# Concurrent Execution of <br> Mutually Exclusive Alternatives 

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## Abstract

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We examine the task of concurrently computing alternative solutions to a problem. We restrict our interest to the case where only one solution is needed: in this case we need some rule for selecting between the solutions. We use "fastest first," where the first successful alternative is selected. For problems where the required execution time is unpredictable this method can show substantial execution time performance increases. These increases are dependent on the mean execution time of the alternatives, the fastest execution time, the overhead involved in concurrent computation, and the overhead of selecting and deleting alternatives. Rather than using the traditional approach of multiple computers cooperating on the solution to a problem, this method achieves a solution competitively.

Among the problems with exploring multiple alternatives in parallel are side-effects and combinatorial explosion in the amount of state which must be preserved. These are solved by process management and an application of "copy-on-write" virtual memory management. The side effects resulting from interprocess communication are handled by a specialized message layer which interacts with process management.

We show how the scheme for parallel execution can be applied to several application areas. The applications are distributed execution of recovery blocks, OR-parallelism in Prolog, and polynomial root-finding.

## Table of Contents

Page

1. Introduction ..... 1
1.1. Thesis Summary ..... 4
2. Theoretical Underpinnings ..... 8
2.1. Extremal Behavior ..... 8
2.2. Performance Analysis ..... 10
2.2.1. Overhead ..... 11
2.2.2. Analytic Description ..... 11
2.2.3. Parallel Speedup ..... 14
2.2.4. Domain-wide performance indices ..... 19
2.3. Conclusions ..... 21
3. Algorithms for Parallel Execution ..... 22
3.1. System Model ..... 22
3.2. Process Management ..... 23
3.2.1. Synchronization ..... 25
3.2.2. Atomicity ..... 26
3.3. Predicates ..... 26
3.3.1. Representation of Predicatcs ..... 27
3.4. Interprocess Communication ..... 32
3.4.1. Messages ..... 32
3.4.2. Multiple Worlds ..... 33
3.5. Discussion ..... 37
4. Implementation, Applications and Experiments ..... 40
4.1. Measurement of Overhead Costs ..... 40
4.2. Copy-on-write ..... 41
4.2.1. Motivation ..... 42
4.2.2. Data Acquisition ..... 42
4.2.3. Data Analysis ..... 44
4.2.4. Relationships ..... 50
4.2.5. Write Fraction for Real Programs ..... 54
4.2.5.1. Franz Lisp ..... 55
4.2.5.2. GNU Emacs ..... 57
4.2.6. Conclusions about copy-on-write ..... 58
4.3. Remote fork() ..... 60
4.3.1. Further process migration ideas ..... 60
4.4. Disk Response Time ..... 62
4.5. Network Response Time ..... 63
4.6. Sibling Elimination ..... 64
4.6.1. Real Time ..... 66
4.6.2. System Time ..... 74
4.6.3. Real Time, 16 Procs only ..... 82
4.6.4. System Time, 16 Procs only ..... 88
4.6.5. Correction for process scheduling ..... 94
4.7. Possible Sources of Error in Measurements ..... 100
4.7.1. UNIX Clock Facility ..... 101
4.7.2. Experimental Apparatus ..... 102
4.8. Discussion ..... 104
4.9. Applications ..... 104
4.9.1. Distributed Execution of Recovery Blocks ..... 104
4.9.2. Polynomial Root-finding ..... 106
4.9.2.1. Example ..... 107
4.9.2.2. Parallel Execution ..... 113
4.9.3. Other Applications of the Technique ..... 117
4.9.3.1. OR-parallelism in Prolog ..... 117
4.9.3.1.1. Tutorial Example ..... 118
4.9.3.1.2. Existing Solutions ..... 120
4.9.3.1.3. Discussion ..... 126
4.9.3.1.4. Measurements of Published Prolog Programs ..... 128
4.9.3.2. Polyalgorithms ..... 130
4.9.3.3. Simulation ..... 131
5. Related Work ..... 132
6. Conclusions ..... 135
7. Directions for Future Work ..... 141
8. References ..... 142
9. Appendix I: do fork.c ..... 153
10. Appendix II: netrand.c ..... 155
11. Appendix III: Further Jenkins-Traub executions ..... 157
11.1. Polynomial \#1 ..... 157
11.2. Polynomial \#2 ..... 158
11.3. Polynomial \#3 ..... 160
11.4. Polynomial \#4 ..... 161
11.5. Polynomial \#5 ..... 163
11.6. Polynomial \#6 ..... 164
11.7. Polynomial \#7 ..... 165
11.8. Polynomial \#8 ..... 166
12. Appendix IV: Source, Prolog Sorts ..... 168
13. Appendix V: Timings, Prolog Sorts, Large Lists ..... 170
14. Appendix VI: Sort Performance, Small Lists ..... 173
15. Appendix VII: Naive Sort Performance ..... 174
16. Appendix VIII: Performance on Small, Ordered Lists ..... 175
17. Appendix IX: Program to estimate memory speeds ..... 176
18. Appendix X: Lower bound affects dispersion ..... 178
19. Appendix XI: do_elim Script ..... 184
20. Appendix XII: do_elim.c ..... 186
21. Appendix XIII: " $C$ '" version of Jenkins-Traub algorithm ..... 194
22. Appendix XIV: cvaryangle.c ..... 208
23. Appendix XV: r2p.c ..... 210
24. Appendix XVI: cmach.c ..... 215
25. Appendix XVII: Extremal exploitation of randomness ..... 220
26. Biography ..... 224

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## 1. Introduction

This thesis examines the problem of concurrently executing alternative solutions to a problem. There are four major questions which scientific research can answer. The first question is "can we do this?;" the second is "if so, how?," the third is "when do we want to?," and the fourth, "where will opportunities exist?."

This thesis answers each of these questions for this problem; a summary of the thesis results can be found at the end of the introduction.

Over time, the ratio of people to computers has decreased ${ }^{1}$. In the mainframe environments of the 1960s, there were often several thousand users per computer. As time advanced, departmental minicomputers which became popular in the 1970s had an order of magnitude fewer users for each computer, even though these computers were roughly comparable in power to the earlier generation. The 1980s have seen the introduction of supermicrocomputers where the number of users is a dozen or less, and workstations and personal computers which may support only a single user. Thus, it is reasonable to expect that in the near future, rich computational environments will offer use of several computers at the same time. These computers are typically connected through a bus, where the communications abstraction is a shared memory, or a network, where the communications abstraction is message-passing. A description of some computer organizations which incorporate multiple processors can be found in Smith [Smith1986a].

