

Approaches to Parallel Quantifier Elimination

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Abstract

Special-purpose quantifier elimination procedures for problems of low degree using virtual substitution of test terms have recently turned out to be applicable to a variety of non-trivial non-academic problems. We study parallel algorithms based on these methods for several parallelization environments: a Cray YMP4/T3D, a workstation cluster, and a multi-processor Sparc. Our implementations show remarkable though sublinear speed-ups in all these environments.

1 Introduction

The basic motivation of real quantifier elimination is to “eliminate” unwanted variables representing real quantities from an algebraic description of some situation. As an example, consider conditions for the solvability of the equation $ax^2 + b = 0$ where a and b are real parameters: This equation has a real solution if and only if $ab < 0$ or $b = 0$. Formally, we have eliminated the quantifier “ $\exists x$ ” from $\exists x(ax^2 + b = 0)$ yielding the equivalent $ab < 0 \vee b = 0$. The latter provides a better idea of the range of a and b described by the two formulas.

For a long time the use of quantifier elimination in application problems outside pure mathematics has been fairly limited due to the poor efficiency of the implemented methods. Recently, implementations of quantifier elimination have been able to solve problems of interesting size in science, engineering, and also in economics, namely in operations research, cf. [16] and the references there. Quantifier elimination by *virtual substitution* of test terms [39, 30, 42, 43] plays a crucial role for this success, in particular for problems involving numerous parameters.

The purpose of our work is to study parallelization and distribution of the virtual substitution method. Here, *distribution* refers to parallelization on distributed-memory hosts or network clusters of several hosts. We have worked with the following environments, of which the first two are distributed ones:

- PVM and RR on a Cray YMP4/T3D supercomputer using a REDUCE implementation of a special case of the

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method for non-parametric linear optimization.

- PVM based DTS on a workstation cluster.
- Multithreading on a multi-processor Sun Sparc-10 running Solaris 2.

For the latter two cases, we have realized a SACLIB/PARSAC-2 based C implementation “QEDTS.”

In all these environments we obtain remarkable speed-ups. Since our parallelization proceeds in such a way that we perform essentially the same computations as in the sequential case, the speed-ups are limited by the number of processors. For decision problems, there are straightforward extensions of our approaches which will obviously give rise to superlinear speed-ups.

2 History and Related Work

The first real quantifier elimination procedure was found by Tarski in the 1930's. It remained unpublished until 1948 [38]. Tarski's procedure was very inefficient, more precisely it was not elementary recursive.

In 1975 Collins introduced a new method called *cylindrical algebraic decomposition* (CAD) [7], which is worst-case doubly exponential in the number of variables. A full implementation of CAD within the computer algebra system SAC-2 [8] has been finished by Arnon in 1981. In the past 20 years of research, CAD has gone through numerous improvements [31, 21, 22], resulting in *partial* CAD [9] implemented in Hong's QEPCAD program based on SACLIB [6], a C version of SAC-2.

Around 1988 it has been shown that real quantifier elimination is inherently hard for some problem classes [10, 39]. Thus the attention turned to special procedures for restricted problem classes, where the elimination procedures can be tuned to the structure of the problem. The focus was on considering formulas in which the occurrence of quantified variables is restricted to low degrees [24, 25, 19]. This was initiated by a theoretical paper of Weispfenning in 1988 [39], who introduced the method of *virtual disjunctive term substitution*, which has been continuously extended and improved [30, 42, 43, 15, 14].

The worst-case complexity of this method is doubly exponential only in the number of the *quantifier blocks* of the input formula, but singly exponential in the number of *quantified* variables. This makes the method attractive for problems containing many parameters. It is implemented in the REDUCE [20] package REDLOG [12, 13]. In addition,

there is an implementation in the computer algebra system Risa/Asir [37]. For the purpose of this paper, the authors have reimplemented the method once more in C based on SACLIB/PARSAC-2 [6, 27, 28, 29, 18]. All these implementations are currently restricted to input obeying certain degree restrictions with respect to the quantified variables. In principle, the method can be extended to arbitrary degrees [43].

