hides the determinism by twice unifying an unbound variable with a nonvariable. Unraveling the head unifications again after analysis results in:

\[
a(X, Y, Z) :- Y = Z, \ X = Y.
\]

In this version, the nonvariables \(Y\) and \(Z\) are unified together, better exposing the deterministic check that is done, and the unbound variable \(X\) is only unified once.

6. Determinism transformation

This section groups four transformations that expose the determinism inherent in a predicate. The purpose of the first three transformations is to make the determinism in the predicate easily visible, so that the fourth transformation, determinism extraction, is as successful as possible in generating case statements. The following transformations are done in order:

(1) **Head-body segmentation.** By separating the heads of clauses from the clause bodies, this reduces the code expansion caused by type enrichment and determinism extraction.

(2) **Type enrichment.** This adds types to predicates for which global analysis is not able to determine the type. The compiler creates different versions of the predicate assuming different input types. This increases code size, but improves performance since often a predicate is deterministic at runtime even though this could not be detected at compile-time.

(3) **Goal reordering.** This reorders goals in a clause to expose more determinism. Tests (such as arithmetic relations) are moved to the left and predicates guaranteed to succeed (such as unifications with uninitialized variables) are moved to the right.
(4) Determinism extraction with test sets. This transformation converts the predicate into a nested case statement that makes its determinism explicit, so that a straightforward compilation to BAM code is possible.

6.1. Head-body segmentation

This transformation reduces the code expansion resulting from enrichment and determinism extraction. A predicate is split into a new predicate and a set of clause bodies. The new predicate contains only the goals of the original predicate that are useful for determinism extraction, i.e. all explicit unifications and tests (including type checking and arithmetic comparisons, see Table 4.11) in each clause starting from the head up to the first goal that is not in this category. The rest of the clause bodies are separated from the predicate. This is done to avoid code duplication in determinism extraction, since the same clause may occur in several leaves of the decision tree.

For example, the predicate:

\[
p(A, B) :-
\begin{array}{l}
\{ \text{var}(A), p(A), q(A, C), t(C, D), u(D, B) \\
; \ A \neq B, \ r(A), \ s(A) \}
\end{array}
\]

is transformed into:

\[
p(A, B) :-
\begin{array}{l}
\{ \text{var}(A), \ 'Sd1'(A, B) \\
; \ A \neq B, \ 'Sd2'(A) \}
\end{array}
\]

\[ 'Sd1'(A, B) :- p(A), q(A, C), t(C, D), u(D, B). \]

\[ 'Sd2'(A) :- r(A), s(A). \]

The new predicate consists only of those parts of the original predicate that are useful for extracting determinism. The determinism extraction is free to create a decision tree from the new predicate without worrying about duplicating the clause bodies at the leaves of the tree. The separated clause bodies are compiled once only, and the BAM transformation stage (Chapter 6) merges them with the decision tree, thus creating a decision graph.
The decision exactly where to split the clause bodies depends on several factors. All goals in the body are classified into two kinds: goals that are useful for extracting determinism (called "tests"), and other goals. Then the split follows these rules: (1) Only those tests all of whose variables are in the head become part of the new predicate. (2) If the length of the clause body is less than a given threshold, then all of it becomes part of the new predicate.

Head-body segmentation interacts with type propagation. It often occurs that a clause body is called from several leaves in the decision tree with different types. In that case, it is compiled with a type that is the intersection of the types of the entry points. A complication arises when one of the leaves considers a variable to be uninitialized, and another leaf does not. In that case, the first leaf jumps to a piece of code to initialize the variable, and only afterwards jumps to the clause body.

6.2. Type enrichment

By looking at the type or the value of one or more arguments it is possible to reduce the set of clauses that have to be tried. Often the dataflow analysis is able to derive sufficiently strong types so that a good selection can be done, i.e. a deterministic predicate can be compiled efficiently. However, if the types given for the predicate are weak then a source transformation is done to enrich them. The enrichment consists of adding a test to check at run-time whether an argument is a variable or a nonvariable, and to branch to different copies of the predicate in each case.

The number of arguments that are enriched is given by the argument \( S \) of the compiler option `select_limit(S)`. Define a good predicate argument as one that is an argument of a unification not known to succeed always, i.e. in the unification neither argument is known to be unbound. An argument is known to be of a given type if the type is implied by the type formula. Whether or not enrichment is done is based on the following heuristic:

**Enrichment Heuristic 1:** If the number of good arguments known to be nonvariable is less than the selection limit \( S \), then choose the lowest numbered good argument that is not known to be nonvariable. Otherwise choose only the first argument, if it is a good argument and it is not known to be nonvariable.

This heuristic is applied recursively on enriched predicates. The default selection limit is always \( S=1 \).

This default is justified given that (1) a selection limit \( S=1 \) already generalizes the first argument selection
of the WAM, and (2) compilation time and object code size increase rapidly with the selection limit. Even with \( S = 1 \), the source transformation occasionally results in some duplicate code being generated. This is removed by the BAM transformation stage. When \( S = 1 \) the heuristic is simpler.

**Enrichment Heuristic 2:** If there exist no good arguments known to be nonvariable, then choose the lowest numbered good argument that is not known to be nonvariable. Otherwise choose the first argument, if it is a good argument and it is not known to be nonvariable.

This heuristic generalizes the first-argument selection of the WAM, i.e., it always does at least a first-argument selection, but depending on the types that the predicate has (often derived from dataflow analysis) and the predicate itself (what kinds of head unifications it does), the amount of selection can be vastly greater.

The heuristic may seem complex, but it is a natural way to make a predicate deterministic.

To show how enrichment works, consider the following predicate without type declarations:

\[
\begin{align*}
a(a) \\
a(b)
\end{align*}
\]

It is transformed into:

\[
\begin{align*}
a(A) & :- \text{var}(A), a_v(A). \ % \text{If } A \text{ is unbound.} \\
a(A) & :- \text{nonvar}(A), a_n(A). \ % \text{If } A \text{ is nonvariable.} \\
a_v(a) & \ \\
a_v(b) & \\
a_n(a) & \\
a_n(b)
\end{align*}
\]

The predicate \( a/1 \) has been enriched with an unbound type (in \( a_v/1 \)) and with a nonvariable type (in \( a_n/1 \)). As another example, consider the definition without any type declarations:

\[
\begin{align*}
\text{member}(X, [X|_1]). \\
\text{member}(X, [_|_1]).
\end{align*}
\]

In this case the heuristic picks the second argument, since the first one does no useful unifications. After enrichment, the predicate becomes:

\[
\begin{align*}
\text{member}(X, L) & :- \text{var}(L), \text{member}_v(X, L). \\
\text{member}(X, L) & :- \text{nonvar}(L), \text{member}_n(X, L). \\
\text{member}_v(...) & :- \text{(same as original definition)} \\
\text{member}_n(...) & :- \text{(same as original definition)}
\end{align*}
\]

The two tests \( \text{var}(L) \) and \( \text{nonvar}(L) \) determine which of the two dummy predicates to execute, \( \text{member}_v/2 \) or \( \text{member}_n/2 \), and are compiled into a single conditional branch. This is a
consequence of the fact that the two tests are mutually exclusive, i.e., if one succeeds then the other fails and vice versa. Both \texttt{member_v/2} and \texttt{member_n/2} have the same definition as the original predicate, but they have different types for the second argument. The predicate \texttt{member_v/2} is compiled assuming the second argument is a variable. The predicate \texttt{member_n/2} is compiled assuming the second argument is a nonvariable. Both \texttt{member_v/2} and \texttt{member_n/2} are also targets of the factoring transformation (section 4).

Type enrichment can introduce a significant increase in code size if it is not handled carefully. In practice, the code size is kept small because: (1) the added types result in significantly smaller code for clause selection in each of the two dummy predicates, (2) before doing enrichment, head-body segmentation separates clause heads from the bodies, so that long clause bodies are not duplicated, and (3) the BAM transformation stage (Chapter 6) removes any remaining duplicate code. In a sense, the definitions are first “loosened up” by head-body segmentation and type enrichment to allow more optimization, and then later “tightened up.”

6.3. Goal reordering

This transformation reorders goals in a clause to increase determinism and to reduce the number of superfluous unifications that are done. Goals that are useful in determinism extraction are put as early as possible, and goals that are certain to succeed (such as unifications with uninitialized variables) are put later.

The goals in a clause are classified in four categories: tests (Table 4.11), unifications with unbound variables, unifications with uninitialized variables, and other goals. The goals are reordered so that tests are first (for deterministic selection), followed by unifications with unbound variables (may be affected by aliasing), unifications with uninitialized variables (unaffected by aliasing, so they can safely be put last), and the other goals. The reordering takes into account the fact that unification is commutative, i.e., that unification goals can be permuted in any way without changing the semantics. Some reorderings are better than others because aliasing can worsen the type formula, e.g., if \( X \) is unbound (\texttt{var(X)}) then after performing the unification \( Y = Z \) it may not be unbound any more, if it is aliased to \( Y \) or \( Z \). The reordering is
constrained so that aliasing does not change the operational semantics.

For example, consider a predicate that has an uninitialized argument:

\[
:a(A, B, C) :- \text{uninit}(C).
\]

\[
a(X, Y, Z) :- Z=X, X<Y, \ldots
\]

The transformation knows that the unification \( Z=X \) does not instantiate \( X \) or \( L \) because \( Z \) is unbound and unaliased. Therefore the unification is moved back:

\[
a(X, Y, Z) :- X<Y, Z=X, \ldots
\]

This has two advantages: (1) the test \( X<Y \) is brought forward so that it can be used by determinism extraction, and (2) the unification \( Z=X \) is not done if the test \( X<Y \) fails.

This transformation compensates for the popular programming style which puts all unifications in the head and all tests in the body, e.g. people prefer to write:

\[
a([X|L], [X|\lambda]) :- \text{var}(X), \ldots
\]

instead of:

\[
a([X|L], Z) :- \text{var}(X), Z=[X|\lambda], \ldots
\]

The first version does not imply anything about the instantiation pattern of the arguments, whereas the transformed version does.

6.4. Determinism extraction with test sets

The majority of predicates written by human programmers are intended to be executed in a deterministic way. These predicates are in effect case statements, yet they are too often compiled in an inefficient manner, by means of shallow backtracking (i.e. saving the machine state, unification with the clause heads, and repeated failure and state restoration). This section describes the general technique used in the compiler to convert shallow backtracking into conditional branching.
6.4.1. Definitions

Predicates are compiled into code which is as deterministic as possible through the concept of the test set. Two definitions are useful:

Definition ST: A goal $G$ is a simple test with respect to the kernel Prolog predicate $P$ and the type formula $F$ if it satisfies the following conditions:

- $G$ uses only variables that occur in the head of $P$.
- The implementation of $G$ does not change any state in the execution model, i.e. $G$ does not cause side-effects (I/O or database operations), $G$ does not create choice points, and $G$ does not bind any variables.
- $G$ does not always succeed.

Definition TS: A set of goals is a test set with respect to the kernel Prolog predicate $P$ and the type formula $F$ if it satisfies the following conditions:

- Each goal in the set is a simple test according to definition ST.
- With a given set of variable values, at most one goal in the set can succeed.
- A multi-way branch in which each destination corresponds to the success of one of the goals in the set can be implemented in the target architecture.

The tests in the set need not actually be present in the definition of $P$. Whether or not a given set of goals is a test set depends on the architecture and the predicate $P$. 

---

**Figure 4.10** - Some examples of test sets

Test set

- branch if less than $A < B$, $A \geq B$
- four-way branch on type $\{\text{var}(A), \text{atomic}(A), \text{cons}(A), \text{structure}(A)\}$
6.4.2. Some examples

Most conditional branches in an architecture correspond to a test set. For example, a branch-if-less-than instruction corresponds to the test set \((A < B, A \geq B)\). More complex conditions such as an n-way branch implemented by hashing can also be represented as test sets. Figure 4.10 shows some examples of test sets. The second and third examples correspond to WAM instructions.

To illustrate the use of test sets, consider the predicate:

\[
\text{max}(A, B, C) :\begin{align*}
\text{if } A & > B \text{ then } C = A \\
\text{else if } A & < B \text{ then } C = B \\
\text{else } & (C = B \text{ or } C = A)
\end{align*}
\]

which is one way to calculate the maximum of \(A\) and \(B\). It is compiled as:

\[
\text{max}(A, B, C) :\begin{align*}
\text{if } A & > B \text{ then } C = A \\
\text{else if } A & < B \text{ then } C = B \\
\text{else } & (C = B \text{ or } C = A)
\end{align*}
\]

(The Prolog notation is simplified for readability.) The predicate is executed completely deterministically if \(A > B\) or \(A < B\); a choice point is created only when \(A = B\). The choice point maintains the operational semantics: since both clauses of the original predicate succeed when \(A = B\), there are two identical solutions.

---

type testset = testset(testset_name, testset_ident, set of goal);

function determinism(D : disjunction; H : goal; F : formula; Previous : set of testset) : disjunction;
var TS : testset;
TS_set : set of testset;
begin
if length(D) \leq 1 then return D;
TS_set := find_testset(D, H, F, Previous);
if TS_set = \(\emptyset\) then return D;
TS := pick_testset(TS_set);
return code_testset(TS, D, H, F, Previous)
end;

Figure 4.11 - The determinism extraction algorithm
function find_testsets(D : disjunction, H : goal; F : formula; Previous : set of testset) : set of testset:
var
  TS : testset;
  TS_set : set of testset;
  i, j : integer;
begin
  "At this point D = (C_1, ..., C_n) where D has n choices",
  TS_set := 0;
  for i := 1 to n do begin
    "C_i = (G_{ij}, ..., G_{ik}) where C_i has n_i goals",
    for j := 1 to n_i do
      if G_{ij} = "!" then exit inner loop
      else for all testsets TS from table do begin
        "TS = testset(Name, Ident, Tests) from Table 4.11",
        if TS \in Previous and \text{vars}(G_{ij}) \subseteq \text{vars}(H) and \text{bindset}(G_{ij}, F) = 0 then
          if \exists T \in Tests : (G_{ij} \text{ implies } T \text{ and not}(F \text{ implies } T)) then
            TS_set := TS_set \cup \{TS\}
        end
      end
  end
  return TS_set;
end;

Figure 4.12 – Finding all test sets in a predicate

function pick_testset(TS_set : set of testset) : testset:
var
  TS : testset;
begin
  pick TS in TS_set such that
  \forall U in TS_set : \text{goodness}(TS) \geq \text{goodness}(U); /* From Equation (G) */
  return TS
end;

Figure 4.13 – Picking the best test set

6.4.3. The algorithm

Given a predicate, the compiler proceeds by first finding all test sets that contain tests that are implied by goals in the predicate. This depends on the type formula that is known for the predicate; for example, the unification X=a is only a test if X is nonvariable, i.e. if the type formula implies nonvar(X). Then a "goodness" measure is calculated for each test set, and the test set with the largest goodness is used first. The goodness measure is calculated heuristically; in the current implementation each test set is weighted by an architecture-dependent goodness (which depends on how efficiently it is
function code_testset(TS : testset; D : disjunction; H : goal; F : formula; Previous : set of testset) : disjunction;
var T : goal;
    Choices : disjunction;
begin
    Choices := [ ];
    "At this point TS = testset(Name, Ident, Tests) */
    for all T ∈ Tests do begin
        D_test := subsume(T, D);
        D_test := determinism(D_test, H, update_formula(T, F), Previous ⊔ {TS});
        append 'Test' (T, D_test) to Choices;
        D := subsume(not(T), D)
    end;
    D := determinism(D, H, F, Previous ⊔ {TS});
    append 'Selse' (D) to Choices;
    return 'Sease' (Name, Ident, Choices)
end;

Figure 4.14 – Converting a disjunction into a case statement

implemented in the architecture) and by the number of possible outcomes (e.g. hashing with a large number
of cases is considered better than a two-way branch). The predicate is converted into a case statement
using the best test set. The algorithm is called recursively for each arm of the case statement to build a
decision tree. This tree is collapsed into a graph by the BAM transformation stage.

Figures 4.11 through 4.14 give a pseudocode definition of this algorithm. The figures define the
function determinism(D, H, F, Previous) that performs the determinism extraction. Given a predicate
written as a head H and a disjunction D, along with the type formula F that is true for that predicate, the
function finds as many test sets as possible in the disjunction and converts them into case statements. It
returns a new disjunction that contains these case statements. The parameter Previous is used to avoid
infinite recursion. It contains all test sets that have already been used to make sure each test set is only
used once.

The function find_testsets(D, H, F, Previous) returns a list of all test sets in the disjunction (Figure
4.12). It picks a test set if there is a goal in the predicate which implies a test in the test set. It limits the
goals to those that do not bind any variables (bindsel(GH, F) = Ø) and those that use only variables that
occur in the head (var(GH) ⊂ var(H)). The function pick_testset(TS_sel) returns the test set with the
greatest measure of goodness, as given by Equation (G) (Figure 4.13). The function code_testset(TS, D,
II. F. Previous) converts the disjunction \( D \) into a case statement when given a test set \( TS \) (Figure 4.14).

It uses the functions \( \text{subsume}(F, F_1) \) and \( \text{update}_\text{formula}(F, F_1) \), which are defined in section 3.

### Table 4.11 — Test sets

<table>
<thead>
<tr>
<th>Name</th>
<th>Example Test</th>
<th>Example BAM translation</th>
</tr>
</thead>
<tbody>
<tr>
<td>equal</td>
<td>( X=Y ) (X or Y is simple at run-time)</td>
<td>( \text{equal}(X,Y,Lbl) )</td>
</tr>
<tr>
<td>equal(atomic,A) (A is an atom)</td>
<td>( X=A )</td>
<td>( \text{equal}(X,A,Lbl) )</td>
</tr>
<tr>
<td>equal(structure,F/N) (F/N is name/arity)</td>
<td>( 'Sname_arity' (X,F,N) )</td>
<td>( \text{equal}([X],F/N,Lbl) )</td>
</tr>
<tr>
<td>hash(atomic)</td>
<td>( X=A ) (A is atomic)</td>
<td>( \text{hash}(\text{atom},X,N,Lbl) )</td>
</tr>
<tr>
<td>hash(structure)</td>
<td>( X=S ) (S is a structure)</td>
<td>( \text{hash}(\text{structure},X,N,Lbl) )</td>
</tr>
<tr>
<td>comparison(Class,Kind) (Class ( \in ) {eq, lst, gts}) (Kind ( \in ) {arith, unify, stand})</td>
<td>( X&lt;Y )</td>
<td>( \text{jump}(\text{lts},X,Y,Lbl) )</td>
</tr>
<tr>
<td>Type (Type ( \in ) AllTypes)</td>
<td>( \text{var}(X) )</td>
<td>( \text{test}(\text{eq},\text{tvar},X,Lbl) )</td>
</tr>
<tr>
<td>switch(Type) (Type ( \in ) TagTypes - {var})</td>
<td>( \text{atom}(X) )</td>
<td>( \text{switch}(\text{atom},X,L1,L2,L3) )</td>
</tr>
</tbody>
</table>

Table 4.11 lists the test sets currently recognized by the compiler. This includes unification goals, all type checking predicates, and all arithmetic comparisons. For each test set it gives the name, a representative test in the test set (only one is given, although usually there are several others), and the translation of that test into a conditional branch of the BAM instruction set. For the test sets hash(atomic) and hash(structure) the BAM code includes a hash table (not shown) in addition to the hash instruction. The following definitions simplify the table:

- \( \text{TagTypes} = \{\text{var}, \text{atom}, \text{structure}, \text{cons}, \text{negative}, \text{nonnegative}, \text{float}\} \), i.e. all types that correspond to one tag in the VLSI-BAM architecture.
- \( \text{AllTypes} = \text{TagTypes} \cup \{\text{atomic}, \text{integer}, \text{simple}, \text{compound}\} \), i.e. it includes types that correspond to more than one tag.

The goodness measure for a test set in a predicate is calculated using the following rule:

\[
\text{Goodness} = 1000 \cdot D + G
\]  

(G)

where \( D \) is the number of directions of the test set that occur in the predicate and \( G \) is the raw goodness measure of the test set. This rule ensures that the number of useful directions in the test set is most important. The raw goodness is used only when the number of directions is the same. Table 4.12 gives the raw goodness of all test sets in the VLSI-BAM architecture [34], with a brief justification of the ranking. The
Table 4.12 – Raw goodness measure of test sets in the VLSI-BAM

<table>
<thead>
<tr>
<th>Test set</th>
<th>Rank</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>switch(cons)</td>
<td>131</td>
<td>Switch is best because it is fast and it is a three-way branch, so it gives the most information. Switch of compound terms is better than other switches because it makes traversing a recursive term (like a list or a tree) fast.</td>
</tr>
<tr>
<td>switch(structure)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>switch(negative)</td>
<td>130</td>
<td>Switch of atomic terms is worse because it penalizes the case of traversing a recursive term.</td>
</tr>
<tr>
<td>switch(nonnegative)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>switch(atom)</td>
<td>129</td>
<td>Switch of integer is worse because the VLSI-BAM has separate negative and nonnegative (t.pos and t.neg) tags, requiring two branches.</td>
</tr>
<tr>
<td>var</td>
<td>120</td>
<td>These test sets are types that correspond directly to tags, and there exist fast two-way branches on tags.</td>
</tr>
<tr>
<td>atom</td>
<td></td>
<td></td>
</tr>
<tr>
<td>cons</td>
<td></td>
<td></td>
</tr>
<tr>
<td>structure</td>
<td></td>
<td></td>
</tr>
<tr>
<td>negative</td>
<td></td>
<td></td>
</tr>
<tr>
<td>nonnegative</td>
<td></td>
<td></td>
</tr>
<tr>
<td>equal</td>
<td>85</td>
<td>This test set requires two instructions—a compare and branch, and also possibly loading its arguments into registers.</td>
</tr>
<tr>
<td>equal(atomic,_)</td>
<td>80</td>
<td>These test sets each require two instructions—a compare and branch.</td>
</tr>
<tr>
<td>comparison(...)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>integer</td>
<td>79</td>
<td>These test sets are types that each correspond to two tags, so they need two tag checks.</td>
</tr>
<tr>
<td>atomic</td>
<td></td>
<td></td>
</tr>
<tr>
<td>compound</td>
<td></td>
<td></td>
</tr>
<tr>
<td>equal(structure,_)</td>
<td>60</td>
<td>Equality comparison of a structure's functor &amp; arity needs a memory reference.</td>
</tr>
<tr>
<td>simple</td>
<td>50</td>
<td>This test set corresponds to a type that needs five tag checks (four without floating point).</td>
</tr>
<tr>
<td>hash(atomic)</td>
<td>41</td>
<td>Hashing is the slowest because it needs to calculate the hash address.</td>
</tr>
<tr>
<td>hash(structure)</td>
<td>40</td>
<td>Hashing on a structure is slightly slower than hashing on an atomic term because a memory load is needed to access the main functor of the structure, whereas the atomic term is directly available in the register.</td>
</tr>
</tbody>
</table>

The value of the rank is not important; only the relative order is important. Architectures rank the test sets according to how efficiently they are implemented in the architecture. To compile for a different architecture, only the ranking is changed in the compiler. The ranking is modified for other processors by a compiler option. For example, for the MIPS processor, the option mips changes the ranking to make the test set equal(atomic, [1]) best, i.e. a comparison with the atom [1] (nil), because it can be implemented with a single-cycle conditional branch instruction. The MIPS does not have separate tags for negative and nonnegative integers, so the test sets negative and nonnegative are not implemented as efficiently as on the VLSI-BAM. These two test sets have lower ranks.
Chapter 5
Compiling Kernel Prolog to BAM Code

1. Introduction

The previous chapters described the conversion of standard Prolog to kernel Prolog and the optimizing kernel transformations. This chapter shows how the optimized kernel Prolog is compiled to BAM code. The compilation to BAM is performed in two steps for each predicate. In the first step, the control instructions that make up the framework of the predicate are compiled by the predicate compiler. This includes compiling the deterministic case statements into conditional branches and the disjunctions with choice point instructions.

In the second step, the clauses that make up the body of the predicate are compiled by the clause compiler. The clause compiler uses two primitives, the goal compiler and the unification compiler, to compile goals and explicit unifications. The clause compiler also does register allocation, entry specialization (replacing built-in predicates by faster entry points), and performs the write-once transformation (for fast trailing), and the dereference chain transformation (to maintain consistency with the dataflow analysis). These transformations are explained in detail in the sections below.

2. The predicate compiler

In the kernel transformation stage (Chapter 4), determinism extraction attempts to convert each predicate into a series of nested case statements. This is not always successful; sometimes the case statements still retain disjunctions (OR choices) that could not be converted into deterministic code. The predicate compiler compiles both the case statements and the disjunctions into BAM code. The case statements are compiled into conditional branches. The disjunctions are compiled into choice point instructions. The predicate compiler uses two primitives, the determinism compiler and the disjunction compiler, to compile the predicate's case statements and disjunctions.
2.1. The determinism compiler

Compiling a kernel Prolog predicate into deterministic BAM code is done in two steps. First, the determinism transformation (a kernel Prolog transformation, Chapter 4) converts a kernel Prolog predicate into a series of nested case statements. Then the determinism compiler compiles the nested case statements into BAM code. A case statement may contain any test set, and each test set is mapped to a conditional branch. The test sets and their corresponding conditional branches are given in Table 4.11.

2.2. The disjunction compiler

A disjunction (an OR formula) is a list of clauses that encapsulates a choice. The first clause is executed the first time the disjunction is encountered. The remaining clauses are executed in order on backtracking—each time backtracking returns to the disjunction the next clause is tried. This is implemented by code which generates choice points. A choice point encapsulates the state of the abstract machine at the time it is created. Backtracking restores machine state from a choice point to let execution continue from the point at which the choice point was created.

Creating and restoring machine state in choice points is time-consuming. To minimize the size of the choice points (and hence the time required to create them), the choice point management instructions in the BAM are streamlined to perform the least amount of data movement. They save only those registers that are needed in the clauses of the disjunction after the first, and for each clause of the disjunction they restore only those registers that are needed in that clause. Argument registers are restored in the clause itself and not in the fail instruction. Therefore the size of the choice point does not have to be stored in the choice point and decoded in the fail instruction. A disadvantage is a slightly larger code size.† Consider the following kernel Prolog for a predicate P with n clauses:

\[
\text{Head} := (\ C_1 ; C_2 ; \ldots ; C_n ; \text{fail}).
\]

A single choice point is created for each invocation of P. The set of registers saved in the choice point is the set of all head arguments that are used in clauses after the first, i.e. C_2 through C_n. Arguments that

† This is less of a problem in the VLSI-BAM since the instruction reorderer merges pairs of single-word loads into double-word loads.
occur only in clause \( C_1 \) do not have to be stored in the choice point. The set of registers that is restored for each clause is the set of arguments used in that clause.

Before creating the choice point, the compiler dereferences those arguments that it can deduce will be dereferenced later. This avoids dereferencing the same argument more than once. The set of arguments to be dereferenced is derived by checking the type formula corresponding to each goal in the body of the predicate's definition, and noting whether its arguments have to be dereferenced. For example, arithmetic operations and relational tests are goals that require their arguments to be dereferenced.

To illustrate the compilation scheme, consider the following predicate:

\[
p(A, B, C, D) := \{ \begin{align*}
& a(A), \\
& c(C), \\
& d(D), \\
& \text{fail}
\end{align*}
\]

It is compiled as:

```
procedure(p/4).
    choice(1/3, [2, 3], l(p/4, 2)). ; Save registers r(2) and r(3).
    jump(a/1).
label(1(p/4, 2)).
    choice(2/3, [2, no], l(p/4, 3)). ; Restore only register r(2).
    move(r(2), r(0)).
    jump(c/1).
label(1(p/4, 3)).
    choice(3/3, [no, 3], fail). ; Restore only register r(3).
    move(r(3), r(0)).
    jump(d/1).
```

The choice instructions do all the choice point manipulation: `choice(1/3,...)` creates the choice point, `choice(2/3,...)` modifies the address to return to on backtracking, and `choice(3/3,...)` removes the choice point. Register \( r(0) \) is not saved in the choice point because it is not needed in clauses beyond the first. The second and third clauses restore only the registers they need. Register \( r(1) \) is not saved because it is not needed at all.

Each choice instruction contains a list of the registers that it used. The length of the list is the same for all choice instructions in a predicate. For choices after the first, the atom `no` is put in the positions of registers that do not have to be restored. For example, the list \( \{0, \text{no}, 5\} \) means that registers \( r(0) \) and \( r(5) \) are restored from the first and third locations in the choice point, and the second location is not
accessed.

In this example a further optimization can be done by merging the move instructions with the choice instructions, i.e.:

\[
\begin{align*}
\text{choice}(3/3, &\text{[no,3]}, \text{fail}). \\
\text{move}(r(3), r(0)).
\end{align*}
\]

becomes:

\[
\text{choice}(3/3, \text{[no,0]}, \text{fail}).
\]

This is possible because the value loaded in a register is determined by its position in the list, not by its number, and because register \( r(3) \) is only used to load \( r(0) \).
3. The clause compiler

The clause compiler converts a clause from kernel Prolog form (with type annotations) to BAM code. The structure of the clause compiler is given in Figure 5.1. After compiling the goals in the body there are two intermediate results: (1) BAM code in which variables have not yet been allocated to registers (skeleton code) and (2) a variable occurrence list (the varlist), that contains all unallocated variables in the skeleton code. The final BAM code is obtained by passing the varlist to the register allocator.

Each goal in the clause body is compiled in four steps. First, three transformations are performed on the goal: entry specialization, the write-once transformation, and the dereference chain transformation. Then the goal is compiled into BAM code by one of two routines, the unification compiler or the goal compiler, depending on whether the goal is a unification or not.

These are the important blocks in the clause compiler:

(1) The goal compiler. Its main task is to handle argument passing. Because of the interaction between the different kinds of unbound variables, initialized and uninitialized, this results in a case analysis. In addition, the goal compiler compiles in-line some built-in predicates and the dummy predicates that were created in the transformation to kernel Prolog.

(2) The unification compiler. Its task is, given a type, to compile an explicit unification into the simplest possible code.

(3) The register allocator. Its task is to allocate variables to registers in such a way that the number of superfluous move instructions is minimized. It uses a data structure called the varlist which is generated by the clause body compiler.

(4) Entry specialization. This attempts to replace each goal in the clause by a faster entry point, depending on the types known at the call.

(5) Write-once transformation. This transformation is part of a technique for reducing the overhead of trailing.

(6) Dereference chain transformation. This transformation is necessary to keep the dataflow analysis and the clause compiler consistent.
The following sections give more details about these each of these blocks. First, an example of a clause compilation is given, with emphasis on the skeleton code, the varlist, and a specification of the register allocator. This is followed by discussions of the goal compiler, the unification compiler, entry specialization, the write-once transformation, and the dereference chain transformation.

3.1. Overview of clause compilation and register allocation

This section gives an example of how a clause is compiled. Consider the following clause with no types:

\[ a(A, B) :- b(A, C), d(C, B). \]

Compilation of this clause proceeds in three steps: First the kernel Prolog is compiled to BAM code and a variable occurrence list, or varlist. In this example, most of the work in this step is done in the goal compiler. The resulting BAM code is referred to as skeleton code since variables have not yet been allocated to registers. The varlist is derived from the skeleton code and contains the list of variables and registers in it. Second, the register allocator uses the varlist to allocate variables to registers. Third, after all predicates and all clauses are compiled, the BAM optimization stage improves the code (Chapter 6). The skeleton code for this clause is:

```
allocate(X). ; Create an environment (its size is still unknown).
move(r(0), A). ; Load the head arguments into variables A and B.
move(r(1), B).
move(tvar-r(h), C). ; Create an unbound variable and put it in C and D.
move(tvar-r(h), D). ; C may exist beyond a call, D exists between calls.
pragma(push(variable)).
push(D, r(h), 1).
move(A, r(0)). ; Load the parameters of the first call.
move(D, r(1)).
call(b/2).
pragma(tag(C, tvar)). ; C has an extra link, with a tvar tag.
move([C], r(0)). ; Extra indirection to remove the extra link.
move(B, r(1)).
call(d/2).
deallocate(X). ; No last call optimization in the skeleton code.
return.
```

The varlist for this clause is:
Corresponds to move(r(0),A).

Corresponds to the unbound variable in C and D.

Corresponds to call(b/2).

Corresponds to call(d/2).

3.1.1. Construction of the varlist

The varlist is constructed to satisfy these conditions:

1. The only contents of the varlist are unbound variables, temporary registers, and the atoms fence and pref.

2. The order of variable occurrences is the same in the skeleton code and the varlist.

3. The atom fence is inserted as a marker at each point where temporary variables do not survive. This corresponds to each call(...) instruction in the skeleton code.

4. Two variables that are preferably allocated to the same register are preceded by the atom pref and called a pref pair. A pref pair is created when allocating the variables to the same register allows an instruction to be removed. For example, the move(A,r(0)) instruction can be removed if the variable A is allocated to register r(0).

5. A variable occurs exactly once in the varlist if and only if it occurs exactly once in the skeleton code. Such a variable is called a void variable. An instruction containing a void variable may be removed.

6. A variable occurs more than once in the varlist if and only if it occurs more than once in the skeleton code.

3.1.2. The register allocator

The register allocator assigns a register to each variable in the varlist such that there are no conflicts, i.e. a single register never holds two values at the same time. The allocator also calculates the size of the environment (the number of permanent registers) for the allocate and deallocate instructions.

The algorithm is defined in Figure 5.2. It assumes that variables are represented as logical variables, i.e.
procedure register_allocator(VL : varlist);
var \ V_{end}, V_{temp}, V_{pref}, V_{perm} : set of variable;
begin
\ V_{end} := \{ variable Y | Y occurs exactly once in VL \} ;
\forall X \in \ V_{end} \ do \ Allocate \ each \ X \ to \ r (\text{void});
\ V_{perm} := \{ variable Y | The \ sequence \ \{ Y, ..., fence, ..., Y \} \ occurs \ in \ VL \} ;
\forall X \in \ V_{perm} \ do \ Allocate \ each \ X \ to \ a \ different \ p ( I ) ;
Environment \ size := \ number \ of \ elements \ in \ V_{perm} ;
\ V_{temp} := \{ variable Y | Y \ occurs \ more \ than \ once \ in \ VL \} ;
\ V_{pref} := prefer( VL ) ;
while \ V_{temp} \neq \emptyset \ do \ begin
\ while \ \exists X \in \ V_{pref} : X \ is \ allocatable \ to \ r ( I ) \ without \ conflict \ do \ begin
Allocate \ X \ to \ its \ preferred \ register \ r ( I ) ;
\ V_{pref} := \ V_{pref} - \{ X \} ;
\ V_{temp} := \ V_{temp} - \{ X \} ;
\ V_{pref} := prefer( VL )
\ end ;
if \ \exists X \in \ V_{temp} \ then \ begin
Allocate \ X \ to \ the \ lowest \ r ( I ) \ possible \ without \ conflict ;
\ V_{pref} := \ V_{pref} - \{ X \} ;
\ V_{temp} := \ V_{temp} - \{ X \} ;
\ V_{pref} := prefer( VL )
\ end ;
end ;
\end;
function prefer( VL : varlist ) : set of variable;
begin
return \{ variable Y | The \ sequence \ \{ pref,Y,...\} \ or \ \{ pref,_,Y \} \ occurs \ in \ VL \} ;
\end ;
Figure 5.2 – The register allocator

that allocating a variable to a register binds that variable in all sets that contain it. It assumes that there are
an infinite number of temporary and permanent registers. It uses the following correspondence between
variable lifetimes and registers:

(1) A variable that occurs exactly once is allocated to \ r (\text{void}).

(2) A variable occurring on both sides of a fence marker (it crosses a fence) is allocated to a per-
manent register \ p ( I ) (a location in the environment).

(3) A variable that does not cross a fence and that occurs more than once is allocated to a temporary
register \ r ( I ).

The algorithm is independent of the write-once transformation and the dereference chain transformation.
This is possible because the clause compiler is careful to feed the allocator a varlist that takes the two transformations into account.

In the example of the previous section, the allocator assigns the following values to the variables:

- \( A = r(0) \)
- \( B = p(0) \)
- \( C = p(1) \)
- \( D = r(1) \)
- \( X = 2 \)

Since both \( B \) and \( C \) cross a fence, they are allocated to permanent registers. Both \( A \) and \( D \) are allocated to their preferred registers. The number of permanent variables, \( X \), is 2.

3.1.3. The final result

The final BAM code output by the compiler after all transformations and optimizations (including the BAM transformations of chapter 6) is:

```plaintext
allocate(2). ; Allocate space for two permanent variables.
move(r(1), p(0)).
move(tvar"r(h), r(1))", ; Create an unbound variable and put it in r(1) and p(1).
move(r(1), p(1)).
pragma(push(variable)).
push(r(1), r(h), 1).
call(b/2).
pragma(tag(p(1), tvar)).
movem(p(1), r(0)). ; Indirection due to dereference chain transformation.
movem(p(0), r(1)).
deallocate(2).
jump(d/2). ; Last call optimization converts 'call' to 'jump'.
```

3.2. The goal compiler

Given a goal and type information about the goal, this module sets up the arguments to call the goal, does the call, and sets up the return arguments. The main task of the goal compiler is to handle the complexities that arise when supporting combinations of uninitialized and initialized parameters. The following situations are also handled:

1. Duplicate variables. An uninitialized variable that occurs twice in a goal must be initialized before calling the goal.
(2) **Uninitialized register variables.** Passing arguments as uninitialized register variables requires some care. These variables are not passed into a predicate, but are outputs returned in registers.

(3) **Dummy predicates.** Several compiler transformations create new predicates as part of the transformation. These predicates are only called once, so they are compiled in-line.

(4) **Built-in predicates.** Some built-in predicates are translated into in-line code (Table 5.5).

---

function compile_goal(G :goal; F : formula; V_in : set) : return (Code : list; F_out : formula; V_out : set);

var V_uninit, V_init : set of variable;

Initcode, Precode, Call, Postcode : list of instruction;

A : term;

g, r : [init, mem, reg];

i : integer;

begin

/* Initialize all uninitialized variables that are duplicated */

V_uninit := \{ X \mid F \implies (\text{uninit}_{\text{mem}}(X) \text{ or } \text{uninit}_{\text{reg}}(X)) \};

V_init := (\text{vars}(G) \setminus V_{in}) \cup V_{uninit} \cap \text{dups}(G); /* Table 4.6*/

Initcode := list of (\forall X \in V_{init} : \text{Code to initialize the variable } X);

/* Pass arguments to the goal and clean up afterwards */

Pocode := [];

Postcode := [];

for i := 1 to arity(G) do begin

A := (argument i of goal G);

g_i := \text{given\_flag}(A, F, V_{in}); /* Table 5.1*/

r_i := \text{require\_flag}(A, G); /* Table 5.2*/

Append precod[ g_i, r_i ] to Precode; /* Table 5.3*/

Append postcode[ g_i, r_i ] to Postcode /* Table 5.4*/
end;

/* Call the goal */

if (G can be expanded in-line) then

Call := (in-line expansion of G) /* Table 5.5*/

else if (G is a dummy predicate) then

Call := (in-line compilation of G's definition)

else if (G does not alter temporary registers) then

Call := (a simple call instruction for G) /* Table 3.7*/

else

Call := (a call instruction for G);

Code := append(Initcode, Precode, Call, Postcode)

end;

end;

---

The function compile_goal(G, F, V_{in}) defines the goal compiler (Figure 5.3). Its inputs are the goal (G), a type formula (F), and the set of variables that have a value on input (V_{in}). Its outputs are a list of BAM instructions (Code), the type formula true on output (F_{out}), and the set of variables that have a value on input (V_{out}).
Each goal has three type formulas associated with it: a Require type, a Before type, and an After type. These types are optionally given by programmer input and are supplemented by dataflow analysis. The compiler maintains a table of these types for all predicates including built-ins and internals. The Require type gives the types that the arguments being passed to the goal must have, i.e. the goal compiler is required to make them true in all cases. The Before type gives the types that are true before the call. The After type gives the types that are true after the call returns. No special action is needed by the goal compiler to ensure the validity of the Before and After types.

Compiling a goal is made more complex because the kind of argument needed by the goal may not be the same as the one that is given to it. The goal's Given type (which is valid before the goal and given by $F$ in Figure 5.3) must be reconciled with the goal's Require type. The most common Require and Given types are the three varieties of unbound variables: uninitialized memory and register variables and initialized variables. This requires a case analysis with $3 \times 3$ cases for each argument of the goal to properly match the Require and Given types.