A question which has intrigued many researchers is how this increasing supply of computational resources, in the form of multiple computers, can be used to solve bigger problems, to solve problems faster, and to solve problems more reliably. Traditional approaches have been cooperative, in the sense that the multiple computers cooperate in developing the problem solution. Here, we look at a competitive approach, that of pursuing alternative solution methods.

There are many situations where there exist several alternative methods for computing a result, where a result in the most general case is a state change. Our designs show what can be done to execute instances of this problem type, speculatively, in parallel.

We are interested in what performance gains can be achieved. We measure

[^0]performance using the metric of execution time, which is the amount of wall clock time necessary to carry out a computation. Thus, we may increase performance by this measure, while decreasing performance by measures such as throughput, which is a measure of the amount of useful work done per unit time.

This thesis demonstrates that the speedup promised by parallel exploitation of randomness is achievable in practice.

We begin by describing the computations to be analyzed. These are essentially a set of alternative methods for causing a state change to take place, with the additional constraint that at most one alternative state change occurs. These might be denoted a, b, and $c$, and expressed as

SELECT
a
OR
b
OR
c
TCELES
a, b. and c comprise a block. The block's semantics are simple: one of its components is executed.

Use of other language features allows control of the execution, if necessary. For example, if one wanted to execute the alternatives in order $a, b, c$, loops and guards could be applied. e.g..

```
FOR I=1 TO 3
    SELECT
            WHEN I=1: a
        OR
            WHEN I=2: b
        OR
            WHEN I=3: C
    TCELES
ROF
```

Only alternatives with open guards can complete. Some of the alternatives may compute an acceptable result. while others may not. The essential problem is the choice between successful alternatives, or an indication of failure if there are no such alternatives. An
error condition is raised when no altemative is successful. An ALGOL-like language construct embodying this situation is:

```
ALTBEGIN
    ENSURE guard 1 WITH method ( OR
    ENSURE guard}2\mathrm{ WITH method}2\mathrm{ OR
    .
    ENSURE \mp@subsup{guard}{N}{}\mathrm{ WITH method}N\mathrm{ OR}
    FAIL /* no method succeeded */
END
```

Figure 1: Alternative Block
What we want is for at most one method to be applied to our problem, or for whatever conditions constitute failure to be indicated. Each method, $1 . N$, has associated with it a guard condition, which it must satisfy to be considered successful. A method is called an alternative. When the alternatives are composed into a block, as illustrated in figure 1 , the interpretation is that one alternative (possibly failure) is selected nondeterministically. The non-determinism defined in the semantics of alternative selection can often be exploited with parallelism for higher-performance computing. Nondeterministic algorithms have been discussed in the literature; Cohen [Cohen1979a] provides a good survey; he mentions the possibility of parallel execution, algorithms for which we will describe in Chapter 3.

The selection is non-deterministic and unfair, in that the selection of alternates is not equiprobable, and should not be: it's clear that the alternative of failure should be given as low a probability of success as is possible, noting that when all the alternatives fail its conditional probability must be 1 . The semantics of the construct are similar to Dijkstra's [Dijkstra1976a] guarded commands, in the special case where the same guard is used for all the statements. In an implementation setting, the construct resembles the Ada [Ledgard1981a] select with guarded alternatives; the selection of open (i.e.. have satisfied the guard) alternatives is arbitrary.

Once our model is defined, and the semantics thus fixed, we can apply semantics-
preserving transformations to increase performance or achieve other goals. A successful transformation, then, has two requirements. First, it must correctly preserve the semantics. Second, it must achieve the goal set for it, e.g., a performance increase.

We present (1) a model for selection of alternatives in a sequential setting, (2) a transformation which allows alternatives to execute concurrently, (3) a description of the semantics-preservation mechanism, and (4) parameterization of where the performance improvements can be expected. Additionally, we show example application areas for our method.

### 1.1. Thesis Summary

This thesis examines methods for solving problems for which there are alternative solution methods. When all such solution methods are equally acceptable. executing the alternatives concurrently and selecting the first successful computation can improve response time. The particular solution method which is fastest often depends on the data. The major contributions of this work are:

- A scheme for parallel execution of nondeterministic algorithms. Consistency and correctness are ensured with "Multiple Worlds," where "copy-on-write" pagemanaged storage prevents side-effects and inhibits combinatorial explosion in state storage requirements.
- An analysis of the possible speedup which identifies necessary properties of the alternatives.
- An analysis of the overhead involved in the proposed parallel execution scheme. tools for measuring this overhead, and measurements derived with these tools.
- Use of the rfork() remote fork mechanism to create a copy of a running process; this extends the idea of process migration.
- Use of sibling elimination to reduce the scheme's effect on throughput.
- Predicates to extend the scheme to allow interprocess communication. where interacting processes are also "copy-on-write."
- Example applications from three areas: parallel execution of logic programs. concurrent execution of recovery block altemates expressed in our language (RB), and numerical analysis, particularly polynomial root-finding.

The scheme for parallel execution (Chapter 3) relies on the availability of alternative solution methods. Nondeterministic algorithms give rise to alternatives by making several choices, each one of which gives rise to an alternative. These alternatives are executed in parallel; the results of the first successful computation are chosen. At this point, the slower computations can be eliminated. The alternatives share a common parent process, from which their state is inherited "copy-on-write" so that unchanged state can be shared, while internal consistency and correctness of each alternative are maintained. The rfork() primitive extends the spawning of alternatives to a distributed setting. To ensure correctness of interprocess communication, completion predicates are attached to all messages sent by a process which has siblings; these ensure that receiving processes do not make updates which depend on state which may be eliminated.

We identify the sources of overhead for concurrent execution using this scheme, and provide a measure of the performance improvement ( $P I$ ) which can be used to compare execution strategies. We use this measure to identify opportunities for response time improvement. The best sort of situation (discussed formally in Chapter 2) for our approach is one where: the alternatives require a significant amount of computation time, as encapsulated in the mean of their execution times; each alternative changes a small amount of the state of the calling process. thus reducing the overhead due to copying state; there is enough difference between the execution times of the alternatives that choosing the fastest and killing the others is worth the overhead of spawning the copies and deleting the slower siblings. The speedup is dependent on both the fraction of execution time devoted to overhead and the variance exhibited by the execution times of the alternatives. The implication is that if overhead can be understood and controlled, there is an opportunity for speedup roughly proportional to the variance; thus superlinear speedups, where the execution time is less than $O(1 / N)$ for $N$ processors, are possible under this scenario.