In 1993 Weispfenning introduced a new complete elimination procedure based on comprehensive Gröbner bases in combination with multivariate real root counting, cf. [41] and [40, 4, 33]. It has been implemented within the computer algebra system MAS [11].

In recent years there have been impressive theoretical results on asymptotically fast real elimination algorithms [34, 3]. Since there is no implementation available, the question whether these methods are of practical relevance remains unanswered.

The first work in parallelizing quantifier elimination has been done by Saunders et al. in 1989 [35]. They have parallelized the original Collins CAD algorithm based on SAC-2 on an 8-processor Sequent Symmetry shared-memory machine. They report a speed-up of about factor 3 using 7 processors.

In 1993 Hong has parallelized his SACLIB based QEPCAD on a workstation network based on an apparently unpublished communication protocol [23]. Using a master-slave model on 6 hosts, he obtains speed-ups ranging from factor 4 up to superlinear.

Hong has also developed a parallel extension of SACLIB named PACLIB [26], which runs on shared-memory machines. There are numerous symbolic algorithms implemented based on PACLIB [36], not including quantifier elimination, however.

Another independent parallel extension of SACLIB is PARSAC-2 [27, 28, 18], originally developed by Küchlin. PARSAC-2 is a parallel Computer Algebra library targeted for execution on multi-processor workstations. Up to now, the applications of PARSAC-2 have been limited to the area of integer and polynomial arithmetic and Gröbner basis computation [29, 1, 2].

3 Components of our Parallelization Environments

3.1 PVM

PVM [17] stands for Parallel Virtual Machine. It is a software package that allows a network of parallel and serial computers to appear as single concurrent distributed-memory machine. Under PVM, C or Fortran-77 applications can be parallelized using common message-passing constructs. It is available for a variety of architectures including the Cray YMP4/T3D and Sun Workstations used for our studies.

3.2 RR

RR [32] provides an extension of the Portable Standard Lisp (PSL) system underlying REDUCE. It supports both the Cray YMP4/T3D and workstation clusters. Communication and control are implemented on top of the PVM manager. RR supports the fork/join paradigm by means of remote procedure calls and remote-wait/remote-receive primitives.

3.3 The S-thread System

PARSAC-2 contains the S-threads parallel symbolic programming environment [27]. S-threads is *middle-ware* between the operating system below and the application software above. Most importantly, an S-thread extends an operating system thread by a control block that supports memory management. Furthermore, the memory allocation mechanism as well as the mark-sweep garbage collection of SACLIB are modified in such a way that concurrent allocation is enabled. Practically all sequential procedures of the SACLIB library can now be executed concurrently, by forking them on a separate S-thread.

S-threads support *virtual parallel programming*: we parallelize an algorithm according to its inherent logical parallel structure. The resulting logical threads of control are mapped at run-time to real concurrency provided by the hardware and the operating system. Thus the same executable can run on 1, 2, or more processors.

The implementation of S-threads is based on operating system threads. For the current work, we used the S-thread system on Sparc workstations under Solaris 2, which in turn relies on Solaris/POSIX threads [18].

The following parallelization paradigms of the S-thread system are of importance for our quantifier elimination algorithm:

Work Parallelism in its pure form (\forall -Parallelism) is the concept supported most directly by the fork/join paradigm of multithreading. A given amount of work is divided into multiple threads of control and forked onto a number of threads. Each thread is finally collected (joined) again by the parent, and its result is retrieved.

In our quantifier elimination algorithm, we employ work parallelism by applying quantifier elimination to subformulas. The final result is then obtained as the disjunction over the quantifier-free equivalents obtained for the subformulas.

Search Parallelism in its pure form (\exists -Parallelism) reflects a parallel search for some item. A number of search threads are forked, and the first child to find the item synchronizes with the parent to deliver the result. The parent subsequently terminates the other children, not being interested in their results.