<table>
<thead>
<tr>
<th>Condition on argument $A$</th>
<th>$g_i$</th>
</tr>
</thead>
<tbody>
<tr>
<td>nonvar($A$)</td>
<td>ini</td>
</tr>
<tr>
<td>$\text{var}(A) \land (F \implies \text{uninit_mem}(A))$</td>
<td>mem</td>
</tr>
<tr>
<td>$\text{var}(A) \land ((A \in V_{sf}) \lor (F \implies \text{uninit_reg}(A)))$</td>
<td>reg</td>
</tr>
<tr>
<td>$\text{var}(A) \land (A \in V_{sf})$</td>
<td>ini</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Condition on argument $A$</th>
<th>$r_i$</th>
</tr>
</thead>
<tbody>
<tr>
<td>require($G$) implies uninit_mem($A$)</td>
<td>mem</td>
</tr>
<tr>
<td>require($G$) implies uninit_reg($A$)</td>
<td>reg</td>
</tr>
<tr>
<td>otherwise</td>
<td>ini</td>
</tr>
</tbody>
</table>
Table 5.3 – Calculating the precode from the flags

<table>
<thead>
<tr>
<th>$g_1$</th>
<th>$r_1$</th>
<th>precode [$g_1, r_1$]</th>
</tr>
</thead>
<tbody>
<tr>
<td>reg</td>
<td>reg</td>
<td>[]</td>
</tr>
<tr>
<td>mem</td>
<td>reg</td>
<td>[]</td>
</tr>
<tr>
<td>ini</td>
<td>reg</td>
<td>[]</td>
</tr>
<tr>
<td>reg</td>
<td>mem</td>
<td>[move(tvar, r(h), B), adda(r(h), 1, r(h))]</td>
</tr>
<tr>
<td>mem</td>
<td>mem</td>
<td>[]</td>
</tr>
<tr>
<td>ini</td>
<td>mem</td>
<td>[move(tvar, r(h), B), adda(r(h), 1, r(h))]</td>
</tr>
<tr>
<td>reg</td>
<td>ini</td>
<td>[move(tvar, r(h), B), push(B, r(h), 1)]</td>
</tr>
<tr>
<td>mem</td>
<td>ini</td>
<td>[move(A, [A]), move(A, B)]</td>
</tr>
<tr>
<td>ini</td>
<td>ini</td>
<td>[]</td>
</tr>
</tbody>
</table>

Table 5.4 – Calculating the postcode from the flags

<table>
<thead>
<tr>
<th>$g_1$</th>
<th>$r_1$</th>
<th>postcode [$g_1, r_1$]</th>
</tr>
</thead>
<tbody>
<tr>
<td>reg</td>
<td>reg</td>
<td>[]</td>
</tr>
<tr>
<td>mem</td>
<td>reg</td>
<td>[move(B, [A])]</td>
</tr>
<tr>
<td>ini</td>
<td>reg</td>
<td>unify(A, B)</td>
</tr>
<tr>
<td>reg</td>
<td>mem</td>
<td>[move(B, A)]</td>
</tr>
<tr>
<td>mem</td>
<td>mem</td>
<td>[]</td>
</tr>
<tr>
<td>ini</td>
<td>mem</td>
<td>unify(A, B)</td>
</tr>
<tr>
<td>reg</td>
<td>ini</td>
<td>[move(B, A)]</td>
</tr>
<tr>
<td>mem</td>
<td>ini</td>
<td>[]</td>
</tr>
<tr>
<td>ini</td>
<td>ini</td>
<td>[]</td>
</tr>
</tbody>
</table>

Require and Given flags $r_1$ and $g_1$ (with values in {ini, mem, reg}) are associated with each goal argument for the Require and Given types. Tables 5.1 and 5.2 define how the Require and Given flags are calculated. The function require($G$) in Table 5.2 is a defined predicate in the compiler that returns the Require type for any goal. It knows all about built-in and internal predicates and the results of dataflow analysis.

Duplicate arguments (e.g. $A$ in the call $p(A, A)$) are treated specially. An argument that is duplicate cannot be uninitialized—it occurs in more than one place, so it is not unaliased any more. The goal compiler initializes these arguments before doing the case analysis.

Table 5.3 gives the precode, i.e. the code that is generated before the call to set up, and Table 5.4 gives the postcode, i.e. the code that cleans up after the call. To enforce the Require type, in seven of the nine cases a different argument $B$ is passed to the call instead of the goal’s original argument $A$. For example, if the Given flag is mem and the Require type is reg, then the compiler must create a new variable $B$ of type uninitt_reg($B$) to pass to the goal. After the goal returns, the original argument $A$ and the returned argument $B$ are unified together. The new variable $B$ is created for all combinations of Given and
Require flags except (reg, reg) and (mem, mem). In these two cases no precode or postcode is needed.

To simplify the presentation, Figure 5.3 only does part of what the algorithm implemented in the compiler does. The definition of compile_goal in the figure only handles Require and Given types that are all uninitialzied variables. The actual algorithm handles any types. The type formula $F$ and the variable set $V_{ij}$ are updated continuously during the execution of compile_goal. A variable occurrence list is calculated for the register allocator. The actual algorithm handles 12 cases for parameter passing instead of 9—as an optimization, two varieties of Given uninitialized register types are recognized.

<table>
<thead>
<tr>
<th>Kernel Prolog</th>
<th>BAM instruction</th>
</tr>
</thead>
<tbody>
<tr>
<td>'Scut_load' (X)</td>
<td>move(r(b), X)</td>
</tr>
<tr>
<td>'Scut' (X)</td>
<td>cut(X)</td>
</tr>
<tr>
<td>'Sname_arity' (X, '., 2)</td>
<td>test(ne, t1st, X, fail)</td>
</tr>
<tr>
<td>'Sname_arity' (X, Na, Ar)</td>
<td>equal([X], tatm&quot;Na/Ar&quot;, fail)</td>
</tr>
<tr>
<td>'Sname_arity' (X, Na, 0)</td>
<td>equal(X, tatm&quot;Na&quot;, fail)</td>
</tr>
<tr>
<td>'Stest' (X, Types)</td>
<td>(a sequence of test instructions)</td>
</tr>
<tr>
<td>'Sequal' (X, Y)</td>
<td>equal(X, Y, fail)</td>
</tr>
<tr>
<td>'Sadd' (A, B, C)</td>
<td>add(A, B, C)</td>
</tr>
<tr>
<td>'Ssub' (A, B, C)</td>
<td>sub(A, B, C)</td>
</tr>
<tr>
<td>'Smod' (A, B, C)</td>
<td>mod(A, B, C)</td>
</tr>
<tr>
<td>'Smul' (A, B, C)</td>
<td>mul(A, B, C)</td>
</tr>
<tr>
<td>'Sdiv' (A, B, C)</td>
<td>div(A, B, C)</td>
</tr>
<tr>
<td>'Sand' (A, B, C)</td>
<td>and(A, B, C)</td>
</tr>
<tr>
<td>'Sor' (A, B, C)</td>
<td>or(A, B, C)</td>
</tr>
<tr>
<td>'Sxor' (A, B, C)</td>
<td>xor(A, B, C)</td>
</tr>
<tr>
<td>'Ssll' (A, B, C)</td>
<td>sll(A, B, C)</td>
</tr>
<tr>
<td>'Ssra' (A, B, C)</td>
<td>sra(A, B, C)</td>
</tr>
<tr>
<td>'Snot' (A, C)</td>
<td>not(A, C)</td>
</tr>
</tbody>
</table>

3.2.1. An example of goal compilation

This section gives a simple example of compilation to show how the goal compiler works in practice.

Consider the following predicate in standard Prolog:

\[ a(X, Y) :- Y \text{ is } X+1. \]

This is converted to kernel Prolog:

\[ a(X, Y) :- 'Sadd'(X, 1, Y). \]

To compile the call to $'Sadd'/3$ it is necessary to pass parameters in the right way. In particular, it is necessary to pass the output of the addition into variable $Y$. The built-in $'Sadd'(A, B, C)$ has the
following types associated with it:

\[
\text{Require} = (\text{deref}(A), \text{deref}(B), \text{uninit\_reg}(C)).
\]
\[
\text{After} = (\text{integer}(A), \text{integer}(B), \text{integer}(C), \text{rderef}(A), \text{rderef}(B), \text{rderef}(C)).
\]

From the Require type, the first two arguments \(X\) and \(I\) of \(\text{Sadd} / 3\) must be dereferenced and the third argument \(Y\) must be an uninitialized register. The Given types of \(X\) and \(Y\) depend on the type formula for \(a(X, Y)\). Assume first that no type is given for \(a(X, Y)\). From Tables 5.1 and 5.2, the Given flag for \(Y\) is \text{ini} and the Require flag for \(Y\) is \text{reg}. From Tables 5.3 and 5.4, the precode in this case is empty and the postcode is a call to \(\text{unify}(A, B)\) to generate unification code. The compiled BAM code is:

```baml
procedure(a/2).
  deref(r(0), r(0)). ; Dereference \(X\).
  add(r(0), 1, r(0)). ; Perform the addition.
  deref(r(1), r(1)). ; Dereference \(Y\).
  unify(r(0), r(1), nonvar, ?, fail). ; Unify \(Y\) with the result of the addition.
return.
```

If \(a(X, Y)\) has a type then the code can often be simplified. For example, assume that its type is \((\text{deref}(X), \text{uninit\_mem}(Y))\), i.e. \(X\) is dereferenced and \(Y\) is an uninitialized memory variable.

Then the Given flag for \(Y\) is \text{mem}. The compiled BAM code is:

```baml
procedure(a/2).
  add(r(0), 1, r(0)). ; Perform the addition (\(X\) is dereferenced).
  pragma(tag(r(1), tvar)). ; Bind \(Y\) to the result of the addition.
  move(r(0), [r(1)]).
return.
```

3.3. The unification compiler

This section gives an overview of the compilation of unification, the optimizations that are done, and several examples.

3.3.1. The unification algorithm

Given a unification goal and type information about its arguments, this algorithm generates the simplest possible code to implement the unification. In the general case, the algorithm builds a tree of instructions. Each node of the tree has three branches—one each for read mode and write mode unification, and one for failure. The algorithm generates dereference instructions if necessary and trail instructions to undo
variable bindings when backtracking. It does other optimizations including optimal write mode unification, type propagation, and depth limiting.

Write mode unification of a term generates a block of push instructions that builds the term on the heap. Read mode unification of a term is done sequentially for each of the term's arguments. First it checks the name and arity of the term. Then the arguments are unified. For arguments that are simple terms this consists of a single move, equal, or unify instruction. For arguments that are compound terms the unification algorithm is called recursively.

The function unify(X, Y, F, V_{sf}) defines the unification algorithm (Figure 5.4 and 5.5). Its inputs are the two terms to be unified (X and Y), the type formula true on input (F), and the set of variables that have a value on input (V_{sf}). Its outputs are a list of BAM instructions (Code), the type formula true on output (F_{out}), and the set of variables that have a value on output (V_{sf, out}).

The algorithm does several tasks that are not shown in the figure since they would unnecessarily complicate the presentation. The instruction list, the type formula, and the variable set are updated continuously during the compilation. Before using the value of a variable, it is dereferenced if necessary. Before binding a value to a variable, it is trailed if necessary. A variable occurrence list (varlist) is calculated for the register allocator (Figure 5.2).

3.3.2. Optimizations

The actual implementation does four optimizations not shown in Figure 5.4 and 5.5. It does optimal write mode unification. It keeps track of terms that are ground and recursively dereferenced to avoid compiling superfluous write mode unifications and dereferences. To reduce code size, it performs the last argument optimization and the depth limiting transformation.

3.3.2.1. Optimal write mode unification

The algorithm is modified to build a compound term in write mode with the least number of move instructions. First the code for building the main functor with empty slots for its arguments is generated. This is followed by the code for building the arguments and filling in the slots with the correct heap offsets.
function unify(X, Y: term; F: formula; V, \text{out}: set) return \langle Code : list; F_{\text{out}} : formula; V_{\text{out}} : set \rangle;
begin
    Code := [];
    if (\text{var}(X) \&\& \text{var}(Y)) then begin
        if (F \implies (\text{unbound}(X) \lor \text{unbound}(Y))) then
            Compile a store instruction
        else
            Compile a call to a general unification subroutine;
    end else if (\text{nonvar}(X) \&\& \text{nonvar}(Y)) then begin
        Compile a check that X and Y have the same functor and arity \( a \);
        for \( i := 1 \) to \( a \) do begin
            Append \( \text{unify}(X, Y, F, V, 1) \) to Code
        end;
        return
    end if (\text{nonvar}(X) \&\& \text{var}(Y)) then Swap X and Y
    else if (\text{var}(X) \&\& \text{nonvar}(Y)) then Do nothing;
    if (X \in V_{\text{out}}) then return \( \text{unify} \_\text{write}(X, Y, F, V_{\text{out}}) \);
    else begin /* At this point \( X \in V_{\text{out}} */
        if (F \implies \text{nonvar}(X)) then return \( \text{unify} \_\text{read}(X, Y, F, V_{\text{out}}) \)
        else if (F \implies \text{var}(X)) then return \( \text{unify} \_\text{write}(X, Y, F, V_{\text{out}}) \)
        else begin
            Compile a three-way conditional branch comparing the tags of X and Y;
            Call \( \text{unify} \_\text{read} \) and \( \text{unify} \_\text{write} \) to compile the read and write mode branches
        end
    end
end;

Figure 5.4 – The unification compiler: the main routine

This technique was proposed as an optimization over the WAM by André Mariën [44]. The examples of unification given later use this technique. The justification of the BAM instructions needed for unification was done with this technique (Chapter 3).

3.3.2.2. Last argument optimization

This is an important optimization that significantly reduces the code size. It can be performed whenever a compound term has a compound term in its last argument. Without this optimization, the tree generated by the algorithm has the same depth as the term that is compiled. For each level in the tree a new block of write mode code is generated. For lists of \( n \) elements this results in \( O(n^2) \) move instructions.

The optimization reduces the code size to \( O(n) \) by creating only a single write mode block, and letting all depths of the tree jump into it. This optimization was proposed by Mats Carlsson [14]. The code for write
function unify_write(X: term; Y: term; F: formula; V_{nf}: set; V_{out}: set) return (Code : list; F_{out}: formula; V_{nf_out}: set);
begin
   /* At this point X is an unbound variable */
   Generate a block of instructions to create the term Y on the heap;
   Bind X to this block (i.e. generate code to dereference X if necessary,
   store a pointer to this block in X, and trail X if necessary)
end;

function unify_read(X: term; Y: term; F: formula; V_{nf}: set; V_{out}: set) return (Code : list; F_{out}: formula; V_{nf_out}: set);
begin
   /* At this point Y is a nonvariable and F implies nonvar(X) */
   Code := [ ];
   Compile a check that X contains a structure of same functor and arity as Y;
   for i := 1 to arity(Y) do begin
      Append unify(X, Y; F, V, i) to Code
   end
end;

Figure 5.5 – The unification compiler: read and write mode unification

mode unification of a nested term is replaced by a single jump instruction to the write mode code block of
the outermost term. An example of unification given below uses this optimization.

3.3.2.3. Type propagation

There are two ways in which propagating type information during the compilation of unification
improves the code. First, during the unification, the algorithm keeps track of the variables that are ground,
uninitialized, and recursively dereferenced. This information is propagated into the arguments of com-
 pound terms. The propagation of ground and recursively dereferenced types was added after measure-
 ments of the dataflow analyzer showed that these types are numerous.

Second, when a new variable is encountered in a term, then the unification compiler has the choice
whether to create it as an initialized variable or as an uninitialized variable. It is not always best to create
new variables as uninitialized, since this often makes it impossible to apply last call optimization. To solve
this problem it is necessary to look ahead in the clause. The variable is created as uninitialized only if there
is a goal later in the clause with this variable in an argument position that must be uninitialized.
3.3.2.4. Depth limiting

Because the unification compiler generates a separate read and write mode branch for each functor in the term that is unified, deeply nested terms result in a code size explosion. The last argument optimization (see above) reduces the code size when the nesting occurs in the last argument. For other cases, a different technique is necessary. The unification compiler replaces a deeply nested subterm by a variable, creates the subterm with write mode unification and does a general unification with the variable. The depth limit is set by the compiler option `depth_limit(N)`, and the default depth is `N=2`. For example, consider the following unification where the complicated term \( z(\ldots) \) is nested deeply:

\[
X\leftarrow s\left(t\left(u\left(z\left(...\right)\right)\right)\right)
\]

It is replaced by a sequence of three unifications:

\[
X\leftarrow s\left(t\left(u\left(...A\ldots\right)\right)\right), \quad B\leftarrow z\left(\ldots\right), \quad A\leftarrow B
\]

The variable \( B \) does not yet have a value, so the unification \( B\leftarrow z\left(\ldots\right) \) is executed in write mode. A general unification is performed for \( A\leftarrow B \). Since the size of a write mode unification is linear in the size of the compound term, this considerably shortens the code for deeply nested terms. Measurements were done to determine the effect of this transformation on execution time. In most cases it is insignificant, e.g., for the `nand` benchmark (Chapter 7), a program that contains deeply nested structures, the difference in execution time between depth limits of two and three is insignificant (i.e., only a few cycles out of several hundred thousand).

3.3.3. Examples of unification

Consider the following sample clause:

\[
a(A, \ s(A, [X]X))
\]

The WAM code for this clause is (assuming the two arguments of the clause are in registers \( r(0) \) and \( r(1) \)):
procedure a/2
get_structure s/2, r(1)
unify_value r(0)
unify_variable r(3)
get_list r(3)
unify_variable r(2)
unify_value r(2)
proceed

;; the clause has two arguments.
;; unify r(1) with s(A, [X|X]).
;; unify the first argument with r(0).
;; load the second argument into r(3).
;; unify r(3) with [X|X].
;; load the first argument into r(2).
;; unify the second argument with r(2).
;; return to caller.

Temporary values are stored in registers r(2) and r(3). The execution time of this code averaged over read and write mode is 63 cycles on the Xenologic X-1 processor [85], an implementation of the PLM architecture [28]. The BAM code generated for the same clause is (the pragmas have been left out for clarity):

procedure (a/2).

deref(r(1), r(1)). ;; dereference r(1).
switch(tstr, r(1), l(a/2,3), l(a/2,4), fail). ;; three-way branch.

label(l(a/2,3)). ;; write mode for s(A, [X|X]).

trail(r(1)). ;; conditionally push r(1) on trail stack.
move(tstr'h', r(1))). ;; bind s(A, [X|X]) to second argument.
push(tatm' s/2, h, 1). ;; create the term s(A, [X|X]).
push(r(0), h, 1).
push(tlist' h+2, h, 1).

pad(1).

label(l(a/2,1)). ;; common code for last arg. opt.

move(tvar'h', r(2)). ;; create the two arguments of [X|X].
push(r(2), h, 1).
push(r(2), h, 1).
return.

label(l(a/2,4)). ;; read mode for s(A, [X|X]).
equal((r(1), tatm' s/2, fail)). ;; check functor & arity of s/2.
move((r(1)+1), r(3)). ;; load first argument into r(3).
deref(r(3), r(3)).
deref(r(0), r(0)).
unify(r(3), r(0), 2, 2, fail). ;; unify first argument with r(0).
move((r(1)+2), r(0)). ;; load second argument into r(0).
deref(r(0), r(0)).
switch(tlist, r(0), l(a/2,6), l(a/2, 7), fail). ;; three-way branch.

label(l(a/2,6)). ;; write mode for [X|X].

trail(r(0)).
move(tlist'h', r(0))).
jump(l(a/2,1)). ;; jump to common code (last arg. opt.).

label(l(a/2,7)). ;; read mode for [X|X].

move((r(0), r(2)).
move((r(0)+1), r(0)).
deref(r(0), r(0)).
deref(r(2), r(2)).
unify(r(0), r(2), 2, 2, fail). ;; unify arguments of [X|X].
return.

Again, the two arguments of the clause are in registers r(0) and r(1) and temporary values are stored in registers r(2) and r(3). To reduce the code size, the write mode code for [X|X] jumps into the
middle of the code for \( s(A, [X|X]) \). With this optimization the code is 29 BAM instructions long (after translation and instruction reordering, this is 264 bytes on the VLSI-BAM). The WAM code is only 7 instructions long (17 bytes on the PLM) because each instruction encapsulates a choice. WAM instructions for unification assume the existence of a read/write mode bit in the implementation, which collapses the execution tree onto itself.

The code size ratio VLSI-BAM/PLM is large for this example. It was hoped during development that (1) code expansion would be less for other kinds of Prolog code (e.g. calls, parameter passing, backtracking), and (2) dataflow analysis would reduce the complexity of unifications. These intuitions have been borne out (Chapter 7): the static code size in VLSI-BAM bytes measured for large programs is only three times that of the PLM, a microcoded WAM with a byte-coded instruction set.

The execution time of the above code on the VLSI-BAM is 25 cycles (measured with a simulator taking pipeline delays into account and averaged over read and write mode). This is about 40% of the cycles needed for the X-1. This time can be estimated by taking the average execution times of BAM instructions when translated to the VLSI-BAM architecture: unify takes 5 cycles, equal takes 3 cycles, switch, deref, trail, and move from memory take 2 cycles each, push, adda, and all other move instructions take 1 cycle each, and pad instructions take 0 cycles because they are collapsed into the pushes. These estimates are only approximately correct because of instruction reordering optimizations performed on VLSI-BAM code.

Through programmer annotation or dataflow analysis it is sometimes possible to know the type of an argument at compile-time. For example, sometimes it is known whether an argument is unbound or bound. Consider the same sample clause again:

\[ a(A, s(A, [X|X])). \]

Assume it is known that the second argument is an uninitialized memory variable. This is expressed with the following type declaration:

\[ :- \text{mode}(\text{a}(A,B):-\text{uninit\_mem}(B)). \]

With this type the clause's code is only 9 BAM instructions long (36 bytes on the VLSI-BAM):
procedure(a/2)
move(tstr"h,[r(1)]) :: bind s(A,[X|X]) to second argument.
push(tatm"(s/2),h,1).
push(r(0),h,1).
push(tlist"(h+2),h,1).
pad(1).
move(tvar"h,r(0)) :: create the two arguments of [X|X].
push(r(0),h,1).
push(r(0),h,1).
return. :: return to caller.

The execution time of this example is 11 cycles.

3.4. Entry specialization

For each goal in the clause, the clause compiler attempts to replace it with a faster entry point, depending on the types existing at that point. For example, if it is known that the arguments N and A of the predicate functor (X, N, A) are atomic then a faster version can be compiled.

Entry specialization is done in both the clause compiler and the dataflow analysis. Doing it in both places is complementary since the analysis only keeps track of a limited set of types: ground, nonvariable, uninitialized, and recursively dereferenced. During clause compilation more information is known, for example, if the goal X<Y occurs in a clause, then afterwards it is known that X<Y is true. Analysis does not have a representation for this information, but it could be useful for entry specialization.
Entry specialization can be done for any predicate whose definition is not in the program. The system has implemented this for the built-in predicates, but it can be used by the programmer for any library predicate. For each predicate that has faster entry points, a modal_entry declaration is given, along with type declarations for the fast entry points. These declaration are used in the dataflow analysis and the clause compiler to replace any call to the predicate with a faster entry point. For example, here is the modal entry declaration for the name(A,B) built-in predicate:

```
:- modal_entry(name(A,B),
    mode(atomic(A),
        mode(uninit(B),
            entry('S_name < *1' (A,B)),
            entry('S_name < *1' (A,B))
        ),
        mode(uninit(A),
            entry('S_name > *2' (A,B)),
            mode(uninit(B),
                entry('S_name > *2' (A,B)),
                entry(name(A,B))
            )
    )
).
```

This declaration defines a binary tree, depicted in Figure 5.6. The nodes of the tree are decision points containing a type. If the type is valid then the left subtree is chosen, otherwise the right subtree is chosen. The leaves of the tree are the entry points. If none of the types are valid then the leftmost leaf is chosen, which
usually is the same predicate as the original one. Each of the four fast entry points also has a type declaration:

\[
\begin{align*}
&\text{:- mode('S name > 1' (A,B), (derf(A),derf(B)), atomic(A), (list(B),ground(B)), n).} \\
&\text{:- mode('S name < '1' (A,B), (uninit(A),derf(B)), true, (atomic(A),derf(A),list(B),ground(B)), n).} \\
&\text{:- mode('S name > '2' (A,B), (derf(A),uninit(B)), true, (atomic(A),list(B),ground(B),rderef(B)), n).} \\
&\text{:- mode('S name > 1 *2' (A,B), (derf(A),uninit(B)), atomic(A), (list(B),ground(B),rderef(B)), n).}
\end{align*}
\]

These declarations are written in a five-argument form that is more general than a standard type declaration (Appendix A): it gives the entry types (both Require and Before) and the exit (After) types for the predicate.

3.5. The write-once transformation

In the BAM all unbound variables are kept on the heap. This makes trail checking significantly faster. However, when combined with the ability to destructively modify the value of permanent variables (e.g. to dereference them and save the dereferenced value in the permanent) it leads to several problems. These problems are all neatly resolved by the write-once transformation.

Putting all unbound variables on the heap means that there are no pointers to the environment/choice point stack; all pointers point to the heap. This reduces trail checking to a single comparison with the heap backtrack pointer \(\pi(hb)\) and a conditional push to the trail stack. It is not necessary to do another comparison to decide whether the variable is on the heap or in an environment. In addition, since all unbound variables are created on the heap there are no "unsafe variables" as in the WAM. An unsafe variable is an unbound variable that is created on the environment and that must be moved to the heap ("globalized") before last call optimization deallocates its memory.

Modifying the value of a permanent variable (e.g. by dereferencing or binding it) cannot be done without a trail operation. Indeed, consider the case where a permanent dereferences to a nonvariable term. If the dereferenced value overwrites the original value, then both the original value and its address have to be trailed since backtracking has to restore the original value. This is expensive, since it has to be done
every time a permanent is bound or dereferenced.

One solution to this problem is never to store a dereferenced permanent back in the environment. This solves the problem but it is inefficient since a permanent may have to be dereferenced several times in a clause.

A better solution is to allocate a new permanent on the environment whenever the value of an old one needs to be changed. The new permanent gets the new value and the old permanent is unchanged. As a result, all permanent variables are only given values once, so they are called “write-once” permanents. Because it is not changed, the old permanent does not have to be trailed. At the cost of a slightly bigger environment, this completely eliminates the need to trail permanent variables. This allocation scheme is implemented in the clause compiler.

To summarize:

1. All unbound variables are created on the heap, and unbound permanent variables in an environment always point to the heap.

2. The trail check is a single comparison with r (hb) and a conditional push to the trail stack (2 cycles on the VLSI-BAM).

3. Permanent variables are only given a single value in a clause. Whenever a permanent would be changed, a new one is allocated and given the modified value.

4. Register allocation must allocate a different permanent register for each permanent variable in the clause. It is not allowed to use the same register for two variables whose lifetimes do not overlap.

This solution is implemented in the clause compiler by mapping a permanent variable onto a new variable whenever its value would change. The register allocator treats the new variables just like any other, and allocates them to temporary or permanent registers.

The main disadvantage of this technique is that environments are larger. For example, consider a clause of the form:

\[e(A, E) :- a(A, B), b(B, C), c(C, D), d(D, E).\]

where variables are chained from one predicate to the next. In the WAM, it is allowed to allocate
permanent variables such that variables whose lifetimes do not overlap are allocated to the same permanent register. For the above example, this requires just two permanent registers, so the total environment size is four words (it also includes registers \( r(e) \) and \( r(cp) \)). Only two permanents are needed no matter how long the chain of body goals is. This method requires trailing of the permanent's values, because backtracking must see the original values. This scheme is consistent with the original implementation of the WAM, i.e. binding permanent variables on the environment and globalizing unsafe variables to ensure correctness.

In contrast, the number of permanent variables needed by the write-once technique increases linearly with the length of the chain. For the above example, this requires four permanent variables, so the total environment size is six words. The total memory usage is increased by less than this amount because no trailing of permanents is needed.

This is an example of a trade-off between memory space and execution time. The extra memory space needed is comparable to the increased size of the trail stack if there is no trail check for permanent variables. Since this is small, I have opted to decrease execution time at the expense of larger environments. By keeping all unbound variables on the heap and by implementing permanent variables as write-once variables, permanent variables can be dereferenced and bound without trailing, and the cost of trailing heap variables is reduced to a single comparison and conditional push.

### 3.6. The dereference chain transformation

This transformation is needed to maintain consistency between the dataflow analysis and the clause compiler. A new unbound variable (of either initialized type or uninitialized memory type) is created as a pointer to a memory location. Binding the variable stores the new value in the location. However, the register(s) that originally contained the unbound variable still have pointers to the location. One level of indirection is needed to access the value.
To see why this is necessary and what it implies, consider the execution of the clause `main` (Figure 5.7):

```
main :- (i) a(A), (ii) write(A).
```

```
a(A) :- A = t(s(t(a), u(b), v(c))).
```

The relevant situation can be seen in the transition from (i) (just before the call to `a(A)` to (ii) (just after the call to `a(A)`). At (i) a new unbound variable A is created on the heap. At (ii) the variable A has been bound to a value. The important point is that A still has a `tvar` tag, and that one indirection is needed to access the `tstr` pointer. The extra link exists because the creation of A and its binding are done in separate steps. This is true for both initialized unbound variables and uninitialized memory variables.
This situation is not a problem unless dataflow analysis determines that A is returned as a dereferenced value. In that case there is a conflict between what the analysis deduces and what the clause compiler thinks is true. There are two ways to solve this problem: either weaken the analysis so that it will not deduce a dereference type in this case, or modify the clause compiler to ensure that the variable is dereferenced by doing an extra indirection whenever the variable is accessed after it is bound. The compiler implements the second solution since dereferencing is a time-consuming operation and it is important to derive as many dereference types as possible. The trade-off between doing an extra indirection for a value that may not be accessed later and doing an extra dereference loop seemed to be a fair one.

The compiler inserts code to do this indirection whenever the variable is accessed after it is bound. In addition to maintaining consistency with the analysis, this speeds up later dereferencing. There is a minor interaction with the register allocator—for correctness, variables that get an extra indirection are not allowed to be pref pairs.
Chapter 6
BAM Transformations

1. Introduction

After compiling the program from kernel Prolog into BAM code, a series of optimizing transformations is performed. The transformations performed are: (1) duplicate code elimination, (2) dead code elimination, (3) jump elimination, (4) label elimination, (5) synonym optimization, (6) peephole optimization, and (7) determinism optimization. This chapter first gives two definitions and then presents the transformations.

2. Definitions

The following two definitions are useful:

Definition DB: A distant branch is a branch that always transfers control to an instruction other than the next in the instruction stream.

According to this definition, there are exactly four distant branches in the BAM: fail, return, jump, and switch. All other branches do not satisfy the definition since they can fall through to the next instruction.

Definition BB: A contiguous block is any sequence of instructions that terminates with a distant branch.

According to this definition, a contiguous block can start with any instruction and can contain conditional branches with a fall through case. Therefore the code contains a large number of overlapping contiguous blocks. This is useful to get maximum optimization when looking for contiguous blocks that satisfy some property. The individual transformations mentioned in this chapter will usually only look at contiguous blocks satisfying certain constraints, for example, the contiguous blocks that begin with a label.

3. The transformations

Seven transformations (Figure 6.1) are done on the BAM code generated for each predicate by the kernel to BAM compilation stage. A transitive closure is performed on the sequence of seven transformations, i.e., they are applied repeatedly until there are no more changes. Each transformation is carefully
coded to result in code that is better (i.e. faster or shorter) than its input, so the closure operation terminates.

![Diagram of BAM Transformations]

3.1. Duplicate code elimination

All duplicate contiguous blocks except the last occurrence are replaced by a jump to the last one. This optimization is also known as *cross-jumping*. It tightens up loose code generated by the type enrichment transformation (Chapter 4). It is implemented by first creating a table indexed by all contiguous blocks that (1) begin with a label, (2) do not contain any other labels (but they are allowed to contain branches), and (3) are not degenerate blocks that consist of only a single jump, return, or fail instruction (but a single switch is allowed). The table contains the label of the last occurrence of the block. All con-
tiguous blocks in the code, including those that do not begin with labels, are looked up in the table and replaced by jumps if they are not the last occurrence. The result of this optimization is to reduce code size at the price of slightly slowing down execution.

3.2. Dead code elimination

All code that is not reachable from the entry point of a predicate is removed. This is done in two steps: First, all the labels that are reachable through any number of branches are calculated by doing a transitive closure. Second, a linear traversal of the code is done and the instructions following a distant branch up to the next reachable label are eliminated.

3.3. Jump elimination

Rearrange contiguous blocks to minimize the number of jump, call, and return instructions. This optimization is a variant of the jump chaining optimization. A transitive closure is done on the following replacements:

1) Replace a jump by the contiguous block it points to if the block is only pointed to by one branch or if the block is shorter than a preset threshold. The threshold can be changed by a compiler directive. The replacement is not done if the block is part of write mode unification or unification with an atom, since these two cases are hurt by the transformation.

2) Replace a call to a dummy predicate by the code for the predicate if it is straightline code, i.e. its code consists only of non-branches, call instructions, and branches all of whose destinations are fail. The predicate's code must be terminated by a return or fail instruction.

3) Replace a conditional branch to a conditional branch by a new conditional branch if possible. The only case currently recognized is:

\[
\text{test}(\text{ne}, \text{tvar}, V, L).
\]

\[
\ldots
\]

\[
\text{label}(L).
\]

\[
\text{switch}(\text{Tag}, V, \text{fail}, L2, L3).
\]

which causes the test instruction to be replaced by:
4. Replace a branch one of whose destinations is a jump or fail instruction by a new branch identical to the original one except that the destination label is replaced by the destination label of the jump or by fail.

3.4. Label elimination

Remove all labels that are not jumped to by any branch in the code. This is done in two steps: First, the set of all destinations of all branch instructions is collected. Second, the labels not in this set are removed from the code.

3.5. Synonym optimization

This transformation is similar to strength reduction. It does a linear traversal of the code and replaces every addressing mode by the cheapest addressing mode that contains the same value. For example, if \( p(1) \) and \( r(0) \) contain the same value, then an occurrence of \( p(1) \) can be replaced by \( r(0) \). The following cost order (from cheapest to most expensive) is used by default and is based on the cost in the VLSI-BAM architecture:

<table>
<thead>
<tr>
<th>Addressing mode</th>
<th>Reason for cost</th>
<th>Overhead (cycles)</th>
</tr>
</thead>
<tbody>
<tr>
<td>( r(b) )</td>
<td>Promotes creation of cut ((r(b))) which is a no-op</td>
<td>0</td>
</tr>
<tr>
<td>( r(I) )</td>
<td>Usable without overhead</td>
<td>0</td>
</tr>
<tr>
<td>Atom</td>
<td>Requires ldi (load immediate) instruction</td>
<td>1</td>
</tr>
<tr>
<td>Tag(^X)</td>
<td>Tagged pointer creation needs lea (load effective address) instruction</td>
<td>1</td>
</tr>
<tr>
<td>( p(I) )</td>
<td>Permanent variable needs ld (load) instruction</td>
<td>1</td>
</tr>
<tr>
<td>( {r(I)} )</td>
<td>Indirection needs ld (load) instruction</td>
<td>1</td>
</tr>
<tr>
<td>( {r(I)+N} )</td>
<td>Offset indirect needs ld (load) instruction</td>
<td>1</td>
</tr>
<tr>
<td>( {p(I)} )</td>
<td>Indirect permanent needs 2 ld (load) instructions</td>
<td>2</td>
</tr>
<tr>
<td>( {p(I)+N} )</td>
<td>Offset indirect permanent needs 2 ld (load) instructions</td>
<td>2</td>
</tr>
<tr>
<td>( r(void) )</td>
<td>Most expensive because it must not be changed</td>
<td>-</td>
</tr>
</tbody>
</table>

The reason given for the cost describes the instructions necessary to implement the addressing mode for the VLSI-BAM. More information on the instruction set of the VLSI-BAM is given in [34]. The addressing mode \( r(void) \) is created by the register allocator. It corresponds to a void variable, i.e. a variable that occurs only once in a clause and whose value may therefore be ignored. It is made the most
expensive because it must remain unchanged so that peephole optimization can remove the instruction containing it.

The synonym optimization is implemented by maintaining a set of equivalence classes at all points of the program, where each equivalence class is a set of addressing modes whose values are identical. Labels in the code cause the set of equivalence classes to be reset to empty. A future extension of this module could eliminate this restriction by following the labels and performing a transitive closure, resulting in a slight performance gain.

3.6. Peephole optimization

A transitive closure is performed on a peephole transformation with a window of three instructions. The set of patterns was determined empirically by looking at the compiler's output and adding patterns to fix obvious inefficiencies. Each pattern is implemented as a single clause in the optimizer. The patterns are one, two, and three instructions long. However, the window is extended to arbitrary size for one pattern, a generalized last call optimization:

\[
\begin{align*}
call(N/A), \\
deallocate(I). \text{ Arbitrary number of deallocate instructions.} \\
\ldots \\
deallocate(J). \\
return.
\end{align*}
\]

which is transformed to:

\[
\begin{align*}
deallocate(I). \text{ Same sequence as above.} \\
\ldots \\
deallocate(J). \\
jump(N/A).
\end{align*}
\]

3.7. Determinism optimization

A choice instruction is removed if it is followed by a sequence of instructions that cannot fail and a cut instruction. This simple-looking optimization significantly increases determinism—many predicates (e.g. Warren's quicksort benchmark) containing a cut become deterministic that would otherwise be compiled with a choice point.
A similar optimization is performed by the simplification transformation of kernel Prolog (Chapter 4). For example, it transforms $(!, p : q)$ into $(!, p)$. The determinism optimization extends simplification—if the goal $s$ compiles into instructions that cannot fail then it is able to successfully optimize the BAM code of $(s, !, p : q)$ even when simplification cannot determine that $s$ always succeeds.

Consider this predicate, which contains no cut:

```
:- mode((max(A, B, C) :- uninit(C))).  % C is unbound and unaliased.

max(A, B, C) :- A<=B, B=C.          % No cut here.
max(A, B, C) :- A=C.
```

It is compiled into the following BAM code (slightly simplified for readability):

```
procedure(max/3).
  deref(r(0),r(0)).
  deref(r(1),r(1)).
  jump(lts,r(0),r(1),l(max/3,1)).  % Conditional branch A<B.
  move(r(0),[r(2)]).                 % A<B is false.
  return.
label(l(max/3,1)).
  choice(1/2,[0,2],l(max/3,4)).      % A<B is true.
  move(r(1),[r(2)]).
  return.
label(l(max/3,4)).
  choice(2/2,[0,2],fail).
  move(r(0),[r(2)]).
  return.
```

When $A<B$ is true, a choice point is created to try both clauses. If a cut is inserted into the first clause:

```
:- mode((max(A, B, C) :- uninit(C))).  % C is unbound and unaliased.
max(A, B, C) :- A<=B, !, B=C.          % Cut is added here.
max(A, B, C) :- A=C.
```

then the code becomes deterministic:
procedure (max/3).
    move (b, r(3)).
    deref (r(0), r(0)).
    deref (r(1), r(1)).
    jump (lts, r(0), r(1), l(max/3, 4)). // Conditional branch A<B.
    move (r(0), (r(2))).
    return.
label (l(max/3, 4)).
cut (r(3)).
move (r(1), (r(2))).
return.

Measurements done by Touati [70] justify this optimization. He finds that it makes about half of all choice point operations avoidable.
Chapter 7
Evaluation of the Aquarius system

1. Introduction

This chapter attempts to quantify some of the ideas that were introduced in previous chapters. The evaluation process is as important as any other part of the implementation of a large software system. During the design phase it guides the design decisions. After the design is complete, it shows what features of the design contributed most to its effectiveness and it gives a foundation for starting the next design. Quantitative measurements are the most reliable guideposts one has during the design. For example, it is easy to imagine many possible compiler optimizations, but most of these have an insignificant effect on performance. It is more difficult to discover optimizations that are widely applicable.

Five evaluations are performed in this chapter:

1. The absolute performance of the system.
2. The effectiveness of the dataflow analysis.
3. The effectiveness of the determinism transformation.
4. A brief comparison with a high performance implementation of the C language.
5. A bug analysis, summarizing the number and types of bugs encountered during development.

Table 7.1 describes the benchmarks used in this chapter and their size in lines of code (not including comments). The benchmarks were chosen as examples of realistic programs doing computations representative of Prolog. This includes benchmarks that spend much of their time executing built-in predicates because this behavior is common in real-world programs. The benchmarks are divided into two classes, small and large, depending on whether the compiled code with analysis is smaller or larger than 1000 words. The benchmarks log10, ops8, times10, and divide10 are grouped together and referred to as deriv because they are closely related. The benchmarks are available by anonymous ftp to arpa.berkeley.edu.