Since the potential for speedup is sensitive to overheads, we examined these overheads in Chapter 4. Copying is the major overhead in the creation and maintenance of the concurrent executions. Although there had been no previous work examining the efficacy of "copy-on-write," our results indicate that the technique is extremely effective in practice. We provide several useful measurement tools to gather execution time data. These tools are used to adapt our measurements to new computers; thus, with a few
measurements the domain for performance improvement using this method can be delimited. We implemented a system which performs a remote fork, which is similar to process migration with the exception that two processes exist when the operation is successful, rather than one migrated process. Our measurements confirm that the major overhead cost is copying, and further, offer empirical proof that child processes can be "spawned" in a distributed execution environment. Response time is affected by the response times of several system components; we examine two subcomponents. the disk and the network, in chapter 4. The final overhead is sibling elimination, which we modeled on a multiprocess timesharing system. This should represent the worst case execution, for which the results are encouraging, as they show the elimination to be remarkably cheap, and insensitive to process behavior. Given that the overheads are predictable and parameterizable these parameters can be used in deciding whether to apply the concurrent execution scheme.

After identifying the opportunities for response time improvement with analytic work and measurements, some example applications for concurrent execution are described. These example applications are drawn from several different areas of computer science, and illustrate the general utility of the scheme.

- Parallel implementation of logic programming languages, particularly ORparallelism in Prolog, provides an appropriate environment, because the computation is data-driven. The execution time and control flow can vary greatly with the input. The way in which unification operates (as a "sophisticated pattern matcher`') leads to an overwhelming preponderance of read references made to page-managed memory: while a high percentage of references are writes, these references are mainly to the stack, and thus locality is high; stack "growth" can be handled locally, reducing copying. Many logic programs have a great degree of parallelism, so that appropriate opportunities must be identified with respect to the overheads implied by our scheme. In particular, coarse-grained parallelism is better to exploit than fine-grained parallelism at the level of overhead we have observed. Our scheme deals with side-effects other than variable binding, and can run efficiently on general-purpose hardware.
- Distributed execution of recovery block alternates uses the "fastest-first" behavior
in an attempt to find a rapid failure-free path through the computation. Recovery blocks are designed for fault-tolerance, thus, there may be further requirements beyond fast execution time; we describe our RB language and suggest modifications to the concurrent execution scheme to increase robustness. These changes increase the amount of state available in a system to facilitate recovery.
- Polynomial Root-Finding with the Jenkins-Traub algorithm for finding complex roots of polynomials with complex coefficients was chosen as the third example. The sequential algorithm chooses an angle at random in its search for a root; we choose multiple angles and execute the choices in parallel. This application has shown speedups of a factor of 3 , and some examples demonstrate desirable properties of our execution scheme in the face of alternative failures.

In summary, we examined a problem setting, introduced and applied a parallel execution scheme for improved response time, and demonstrated application areas and methods for determining appropriate applications through measurement of overhead.

## 2. Theoretical Underpinnings

### 2.1. Extremal Behavior

Consider the $n$ random variables $X_{1}, \ldots, X_{n}$, each representing the execution time of a computation. The order statistics are computed by permuting the values of $X_{i}$ to form a new set of values $X_{(1)}, \ldots X_{(n)}$ such that $X_{(i-1)} \leq X_{(i)} \leq X_{(i+1)}$. This ascending order implies that $X_{(n)}=\max _{i=1}^{n} X_{i}$ and $X_{(1)}=\min _{i=1}^{n} X_{i} . X_{(1)}$ can be interpreted as the random variable defined by the first event to complete. Then, the distribution of $X_{(1)}, F(t)=P\left(X_{(1)} \leq t\right)$ is given by $1-\prod_{i=1}^{n}\left[1-F_{i}(t)\right]$ where $F_{i}(t)$ is the distribution of $X_{i}$. The random variables are assumed to be independent. If the random variables are exponentially distributed each with parameter $G_{i}$. then $1-F_{i}(t)=\exp \left(G_{i} \cdot t\right)$. Thus, $X_{(1)}$ is also exponentially distributed with rate (parameter) $\sum_{i=1}^{n} G_{i}$. This means that the average value of $X_{(1)}$ is given by $\frac{1}{\sum_{i=1}^{n} G_{i}}$. In particular, if the random variables are identically distributed (all the $G_{i}$ are the same), the average time to completion is given by $\frac{1}{n \cdot G}$ or, in other words, the average time to completion $(\min )$ is $\frac{1}{n}$ times the average execution time of a replica, $\frac{1}{G}$. Thus. the performance gain in the extreme is $O\left(\frac{1}{n}\right)$, that is. linear in the number of processes. A short discussion of limit theorems for order statistics is available in Feller [Feller1970a] and a detailed treatise on the behavior of these extremes is given by Galambos [Galambos1987a]. While the exponential distribution eases analysis and leads readily to a linear speedup, there remains the possibility of arbitrary speedups. since the speedup is a function of the distribution. We should note as an aside that system throughput is not affected if the linear speedup occurs, since it is a measure of the amount of useful work performed in unit time. We compare $n$ processors executing until first completes versus one executing a random instance. $\langle X\rangle$ is the expectation of random variable $X$. The analysis shows that $n \cdot\left\langle X_{(1)}\right\rangle=\left\langle X_{i}\right\rangle$; thus, there is no decrease in system throughput at the limit, even though the work of $n-1$ processors might be interpreted as wasted.

Since the random variables must be independent for the analysis to hold, the execution times should be random. Two cases exhibiting this behavior are given in Appendices XVII and X. Appendix XVII shows a simple search problem. The experimental apparatus is given; the problem closely approximates many classical search problems where the input is random. For example, there is an obvious mapping between the random search problem and the problem of finding a name in a phone directory, given a number. Measurements are shown in figure 2:


Figure 2: Use of randomness in search (Appendix XVII)
The results provide an existence proof that the tradeoff between randomization and parallelism can be exploited for speedups in practice. Another interesting aspect of our
approach is that the variance of the time to completion $X$ is reduced by order $O\left(\frac{1}{n^{2}}\right)$ when the execution times are exponentially distributed. This implies that the distribution of $X$ tends to be very sharply focused around the best time possible. This fact has important implications for systems where the execution time must be predictable, such as realtime systems. Actual algorithms exhibit this behavior to the extent that you may randomize their execution times to be independent random variables. For example, Appendix X shows that a random choice of partitioning element in quicksort has little effect on the execution time, even though the behavior is as predicted, i.e., decreasing execution times and decreasing variance. The performance is limited in this case since the distribution function. $F()$. is bounded on the bottom by $O(n \log n)$, and the mean execution time of the algorithm approximates this quite closely.