In our quantifier elimination algorithm, in particular with decision problems, i.e. parameter-free elimination problems, we could use a form of search parallelism: As soon as one of the forked subproblems yields “true,” all other subproblems become irrelevant and these computations can be aborted.

The synchronization needed by search parallelism is now supported in S-threads by the concept of *thread groups*. A thread group is a collection of threads, similar to a process group in the traditional Unix world. We designed our thread groups to support both work and search parallelism in a uniform way.

After a thread group has been opened, threads can be forked into the group at any time. The parent can wait for the next thread of the group, or it can send a signal to the entire group, for example to terminate or to suspend all threads (and their children, of course).

On shared-memory machines with fast context switching, the main use of a thread group is in supporting search parallelism. Then, all threads of the group execute as concurrent operating system threads. On a uniprocessor machine, execution of these threads will be interleaved to maintain logical concurrency.

Of course, all these features are recursively inherited to forked threads. This is of crucial importance for the parallelization of our quantifier method as we use a recursive approach in our parallelization.

3.4 DTS

The fork/join concept of shared-memory parallel programming can be carried over to the distributed-memory network, provided all global access to data-structures can be confined to parameters that can be packed with fork/join. An extension of the S-threads called DTS (Distributed Thread System [5]) is based on PVM which transports fork/join calls across the net. Thus, we can use the same parallelization paradigm (and essentially the same code), regardless whether we compute on a shared-memory workstation or on a network of single-processor or even multi-processor workstations. In addition, as DTS relies on S-threads, we do not take care about the number of forked tasks. That is, we usually overload the processors and thus overlap the network communication by actual computation tasks.

4 Quantifier Elimination by Virtual Substitution

The applicability of the virtual substitution method in the form described here is restricted to formulas in which the quantified variables occur at most quadratically. Moreover, quantifiers are eliminated one by one, and the elimination of one quantifier can increase the degree of other quantified variables.

For eliminating the quantifiers from an input formula

$$\varphi(u_1, \dots, u_m) \equiv \mathbf{Q}_1 x_1 \dots \mathbf{Q}_n x_n \psi(u_1, \dots, u_m, x_1, \dots, x_n),$$

where $\mathbf{Q}_i \in \{\exists, \forall\}$, the elimination starts with the innermost quantifier regarding the other quantified variables within ψ as extra parameters. Universal quantifiers are handled by means of the equivalence $\forall x \psi \longleftrightarrow \neg \exists x \neg \psi$. We may thus restrict our attention to a formula of the form

$$\varphi^*(u_1, \dots, u_k) \equiv \exists x \psi^*(u_1, \dots, u_k, x),$$

where the u_{m+1}, \dots, u_k are actually x_i quantified from further outside.

We fix real values a_1, \dots, a_k for the parameters u_1, \dots, u_k . Then all polynomials occurring in ψ^* become linear or quadratic univariate polynomials in x with real coefficients. So the set

$$M = \{b \in \mathbb{R} \mid \psi^*(a_1, \dots, a_k, b)\}$$

of all real values b of x satisfying ψ^* is a finite union of closed, open, and half-open intervals on the real line. The endpoints of these intervals are among $\pm\infty$ together with the real zeros of the linear and quadratic polynomials occurring in ψ^* . Candidate terms $\alpha_1, \dots, \alpha_r$ for the zeros can be computed uniformly in u_1, \dots, u_k by the solution formulas for linear and quadratic equations.

If all inequalities in ψ^* are weak, then all the intervals constituting M will, into each direction, be either unbounded or closed. In the latter case, such an interval will contain its real endpoint. Thus M is non-empty if and only if the substitution of $\pm\infty$ or of one of the candidate solutions α_j for x satisfies ψ^* . The substitution of $\pm\infty$ into a

polynomial equation or inequality is evaluated in the obvious sense. The substitution of expressions in u_1, \dots, u_k of the form $(a + b\sqrt{c})/d$ among the α_j can be rewritten in such a way that all denominators involving the u_i and all square-root expressions are removed from the result [43]. By disjunctively substituting all candidates into ψ^* we obtain a quantifier-free formula equivalent to $\exists x \psi^*$.