All VLSI-BAM numbers in this chapter were obtained from the VLSI-BAM instruction-level simulator and include cache effects [17]. The simulated system has 128 KB instruction and data caches. The
### Table 7.1 - The benchmarks

<table>
<thead>
<tr>
<th>Benchmark</th>
<th>Lines</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>nreverse</td>
<td>10</td>
<td>Naive reverse of a 30-element list.</td>
</tr>
<tr>
<td>tak</td>
<td>15</td>
<td>Recursive integer arithmetic.</td>
</tr>
<tr>
<td>qsort</td>
<td>19</td>
<td>Quicksort of a 50-element list.</td>
</tr>
<tr>
<td>log10</td>
<td>27</td>
<td>Symbolic differentiation.</td>
</tr>
<tr>
<td>ops8</td>
<td>27</td>
<td>Symbolic differentiation.</td>
</tr>
<tr>
<td>times10</td>
<td>27</td>
<td>Symbolic differentiation.</td>
</tr>
<tr>
<td>divide10</td>
<td>27</td>
<td>Symbolic differentiation.</td>
</tr>
<tr>
<td>serialisc</td>
<td>29</td>
<td>Calculate serial numbers of a list.</td>
</tr>
<tr>
<td>queens_8</td>
<td>31</td>
<td>Solve the eight queens puzzle.</td>
</tr>
<tr>
<td>mu</td>
<td>33</td>
<td>Prove a theorem of Hofstadter’s &quot;mu-math.&quot;</td>
</tr>
<tr>
<td>zebra</td>
<td>36</td>
<td>A logical puzzle based on constraints.</td>
</tr>
<tr>
<td>sendmorec</td>
<td>43</td>
<td>The SEND+MORE=MONEY puzzle.</td>
</tr>
<tr>
<td>fast_mu</td>
<td>54</td>
<td>An optimized version of the mu-math prover.</td>
</tr>
<tr>
<td>query</td>
<td>68</td>
<td>Query a static database (with integer arithmetic).</td>
</tr>
<tr>
<td>poly_10</td>
<td>86</td>
<td>Symbolically raise a polynomial to the tenth power.</td>
</tr>
<tr>
<td>crypt</td>
<td>64</td>
<td>Solve a simple cryptarithmic puzzle.</td>
</tr>
<tr>
<td>meta_qsort</td>
<td>74</td>
<td>A meta-interpreter running qsort.</td>
</tr>
<tr>
<td>prover</td>
<td>81</td>
<td>A simple theorem prover.</td>
</tr>
<tr>
<td>browse</td>
<td>92</td>
<td>Build and query a database.</td>
</tr>
<tr>
<td>unify</td>
<td>125</td>
<td>A compiler code generator for unification.</td>
</tr>
<tr>
<td>flatten</td>
<td>158</td>
<td>Source transformation to remove disjunctions.</td>
</tr>
<tr>
<td>sdda</td>
<td>273</td>
<td>A dataflow analyzer that represents aliasing.</td>
</tr>
<tr>
<td>reducer_nowrite</td>
<td>298</td>
<td>A graph reducer based on combinators.</td>
</tr>
<tr>
<td>reducer</td>
<td>301</td>
<td>Same as above but writes its answer.</td>
</tr>
<tr>
<td>boyer</td>
<td>377</td>
<td>An extract from a Boyer-Moore theorem prover.</td>
</tr>
<tr>
<td>simple_analyzer</td>
<td>443</td>
<td>A dataflow analyzer analyzing qsort.</td>
</tr>
<tr>
<td>nand</td>
<td>493</td>
<td>A logic synthesis program based on heuristic search.</td>
</tr>
<tr>
<td>chat_parser</td>
<td>1138</td>
<td>Parse a set of English sentences.</td>
</tr>
<tr>
<td>chat</td>
<td>4801</td>
<td>Natural language query of a geographical database.</td>
</tr>
</tbody>
</table>

Caches are direct mapped and use a write-back policy. They are run in warm start; each benchmark is run twice and the results of the first run are ignored. The cache overhead is greatest for tak compiled without analysis, and for poly_10, simple_analyzer, chat, and boyer. For these programs it ranges from 9% to 24%. For meta_qsort, reducer, and chat_parser the overhead ranges from 2% to 3%. For all other programs the overhead is less than 0.5%.

#### 2. Absolute performance

This section compares the performance of Aquarius Prolog with Quintus Prolog. Tables 7.2 and 7.3 compare the performance of Quintus Prolog version 2.5 running on a Sun 4/65 (25 MHz SPARC) with that of Aquarius Prolog running on the VLSI-BAM (30 MHz). The "Raw Speedup" column gives the ratio of the speeds. The "Normalized Speedup" column divides this ratio by 1.8. Our group is in the process of...
porting the Aquarius system to the MIPS, MC68020, and SPARC processors. It was not possible to get numbers for these systems in time for the final version of this dissertation.

The normalization factor of 1.8 takes into account the Prolog-specific extensions of the VLSI-BAM (a factor of 1.5) and the clock ratio (a factor of 30/25 = 1.2). The general-purpose base architecture of the VLSI-BAM is very similar to the SPARC. The effect of the architectural extensions of the VLSI-BAM [34] has been carefully measured to be about 1.5 for large programs. However, for the small programs the compiler is able to remove many Prolog-specific features, so that the normalized speedup numbers in Table 7.2 are an underestimate.

<table>
<thead>
<tr>
<th>Benchmark</th>
<th>Size (lines)</th>
<th>Quintus v2.5 (Sun 4/65)</th>
<th>Aquarius (VLSI-BAM)</th>
<th>Normalized Speedup</th>
<th>Raw Speedup</th>
</tr>
</thead>
<tbody>
<tr>
<td>deriv</td>
<td>1.143</td>
<td>0.0913</td>
<td>7.0</td>
<td>12.5</td>
<td></td>
</tr>
<tr>
<td>log10</td>
<td>27</td>
<td>0.153</td>
<td>0.0168</td>
<td>1.143</td>
<td></td>
</tr>
<tr>
<td>ops8</td>
<td>27</td>
<td>0.239</td>
<td>0.0189</td>
<td>1.239</td>
<td></td>
</tr>
<tr>
<td>times10</td>
<td>27</td>
<td>0.345</td>
<td>0.0257</td>
<td>1.345</td>
<td></td>
</tr>
<tr>
<td>divide10</td>
<td>27</td>
<td>0.406</td>
<td>0.0299</td>
<td>1.406</td>
<td></td>
</tr>
<tr>
<td>reverse</td>
<td>10</td>
<td>1.62</td>
<td>0.136</td>
<td>6.6</td>
<td></td>
</tr>
<tr>
<td>qsort</td>
<td>19</td>
<td>4.820</td>
<td>0.173</td>
<td>15.5</td>
<td></td>
</tr>
<tr>
<td>serialise</td>
<td>29</td>
<td>3.10</td>
<td>0.447</td>
<td>3.9</td>
<td></td>
</tr>
<tr>
<td>query</td>
<td>68</td>
<td>23.7</td>
<td>3.57</td>
<td>3.7</td>
<td></td>
</tr>
<tr>
<td>mu</td>
<td>33</td>
<td>7.04</td>
<td>0.808</td>
<td>4.8</td>
<td></td>
</tr>
<tr>
<td>fast_mu</td>
<td>54</td>
<td>9.08</td>
<td>0.932</td>
<td>5.4</td>
<td></td>
</tr>
<tr>
<td>queens_8</td>
<td>31</td>
<td>21.2</td>
<td>1.13</td>
<td>10.4</td>
<td></td>
</tr>
<tr>
<td>tak</td>
<td>15</td>
<td>1120.</td>
<td>25.4</td>
<td>24.5</td>
<td></td>
</tr>
<tr>
<td>poly_10</td>
<td>86</td>
<td>417.</td>
<td>35.5</td>
<td>6.5</td>
<td></td>
</tr>
<tr>
<td>sendmore</td>
<td>43</td>
<td>490.</td>
<td>38.4</td>
<td>7.1</td>
<td></td>
</tr>
<tr>
<td>zebra</td>
<td>36</td>
<td>423.</td>
<td>84.1</td>
<td>2.8</td>
<td></td>
</tr>
<tr>
<td>geometric mean</td>
<td>6.7</td>
<td>12.1</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>standard deviation of mean</td>
<td>1.9</td>
<td>3.3</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

For the small benchmarks, the normalized speedup is somewhere between 6.7 and 12.1 (Table 7.2). The normalized speedup of the large benchmarks without built-in predicates is about 5.2 (Table 7.3). Speedup is better for the small benchmarks because dataflow analysis is able to derive better types for many of them. For some of them (such as tak and reverse) it derives essentially perfect types. The small programs show a large variation in speeds. The tak benchmark does well because it relies on integer arithmetic, which is compiled efficiently using uninitialized register types. The zebra benchmark does poorly for two reasons. First, it does a large amount of backtracking, which is inherently limited by memory bandwidth. Second, it works by successively instantiating arguments of a compound data
<table>
<thead>
<tr>
<th>Benchmark</th>
<th>Size (lines)</th>
<th>Quintus v2.5 (Sun 4/65)</th>
<th>Aquarius (VLSI-BAM)</th>
<th>Normalized Speedup</th>
<th>Raw Speedup</th>
</tr>
</thead>
<tbody>
<tr>
<td>No built-ins</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>prover</td>
<td>81</td>
<td>8.67</td>
<td>0.921</td>
<td>5.2</td>
<td>9.4</td>
</tr>
<tr>
<td>meta_qsort</td>
<td>74</td>
<td>49.6</td>
<td>4.71</td>
<td>5.8</td>
<td>10.5</td>
</tr>
<tr>
<td>nand</td>
<td>493</td>
<td>173.3</td>
<td>13.7</td>
<td>7.0</td>
<td>12.7</td>
</tr>
<tr>
<td>reducer_nowrie</td>
<td>298</td>
<td>312.</td>
<td>37.2</td>
<td>4.6</td>
<td>8.4</td>
</tr>
<tr>
<td>chat_parser</td>
<td>1138</td>
<td>1157.</td>
<td>129.5</td>
<td>5.0</td>
<td>8.9</td>
</tr>
<tr>
<td>browse</td>
<td>92</td>
<td>5450.</td>
<td>741.</td>
<td>4.1</td>
<td>7.4</td>
</tr>
<tr>
<td>geometric mean</td>
<td></td>
<td></td>
<td></td>
<td>5.2</td>
<td>9.4</td>
</tr>
<tr>
<td>standard deviation of mean</td>
<td></td>
<td></td>
<td></td>
<td>0.5</td>
<td>0.8</td>
</tr>
<tr>
<td>Including built-ins</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>unify</td>
<td>125</td>
<td>18.3</td>
<td>1.40</td>
<td>7.2</td>
<td>13.0</td>
</tr>
<tr>
<td>flatten</td>
<td>158</td>
<td>13.6</td>
<td>1.42</td>
<td>5.3</td>
<td>9.6</td>
</tr>
<tr>
<td>sdda</td>
<td>273</td>
<td>29.5</td>
<td>2.94</td>
<td>5.6</td>
<td>10.0</td>
</tr>
<tr>
<td>crypt</td>
<td>64</td>
<td>21.7</td>
<td>4.00</td>
<td>3.0</td>
<td>5.4</td>
</tr>
<tr>
<td>simple_analyzer</td>
<td>443</td>
<td>180.</td>
<td>33.4</td>
<td>3.0</td>
<td>5.4</td>
</tr>
<tr>
<td>reducer</td>
<td>301</td>
<td>405.</td>
<td>44.9</td>
<td>5.0</td>
<td>9.0</td>
</tr>
<tr>
<td>chat</td>
<td>4801</td>
<td>3100.</td>
<td>699.</td>
<td>2.5</td>
<td>4.4</td>
</tr>
<tr>
<td>boyer</td>
<td>377</td>
<td>4870.</td>
<td>1360.</td>
<td>2.0</td>
<td>3.6</td>
</tr>
<tr>
<td>geometric mean</td>
<td></td>
<td></td>
<td></td>
<td>3.8</td>
<td>6.9</td>
</tr>
<tr>
<td>standard deviation of mean</td>
<td></td>
<td></td>
<td></td>
<td>0.7</td>
<td>1.3</td>
</tr>
<tr>
<td>geometric mean (all large programs)</td>
<td></td>
<td></td>
<td></td>
<td>4.4</td>
<td>7.9</td>
</tr>
</tbody>
</table>

Table 7.4 – Time spent in built-in predicates

<table>
<thead>
<tr>
<th>Benchmark</th>
<th>Time (%)</th>
<th>Most used built-ins</th>
</tr>
</thead>
<tbody>
<tr>
<td>prover</td>
<td>0</td>
<td>-</td>
</tr>
<tr>
<td>meta_qsort</td>
<td>0</td>
<td>-</td>
</tr>
<tr>
<td>chat_parser</td>
<td>0</td>
<td>-</td>
</tr>
<tr>
<td>nand</td>
<td>&lt;1</td>
<td>-</td>
</tr>
<tr>
<td>browse</td>
<td>1</td>
<td>length/2</td>
</tr>
<tr>
<td>reducer</td>
<td>40</td>
<td>write/1, compare/3, arg/3</td>
</tr>
<tr>
<td>unify</td>
<td>40</td>
<td>arg/3, functor/3, compare/3</td>
</tr>
<tr>
<td>crypt</td>
<td>50</td>
<td>div/2, mod/2, */2</td>
</tr>
<tr>
<td>boyer</td>
<td>60</td>
<td>arg/3, functor/3</td>
</tr>
<tr>
<td>simple_analyzer</td>
<td>70</td>
<td>compare/3, sort/2, arg/3</td>
</tr>
<tr>
<td>sdda</td>
<td>70</td>
<td>write/1, =/=2, compare/3</td>
</tr>
<tr>
<td>flatten</td>
<td>80</td>
<td>write/1, sort/2, compare/3, name/2, functor/3, arg/3</td>
</tr>
</tbody>
</table>

The analysis algorithm does not have a representation for this operation, so it cannot be optimized.

The built-in predicates in Aquarius Prolog are not greatly faster than those in Quintus Prolog, since many of the Quintus built-ins are not written in Prolog, but in hand-crafted assembly. The Aquarius system shows better speedup over Quintus built-ins written in Prolog (such as read/1 and write/1) and the entry specialization transformation also speeds up the built-ins. Table 7.4 gives the percentage of time that
the benchmarks spend executing inside built-in predicates. This number does not take into account built-ins that are implemented as in-line code (arithmetic test, addition and subtraction, and type checking). The table also gives the most often used built-in predicates for each benchmark in decreasing order of usage.

Several benchmarks use built-in predicates significantly. The normalized speedup for these programs is 3.8, somewhat less than programs without built-ins (Table 7.3). The normalized speedup for all large programs is 4.4 (the reducer benchmark is counted only once in this average). The boyer benchmark does poorly because it relies heavily on the arg/3 and functor/3 built-in predicates. The chat benchmark uses these built-ins as well as others including setof/3, but it was not possible to measure the fraction of execution time spent in them. The sdda and flatten benchmarks do well partly because the write/1 built-in is much faster in Aquarius than in Quintus.

3. The effectiveness of the dataflow analysis

This section evaluates the effectiveness of the dataflow analysis with three kinds of measurements. Tables 7.5, 7.6, and 7.7 give the effect of the dataflow analyzer on performance and code size, and the efficiency of the analyzer both in terms of its execution time and the fraction of arguments for which types can be deduced.

For a representative set of realistic Prolog programs of various sizes up to 1,100 lines, the analyzer is able to derive type information for 56% of all predicate arguments. It finds that on average 23% of all predicate arguments are uninitialized, 21% of arguments are ground, 10% of arguments are nonvariables, and 17% of arguments are recursively dereferenced. The sum of these three numbers is greater than 56% since it is possible for an argument to have multiple types, e.g. it can be ground and recursively dereferenced at the same time. Doing analysis reduces execution time on the VLSI-BAM by 18% for programs without built-ins and static code size by 43% for all programs.

Table 7.5 gives the execution time in microseconds of the benchmarks for the VLSI-BAM compiled without analysis (No Modes) and with analysis (Auto Modes). The last three columns give the ratios of the auto modes to the no modes times. To give an idea how built-ins affect the results of analysis, Table 7.5 gives two performance ratios for the large benchmarks: the first for all programs, and the second for
Table 7.5 - The effect of dataflow analysis on performance

<table>
<thead>
<tr>
<th>Benchmark</th>
<th>No Modes (μs)</th>
<th>Auto Modes (μs)</th>
<th>Auto/No Modes</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Time</td>
<td>Deref</td>
<td>Trail</td>
</tr>
<tr>
<td>deriv</td>
<td>146</td>
<td>18.2</td>
<td>5.5</td>
</tr>
<tr>
<td>log10</td>
<td>25.9</td>
<td>2.3</td>
<td>0.7</td>
</tr>
<tr>
<td>ops8</td>
<td>28.5</td>
<td>3.3</td>
<td>1.0</td>
</tr>
<tr>
<td>times10</td>
<td>39.7</td>
<td>5.1</td>
<td>1.3</td>
</tr>
<tr>
<td>divide10</td>
<td>51.7</td>
<td>7.5</td>
<td>2.5</td>
</tr>
<tr>
<td>nreverse</td>
<td>308</td>
<td>79.7</td>
<td>31.1</td>
</tr>
<tr>
<td>qsort</td>
<td>378</td>
<td>109</td>
<td>25.1</td>
</tr>
<tr>
<td>serialise</td>
<td>512</td>
<td>75.8</td>
<td>12.3</td>
</tr>
<tr>
<td>mu</td>
<td>992</td>
<td>154</td>
<td>48.0</td>
</tr>
<tr>
<td>fast_mu</td>
<td>1120</td>
<td>148</td>
<td>38.0</td>
</tr>
<tr>
<td>queens_8</td>
<td>1700</td>
<td>271</td>
<td>67.9</td>
</tr>
<tr>
<td>query</td>
<td>5180</td>
<td>560</td>
<td>174</td>
</tr>
<tr>
<td>tak</td>
<td>71700</td>
<td>13800</td>
<td>3180</td>
</tr>
<tr>
<td>poly_10</td>
<td>64000</td>
<td>6280</td>
<td>1740</td>
</tr>
<tr>
<td>zebra</td>
<td>84600</td>
<td>11400</td>
<td>8.6</td>
</tr>
<tr>
<td>average</td>
<td>666</td>
<td>29.2</td>
<td>19.0</td>
</tr>
<tr>
<td>average (no built-ins)</td>
<td>82</td>
<td>58.3</td>
<td>39.0</td>
</tr>
</tbody>
</table>

The time spent in dereferencing and trailing, two of the most common Prolog-specific operations, is significantly reduced by analysis. For the small benchmarks analysis reduces dereferencing from 17% to 5% of execution time, and trailing from 4% to 0.6% of execution time. This is because they are simple enough that analysis is able to deduce most relevant modes. For the large benchmarks dereferencing is reduced from 11% to 9% and trailing is reduced from 2.3% to 1.3%. These results are less extreme for two reasons: the large benchmarks use built-ins, which are unaffected by analysis, and the analyzer loses infor-
Table 7.6 - The effect of dataflow analysis on static code size

<table>
<thead>
<tr>
<th>Benchmark</th>
<th>No Modes (instructions)</th>
<th>Auto Modes (instructions)</th>
<th>Auto/No Modes</th>
</tr>
</thead>
<tbody>
<tr>
<td>tak</td>
<td>80</td>
<td>34</td>
<td>0.42</td>
</tr>
<tr>
<td>nreverse</td>
<td>287</td>
<td>139</td>
<td>0.48</td>
</tr>
<tr>
<td>queens_8</td>
<td>472</td>
<td>146</td>
<td>0.31</td>
</tr>
<tr>
<td>qsort</td>
<td>485</td>
<td>215</td>
<td>0.44</td>
</tr>
<tr>
<td>deriv</td>
<td>5891</td>
<td>1123</td>
<td>0.19</td>
</tr>
<tr>
<td>log10</td>
<td>1464</td>
<td>272</td>
<td></td>
</tr>
<tr>
<td>ops8</td>
<td>1469</td>
<td>277</td>
<td></td>
</tr>
<tr>
<td>times10</td>
<td>1479</td>
<td>287</td>
<td></td>
</tr>
<tr>
<td>divide10</td>
<td>1479</td>
<td>287</td>
<td></td>
</tr>
<tr>
<td>query</td>
<td>1425</td>
<td>403</td>
<td>0.28</td>
</tr>
<tr>
<td>serialisc</td>
<td>860</td>
<td>520</td>
<td>0.60</td>
</tr>
<tr>
<td>mu</td>
<td>1169</td>
<td>731</td>
<td>0.63</td>
</tr>
<tr>
<td>fast_mu</td>
<td>1165</td>
<td>718</td>
<td>0.62</td>
</tr>
<tr>
<td>zebra</td>
<td>1271</td>
<td>814</td>
<td>0.64</td>
</tr>
<tr>
<td>poly_10</td>
<td>3023</td>
<td>893</td>
<td>0.30</td>
</tr>
<tr>
<td>average</td>
<td></td>
<td></td>
<td>0.45</td>
</tr>
<tr>
<td>crypt</td>
<td>1239</td>
<td>1027</td>
<td>0.83</td>
</tr>
<tr>
<td>browse</td>
<td>1863</td>
<td>1150</td>
<td>0.62</td>
</tr>
<tr>
<td>prover</td>
<td>4395</td>
<td>1318</td>
<td>0.30</td>
</tr>
<tr>
<td>meta_qsort</td>
<td>2484</td>
<td>1424</td>
<td>0.57</td>
</tr>
<tr>
<td>flatten</td>
<td>4267</td>
<td>2335</td>
<td>0.55</td>
</tr>
<tr>
<td>unify</td>
<td>6326</td>
<td>4210</td>
<td>0.67</td>
</tr>
<tr>
<td>sdda</td>
<td>6526</td>
<td>5031</td>
<td>0.77</td>
</tr>
<tr>
<td>simple_analyzer</td>
<td>9057</td>
<td>5836</td>
<td>0.64</td>
</tr>
<tr>
<td>nand</td>
<td>23406</td>
<td>6654</td>
<td>0.28</td>
</tr>
<tr>
<td>reducer</td>
<td>11726</td>
<td>7682</td>
<td>0.66</td>
</tr>
<tr>
<td>boyer</td>
<td>24862</td>
<td>9136</td>
<td>0.37</td>
</tr>
<tr>
<td>chat_parser</td>
<td>33557</td>
<td>20516</td>
<td>0.61</td>
</tr>
<tr>
<td>average</td>
<td></td>
<td></td>
<td>0.57</td>
</tr>
</tbody>
</table>

Information due to its inability to handle aliasing and its limited type domain.

Table 7.6 gives the static code size (in VLSI-BAM instructions) for the benchmarks compiled without analysis (No Modes) and with analysis (Auto Modes). The effect of analysis on code size is greater than the effect on performance. This follows from the compiler's implementation of argument selection: when no modes are given, the compiler generates more code to handle arguments of different types. If analysis derives the type then the code becomes much smaller. The code size compares favorably with other symbolic processors, and is low enough that there is no disadvantage to having a simple instruction set. With the analyzer, code size on the VLSI-BAM is similar to the KCM [6], about three times the PLM, a micro-coded WAM [28], and about one fourth the SPUR using macro-expanded WAM [8].
Table 7.7 presents data about the efficiency of the dataflow analyzer. For each benchmark it gives the number of predicate arguments (Args) where a predicate of arity N is counted as N, the number of predicates (Preds), the analysis time (Time), the fraction of arguments that are uninitialized (uninit), ground (ground), nonvariable (nonvar), or recursively dereferenced (reref), and the fraction of arguments that have any of these types (any). Analysis time is measured under Quintus release 2.0 on a Sun 3/60. It is roughly proportional to the number of arguments in the program, except for the nand benchmark. The sum of the individual modes columns is usually greater than the any modes column. This is because arguments can have multiple modes—they can be both recursively dereferenced and ground or nonvariable. Uninitialized arguments are present in great quantities, even in large programs such as chat_parser and simple_analyzer. Comparing the small and large benchmarks, the fraction of derived modes decreases for

<table>
<thead>
<tr>
<th>Benchmark</th>
<th>Args</th>
<th>Preds</th>
<th>Time (sec)</th>
<th>Modes (fraction of arguments)</th>
<th>uninit</th>
<th>ground</th>
<th>nonvar</th>
<th>reref</th>
<th>any</th>
</tr>
</thead>
<tbody>
<tr>
<td>dcriv</td>
<td>12</td>
<td>8</td>
<td>11.9</td>
<td>0.33</td>
<td>0.67</td>
<td>0.00</td>
<td>0.67</td>
<td>1.00</td>
<td></td>
</tr>
<tr>
<td>log10</td>
<td>3</td>
<td>2</td>
<td>2.9</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>ops8</td>
<td>3</td>
<td>2</td>
<td>3.0</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>times10</td>
<td>3</td>
<td>2</td>
<td>3.0</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>divide10</td>
<td>3</td>
<td>2</td>
<td>2.9</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>tak</td>
<td>4</td>
<td>2</td>
<td>2.3</td>
<td>0.25</td>
<td>0.75</td>
<td>0.00</td>
<td>0.75</td>
<td>1.00</td>
<td></td>
</tr>
<tr>
<td>nreverse</td>
<td>5</td>
<td>3</td>
<td>2.2</td>
<td>0.40</td>
<td>0.60</td>
<td>0.00</td>
<td>0.60</td>
<td>1.00</td>
<td></td>
</tr>
<tr>
<td>qsort</td>
<td>7</td>
<td>3</td>
<td>3.4</td>
<td>0.43</td>
<td>0.57</td>
<td>0.00</td>
<td>0.57</td>
<td>1.00</td>
<td></td>
</tr>
<tr>
<td>query</td>
<td>7</td>
<td>5</td>
<td>4.2</td>
<td>0.86</td>
<td>0.14</td>
<td>0.00</td>
<td>0.14</td>
<td>1.00</td>
<td></td>
</tr>
<tr>
<td>zebra</td>
<td>10</td>
<td>6</td>
<td>3.5</td>
<td>0.10</td>
<td>0.00</td>
<td>0.50</td>
<td>0.00</td>
<td>0.60</td>
<td></td>
</tr>
<tr>
<td>serialise</td>
<td>16</td>
<td>7</td>
<td>4.2</td>
<td>0.38</td>
<td>0.19</td>
<td>0.06</td>
<td>0.19</td>
<td>0.63</td>
<td></td>
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<tr>
<td>queens_8</td>
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<td>7</td>
<td>6.0</td>
<td>0.31</td>
<td>0.69</td>
<td>0.00</td>
<td>0.69</td>
<td>1.00</td>
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<tr>
<td>mu</td>
<td>17</td>
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<td>9.6</td>
<td>0.12</td>
<td>0.47</td>
<td>0.00</td>
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<td>0.65</td>
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<tr>
<td>poly_10</td>
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<td>11</td>
<td>16</td>
<td>0.33</td>
<td>0.67</td>
<td>0.00</td>
<td>0.67</td>
<td>1.00</td>
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<td>0.05</td>
<td>0.55</td>
<td>0.89</td>
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</tr>
<tr>
<td>average</td>
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<td>0.35</td>
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<td>0.06</td>
<td>0.45</td>
<td>0.89</td>
<td></td>
</tr>
<tr>
<td>meta_qsort</td>
<td>10</td>
<td>7</td>
<td>11</td>
<td>0.30</td>
<td>0.00</td>
<td>0.10</td>
<td>0.00</td>
<td>0.40</td>
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<tr>
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<td>12</td>
<td>0.00</td>
<td>0.61</td>
<td>0.11</td>
<td>0.56</td>
<td>0.72</td>
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<tr>
<td>prover</td>
<td>22</td>
<td>9</td>
<td>13</td>
<td>0.27</td>
<td>0.09</td>
<td>0.27</td>
<td>0.14</td>
<td>0.68</td>
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<tr>
<td>browse</td>
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<td>14</td>
<td>20</td>
<td>0.24</td>
<td>0.45</td>
<td>0.05</td>
<td>0.40</td>
<td>0.74</td>
<td></td>
</tr>
<tr>
<td>boyer</td>
<td>62</td>
<td>25</td>
<td>31</td>
<td>0.27</td>
<td>0.00</td>
<td>0.06</td>
<td>0.00</td>
<td>0.34</td>
<td></td>
</tr>
<tr>
<td>flatten</td>
<td>83</td>
<td>28</td>
<td>34</td>
<td>0.27</td>
<td>0.08</td>
<td>0.16</td>
<td>0.11</td>
<td>0.52</td>
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<tr>
<td>sdda</td>
<td>87</td>
<td>32</td>
<td>45</td>
<td>0.18</td>
<td>0.07</td>
<td>0.17</td>
<td>0.08</td>
<td>0.44</td>
<td></td>
</tr>
<tr>
<td>reducer</td>
<td>134</td>
<td>41</td>
<td>50</td>
<td>0.13</td>
<td>0.10</td>
<td>0.05</td>
<td>0.12</td>
<td>0.29</td>
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<tr>
<td>unify</td>
<td>141</td>
<td>29</td>
<td>84</td>
<td>0.18</td>
<td>0.19</td>
<td>0.14</td>
<td>0.21</td>
<td>0.56</td>
<td></td>
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<tr>
<td>nand</td>
<td>180</td>
<td>43</td>
<td>5900</td>
<td>0.26</td>
<td>0.67</td>
<td>0.00</td>
<td>0.28</td>
<td>0.93</td>
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<td>simple_analyzer</td>
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<td>71</td>
<td>77</td>
<td>0.23</td>
<td>0.10</td>
<td>0.08</td>
<td>0.10</td>
<td>0.41</td>
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</tr>
<tr>
<td>chat_parser</td>
<td>744</td>
<td>156</td>
<td>263</td>
<td>0.44</td>
<td>0.19</td>
<td>0.02</td>
<td>0.09</td>
<td>0.67</td>
<td></td>
</tr>
<tr>
<td>average</td>
<td></td>
<td></td>
<td></td>
<td>0.23</td>
<td>0.21</td>
<td>0.10</td>
<td>0.17</td>
<td>0.56</td>
<td></td>
</tr>
</tbody>
</table>
the large programs for each type except nonvariable. For both the small and large benchmarks the analyzer transforms one third of the uninitialized modes into uninitialized register modes.

4. The effectiveness of the determinism transformation

To show what parts of the determinism transformation of Chapter 4 are the most effective, it is useful to define a spectrum of determinism extraction algorithms ranging from pure WAM to the full mechanism of the Aquarius compiler. To do this, the Aquarius mechanism for extracting determinism is divided into three orthogonal axes:

(1) The kind of tests used to extract determinism. These tests are separated into three classes: explicit unifications (e.g. \(x=a, x=s(Y)\)), arithmetic tests (e.g. \(x<y, x>1\)), and type checks (e.g. \(\text{var}(X), \text{atomic}(X)\)). Pure WAM uses only explicit unifications with nonvariables. Aquarius uses all three kinds.

(2) Which argument(s) are used to extract determinism. Pure WAM uses only the first argument of a predicate. Aquarius uses any argument that it can determine is effective. It uses enrichment heuristic 2 (Chapter 4 section 6.2).

(3) Whether the factoring transformation is performed (Chapter 4). Factoring significantly increases determinism for predicates that contain many identical compound terms in the head. Pure WAM does not assume factoring. Aquarius does factoring by default.

These three parameters define a three-dimensional space of determinism extraction algorithms. Each algorithm is characterized by a 3-tuple depending on its position on each of the axes (Table 7.8). This results in \(3 \times 2 \times 2 = 12\) data points. Pure WAM selection corresponds to the first element in each column, denoted by the 3-tuple (U, ONE, NF). The Aquarius compiler's selection corresponds to the last element in each column, denoted by the 3-tuple (UAT, ANY, F).

For each of these 12 points three parameters were measured: execution time, static code size, and compile time. All programs are compiled with dataflow analysis and executed on the VLSI-BAM. All averages are geometric means. It was only possible to do measurements for nine benchmarks: reverse, qsort, query, mu, fast_mu, queens_8, flatten, meta_qsort, and nand. Therefore the variance of the results is large and
Table 7.8 - Three dimensions of determinism extraction

<table>
<thead>
<tr>
<th>Kind of test</th>
<th>Which argument</th>
<th>Factoring</th>
</tr>
</thead>
<tbody>
<tr>
<td>Explicit unifications only (U).</td>
<td>First argument only (ONE).</td>
<td>No factoring (NF).</td>
</tr>
<tr>
<td>Explicit unifications and arithmetic tests (UA).</td>
<td>Any argument (ANY).</td>
<td>Do factoring (F).</td>
</tr>
<tr>
<td>Explicit unifications, arithmetic tests, and type checks (UAT).</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

they can be relied upon only to indicate trends. The benchmarks were written for the WAM. The measurements compare only the relative powers of different kinds of determinism extraction in the BAM. They do not compare the WAM and BAM directly.

Figure 7.1 - The effectiveness of determinism extraction

Figure 7.1 depicts the 12 points as a lattice. Each vertex denotes one particular combination of determinism extraction. The top element corresponds to Aquarius selection and the bottom element corresponds to WAM selection. Each edge connects two points that differ by one step in one coordinate. The vertices
are marked with the percent slowdown compared to Aquarius selection. The edges are marked with the percent difference in execution time between their two endpoints.

The mean speedup for the nine benchmarks when going from WAM selection (U, ONE, NF) to Aquarius selection (UAT, ANY, F) is 16%. There is no significant change in mean code size for any of the twelve data points. The variance of the compile time is too large to make any conclusions about it.

The mean speedup of factoring is 8%. However, factoring is the only transformation that sometimes slows down execution. The factoring heuristic should be refined to look inside compound arguments to see whether there is any potential determinism there. If there is none, it should not factor that argument.

One way of finding a set of effective extensions for determinism extraction is by traversing the lattice from bottom to top, and picking the edge with the greatest performance increase at each vertex. Starting at WAM selection (U, ONE, NF), the first extension is the ability to use arithmetic tests in selection. This speeds up execution by 3%. The second extension is the ability to select on any argument. This speeds up execution by another 3%. The third extension is the factoring transformation. This speeds up execution by 8%. At this point, the resulting performance is within 2% of Aquarius selection.

The resulting vertex (UA. ANY, F) seems to be a particularly good one, i.e. the ability to select on arithmetic tests in any argument works well together with factoring. Leaving out any one of these three extensions reduces performance by at least 8%. A plausible reason for this result is that the benchmarks do many arithmetic tests on the arguments of compound terms and it is only the combination of the three extensions that is able to compile this deterministically.

5. Prolog and C

The performance of Aquarius Prolog is significantly better than previous Prolog systems. A question one can pose is how the system compares with an implementation of an imperative language. This section presents a comparison of Prolog and the C language on several small programs. The comparison is not exhaustive—there are so many factors involved that I do not attempt to address this issue in its entirety. I intend only to dispel the notion that implementations of Prolog are inherently slow because of its expressive power. A serious comparison of two languages requires answering the following questions:
(1) How can implementations of different languages be compared fairly? This comparison concentrates exclusively on the language and ignores features external to the language itself, such as user interface, development time, and debugging abilities. One method is to pick problems to be solved, and then to write the "best" programs in each language to solve the problems, choosing the algorithms appropriate for each language. The disadvantages of this approach are (a) different languages are appropriate for different problems, (b) how does one decide when one has written the "best" program? To avoid these problems I have chosen to compare algorithms, not programs.

(2) Which algorithms will be implemented in both languages? Ideally one should select a range of algorithms, from those most suited to imperative computations (e.g. array computations) to those most suited to symbolic computation (e.g. large dynamic data objects, pattern matching). Prolog is at an advantage at the symbolic end of the spectrum because to implement symbolic computations in an imperative language we effectively have to implement more and more of a Prolog-like system in that language. The programmer does the work of a compiler. At the imperative end of the spectrum, the efficiency of Prolog depends strongly on the ability of the compiler to simplify its general features.

(3) What programming style will be used in coding the algorithms? I have made an attempt to program in a style which is acceptable for both languages. This includes choosing data types in both languages that are natural for each language. For example, in Prolog dynamic data accessed by pointers is easiest to express, whereas in C static arrays are easiest to express. It is possible to use dynamic data in C, but it requires more effort and is used only for those tasks that need it specifically.

(4) How are architectural features taken into account? For fairness both implementations should run on the same machine. The measurements use the same processor, the MIPS, for both implementations. However, a general-purpose architecture favors the execution of imperative languages, since it has been designed to execute such languages well. This shows up for algorithms whose Prolog implementation makes heavy use of Prolog-specific features. To allow the reader to make an informed judgment, the table does not correct for this fact. It is important to bear in mind that by adding additional architectural features comprising 5% of the chip area to the VLSI-BAM (a pipelined processor similar in many ways to the MIPS), the performance increases by 50% for programs that use
Prolog-specific features (compiled with the current version of the Aquarius compiler). Architectural studies done by our research group suggest that these features could be added to a future MIPS processor.

Table 7.9 compares the execution time of small algorithms coded in both C and Prolog on a 25 MHz MIPS processor. Measurements are given for tak, fib, and hanoi, which are recursion-intensive integer functions; and for quicksort, which sorts a 50 element list 10000 times. Prolog and C source code is available by anonymous ftp to arpa.berkeley.edu. In all cases the user time is measured with the Unix "time" utility. The C versions are compiled with the standard MIPS C compiler using both no optimization and the optimization level that produces the fastest code (usually level 4). The Prolog versions are compiled with dataflow analysis and translated into MIPS assembly by a partial translator. The same algorithms were encoded for both Prolog and C, in a natural style for each. The natural style in C is to use static data, whereas in Prolog all data is allocated dynamically.

<table>
<thead>
<tr>
<th>Benchmark</th>
<th>Aquarius</th>
<th>MIPS C</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Prolog</td>
<td>Unoptimized</td>
</tr>
<tr>
<td>tak(24,16,8)</td>
<td>1.2</td>
<td>2.1</td>
</tr>
<tr>
<td>fib(30)</td>
<td>1.5</td>
<td>2.0</td>
</tr>
<tr>
<td>hanoi(20,1,2,3)</td>
<td>1.3</td>
<td>1.6</td>
</tr>
<tr>
<td>quicksort</td>
<td>2.8</td>
<td>3.3</td>
</tr>
</tbody>
</table>

Recursive functions are fast in Prolog for three reasons: last call optimization converts recursion into iteration, environments (stack frames) are allocated per clause and not per procedure as in C, and outputs are returned in registers (they are of uninitialized register type). Last call optimization allows functions with a single recursive call to execute with constant stack space. This is essential for Prolog because recursion is its only looping construct. The MIPS C compiler does not do last call optimization. C has constructs to denote iteration explicitly (e.g. "for" and "while" loops) so it does not need this optimization as strongly. The time for fib(30), the only recursive integer function that is not able to use last call optimization in Prolog, is closest to C.

The two quicksort implementations are careful to use the same pivot elements. The C implementation uses an array of integers and does in-place sorting. The Prolog implementation uses lists and creates a new sorted list. The list representation needs two words to store each data element. Coincidentally, the
Prolog version is twice as slow as the C version, the same as the ratio of the data sizes.

<table>
<thead>
<tr>
<th>Kind</th>
<th>Description</th>
<th>%</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mistake</td>
<td>A part of the compiler that is incorrect due to an oversight. When many mistakes occur related to one particular area, then they become hotspot bugs.</td>
<td>39</td>
</tr>
<tr>
<td>Local</td>
<td>A problem that can be fixed by changing just a few predicates. For example, it may be due to a typographical error or a simple oversight in a predicate definition.</td>
<td>(37)</td>
</tr>
<tr>
<td>Global</td>
<td>A problem that can be fixed only with many changes throughout the compiler. This kind of mistake is more fundamental. For example, avoiding the generation of BAM instructions, with double indirections requires many small changes.</td>
<td>(3)</td>
</tr>
<tr>
<td>Incomplete</td>
<td>A part of the compiler whose first implementation is incomplete because of incomplete understanding of its purpose. Later use stretches it beyond what it was intended to do, so that it needs to be extended and/or cleaned up. For example, the updating of type formulas when new information is given.</td>
<td>19</td>
</tr>
<tr>
<td>Hotspot</td>
<td>A critical area of the compiler that requires much thinking to get correct. Its importance is much greater than its size would indicate. Such an area gets more than its share of mistakes.</td>
<td>16</td>
</tr>
<tr>
<td>Conceptual</td>
<td>A concept in the compiler design whose implementation is prone to many mistakes. For example, the concept of uninitialized variables.</td>
<td>(13)</td>
</tr>
<tr>
<td>Physical</td>
<td>A part of the compiler's text. For example, symbolic unification in the dataflow analyzer and parameter passing in the clause compiler both resulted in many bugs.</td>
<td>(14)</td>
</tr>
<tr>
<td>Mixture</td>
<td>An undesired interaction between separate parts of the compiler. Despite careful design, often the separate transformations and optimizations are not completely orthogonal, but interact in some (usually limited) way. For example, maintaining consistency between the dataflow analyzer and the clause compiler. This leads to the dereference chain transformation, which in turn leads to the problem of interaction between it and the preferred register allocation.</td>
<td>16</td>
</tr>
<tr>
<td>Improvement</td>
<td>A possible improvement in the compiler. This is not strictly a bug, but it may point to an important optimization that could be added to the compiler. For example, a possible code optimization or reduction in compilation time.</td>
<td>9</td>
</tr>
<tr>
<td>Understanding</td>
<td>A problem due to the programmer misunderstanding the required input to the compiler. This is not strictly a bug, but it may point to difficulties in the compiler's user interface or in the language. For example, the difference between the terms <em>is</em> and <em>&lt;</em> in Prolog. The first is a variable and the second is a structure.</td>
<td>4</td>
</tr>
</tbody>
</table>

6. Bug analysis

This section gives an overview of the number and types of bugs encountered during compiler development. A bug in a program is a problem that leads to incorrect or undesired behavior of the program. In the compiler, this means incorrect or slow compilation, or slow execution of compiled code.
Table 7.10 classifies the bugs found during development [76]. (The percentages do not add up to 100% because bugs can be of more than one type.)