The next section discusses the measurement of speedup we will use to test the success of the method, and shows the conditions necessary for this speedup to occur.

### 2.2. Performance Analysis

The possibility of a performance increase stems from the fact that we can select the fastest alternative by means of the synchronization protocol. The introduction's argument promises linear speedup. There are two facts which frustrate this promise. First, the dispersion of execution times must be significant, and the probability distribution function, $F()$, must have a long "tail." Second, we ignored the overhead involved in concurrent execution such as copying, sibling elimination, and processor contention.

The cost we must pay for obtaining execution time proportional to the time for the fastest alternate is use of available hardware.

Note that the action of continuing execution of the successful alternative and the process of sibling elimination can take place asynchronously. The effects of various overheads and system parameters are analyzed in the next section.

### 2.2.1. Overhead

To understand the overhead implied by the method, we should compare a sequential execution of the construct, in the best case, where the fastest alternative is selected. There are penalties we are paying for parallel execution of all alternatives. We must compare this scheme with sequential execution of the alternative which will be selected in any case. These penalties are:

1. Memory Copying. In the distributed case we must copy state for a remote child so that it can read or write locally. In the shared memory multiprocessor case, the copying overhead (in execution time) is reduced as the interprocessor bandwidth is much higher. There is additional copying to be performed during synchronization, as the changed state is updated in the parent's storage. The parent is constrained to remain blocked while the children are executing.
2. Sibling elimination. This is asynchronous, and naturally parallel, but the instructions to terminate the alternates must still be issued, and they increase with the number of alternates.
3. Effect on throughput, due to wasted work. As our bias has been towards execution time as a performance goal, we were willing to trade away throughput. Users may want to know what the tradeoffs are here, so the effect on system throughput should be analyzed.

### 2.2.2. Analytic Description

Assume that we have $N$ alternative methods of performing a computation. A computation is a transformation from an input set (or Domain) to an output set (or Range); these sets consist of state vectors, intended to describe the relevant state of the world, i.e., the machine state. For Domain $\mathbf{D}$ and Range $\mathbf{R}, \vec{x} \in \mathrm{D}$ is transformed via the computation into some $\vec{y} \in \mathbf{R}$, thus we could write $\vec{y}=C \overrightarrow{(x)}$ There may be several such $C$ which we classify as interesting. Transformations of $C$ which add or remove useless operations are infinitely numerous, but not interesting. Algorithmic differences, random parameters, or significant differences in implementation technique are interesting. Assume that the $N$ alternatives postulated earlier are $N$ such interesting $C$ s, and that they will be applied to some $\vec{x} \in \mathrm{D}$. Each $C$ consists of some series of steps, where $\vec{x}$ is transformed into $\vec{x}^{\prime}, \ldots$ until $\vec{y}$ is achieved. Each step requires some amount of clock time, $\tau$, to complete; for
$C(\vec{x}) \tau(C, \vec{x})$ is the sum of these times. $\tau$, the execution time, gives us a way of comparing the performance of two computational methods on the same input, say $\vec{x}$.

There are many practical situations in which we want to minimize the computation time required for the transformation of $\vec{x}$ to $\vec{y}$. We will denote the $N$ alternatives as $C_{1}, \ldots, C_{N}$. Since our goal is minimizing execution time, let us consider some possible relations between the $C_{i}$ on elements of D .

1. $\tau\left(C_{i}, \vec{x}\right) \leq \tau\left(C_{j}, \vec{x}\right)$ for every $\vec{x} \in \mathrm{D}$ which interests us. It's clear that we should use $C_{i}$ and discard $C_{j}$ for every $i$ and $j$ for which this holds.
2. $\tau\left(C_{i}, \vec{x}\right) \leq \tau\left(C_{j}, \vec{x}\right)$ for some $\vec{x}$ which interest us, and we can accurately predict for which $\vec{x}$ this relation holds. In this case, we can construct a synthetic computation, $C_{N+1}$, which selects $C_{i}$ when this holds. To anchor the relation with an example consider the case of two list-sorting algorithms, $Q$ and $I$. $Q$ is faster than $I$ when the number of elements to be sorted is greater than 10 . Thus, using this knowledge, we can construct a synthetic sorting routine as follows:
```
sort( list, size ) :=
    if( size > 10 )
        Q( list, size )
    else
        I( list, size ).
```

The synthetic routine partitions the input domain by performance, and thus achieves performance superior to either $Q$ or $I$. The tough point here is the partitioning; it's rarely as simple to delimit performance boundaries as " size < 10." If the input set can be partitioned, but only at significant computational cost, the desired property of the synthetic routine, that $\tau\left(C_{N+1}, \vec{x}\right)=\min _{i} \tau\left(C_{i}, \vec{x}\right)$ for all $\vec{x}$ of interest, may be achievable with the following technique.
If all interesting $\vec{x}$ are known in advance, we can associate one $C_{i}$ with each $\vec{x}$ in a precomputed table. Then, $\tau\left(C_{N+1}, \vec{x}\right)$ can be calculated by adding the cost of a table lookup to the cost of executing the table element on $\vec{x}$. In rare cases (rare due to the amount of state under consideration, and the sizes of $\mathbf{R}$ and D for the problems we are
interested in), we may be able to store $\vec{y}$ in the table, thus removing any execution time cost except for a table lookup.
3. $\tau\left(C_{i}, \vec{x}\right) \leq \tau\left(C_{j}, \vec{x}\right)$ for some $\vec{x}$ which interest us, but while interesting, the $\vec{x}$ cannot easily be related to $\tau\left(C_{i}, \vec{x}\right)$ Essentially, this means that the table lookup technique cannot be used, because we cannot reasonably precompute the values of $\tau\left(C_{i}, \vec{x}\right)$. This might be due to the input set, e.g., infinite size. For example, a naive quicksort is not stable, and where the list is ordered the sort is slow. In these cases, a stable sort with good performance, e.g., heapsort. may be preferable. However, it's clear that storing a lookup table of all "interesting" lists is infeasible, and pretesting for the "ordered" property is potentially expensive. Another problem is that $\tau\left(C_{i}, \vec{x}\right)$ may vary due to the execution environment (which may or may not be described by $\vec{x}$; it probably should be, for completeness), e.g., processor type, multiprocessing workload, or interactions with other computations. In these cases, where performance on the $\vec{x} \in \mathrm{D}$ is unpredictable, we might try other schemes:
A. Statistical data can be applied, e.g., quicksort is "almost always" $O(n \log n)$. Thus, we $1 l$ rarely go wrong using it.
B. An algorithm can be selected at random from amongst the $C_{i}$ when given $\vec{x}$.
C. The $C_{i}$ can be applied to $\vec{x}$ concurrently; the first $C_{i}$ which produces $\vec{y}$ is selected. The other $C_{i}$ are irrelevant and can be terminated. There is, however, overhead in setup and synchronization (selection) which cannot be ignored.
Scheme A relies on information which may not be available. Scheme B, when run repeatedly on some uniformly distributed input $\vec{x}$, will perform at the arithmetic means of the computations' performance, i.e., $\frac{\sum_{i=1}^{N} \tau\left(C_{i}, \vec{x}\right)}{N}$. Scheme C offers some opportunity for achieving the best performance on each input $\vec{x}$. We will try to characterize this opportunity. Note that there are two possibilities for concurrent execution, real and virtual. Real concurrency means that the evaluation of $C_{i}(\vec{x})$ is taking place simultaneously with that of $C_{j}(\vec{x})$ virtual means that there is some sharing of hardware, for example through muhtiprocessing.