If ψ^* contains also strict inequalities, we need to add to our candidates for points in M expressions of the form $\alpha \pm \varepsilon$, where α is candidate solution for some left-hand side polynomial occurring in a strict inequality. The symbol ε stands for a positive infinitesimal number. Again the substitution of these expressions into a polynomial equation or inequality can be rewritten in such a form that there occur neither denominators involving any of the u_i , nor any square root expressions, nor the symbol ε in the result [43]. Again this yields a quantifier-free formula equivalent to $\exists x \psi^*$. For practical applications this method, of course, has to be refined by a careful selection of a smaller number of candidate solutions and by a combination with powerful simplification techniques for quantifier-free formulas, cf. [14] for details.

Recall that the well-known solution formula for quadratic equations

$$ax^2 + bx + c = 0$$

requires $a \neq 0$. In our situation, a is a term in u_1, \dots, u_k . So $a \neq 0$ can in general not be decided uniformly, but depends on the interpretation of the u_i . Thus a quadratic polynomial $ax^2 + bx + c$ does not only deliver two square-root expressions α_1 and α_2 as candidate solutions, but also $\alpha_3 = -c/b$, which in turn requires $b \neq 0$. Let t_1, t_2 , and t_3 be the candidate points for M obtained from α_1, α_2 , and α_3 , respectively, by possibly adding or subtracting ε . With the substitution of all the t_i into ψ^* , it is necessary to add the conditions on the non-vanishing of a and b . Formally, we obtain

$$(a \neq 0 \wedge \Delta \geq 0 \wedge (\psi^*[x/t_1] \vee \psi^*[x/t_2])) \vee (a = 0 \wedge b \neq 0 \wedge \psi^*[x/t_3]),$$

where Δ denotes the discriminant of the equation $ax^2 + bx + c = 0$. If, however, a is a rational constant, then the case distinction is superfluous. In particular, if a is non-zero, we drop the second case.

We can obtain dramatic improvements of the general procedure sketched above by reducing the number of test candidates for M depending on the structure of the formula ψ^* [30, 43]. One simple instance for such an improvement is the following natural extension of *Gauss elimination*: Suppose ψ^* is of the form

$$bx + c = 0 \wedge \psi^{**},$$

where at least one of the coefficient terms b, c is a rational non-zero constant. Then we know that, under any interpretation of the u_i , the equation is *non-trivial*, i.e. different from $0 = 0$. Hence the only test candidate required is $-c/b$ substituted, of course, with the condition $b \neq 0$. No additional test candidates that arise from equations or inequalities in the remainder ψ^{**} of ψ^* need be considered. This idea can easily be extended to a quadratic equation instead of a linear one, taking into account again the discriminant.

The notion of *virtual* substitution refers to both adding conditions and resolving non-standard subterms such as surds or infinitesimals. This method is doubly exponential in the number of quantifier changes. For pure “ \exists ” or

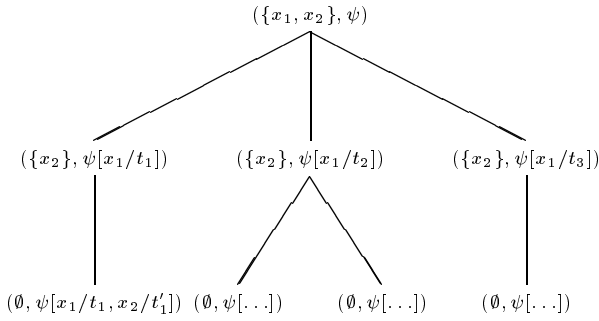


Figure 1: An elimination tree

“ \forall ” blocks, it is singly exponential in the number of quantified variables. In theory, parameters play a minor role for the complexity. They turn in fact out to be very cheap in practice, too.