The development of the compiler started early 1988 and proceeded until late 1990. An extensive suite of test programs was maintained to validate versions of the compiler. The test suite was continually extended with programs that resulted in bugs and with programs from external sources. Records were kept of all bugs reported by users of the compiler other than the developer. A total of 79 bug reports were sent from January 1989 to August 1990 by five users. The frequency of bug reports stayed constant near four per month. Statistical analysis is consistent with the distribution being random with no time dependence, i.e. the number of bug reports fluctuates, but there is no increasing or decreasing trend. Therefore the development introduced bugs at about the same rate as they were reported and fixed. This coincidence can be explained by postulating that the time spent developing was limited by the necessity of having to spend time debugging to maintain a minimum level of robustness in the compiler. This is consistent with my personal experience during the development process.
Chapter 8
Concluding Remarks and Future Work

"So many things are possible just as long as you don't know they're impossible."
-Norton Juster, The Phantom Tollbooth

1. Introduction

In this chapter I recapitulate the main result of this dissertation, distill some practical lessons learned in the design process, talk about the caveats of language design, and give directions for future research.

2. Main result

My thesis is that logic programming can execute as fast as imperative programming. For this purpose I have implemented a new optimizing Prolog compiler, the Aquarius compiler. The driving force in the compiler is to specialize the general mechanisms of Prolog (i.e. the logical variable, unification, dynamic typing, and backtracking) as much as possible. The main ideas in the compiler are: the development of a new abstract machine that allows more optimization, a mechanism to generate efficient code for deterministic predicates (converting backtracking to conditional branching), specialization of unification (encoding each occurrence of unification in the simplest possible way), and the use of global dataflow analysis to derive types.

The resulting system is significantly faster than previous implementations and is competitive with C on programs for which dataflow analysis is able to do sufficiently well. It is about five times faster than Quintus Prolog, a popular commercial implementation.

3. Practical lessons

During the design of this compiler I have found four principles useful.

(1) Simplicity is common. Most of the time, only simple cases of the general mechanisms of the language are used. For example, most uses of unification are memory loads and stores. Many of these simple cases are easily detected at compile-time.
(2) **Use the design time wisely.** There are many possible optimizations that one can implement in a compiler of this sort. To get the best results, rank them according to their estimated performance gain relative to their implementation effort, and only implement the best ones. Do not be distracted by clever ideas unless you can prove that they are effective.

(3) **Keep the design simple.** For each optimization or transformation, implement the simplest version that will do the job. Do not attempt to implement a more general version unless it can be done without any extra effort. It is easy to become entangled in the mechanics of implementing a complex optimization. Often a simple version of this optimization achieves most of the benefits in a fraction of the time.

(4) **Document everything, including bugs.** Documentation is an extension to one’s memory and it pays for itself quickly. The mental effort spent in writing down what one has done results in a better recollection of what happened. In this design, I have maintained two logs. The first is a file in chronological order that documents each change and the reason for it. The second is a directory containing bug reports contributed by the users of the compiler and brief discussions of the fixes.

The first three of these principles are corollaries of what is sometimes called the "80-20 rule": 80% of the results are obtained with 20% of the effort. Using this principle consistently was very important for my work and for the BAM project as a whole.

4. **Language design**

The Prolog language is only an approximation to the ideal of logic programming. During this research, our group has grappled with some of the deficiencies of Prolog. There are deficiencies in the area of logic: Prolog’s approximation to negation (i.e. negation-as-failure) is unsound (i.e. it gives incorrect results) when used in the wrong way. Prolog’s implementation of unification can go into infinite loops when creating circular terms. The default control flow is too rigid for data-driven programming.

There are deficiencies in the area of programming: The correspondence between a program and its execution efficiency is not always obvious. Unification is only able to access the surface of a complex data structure. Because the clauses of a predicate are written separately, many conditions have to be repeated or
extra predicates have to be defined. There is a sense in which Prolog is a kind of assembly language.

All of the above problems have solutions, some of which have been implemented in existing systems and in the Aquarius system. However, for three reasons I have resisted the impulse to change the language more than just a little. First, of all logic languages, the Prolog language has the largest and most vigorous user community, and this is a resource I wanted to tap. There are many programs written in Prolog, in various styles, and I wanted to see if this existing pool of ingenuity could be made to run faster. Second, it is unwise to change more than one component of a system at the same time, especially if they can interact in unpredictable ways. That is, one should not design a new language and a compiler for it at the same time. Third, I do not deem myself competent yet to design a language. I believe in the rule of bootstrapped competence: Before writing a compiler, write programs. Before designing a language, write compilers. Competence in each task is limited by competence in its prerequisite.

The best languages are those which distill great power in a small set of features. This makes such languages useful as tools for thought as well as for implementation. Practical aspects such as how efficient it can be implemented are as important in a good language design as theoretical aspects. A good language is theoretically clean (i.e. easily understood) as well as being efficiently implementable. Examples of such languages are Pascal (many algorithms are specified in an Pascal-like pseudo-code), Scheme, and Prolog. To create such a language, a person must have completely digested a set of ideas as well as have a large amount of practical experience. This is a difficult combination—it is easy to gloss over the areas one does not know well.

5. Future work

The goal of achieving parity with imperative languages has been achieved for the class of programs for which dataflow analysis is able to provide sufficient information, and for which the determinism is accessible through built-in predicates. To further improve performance these limits must be addressed.

To guide the removal of these limits it is important to build large applications and study the interaction between programming style and the implementation. This is a problem of successive refinement. A more sophisticated implementation catalyzes a new style of programming, which in its turn catalyzes a new
implementation, and so forth. The first step in this process was the development of the first Prolog compiler and the WAM. The Aquarius system is only the second step. It is able to generate efficient code from programs written in a more logical style than standard Prolog. However, the limits of this style are not yet understood as they are in the WAM. Further work in this area will lead to a successor to Prolog that is closer to logic and also efficiently implementable.

5.1. Dataflow analysis

When writing a program, a programmer commonly has a definite intention about the data's type (intending predicates to be called only in certain ways) and about the data's lifetime (intending data to be used only for a limited period). Because of this consistency, I postulate that a dataflow analyzer should be able to derive this information and a compiler should be able to take advantage of it.

There has been much good theoretical work on global analysis for Prolog, but few implementations, and fewer still that are part of a compiler that takes advantage of the information. Measurements of the Aquarius system show that a simple dataflow analysis scheme integrated into a compiler is already quite useful. However, the implementation has been restricted in several ways to make it practical. As programs become larger, these restrictions limit the quality of the results. I hope the success of this experiment encourages others to relax these restrictions. For example, it would not be too difficult to:

- Extend the domain to represent common types such as integers, proper lists, and nested compound terms. This is especially important for general-purpose processors.
- Extend the domain to represent variable aliasing explicitly. This avoids the loss of information that affects the analyzer.
- Extend the domain to represent data lifetimes. This is useful to replace copying of compound terms by in-place destructive assignment. In this way dynamically allocated data becomes static. The term "compile-time garbage collection" that has been used to describe this process is a misnomer; what is desired is not just memory recovery, but to preserve as much as possible of the old value of the compound term. Often a new compound term similar to the old one is created at the same time the old one becomes inaccessible. Destructive assignment is used to modify only those parts that are
• Extend the domain to represent types for each invocation of a predicate. For example, the analyzer could keep track not only of argument types for predicate definitions, but of argument types for goals inside the definitions. This is useful to implement multiple specialization, i.e. to make separate copies of a predicate called in several places with different types. For the chat-parser benchmark, making a separate copy of the most-used predicate for each invocation results in a performance improvement of 14%.

5.2. Determinism

The second area in which significant improvement is possible is determinism extraction. The Aquarius compiler only recognizes determinism in built-in predicates of three kinds (unification, arithmetic tests, and type checking). Often this is not enough. In many programs, user-defined predicates are used to choose a clause.
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Appendix A
User manual for the Aquarius Prolog compiler

1. Introduction

The Aquarius Prolog compiler reads clauses and directives from stdin and outputs Prolog-readable compiled code to stdout as one fact per instruction. The output is assembly code for the Berkeley Abstract Machine (BAM). Directives hold starting from the next predicate that is input. Clauses do not have to be contiguous in the input stream, however, the whole stream is read before compilation starts.

This manual is organized into ten sections. Section 2 documents the compiler's directives. Section 3 gives the compiler's options. Section 4 gives a short overview of the dataflow analysis done by the compiler. Section 5 gives the type declarations accepted by the compiler. Section 6 summarizes the differences between Aquarius Prolog and the Edinburgh standard. Section 7 gives an example showing how to use the compiler. Section 8 describes the method used to compile specialized entry points to increase the efficiency of built-ins. Section 9 describes the assembly language interface. Section 10 describes how to define BAM assembly macros.

2. Directives
The directives recognized by the Aquarius compiler are given in Table 1.

3. Options
The Aquarius compiler's options are given in three categories: high-level (these options control actions of the compiler at the Prolog level), architecture-dependent (these options are constant for a particular architecture), and low-level (mainly useful for debugging purposes). The default options are set for the VLSI-BAM processor. The options are given in Tables 2, 3, and 4.

4. Dataflow analysis
Dataflow analysis is enabled with the analyze option. It generates ground, nonvar, recursively dereferenced and uninitialized variable types which are merged with the programmer's types. Both uninitialized memory and uninitialized register types are generated. Entry declarations (given by entry directives) are used to drive the analysis. Predicates of arity zero are always used as entry declarations. The quality of the generated types is such that compilation time, execution time, and code size are all significantly reduced. Therefore it is recommended always to compile with analysis. The whole program is kept in memory during the analysis.

All mode, entry, and op directives are executed before the analysis starts. Other directives are executed after the analysis and before compilation. The directives default and clear interfere with dataflow analysis, so they should be given only when the analyze option is disabled.

4.1. Dataflow analysis and dynamic code
The compiler makes the distinction between static and dynamic code. Static code is completely known at compile-time and is subject to analysis. Dynamic code is created at run-time by the built-in predicates assert/1, retract/1, and their cousins. It is not analyzed. There are two cases to consider:

(1) A dynamic predicate calls a static predicate. In this case, there must be an entry declaration giving the worst-case type of the call for each static predicate that might be called by a dynamic predicate.
Leaving out this declaration may result in incorrect compilation.

(2) A static predicate calls a dynamic predicate. The analyzer will assume worst-case types for the dynamic predicate unless it has a type declaration.

The most common uses of dynamic code are as databases of facts, or as rules that only call a limited set of static predicates. For these uses, there is no problem in integrating analyzed static code with dynamic code.

4.2. Dataflow analysis and the call/1 built-in

The call/1 built-in predicate can call any predicate in the program with any modes, and it is not possible in general to determine these predicates and their modes at compile-time. However, most programs that use call/1 will call one of a known set of predicates or will call a dynamic predicate. There are three cases to consider:

(1) If the set of predicates that may be arguments of call/1 is known by the programmer, then these predicates should be given entry declarations with worst-case modes. (This case can be written more efficiently by writing a new predicate that directly calls one of the set, and avoids calling call/1.)

(2) If the predicates that may be arguments of call/1 are dynamic, then analysis is correct without entry declarations. This is true because dynamic predicates are not analyzed.

(3) If any predicate in the program may be an argument of call/1 and nothing is known about the modes then analysis is useless and it should not be done.

5. Types

The Aquarius compiler accepts type declarations for a predicate. Using types results in a significant improvement in code quality. Types are represented as (Head:-Formula) where Head contains only variables and Formula is a logical conjunction. Almost any Prolog test can be used in a type formula. Possible type formulas are given in Table 5. This representation for types is simple, yet powerful enough to represent much important information in a compact way. The representation generalizes the declarations of Dec-10 Prolog. For example, the Dec-10 declaration:

\[
:- \text{mode}(\text{concat}(+,+,\cdot))
\]

is expressed here as:

\[
:- \text{mode}((\text{concat}(A,B,C) :- \text{nonvar}(A), \text{nonvar}(B), \text{var}(C))).
\]

6. Differences with Edinburgh Prolog

Aquarius Prolog recognizes new type-checking built-ins which are not part of the Edinburgh Prolog standard as embodied by C-Prolog. The new built-ins and their definitions in standard Prolog are given in Table 6.

7. An example of the compiler’s use

The following example shows how the compiler is used:
% Run the compiler.
% Code is entered directly.
% Enter the type.
% Enter a simple two-fact predicate.
% End-of-file.
% The output follows:

Run the compiler.
Code is entered directly.
Enter the type.
Enter a simple two-fact predicate.
End-of-file.
The output follows:

% Cputime between start and finish is 1.383

procedure(a/1).
deref(r(0),r(0)).
hash(atomic,r(0),2,l(a/1,1)).
fail.
label(l(a/1,1)).
pragma(hash_length(2)).
pair(a,l(a/1,3)).
pair(b,l(a/1,4)).
label(l(a/1,3)).
label(l(a/1,4)).
return.

8. Entry specialization for more efficient built-ins

The directive modal_entry(Head,EntryTree) adds a discrimination tree of entry points for the predicate Head. This directive is used by the system to implement more efficient built-ins. It is not normally needed by programmers, although they can take advantage of it for other predicates. The compiler uses the discrimination tree to choose the most efficient entry point for each call of a predicate depending on the type formula that is true at the predicate's call. The syntax of the discrimination tree in modal_entry is:

tree(entry(EntryHead)).
tree(mode(Formula,TrueTree,FalseTree)) :-
tree(TrueTree), tree(FalseTree).

EntryHead is the entry point that replaces Head and Formula is a type formula. Compilation of a the predicate Head proceeds by following a path down the discrimination tree. If the formula valid when Head is called implies Formula then the TrueTree is followed. Otherwise the FalseTree is followed. Tree traversal stops when an entry point entry(EntryHead) is encountered. At that point the original call is replaced by EntryHead.

9. Interfacing with BAM assembly language routines

Prolog predicates can efficiently call routines written in BAM assembly code (the compiler's output) or in the target machine's assembly language (for example, VLSI-BAM, MIPS, or MC68020 assembly code). The interface with both low-level languages is provided through the five-argument type declaration. This declaration has the following form:

:- mode(Head, Require, Before, After, Survive).

Head is the head of the predicate. Require is the required type formula, i.e. the formula made true by the compiler. All uninitialized variable types (both uninitialized memory and uninitialized register) must be part of the required formula. Before is the type formula known to be valid before the call. After is the type formula known to be valid after the call. Survive is the register survive flag. If the flag is y then the predicate must not alter the values of any argument registers (except those used to return a result). It must save and restore any argument registers it needs. The predicate is called with a simple_call instruction and must return with a simple_return instruction (or its equivalent in
VLSI-BAM processor assembly). A simple call may not be nested. It is more efficient than a standard call because it does not need an environment frame around it in the calling routine.

If the survive flag is \textit{true} then the predicate is assumed to invalidate all argument register values. In this case the argument registers are available as scratch registers and the calling routine will create an environment frame.

Efficient parameter passing is implemented by using uninitialized variables. These are of two kinds: uninitialized memory and uninitialized register variables. An uninitialized memory variable is a pointer to an empty memory cell. Binding to it is a store to memory. An uninitialized register variable is an empty register. Binding to it is a move to the register. No trailing or dereferencing is needed in either case.

Declaring an argument to have a uninitialized register type means that the output of the routine is stored in the corresponding argument register. Similarly, an uninitialized memory type requires the output to be stored to the location pointed to by the argument register. Inputs and outputs must be put in separate registers.

10. Defining BAM assembly language macros

It is possible to define macros in the Prolog source that are expanded into BAM assembly instructions. The advantages of macros are that they do not have call-return overhead, that unnecessary shuffling of data between registers is avoided, and that the full range of low-level compiler optimizations is performed on them. A macro definition has the following form:

\begin{verbatim}
:- macro((Head :- Body)).
\end{verbatim}

where \textit{Head} is the head of the predicate that will be expanded and \textit{Body} is a series of BAM instructions. For example:

\begin{verbatim}
:- mode(quad(A,B), uninit_reg(B), true, deref(B), y).
:- macro((quad(A,B) :- add(A,A,X), add(X,X,B))).
\end{verbatim}

The macro definition is preceded by a mode declaration telling that the second argument is the output.

Macro definitions must obey the following rules:

1) All legal BAM instructions and addressing modes are allowed in the macro definition including user instructions, except as noted below. User instructions are never generated by the compiler, but they are recognized and optimized in macro definitions. Labels are given as ground terms or as Prolog variables. The latter are given unique ground values by the compiler. Registers are given as user registers (e.g. \texttt{r(h)} and \texttt{r(t2)}) or as Prolog variables (e.g. \texttt{X} and \texttt{Y}). The latter are allocated by the compiler. Do not use numbered registers (\texttt{r(0)}, \texttt{r(1)}, ...).

2) The macro definition must be preceded by a mode declaration. The exit modes must be valid upon exiting the macro. All head arguments that return results must be of uninitialized register type.

3) The macro may not alter any of the head arguments except those returning a result.

4) The second argument of the \texttt{deref(X,Y)} instruction must be a new variable, i.e. it must not have a value upon entering the macro. Failing to obey this constraint will lead to incorrect behavior on backtracking.

5) It is not recommended to create choice points inside macros since it is not known how many registers are live.
<table>
<thead>
<tr>
<th>Directive</th>
<th>Action</th>
</tr>
</thead>
<tbody>
<tr>
<td>:- help.</td>
<td>Print a summary of these directives.</td>
</tr>
<tr>
<td>:- default.</td>
<td>Set the default options for the VLSI-BAM processor and clear all type declarations and modal entries.</td>
</tr>
<tr>
<td>:- mips.</td>
<td>Ensure compatibility with the MIPS processor. This directive should occur only once in a file. It sets the option align(1), disables the option split_integer, and sets all other options to their default values. It clears all type declarations and modal entries.</td>
</tr>
<tr>
<td>:- vlsi_plm.</td>
<td>Ensure compatibility with the re-microcoded VLSI-PLM. This directive should occur only once in a file. It sets the options high_reg(6) and align(1), disables the option split_integer, and sets all other options to their default values. It clears all type declarations and modal entries. Trail checks and shifts are compiled differently.</td>
</tr>
<tr>
<td>:- clear.</td>
<td>Clear all type declarations and modal entries.</td>
</tr>
<tr>
<td>:- option(Options)</td>
<td>Add the options in Options to the current options. Options may be a single option or a list of options.</td>
</tr>
<tr>
<td>:- notooption(Options)</td>
<td>Remove the options in Options from the current options. Options may be a single option or a list of options.</td>
</tr>
<tr>
<td>:- printoption.</td>
<td>Print a list of the currently active options.</td>
</tr>
<tr>
<td>:- mode((Head:-Formula)).</td>
<td>Type declaration for a predicate. The type information is remembered until new types are given for that predicate or until all type information is cleared. This declaration is not used as a starting point for dataflow analysis. However, the types generated by dataflow analysis are used to supplement the declaration, and an error message is given if there is a contradiction. Type declaration for a predicate—same as above. This declaration is also used as a starting point in dataflow analysis. Detailed type declaration for a predicate. This declaration is useful for interfacing with assembly language. H is the head, R is the required type formula (made true by the compiler before each call), B is the before type formula (assumed true before each call), A is the after type formula (assumed true after each call), S is the survive flag (y/n depending on whether the call lets registers survive). The after type formula is used by dataflow analysis to improve the generated types. Detailed type declaration for a predicate—same as above. This declaration is also used as a starting point in dataflow analysis. Optional discrimination tree of efficient entry points for the predicate H. The tree T contains type formulas used to replace each call of the predicate by a more efficient entry point. Macro definition. The head is expanded into a sequence of BAM assembly instructions. Insert the text of the file FileName. This directive may be nested up to the system limit of simultaneous open files. Pass the input &quot;:- pass (Anything).&quot; unaltered to the output in Prolog-readable form. Print the creation date of this version of the compiler. Declare an operator in Prolog. Pass the input &quot;:- op (A, B, C).&quot; unaltered to the output in Prolog-readable form.</td>
</tr>
</tbody>
</table>
### Table 2 – High-level compiler options

<table>
<thead>
<tr>
<th>Option</th>
<th>Default</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>select_limit(L)</td>
<td>L=1</td>
<td>Perform selection for up to L arguments. Selection is done according to the enrichment heuristic. See Chapter 4 section 6.2.</td>
</tr>
<tr>
<td>analyze</td>
<td>off</td>
<td>Perform dataflow analysis for all predicates in the input stream. This option enables analysis of the entire input stream, no matter where it occurs in the stream. The starting points for analysis are the entry declarations and all predicates of arity zero. The types obtained from the analysis are merged with the programmer's types. The predicates are then compiled with the merged types.</td>
</tr>
<tr>
<td>compile</td>
<td>on</td>
<td>Compile the input. When this option is disabled, the entry types generated by the dataflow analyzer for the source predicates are output as valid Prolog-readable type declarations.</td>
</tr>
<tr>
<td>factor</td>
<td>on</td>
<td>Do factoring source transformation. With this transformation similar compound terms in adjacent heads are only unified once. Often this gives faster code.</td>
</tr>
<tr>
<td>comment</td>
<td>on</td>
<td>Give information about what the compiler is doing.</td>
</tr>
<tr>
<td>same_number_solutions</td>
<td>on</td>
<td>Keep the same number of solutions on backtracking as standard Prolog. Relaxing the semantics by removing this option results in better code in some cases.</td>
</tr>
<tr>
<td>same_order_solutions</td>
<td>on</td>
<td>Keep the same order of solutions on backtracking as standard Prolog. Relaxing the semantics by removing this option results in better code in some cases.</td>
</tr>
<tr>
<td>depth_limit(D)</td>
<td>D=2</td>
<td>Nesting depth limit on unification goals. Unifications deeper than this limit are transformed to remain within this limit. This transformation is used because compilation time and code size for deeply nested unifications would otherwise increase as the square of the size of the unification.</td>
</tr>
<tr>
<td>short_block(S)</td>
<td>S=6</td>
<td>Threshold on basic block length for shuffle optimization.</td>
</tr>
</tbody>
</table>

### Table 3 – Architecture-dependent compiler options

<table>
<thead>
<tr>
<th>Option</th>
<th>Default</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>low_reg(L)</td>
<td>L=0</td>
<td>Lowest numbered machine register.</td>
</tr>
<tr>
<td>high_reg(H)</td>
<td>H=100</td>
<td>Highest numbered machine register. In the VLSI-BAM processor, registers higher than r(15) are mapped into memory.</td>
</tr>
<tr>
<td>low_perm(P)</td>
<td>P=0</td>
<td>Lowest numbered permanent variable.</td>
</tr>
<tr>
<td>hash_size(H)</td>
<td>H=5</td>
<td>Minimum size of a hash table.</td>
</tr>
<tr>
<td>align(K)</td>
<td>K=2</td>
<td>Align all compound terms to start on a multiple of K.</td>
</tr>
<tr>
<td>uni</td>
<td>on</td>
<td>Generate unify_atomic instruction to unify with an atomic term.</td>
</tr>
<tr>
<td>split_integer</td>
<td>on</td>
<td>Use separate tags for negative and nonnegative integers.</td>
</tr>
<tr>
<td>Option</td>
<td>Default</td>
<td>Description</td>
</tr>
<tr>
<td>----------</td>
<td>---------</td>
<td>-----------------------------------------------------------------------------</td>
</tr>
<tr>
<td>system(X)</td>
<td>quintus</td>
<td>The system running the compiler (other value: cprolog).</td>
</tr>
<tr>
<td>write</td>
<td>on</td>
<td>Write the object code when compilation is complete.</td>
</tr>
<tr>
<td>peep</td>
<td>on</td>
<td>Do peephole optimization.</td>
</tr>
<tr>
<td>stats(S)</td>
<td>off</td>
<td>Print timing statistics during compilation. S is one of the following atoms, or a list of them: t (top level of compilation), c (compilation of a single procedure), p (peephole optimization), s (selection algorithm—extraction of determinism), d (deterministic code generation).</td>
</tr>
<tr>
<td>debug</td>
<td>off</td>
<td>Print debugging messages during compilation.</td>
</tr>
</tbody>
</table>

---

<table>
<thead>
<tr>
<th>Type</th>
<th>Meaning</th>
</tr>
</thead>
<tbody>
<tr>
<td>nonvar(A')</td>
<td>A is a nonvariable term, i.e. its main functor is instantiated. Nothing is implied about its arguments.</td>
</tr>
<tr>
<td>ground(A),</td>
<td>A is a ground term, i.e. it contains no unbound variables.</td>
</tr>
<tr>
<td>var(A)</td>
<td>A is an unbound variable.</td>
</tr>
<tr>
<td>uninit(A)</td>
<td>A is an uninitialized memory variable. At the Prolog level, this means that A is an unbound variable known not to be aliased to another variable. In the implementation, A is a pointer to an empty memory cell. Binding to this variable is a simple store, without dereferencing or trailing.</td>
</tr>
<tr>
<td>uninit_reg(A)</td>
<td>A is an uninitialized register variable. At the Prolog level, this has the same meaning as an uninitialized memory variable. In the implementation, A is an empty machine register. This type increases the efficiency of parameter passing by returning a value directly in a register. It is useful for interfacing with assembly language.</td>
</tr>
<tr>
<td>deref(A)</td>
<td>A is dereferenced.</td>
</tr>
<tr>
<td>rderef(A)</td>
<td>A is recursively dereferenced, i.e. A is dereferenced and all subterms of A are recursively dereferenced.</td>
</tr>
<tr>
<td>structure(A)</td>
<td>A is a structure.</td>
</tr>
<tr>
<td>list(A)</td>
<td>A is a list, i.e. a cons cell or nil.</td>
</tr>
<tr>
<td>cons(A)</td>
<td>A is a cons cell, i.e. a non-nil list.</td>
</tr>
<tr>
<td>compound(A)</td>
<td>A is a structure or a cons cell.</td>
</tr>
<tr>
<td>functor(A,F,N)</td>
<td>A is the structure F with arity N.</td>
</tr>
<tr>
<td>atom(A)</td>
<td>A is an atom.</td>
</tr>
<tr>
<td>atomic(A)</td>
<td>A is atomic, i.e. a number or an atom.</td>
</tr>
<tr>
<td>simple(A)</td>
<td>A is atomic or an unbound variable.</td>
</tr>
<tr>
<td>integer(A)</td>
<td>A is an integer.</td>
</tr>
<tr>
<td>float(A)</td>
<td>A is a floating point number.</td>
</tr>
<tr>
<td>number(A)</td>
<td>A is an integer or a float.</td>
</tr>
<tr>
<td>negative(A)</td>
<td>A is a negative integer.</td>
</tr>
<tr>
<td>nonnegative(A)</td>
<td>A is a nonnegative integer.</td>
</tr>
<tr>
<td>A&gt;0</td>
<td>A is a positive integer.</td>
</tr>
<tr>
<td>A==x</td>
<td>A is the atom x.</td>
</tr>
<tr>
<td>true</td>
<td>Nothing is known about the type.</td>
</tr>
<tr>
<td>fail</td>
<td>This means &quot;execution can never reach this point.&quot;</td>
</tr>
<tr>
<td>(F1,F2)</td>
<td>This means &quot;F1 and F2,&quot; where F1 and F2 are type formulas.</td>
</tr>
<tr>
<td>(F1,F2)</td>
<td>This means &quot;F1 or F2,&quot; where F1 and F2 are type formulas.</td>
</tr>
<tr>
<td>not (F)</td>
<td>This means &quot;not F,&quot; where F is a type formula.</td>
</tr>
</tbody>
</table>
Table 6 – New type-checking predicates in Aquarius Prolog

<table>
<thead>
<tr>
<th>Predicate</th>
<th>Prolog Definition</th>
</tr>
</thead>
<tbody>
<tr>
<td>nil(A)</td>
<td>:- nonvar(A), A=[].</td>
</tr>
<tr>
<td>cons(A)</td>
<td>:- nonvar(A), A=[<em>1</em>].</td>
</tr>
<tr>
<td>list(A)</td>
<td>:- nonvar(A), (A=[] ; A=[<em>!</em>]).</td>
</tr>
<tr>
<td>compound(A)</td>
<td>:- nonvar(A), +atomic(A).</td>
</tr>
<tr>
<td>structure(A)</td>
<td>:- nonvar(A), +atomic(A), +A=[<em>1</em>].</td>
</tr>
<tr>
<td>ground(A)</td>
<td>:- nonvar(A), functor(A, _, N), ground(N, A).</td>
</tr>
<tr>
<td>simple(A)</td>
<td>:- (var(A) ; atomic(A)).</td>
</tr>
<tr>
<td>negative(A)</td>
<td>:- integer(A), A&lt;0.</td>
</tr>
<tr>
<td>nonnegative(A)</td>
<td>:- integer(A), A&gt;=0.</td>
</tr>
<tr>
<td>is_list(A)</td>
<td>:- (var(A), ! ; A=[] ; A=_[B], is_list(B)).</td>
</tr>
<tr>
<td>is_partial_list(A)</td>
<td>:- (var(A), ! ; A=_[B], is_partial_list(B)).</td>
</tr>
<tr>
<td>is_proper_list(A)</td>
<td>:- (var(A), !, fail;A=[];A=_[B], is_proper_list(B)).</td>
</tr>
</tbody>
</table>

The following clauses are part of the definition:

```prolog
ground(N, _) :- N=:=0.
ground(N, A) :- N\=\=0, arg(N, A, X), ground(X), N1 is N-1, ground(N1, A).
```
Appendix B

Formal specification of the Berkeley Abstract Machine syntax

%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%
% Formal specification of the Berkeley Abstract Machine (BAM) syntax
% Copyright (C) 1989 Peter Van Roy and Regents of the University of California
% May be used and modified for non-commercial purposes if this notice is kept.
% Written by Peter Van Roy.

% This file is an executable Prolog program that checks the syntactic
% correctness of BAM instructions. The predicate instr(I) is true if I is
% a legal BAM instruction. In addition to instructions output by the Aquarius
% compiler, this predicate also accepts the user instructions of the BAM,
% which allow the run-time system to be written completely in BAM assembly.

%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%
% *** Check correctness of a sequence of BAM instructions ***

% Create saved state:
% Note: In C-Prolog this must be started up in a system
% equal to or larger than the one which created it.
main :- save(check, 1), prompt(_, ''), read(InstrJ, pipe(Instr, 0, 0), halt.
main :- halt.

% Pipe working loop:
pipe(end_of_file, M, N) :- !,
  T is M+N,
  write('*** Checked '),write(T),write(' instructions; '),
  write(M),write(' correct and '),write(N),write(' incorrect.'),nl.
pipe(Instr, M, N) :-
  (instr(Instr)
    -> M1 is M+1, N1=N
    ; M1=M, N1 is N+1,
    write('*** Incorrect: '),write(Instr),nl
  ),
  !, read(NewInstr), pipe(NewInstr, M1, N1).

%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%
% *** BAM Instructions ***

% 1. Unification support instructions:
instr(deref(V,W)) :- var_i(V), var_i(W).
instr(equal(EA,A,L)) :- ea_e(EA), arg_i(A), lbl(L).
instr(unify(V,W,F,G,L)) :- var_i(V), var_i(W), nv_flag(F),nv_flag(G),lbl(L).
instr(trail(V)) :- var_i(V).
instr(move(EA,VI)) :- ea_m(EA), var_i(VI).
instr(push(EA,R,N)) :- ea_p(EA), hreg(R), pos(N).
instr(adda(R,S,T)) :- numreg(R), numreg(S), hreg(T).
instr(pad(N)) :- pos(N).
instr(unify_atomic(V, I, L)) :- var_i(V), an_atomic(I), lbl(L).
instr(fail).

2. Conditional control flow instructions:
instr(switch(T, V, A, B, C)) :- a_tag(T), var_i(V), lbl(A), lbl(B), lbl(C).
instr(choice(I/N, Rs, L)) :- pos(I), pos(N), I=<N, lbl(L), regs(Rs).
instr(test(Eq, T, V, L)) :- eq_ne(Eq), var_i(V), a_tag(T), lbl(L).
instr(jump(C, A, B, L)) :- cond(C), numarg_i(A), numarg_i(B), lbl(L).
instr(move(CH, V)) :- a_var(V), choice_ptr(CH).
instr(cut(V)) :- a_var(V).
instr(hash(T, R, N, L)) :- hash_type(T), reg(R), pos(N), lbl(L).
instr(pair(E, L)) :- an_atomic(E), lbl(L).

3. Arithmetic instructions:
instr(add(A, B, V)) :- numarg_i(A), numarg_i(B), a_var(V).
instr(sub(A, B, V)) :- numarg_i(A), numarg_i(B), a_var(V).
instr(mul(A, B, V)) :- numarg_i(A), numarg_i(B), a_var(V).
instr(div(A, B, V)) :- numarg_i(A), numarg_i(B), a_var(V).
instr(mod(A, B, V)) :- numarg_i(A), numarg_i(B), a_var(V).
instr(and(A, B, V)) :- numarg_i(A), numarg_i(B), a_var(V).
instr(or(A, B, V)) :- numarg_i(A), numarg_i(B), a_var(V).
instr(xor(A, B, V)) :- numarg_i(A), numarg_i(B), a_var(V).
instr(not(A, V)) :- numarg_i(A), a_var(V).
instr(sll(A, B, V)) :- numarg_i(A), numarg_i(B), a_var(V).
instr(sra(A, B, V)) :- numarg_i(A), numarg_i(B), a_var(V).
instr(sll). /* vlsi_plm only */
instr(sra). /* vlsi_plm only */

4. Procedural instructions:
instr(procedure(N/A)) :- atom(N), natural(A).
instr(call(N/A)) :- atom(N), natural(A).
instr(return).
instr(simple_call(N/A)) :- atom(N), natural(A).
instr(simple_return).
instr(label(L)) :- lbl(L).
instr(jump(L)) :- lbl(L).
instr(allocate(Perms)) :- natural(Perms).
instr(deallocate(Perms)) :- natural(Perms).
instr(nop).

5.Pragma information for translator and reorderer:
instr(pragha(P)) :- pragma(P).

6. Additions to BAM for the assembly language programmer:
instr(I) :- user_instr(I).

Additions to BAM for the assembly language programmer

This section describes the parts of the BAM language that are never output by the compiler, but only used by the BAM assembly programmer. This is used to write the run-time system in BAM code, so that it is as portable as
% possible. Additional instructions are jump to register address, convert
% tagged atom or integer to untagged integer (ord), its inverse (val), and
% non-trapping full-word unsigned comparison, non-trapping full-word
% arithmetic, and trailing for backtracking destructive assignment.

user_instr(jump_reg(R)) :- reg(R).
user_instr(jump_nt(C, A, B, L)) :- cond(C), numarg_i(A), numarg_i(B), lbl(L).
user_instr(ord(A, B)) :- arg(A), a_var(B).
user_instr(val(T, A, V)) :- a_tag(T), numarg_i(A), a_var(V).
user_instr(add_nt(A, B, V)) :- numarg_i(A), numarg_i(B), a_var(V).
user_instr(sub_nt(A, B, V)) :- numarg_i(A), numarg_i(B), a_var(V).
user_instr(and_nt(A, B, V)) :- numarg_i(A), numarg_i(B), a_var(V).
user_instr(or_nt(A, B, V)) :- numarg_i(A), numarg_i(B), a_var(V).
user_instr(xor_nt(A, B, V)) :- numarg_i(A), numarg_i(B), a_var(V).
user_instr(sll_nt(A, B, V)) :- numarg_i(A), numarg_i(B), a_var(V).
user_instr(sra_nt(A, B, V)) :- numarg_i(A), numarg_i(B), a_var(V).
user_instr(trail_bda(X)) :- a_var(X).

% Additional registers:
% See Implementation Manual for list of existing registers.
user_reg(r(All :- atom(A).

% *** Pragmas ***

% A variable is a multiple of N.
% Inserted just before loads in readmode unification.
pragma(align(V, N)) :- a_var(V), pos(N).

% Inserted just before a sequence of pushes in writemode unification.
% (The pushes may be interleaved with non-memory moves.)
pragma(push(term(Size))) :- pos(Size).
pragma(push(cons)).
pragma(push(structure(A))) :- pos(A).
pragma(push(variable)).

% Specify the tag of a variable.
% (This is useful for processors without explicit tag support.)
pragma(tag(V, T)) :- a_var(V), a_tag(T).

% Length of a hash table.
pragma(hash_length(Len)) :- pos(Len).

% *** Tags ***
a_tag(tatm). /* atom */
a_tag(tint). /* integer */
a_tag(tneg). /* negative integer */
a_tag(tpos). /* nonnegative integer */
a_tag(tstr). /* structure */
/* cons cell */
a_tag(tlst).  
/* variable */

atom_tag(tatm).

pointer_tag(tstr).
pointer_tag(tlst).
pointer_tag(tvar).

% *** Addressing modes ***

heap_ptr(r(h)).
choice_ptr(r(b)).

reg(r(I)) :- int(I).
reg(U) :- user_reg(U).

hreg(R) :- reg(R).
hreg(R) :- heap_ptr(R).

perm(p(I)) :- natural(I).

an_atomic(I) :- int(I).
an_atomic(T-A) :- atom(A), atom_tag(T).
an_atomic(T-(F/N)) :- atom(F), pos(N), atom_tag(T).

a_var(Reg) :- reg(Reg).
a_var(Perm) :- perm(Perm).

arg(Arg) :- a_var(Arg).
arg(Arg) :- an_atomic(Arg).

var_i(Var) :- a_var(Var).
var_i([Var]) :- a_var(Var).

arg_i(Arg) :- var_i(Arg).
arg_i(Arg) :- an_atomic(Arg).

numreg(Arg) :- reg(Arg).
numarg(Arg) :- int(Arg).

numarg_i(Arg) :- var_i(Arg).
numarg_i(Arg) :- int(Arg).

var_off([Var]) :- a_var(Var).
var_off([Var+I]) :- a_var(Var), pos(I).

% Effective address for equal:
ea_e(Var) :- a_var(Var).
ea_e(VarOff) :- var_off(VarOff).

% Effective address for move:
ea_m(Arg) := arg(Arg).
ea_m(VarOff) := var_off(VarOff).
ea_m(Tag'H) := pointer_tag(Tag), heap_ptr(H).

% Effective address for push:
ea_p(Arg) := arg_i(Arg).
ea_p(Tag'H) := pointer_tag(Tag), heap_ptr(H).
ea_p(Tag"(H+D)) := pointer_tag(Tag), pos(D), heap_ptr(H).

% *** Miscellaneous ***
eq_ne(eq). /* Equal */
eq_ne(ne). /* Not equal */

cond(lts). /* Signed less than */
cond(les). /* Signed less than or equal */
cond(gts). /* Signed greater than */
cond(ges). /* Signed greater than or equal */
cond(eq). /* Equal */
cond(ne). /* Not equal */

hash_type(atomic).
hash_type(structure).

lbl(fail).
lbl(N/A) := atom(N), natural(A).
lbl(l(N/A, I)) := atom(N), natural(A), natural(I).

nv_flag(nonvar).
nv_flag(var).
nv_flag('?').

% A list of register numbers:
% (May contain the value 'no' as well)
regs([]).
regs([R|Set]) := (int(R); R=no), regs(Set).

% *** Utilities ***
ground(X) := nonvar(X), functor(X, _, N), ground(N, X).

ground(N, _) := N=:=0.
ground(N, X) := N\=\leq0, arg(N, X, A), ground(A), N1 is N-1, ground(N1, X).

int(N) := integer(N).
natural(N) := integer(N), N\geq0.
pos(N) := integer(N), N>0.
Appendix C

Formal specification of the Berkeley Abstract Machine semantics

%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%
% Formal specification of the Berkeley Abstract Machine (BAM) semantics
% Copyright (C) 1990 Peter Van Roy and Regents of the University of California
% May be used and modified for non-commercial purposes if this notice is kept.
% Written by Peter Van Roy.

% The specification is a Prolog program that defines the meaning of BAM in
% terms of its execution in a simple memory model. It runs BAM code directly
% from the output of the Aquarius compiler.

% The specification does not include the user instructions of the BAM since
% their behavior depends on the target machine.

% The specification is written in the Extended DCG notation.