### 2.2.3. Parallel Speedup

Our analysis must begin with semantics, as otherwise we are subject to criticism of the "apples and oranges' type. Such criticism stems from the observation that changing the problem in order to apply a program transformation makes performance results incomparable; we are comparing unlike programs.

To an observer, the concurrent execution of the $C_{i}$ must look like Scheme B. (as discussed above); that is, that we have followed a single thread of computation, chosen arbitrarily from amongst $C_{1}, \ldots, C_{N}$. Since the $C_{1}, \ldots, C_{N}$ may update shared state described by $\vec{x}$, we solve the problem by copying state when needed and by selecting some $C_{i}$ by virtue of its state changes. Thus, since the observer sees non-deterministic selection of one alternative, we must compare concurrent execution to sequentially performing some $C_{i}$, chosen arbitrarily (we'll assume randomly). Since, as stated previously, execution time is our figure of merit, we'll analyze with that intent, ignoring measures such as throughput. Arbitrary selection is trivial to implement; it costs no execution time for purposes of our analysis. The execution of the selected alternative $\operatorname{costs} \tau\left(C_{i}, \vec{x}\right)$ for the $\stackrel{\rightharpoonup}{x}$ under study. Thus, we can expect the mean cost to be $\frac{\sum_{i=1}^{N} \tau\left(C_{i} \stackrel{\rightharpoonup}{x}\right)}{N}$, the average of the $C_{i}$ 's times when applied to $\vec{x}$. For notational convenience. define $C_{\text {mean }}$ such that

$$
\tau\left(C_{\text {mean }}, \vec{x}\right)=\frac{\sum_{i=1}^{N} \tau\left(C_{i}, \vec{x}\right)}{N}
$$

By executing the $C_{i}$ concurrently, we will expect the cost of execution to be

$$
\tau\left(C_{\text {best }}, \stackrel{\rightharpoonup}{x}\right)+\tau(\text { overhead })
$$

where

$$
\tau\left(C_{\text {best }}, \stackrel{\rightharpoonup}{x}\right) \leq \cdots \leq \tau\left(C_{\text {worst }}, \stackrel{\rightharpoonup}{x}\right)
$$

and overhead is complex. Overhead consists of operations performed to support concurrent execution which would not be necessary in the nondeterministic sequential case. It consists of the following components:
setup: Instead of simply calling $C_{i}$, we must now spend cycles creating execution
environments for $C_{1}, \ldots, C_{N}$ : for example, setting up process table entries and page map tables.
runtime: This consists of copying memory areas which are shared between the $C_{1}, \ldots, C_{N}$ when updates are attempted. This performance is strongly influenced by locality of reference. Additionally, if $C_{\text {best }}$ is sharing resources, e.g., CPU time, with some $C_{i}, i \neq b e s t$, then all such $C_{i}$ 's runtimes must be added to the runtime overhead of $C_{\text {best }}$, as cycles spent processing $C_{i}$ are not spent processing $C_{b e s t}$.
selection: This is the cost involved in selecting $C_{\text {best }}$, e.g., deleting $C_{i}$ such that $i \neq b e s t$, cleaning up system state, such as performing the updates made by $C_{\text {best }}$, e.g., writing checks or bottling beer.

Thus, for a given $C_{1}, \ldots, C_{N}$ and $\vec{x}$,

$$
\begin{aligned}
& \tau(\text { overhead })= \\
& \tau\left(\text { setup }\left(C_{1} \cdots C_{N}, \vec{x}\right) \nrightarrow\right. \\
& \tau\left(\text { runtime }\left(C_{\text {best }}, \vec{x}\right) \oplus\right. \\
& \tau\left(\text { selection }\left(C_{\text {best }}, C_{1} \ldots, C_{N} \cdot \vec{x}\right)\right)
\end{aligned}
$$

and the parallel execution wins at $\vec{x}$ iff

$$
\tau\left(C_{\text {best }}, \vec{x}\right)+\tau(\text { overhead })<\tau\left(C_{\text {mean }}, \vec{x}\right)
$$

Thus, we can calculate the performance improvement $(P I)$ as:

$$
P I=\frac{\tau\left(C_{\text {mean }}, \vec{x}\right)}{\tau\left(C_{\text {best }} \cdot \vec{x}\right)+\tau(\text { overhead })}
$$

essentially a ratio of execution times. For illustration, consider a case where $N=3$, on input $\vec{x}$. Thus, we have three methods $C_{1}, C_{2}$, and $C_{3}$. Let $\tau$ (overhead) be 5 . Some possible relations are tabulated:

|  | $\tau\left(C_{1}, \vec{x}\right)$ | $\tau\left(C_{2}, \vec{x}\right)$ | $\tau\left(C_{3}, \stackrel{\rightharpoonup}{x}\right)$ | $P I$ |
| :--- | :---: | :---: | :---: | :--- |
| $(1)$ | 10 | 20 | 30 | 1.33 |
| $(2)$ | 1 | 19 | 106 | 7.0 |
| $(3)$ | 20 | 20 | 20 | 0.8 |
| $(4)$ | 1 | 2 | 3 | 0.33 |
| $(5)$ | 115 | 120 | 125 | 1.0 |
| $(6)$ | 100 | 200 | 300 | 1.9 |