5 The Elimination Tree

In the sequel, we restrict our attention to the elimination of pure prenex “ \exists ” blocks of quantifiers.

Suppose we have eliminated an existential quantifier. Then we have in general obtained a disjunction $\psi'_1 \vee \dots \vee \psi'_r$. If the next quantifier to be eliminated is also an existential one, then we make use of the equivalence

$$\exists x_{n-1}(\psi'_1 \vee \dots \vee \psi'_r) \longleftrightarrow \exists x_{n-1}(\psi'_1) \vee \dots \vee \exists x_{n-1}(\psi'_r)$$

and eliminate all $\exists x_{n-1}(\psi'_j)$ independently. As a consequence, no candidate solutions obtained from, say, ψ'_1 are substituted into the other ψ'_j . This decreases the complexity class of our procedure for single quantifier blocks from doubly exponential to singly exponential in the number of quantifiers [39].

Proceeding this way, the elimination procedure for

$$\exists x_1 \dots \exists x_n \psi_0(u_1, \dots, u_m, x_1, \dots, x_n)$$

can be considered as the computation of an *elimination tree* of the following form:

- Each node consists of a quantifier-free formula ψ plus a list V of variables. The corresponding partial elimination problem is $\exists V \psi$.
- The root is $(\psi_0, \{x_1, \dots, x_n\})$.
- For each node $(\psi, \{x_i, \dots, x_n\})$, the children are determined as

$$\begin{aligned} &(\psi[x_i/t_1], \{x_{i+1}, \dots, x_n\}) \\ &\quad \vdots \\ &(\psi[x_i/t_k], \{x_{i+1}, \dots, x_n\}), \end{aligned}$$

where $\{t_1, \dots, t_k\}$ is a set of candidate points with respect to x_i for ψ .

Figure 1 shows an example.

Elimination trees have the following properties: On each level all variable lists are identical. The number of variables in these lists is decreased by one with each new level. All

```

W := {(\psi_0, \{x_1, \dots, x_n\})}
R := \emptyset
while W \neq \emptyset do
  select (\psi, V) from W
  W := W \setminus \{(\psi, V)\}
  if V = \emptyset then
    R := R \cup \{(\psi, V)\}
  else
    determine the children C of (\psi, V)
    W := W \cup C
fi
od

```

Figure 2: The sequential elimination algorithm

leaves of the tree contain the empty list. For each level with nodes (ψ'_1, V') , \dots , (ψ'_k, V') we have

$$\left(\bigvee_{i=1}^k \exists V' \psi'_i \right) \longleftrightarrow \exists x_1 \dots \exists x_n \psi_0.$$

This holds in particular for the last level for which $V' = \emptyset$. Hence the left hand side of the above equivalence is then quantifier-free.

6 The Sequential Algorithm

Figure 2 shows a sequential algorithm for constructing the elimination tree. It maintains a set W of working nodes, and another set R of result nodes, i.e., leaves. The set W is implemented either as a queue or as a stack, with the elimination tree being constructed breadth-first or depth-first, respectively. Both approaches have certain advantages: Using a queue allows us to detect identical sibling nodes which are known to come into existence systematically [44]. Using a stack saves space during the computation, and whenever a leaf happens to become “true,” we can immediately abort the computation and prune the remaining tree. As a rule of thumb, it has turned out reasonable to use the depth-first approach only for decision problems and parameter-free optimization problems.

7 A Master-Slave Approach on the Cray

The first approach to distributing the virtual substitution method has been done by the first and the third author in cooperation with W. Neun at the Konrad Zuse Center in Berlin. This implementation is restricted to conjunctions of parameter-free constraints involving only “ \geq ,” “ \leq ,” and “ $=$.” It is part of the REDUCE package REDLOG. It runs on a Cray YMP4/T3D using PVM plus a special version of REDUCE based on PSL extended by RR. The idea is to have a master node generate by breadth-first elimination a *problem base*, which is then distributed to several slaves in order to be processed completely by depth-first elimination. A corresponding algorithm is given in Figure 3. We denote by P the list of available processor nodes, and by $H \subseteq P$ the list of busy processors. The initial problem base size $3|P|$ has been found by practical elimination experiments.