%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%

% Meaning of registers:
% r(b) Index to most recent choice point.
% r(e) Index to current environment.
% r(tr) Top of trail stack.
% r(h) Top of heap stack.
% r(hb) Value of r(h) at last choice point creation.
% r(pc) Code address.
% r(cp) Continuation pointer for code.
% r(tmp_cp) Temporary continuation pointer for code, used only in simple_call.
% r(retry) Retry address for backtracking, only exists inside choice points.
% r(I) Argument and temporary register.
% p(I) Location on current environment.

% Types stored in registers:
% r(e) Contains values of registers \{r(e),r(cp)\} \cup \{p(0), \ldots, p(N-1)\},
% where N is the size of the environment.
% r(b) Contains values of registers \{r(e),r(cp),r(tr),r(h),r(hb),r(retry)\} \cup RS,
% where RS is a subset of \{r(0), r(1), \ldots\}.
% r(tr) Contains a natural number.
% r(h), r(hb) Contain words with a pointer tag.
% r(pc), r(cp) Contain natural numbers or symbolic labels.
% r(tmp_cp) Contains a symbolic label.
% r(retry) Contains a symbolic label.
% r(I) Contains a word.
% p(I) Contains a word.

% Comments:
% A word is either an integer or a structure of the form Tag·Value where Value
% is a natural number except if Tag·tatm, in which case Value is an atom or a
% structure (F/N) where F is an atom and N is a natural number.
% A symbolic label is either the atom 'fail', or the structure F/N, or the
% structure I(F/N,I), where F is an atom and N and I are natural numbers.
% r(cp) is stored in environments, allowing nested calls.
% r(tmp_cp) is not stored in environments, allowing only one level of call.
% However, no environment is needed in a predicate containing a simple_call.
% There are no explicit stacks for environments or choice points: registers
% r(e) and r(b) each contain a set of register values.

%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%

% Accumulator declarations:

% Accumulators:

acc_info(code, T, In, Out, table_command(T,In,Out)).
acc_info(lblmap, T, In, Out, table_command(T,In,Out)).
acc_info(regs, T, In, Out, table_command(T,In,Out)).
acc_info(trail, T, In, Out, table_command(T,In,Out)).
acc_info(heap, T, In, Out, heap_table_command(T,In,Out)).
acc_info(count, T, In, Out, (Out is T+In)).

% Predicate declarations:

% Top level:
pred_info(execute, 0, [regs,heap,trail,code,lblmap,count]).
pred_info(instr_loop, 0, [regs,heap,trail,code,lblmap,count]).
pred_info(instr_loop_end, 1, [regs,heap,trail,code,lblmap,count]).
pred_info(instr, 1, [regs,heap,trail,code,lblmap<count]).

% Addressing modes:
pred_info(heap, 3, [heap]).
pred_info(reg, 3, [regs]).
pred_info(perm, 3, [regs]).
pred_info(a_var, 3, [regs]).
pred_info(var_i, 3, [regs,heap]).
pred_info(arg, 2, [regs]).
pred_info(arg_i, 2, [regs,heap]).
pred_info(numreg, 2, [regs]).
pred_info(numarg, 2, [regs,heap]).
pred_info(var_off, 2, [regs,heap]).
pred_info(imm_tag, 2, [regs]).
pred_info(ea_e, 2, [regs,heap]).
pred_info(ea_m, 2, [regs,heap]).
pred_info(ea_p, 2, [regs,heap]).

% Instruction utilities:
pred_info(deref_rtn, 2, [regs,heap,trail]).
pred_info(deref_rtn_cont, 3, [regs,heap,trail]).
pred_info(equal_rtn, 3, [regs,heap,trail]).
pred_info(switch_rtn, 5, [regs,heap,trail]).
pred_info(test_rtn, 4, [regs,heap,trail]).
pred_info(jump_cond_rtn, 4, [regs,heap,trail]).
pred_info(hash_lookup, 3, [regs,heap,trail,lblmap,code]).
% Implement the accumulator commands:
\[
\text{table\_command}(\text{ins}(l, Val), \text{In}, \text{In}) \leftarrow \text{ins}(\text{In}, l, Val).
\]
\[
\text{table\_command}(\text{get}(I, Val), \text{In}, \text{In}) \leftarrow \text{get}(\text{In}, I, Val).
\]
\[
\text{table\_command}(\text{set}(I, Val), \text{In}, \text{Out}) \leftarrow \text{set}(\text{In}, I, Val, \text{Out}).
\]

% Mask off tag before looking up heap entry:
\[
\text{heap\_table\_command}(\text{ins}(\_\_I, Val), \text{In}, \text{In}) \leftarrow \text{ins}(\text{In}, I, Val).
\]
\[
\text{heap\_table\_command}(\text{get}(\_\_I, Val), \text{In}, \text{In}) \leftarrow \text{get}(\text{In}, I, Val).
\]
\[
\text{heap\_table\_command}(\text{set}(\_\_I, Val), \text{In}, \text{Out}) \leftarrow \text{set}(\text{In}, I, Val, \text{Out}).
\]

% *** Initialization and runtime options ***
:- dynamic(bamspec\_option/1).

main :-
  save(bamspec, 1),
  prompt(_, ''),
  ( copyright,
    execute
    ; error(['Sorry, the executable BAM specification has failed.']))
  ),
  halt.
main :-
  halt.

copyright :-
  write('Berkeley Abstract Machine (BAM) Executable Specification'), nl,
  write('Copyright (C) 1990 Peter Van Roy and '),
  write('Regents of the University of California'), nl, nl.
flag_print(I) :- bamspec_option(print), !, write('Executing '), write(I), nl. flag_print(_).

% Look up symbolic label to get a numeric PC:
numeric_pc(PC, PC) --> (integer(PC)), !.
numeric_pc(PC, NPC) --> (get(PC, NPC) : lblmap).
% Read in the instructions and create the code array and label map:
% The code array gives the instruction corresponding to each PC value.
% The label map gives the PC value corresponding to each symbolic label.
read_code(Code, LblMap) :-
  read(Instr),
  read_code(Instr, 0, Code, LblMap).
read_code(end_of_file, _, Code, LblMap) :- !, seal(Code), seal(LblMap).
read_code((:-Option), PC, Code, LblMap) :- !,
  asserta(bamspec_option(Option)),
  read(NextInstr),
  read_code(NextInstr, PC, Code, LblMap).
read_code(Instr, PC, Code, LblMap) :-
  ins(Code, PC, Instr),
  insert_lblmap(Instr, LblMap, PC),
  PCI is PC+1,
  read(NextInstr),
  read_code(NextInstr, PCI, Code, LblMap).

% Add an entry to the label map:
insert_lblmap(label(L), LblMap, PC) :- !, ins(LblMap, L, PC).
insert_lblmap(procedure(P), LblMap, PC) :- !, ins(LblMap, P, PC).
insert_lblmap(_, _, _) ..

% *** Top level execution ***
execute :-
  write('Reading BAM code'), nl,
  read_code(Code, LblMap),
  write('Starting execution'), nl,
  execute(leaf, Regs, leaf, _, leaf, _, Code, _, LblMap, _, 0, N),
  write('Executed '), write(N), write(' instructions.'), nl,
  print_array(Regs).
execute(File) :-
  seeing(OldFile),
  see(File),
  read_code(Code, LblMap),
  seen,
  see(OldFile),
  execute(leaf, Regs, leaf, _, leaf, _, Code, _, LblMap, _, 0, N),
  write('Executed '), write(N), write(' instructions.'), nl,
  print_array(Regs).
execute -->>
[set (r(e), leaf)]: regs,
[set (r(b), leaf)]: regs,
[set (r(h), tvar'0)]: regs,
[set (r(tr), 0)]: regs,
[set (r(pc), 0)]: regs,
[set (r(cp), global_success/0)]: regs,
instr (choice [1/2, [], global_failure/0]),
instr_loop.

% Instruction execution loop:
instr_loop -->>
(get (r(pc), PC)]: regs,
',
instr_loop_end(PC).
instr_loop -->>
error('Attempt to execute beyond existing code.').

instr_loop_end(write/1) --> !, write_rtn, instr(return), instr_loop.
instr_loop_end(nl/0) --> !, nl, instr(return), instr_loop.
instr_loop_end(global_success/0) --> !,
write('*** Global success ***'), nl.
instr_loop_end(global_failure/0) --> !,
write('*** Global failure ***'), nl.
instr_loop_end(fail) -->
instr(fail),
',
instr_loop.
instr_loop_end(PC) -->>
numeric.pc(PC, NPC),
% Fetch: 
[get (NPC, Instr)]: code,
NPC1 is NPC+1,
[set (r(pc), NPC1)]: regs,
% Execute: 
[1]: count,
flag_print(Instr),
instr(Instr),
',
instr_loop.
instr_loop_end(PC) -->>
error('Program counter is ', PC).

$~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~$ 

% *** BAM Instructions *** 

% 1. Unification support instructions: 
instr(deref(V, W)) -->
 var_i(get, V, X), 
deref_rtn(X, Y), 
 var_i(set, W, Y).
instr(equal(EA, A, L)) -->
eq_e(EA, X),
arg_i(A, Y),
(lbl(L)),
equal_rtn(X, Y, L).
instr(unify(V, W, F, G, L)) -->>
var_i(get, V, X),
var_i(get, W, Y),
(nv_flag(F)),
(nv_flag(G)),
(lbl(L)),
equal_rtn(X, Y, L).
instr(unify_atomic(V, I, L)) -->>
var_i(get, V, X),
(atomic(I)),
(lbl(L)),
equal_rtn(X, I, L).
instr(trail(V)) -->>
var_i(get, V, X),
trail_rtn(X).
instr(move(EA, VI)) -->>
eam(EA, X),
var_i(set, VI, X), !.
instr(push(EA, R, N)) -->>
ea_p(EA, X),
(hreg(R)),
{get(R, Y)}:regs,
{set(X, X)}:heap,
{pos(N)},
add_word(Y, N, YN),
{set(R, YN)}:regs.
instr(adda(R, S, T)) -->>
(hreg(R)),
{get(R, X)}:regs,
numreg(S, Off),
add_word(X, Off, NX),
(hreg(T)),
{set(T, NX)}:regs.
instr(pad(N)) -->>
{get(r(h), H)}:regs,
{pos(N)},
add_word(H, N, NewH),
{set(r(h), NewH)}:regs.

% 2. Conditional control flow instructions:
instr(choice(1/N, Rs, L)) -->> {pos(N), N>1, regs(Rs), lbl(L)}, !,
save_choice_regs(Rs, NewB),
{ins(NewB, r(retry), L)},
{get(r(tr), TR)}:regs, {ins(NewB, r(tr), TR)},
{get(r(e), E)}:regs, {ins(NewB, r(e), E)},
{get(r(cp), CP)}:regs, {ins(NewB, r(cp), CP)},
{get(r(b), B)}:regs, {ins(NewB, r(b), B)},
{get(r(h), H)}:regs, {ins(NewB, r(h), H)},
{seal(NewB)},
{set(r(hb), H)}:regs,
{set(r(b), NewB)}:regs.
instr(choice(I/N,Rs,L)) -->> {pos(N), pos(L), I<1, 1<N, regs(Rs), lbl(L)}, !,
  [get(r(b),B)]:regs,
  restore_choice_regs(Rs, B),
  [set(B,r(retry),L,NewB)],
  [set(r(b),NewB)]:regs.
instr(choice(N/N,Rs,L)) -->> {pos(N), regs(Rs), lbl(L)}, !,
  [get(r(b),B)]:regs,
  restore_choice_regs(Rs, B),
  [set(B,r(retry),L,NewB)],
  [set(r(b),NewB)]:regs.
instr(fail) -->>
  [get(r(b),B)]:regs,
  [get(B,r(h),H)]:regs,
  [set(r(h),N)]:regs,
  [get(B,r(e),E)]:regs,
  [set(r(e),E)]:regs,
  [get(B,r(cpl,CP))]:regs,
  [set(r(cp),CP)]:regs,
  [get(r(tr),CurTR)]:regs,
  [get(B,r(tr),OldTR)],
  detrail rtn(CurTR, OldTR),
  [get(B,r(retry),L)],
  [set(r(pc),L)]:regs.
instr(switch(T,V,A,B,C)) -->>
  {a_tag(T)},
  var_i(get, V, X),
  extract_tag(X, TX),
  {lbl(h), lbl(B), lbl(C)},
  switch_rtn(T, TX, A, B, C).
instr(test(Eq,T,V,L)) -->>
  {a_tag(T)},
  var_i(get, V, X),
  extract_tag(X, TX),
  {eq_ne(Eq)},
  {lbl(L)},
  test_rtn(Eq, T, TX, L).
instr(jump(C,A,B,L)) -->>
  {cond(C)},
  numarg(A, XA), {extract_value(XA, VA), check_int(XA)},
  numarg(B, XB), {extract_value(XB, VB), check_int(XB)},
  {lbl(L)},
  jump_cond_rtn(C, VA, VB, L).
instr(move(r(b),V)) -->>
  [get(r(b),B)]:regs,
  a_var(set, V, B).
instr(cut(V)) -->>
  a_var(get, V, X),
  [set(r(b),X)]:regs,
  [get(X,r(h),H)],
  [set(r(hb),H)]:regs.
instr(hash(T,R,N,L)) -->> hash_type(T), pos(N), lbl(L),
  reg(get, R, X),
hash_indirect(T, X, Y),
[get(L,PC)]:lblmap,
hash_lookup(PC, Y, N).

instr(pair(_,_)) -->>
  {error(['Attempt to execute inside a hash table.'])}.

% 3. Arithmetic instructions:
instr(add(A,B,V)) -->> arith(add, A, B, V).
instr(sub(A,B,V)) -->> arith(sub, A, B, V).
instr(mul(A,B,V)) -->> arith(mul, A, B, V).
instr(div(A,B,V)) -->> arith(div, A, B, V).
instr(mod(A,B,V)) -->> arith(mod, A, B, V).
instr(and(A,B,V)) -->> arith(and, A, B, V).
instr(or(A,B,V)) -->> arith(or, A, B, V).
instr(xor(A,B,V)) -->> arith(xor, A, B, V).
instr(not(A,V)) -->> arith(not, A, 0, V).
instr(sll(A,B,V)) -->> arith(sll, A, B, V).
instr(sra(A,B,V)) -->> arith(sra, A, B, V).

% 4. Procedural instructions:
instr(procedure(N/A)) -->> {atom(N), natural(A)}.
instr(call(N/A)) -->> {atom(N), natural(A)},
  [get(r(pc),PC)]:regs,
  [set(r(cp),PC)]:regs,
  [set(r(pc),N/A)]:regs.
instr(return) -->>
  [get(r(cp),PC)]:regs,
  [set(r(pc),PC)]:regs.
instr(simple_call(N/A)) -->> {atom(N), natural(A)},
  [get(r(pc),PC)]:regs,
  [set(r(tmp_cp),PC)]:regs,
  [set(r(pc),N/A)]:regs.
instr(simple_return) -->>
  [get(r(tmp_cp),PC)]:regs,
  [set(r(pc),PC)]:regs.
instr(label(L)) -->> {lbl(L)}.
instr(jump(L)) -->> {lbl(L)},
  [set(r(pc),L)]:regs.
instr(allocation(N)) -->>
  [natural(N)],
  [get(r(e),E)]:regs,
  [ins(NewE, r(e), E)],
  [get(r(cp),CP)]:regs,
  [ins(NewE, r(cp), CP)],
  [seal(NewE)],
  [set(r(e),NewE)]:regs.
instr(deallocation(N)) -->>
  [natural(N)],
  [get(r(e),E)]:regs,
  [get(E,r(e),NewE)],
  [get(E,r(cp),NewCP)],
  [set(r(e),NewE)]:regs,
  [set(r(cp),NewCP)]:regs.
instr(nop) -->> [].
5. Pragma information for translator and reorderer:
Pragma are no-ops in the execution.
instr(pragmas(P)) --> [pragma(P)], !.

6. Additions to BAM for the assembly language programmer:
The meaning of these instructions depends on the underlying architecture,
so they are not included in this specification. See the Implementation
Manual for a discussion of their use.

Pragma information for translator and reorderer:
Pragma are no-ops in the execution.
instr(pragmas(P)) --> [pragma(P)], !.

** Pragmas **

A variable is a multiple of N.
Inserted just before loads in readmode unification.
pragma(align(V,N)) :- a_var(V), pos(N).

Inserted just before a sequence of pushes in writemode unification.
(The pushes may be interleaved with non-memory moves.)
pragma(push(term(Size))) :- pos(Size).
pragma(push(cons)).
pragma(push(structure(A))) :- pos(A).
pragma(push(variable)).

Specify the tag of a variable.
(This is useful for processors without explicit tag support.)
pragma(tag(V,T)) :- a_var(V), a_tag(T).

Length of a hash table.
pragma(hash_length(Len)) :- pos(Len).

** Tags **

a_tag(tatm). /* atom */
a_tag(tint). /* integer */
a_tag(tneg). /* negative integer */
a_tag(tpos). /* nonnegative integer */
a_tag(tstr). /* structure */
a_tag(tltl). /* cons cell */
a_tag(tvar). /* variable */

atom_tag(tatm).
atomic_tag(tatm).
atomic_tag(tint).
atomic_tag(tneg).
atomic_tag(tpos).

pointer_tag(tstr).
pointer_tag(tltl).
pointer_tag(tvar).
% *** Addressing modes ***

% Both read and write access:

heap(get, W, X) --> {ptr_word(W)}, [get(W,X)]:heap.
heap(set, W, X) --> {ptr_word(W)}, [set(W,X)]:heap.

ptr_word(T_) :- pointer_tag(T).

reg(get, R, X) --> {reg(R)}, [get(R,X)]:regs.
reg(set, R, X) --> {reg(R)}, [set(R,X)]:regs.

reg(r(I)) :- int(I), !.

hreg(R) :- reg(R), !.

perm(get, P, X) --> {perm(P)}, [get(r(e),E)]:regs, [get(E,P,X)].
perm(set, P, X) --> {perm(P)}, [get(r(e),E)]:regs, [set(E,P,X,NewE)],
                [set(r(e),NewE)]:regs.

perm(p(I)) :- natural(I).

a_var(WR, V, X) --> reg(WR, V, X), !.
a_var(WR, V, X) --> perm(WR, V, X).

a_var(Reg) :- reg(Reg), !.
a_var(Perm) :- perm(Perm).

var_i(WR, [V], X) --> a_var(get, V, W), heap(WR, W, X), !.
var_i(WR, V, X) --> a_var(WR, V, X).

% Read access only:

% An int is its own value:
int(N) :- integer(N).

% An atomic is its own value:
an_atomic(I) :- int(I), !.
an_atomic(T\A) :- atom(A), atom_tag(T), !.
an_atomic(T\(F/N)) :- atom(F), pos(N), atom_tag(T).

arg(Arg, Arg) --> {an_atomic(Arg)}, !.
arg(Arg, X) --> a_var(get, Arg, X).

arg_i(Arg, Arg) --> {an_atomic(Arg)}, !.
arg_i(Arg, X) --> var_i(get, Arg, X).

numreg(Arg, Arg) --> {int(Arg)}, !.
numreg(Arg, X) --> reg(get, Arg, X).
numarg(Arg, Arg) -->> \{int(Arg)\}, !.
numarg(Arg, X) -->> var_i(get, Arg, X).

var_off([Var+I], X) -->> a_var(get, Var, I), !,
(pos(I)), add_word(T, I, T2), \{get(T2, X)\}:heap.
var_off([Var], X) -->> a_var(get, Var, I), \{get(I, X)\}:heap.

% Creating immediate tagged pointer objects:
imm_tag(Tag \cdot r(h) + D), W -->> \{pointer_tag(Tag)\}, !,
[get(r(h), T)]:regs,
[pos(D)], add_word(T, D, X),
insert_tag(Tag, X, W).
imm_tag(Tag \cdot r(h), W) -->> \{pointer_tag(Tag)\}, !,
[get(r(h), X)]:regs,
insert_tag(Tag, X, W).

% Effective address for equal:
ea_e(Var, X) -->> a_var(get, Var, X), !.
ea_e(VarOff, X) -->> var_off(VarOff, X).

% Effective address for move:
ea_m(Arg, X) -->> arg(Arg, X), !.
ea_m(VarOff, X) -->> var_off(VarOff, X), !.
ea_m(T \cdot r(h), X) -->> imm_tag(T \cdot r(h), X).

% Effective address for push:
ea_p(Arg, X) -->> arg_i(Arg, X), !.
ea_p(T \cdot Y, X) -->> imm_tag(T \cdot Y, X).

%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%

% *** Miscellaneous ***

eq_ne(eq). /* Equal */
eq_ne(ne). /* Not equal */

cond(lts). /* Signed less than */
cond(les). /* Signed less than or equal */
cond(gts). /* Signed greater than */
cond(ges). /* Signed greater than or equal */
cond(eq). /* Equal */
cond(ne). /* Not equal */

hash_type(atomic).
hash_type(structure).

lbl(fail).
lbl(N/A) :- atom(N), natural(A).
lbl(l(N/A, I)) :- atom(N), natural(A), natural(I).

nv_flag(nonvar).
nv_flag(var).
nv_flag('?').
% A list of register numbers:
% (May contain the value 'no' as well)
regs([]).
regs([R|Set]) :- (int(R); R=no), !, regs(Set).

% Dereference utilities:
deref_rtn(X, X) --> {nonvartag(X)}, !.
deref_rtn(X, Y) -->
  [get(X,X2)]:heap,
  deref_rtn_cont(X, X2, Y).
deref_rtn_cont(X, X, Y) --> !, [Y=X].
deref_rtn_cont(_, X, Y) --> deref_rtn(X, Y).

% Equal routine:
equal_rtn(X, X, _) --> !.
equal_rtn(_, _, L) --> [set(r(pc),L)]:regs.

% Switch and test routines:
switch_rtn(_, tvar, A, _, _) --> !, [set(r(pc),A)]:regs.
switch_rtn(T, TX, _, B, _) --> {equivalent_tag(T,TX)}, !, [set(r(pc),B)]:regs.
switch_rtn(_, _, _, _, C) --> [set(r(pc),C)]:regs.
test_rtn(Eq, T, TX, L) --> {test_true(Eq, T, TX)}, !, [set(r(pc),L)]:regs.
test_rtn(_, _, _, _, _) --> [].
test_true(eq, T, TX) := equivalent_tag(T, TX).
test_true(ne, T, TX) := \+equivalent_tag(T, TX).

% Arithmetic utilities:
arith(Op, A, B, V) -->
  numarg(A, XA), {extract_value(XA, VA), check_int(XA)},
  numarg(B, XB), {extract_value(XB, VB), check_int(XB)},
  arith_operation(Op, VA, VB, VC),
  a_var(set, V, VC).
arith_operation(add, VA, VB, VC) :- VC is VA+VB.
arith_operation(sub, VA, VB, VC) :- VC is VA-VB.
arith_operation(mul, VA, VB, VC) :- VC is VA*VB.
arith_operation(div, VA, VB, VC) :- VC is VA//VB.
arith_operation(mod, VA, VB, VC) :- VC is VA mod VB.
arith_operation(and, VA, VB, VC) :- VC is VA /
  B.
arith_operation( or, VA, VB, VC) :- VC is VA \ VB.
arith_operation( xor, VA, VB, VC) :- VC is (VA \ (VB)) \ (VB \ (VA)).
arith_operation( not, VA, _ , VC) :- VC is \ (VA).
arith_operation( sll, VA, VB, VC) :- VC is VA<<VB.
arith_operation( sra, VA, VB, VC) :- VC is VA>>VB.

% Conditional jump:
jump_cond_rtn(C, VA, VB, L) --> [jump_true(C, VA, VB)), !, [set(r(pc),L)]:regs.
jump_cond_rtn(_, _, _, _) --> !.

jump_true(lts, VA, VB) :- VA< VB.
jump_true(gts, VA, VB) :- VA> VB.
jump_true(les, VA, VB) :- VA\= VB.
jump_true(ges, VA, VB) :- VA\= VB.
jump_true(eq, VA, VB) :- VA\= VB.
jump_true(ne, VA, VB) :- VA\= VB.

% Hash table utilities:
hash_lookup(PC, X, N) -->
  (PC1 is PC+1),
  [get(PC1,pragma(hash_length(N)))]:code,
  (PC2 is PC1+1),
  (PCN is PC1+N),
  hash_lookup_2(PC2, PCN, X).

hash_lookup_2(PC, PCN, _) --> !.
hash_lookup_2(PC, PCN, X) -->
  [get(PC,pair(E,L))]:code,
  E\=X,
  !,
  [set(r(pc),L)]:regs.
hash_lookup_2(PC, PCN, X) -->
  [get(X,Y)] :heap.

% Indirection needed for structures because main functor is in memory:
hash_indirect(atomic, X, X) --> !.
hash_indirect(structure, X, Y) --> [get(X,Y)] :heap.

% Choice point and fail utilities:
save_choice_regs([], _) --> !.
save_choice_regs([Io|Rs], B) --> !,
  save_choice_regs(Rs, B).
save_choice_regs([II|Rs], B) -->
  [get(r(I),R)]:regs,
% Trailing and detrailing:

trail_rtn(X) -->
   [get(r{hb},HB):regs, cmp_trail(X, HB).
cmp_trail(X, HB) --> {less_trail(X, HB)}, !,
   [get(r{tr},TR):regs, [set(TR,X)]:trail,
   [TR is TR+1], [set(r{tr},TR1)]:regs.
cmp_trail(_, _) --> [].

less_trail(_X, _'Y) :- X<Y.

% Restore to unbound the variables on the trail between OldTR and CurTR.
detrail_rtn(CurTR, OldTR) --> (CurTR=OldTR), !.
detrail_rtn(CurTR, OldTR) --> (CurTR>OldTR),
   (CurTR is CurTR-1), [get(CurTR1,V)]:trail,
   [set(V,V)]:heap, detrail_rtn(CurTR1, OldTR).

% General unification routine:

unify_rtn(W1, W2, L) -->
   unify_rtn_2(W1, W2, Flag), unify_end(Flag, L).

unify_end(success, _) --> [].
% For later: detrailing if L=success.
unify_end(fail, L) --> [set(r{pc},L)]:regs.

unify_rtn_2(W1, W2, Flag) -->
   {extract_tag_value(W1, T1, V1)},
   {extract_tag_value(W2, T2, V2)},
   unify_rtn_2(T1, V1, T2, V2, Flag).

unify_rtn_2(tvar, V1, NTag, V2, success) --> [NTag=tvar], !,
trail_rtn(tvar, V1),
(make_word(NTag, V2, Word)),
(set(tvar, V1, Word)):heap.

unify_rtn_2(NTag, V2, tvar, V1, success) --> (NTag == tvar), !,
trail_rtn(tvar, V1),
(make_word(NTag, V2, Word)),
(set(tvar, V1, Word)):heap.

unify_rtn_2(tvar, V1, tvar, V2, success) --> !,
unify_varvar(V1, V2).

% Matching atomic tags:
unify_rtn_2(ATag, V1, BTag, V2, Flag) -->
{atomic_tag(ATag)},
{atomic_tag(BTag)},
{equivalent_tag(ATag, BTag)},{},
unify_atm(V1, V2, Flag).

% Non-matching nonvariable tags:
unify_rtn_2(ATag, _, BTag, _, fail) -->
{ATag == tvar, BTag == tvar},
{\+equivalent_tag(ATag, BTag)},{},

% Matching pointer tags (recursive case):
unify_rtn_2(ATag, V1, ATag, V2, Flag) -->
{pointer_tag(ATag)},
get_size(ATag, V1, Sz),
unify_rtn_args_2(0, Sz, ATag, V1, V2, Flag).

% The term's Size is the maximum offset needed to traverse the term in memory.
get_size(tlst, _, 1) --> [].

get_size(tstr, V, N) -->
{get(tstr, V, Func) : heap,
{Func is(tatm(_) / N)}).

unify_rtn_args_2(N, Sz, _, _, _, success) --> (N > Sz), !.
unify_rtn_args_2(N, Sz, T, V, W, Flag) --> (N < Sz), !,
{VN is V + N},
{WN is W + N},
{get(T \ VN, VX) : heap, deref_rtn(VX, DVX),
{get(T \ WN, WX) : heap, deref_rtn(WX, DWX),
unify_rtn_2(DVX, DWX, F),
{N1 is N + 1},
unify_rtn_args_3(F, N1, Sz, T, V, W, Flag).

% Continue with other arguments if argument unification succeeded:
unify_rtn_args_3(fail, _, _, _, _, _, fail) --> [].
unify_rtn_args_3(success, N1, Sz, T, V, W, Flag) -->
unify_rtn_args_2(N1, Sz, T, V, W, Flag).

% Unifying value parts of two atomic terms with equivalent tag:
unify_atm(V, V, success) --> !.
unify_atm(_, _, fail) --> [].

% Unifying two variables: bind youngest to oldest, trail youngest.
unify_varvar(V1, V2) --> (V1 > V2), !.
trail_rtn(tvar"Vl),
(set(tvar"Vl,tvar"V2)):heap.
unify_varvar(Vl, V2) --> (Vl=<V2), !,
trail_rtn(tvar"V2),
(set(tvar"V2,tvar"V1)):heap.

% Simple type utilities:

% This takes into account the relationship between tpos, tneg and tint.
% For integers it extracts tpos or tneg tags and the absolute value
% of the integer. It creates the correct integer, given the tpos, tneg
% or tint tags.

equivalent_tag(T, T) :- !.
equivalent_tag(tint, tpos) :- !.
equivalent_tag(tint, tneg).

extract_tag(N, tpos) :- integer(N), N>=0, !.
extract_tag(N, tneg) :- integer(N), N<0, !.
extract_tag(T, T).

extract_value(N, N) :- int(N), N>=0, !.
extract_value(N, M) :- int(N), N<0, !, M is -N.
extract_value(_-V, V).

extract_tag_value(W, T, V) :-
  extract_tag(W, T),
  extract_value(W, V).

nonvartag(_) :- int(I), !.
nonvartag(T) :- \+T\=tvar.

% Only used for pointer tags:
insert_tag(T, _-V, T\='V).

make_word(tint, I, I) :- !.
make_word(tpos, I, I) :- !.
make_word(tneg, N, I) :- !, I is -N.
make_word(T, V, T\='V).
add_word(T\textsuperscript{T}, I, J, T\textsuperscript{K}) :- K is I+J.

% Eventually, print out value of PC:
check_int(I) :- int(I), !.
check_int(_) :-
  error(["Operand of conditional is not an integer."]).

% Table utilities:

% This code implements a mutable array, represented as a binary tree.

% Insert a value in logarithmic time and constant space:
% This predicate is used in this program only to create the array,
% although it can also be used to access array elements.
ins(T, I, V) :- hash(I, H), ins_2(T, H, V).


ins_2(N, V, _, _, I, V) :- I=N, !.
ins_2(N, _, L, R, I, V) :-
  compare(Order, I, N),
  ins_2(Order, I, V, L, R).

ins_2(<, I, V, L, _) :- ins_2(L, I, V).

% Access a value in logarithmic time and constant space:
% This predicate cannot be used to create the array incrementally,
% but it is faster than ins/3.
get(T, I, V) :- hash(I, H), get_2(T, H, V).

get_2(node(N,W,L,R), I, V) :-
  compare(Order, I, N),
  get_3(Order, I, V, W, L, R).

get_3(<, I, V, _, L, _) :- get_2(L, I, V).
get_3(>, __, V, W, _, _) :- V=W.
get_3(>, I, V, _, _, R) :- get_2(R, I, V).

% Update an array in logarithmic time and space:

set_2(leaf, I, V, node(I,V,leaf,leaf)).
set_2(node(N,W,L,R), I, V, node(N,NW,NL, NR)) :-
  compare(Order, I, N),
  set_3(Order, I, V, W, L, R, NW, NL, NR).


% Prevent any further insertions in the array:
seal(leaf).
seal(node(_,_,L,R)) :- seal(L), seal(R).

% Print values of array in sorted order:
print_array(Term) :-
    flat_array(Term, 2, Flat),
    print_list(Flat).

print_list([]).
print_list([(A->B) | L]) :-
    write(A), put(9), write(‘= ’), write(B), nl,
    print_list(L).

flat_array(Term, N, Sort) :-
    N>0, N is N-1,
    flat_array(Term, N1, Flat, []), !,
    sort(Flat, Sort).

flat_array(leaf, N, []) :- N=:=0, !.
flat_array(node(_,_,_), N, ‘...’) :- N=:=0, !.
flat_array(Term, _, Term).

flat_array(leaf, _) --> [].
flat_array(node(H,T,L,R), N) -->
    flat_array(L, N),
    {hash(H, I)},
    {flat_array(T, N, F)},
    [(I->F)],
    flat_array(R, N).

% Invertible hash function:
% Bit inversion of the integer components of a ground term. Other parts are
% unchanged. This one inverts the low 16 bits. It can be changed by changing
% the last argument of bit_invert/3.
hash(I, H) :- integer(I), !, bit_invert(I, H, 16).
hash(T, H) :- functor(T, Na, Ar), functor(H, Na, Ar), hash_2(Ar, T, H).

hash_2(0, _, _) :- !.
hash_2(N, T, H) :- N>0,
    arg(N, T, X),
    arg(N, H, X),
    hash(X, Y),
    N1 is N-1,
    hash_2(N1, T, H).

bit_invert(0, 0, _) :- !.
bit_invert(N, I, B) :- N>0,
    L is N>>1,
    R is N\l1,
    B1 is B-1,
    bit_invert(L, LI, B1),
    I is R*(1<<B) + LI.

%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%
% Error handling:

error(L) :-
    write('*** Error: '),
    error_loop(L),
    write('***'), nl.

error_loop([]).
error_loop([M|L]) :- write(M), error_loop(L).

%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%

% Primitive version of write:

comment guard: %
write_rtn -->>
    [get(r(0), X):regs,
     write_rtn(X)]

write_rtn(tvar~V) -->> !, (write('_'), write(V)).
write_rtn(int(I)) -->> {write(I)}, !, {write(I)}.
write_rtn(tatm~(F/N)) -->> !, (write('"'), write(F/N), write('"')).
write_rtn(tatm~A) -->> !, {write(A)}.
write_rtn(tlst~V) -->> !,
    {W is V+1},
    [get(tlst~V, Head):heap,
     get(tlst~W, Tail):heap,
     deref_rtn(Head, DHead),
     deref_rtn(Tail, DTail),
     {write('['),
      write_rtn(DHead),
      {write('"'),
       write_rtn(DTail),
       write('"')}).
write_rtn(tstr~V) -->> !,
    [get(tstr~V, tatm~(F/N))]:heap,
    {write(F), write('('),
     write_arg(V),
     write_args(2, N, V),
     {write(')')}).
write_args(I, N, _) -->> {I>N}, !.
write_args(I, N, V) -->> {I<N}, !,
    {I is I+1},
    {write(','),
     write_arg(V),
     write_args(I, N, V)}.
write_arg(V, I) -->>
    {W is V+1},
    [get(tstr~W, X):heap,
     deref_rtn(X, DX),
     write_rtn(DX).
Appendix D

Semantics of the Berkeley Abstract Machine

1. Introduction

This appendix gives an English-language description of the semantics of the Berkeley Abstract Machine (BAM) as comments attached to a Prólog specification of its syntax. The BAM is intended to operate on the same data structures as the Warren Abstract Machine (WAM), therefore some familiarity with the WAM is an advantage. The semantics are represented by short descriptions supplemented by pseudo-code and examples where necessary.

The BAM is designed to be simple and easily translated to most general-purpose processors. Many of its optimizations apply to any processor, for example the streamlined choice point management and the use of write-once permanent variables to simplify trailing. Although the first target is the VLSI-BAM processor, we have built translators for other processors including the MIPS and the MC68020. Pragmas give information that is used to obtain the best translation for different processors.

The instruction set is divided in six categories, each in a different section. Each section starts with a box giving the syntax of the instructions presented in that section. This is followed by a description of the instructions' actions. Section 2 gives the unification instructions. Section 3 gives the conditional control flow instructions. Section 4 gives the arithmetic instructions. Section 5 gives the procedural control flow instructions. Section 6 gives the pragmas, which contain information that allows better translation. Section 7 gives the user instructions, additions to the BAM that are never output by the compiler but are intended for the BAM assembly programmer. The last section defines the syntax and semantics of the addressing modes used in the instructions.

In explaining the semantics, a few assumptions are made about the data representation. An infinite number of registers is assumed; the translator should map registers of sufficiently large index to memory. A tagged architecture is assumed; i.e. each word contains a tag and a value field which are treated as separate entities in some instructions and as a unit in other instructions. A load-store architecture is assumed; almost any architecture has a subset of instructions that satisfy this assumption. The actual details of the translation to the target architecture are not given since they depend on the characteristics of the architecture. These characteristics include the number of registers, the addressing modes, hardware support for certain features (tagging, dereferencing, trailing, etc.), the precise format of choice points and environments, and so forth.
<table>
<thead>
<tr>
<th>Unification instructions</th>
<th>Unification syntax</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>instr(deref(V,W))</code></td>
<td>( \texttt{var_i(V), var_i(W)} )</td>
</tr>
<tr>
<td><code>instr(equal(EA,A,L))</code></td>
<td>( \texttt{ea_e(EA), arg_i(A), lbl(L)} )</td>
</tr>
<tr>
<td><code>instr(unify(V,W,F,G,L))</code></td>
<td>( \texttt{var_i(V), var_i(W), nv_flag(F), nv_flag(G), lbl(L)} )</td>
</tr>
<tr>
<td><code>instr(trail(V))</code></td>
<td>( \texttt{var_i(V)} )</td>
</tr>
<tr>
<td><code>instr(move(EA,VI))</code></td>
<td>( \texttt{ea_m(EA), var_i(VI)} )</td>
</tr>
<tr>
<td><code>instr(push(EA,R,N))</code></td>
<td>( \texttt{ea_p(EA), hreg(R), pos(N)} )</td>
</tr>
<tr>
<td><code>instr(pad(N))</code></td>
<td>( \texttt{pos(N)} )</td>
</tr>
<tr>
<td><code>instr(unify_atomic(V,l,L))</code></td>
<td>( \texttt{var_i(V), an_atomic(I), lbl(L)} )</td>
</tr>
<tr>
<td><code>instr(fail)</code></td>
<td></td>
</tr>
</tbody>
</table>

- **deref(V,W)**: Dereference the argument V and store the result in W. The argument V is unchanged. This is the only instruction which dereferences its arguments. All other instructions assume that their arguments are dereferenced. Giving the dereference instruction two arguments simplifies the implementation of write-once permanent variables and makes a fast implementation of trailing possible.

- **equal(X,Y,L)**: Compare X to Y and branch to L if they are not equal. The comparison is a full word operation, equivalent to "eq" in Lisp. It is assumed that X and Y are dereferenced.

- **unify(X,Y,T,U,L)**: Perform a general unification of X and Y, and branch to L if it fails. Always binds oldest variables to the youngest. In the failure case all bindings are undone. It is assumed that X and Y are dereferenced. The two parameters T and U are added as an optimization, and may be safely ignored. They are flags (with values ‘?’, var, or nonvar) that say whether it is known if X and Y are variables or nonvariables. With this information a better translation to the target processor can be done.

- **trail(X)**: Push the address of X on the trail stack if the trail condition \( X<r(hb) \) is satisfied. It is assumed that X is a dereferenced unbound variable, i.e. it has a tvar tag. Only one comparison is necessary for the trail check. The static register \( r(hb) \) points to the heap location which was the top of the heap when the most recent choice point was created.

- **move(X,Y)**: Move X to Y. Depending on the addressing mode, this instruction does a load or store or creates a tagged value.

- **push(X,R,N)**: Push X on the stack with stack pointer R, then increment R by N. This instruction is used for write-mode unification.

- **adda(X,Y,R)**: Add X and Y into R. This is a full word operation which never traps, unlike the arithmetic instructions in section 4. This instruction is used to allocate space for uninitialized variables. The second argument Y is an offset which is scaled properly by the translator (i.e. it is unchanged for the VLSI-BAM since it is word-addressed, and it is multiplied by 4 for the MIPS, since it is byte-addressed).
pad(N)  Add N words to the heap pointer \( r(h) \). This is a full word operation which never traps, unlike the arithmetic instructions in section 4. It is used to ensure the correct alignment of compound terms. The space reserved by pad will never be stored to. If the increment is a multiple of the alignment then the pad disappears. The increment is scaled properly by the translator (see previous description of adda).

unify_atom (X, Y, L)  Unify the variable X with the atomic term Y, and branch to L if it fails. It is assumed that X is dereferenced. The \texttt{unify} \_\texttt{atomic} instruction is a special case of general unification that is added to reduce code size in the VLSI-BAM processor. There is a compiler option to enable or disable the generation of this instruction.

fail  Untar all variable bindings and jump to the retry address. Do not restore argument registers. Argument registers are restored by the choice point management instructions.