What can we infer from the examples? (3) indicates, along with (5), that the size of the differences matters. (4) shows that the relative magnitudes of the execution times and the overhead matters. (6) shows that the effects of the overhead (under our assumptions) diminish with increasing relative execution time. (2) illustrates a good situation, where the difference

$$
\tau\left(C_{\text {worst }}, \vec{x}\right)-\tau\left(C_{\text {best }}, \stackrel{\rightharpoonup}{x}\right)
$$

is large. This magnitude of difference is well-encapsulated by such a statistical measure of dispersion [Robbins1975a] (letting values of $\tau$ serve as the random variable) as the variance, and the variance is easily computed as:

$$
\text { variance }=\text { mean value }\left(\tau\left(C_{i}, \stackrel{\rightharpoonup}{x}\right)^{2}\right)-P I^{2} \cdot\left(\tau\left(C_{\text {best }}, \stackrel{\rightharpoonup}{x}\right)+\tau(\text { overhead })\right)^{2}
$$

However this re-expression seems to serve little purpose, as we can manipulate the simple relationships describing PI into forms which genuinely ease analysis. Additionally, the variance, although a good measure of dispersion is not precisely what we want to predict speedup. This is because values other than the best execution time can be altered with no effect on the mean execution time. Altering these values can increase the value of the variance but not change the potential for speedup. In the remainder of this thesis, when we use the term 'variance', we will mean the layman's variance, a measure of dispersion, rather than the statistician's variance.

We can analyze precisely the domains in which there is a performance improvement $(P I>1)$. Letting $R_{\mu}=\frac{\tau\left(C_{\text {mean }}, \vec{x}\right)}{\tau\left(C_{\text {best }}, \vec{x}\right)}$ and $R_{o}=\frac{\tau(\text { overhead })}{\tau\left(C_{\text {best }}, \vec{x}\right)}$, we can calculate $P I$ as:

$$
P I=\left[\frac{1}{1+R_{o}}\right] \cdot R_{\mu}
$$

This re-expression isolates the effect of the dispersion, encapsulated in $R_{\mu}$, from the effect of the overhead, encapsulated in $R_{o}$. Holding one of $R_{\mu}$ or $R_{o}$ fixed allows us to estimate the effects on Pl caused by the other. The behaviors are illustrated in figures 3 and


Figure 3: $P I$ as a function of $R_{\mu}\left(R_{\mathrm{o}}=0.5\right)$
4. The relationship illustrated by the first figure is with $R_{o}$ set to the constant value ${ }^{2} 0.5$;

[^1]$R_{\mu}$ is varied between 0 and 5 , and the values can easily be scaled. The curve is not interesting, as it's a direct proportion for fixed $R_{o} ; R_{o}$ determines the slope of the line. with $R_{o}=0$ the best case giving a slope of 1 . This tells us that the performance improvement we can expect will be proportional to the variance of $\tau\left(C_{i}, \vec{x}\right)$ damped by whatever effect $\tau$ (overhead) exhibits. Holding $R_{\mu}$ fixed and varying the overhead is more interesting, as figure 4 illustrates. The $y$ axis has $P I$ scaled proportional to $R_{\mu}=\exp (1.0)$, and the scales are $\log -\log$ to view a wide range of values.


Figure 4: $P I$ as a function of $R_{o}\left(R_{\mu}=\exp (1.0)\right)$
This tells us that varying the overhead has a significant effect on the performance improvement we achieve, when scaled against the variance in execution times. An important fact which we can deduce from this performance analysis is that with sufficient variance, and small enough overhead, N processors can exhibit superlinear speedup by
parallel execution of N serial algorithms, as opposed to parallel execution of one serial algorithm which has been "parallelized."

### 2.2.4. Domain-wide performance indices

In the previous section, we constrained our analysis of the performance of concurrent execution to the performance at a single input state, encapsulated symbolically in $\vec{x}$. Thus, we have in our performance index. PI, a function of $\vec{x}$. This allows us to identify states at which the execution of alternatives will provide a performance improvement. Now, were we to consider the entirety of the input domain $\mathbf{D}$, we might parameterize the performance at each $\vec{x}$ as $P I \vec{x})$.Our goal in securing a performance improvement is maximizing the "overall" performance improvement. How might we measure this? Let the overall performance improvement be calculated as

$$
O P I(\mathrm{D})=\sum_{\vec{x} \in \mathrm{D}} P l(\vec{x}) \cdot \operatorname{prob}(\stackrel{\rightharpoonup}{x})
$$

$\operatorname{Prob}(\vec{x})$ is the probability that a given input state $\vec{x}$ will occur: we assume that $1=\sum_{\vec{x} \in \mathrm{D}} \operatorname{prob}(\vec{x})$ We note that special care must be taken when $|\mathrm{D}|$ is $+\infty$, i.e., the input set is infinitely large, as (1) this case is common; and (2) we want a useful measure. Thus, our formulation of $O P I$ serves as a weighted value of the $P I(\vec{x})$ values over $\vec{x}$ selected from domain D. This is just the expectation of $P I(\vec{x}) \leqslant P I(\vec{x})>$. If we assume the $\vec{x}$ are equiprobable, then

$$
O P I=\sum_{\vec{x} \in \mathbf{D}} \frac{P I(\vec{x})}{|\mathbf{D}|}
$$

since $\operatorname{prob}(\vec{x})$ is $\frac{1}{|\mathrm{D}|}$. How good a measure is this? First, examine the case where there is no improvement in the performance, i.e. $P I=1$. Then,

$$
O P I=\frac{\sum_{\vec{x} \in \mathrm{D}} 1}{|\mathbf{D}|}=1
$$

which certainly makes sense, given the situation that this symbolic representation describes. If we expand the expression of $P I(\vec{x})$ we get

$$
O P I(\mathrm{D})=\sum_{\vec{x} \in \mathrm{D}}\left[\frac{\tau\left(C_{\text {mean }}, \vec{x}\right)}{\tau\left(C_{\text {best }}, \vec{x}\right)+\tau(\text { overhead })}\right] \cdot \operatorname{prob}(\vec{x})
$$

The value of $O P I(\mathrm{D})$ is maximized when the weighted average of the $P I(\vec{x})$ values is maximized. What factors are there? First, there is the input distribution, which for many interesting problems we know little about ${ }^{3}$. Second, there is the overhead, and the dispersion of values exhibited by $\tau\left(C_{i}, \vec{x}\right)$ at a given $\vec{x}$. It's clear that reducing the overhead always helps; for the limiting case of $\tau($ overhead $)=0$,

$$
\operatorname{PI}(\vec{x})=R_{\mu}=\frac{\tau\left(C_{\text {mean }}, \vec{x}\right)}{\tau\left(C_{\text {best }}, \vec{x}\right)} \geq 1
$$

Now, any dispersion of values of $\tau\left(C_{i}, \vec{x}\right)$ causes $\tau\left(C_{\text {mean }}, \vec{x}\right)>\tau\left(C_{\text {best }}, \vec{x}\right)$ so that $R_{\mu}>1$, and thus $P I>1$.