This implementation has been applied to three benchmark problems from linear programming: Using 8 processors, i.e. 1 master plus 7 slaves, with 64 MB of memory each, a speed-up factor of about 3 compared to sequential

```

W := {(ψ0, {x1, ..., xn})}
R := ∅
run breadth-first elimination until |W| = 3|P|
while W ≠ ∅ or H ≠ ∅ do
  while W ≠ ∅ and H ≠ P do
    select h from P \ H
    H := H ∪ {h}
    select (ψ, V) from W
    W := W \ {(ψ, V)}
    start depth-first elimination of (ψ, V) on h
  od
  if H ≠ ∅ then
    wait until H' ⊆ H have finished
    H := H \ H'
    for each h in H' do
      receive the result C*
      R := R ∪ C*
    od
  fi
od
return R

```

Figure 3: A parallel master-slave elimination algorithm

Problem	# Var.	# Constr.	Seq.	Par.	Factor
afiro	32	59	333 s	95 s	3.5
sc50a	48	97	>900 s	272 s	>3.3
sc50b	48	96	70 s	25 s	2.8

Table 1: Timings (in seconds) obtained on the Cray

depth-first elimination was obtained, cf. Table 1. The input problems there are taken from the Konrad Zuse Center’s NETLIB online library of benchmark examples. Our speed-ups are very similar to those obtained by Saunders et. al with their shared-memory parallelization of CAD [35].

8 Divide-and-Conquer on a Workstation Cluster using DTS

The S-thread system and DTS allow us to fork essentially arbitrary many threads over the net. Therefore, we can apply the concept of divide-and-conquer, where there is no a priori bound for the number of concurrent tasks. Divide-and-conquer algorithms distribute the problem over the computational resources in the fastest possible manner. When the elimination is distributed over several networked hosts, however, any fork of a function causes its arguments to be sent over the net. Therefore we have to take care about the communication costs. This means that whenever a host processes some forked function, all the data passed to this function should actually be processed there at least partially. The algorithm “first_process” of Figure 4 has been carefully designed to meet this issue.

Compare, in contrast, the algorithm “first_divide” of Figure 5, where half of the nodes contained in the argument W are simply passed to yet another host.

Problem	Seq.	1 Host	2 Hosts	3 Hosts
rp-2	198.05 s	320.0 s	211.7 s	175.6 s

Table 2: Timings (in seconds) with “first_process” on a workstation cluster

```

procedure first_process(W)
  L := ∅
  for each (ψ, V) in W do
    determine the children C of (ψ, V)
    L := L ∪ C
  od
  if the V in L are empty then
    return L
  else
    split L into L1 and L2 of equal lengths
    h := fork(first_process, L1)
    L'2 := first_process(L2)
    L'1 := join(h)
    return L'1 ∪ L'2
  fi
end

```

Figure 4: The “first_process” divide-and-conquer algorithm

Problem	Seq.	1 Proc.	2 Proc.	3 Proc.	4 Proc.
period-9	50.4 s	53.4 s	29.0 s	21.4 s	17.4 s
schedule	93.2 s	109.0 s	60.9 s	44.6 s	36.1 s
rp-2	266.1 s	289.9 s	168.2 s	156.4 s	154.2 s

Table 3: Timings (in seconds) with “first_process” on a 4-processor Sparc

Our first set of distributed timings is collected in Table 2. It has been obtained with three 140 MHz Sun Ultra Sparc-1 connected by a conventional 10 Mbit Ethernet. All hosts are running Solaris 2. We observe an initial overhead of about 60% caused by the parallelization environment, i.e. pvm and DTS, and the fact that the distributed version on one host does not construct the tree in a breadth-first manner, and hence we cannot identify as many identical nodes. On more than one host, we reach moderate speed-ups, with three hosts even compared to the sequential version.