3. Conditional control flow instructions

<table>
<thead>
<tr>
<th>Clause selection syntax</th>
</tr>
</thead>
<tbody>
<tr>
<td>instr(switch(T,V,A,B,C)) :- a_tag(T), var_i(V), lbl(A), lbl(B), lbl(C).</td>
</tr>
<tr>
<td>instr(choice(I/N,Rs,L)) :- pos(I), pos(N), I=&lt;N, lbl(L), regs(Rs).</td>
</tr>
<tr>
<td>instr(test(Eq,T,V,L)) :- eq_ne(Eq), var_i(V), a_tag(T), lbl(L).</td>
</tr>
<tr>
<td>instr(jump(C,A,B,L)) :- cond(C), numarg_i(A), numarg_i(B), lbl(L).</td>
</tr>
<tr>
<td>instr(move(CH,V)) :- a_var(V), choice_ptr(CH).</td>
</tr>
<tr>
<td>instr(hash(T,R,N,L)) :- hash_type(T), reg(R), pos(N), lbl(L).</td>
</tr>
<tr>
<td>instr(pair(E,L)) :- an_atomic(E), lbl(L).</td>
</tr>
</tbody>
</table>

\texttt{switch(T,R,A,B,C)}  A three-way branch: branch to the label A, B, C depending on whether the tag of R is \texttt{tvar}, T, or any other value. The label \texttt{fail} is not an address, but denotes a branch to the global failure routine. It is assumed that R is dereferenced.

\texttt{choice(I/N,RS,L)}  The choice point management instruction for choosing clause I out of N clauses. Choice points are of variable size. The semantics of choice depends on I as follows:

\texttt{I=1}  Create a choice point with retry address L. Save in it the registers listed in RS.

\texttt{1<\textless I<\textless N}  Restore the registers mentioned in RS from the choice point, ignoring no terms. The no terms make it possible to know the position of the registers in the choice point without an explicit size field in the choice point. Update the retry address to L.

\texttt{I=N}  Restore the registers mentioned in RS, ignoring no terms. Remove the choice point. (L will always be \texttt{fail} when I=N.)
The above notation is consistent with three possible implementations (in order of decreasing efficiency): (1) The implementation given above, in which only those registers listed in RS are saved and restored, and the choice point does not have a size field. Restoring registers is done by the choice instructions, not by the fail instruction. The compiler does an effort to minimize the set of registers mentioned in RS. (2) Saving all registers up to the maximum register listed in RS. In this case the choice points are of variable size, and the no terms in RS are ignored. The notation is consistent with choice points containing a size field. (3) Always saving and restoring all registers. In this case the choice points are of fixed size, the RS argument is ignored, and the fail instruction restores the registers. In this case the semantics correspond to the try, retry, and trust instructions of the WAM.

\[ \text{test}(E, T, X, L) \]

Branch to label L if the tag of X is equal/not equal to T. Equality/nonequality is selected by the value of E. The label fail is not an address, but denotes a branch to the global failure routine. It is assumed that X is dereferenced.

\[ \text{jump}(C, X, Y, L) \]

Compare X and Y and jump to L if the comparison is true. The kind of comparison is given by C. This instruction traps if either argument is not an integer. The label fail is not an address, but denotes a branch to the global failure routine.

\[ \text{cut}(X) \]

Implement the cut operation. Move X into the \( r(b) \) register; also move the value of \( r(h) \) in this choice point into the \( r(hb) \) register. The latter move is an optimization that reduces the number of trailed variables, but is not needed for correctness. The compiler ensures that X contains a pointer to the choice point which was most recent when the current predicate was entered.

\[ \text{hash}(T, R, N, L) \]

Look up register R in a hash table located at label L. The hash table contains atomic terms (when \( T=\text{atomic} \)) or the main functors of structures (when \( T=\text{structure} \)). If R is not in the hash table, then execution falls through to the next instruction. Otherwise execution continues at the label contained in the hash table. When \( T=\text{structure} \) the compiler guarantees that R points to a structure. The following is an example of hash table code:

```plaintext
... hash(Type, Reg, N, Lbl).  ; Hash Reg into table at Lbl
                        ; Fall through if not present
... label(Lbl).          ; The hash table
hash_length(N).         ; Length of the hash table
pair(E1, Li).           ; N entries
pair(E2, L2).           ...
pair(Ei, Li).           ; Jump to Li if Reg = Ei
... pair(EN, LN).       ...
pair(E, L)              ; A hash table entry. E is either an atom or the main functor of a structure. The label L is the address where execution continues if the supplied value matches E.
```
4. Arithmetic instructions

<table>
<thead>
<tr>
<th>Arithmetic syntax</th>
</tr>
</thead>
<tbody>
<tr>
<td>instr(add(A, B, V)) :- numarg_i(A), numarg_i(B), a_var(V).</td>
</tr>
<tr>
<td>instr(sub(A, B, V)) :- numarg_i(A), numarg_i(B), a_var(V).</td>
</tr>
<tr>
<td>instr(mul(A, B, V)) :- numarg_i(A), numarg_i(B), a_var(V).</td>
</tr>
<tr>
<td>instr(div(A, B, V)) :- numarg_i(A), numarg_i(B), a_var(V).</td>
</tr>
<tr>
<td>instr(and(A, B, V)) :- numarg_i(A), numarg_i(B), a_var(V).</td>
</tr>
<tr>
<td>instr(or(A, B, V)) :- numarg_i(A), numarg_i(B), a_var(V).</td>
</tr>
<tr>
<td>instr(xor(A, B, V)) :- numarg_i(A), numarg_i(B), a_var(V).</td>
</tr>
<tr>
<td>instr(not(A, V)) :- numarg_i(A), a_var(V).</td>
</tr>
<tr>
<td>instr(sll(A, B, V)) :- numarg_i(A), numarg_i(B), a_var(V).</td>
</tr>
<tr>
<td>instr(sra(A, B, V)) :- numarg_i(A), numarg_i(B), a_var(V).</td>
</tr>
</tbody>
</table>

All arithmetic instructions assume that their operands are dereferenced and destructively overwrite the result register. All perform operations on integers with correct tag and return a result with correct tag, trapping if either operand or the result is not a integer. Arithmetic semantics are:

- add(X, Y, Z) \[ Z \leftarrow X + Y \]
- sub(X, Y, Z) \[ Z \leftarrow X - Y \]
- mul(X, Y, Z) \[ Z \leftarrow X \times Y \]
- div(X, Y, Z) \[ Z \leftarrow X / Y \]
- and(X, Y, Z) \[ Z \leftarrow X \text{ and } Y \] (bitwise and)
- or(X, Y, Z) \[ Z \leftarrow X \text{ or } Y \] (bitwise or)
- xor(X, Y, Z) \[ Z \leftarrow X \text{ xor } Y \] (bitwise exclusive or)
- sll(X, Y, Z) \[ Z \leftarrow X \ll Y \] (logical shift of X left Y places)
- sra(X, Y, Z) \[ Z \leftarrow X \gg Y \] (arithmetic shift of X right Y places)
- not(X, Z) \[ Z \leftarrow \text{not } X \] (bitwise invert X into Z)

5. Procedural control flow instructions

<table>
<thead>
<tr>
<th>Procedural syntax</th>
</tr>
</thead>
<tbody>
<tr>
<td>instr(procedure(N/A)) :- atom(N), natural(A).</td>
</tr>
<tr>
<td>instr(call(N/A)) :- atom(N), natural(A).</td>
</tr>
<tr>
<td>instr(return).</td>
</tr>
<tr>
<td>instr(simple_call(N/A)) :- atom(N), natural(A).</td>
</tr>
<tr>
<td>instr(simple_return).</td>
</tr>
<tr>
<td>instr(label(L)) :- lbl(L).</td>
</tr>
<tr>
<td>instr(jump(L)) :- lbl(L).</td>
</tr>
<tr>
<td>instr(allocate(Perms)) :- natural(Perms).</td>
</tr>
<tr>
<td>instr(deallocate(Perms)) :- natural(Perms).</td>
</tr>
</tbody>
</table>

procedure(P) \hspace{1cm} The entry point of procedure P.

call(N/A) \hspace{1cm} Call the procedure N/A, assuming a fixed location for the arguments. The arguments of N/A are sequentially loaded into argument registers. By default the registers used are numbered from zero, i.e. \[ r(0), r(1), \ldots \]. This call is used for all user-defined predicates. It may be nested, but must be surrounded by an allocate-deallocate pair when used in the body of a predicate.

return \hspace{1cm} Return from a call.
simple_call(N/A) Simple call of the procedure N/A, assuming the same argument passing as call(N/A). This is a one-level call; it may not be nested. It does not require a surrounding allocate-deallocate pair. It can be implemented by saving the return address in a fixed register. This instruction is useful for interfacing with assembly routines.

simple_return Return from a simple call.

label(L) Denotes a branch destination. The label fail is not an address, but denotes a branch to the global failure routine.

jump(L) Jump unconditionally to label L. The label may be to the first instruction of another procedure N/A or it may be internal to the current procedure. The label fail is not an address, but denotes a branch to the global failure routine.

allocate(N) Create an environment of size N on the local stack, i.e. a new set of N permanent variables which are denoted by p(I). Typically, the only state registers stored in the environment are r(e) and r(cp). The environment must NOT contain the r(b) register.

deallocate(N) Remove the top-most environment (which is of size N) from the local stack.

6. Pragmas

<table>
<thead>
<tr>
<th>Pragma syntax</th>
</tr>
</thead>
<tbody>
<tr>
<td>instr(pragma(Pragma)) := pragma(Pragma).</td>
</tr>
<tr>
<td>pragma(align(V,N)) := a_var(V), pos(N).</td>
</tr>
<tr>
<td>pragma(push(term(Size))) := pos(Size).</td>
</tr>
<tr>
<td>pragma(push(cons)).</td>
</tr>
<tr>
<td>pragma(push(structure(A))) := pos(A).</td>
</tr>
<tr>
<td>pragma(push(variable)).</td>
</tr>
<tr>
<td>pragma(tag(V,T)) := a_var(V), a_tag(T).</td>
</tr>
<tr>
<td>pragma(hash_length(Len)) := pos(Len).</td>
</tr>
</tbody>
</table>

align(V,N) At this point the contents of register or permanent V are a multiple of N. This information helps the reordering stage to generate double-word load instructions for the VLSI-BAM processor.

hash_length(N) N is the length of the hash table starting at this point.

push(term(S)) At this point a block of push instructions is about to create a term of size S on the heap.

push(cons) At this point a cons cell (of size two words) is about to be created on the heap. This information helps the reordering stage to generate double-word push instructions for the VLSI-BAM processor.

push(structure(A)) At this point a structure of arity A is about to be created on the heap. This information helps the reordering stage to generate double-word push instructions for the BAM processor.

push(variable) At this point an unbound, initialized variable is about to be created on the heap.

hash_length(N) This is the start of a hash table of length N.
tag(V,T)

The contents of variable V have tag T. This pragma precedes a load or a store with address V. It is used to make loads and stores efficient for processors which do not have explicit tag support.

7. User instructions

This section describes the parts of the BAM language that are never output by the compiler, but only used by the BAM assembly programmer. This is used to write the run-time system in BAM code, so that it is as portable as possible. Additional instructions are jump to register address, creating and decomposing tagged words, non-trapping full-word arithmetic, non-trapping full-word unsigned comparison, and trailing for backrackable destructive assignment. Additional registers are used in implementing the run-time system, and can be mapped to memory locations.

<table>
<thead>
<tr>
<th>Additional instructions</th>
</tr>
</thead>
<tbody>
<tr>
<td>instr(I) :- user_instr(I).</td>
</tr>
<tr>
<td>user_instr(jump_reg(R)) :- reg(R).</td>
</tr>
<tr>
<td>user_instr(jump_nt(C,A,B,L)) :- cond(C), numarg_i(A), numarg_i(B), lbl(L).</td>
</tr>
<tr>
<td>user_instr(ord(A,B)) :- arg(A), a_var(B).</td>
</tr>
<tr>
<td>user_instr(val(T,A,V)) :- a_tag(T), numarg_i(A), a_var(V).</td>
</tr>
<tr>
<td>user_instr(add_nt(A,B,V)) :- numarg_i(A), numarg_i(B), a_var(V).</td>
</tr>
<tr>
<td>user_instr(sub_nt(A,B,V)) :- numarg_i(A), numarg_i(B), a_var(V).</td>
</tr>
<tr>
<td>user_instr(and_nt(A,B,V)) :- numarg_i(A), numarg_i(B), a_var(V).</td>
</tr>
<tr>
<td>user_instr(or_nt(A,B,V)) :- numarg_i(A), numarg_i(B), a_var(V).</td>
</tr>
<tr>
<td>user_instr(xor_nt(A,B,V)) :- numarg_i(A), numarg_i(B), a_var(V).</td>
</tr>
<tr>
<td>user_instr(not_nt(A,V)) :- numarg_i(A), a_var(V).</td>
</tr>
<tr>
<td>user_instr(sll_nt(A,B,V)) :- numarg_i(A), numarg_i(B), a_var(V).</td>
</tr>
<tr>
<td>user_instr(sra_nt(A,B,V)) :- numarg_i(A), numarg_i(B), a_var(V).</td>
</tr>
<tr>
<td>user_instr(trail_bda(X)) :- a_var(X).</td>
</tr>
<tr>
<td>user_reg(r(A)) :- atom(A).</td>
</tr>
</tbody>
</table>

jump_reg(R) Jump unconditionally to the address stored in register R.

jump_nt(C,A,B,L) Compare A and B and jump to L if the comparison is true. The kind of comparison is given by C. This instruction does a full word comparison and never traps. The label fail is not an address, but denotes a branch to the global failure routine.

ord(A,B) Store in B the machine integer that corresponds to the atom or integer in A. This function strips the tag from A, and therefore depends on the target machine and the program that is compiled. It is used to convert atoms and integers into table indices.

val(T,A,V) Create a tagged word in B by combining the tag T and the machine integer in A. This function is the inverse of ord(A,B): In the sequence ord(A1,B), val(T,B,A2) the argument A2 will receive an identical value to A1 if T is the tag of A1.
add_nt(A, B, V)  These arithmetic instructions destructively overwrite the result register
sub_nt(A, B, V)  All perform operations on full words, return a full word, and never
and_nt(A, B, V)  trap. See the previous section on arithmetic for a description of the
or_nt(A, B, V)  operations performed.
xor_nt(A, B, V)
not_nt(A, B, V)
sll_nt(A, B, V)
sra_nt(A, B, V)

trail_bda(X)  Push the address and value of X on the trail stack if the trail condition
               X<r(hb) is satisfied. It is assumed that X is dereferenced. When
detailing, the old value of X is restored. This is used to implement
backtrackable destructive assignment. Only one comparison is neces-
sary for the trail check. The state register r(hb) points to the heap
location which was the top of the heap when the most recent choice
point was created.

8. Instruction arguments

This section defines the syntax of the instructions' arguments.

<table>
<thead>
<tr>
<th>Addressing modes for equal, move and push</th>
</tr>
</thead>
<tbody>
<tr>
<td>% Effective address for equal:</td>
</tr>
<tr>
<td>ea_e(Var)       :- a_var(Var).</td>
</tr>
<tr>
<td>ea_e(VarOff)    :- var_off(VarOff).</td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td>% Effective address for move:</td>
</tr>
<tr>
<td>ea_m(Arg)       :- arg(Arg).</td>
</tr>
<tr>
<td>ea_m(VarOff)    :- var_off(VarOff).</td>
</tr>
<tr>
<td>ea_m(Tag-H)     :- pointer_tag(Tag), heap_ptr(H).</td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td>% Effective address for push:</td>
</tr>
<tr>
<td>ea_p(Arg)       :- arg_i(Arg).</td>
</tr>
<tr>
<td>ea_p(Tag-H)     :- pointer_tag(Tag), heap_ptr(H).</td>
</tr>
<tr>
<td>ea_p(Tag-H+D)   :- pointer_tag(Tag), pos(D), heap_ptr(H).</td>
</tr>
</tbody>
</table>
Other addressing modes

\[
\begin{align*}
\text{heap_ptr}(r(h)) & : - \text{int}(I) . \\
\text{choice_ptr}(r(b)) & : - \text{user_reg}(T) . \\
\text{reg}(r(I)) & : - \text{int}(I) . \\
\text{reg}(T) & : - \text{user_reg}(T) . \\
\text{hreg}(R) & : - \text{reg}(R) . \\
\text{hreg}(F) & : - \text{heap_ptr}(R) . \\
\text{perm}(p(I)) & : - \text{natural}(I) . \\
\text{an_atomic}(I) & : - \text{int}(I) . \\
\text{an_atomic}(T^A) & : - \text{atom}(A) , \text{atom_tag}(T) . \\
\text{an_atomic}(T'(F/N)) & : - \text{atom}(F) , \text{pos}(N) , \text{atom_tag}(T) . \\
\text{a_var}(Reg) & : - \text{reg}(Reg) . \\
\text{a_var}(Perm) & : - \text{perm}(Perm) . \\
\text{arg}(Arg) & : - \text{a_var}(Arg) . \\
\text{arg}(Arg) & : - \text{an_atomic}(Arg) . \\
\text{var_i}(Var) & : - \text{a_var}(Var) . \\
\text{var_i}([Var]) & : - \text{a_var}(Var) . \\
\text{arg_i}(Arg) & : - \text{var_i}(Arg) . \\
\text{arg_i}(Arg) & : - \text{an_atomic}(Arg) . \\
\text{numreg}(Arg) & : - \text{reg}(Arg) . \\
\text{numreg}(Arg) & : - \text{int}(Arg) . \\
\text{numarg_i}(Arg) & : - \text{var_i}(Arg) . \\
\text{numarg_i}(Arg) & : - \text{int}(Arg) . \\
\text{var_off}([Var]) & : - \text{a_var}(Var) . \\
\text{var_off}([Var+I]) & : - \text{a_var}(Var) , \text{pos}(I) . \\
\end{align*}
\]

Tag syntax

\[
\begin{align*}
\text{a_tag}(\text{tatm}) & : * \text{atom} */ \\
\text{a_tag}(\text{tint}) & : * \text{integer} */ \\
\text{a_tag}(\text{tneg}) & : * \text{negative integer} */ \\
\text{a_tag}(\text{tpos}) & : * \text{nonnegative integer} */ \\
\text{a_tag}(\text{tstr}) & : * \text{structure} */ \\
\text{a_tag}(\text{tlst}) & : * \text{cons cell} */ \\
\text{a_tag}(\text{tvar}) & : * \text{variable} */ \\
\text{atom_tag}(\text{tatm}) & : \\
\text{pointer_tag}(\text{tstr}) & : \\
\text{pointer_tag}(\text{tlst}) & : \\
\text{pointer_tag}(\text{tvar}) & : \\
\end{align*}
\]
### Conditionals syntax

<table>
<thead>
<tr>
<th>Condition</th>
<th>Syntax</th>
</tr>
</thead>
<tbody>
<tr>
<td>eq</td>
<td>eq(e).</td>
</tr>
<tr>
<td>eq_ne</td>
<td>eq_ne(ne).</td>
</tr>
<tr>
<td>cond</td>
<td>/* Equal */</td>
</tr>
<tr>
<td>cond(e)</td>
<td>cond(ne).</td>
</tr>
<tr>
<td>cond(ne)</td>
<td>/* Not equal */</td>
</tr>
<tr>
<td>cond(lts)</td>
<td>/* Signed less than */</td>
</tr>
<tr>
<td>cond(les)</td>
<td>/* Signed less than or equal */</td>
</tr>
<tr>
<td>cond(gts)</td>
<td>/* Signed greater than */</td>
</tr>
<tr>
<td>cond(ges)</td>
<td>/* Signed greater than or equal */</td>
</tr>
</tbody>
</table>

### Miscellaneous syntax

<table>
<thead>
<tr>
<th>Syntax</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>hash_type</td>
<td>hash_type(atomic).</td>
</tr>
<tr>
<td>hash_type</td>
<td>hash_type(structure).</td>
</tr>
<tr>
<td>lbl</td>
<td>lbl(fail).</td>
</tr>
<tr>
<td>lbl(N/A)</td>
<td>lbl(N/A) :- atom(N), natural(A).</td>
</tr>
<tr>
<td>lbl(N/A/I)</td>
<td>lbl(N/A/I) :- atom(N), natural(A), int(I).</td>
</tr>
<tr>
<td>nv_flag</td>
<td>nv_flag(nonvar).</td>
</tr>
<tr>
<td>nv_flag</td>
<td>nv_flag(var).</td>
</tr>
<tr>
<td>nv_flag</td>
<td>nv_flag('?').</td>
</tr>
<tr>
<td>regs</td>
<td>% A list of register numbers:</td>
</tr>
<tr>
<td></td>
<td>% (May contain the value 'no' as well)</td>
</tr>
<tr>
<td></td>
<td>regs([]).</td>
</tr>
<tr>
<td></td>
<td>regs([R</td>
</tr>
</tbody>
</table>

### Utility predicates

<table>
<thead>
<tr>
<th>Predicate</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>ground(X)</td>
<td>ground(X) :- nonvar(X), functor(X, _, N), ground(N, X).</td>
</tr>
<tr>
<td></td>
<td>ground(N, _) :- N=:=0.</td>
</tr>
<tr>
<td></td>
<td>ground(N, X) :- N=:=0, arg(N,X,A), ground(A), N1 is N-1, ground(N1,X).</td>
</tr>
<tr>
<td>int(N)</td>
<td>int(N) :- integer(N).</td>
</tr>
<tr>
<td>natural(N)</td>
<td>natural(N) :- integer(N), N&gt;=0.</td>
</tr>
<tr>
<td>pos(N)</td>
<td>pos(N) :- integer(N), N&gt;0.</td>
</tr>
</tbody>
</table>
Appendix E

Extended DCG notation:
A tool for applicative programming in Prolog

1. Introduction

This appendix describes a preprocessor that simplifies purely applicative programming in Prolog. The preprocessor generalizes Prolog's Definite Clause Grammar (DCG) notation to allow programming with multiple accumulators. It has been an indispensable tool in the development of the Aquarius Prolog compiler. Its use is transparent in versions of Prolog that conform to the Edinburgh standard. The preprocessor and a user manual are available by anonymous ftp to arpa.berkeley.edu.

It is desirable to program in a purely applicative style, i.e. within the pure logical subset of Prolog. In that case a predicate's meaning depends only on its definition, and not on any outside information. This has two important advantages. First, it greatly simplifies verifying correctness. Simple inspection is often sufficient. Second, since all information is passed locally, it makes the program more amenable to parallel execution. However, in practice the number of arguments of predicates written in this style is large, which makes writing and maintaining them difficult. Two ways of getting around this problem are (1) to encapsulate information in compound structures which are passed in single arguments, and (2) to use global instead of local information. Both of these techniques are commonly used in imperative languages such as C, but neither is a satisfying way to program in Prolog, for the following reasons:

- Because Prolog is a single-assignment language, modifying encapsulated information requires a time-consuming copy of the entire structure. Sophisticated optimizations could make this efficient, but compilers implementing them do not yet exist.
- Using global information destroys the advantages of programming in an applicative style, such as the ease of mathematical analysis and the suitability for parallel execution.

A third approach with neither of the above disadvantages is extending Prolog to allow an arbitrary number of arguments without increasing the size of the source code. The extended Prolog is translated into standard Prolog by a preprocessor. This report describes an extension to Prolog's Definite Clause Grammar notation that implements this idea.

2. Definite Clause Grammar (DCG) notation

DCG notation was developed as the result of research in natural language parsing and understanding [Pereira & Warren 1980]. It allows the specification of a class of attributed unification grammars with semantic actions. These grammars are strictly more powerful than context-free grammars. Prologs that conform to the Edinburgh standard [Clocksin & Mellish 1981] provide a built-in preprocessor that translates clauses written in DCG notation into standard Prolog.

An important Prolog programming technique is the accumulator [Sterling & Shapiro 1986]. The DCG notation implements a single implicit accumulator. For example, the DCG clause:

\[
\text{term}(S) \rightarrow \text{factor}(A), \ [+], \ \text{factor}(B), \ \{S \text{ is } A+B\}.
\]

is translated internally into the Prolog clause:

\[
\text{term}(S, X1, X4) \leftarrow \text{factor}(A, X1, X2), \ X2=[+1X3], \ \text{factor}(B, X3, X4), \ S \text{ is } A+B.
\]

Each predicate is given two additional arguments. Chaining together these arguments implements the accumulator.
3. Extending the DCG notation

The DCG notation is a concise and clear way to express the use of a single accumulator. However, in the development of large Prolog programs I have found it useful to carry more than one accumulator. If written explicitly, each accumulator requires two additional arguments, and these arguments must be chained together. This requires the invention of many arbitrary variable names, and the chance of introducing errors is large. Modifying or extending this code, for example to add another accumulator, is tedious.

One way to solve this problem is to extend the DCG notation. The extension described here allows for an unlimited number of named accumulators, and handles all the tedium of parameter passing. Each accumulator requires a single Prolog fact as its declaration. The bulk of the program source does not depend on the number of accumulators, so maintaining and extending it is simplified. For single accumulators the notation defaults to the standard DCG notation.

Other extensions to the DCG notation have been proposed, for example Extraposition Grammars [Pereira 1981] and Definite Clause Translation Grammars [Abramson 1984]. The motivation for these extensions is natural-language analysis, and they are not directly useful as aids in program construction.

4. An example

To illustrate the extended notation, consider the following Prolog predicate which converts infix expressions containing identifiers, integers, and addition (+) into machine code for a simple stack machine, and also calculates the size of the code:

```prolog
expr_code(A+B, S1, S4, Cl, C4) :-
    expr_code(';', S1, S2, Cl, C2),
    expr_code(B, S2, S3, C3, C4),
    C3-[plus](C4),  /* Explicitly accumulate 'plus' */
    S; is S3+1.  /* Explicitly add 1 to the size */
expr_code(I, S1, S2, Cl, C2) :-
    atomic(I),
    Cl=[push(I)],
    S2 is S1+1.
```

This predicate has two accumulators: the machine code and its size. A sample call is `expr_code(a+3+b, 0, Size, Code, [])`, which returns the result:

Size = 5
Code = [push(a), push(3), plus, push(b), plus]

With DCG notation it is possible to hide the code accumulator, although the size is still calculated explicitly:

```prolog
expr_code(A+B, S1, S4) -->
    expr_code(A, S1, S2),
    expr_code(B, S2, S3),
    [plus],  /* Accumulate 'plus' in a hidden accumulator */
    {S4 is S3+1}.  /* Explicitly add 1 to the size */
expr_code(I, S1, S2) -->
    {atomic(I)},
    [push(I)],
    {S2 is S1+1}.
```

The extended notation hides both accumulators:
expr_code(A + B) -->>
  expr_code(A),
  expr_code(B),
  [plus]:code,  /* Accumulate 'plus' in the code accumulator */
  [1]:size,  /* Accumulate 1 in the size accumulator */
expr_code(I) -->>
  {atomic(I),
   [push(I)]:code,
   [1]:size.
The translation of this version is identical to the original definition. The preprocessor needs the following declarations:

acc_info(code, T, Out, In, {Out=[T|In]}). /* Accumulator declarations */
acc_info(size, T, In, Out, {Out is In+T}).
pred_info(expr_code, 1, {size,code}). /* Predicate declaration */

For each accumulator this declares the accumulating function, and for each predicate this declares the name, arity (number of arguments), and accumulators it uses. The order of the In and Out arguments determines whether accumulation proceeds in the forward direction (see size) or in the reverse direction (see code). Choosing the proper direction is important if the accumulating function requires some of its arguments to be instantiated.

5. Concluding remarks
   An extension to Prolog's DCG notation that implements an unlimited number of named accumulators was developed to simplify purely applicative Prolog programming. Comments and suggestions for improvements are welcome.

6. References
   [Abramson 1984]
   [Clocksin & Mellish 1981]
   [O'Keefe 1988]
   [Pereira 1981]
   [Pereira & Shieber 1981]
   [Pereira & Warren 1980]
   [Sterling & Shapiro 1986]
Extended DCG notation:
A tool for applicative programming in Prolog

User Manual

1. Introduction

This manual describes a preprocessor for Prolog that adds an arbitrary number of arguments to a predicate without increasing the size of the source code. The hidden arguments are of two kinds:

(1) Accumulators, useful for results that are calculated incrementally in many predicates. An accumulator expands into two additional arguments per predicate.

(2) Passed arguments, used to pass global information to many predicates. A passed argument expands into a single additional argument per predicate.

The preprocessor has been tested under C-Prolog and Quintus Prolog. It is being used by the author in program development, and is believed to be relatively bug-free. However, it is still being refined and extended. The most recent version is available by anonymous ftp to arpa.berkeley.edu or by contacting the author. Please let me know if you find any bugs. Comments and suggestions for improvements are welcome.

2. Using the preprocessor

The preprocessor is implemented in the file accumulator.pl. It must be consulted or compiled before the programs that use it. In Prologs that conform to the Edinburgh standard, such as C-Prolog or Quintus Prolog, the user-defined predicate term_expansion/2 is called when consulting or compiling each clause that is read. With this hook the use of the preprocessor is transparent.

Clauses to be expanded are of the form (Head–>Body) where Head and Body are the head and body of the clause. The head is always expanded with all of its hidden arguments. Table 1 summarizes the expansion rules for body goals. In the table, Goal denotes any goal in a clause body, Acc denotes an accumulator, Pass denotes a passed argument, and Arg denotes either an accumulator or a passed argument. Hidden arguments of body goals that are not in the head have default values which can be overridden. For compatibility with DCG notation the accumulator dcg is available by default. If-then-else is not handled in this version.

The preprocessor assumes the existence of a database of information about the hidden parameters and the predicates to be expanded. Three relations are recognized: a declaration for each predicate, each accumulator, and each passed argument. These relations can be put at the beginning of each file (in which case their scope is the file) or stored in a separate file that is consulted first (in which case their scope is the whole program).

A short example gives a flavor of what the preprocessor does:

```
% Declare the accumulator 'castor':
acc_info(castor, _, _, _, true).

% Declare the passed argument 'pollux':
pass_info(pollux).

% Declare three predicates using these hidden arguments:
pred_info(p, 1, [castor, pollux]).
pred_info(q, 1, [castor, pollux]).
```
Table 1 — Expansion rules for the preprocessor

<table>
<thead>
<tr>
<th>Body goal</th>
<th>Action</th>
</tr>
</thead>
<tbody>
<tr>
<td>(Goal)</td>
<td>Don't expand any hidden arguments of Goal.</td>
</tr>
<tr>
<td>Goal</td>
<td>Expand all of the hidden parameters of Goal that are also in the head. Those hidden parameters not in the head are given default values.</td>
</tr>
<tr>
<td>Goal:L</td>
<td>If Goal has no hidden arguments then force the expansion of all arguments in L in the order given. If Goal has hidden arguments then expand all of them, using the contents of L to override the expansion. L is either a term of the form Acc, Acc(Left, Right), Pass, Pass(Value), or a list of such terms. When present, the arguments Left, Right, and Value override the default values of arguments not in the head.</td>
</tr>
<tr>
<td>List:Acc</td>
<td>Accumulate a list of terms in the accumulator Acc.</td>
</tr>
<tr>
<td>List</td>
<td>Accumulate a list of terms in the accumulator deg.</td>
</tr>
<tr>
<td>X/Arg</td>
<td>Unify X with the left term for the accumulator or passed argument Arg.</td>
</tr>
<tr>
<td>Acc/X</td>
<td>Unify X with the right term for accumulator Acc.</td>
</tr>
<tr>
<td>X/Acc/Y</td>
<td>Unify X with the left and Y with the right term for the accumulator Acc.</td>
</tr>
<tr>
<td>insert(X,Y):Acc</td>
<td>Insert the arguments X and Y into the chain implementing the accumulator Acc. This is useful when the value of the accumulator changes radically because X and Y may be the arguments of an arbitrary relation.</td>
</tr>
<tr>
<td>insert(X,Y)</td>
<td>Insert the arguments X and Y into the chain implementing the accumulator deg. This inserts the difference list X-Y into the accumulated list.</td>
</tr>
</tbody>
</table>

Pred_info(r, l, [castor,pollux]).

The program:
p(X) --> Y is X+1, q(Y), r(Y).

This example declares one accumulator, one passed argument, and three predicates using them. The program consists of a single clause. The preprocessor is used as follows: (bold-face denotes user input)

\c prolog
C-Prolog version 1.5
?- ['accumulator.pl'].
accumulator.pl consulted 9780 bytes 1.7 sec.
yes
?- ['example.pl'].
example.pl consulted 668 bytes 0.25 sec.
yes
?-

Now the predicate p(X) has been expanded. We can see what it looks like with the listing command:

?- listing(p).

p(X, S1, S3, P) :- Y is X+1, q(Y, S1, S2, P, r(Y, S2, S3, P)).

(Variable names have been changed for clarity.) The arguments S1, S2, and S3 which implement the
accumulator castor, are chained together. The argument \( P \) implements the passed argument. It is added as an extra argument to each predicate.

In object-oriented terminology the declarations of hidden parameters correspond to classes with a single method defined for each. Declarations of predicates specify the inheritance of the predicate from multiple classes, namely each hidden parameter.

3. Declarations

3.1. Declaration of the predicates

Predicates are declared with facts of the following form:

\[
\text{pred_info(} \text{Name, Arity, List)}
\]

The predicate \( \text{Name/Arity} \) has the hidden parameters given in \( \text{List} \). The parameters are added in the order given by \( \text{List} \) and their names must be atoms.

3.2. Declaration of the accumulators

Accumulators are declared with facts in one of two forms. The short form is:

\[
\text{acc_info(} \text{Acc, Term, Left, Right, Joiner)}
\]

The long form is:

\[
\text{acc_info(} \text{Acc, Term, Left, Right, Joiner, LStart, RStart)}
\]

In most cases the short form gives sufficient information. It declares the accumulator \( \text{Acc} \), which must be an atom, along with the accumulating function, \( \text{Joiner} \), and its arguments \( \text{Term} \), the term to be accumulated, and \( \text{Left} \) & \( \text{Right} \), the variables used in chaining.

The long form of \( \text{acc_info} \) is useful in more complex programs. It contains two additional arguments, \( \text{LStart} \) and \( \text{RStart} \), that are used to give default starting values for an accumulator occurring in a body goal that does not occur in the head. The starting values are given to the unused accumulator to ensure that it will execute correctly even though its value is not used. Care is needed to give correct values for \( \text{LStart} \) and \( \text{RStart} \). For DCG-like list accumulation both may remain unbound.

Two conventions are used for the two variables used in chaining depending on which direction the accumulation is done. For forward accumulation, \( \text{Left} \) is the input and \( \text{Right} \) is the output. For reverse accumulation, \( \text{Right} \) is the input and \( \text{Left} \) is the output.

To see how these declarations work, consider the following program:

\[
\text{Example illustrating the difference between forward and reverse accumulation:}
\]

\[
\text{Declare the accumulators:}
\]

\[
\text{acc_info(fwd, T, In, Out, Out=[T|In])} \quad \text{Forward accumulator.}
\]
\[
\text{acc_info(rev, T, Out, In, Out=[T|In])} \quad \text{Reverse accumulator.}
\]

\[
\text{Declare the predicates using them:}
\]

\[
\text{pred_info(flist, 1, [fwd]).}
\]
\[
\text{pred_info(rlist, 1, [rev]).}
\]

\[
\text{flist(N, [). List) creates the list \{1. 2, ..., N\}}
\]
\[
\text{flist(0) --> [1].}
\]
\[
\text{flist(N) --> N>0. \{N|fwd. N1 is N-1, flist(N1).}
\]

\[
\text{rlist(N, List, []) creates the list \{N, ..., 2, 1\}}
\]
\[
\text{rlist(0) --> [1].}
\]
\[
\text{rlist(N) --> N>0. \{N|rev. N1 is N-1, rlist(N1).}
\]

This defines two accumulators \( \text{fwd} \) and \( \text{rev} \) that both accumulate lists, but in different directions. The
joiner of both accumulators is the unification Out = [T | In], which adds T to the head of the list In and creates the list Out. In accumulator fwd the output Out is the left argument and the input In is the right argument. This builds the list in ascending order. Switching the arguments, as in the accumulator rev, builds the list in reverse. A sample execution gives these results:

```
1 ?- flist(10, [], List).
   List = [1,2,3,4,5,6,7,8,9,10]
   yes
1 ?- rlist(10, List, []).
   List = [10,9,8,7,6,5,4,3,2,1]
   yes
1 ?-
```

If the joining function is not reversible then the accumulator can only be used in one direction. For example, the accumulator add with declaration:

```
acc_info(add, I, In, Out, Out is I+In).
```

It can only be used as a forward accumulator. Attempting to use it in reverse results in an error because the argument In of the joiner is uninstantiated. The reason for this is that the predicate is/2 is not pure logic: it requires the expression in its right-hand side to be ground.

3.3. Declaration of the passed arguments

Passed arguments are declared as facts in one of two forms. The short form is:

```
pass_info(Pass)
```

The long form is:

```
pass_info(Pass, PStart)
```

In most cases the short form is sufficient. It declares a passed argument Pass, that must be an atom. The long form also contains the starting value PStart that is used to give a default value for a passed argument in a body goal that does not occur in the head. Most of the time this situation does not occur.

4. Tips and techniques

Usually there will be one clause of pred_info for each predicate in the program. If the program becomes very large, the number of clauses of pred_info grows accordingly and can become difficult to keep consistent. In that case it is useful to remember that a single pred_info clause can summarize many facts. For example, the following declaration:

```
pred_info(_, _, List).
```

 gives all predicates the hidden parameters in List. This keeps programming simple regardless of the number of hidden parameters.
Appendix F

Source code of the C and Prolog benchmarks

/* C version of tak benchmark */

#include <stdio.h>

int tak(x, y, z)
int x, y, z;
{
    int a1, a2, a3;
    if (x <= y) return z;
    a1 = tak(x-1, y, z);
    a2 = tak(y-1, z, x);
    a3 = tak(z-1, x, y);
    return tak(a1, a2, a3);
}

main()
{
    printf("%d\n", tak(24, 16, 8));
}

/* Prolog version of tak benchmark */

main :- tak(24,16,8,X), write(X), nl.

tak(X,Y,Z,A) :- X =< Y, Z = A.
tak(X,Y,Z,A) :- X > Y,
    X1 is X - 1, tak(X1,Y,Z,A1),
    Y1 is Y - 1, tak(Y1,Z,X,A2),
    Z1 is Z - 1, tak(Z1,X,Y,A3),
    tak(A1,A2,A3,A).

/* C version of fib benchmark */

#include <stdio.h>

int fib(x)
int x;
{
    if (x <= 1) return 1;
    return (fib(x-1)+fib(x-2));
}
main()
{
    printf("%d\n", fib(30));
}

/* Prolog version of fib benchmark */
main :- fib(30,N), write(N), nl.

fib(N,F) :- N =< 1, F = 1.
fib(N,F) :- N > 1,
    N1 is N - 1, fib(N1,F1),
    N2 is N - 2, fib(N2,F2),
    F is F1 + F2.

/*-------------------------------------------------------------------------------
*/

finclude <stdio.h>

han(n,a,b,c)
{
    int nl;
    if (n<=0) return;
    nl = n-1;
    han(n1,a,c,b);
    han(n1,c,b,a);
}

main()
{
    han(20,1,2,3);
}

/*-------------------------------------------------------------------------------
*/

/* Prolog version of hanoi benchmark */
main :- han(20,1,2,3).

han(N,_,_,_) :- N=<0.
han(N,A,B,C) :- N>0,
    N1 is N - 1,
    han(N1,A,C,B),
    han(N1,C,B,A).

/*-------------------------------------------------------------------------------
*/

/* C version of quicksort benchmark */
# include <stdio.h>

int ilist[50] = {27, 74, 17, 33, 94, 18, 46, 83, 65, 2, 32, 53, 28, 85, 99, 47, 28, 82, 6, 11, 55, 29, 39, 81, 90, 37, 10, 0, 66, 51, 7, 21, 85, 27, 31, 63, 75, 4, 95, 99, 11, 28, 61, 74, 18, 92, 40, 53, 59, 8};

int list[50];

qsort(l, r)
int l, r;
{
    int v, t, i, j;
    if (l<r) {
        v=list(l]; i=l; j=r+1;
        do {
            do i++; while (list[i]<v);
            do j--; while (list[j]>v);
            t=list[j]; list[j]=list[i]; list[i]=t;
        ) while (j>i);
        list[i]=list[j]; list[j]=list[l]; list[l]=t;
        qsort(l, j-1);
        qsort(j+1, r);
    }
}

main()
{
    int i, j;
    for(j=0; j<10000; j++) {
        for(i=0; i<50; i++) list[i]=ilist[i];
        qsort(0, 49);
    }
    for(i=0; i<50; i++) printf("%d ", list[i]);
    printf("\n");
}

/* Prolog version of quicksort benchmark */

main :- range(l, I, 9999), qsort(_), fail.
main :- qsort(S), write(S), nl.

range(L, L, H).
range(L, I, H) :- L<H, L1 is L+1, range(L1, I, H).

qsort(S) :- qsort([27, 74, 17, 33, 94, 18, 46, 83, 65, 2, 32, 53, 28, 85, 99, 47, 28, 82, 6, 11, 55, 29, 39, 81, 90, 37, 10, 0, 66, 51, 7, 21, 85, 27, 31, 63, 75, 4, 95, 99, 11, 28, 61, 74, 18, 92, 40, 53, 59, 8], S).
qsort ([X|L], R, R0) :-
  partition(L, X, L1, L2),
  qsort (L2, R1, R0),
  qsort (L1, R, [X|R1]).
qsort ([], R, R).

partition([Y|L], X, [Y|L1], L2) :- Y<X, partition(L, X, L1, L2).
partition([Y|L], X, L1, [Y|L2]) :- Y>X, partition(L, X, L1, L2).
partition([], _, [], []).
Appendix G

Source code of the Aquarius Prolog compiler

Due to the size of the source code, it has not been included here. The complete Aquarius system including source code will be distributed in Spring 1991. The source code of the compiler may also be obtained from the author.