It is worthwhile to discuss the value of $\operatorname{prob}(\vec{x})$ further. We estimated the sequential performance by assuming that a random choice be made from among the alternatives. This represents the "best guess" that could have been made using our statement of the problem. What if we relax the constraints on information available, so that rather than equiprobable inputs, we have some data gathered about the probability distributions of the input. The data may have been gathered through a small statistical sample, such as might be possible from a short historical record of the inputs ${ }^{4}$. How can this data help?

If $\tau\left(C_{i}, \vec{x}\right)<\tau\left(C_{\text {mean }}, \vec{x}\right)$ with probability $P_{i}>\frac{1}{|\mathrm{D}|}$, the expected performance of the sequential algorithm might be improved by always choosing $C_{i}$ rather than choosing at random. Unfortunately, we also need to know the values of $\tau\left(C_{i}, \vec{x}\right)$ to be effective. The difference in expected values amortized over the input domain can be represented as

$$
\sum_{\vec{x} \in \mathrm{D}}\left(\tau\left(C_{i}, \vec{x}\right)-\tau\left(C_{\text {mean }}, \vec{x}\right)\right]
$$

If we refer to the inner term as $\Delta_{i}$, we know that $\operatorname{prob}\left(\Delta_{i}<0\right)=P_{i}$. But knowing this value, and not knowing $\left|\Delta_{i}\right|$, we can not predict whether a performance increase will

[^2]result. With the advance of systems for process migration and load-balancing, determination and control of this value becomes more computationally difficult. The "loadbalancing'' can contribute to the observed execution times in such a way that it generates an unpredictable domain-wide dispersion [Barak 1985a].

### 2.3. Conclusions

We have analyzed the properties necessary for a set of alternatives to show a performance improvement, in particular the relationship between the dispersion of execution times and the overhead associated with the parallel execution. These properties were general; for example, $\vec{x}$ may encapsulate the execution environment so that processors with heterogeneous performance can be accounted for.

The factors contributing to $\tau$ (overhead) were dissected; they are mainly a property of the execution environment and not of the application. Thus, measurements of overhead may allow us to calculate the performance improvement, $P I$, based solely on the application characteristics. thus removing a variable from any analysis.

## 3. Algorithms for Parallel Execution

### 3.1. System Model

A process is an independently schedulable stream of instructions. In implementations, it is often associated with some unit of state, e.g., an address space, and a set of operations provided by a kernel to manage that state. Interprocess communication is done solely through passing messages. Thus, a message is the only means by which:

- $P_{m}$ can make $P_{j}$ aware of a change in $P_{m}$ 's state.
- $P_{m}$ can cause a change in $P_{j}$ 's state.

Interprocess communication (IPC) is assumed to behave reliably (no lost or duplicated messages) and FIFO (no out of order messages).

System state is divided into two types, source and sink. The division is made on the basis of idempotence; operations on sink devices can be retried without the effects being visible, while operations on sources cannot be retried. For definiteness, consider a page of backing store and a teletype device, respectively. Side effects which affect sink state can be hidden; this is a common technique in the implementation of such abstract operations as transactions; the idea is that the transaction has the property of atomicity, meaning that either none or all the transactions component actions occur, and that intermediate states are not observable external to the transaction. Complex transactions may involve reads, which can occur unhindered, or writes, which must be done to a temporary copy until the transaction commits, or in other words, makes its changes permanent. Reads intended for the recently written copy are satisfied by that copy so that the transaction is internally consistent, i.e., it can read what was written.

Sink state is manipulated as fixed-size pages. All sink state can be represented in this fashion; this is clear from implementations of a single-level store, as in MULTICS [Organick1972a]. Thus we bury the entire memory hierarchy under the page abstraction; files are named sets of pages, and thus mechanisms which are used to transparently access files over networks [Sandberg1985a, Weinberger1984a] can be used to hide the network through the page management abstraction. This has been successfully done in at least one commercial system, Apollo’s Domain ${ }^{\text {TM }}$ [Nelson1984a, Leach1983a, Leach 1982a].

### 3.2. Process Management

Two primitives encapsulate the entire semantics of the process management component. The process management component is concerned with the mutually oblivious alternatives. To spawn the alternatives, the parent uses alt_spawn ( $n$ ). which returns numbers from 1 to $n$ in the alternates and 0 to the parent. Thus a language preprocessor applied to a program with mutually exclusive alternatives would generate (in pseudo-C):

```
switch( alt_spawn( n ) )
I
    case 0:
        alt_wait( TIMEOUT );
        fail(); /* if returned */
    case 1:
        /* Eirst alternate */
    case n:
        /* n-th alternate */
        alt_wait( 0 );
}
```

Figure 5: Use of alt_spawn() and alt_wait()
The purpose of alt_wait () is manifold: the essence is establishing a single path through the tree of possible computations which is reflected in the execution history of the running process.

Alt_wait() is the synchronization locus. Alt_wait() takes a TIMEOUT value as an argument: the point is that this value should be chosen such that if TIMEOUT time units have elapsed, it is highly probable that no alternative succeeded. While choosing such a value is hard, most computations have an execution time which is clearly unacceptable to the application: this value can then be used. The point of passing such a timeout value will be seen shortly.

When a spawned alternate calls alt_wait() at the termination of its computation, a rendezvous between the alt_wait () ing parent and the child is effected. The behavior is much like that of the UNIX exec () system call, where the new data and executable code are read in from a named file. For alt_wait (), the parent process absorbs the state changes made by its child by atomically replacing its page pointer(s) with that of the child. Thus, the flow of control through the child appears to have been seamless, up to and including maintenance of the process id. It is as if the parent process was "lucky," and performed the execution of the fastest alternative itself.

Use of these primitives is shown by concurrent execution of the program segment in figure 5 shown in figure 6:


Figure 6: Concurrent Execution of Alternates
If all the GUARD conditions have been satisfied, a process which completes its program segment attempts to synchronize. If any of the conditions required by the GUARD were not satisfied, the process aborts without synchronizing. Note that the GUARD can be executed before spawning the alternative, in the child process, at the synchronization point, or at any combination of these places, for redundancy. We currently expect the
child process to execute it, thus speeding up spawning and synchronization.