We shall turn back to the distributed case once more in the next section.

9 Divide-and-Conquer on a Multi-processor Sparc

Our shared-memory parallel computations based on S-threads have been performed on a multi-processor Sun Sparc-10 with four 90 MHz Ross Hyper-Sparc processors running Solaris 2. Such a Hyper-Sparc processor is roughly equivalent to a 50 MHz Super-Sparc.

Table 3 collects parallelization timings for several input problems using the algorithm “first_process” of Figure 4 already discussed in the previous section.

Here, the system overhead of the concurrent computation using 1 processor compared to the sequential one is much smaller than in the distributed case. In fact, the overhead of the S-thread system can be neglected. The speed-ups obtained with more processors are similar to those in the distributed case. The absolute running times are, of course, much better. Note that the hosts used for the distribution are significantly faster than 1 processor on the parallel Sparc.

In the example “rp2,” we observe almost no additional speed-up by the 4th processor. Traced output indicates that there are not enough threads forked to exploit this additional resource. Therefore, we have devised the algorithm “first_divide” shown in Figure 5. It forks half of the work

```

procedure first_divide( $W$ )
  if  $|W| = 1$  then
    determine the children  $C$  of  $(\psi, V) \in W$ 
    if the  $V$  in  $C$  are empty then
      return  $C$ 
    else
      return first_divide( $C$ )
    fi
  else
    split  $W$  into  $L_1$  and  $L_2$  of equal lengths
     $h := \text{fork}(\text{first\_divide}, L_1)$ 
     $L'_2 := \text{first\_divide}(L_2)$ 
     $L'_1 := \text{join}(h)$ 
    return  $L'_1 \cup L'_2$ 
  fi

```

Figure 5: The “first_divide” divide-and-conquer algorithm

Problem	Seq.	1 Proc.	2 Proc.	3 Proc.	4 Proc.
period-9	50.4 s	53.8 s	29.4 s	21.9 s	17.6 s
schedule	93.2 s	110.6 s	59.2 s	43.1 s	35.8 s
rp-2	266.1 s	297.1 s	170.2 s	130.3 s	111.2 s

Table 4: Timings (in seconds) with “first_divide” on a 4-processor Sparc

at the earliest possible time before doing any computation itself.

Table 4 shows the timings obtained with “first_divide.” It actually yields additional speed-up with 3 and 4 processors. On the other hand we loose performance with 1 and 2 processors caused by the fact that the construction of the elimination tree by “first_process” is much closer to breadth-first than that by “first_divide.”

Finally, we have combined distribution with parallelization by connecting two 4-processor Sparcs. Since the S-thread system and DTS cooperate perfectly, there is no additional code necessary for doing so, i.e., the same executable as in the previous section is running. The timings collected in Table 5, which we obtained by the algorithm “first_process,” suggest that also with a distributed approach, one can make good use of multi-processor shared-memory machines.

10 Conclusions and Future Work

We have demonstrated that remarkable speed-ups are obtained with parallel quantifier elimination. This holds for shared-memory multi-processor workstations as well as for distributed-memory hosts and networks of workstations.

Our parallel algorithms retain the structures of their corresponding sequential variants. Therefore, we can distinguish between the system effects and the algorithmic effects on the speed-ups.

In the case of decision problems, one wishes to determine

	1 Host	2 Hosts
1 Processor	459.5 s	290.7 s
4 Processors	206.6 s	149.8 s

Table 5: Timing (in seconds) obtained for “rp2” by connecting multi-processor Sparcs with “first_process”

as fast as possible whether there is a leaf containing “true.” Search parallelism allows us to abort the computation immediately when such a leaf is found. We expect superlinear speed-ups from a corresponding implementation.

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