<table>
<thead>
<tr>
<th>File</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>accumulator.pl</td>
<td>Extended DCG preprocessor</td>
</tr>
<tr>
<td>accumulator_cleanup.pl</td>
<td>Cleanup file needed for preprocessor</td>
</tr>
<tr>
<td>analyze.pl</td>
<td>Dataflow analyzer</td>
</tr>
<tr>
<td>clause_code.pl</td>
<td>Clause compiler</td>
</tr>
<tr>
<td>conditions.pl</td>
<td>Formula manipulation utilities</td>
</tr>
<tr>
<td>compiler.pl</td>
<td>Top level of compiler, includes type enrichment</td>
</tr>
<tr>
<td>expression.pl</td>
<td>Compile arithmetic expressions</td>
</tr>
<tr>
<td>factor.pl</td>
<td>Factoring transformation</td>
</tr>
<tr>
<td>flatten.pl</td>
<td>Flattening transformation</td>
</tr>
<tr>
<td>inline.pl</td>
<td>In-line replacement</td>
</tr>
<tr>
<td>mutex.pl</td>
<td>Mutual exclusion and implication of formulas</td>
</tr>
<tr>
<td>peephole.pl</td>
<td>BAM transformations (except synonym)</td>
</tr>
<tr>
<td>preamble.pl</td>
<td>Part of standard form transformation</td>
</tr>
<tr>
<td>proc_code.pl</td>
<td>Predicate compiler</td>
</tr>
<tr>
<td>regalloc.pl</td>
<td>Register allocator</td>
</tr>
<tr>
<td>segment.pl</td>
<td>Head-body segmentation and goal reordering</td>
</tr>
<tr>
<td>selection.pl</td>
<td>Determinism extraction</td>
</tr>
<tr>
<td>standard.pl</td>
<td>Standard form transformation</td>
</tr>
<tr>
<td>synonym.pl</td>
<td>Synonym optimization</td>
</tr>
<tr>
<td>tables.pl</td>
<td>Compilation tables</td>
</tr>
<tr>
<td>testset.pl</td>
<td>List of test sets</td>
</tr>
<tr>
<td>transform_cut.pl</td>
<td>Cut transformation</td>
</tr>
<tr>
<td>unify.pl</td>
<td>Unification compiler</td>
</tr>
<tr>
<td>utility.pl</td>
<td>Utility predicates</td>
</tr>
</tbody>
</table>
Appendix 2

"Fast Prolog With an Extended General Purpose Architecture"

Bruce Holmer, et al.

Fast Prolog with an Extended General Purpose Architecture

Bruce K. Holmer, Barton Sano, Michael Carlson, Peter Van Roy, Ralph Haygood, William R. Bush, Alvin M. Despain

Computer Science Division
University of California, Berkeley

Joan M. Pendleton
Harvest VLSI Design Center, Inc.

ABSTRACT

Most Prolog machines have been based on specialized architectures. Our goal is to start with a general purpose architecture and determine a minimal set of extensions for high performance Prolog execution. We have developed both the architecture and optimizing compiler simultaneously, drawing on results of previous implementations. We find that most Prolog specific operations can be done satisfactorily in software; however, there is a crucial set of features that the architecture must support to achieve the best Prolog performance. The emphasis of this paper is on our architecture and instruction set. The costs and benefits of the special architectural features and instructions are analyzed. Simulated performance results are presented and indicate a peak compiled Prolog performance of 3.68 million logical inferences per second.

1. Introduction

Logic programming in general and Prolog [1] in particular have become popular for rapid software prototyping, natural language translation, and expert system programming. Prolog's use of dynamic typing, backtracking, and unification place heavy computational demands on general purpose computers. In an attempt to achieve ever higher performance, several special purpose architectures have been proposed and built. Early Prolog architectures [2] were microcoded interpreters. Because no compilation was done, performance was disappointing. Higher performance processors [3-6] have since been based on the Warren Abstract Machine (WAM) [7]. Their instruction sets were derived from the WAM to support execution of Prolog programs. These processors are special purpose, microcoded engines which depend on parallel execution of operations within each relatively coarse-grained instruction for high performance. Initial designs implemented only the instructions that supported the WAM and depended on a host processor for non-WAM computations. To support Prolog built-ins (primitive Prolog operations provided by the system) and system I/O, newer designs incorporate general purpose instructions to minimize dependence on a host. Alternatively, the use of a simple, non-WAM instruction set better supports compiler optimization. Several such special purpose reduced instruction set architectures have been proposed for logic programming [8-11]. These architectures include primitives which support the use of tagged data, pointer dereference, and multi-way branches. Our hypothesis is that providing support for both compiler optimization and low-level operations can best be accomplished by extending a simple general purpose architecture to support Prolog without compromising the general purpose performance.

The performance improvements of recent general purpose architectures over older architectures can be traced to research in which both the compiler and architecture were developed together [12-14]. Architectural features that cannot be used by the compiler or which cannot demonstrate performance improvement are not included. Likewise, architectural features are added which support often used primitive operations. We have adopted this approach from the beginning of our project.

It has been conjectured that commercial special purpose symbolic processing architectures are doomed because they are not commodity items, and consequently, economics prevent them from staying on the leading edge of implementation technology. However, if the architectural features necessary to improve symbolic performance are modest and do not interfere with the general purpose architecture, then as more chip area becomes available, future implementations of general purpose processors can deliver high performance symbolic computing in a standard product. We hope that our work is a step towards this result.

This paper presents the design of a processor based on the Berkeley Abstract Machine (BAM) architecture and motivates its design with the results of our preliminary studies. We also present a brief discussion of the optimizing compiler, a cost/benefit analysis of the architectural features, and the simulated performance. Familiarity with the WAM is helpful. Section 2 summarizes the processor architecture and hardware implementation. Section 3 presents the instruction set along with the results of our studies which motivated instruction selection. The compilation of Prolog programs is described in section 4, and in section 5 we present a cost/benefit analysis of the special features and instructions. Section 6 gives the performance results. The final section concludes with a summary of our results.

2. Processor Architecture and Implementation

The BAM processor is a general purpose, single chip, pipelined processor with extensions to support Prolog execution (Figure 1). Both data and instruction words are 32 bits, and most instructions execute in a single cycle. The main features for Prolog are tag manipulation (integrated into arithmetic and the memory system), a double-word data port to memory, special branch on tag support, and several instructions to support our execution model for Prolog.

The architecture is presented in detail along with our motivations in the subsections below. Retaining a core general purpose architecture imposes constraints on the symbolic extensions. For example, the processor should be able to handle tagged data items as single entities, with no special treatment for the tags. We discuss the ramifications of this on the word format and the virtual memory system. Then we present the architecture's register structure and memory interface. Finally, we present some details of the implementation such as the pipeline structure and our mechanism for

2.1. Word Format

Prolog does not require the user to specify the type of a data item. This requires that run time checking be implemented by adding a tag to each data item to encode the type of that item. Many Prolog processors handle the tag and value fields separately. This approach does not satisfy our goal of integrating tagging into a general purpose architecture. Instead, we use a standard 32-bit word length and place the tag in the most significant four bits of the word. Arithmetic computations and addresses, however, use the entire 32-bit word, so general purpose computations are not affected by Prolog's use of tags. Tag values fixed by the hardware are those for non-negative integers (0000) and negative integers (1111). This selection of tags for integers is a common technique used by Lisp implementations on general purpose machines [15]. We have also fixed the tag value for variable pointers (vvar = 0001) to increase the number of bits available for branch displacements in several Prolog specific instructions. All other tag values are software defined. Our Prolog implementation uses tags similar to those of the WAM.

2.2. Segmented Virtual Addresses

One consequence of using both the tag and value as an address is that each data type is mapped into its own area of virtual memory. For Prolog's execution model one wishes to place several data types in the same stack or heap. One possible solution is to mask (zero) the tag bits of the address before using it to access memory. This solution is not satisfactory when applied to applications not using tags (for example, C programs). To avoid this difficulty, we have introduced a segment table which maps the most significant six bits of an address to a twelve-bit value (Figure 2). An address before mapping is referred to as a short virtual address (SVA), and the 38-bit address resulting from the mapping is referred to as a long virtual address (LVA). This memory segmentation scheme is similar to the segmentation used in the 801 processor [16]. The 801 uses segmentation to extend the virtual address space; however, our primary motivation for using segmentation is to allow multiple data types to be mapped to the same LVA segment. Mapping two bits in addition to the tag bits allows the use of several memory areas for a given data type, each area using a different mapping. At one extreme all data types can be mapped to the same LVA segment (this is equivalent to masking the most significant six address bits). At the other extreme, all SVA segments can be mapped to distinct LVA segments. In our current implementation of Prolog, variable, list, and structure pointers are mapped to the same LVA segment, whereas the environment/choice point stack, the trail stack, and the symbol table are mapped to separate segments.

Another use of segmentation is for sharing data in a multiprocessor system. In this case the 38-bit LVA is used as the global virtual address and sharing of data by cooperating processes is done at the segment level.

2.3. Memory Interface

The high memory bandwidth requirement of Prolog dictates separate instruction and data buses (Figure 1). In addition, we have expanded the data bus to double-word width. A double-word data bus is motivated by Carlson's study [17] of the architectural requirements of high performance Prolog processors. Carlson compiled Prolog programs into basic register transfer level operations and then compacted them into more complex instructions while enforcing microarchitectural constraints. His results show that the best performance/cost tradeoff occurs when the architecture provides a double-word port to data memory.

A double-word memory port improves the performance of term creation and speeds block transfers to and from environments and choice points. Some previous Prolog processors support fast choice point creation and restoration through the use of specialized buffers or shadow registers [3,9]. Such hardware solutions are costly and do not fit our goal of maintaining a general purpose architecture. Instead, we rely on double-word memory operations and on compiler optimization to minimize shallow backtracking [18].

Our processor design is tightly coupled with the cache design. We decided against on-chip caches since, in our case, it is more appropriate to use processor chip area for architectural features and use fast, dense static RAM chips for large caches. To speed cache accesses, however, protection violation and consistency checks and address tag comparisons are done on-chip. More details about the cache interface are given in [19].

2.4. Base Architecture

All programmer visible processor registers are accessed as two sets of 32 registers: the general purpose register set and the special register set. The general purpose registers are used for procedure argument passing, temporary storage, and as stack pointers. The only general purpose register with a preassigned use is the continu
The stage PLA and tion stage or ters are defined push). vide for and opcode. Our design 3.

2.5. Implementation Details

The execution pipeline consists of five stages (Figure 3). All instructions which modify registers or memory do so in the last pipeline stage. Bypassing forwards available results of calculations to instructions following in the pipeline. Hardware interlocks are provided for both load and store delays. If data from a load instruction is used by the next instruction, then the next instruction is delayed by a cycle. Also, memory instructions immediately following a store are delayed by a cycle.

<table>
<thead>
<tr>
<th>Instruction Fetch</th>
<th>R Instruction Read</th>
<th>A ALU</th>
<th>M Memory Read</th>
<th>W Register/Memory Write</th>
</tr>
</thead>
</table>

Figure 3
BAM Processor Execution Pipeline

All instructions are 32 bits with a 6-bit opcode and fixed source register format. Instruction execution is controlled by an opcode pipeline which operates in parallel with the execution pipeline. Each stage of the opcode pipe decodes the opcode associated with that stage of the execution pipeline. Multi-cycle instructions and conditional instructions are implemented using "internal opcodes" [20]. The internal opcodes of multi-cycle instructions are fetched from a PLA and inserted into the opcode pipeline. When an internal opcode is inserted, no instruction is fetched during that cycle. Thus a single external opcode can invoke a sequence of internal opcodes to provide for often used complex operations (for example, pointer dereferencing). Internal opcode insertion is also used for: atomic synchronization operations, for pipeline interlock delays, and for trap and interrupt handling. Conditional execution is implemented by conditionally replacing an opcode in the opcode pipe with an internal opcode. Our design uses 55 external opcodes and 24 internal opcodes; of the internal opi­codes, nine are related to traps (trap, TIP), 13 implement multi-cycle instructions (dref, sex, std, pushd, ldr, jmpcr), and two implement conditional operation instructions (sav, pushr).

"Fast tag logic" is used to implement single-cycle tag-compare-and-branch instructions. The fast tag logic consists of an extra register file which duplicates the tag portion of the general purpose register file and special tag comparison logic which allows quick tag comparison and branch. Previous Prolog processors [3] have also duplicated tag bits to accelerate branching on tag value.

The general purpose register file has two read ports (one single-word and one double-word) and two write ports (both single-word). This port structure provides the bandwidth required by single-cycle double-word memory accesses without greatly increasing the complexity of the register file design.

3. Instruction Set

In this section we present the BAM instruction set. The instructions are divided into three groups: general purpose, Prolog inspired general purpose, and Prolog specific. The general purpose instructions are those which can be found in typical processors. The Prolog inspired instructions are those which are not often present in general purpose processors, but which can still be used for general computation. The remaining instructions are tailored specifically to the requirements of Prolog execution.

The general purpose instructions are summarized in Table 1. It is important to point out that all arithmetic and logical operations operate on the full 32-bit word. Also, conditional branches consist of separate compare and branch instructions. Compare instructions set or clear the TF (true-false) condition code bit, and the branch instructions take the branch when TF is set. Branches, jumps, and calls are delayed by one instruction. The instruction in a branch delay slot can always be executed (br), annulled (turned into a nop) if the branch is taken (bran), or annulled if the branch is not taken (bran). Both directions of annulling are included because Prolog often favors annulling when the branch is taken (for example, branching out of straight-line code to the unification failure routine), whereas conditional branches to the top of a loop (common in procedural languages) favor annulling when the branch is not taken.

The remainder of this section motivates and presents our exten­sions to the general purpose instruction set. A major influence on the design of these extensions was the simultaneous development of an optimizing Prolog compiler. The abstract machine used by the compiler was initially designed using a top-down approach [21]. We assumed a set of data structures similar to those used by the WAM. Knowledge of possible compiler optimizations was applied to the semantics of Prolog to decompose Prolog's general operations into their components. These components, the abstract instruction set, are the instructions and addressing modes required to compile Prolog operations into efficient code. Efficient translation of abstract instructions into the architectural instruction set was a prime influence in the first pass of the instruction set design.

In addition to our studies of abstract instruction sets, we inves­tigated the microarchitectural requirements for high performance Prolog [17] and gathered execution statistics for the VLSI-PLM, a microcoded implementation of the WAM [4]. These investigations pointed out those microarchitectural features that would give the greatest performance gains and the Prolog operations that most need instruction set support.

3.1. Prolog Inspired General Purpose Instructions

Prolog inspired general purpose instructions are those instruc­tions which support Prolog and which also may be useful in the implementation of other languages (Table 2). These instructions include load and store of immediates, single-cycle double-word load and store, and push and pop memory operations.

Immediate can be loaded, stored, or used in a comparison (ldi, sti, std, cmp)). The immediates are tagged and are created by sign-extending a 12 or 17-bit immediate and replacing the four most significant bits with an immediate tag. Load immediate (ldi) is used for creating integers and atoms. Store immediate (sti) is an optimization of a ldi. st sequence and is used to bind an atom with a vari­able that is known at compile time to be unbox.

Double-word memory operations (ldd, std, stdc, pushd, pushdc) are motivated by Prolog's large memory bandwidth require­ments. A double-word store or push is single-cycle only if the source registers form a consecutive, even/odd register pair, because only three registers, two of which must be adjacent, can be read per cycle from the register file. Although non-consecutive double store and push (std, pushd) are two-cycle instructions, this is offset by the
absence of a pipeline stall when they are immediately followed by a memory operation.

Push instructions are included to support compound term creation. Using branch-and-bound search techniques, we determined an optimal set of single-cycle instructions for creation of all possible two and three-word structures. This set of instructions is optimal in the sense that, for our microarchitecture, each structure is created in the smallest number of cycles. The resulting "compound term creation instruction set" favors the idiom of placing two words of data in registers and then moving them to memory using a double-word push. Push operations also allow the fill of the cache line from memory to be skipped if a push incurs a cache miss and also refers to the first word of the cache line [19]. This optimization has been used in a previous Prolog design [5]. The push instructions allow the amount of the increment to be specified, and any general purpose register can be used as a stack pointer.

Prolog requires that variable assignment be undone on backtracking. This unbinding of variables is implemented by recording variable addresses on a "trail" stack. The original WAM model requires several pointer comparisons to determine if trailing is necessary. Our implementation restricts variables to the global stack (which reduces the number of comparisons to one) and uses a compare instruction followed by a conditional push (pushc). The pop instruction is used during backtracking to retrieve variable addresses from the trail stack. The compiler can reduce the amount of trailing and detailing through the use of flow analysis to determine when uninitialized variables [22] can be used (our use of uninitialized variables is different from [22] — we use the same tag for both initialized and uninitialized variables and determine at compile time when destructive assignment is safe).

Unsigned maximum (umax) is provided to simplify the management of the environment and choice point stack pointers. Because these stacks are intermixed, allocation occurs at the maximum of the two stack pointer values.

3.2. Prolog Specific Instruction Set Support

Prolog specific instructions are those instructions which are tailored specifically for efficient execution of Prolog (Table 3). These instructions support tagged pointer creation, two and three-way branch on tag, pointer dereferencing, and unification of atoms.

3.2.1. Tagged Data Support

Pointer creation is accomplished by the load effective address (leac) instruction which calculates an address and then replaces the most significant four bits with an immediate tag. This instruction is used to create pointers to unbound variables and compound terms (lists and structures).

Type checking built-ins are supported with single-cycle compare-and-branch-on-tag instructions (beqeq and bgneq). These instructions also allow the compiler to replace shallow backtracking with a conditional branch on an argument's tag.

Prolog allows unbound variables to be bound together. The resulting reference chain must be dereferenced before subsequent variable binding. WAM instructions always dereference their operands, often resulting in superfluous dereferencing. However, our
optimizing compiler keeps track of which variables are dereferenced and generates explicit dereferences only when necessary. Implementing dereference as a single instruction reduces static code size and allows dereference memory reads to be pipelined, resulting in a tighter loop than the equivalent assembly code [9, 10]. We use the same tag value for both unbound variables and reference pointers (unbound variables are self-referential). The dereference instruction (dref) is implemented as a sequence of internal opcodes.

All of the basic arithmetic and compare instructions (add, sub, and, or, xor, cmp) have a version which traps on 28-bit overflow. These instructions operate on the full 32-bit word, but 28-bit overflow occurs if either of the sources or the result do not have integer tags (0000 or 1111). The trap on 28-bit overflow allows Prolog arithmetic operations to be compiled to fast, safe code which avoids extra instructions for tag overflow checking. If a 28-bit overflow does occur, the trap routine can signal an overflow error or convert the data into an alternative representation.

3.2.2. Unification Support

Unification is one of the primary operations of Prolog: it is used for argument passing, structure creation, structure decomposition, and pattern matching. Although general unification is a complex algorithm, if one is given information about the arguments being unified, the general algorithm can be greatly simplified. This is one of the advantages of the WAM instruction set over an interpreter. Our compiler takes this principle further and propagates information to simplify unification as much as possible.

Analysis of the primitives necessary to support unification of a Prolog variable with an atom [21] motivates the single-cycle unified intermediate instruction (unif) which binds the atom to the variable if the variable is unbound, and otherwise tests them for equality.

Unification of a Prolog variable with a compound term also benefits from special support. Analysis of the primitives necessary to support unification of a Prolog variable with a list or structure [21]
motivates the switch-tag instruction (swt), a three-way branch based on the tag of one register. One direction of the branch is taken if the tag is an unbound variable; a second direction is taken if the tag matches a specified immediate tag (usually list or structure); and a third direction is taken for all other tags. The three-way branch could be implemented using two two-way branches, however, WAM execution statistics (Table 4) show that there is a small but significant performance advantage to the three-way branch.

The LOW RISC processor [8] provides a 5-way branch and the Carmel-2 processor [10] provides a 10-way branch based on the tag of a single register. WAM execution statistics show that such generality is unnecessary for unification of a Prolog variable with a compound term.

When the compiler cannot determine any information about the types of the arguments to be unified, then general unification must be used. In this case one can still take advantage of dynamical properties of the argument types. The common cases of general unification should be done quickly in-line and infrequent cases passed to a general unification subroutine. Analysis of WAM execution (Table 5) indicates that about 70% of all general unifications are simple bindings of an unbound variable with a nonvariable. These statistics motivate the switch-bind instruction (swb), a three-way branch based on the tags of two registers. The conditions of the three branch directions are: variable/nonvariable, nonvariable/variable, and otherwise (order of the arguments matters). This allows the common cases of variable/nonvariable and nonvariable/variable to be done in-line. A general unification subroutine is called for all other cases. Note that although the quick success and quick failure cases are simple to check for, their execution frequency is low enough that we have chosen not to do these checks in-line.

The Pegasus processor [9] supports general unification with a 16-way branch based on two tag bits from each of two registers. The LIBRA processor [11] has a "partial unify" instruction. This single-cycle instruction performs either a nop, a store, a call, or a branch depending on the tags and comparison of the two arguments. It executes the variable/nonvariable case of general unification in four cycles (not counting dereferencing of the arguments). Using switch-bind (swb), BAM executes this case in five cycles. Although the partial unify instruction of the LIBRA has a slight performance advantage, its complexity does not fit with our goal of minimally extending a general purpose architecture.

4. Compilation of Prolog

A significant aspect of our project was the simultaneous development of an optimizing Prolog compiler [21, 23]. The compiler incorporates techniques for determinism extraction and use of destructive assignment. The compiler accepts standard Prolog and produces code for a simple non-WAM abstract machine. Although the compiler uses stacks and data structures similar to WAM implementations, it does not use the WAM during compilation, but instead directly compiles to its own abstract machine. Automatic mode generation (type inferencing) is implemented using abstract interpretation [24]. It derives ground, uninitialized variable [22], and dereference modes. Optimizations are still being implemented, and we expect our performance numbers to improve compared to the numbers listed in the following sections.

Compilation of Prolog is done in three stages. First, the compiler produces code for its abstract machine. Second, this code is macro-expanded into the BAM instruction set. Finally, the BAM code is optimized by a peephole optimizer and instruction reordering stage that maximizes the use of the double-word bus and minimizes the number of nops and pipeline stalls.

5. Cost/Benefit Analysis of Architectural Features and Instructions

In section 3 we motivated our instruction selection based on several sources of information: work on abstract instruction sets for compilers, bottom-up analysis of microarchitectural requirements for high performance Prolog, and analysis of WAM execution statistics. In this section we give a more rigorous validation of the architectural design and instruction selection by analyzing the cost and performance benefits of each special purpose feature and instruction. There has been some work to determine such results for other designs [9, 10, 15], but no complete analysis has been done.

5.1. Cost of Features

Table 6 shows the implementation cost of those features which
extend the BAM beyond a general purpose architecture. Implementation cost is expressed in terms of chip area required to implement the feature and in terms of VLSI design effort required. The chip area is measured in percent of total active area which includes both transistor and wiring area. The chip contains approximately 110,000 transistors, and the total active area is 91 square millimeters using 1.2 μ CMOS. The VLSI layout was done using a symbolic layout editor with custom designed, transistors. and the feature and in terms of VLSI layout mapping. A smaller register file, however, shows less benefit than the compiler, since the microarchitecture and logic design were done carefully to prevent these features from being on the critical path.

Segment mapping requires the greatest area of the special features. This area is primarily due to the 32 by 24-bit register file which contains the segment map. This register file is used to extend the address space as well as perform tag mapping. A smaller register file tailored to tag mapping alone would take less area. The next greatest area consuming feature is the tagged-immediate generation circuitry. This is due in part to the use of three distinct instruction formats for tagged-instantiates. The double-word memory port requires extra ports on the general purpose register file to support the increased bandwidth. The area listed is the difference in size between our four/five-port register file and the more usual three-port register file. The extra pads required by the double-word bus are not included in the cost. After the fast tag logic, the remaining features use a very small portion of the total active area.

<table>
<thead>
<tr>
<th>Feature</th>
<th>Active area</th>
<th>Design complexity</th>
<th>Instructions affected</th>
</tr>
</thead>
<tbody>
<tr>
<td>segment mapping</td>
<td>4.8%</td>
<td>&quot;100% compiled&quot;</td>
<td>idi, cmp, sti, std, lea, uni</td>
</tr>
<tr>
<td>tagged-immediate</td>
<td>2.2%</td>
<td>100% compiled</td>
<td>idi, cmp, sti, std, lea, uni</td>
</tr>
<tr>
<td>double-word memory port</td>
<td>1.9%</td>
<td>95% compiled; 5% by hand</td>
<td>idi, std, stdc, pushd, pushdc</td>
</tr>
<tr>
<td>fast tag logic</td>
<td>1.6%</td>
<td>&quot;100% compiled&quot;</td>
<td>bgeq, bgt, bne, swt, swb, dref, uni</td>
</tr>
<tr>
<td>multi-cycle/conditional</td>
<td>0.1%</td>
<td>100% compiled</td>
<td>stx, std, pushd, push, dref, uni</td>
</tr>
<tr>
<td>tag overflow detect</td>
<td>0.0%</td>
<td>100% by hand (10 gates)</td>
<td>cmp28, add28, sub28, and28, or28, xor28</td>
</tr>
<tr>
<td>total special features</td>
<td>10.6%</td>
<td>99% compiled; 1% by hand</td>
<td></td>
</tr>
</tbody>
</table>

Table 6. Cost of Special Architectural Features

For each special feature of the BAM processor, this table gives the percentage of active area (transistors and wires) required to implement the feature, the design complexity of the layout, and a list of instructions which depend on the feature. The design complexity is given as a percentage of the layout that was automatically generated (using nizers, routers, etc.) and the percentage that was laid out by hand. "100% compiled indicates that less than 30 gates were placed by hand. Multi-cycle/conditional is a subset of internal opcodes—the 0.1% active area refers to the entire internal opcode implementation.

5.3. Benefits of Individual Instructions

Table 8 provides a similar analysis applied to individual instructions or instruction groups, rather than to architectural features. Significant (greater than one percent) performance benefit is obtained from a majority of the special purpose instructions (dref, min, max, lea, pushd, cmp, and bgeq). The multi-cycle pointer dereference instruction (dref) has an average execution time of 1.6 cycles. Macro-expansion of dref into an explicit loop increases the average dereference time to 2.2 cycles. Although the benefit of dref per dereference is only 0.6 cycle, the total performance benefit is significant because of its frequent use. Some of the smaller benchmarks, however, show no benefit for dref due to the complete elimination of dereferencing by compiler optimization. Unsigned maximum (umax) is used during environment and choice point creation. Omission of umax causes the time to increase from one to three cycles. Tagged-pointer creation (lea) is a frequent operation, and its omission adds an extra cycle for tag insertion (using or). Elimination of auto-increment addressing (push, pushd, pushdc) requires one extra cycle for each block allocation. The three-way branch on tag (swt) can be replaced by two bgeq instructions, adding an extra cycle in two of the branch directions. Elimination of the two-way branch on tag (bgeq/ ne) would require a two instruction compare and branch.

The remaining instructions have less than one percent average performance benefit. Because the VLSI-PLM spends about 5% of its time trailing variable addresses, we included special support in the BAM (push). However, due to the compiler’s use of uninitialized variables, which do not have to be trailed, trailing time is reduced to 1.4% in the BAM. Omitting push decreases slow down of 0.7%, which corresponds to trail time increasing from 2 to 3 cycles. Preliminary analysis using macro-expanded WAM for the chat_parrer benchmark indicated that the benefit for pop would be 1.5%. Compiler optimization reduces the number of general unplications.
minimizing the benefit of swb. Our initial studies also overestimated the benefits of special support for unification of atoms (uni, sst, std). Although push, swb, pop, uni, sst, and std provide marginal performance benefit, their implementation uses only features already required by other instructions.

An interesting conclusion about the number of directions needed in multi-way branches can be made from these measurements. Multi-way branches are implemented in the BAM with the swt and swb instructions, which are both single-cycle three-way branches (Table 3). Swt is used for unification of compound terms, for which greater than a three-way branch is not needed (Table 4 and 21). Swb is used for unification of terms whose types are unknown at compile time. It takes care of 70% of these cases (Table 3), which gives an 0.6% execution time improvement (Table 8). If some single-cycle branch took care of 100% of these cases, we calculate the further improvement would be about 0.7%. Given the additional complexity that such a branch implies, we conclude that a multi-way branch with more than three directions is not effective for Prolog.

### 6. Performance Results

Table 9 compares the performance of the BAM processor to that of other Prolog systems. The results for BAM are simulated assuming a 30 MHz clock and include overhead due to cache misses [19]. The simulated system has 128 KB instruction and data caches. The caches are direct mapped and use a write-back policy. They are run in warm start, that is, each benchmark is run twice and the results of the first run are ignored. Cache effects are significant only for the last five programs in Table 9. The cache overhead is greatest for
simple_analyzer, poly_10, and tak; for these programs the overhead ranges from 11% to 38%. For meta_qsort and chat_parser the overhead is less than 3%.

Although programs are usually compiled with automatic mode generation, we have included numbers without modes to show the effect on performance. The average performance improvement due to automatic mode generation is 1.44. The number is higher for some of the smaller benchmarks because mode generation is able to do a better job for them. For example, the qsort and queens_8 benchmarks perform well because the mode information allows the compiler to eliminate most choice point creation and replace variable binding with destructive assignment. The number is lower for the simple_analyzer benchmark because it uses built-in predicates heavily.

The KCM [6], one of the best WAM implementations, has a relatively large amount of specialized hardware to execute a WAM-like instruction set efficiently, whereas the BAM processor uses modest hardware to support an optimizing compiler. We find that the speed advantage of the BAM over the KCM is equal to or greater than the cycle time ratio.

A common measure of Prolog speed is logical inferences per second (LIPS). In general this quantity is ambiguous; however, it is well defined for the naive reverse benchmark. The execution time for naive reverse with automatic modes (Table 9) gives a performance of 3.68 million LIPS.

Table 10 compares the static code sizes of the BAM, the KCM [6], and the SPUR [27] relative to the PLM [3]. Macro expansion of WAM code into SPUR instructions causes the large code size of the SPUR. Static code size for the BAM is surprisingly small, only slightly larger than that of the KCM. This is due to direct compilation into simple instructions, the success of flow analysis in reducing code size, and the appropriateness of the BAM instruction set for Prolog.

<table>
<thead>
<tr>
<th>Benchmark</th>
<th>Quintus</th>
<th>VLSI-PLM</th>
<th>KCM</th>
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<th>auto modes</th>
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<td>(10.3)</td>
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<td>(1.00)</td>
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</table>

Table 9
Performance Results

This table compares the performance of BAM with that of several other Prolog implementations for which benchmark results are available—Quintus Prolog, the VLSI-PLM, and the KCM. Each result is presented as a time in milliseconds followed in parentheses by the ratio to the next BAM time. The Quintus Prolog results are for compiled code executing under Quintus Prolog Release 2.0 on a Sun 3/60. The VLSI-PLM [4] results are simulated assuming a cycle time of 100 ns with no cache misses. The KCM results [5] are derived from actual measurements of a system with a cycle time of 80 ns. The BAM results are simulated assuming a 30 MHz clock and 128 KB instruction and data caches [19]. For BAM, the auto modes and no modes columns give results with and without automatic mode generation. Results are presented for the well-known Warren benchmarks (the first eight in the table), of which query is modified to use integer division in place of the original floating point; for mu, which proves a theorem of Hofstadter's "mu-math"; for prover, a simple theorem prover; for queens_8, which solves the eight queens problem using an incremental generate-and-test strategy; for meta_qsort, a meta-interpreter running Warren's qsort; for simple_analyzer, a flow analyzer analyzing Warren's qsort; for poly_10, which symbolically raises a polynomial to the tenth power; for tak, which executes recursive integer arithmetic; and for chat_parser, which parses a set of English sentences. Further information about the benchmarks may be found in [28]. The benchmarks are available by anonymous ftp from arpa.berkeley.edu

<table>
<thead>
<tr>
<th>Benchmark</th>
<th>BAM / PLM</th>
<th>KCM / PLM</th>
<th>SPUR / PLM</th>
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<tr>
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<td>12.0</td>
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</table>

Table 10
Static code size ratios

This table gives the static code sizes of the BAM, the KCM, and the SPUR relative to the PLM, a micro-coded implementation of the WAM [3]. The BAM code size is calculated from prover, meta_qsort, simple_analyzer, and chat_parser. The KCM code size is from [6]. The SPUR code size is from [27].

7. Conclusions

The primary goal of our research has been to determine a minimal set of extensions to a general purpose architecture necessary for achieving high performance logic programming. At the same time, however, performance of the general purpose architecture has not been compromised. We have identified tagged-immediate support, segment mapping, double-word memory bus, special logic for fast branch on tag, and multi-cycle instruction support as important Prolog specific features. Our measurements justify the utility of push, pointer dereference, and tagged-pointer creation instructions. Our special instructions for trailing and unification of atoms, how-
ever, are of marginal benefit. Finally, we conclude that a multi-way branch with more than three directions is not effective for Prolog.

We have demonstrated that one can extend a general purpose architecture to include explicit support for symbolic languages such as Prolog with modest increase in chip area (11%) and yet attain a significant performance benefit (70%).

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References
Appendix 3

"A CAD Design Environment Based Upon Prolog"

Gino Cheng, William R. Bush, and Alvin M. Despain

Proceedings of ICCAS 1989
A CAD Design Environment Based Upon Prolog

Gino Cheng, William R. Bush, Alvin M. Despain

Computer Science Division
Department of Electrical Engineering and Computer Science
University of California, Berkeley 94720

Abstract

The rapid prototyping of microprocessors requires a high level of automation. An environment suitable for developing application programs which accelerate the design process should provide an efficient method for manipulating data and a powerful programming environment. This paper describes the benefits we have discovered by using PROLOG as the foundation for ASP, a suite of CAD tools tailored towards the automatic generation of microprocessors. PROLOG provides an inherent relational database which is ideal for describing and manipulating a host of elements at all phases of a design, from a behavioral description to a circuit layout. PROLOG also lends itself to heuristical as well as algorithmic programming styles.

1. Introduction

There are many characteristics inherent to data elements in Computer Aided Design (CAD) that make them difficult to represent in a database [1-2]. The difficulty lies in expressing the many different relationships between elements. For example, a wire element may be related to other wire elements by node, by layer, and by location. A CAD tool should be able to generate a set of elements by any of these relations. This paper will show that the relational database inherent in Prolog is well suited for the requirements of a CAD database. An implementation of objects which cover the entire design process is presented.

Although some CAD problems are well understood, most of the problems in CAD are only partially understood or not well defined. Problems of this nature are solved by employing heuristics such as simulated annealing, and rule based expert systems. Problems that are well understood such as the simulation and channel routing are solved by proven algorithms. Problems that are partially understood may have heuristics imbedded within algorithms. Prolog supports both algorithmic as well as heuristic programming techniques which make it an ideal candidate for CAD programming. This paper will illustrate many of the Prolog programming techniques employed in ASP.

ASP [3] is a full-range synthesis system tailored for the development of microprocessors. It produces VLSI masks from instruction set architecture specifications written in Prolog. The system is composed of several hierarchical components that span behavioral, circuit, and geometric synthesis. Behavioral descriptions are transformed into register transfer level descriptions by VIPER [4]. Controller and datapath are realized in sticks by a suite of layout tools in VENOM. The blocks are compacted, placed, and routed by Sticks Pack [5].

This paper will reveal some of the problems associated with representing data for CAD while illustrating the solutions that we have discovered using Prolog. An application of these philosophies, Sticks in Prolog (SIP) is explained in detail and the other abstract levels in ASP...
are introduced. Advantages for using a clause based language for CAD development will be presented by describing the programming methodology employed by ASP.

2. Design Considerations For Implementing CAD Objects

To model the many complex CAD structures as well as the relationships between structures, many CAD environments use object oriented databases. CAD elements, whether they be geometry for a compactor, transition states for a simulator, or logic expressions for a logic minimizer can all be expressed in terms of objects. There are two strategies for representing CAD elements as objects.

In one approach the database provides a set of primitive objects (objects such as polygons, properties, containers, and paths) that model CAD relationships with a representation policy. For example, a container object can be used to describe a common node relationship by placing all objects belonging to a node within the container object. Similarly, a common layer relationship can be represented by placing all objects that share a common layer within the container object [6]. The primitive objects must be capable of representing every data element and relationship that will be necessary for any design. A policy to represent CAD elements with the primitive objects must be chosen. There may be several possible representations of an element within a given set of objects. For example, given a data object of type BOX containing four integer value fields, a box can be represented as a center coordinate with width and length measurements as in CIF, or as a pair of coordinates denoting two opposite corners. Relationships between objects must be explicitly defined. Once established, all CAD applications must adhere to the well defined set of policies.

In another approach, each CAD element is expressed as an object [7-8]. For example, elements such as wires, nets, contacts, transistors, and waveforms are all expressed as tailored objects. Relationships can be expressed implicitly within the objects by adding data fields. For example, a wire element may contain a field describing the layer of the wire or by a pointer to another of the same layer. With this methodology, the representation policy is deeply imbedded within the data objects. Provisions must be made for adding new objects. For example, assume that a system tailored for CMOS circuits must be modified to handle bipolar transistors for a BIMOS circuit. If the data fields chosen for the transistor element are incapable of representing the bipolar transistor, a new data type must be added to the system. Furthermore, all programs that process transistors must be modified to support the new data type. The primary issue in developing a set of data objects to represent CAD elements is determining how much inherent support to offer [9].

2.1. PROLOG as a Database

Relationships between the elements can be expressed in terms of groups. For example, elements in a cell can be grouped by node, by location and by layer. Current object oriented databases for CAD have strict set relations [6-8]. For example, many databases categorize wires by layer, but not location. To find wires of the same layer, one simply calls a generator that returns instances of wires that are of the queried layer. But to find wires of the same grid, one cannot simply generate wires based upon the grid information, but must generate wires by layer and filter out the wires that are not of a common grid. Data in Prolog is linked by structure and by value. Thus, the procedure for generating all wires on the metal 1 layer is the same as the procedure for generating all wires on row 5, or generating all wires of node vdd, or generating all the wires of row 5 and node vdd in metal 1. Prolog also provides
structures such as binary trees and sorted lists. These constructs make accesses to the ASP Prolog database very uniform.

In ASP, each CAD element is expressed as an object. Elements ranging from behavioral descriptions of architectures to logic equations for a module generator to offset contacts in an ALU layout are all directly expressed in and referenced through Prolog. In Prolog there is no syntactic or semantic difference between a procedure call and a database query. This makes the introduction of new data types very simple. Clauses that process new data types can be easily integrated into the system. There are thirty different representations of a design, each with a set of data objects. One of the lowest levels, Sticks in Prolog, will be described in detail in the next section.

2.2. Sticks in PROLOG

Sticks in Prolog (SIP) is a grid based sticks representation in Prolog that supports hierarchy and parameterized elements. Module generators or human designers generate SIP files which are converted to mask geometry by the STICKS-PACK compactor. In SIP, VLSI elements are modeled as facts. Attributes for the elements are represented as atoms within the facts. Currently, the SIP language consists of four facts representing VLSI elements:

- wire(Layer, pt(X1, Y1), pt(X2, Y2), Width, Net).
- cont(Type, pt(X1, Y1), Offset, Net).
- transistor(Type, pt(X1, Y1), pt(X2, Y2), pt(X3, Y3), Width, Net).
- pin(pt(X1, Y1), Layer, Width, Label, Cell).

Layer are of the atoms: \( m1, m2, p, pd, nd \)

These represent the physical layers of the element (metal1, metal2, poly, P-diffusion, or N-diffusion).

Contact offsets are of the atoms: \( nw, nn, ne, ww, nof, ee, sw, ss, se \)

Contact types are of the atoms: \( m1m2, mLpd, mLnd, mlp \)

Width, XY coordinates, W, and L are integers. Nets are atoms that represent the connectivity node of the element. Elements of the same node are electrically connected. Nodal information is extracted by a net extracting program. \( pt(X, Y) \) represents a point location at \((X, Y)\). Transistors have 3 point locations, one for the source, one for the gate, and one for the drain. Each location has a separate node.

Example: An Inverter in SIP:

- wire(m1, pt(0,0), pt(0,3),2,vdd).
- wire(m1, pt(0,1), pt(1,1),2,vdd).
- wire(m1, pt(10,0), pt(10,3),2,vss).
- wire(m1, pt(10,1), pt(11,1),2,vss).
- wire(m1, pt(8,3), pt(2,3),2,vss).
- wire(m1, pt(6,3), pt(6,3),2,vss).
Different CAD applications often generate different sets of elements. For example, the simulator may generate all of the elements that are of nodes adjacent to a given node. The compactor may generate all of the elements that are of the same grid and layer as a given element. The floorplanner may generate all of the terminals of a given cell side. With the SIP representation, data elements can be generated by any combination of characteristics very easily. For example all of the wires that are of ml of node vdd which have a width greater than 3 can be generated in two lines of Prolog:

```
wire(ml, P1, P2, Width, vdd),
Width > 3,
```

This representation also allows fields to be easily parameterized within a cell. For example,

In a cell definition we have parameterized an output transistor with the statement:

```
parameter(outputtrans, pt(2, 3)).
```

A call to the following clause would permit the modification of the W/L ratio of any transistor that has been parameterized.

```
modsize(Name, Neww, Newl):-
  parameter(Name, pt(Xloc, Yloc)),
  retract(trans(Layer, pt(Sy, Sy), pt(Xloc, Yloc), pt(Dx, Dy), _, _, Ns, Ng, Nd)),
  assert(trans(Layer, pt(Sy, Sy), pt(Xloc, Yloc), pt(Dx, Dy), Neww, Newl, Ns, Ng, Nd)), !.
```

```
modsize(Name, Neww, Newl):-
  writeln("transistor not found"), !.
```
This flexibility allows tools to address and modify specific elements within any context. For example, a program that tries to optimize the performance of a circuit containing many cells can do so by adjusting the W/L ratio of the output transistors. With the output transistors parameterized, the program can reference the output transistors from any cell simply as "outputrans" regardless of the transistor's environment.

SIP provides an excellent abstraction of VLSI layout for an automated module generators that produce sticks layout, for example, the following clause:

```
makeinver(Vddgrid, Vssgrid, Ingrid, Outgrid, Pw, Pl, Nw, Nl):-
    Pdgrid is Vddgrid - 1,
    Ndgrid is Vssgrid + 1,
    assert(wire(m1, p(2, Vddgrid), p(3, Pdgrid), 1, unk)),
    assert(wire(m1, p(3, Vddgrid), p(2, Ndgrid), 1, unk)),
    assert(wire(m1, p(1, Vddgrid), p(5, Vddgrid), 1, unk)),
    assert(wire(m1, p(4, Pdgrid), p(4, Ndgrid), 1, unk)),
    assert(wire(m1, p(4, Outgrid), p(5, Outgrid), 1, unk)),
    assert(wire(p, p(3, Pdgrid), p(3, Ndgrid), 1, unk)),
    assert(wire(p, p(0, Ingrid), p(3, Ingrid), 1, unk)),
    assert(wire(c0111(m1d, p(2, Pdgrid), nof, unk)),
    assert(wire(c0111(m1d, p(2, Ndgrid), nof, unk)),
    assert(wire(c0111(m1d, p(4, Pdgrid), nof, unk)),
    assert(wire(c0111(m1d, p(4, Ndgrid), nof, unk)),
    assert(wire(c0111(m1d, p(1, Pdgrid), p(2, Pdgrid), p(3, Pdgrid), Pw, Pl, unk, unk, unk)),
    assert(wire(c0111(m1d, p(1, Ndgrid), p(2, Ndgrid), p(3, Ndgrid), Nw, NI, unk, unk, unk)), !.
```

will generate an arbitrarily sized inverter with variable input and output locations. Nodal information is deduced by the extractor. ROMs, PLAs, and other regular layout structures can be generated in a similar fashion.

3. PROLOG Programming for CAD

There has been a growing trend in CAD to develop tools that use both algorithmic and rule-based programming styles [10-11]. Algorithms are generally fast, but are inefficient at handling problems that have many special cases. Rule-based systems are well suited for solving problems with many special cases or problems that are not well defined. Rule-based systems have generally been slow. Rules in such a system must be looked up and efficient management systems have not yet been developed. Many CAD problems, such as simulation, have algorithmic solutions, but most problems, such as routing and logic minimization, can be solved by a host of methods.

Prolog provides an environment for both algorithmic and rule-based programming styles. Several examples of both styles have been implemented in ASP. An example of how simulated annealing is implemented in Prolog is illustrated in the Appendix. The clausal nature of Prolog allows rules to be easily updated or modified. Algorithms can also be expressed in a simple and intuitive manner which makes Prolog a language ideal for rapid prototyping.

Prolog source code is typically 10-100 times more dense than C or Fortran source code performing the same function. This makes Prolog systems much more readable and
maintainable. For a large system such as a silicon compiler, this has turned out to be essential.

3.1. PROLOG Programming Methodology Employed by ASP

There are three basic formats for Prolog clauses are employed by ASP:

Procedural Clauses: These clauses work to achieve a certain value or state without failing. Examples of such clauses include arithmetical functions and list manipulations.

/* The mindist routine finds the minimum spacing distance between two objects of layer and width. The 'space' routine returns the minimum spacing distance between two layers, and the width routine determines the minimum width of a layer */

mindist(Layer1, Width1, Layer2, Width2, Distancebetweenobjects):
  space(Layer1, Layer2, Distance),
  width(Layer1, Widthspace1),
  Widthmod1 is Width1*Widthspace1,
  width(Layer2, Widthspace2),
  Widthmod2 is Width2*Widthspace2,
  Distancebetweenobjects is Widthmod1 + Widthmod2 + Distance.

Filtering Clauses: These clauses interpret a given set of data elements differently depending upon the values of certain data fields. If-Then, and Case constructs can be expressed through these clauses.

/* checkconstr determines how to space two elements. Each sub-clause filters out a certain condition. If the elements are on the same row, the spacing is irrelevant. If the elements are contacts, they can not be stacked upon each other and must be spaced accordingly. If the elements are not contacts and of the same node, the spacing doesn't matter, otherwise the elements must be spaced */

checkconstr(Layer1, Width1, Node1, Row1, Layer2, Width2, Node2, Row2, Layer2):
  Row1=Row2.
checkconstr(Layer1, Width1, Node1, Row1, Layer2, Width2, Node2, Row2, Layer2):
  contacts(Layer1, Layer2).
  
Generator Clauses: These clauses generate sets of elements through backtracking or the bagof construct in Prolog.
Malcebox, a routine that creates boxes from various elements, first processes wires, followed by contacts and transistors. /*

makebox:-
    wire(Layer, pt(X1, Y1), pt(X2, Y2), Wid, Node),
    ..... fail.
makebox:-
    con(Type, pt(Row, Y), Oset, _),
    ..... fail.
makebox:-
    trans(Type, pt(Sx, Sy), pt(Gx, Gy), pt(Dx, Dy), W, L, Sn, Gn, Dn),
    ..... fail.
makebox.

4. Conclusion

Prolog provides a relational database and a powerful programming environment. The relational database is easy to use, can represent all CAD objects, and provides a flexible interface to the programming environment. The clausal nature of Prolog provides an environment suitable for algorithmic and rule based programming styles. The success of ASP has shown that Prolog is a robust language well suited for CAD development.

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5. Appendix

% Simulated Annealing package
% You provide the move set, stopping criterion, and number of inner loop iterations.

siman(InitTemp, State0, Cost, Finalstate, Finalcost):
   doOuter(InitTemp, State0, Cost0, Finalstate, Finalcost).

% Outer Loop
doOuter(Temp, State0, Cost, State0, Cost):
   endhere(Temp, State0, Cost). % Outer loop complete by criterion endhere

doOuter(0, Temp, State, Cost, Finalstate, Finalcost):
   doInner(0, Temp, State, Cost, Newstate, Newcost),
   updatetemp(Temp, NewT),

% Inner Loop
doInner(Count, Temp, State, Cost, State, Cost):
   maxinnercount(Mcount),
   Count > Mcount. % inner loop complete

doInner(Count, Temp, State, Cost, State, Cost):
   gennewstate(State, Newstate, Newcost), % create a new state by move
   Deltacost is Cost - Newcost,
   accept(Deltacost, Temp),
   Nextcount is Count + 1,
   doInner(Nextcount, Temp, Newstate, Newcost, Finalstate, Finalcost).

doInner(Count, Temp, State, Cost, Finalstate, Finalcost):
   % new state not accepted
   Nextcount is Count + 1,
   doInner(Nextcount, T, State, Cost, Finalstate, Finalcost).

accept(Deltacost, Temp): % Good move
   Deltacost <= 0.

accept(Deltacost, Temp): % Random factor
   Aexp is -\*Deltacost/Temp,
   Afactor is exp(Aexp),
   random(Randnum),
   Randnum < Afactor.

updatetemp(Temp, Newtemp):
   Newtemp is Temp *0.04, 1.

maxinnercount(100).
6. References


Appendix 4

"The Validation of a Multiprocessor Simulator"

Tam Nguyen and Vason Srini

July 25, 1989
The Validation of a Multiprocessor Simulator

Tam M. Nguyen     Vason P. Srinivasan

Computer Science Division
University of California
Berkeley, CA 94720

Abstract

One of the key steps in performance prediction of multiprocessor systems using simulations is the validation process. A step in the validation process consists of sequential execution of benchmark programs on the multiprocessor simulator and a uniprocessor simulator, and comparing the results and performance measurements data. The simulated cycle count, simulator overhead, operation count, and memory access count are identified to be the key performance data needed for the comparison. This process is illustrated using the multiprocessor NuSim for the parallel execution of Prolog programs and the uniprocessor simulator VPsim. For large programs, the counts obtained from the two simulators are within 10% of each other.

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July 25, 1989
1 Introduction

Simulation is an accurate and effective approach in predicting performance of a new multiprocessor system. Taking into account the many intricate details in the hardware and software designs, the degree of accuracy depends on how much detail is included in the simulator. To ensure that the simulator accurately reflects the real system (yet to be built), the simulator must be carefully validated for correct functional as well as timing results.

The validation process is carried out primarily by comparing performance data from the new simulator with known data obtained from previously validated sources. The validation process itself can be quite tedious and difficult, with massive amounts of information that need to be analyzed. In this paper, we present our approach to validation. The process involves sequential execution of benchmark programs on the multiprocessor simulator and a uniprocessor simulator, comparing results and performance data.

2 Validation Methodology

There are many approaches to the validation of a simulation model [Sar88]. The concept of our approach to validation is quite simple: comparing new, unverified results with previously known answers. The more difficult task is the careful consideration of the many different factors that can affect the results and the degree of these effects. The validation process for a computer system simulator is best done in a stepwise fashion. The exact details of the necessary steps depends on the availability of the known result, or the basis, used for comparison.

In this paper, the term host designates the machine on which the simulator is run and target refers to the computer architecture/system being simulated. Validation refers to the process of ensuring that the simulator is coded correctly and that it accurately models the target.

In the initial phase, where a paper design is the only basis available, validation of the simulator usually consists of:

1. Manually checking for correct coding according to the paper design.
2. Running the simulator and checking for functional correctness, comparing the results with manually worked out solutions.
3. Manually checking the timing of sub-blocks in the simulator.
4. Running the simulator to obtain timing estimates.
5. Running simulator with instrumentation turned on to capture dynamic execution statistics.
The term *manually* used above refer to the *ad hoc* approach of eyeballing (for steps 1 and 3), hand calculations (step 2), or writing small, very special purpose software tools to accomplish the tasks. This approach is very tedious and error prone, but is often the only possible way at this phase since a paper design is the only available basis. In the last step, the monitor facility for instrumentation should not affect the timing.

Once the initial simulator is validated, it may be used as a basis for validating other simulation systems. The validation process can now be done with a greater degree of automation, and thus achieving greater efficiency. However, great care must still be taken to understand the factors that cause discrepancies.

The validation process of a multiprocessor system\(^1\) simulator involves the following steps:

1. *Sequential execution on one processor.* This is done to test the processor module of the simulator and the relevant support modules such as assembler and loader.

2. *Parallel execution on one processor.* This is a degenerate case, done to measure the overhead of parallel execution.

3. *Parallel execution on two processors.* This is a special case for testing interprocessor communication with no interference since there is exactly one sender and one receiver.

4. *Parallel execution on three or more processors.* This is the general case of parallel execution, with potential for interference on shared resources such as the memory and communication channels. It is also used to test the full extent of the parallel execution model. As more processors are added to the configuration, the saturation of shared resources will occur and bottlenecks will appear.

In this paper, we present the application of the first step of validation of a multiprocessor simulator, using a previously validated uniprocessor simulator as a basis. Since there are architecture and execution model variations in the two simulators, their results are compared for proximity, not for exact equality. The following sections provide details on the simulators and the validation approach.

3 Simulator Descriptions

The validation process is demonstrated using two simulators: VPsim and NuSim. Both simulators provide an abstract machine engine for fast execution of the Prolog language. VPsim is a previously validated simulator to be used as the basis of comparison for NuSim.

\(^1\)The term *multiprocessor system* is used to include both the multiprocessor architecture and the parallel execution model.
3.1 VPsim

VPsim is a register transfer level simulator for the VLSI-PLM [STN88]. This chip is a VLSI implementation of a high performance engine for Prolog, a modified version of the abstract machine proposed by Warren [War83]. VPsim is written in the C language, consisting of 4500 lines of C code and 9000 lines of microcode operations (register transfers, CPU operations and microbranches).

To verify VPsim's functional correctness, a wide variety of Prolog programs were run on VPsim and compared with those obtained from runs on software Prolog environments such as Quintus Prolog. Because VPsim is microcode driven, the microstates automatically provide accurate timing, with each microstate being executed in exactly one processor cycle. Gate and transistor level simulations of the VLSI-PLM chip are compared against the results from VPsim. The fabricated chip has passed an extensive testing process and has successfully executed a number of benchmark programs. Work is in progress to interface the chip with a cache and memory board to be used as a coprocessor for the SUN workstation.

From the perspective of this paper, VPsim is a solid simulator that has been well tested and has been verified by the hardware. It is an available resource that can be used as a basis for testing other simulation systems.

3.2 NuSim

To carry out our study in parallel execution of Prolog, we need an accurate and flexible tool to be used as a testbed for new ideas. We approach our study from a system designer's point of view, working with the complete system from software execution model to hardware support for high performance. We are particularly interested in practical designs that can be built in reasonable time. For these reasons, we base our multiprocessor study on our knowledge and experience with sequential execution of Prolog on the VLSI-PLM. In addition to the Prolog specific instructions, the chip contains a number of simple general purpose instructions and primitive support for synchronization. This makes it a good candidate building block for a multiprocessor system.

A simulator can best serve our interest in hardware support for high performance. The result obtained from a simulation run reflects a composite effect of many intricate details that can not be easily formulated or calculated. By varying the parameters of the simulator, the effect that each parameter has on overall performance can be measured.

We have constructed a new simulation system, called NuSim, to facilitate our studies of parallel execution models and the underlying multiprocessor architectures. This simulator framework allows for the complete system simulation: from the instruction set level to the memory architecture level with caches and coherency protocols. The key feature of this simulator framework is flexibility, which allows for extensive instrumentation and continual updates and changes. The modular design identifies main features of the execution model.
and the architectures being simulated as cleanly separated modules with clearly defined interfaces. This allows for easy modifications to the individual modules to support new execution models and architectures.

NuSim is an event-driven simulator, with the events being memory accesses ordered by time. This technique simulates a multiprocessor using a uniprocessor. NuSim consists of 16000 lines of C code and two small machine dependent routines to save and restore the coroutine stacks. It is fairly portable. currently running under 4.3 BSD Unix on the VAX 735 and Sun 3, and under System V Unix on an Intel 386 personal computer.

Figure 1 shows the structure of the NuSim simulator. Two of the major modules of the simulator are the processor module and the memory system module. The processor module emulates the VLSI-PLM instruction set, and is thus comparable to VPsim. The memory system simulates a multi [Bel85] memory architecture, with each processor having a local cache and all caches communicate with main memory via a single bus. The caches are kept consistent via a hardware consistency protocol. In the context of this paper, these two modules form the core of the simulator to which the validation process is applied. The question at hand is: how well does NuSim simulate a VLSI-PLM?
3.3 Simulator Differences

Although both NuSim and VPsim essentially simulate the VLSI-PLm chip, they were created for very different purposes. VPsim was designed as a simulator for a very specific microarchitecture of a Prolog processor. Details of the VLSI-PLm microarchitecture are “hard-wired” into the microcode, in terms of what micro-operations are possible and the constraints in packing the micro-operations into a micro-state. On the other hand, NuSim was conceived as a more general purpose multiprocessor simulator for system integration, dealing at all levels from hardware architecture to software execution model. It will be used to experiment with different architectures and execution model tradeoffs.

Because of the different goals in creating the simulators, there are a number of differences between them. These differences are identified to help us understand the differences in performance numbers. The following are some differences between VPsim and NuSim (running sequential code):

- **Simulation level.** VPsim is a register-transfer-level, cycle-by-cycle simulation, while NuSim is an event-driven simulator which step by memory access. The clock of VPsim is incremented each cycle, while the clock of NuSim is incremented by a value obtained from table lookup.

- **cdr-coding.** VPsim uses cdr-coding, while NuSim does not. Cdr-coding is a compressed representation for list elements stored in consecutive memory locations. It requires a bit to indicate if the next location is the car of the next element. Cdr-coding is eliminated because its complexity has caused many subtle bugs in the microcode while contributing little to the overall performance [Dob67].

- **Instruction fetch.** NuSim does instruction fetch on demand and accounts time for all fetches. VPsim does prefetching, which does not charge time for all fetches, but may spend time to fetch unnecessarily.

- **Memory system.** Nusim has a cache/memory system with realistic values for memory access time. It accounts time for cache misses and block transfers from memory. VPsim has single (processor) cycle memory.

- **Prolog builtins.** VPsim treats some Prolog builtins (language predefined routines) as external functions, and ships data outside the VLSI-PLm processor for processing by the host. A varying amount of time is charged for the data shipment (3 to 10 cycles), but no time is charged for executing the external function. VPsim also implements some Prolog builtins in the library using VLSI-PLm assembly code. NuSim, on the other hand, executes all Prolog builtins inside the processor, and charges time for them as normal instructions. In NuSim, all builtins are written in C code.
4 The Validation Example

In this section, we will compare the performance results of NuSim to those of VPsim to see how closely NuSim simulates a VLSI-PLM processor. Many benchmarks were run on both NuSim and VPsim, and their execution outputs were compared for functional correctness. A group of benchmarks were chosen for closer timing evaluation. These benchmarks differ widely in static code size and dynamic memory usage and execution time.

We have identified a number of measurements for comparison. They are: static code size, cycle count, simulation overhead, operation count, and memory access count. Each type of measurement provides a different perspective of the simulation results, helping to understand the similarity and differences between the two simulators and at the same time validating the results of NuSim.

<table>
<thead>
<tr>
<th>Benchmark</th>
<th>NS code</th>
<th>VP code</th>
<th>NS/VP</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>bintree</td>
<td>181</td>
<td>198</td>
<td>0.91</td>
<td>build a 6-node binary tree</td>
</tr>
<tr>
<td>compiler_bintree</td>
<td>11409</td>
<td>12488</td>
<td>0.91</td>
<td>compiling the bintree program</td>
</tr>
<tr>
<td>compiler_plml</td>
<td>11613</td>
<td>12750</td>
<td>0.91</td>
<td>compiling portion of the compiler</td>
</tr>
<tr>
<td>hanoi</td>
<td>91</td>
<td>82</td>
<td>1.11</td>
<td>towers of hanoi for 8 disks</td>
</tr>
<tr>
<td>mumath</td>
<td>262</td>
<td>251</td>
<td>1.04</td>
<td>Hofstadter’s mumath problem for multi</td>
</tr>
<tr>
<td>newchat</td>
<td>8018</td>
<td>8446</td>
<td>0.95</td>
<td>parsing sentences with the chat parser</td>
</tr>
<tr>
<td>newvl</td>
<td>164</td>
<td>109</td>
<td>1.11</td>
<td>naive reverse a 30-element list</td>
</tr>
<tr>
<td>palin25</td>
<td>290</td>
<td>239</td>
<td>1.12</td>
<td>palindrome for a 25-character string</td>
</tr>
<tr>
<td>puzzle</td>
<td>1135</td>
<td>1049</td>
<td>1.10</td>
<td>solve a puzzle</td>
</tr>
<tr>
<td>q5</td>
<td>249</td>
<td>163</td>
<td>1.53</td>
<td>quicksort on 50 numbers</td>
</tr>
<tr>
<td>q5-meta</td>
<td>487</td>
<td>397</td>
<td>1.23</td>
<td>Prolog meta interpreter running q5</td>
</tr>
<tr>
<td>queens8</td>
<td>285</td>
<td>304</td>
<td>0.97</td>
<td>8-queens problem</td>
</tr>
<tr>
<td>reducer</td>
<td>2017</td>
<td>2020</td>
<td>1.00</td>
<td>a graph reducer for T-combinators</td>
</tr>
<tr>
<td>sdda</td>
<td>1663</td>
<td>1636</td>
<td>1.02</td>
<td>static data dependency analysis</td>
</tr>
<tr>
<td>talk</td>
<td>69</td>
<td>77</td>
<td>0.90</td>
<td>solves a recursively defined function</td>
</tr>
<tr>
<td>con1</td>
<td>52</td>
<td>46</td>
<td>1.13</td>
<td>concatenation of 3- and 2-element lists</td>
</tr>
<tr>
<td>con6</td>
<td>55</td>
<td>48</td>
<td>1.15</td>
<td>pairwise partition of a 5-element list</td>
</tr>
<tr>
<td>fibo</td>
<td>71</td>
<td>69</td>
<td>1.03</td>
<td>compute 5th fibonacci number</td>
</tr>
</tbody>
</table>

4.1 Static Code Size

Table 1 shows the descriptions and the static code sizes (in number of lines) for the same benchmark compiled using different options for execution under NuSim (NS) and VPsim (VP). The three smallest benchmarks (con1, con6, and fibo) are listed separately at the bottom. The ratios NS/VP show that static NS code and VP code are for the most part well within 10% of one another. The ones that show big variances are due to the lack of
cdr-coding in NuSim, which requires two instructions to build an element (car and cdr) of a list. For example, nrev1 builds a list of 30 elements before reversing it and qs4 builds a list of 50 elements before quick-sorting it.

4.2 Cycle Count (Simulated Time)

Columns VP cycles and NS/VP cycles of Table 2 show the cycle count of VPsim and the ratio of NuSim/VPsim cycles, respectively. The hit ratio column shows results for NuSim configured to a 4-way associative, 64K byte cache with a block size of 16 bytes.

From these columns, we observe that:

- Simulated time of NuSim is quite comparable to VPsim (column NS/VP cycles value is approximately 1) for the large benchmarks (compiler, bintree, compiler.plml, newchat, queens8, reducer, and tak).

- NuSim cycle count is worse than VPsim in the small benchmarks due to low hit ratio (cache cold start). For example, con1, con6, and fibo have the lowest hit ratios among the benchmarks measuring at 88.7%, 95.7%, and 95.6% respectively.

- Non-cdr coded lists also contributes a little to the degradation in performance for a small benchmark such as nrev1 which has a decent hit ratio of 95.3%.

4.3 Simulation Overhead

Although the time that the simulators require to run is largely independent of the correctness of the results, it is interesting to compare simulation overhead of the two simulators because they simulate at two different levels and follow different simulation methodologies.

The following explanations refer to Table 2:

- Column VP systime provides the system simulation time (the time taken to run the simulator on the host in seconds), and column NS/VP systime provides the NuSim to VPsim ratio. These numbers are obtained from running simulations on a SUN 3/60 with 16MB of memory. These values give a feel for the response time of the simulators, ranging from .5 sec to 5920 secs (or 1.64 hours).

- The overhead columns are provided as the ratio of cycle count (discussed in section 4.2) to system simulation time, assuming 100ns cycle time for the NuSim processor and the VLSI-PLM chip. For example, a value such as 2000
Table 2: Cycle Count and Simulation Time

<table>
<thead>
<tr>
<th>Benchmark</th>
<th>VP cycles</th>
<th>NS/VP cycles</th>
<th>NS hit ratio</th>
<th>VP systime</th>
<th>NS/VP systime</th>
<th>VP overhd</th>
<th>NS/VP overhd</th>
</tr>
</thead>
<tbody>
<tr>
<td>bintree</td>
<td>9875</td>
<td>1.30</td>
<td>97.8</td>
<td>3.5</td>
<td>1.43</td>
<td>2544</td>
<td>1.10</td>
</tr>
<tr>
<td>compiler.bin</td>
<td>2208006</td>
<td>0.99</td>
<td>99.5</td>
<td>529.5</td>
<td>0.67</td>
<td>2398</td>
<td>0.57</td>
</tr>
<tr>
<td>compiler.plm</td>
<td>5997896</td>
<td>0.89</td>
<td>99.6</td>
<td>1426.4</td>
<td>0.75</td>
<td>2378</td>
<td>0.65</td>
</tr>
<tr>
<td>hanoi</td>
<td>78884</td>
<td>1.50</td>
<td>99.9</td>
<td>21.4</td>
<td>1.17</td>
<td>2713</td>
<td>0.78</td>
</tr>
<tr>
<td>mumath</td>
<td>96907</td>
<td>1.26</td>
<td>99.8</td>
<td>26.2</td>
<td>0.92</td>
<td>2704</td>
<td>0.73</td>
</tr>
<tr>
<td>newchat</td>
<td>6911008</td>
<td>1.09</td>
<td>99.9</td>
<td>1315.9</td>
<td>1.01</td>
<td>1904</td>
<td>0.92</td>
</tr>
<tr>
<td>nrev1</td>
<td>21192</td>
<td>1.38</td>
<td>98.3</td>
<td>6.1</td>
<td>1.31</td>
<td>2678</td>
<td>0.95</td>
</tr>
<tr>
<td>palin25</td>
<td>25026</td>
<td>1.08</td>
<td>98.6</td>
<td>7.4</td>
<td>1.08</td>
<td>2957</td>
<td>1.00</td>
</tr>
<tr>
<td>puzzle</td>
<td>39456475</td>
<td>0.67</td>
<td>99.9</td>
<td>5920.2</td>
<td>0.43</td>
<td>1500</td>
<td>0.65</td>
</tr>
<tr>
<td>qs4</td>
<td>43190</td>
<td>0.98</td>
<td>98.9</td>
<td>11.9</td>
<td>0.92</td>
<td>2755</td>
<td>0.94</td>
</tr>
<tr>
<td>qs4.meta</td>
<td>348051</td>
<td>1.17</td>
<td>98.9</td>
<td>113.6</td>
<td>0.65</td>
<td>3264</td>
<td>0.56</td>
</tr>
<tr>
<td>queens8</td>
<td>19759942</td>
<td>1.04</td>
<td>100.0</td>
<td>3354.2</td>
<td>1.18</td>
<td>1697</td>
<td>1.13</td>
</tr>
<tr>
<td>reducer</td>
<td>2543554</td>
<td>1.07</td>
<td>99.5</td>
<td>439.6</td>
<td>1.11</td>
<td>1729</td>
<td>1.04</td>
</tr>
<tr>
<td>sdda</td>
<td>85382</td>
<td>1.14</td>
<td>98.5</td>
<td>26.0</td>
<td>0.93</td>
<td>3279</td>
<td>0.82</td>
</tr>
<tr>
<td>tak</td>
<td>9398259</td>
<td>0.96</td>
<td>99.2</td>
<td>2461.5</td>
<td>0.62</td>
<td>2619</td>
<td>0.65</td>
</tr>
<tr>
<td>con1</td>
<td>256</td>
<td>2.96</td>
<td>86.7</td>
<td>0.5</td>
<td>6.00</td>
<td>19331</td>
<td>2.03</td>
</tr>
<tr>
<td>con6</td>
<td>1307</td>
<td>1.52</td>
<td>95.7</td>
<td>0.7</td>
<td>4.29</td>
<td>5356</td>
<td>2.82</td>
</tr>
<tr>
<td>fibo</td>
<td>2225</td>
<td>1.44</td>
<td>95.6</td>
<td>1.2</td>
<td>2.50</td>
<td>5363</td>
<td>1.73</td>
</tr>
</tbody>
</table>

In these columns means that it took 2000 seconds of the SUN 3/60 time to simulate 1 second of the VLSI-PLM.

The worst numbers in the overhead columns appear in the three smallest benchmarks con1. con6. and fibo. This is due to the initial overhead of starting up the simulators. Also in the three smallest benchmarks. the overhead of NuSim is much higher than VPsim (1.73 to 2.82 times worse). This is because NuSim takes more time to startup. being a multiprocessor simulator and having to assemble the benchmark into assembly code. For the larger benchmarks. the NuSim is more efficient than VPsim. Excluding the three smallest benchmarks. the average overheads of NuSim and VPsim are 2203 and 2555. respectively. Thus NuSim is 16% more efficient.

- Even though NuSim simulates the VLSI-PLM at a slightly higher level than the register-transfer level of VPsim. it is not that much more efficient because VPsim microcode is "flat" while NuSim C-routines are hierarchically structured. The cost of structured code depends on the efficiency of the code generated by the C compiler for subroutine calls and returns.

Simulation of the VLSI-PLM on a SUN 3/60 is more than 2000 times slower than actual execution on a VLSI-PLM because of the following reasons:
• Data and control transfers are processed sequentially. In a real machine, it would be done in parallel. The VLSI-PLM has a two stage pipeline, with the data unit and microsequencer executing in parallel. The VLSI-PLM data unit is also capable of doing 8 simultaneous transfers in one cycle.

• The host processor is less powerful than the target processor for symbolic computation and the host memory access time is slower than the target memory access time. The SUN 3/60 that we use has a 20MHz MC68020 and 16MB of main memory (300ns access time). There is no cache. The VLSI-PLM is a complex processor with tag processing capability.

• The code generated by the C compiler affects the execution time of the host. For example, inefficient subroutine calls and returns penalize the hierarchical structure of NuSim C code.

• The presence of extensive instrumentation code in the simulators for extracting performance results slows down execution on the host.

• The operating system characteristic of the host can greatly affect performance. The SUN 3/60 runs 4.3 BSD Unix and virtual memory. The CPU accesses a shared file server connected via Ethernet, and thus page faults are very expensive.

The factors above blend together in the real uniprocessor system and it is difficult to measure them separately. This is the reason why a simulator is needed for experimentation with individual system parameters. For simulating a multiprocessor configuration, the event driven approach of NuSim may be accelerated by use of a faster uniprocessor, or a multiprocessor host, as demonstrated by [Wil87, Jon86]. For the greatest efficiency in simulation, a direct execution approach such as the one proposed by Fujimoto [FC88] may be used, where the benchmark is compiled into code directly executable by the host. Instrumentation counters are inserted by the compiler into the code to measure performance for the target machine.

4.4 Operation Count

In Prolog, the metric Logical Inferences Per Second in units of 1000 (KLIPS) is often used for measuring the performance of Prolog engines. A logical inference can be defined as a Prolog function call, which include VLSI-PLM instructions calls, executes, and escapes for Prolog builtins. This metric is quite inaccurate since the logical inference can not be measured exactly. The amount of work done by a Prolog function call depends on the number and type of arguments in Prolog. For parallel execution, the KLIPS measurement has even less significance. Multiprocessors may do more work but do not necessarily achieve the final result any faster, if the additional computations do not contribute directly to the result.
Table 3: Logical Inference Count

<table>
<thead>
<tr>
<th>Benchmark</th>
<th>NS calls</th>
<th>NS escapes</th>
<th>NS KLIPS</th>
<th>VP calls</th>
<th>VP escapes</th>
<th>VP KLIPS</th>
<th>NS/VP</th>
</tr>
</thead>
<tbody>
<tr>
<td>bintree</td>
<td>77</td>
<td>151</td>
<td>177</td>
<td>128</td>
<td>101</td>
<td>232</td>
<td>0.76</td>
</tr>
<tr>
<td>compiler_bintree</td>
<td>15113</td>
<td>7186</td>
<td>102</td>
<td>20886</td>
<td>2539</td>
<td>106</td>
<td>0.96</td>
</tr>
<tr>
<td>compiler_plml</td>
<td>42597</td>
<td>22318</td>
<td>122</td>
<td>67060</td>
<td>3992</td>
<td>118</td>
<td>1.03</td>
</tr>
<tr>
<td>hanoi</td>
<td>767</td>
<td>765</td>
<td>129</td>
<td>1022</td>
<td>511</td>
<td>194</td>
<td>0.67</td>
</tr>
<tr>
<td>mumath</td>
<td>1211</td>
<td>82</td>
<td>106</td>
<td>1221</td>
<td>73</td>
<td>134</td>
<td>0.79</td>
</tr>
<tr>
<td>newchat</td>
<td>66905</td>
<td>60</td>
<td>89</td>
<td>66911</td>
<td>55</td>
<td>97</td>
<td>0.92</td>
</tr>
<tr>
<td>mrev1</td>
<td>497</td>
<td>2</td>
<td>171</td>
<td>497</td>
<td>3</td>
<td>236</td>
<td>0.72</td>
</tr>
<tr>
<td>palin25</td>
<td>228</td>
<td>97</td>
<td>121</td>
<td>323</td>
<td>3</td>
<td>130</td>
<td>0.93</td>
</tr>
<tr>
<td>puzzle</td>
<td>19796</td>
<td>6016</td>
<td>10</td>
<td>21800</td>
<td>4015</td>
<td>7</td>
<td>1.50</td>
</tr>
<tr>
<td>qs4</td>
<td>361</td>
<td>231</td>
<td>144</td>
<td>610</td>
<td>3</td>
<td>142</td>
<td>1.02</td>
</tr>
<tr>
<td>qs4_meta</td>
<td>2694</td>
<td>720</td>
<td>84</td>
<td>3795</td>
<td>3</td>
<td>109</td>
<td>0.77</td>
</tr>
<tr>
<td>queens8</td>
<td>76457</td>
<td>151736</td>
<td>111</td>
<td>228009</td>
<td>155</td>
<td>115</td>
<td>0.96</td>
</tr>
<tr>
<td>reducer</td>
<td>15091</td>
<td>6305</td>
<td>79</td>
<td>18815</td>
<td>2491</td>
<td>84</td>
<td>0.94</td>
</tr>
<tr>
<td>sdda</td>
<td>552</td>
<td>408</td>
<td>99</td>
<td>715</td>
<td>249</td>
<td>113</td>
<td>0.87</td>
</tr>
<tr>
<td>tak</td>
<td>63609</td>
<td>111317</td>
<td>195</td>
<td>174924</td>
<td>3</td>
<td>186</td>
<td>1.05</td>
</tr>
<tr>
<td>con1</td>
<td>4</td>
<td>2</td>
<td>79</td>
<td>4</td>
<td>3</td>
<td>273</td>
<td>0.29</td>
</tr>
<tr>
<td>con6</td>
<td>6</td>
<td>30</td>
<td>181</td>
<td>6</td>
<td>31</td>
<td>263</td>
<td>0.64</td>
</tr>
<tr>
<td>fibo</td>
<td>15</td>
<td>23</td>
<td>118</td>
<td>36</td>
<td>3</td>
<td>175</td>
<td>0.65</td>
</tr>
</tbody>
</table>

Table 3 shows the number of normal calls/executes and Prolog builtin invocations (or escapes). Since VPsim does calls to library routines for some of the builtns, it has a much higher calls count and fewer escape count than NuSim. In order for KLIPS to be a useful measure, the condition $N_S_{call} + N_S_{escape} \approx V_P_{call} + V_P_{escape}$ should hold true. The following results show that this condition does not hold, due to the implementation variations of NuSim and VPsim (described in section 3.3).

Each of the KLIPS columns is calculated by

$$\frac{calls + escapes}{cycles} \times 10000$$

where cycles is obtained from Table 2. The unit for calls and escape is the logical inference. The constant factor of 10000 comes from the KLIPS unit conversion:

$$1 \text{ KLIP} = \frac{10^6 \text{ nsec}}{1 \text{ sec}} = \frac{1 \text{ cycle}}{100 \text{ nsec}} = \frac{1 \text{ K}}{}$$

The NS KLIPS and VP KLIPS columns differ widely, showing once again the problem with this metric. For comparison purpose, the timing information in table 2 is much more useful than this metric.
4.5 Memory Accesses

Table 4 compares the number of memory accesses made in running the simulations on NuSim and the VPsim. VP total refs gives the total count of memory references to give a sense of the order of magnitude of memory accesses, which range from about 100 to over 12 million. The next 4 columns show the ratios of accesses between NuSim and VPsim for total references, instruction fetches, reads, and writes.

<table>
<thead>
<tr>
<th>Benchmark</th>
<th>VP</th>
<th>NuSim/VP</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>total refs</td>
<td>refs</td>
</tr>
<tr>
<td>butree</td>
<td>5601</td>
<td>1.19</td>
</tr>
<tr>
<td>compiler_bitree</td>
<td>1259778</td>
<td>1.07</td>
</tr>
<tr>
<td>compiler_plim1</td>
<td>3311904</td>
<td>0.96</td>
</tr>
<tr>
<td>hanoi</td>
<td>51811</td>
<td>1.35</td>
</tr>
<tr>
<td>munch</td>
<td>53052</td>
<td>1.26</td>
</tr>
<tr>
<td>newchat</td>
<td>3655155</td>
<td>1.16</td>
</tr>
<tr>
<td>nrevl</td>
<td>8473</td>
<td>1.51</td>
</tr>
<tr>
<td>palin25</td>
<td>12758</td>
<td>1.10</td>
</tr>
<tr>
<td>puzzle</td>
<td>11600446</td>
<td>0.81</td>
</tr>
<tr>
<td>qs</td>
<td>24302</td>
<td>0.93</td>
</tr>
<tr>
<td>qs4_meta</td>
<td>197469</td>
<td>1.13</td>
</tr>
<tr>
<td>queens</td>
<td>12354397</td>
<td>1.09</td>
</tr>
<tr>
<td>reducer</td>
<td>1367055</td>
<td>1.14</td>
</tr>
<tr>
<td>sdds</td>
<td>45313</td>
<td>1.13</td>
</tr>
<tr>
<td>tak</td>
<td>5979238</td>
<td>0.63</td>
</tr>
<tr>
<td>con1</td>
<td>94</td>
<td>2.11</td>
</tr>
<tr>
<td>con2</td>
<td>499</td>
<td>1.55</td>
</tr>
<tr>
<td>fbo</td>
<td>1207</td>
<td>1.10</td>
</tr>
</tbody>
</table>

We observe the following:

- NuSim fetches instructions on demand, while VPsim does prefetching. NuSim instructions are encoded in word streams, with the opcode and each operand taking up one 32-bit word. VPsim has the code stored in string tables, but the microcode generates prefetch signals to simulate an encoding of 8-bit opcode and 32-bit arguments.

- The total reference ratios are for most benchmarks are about 1. The big variations are for con1 (2.11), con2 (1.55), and nrev1 (1.51). The variations are perfect examples of worst case performance without cdr-coding (in NuSim), which would require more instruction fetches, reads and writes. For the larger benchmarks, cdr-coding makes little difference.

- The ifetch ratios show that the word-encoding of NuSim require more fetches, as expected. However, for tak, NuSim fetches much less (ifetch ratio of 0.66)
because many subtractions are done and NuSim use the builtin instruction \( \text{ts}/2 \) while \( \text{VPsim} \) does a call to the library routine \( \text{sub}/3 \) which require a longer sequence of simpler instructions.

5 Discussion

Simulation is an important part of system integration. In this paper, we have shown a methodology for validating the simulator of a multiprocessor system. We applied this scheme to validate the processor and the memory module of a multiprocessor simulator (NuSim) by comparing it with a previously validated uniprocessor simulator (VPsim). Benchmarks of various sizes were executed sequentially on both simulators, and different performance measurements were evaluated and compared against one another.

Because the simulation result is a composite result of many factors, we chose a number of measurements for comparison to obtain different perspectives on performance and to understand the reasons of the variations. The chosen measurements were: code size, cycle count, simulation overhead, operation count, and memory access counts. The different measurements indicate that the variations are significant only for the small benchmarks, where startup time and slight model differences are a big percentage of total execution time. For large programs, NuSim is within 10% of the VLSI-PLM timing. Perhaps more importantly, all variations can be accounted for. We can thus conclude that NuSim is representative of a VLSI-PLM in a multiprocessor system. With NuSim, we can continue our study of implementable multiprocessor systems for parallel execution of numeric and symbolic programs, using logic programming. We also believe that the chosen measurements can be used in validating other simulation systems.

6 Acknowledgement

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References