### 3.2.1. Synchronization

It is at the synchronization point that the data for sibling elimination are available; all processes which assumed that the successful child had failed must be deleted, as they have made an assumption we now know to be false. To minimize the effect on throughput, when an alternative is selected, its "siblings" are eliminated. This is done by informing the scheduler that the process is to be terminated. The deletion can be done synchronously (where the other alternates are deleted before execution resumes in the parent) or asynchronously (where the deletion occurs at some time after the alt_wait () resumes in the parent, but exactly when is not specified); we suspect that asynchronous elimination will give better execution-time performance. This is because the execution time we are concerned with is that of the successful alternative. If the successful alternative cannot continue executing until its siblings are eliminated, then it is waiting, and thus will have increased execution time. Now, on the other hand, if the sibling elimination is started but the successful alternative does not wait for completion, it will result in faster execution, as work (and delay) has been removed from its execution trace. Measurements in Chapter 4 show that these suspicions are well-founded.

Now, communications problems or system failures may prevent this information from reaching the scheduling component of a remote system, yet we must still preserve the 'at most one'" semantics of our design. The backup in this case is that the synchronization action is designed so that it can be done at most once; that is, if the remote system attempts synchronization for the alternative it is executing, it is informed that it is "too late' for the synchronization, and it should terminate itself. In applications where this might create a single point of failure, the synchronization is set up as a majority consensus [Thomas 1979a] decision across several nodes. The engineering tradeoff here is between performance and reliability; the additional communication and protocol of multiple-node synchronization is the price paid for increased robustness of the synchronization.

### 3.2.2. Atomicity

An important question is "when do the alternate's changes become visible?." Clearly, this must occur at some point after the synchronization policy described above has been effected. Since we have taken the trouble to prevent the effects of other alternatives being visible, the update will be, by default, atomic (although we can have an ordered set of intermediate states made visible by the timestamping information we've preserved). Since any state changes must only be visible after the synchronization point, we'll assume that the state changes are made atomically, to simplify the discussion. It should be clear how the intermediate state changes can be made visible, but it seems pointless. How, then, can the state updates be made atomic? The method depends on cooperation between the method for supporting alternatives and the memory manager. Since we have required the predicates (described in the next section) to be stored in such a way that the value of a predicate is stored in one place. we can atomically update that information. Thus, an atomic state change is achieved as follows:

```
begin ATOMIC:
    All pages to be changed are predicated
        with FALSE;
    The predicating conditions (all TRUE) are
        removed from the alternate's pages.
end ATOMIC
```

Thus, it appears to other processes that all changes (to address spaces, or to what could normally be considered files) occur at once. The way in which I/O (e.g., printing a value) takes place must be considered carefully in an implementation, but the point at which the predicates can be removed from the pages is the same point at which I/O can be performed using these pages.

### 3.3. Predicates

Ideally, we would like an alternative to carry on with its computation as much as it can before either blocking or synchronizing. To effect this, we add "predicates" to the messages.

### 3.3.1. Representation of Predicates

Each page object has associated with it some set of predicates; these are the same predicates that are associated with message-sending. In most cases, there will be NO predicates associated with a page, as there is only one timeline, reality, that it is associated with; there are of course no conditions on reality.

Predicates might be stored as a bit-map, indexed by the process id, if the number of possible process ids is small (e.g., if it is limited by the Operating System's process table). If this is so, then predicate matches are essentially performed with existing machine instructions for comparisons, and rules for deciding whether a predicated object is accessible by a predicated process. Alternatively, the predicates can be stored as lists; in this case, there must be a more sophisticated, and possibly slower, comparison procedure. One idea is to store a linked list of process ids as a predicate; elements of this list can then be checked against their process table entries to determine the value of the predicate. If forward and back pointers ${ }^{5}$ are used, we can update the value of these elements as processes change state, with the idea that processes change state much less frequently than they make memory references to objects. It seems both "fairer" and more practical to control state by using software-implemented per-process predicates. The arguments are as follows:

- Any mechanism which stores predicates at the page level must pervade the page management system. Thus, all users must pay the cost for the extra page management data, while not all users may have need for the data and page management facility.
- Predicates are used only in the process control and message transmission activities. Thus, predicate changes and evaluations are performed only at points where processes change state, not on a per-access basis.
- A bit-map for a system with 8192 processes would require 1024 bytes; thus for a system with 1024 bytes per page, there would be a page of predicates per page of data. Additionally, the comparison of predicates would have to take place at each page reference, thus slowing an already time-intensive operation. If the bit-maps are stored with the processes, the storage requirements grow as $O\left(n^{2}\right)$ for a bitmap, thus for 8192
${ }^{5}$ Back pointer here means that attached to the predicate is a list of processes which depend upon its value.
processes, 8 megabytes of bitmap storage would be required, and it is expected that most of it would be unused or sparsely used. There is a possibility for a time/space tradeoff since all bitmaps except for those of running processes can be stored in a compressed form. However, efficient compression algorithms often require significant execution time to compress and uncompress data, and this penalty would be paid at an already busy point, the context switch.
- With a list implementation, processes with no predicates would pay little (2 pointers) overhead, and testing this condition would require about two instructions: as has been pointed out in several performance studies [Leffler1984a, McKusick1985a], several thousand instructions are executed per I/O or context switch; thus the added overhead is significantly less than one per cent for processes which don't use the facility.
- As the system is described, there are three possible values which must be stored: "must complete," "can't complete," and "don't care." Thus, at least two bits per process are necessary, increasing the bitmap storage requirements by a factor of two. With lists, the presence of a process identifier in either the "can't complete" or "must complete" lists indicates that condition, and its non-presence in both lists indicates "don't care."

Thus, we will implement multiple "copy-on-write" [Bobrow1972a] forks to maximize sharing during parallel execution, and keep updated and newly-written pages linked in a per-process descriptor table. Pictorially,


One important notion is that the process-stored information must be dependencies, so that storage of the predicate values themselves can be logically centralized, even if this is
not the case in reality.
Two operations, READ_PAGE and WRITE_PAGE are defined: it's obvious that any value manipulation can be performed with these primitives. They provide the point of interaction between the process state, as defined by its predicate values, and the state of external values. These operations can be implemented as follows:

```
READ_PAGE:
    Obtain the PREDICATES of the calling PROCESS.
    Examine the PAGE TABLE for a PAGE
        such that No PREDICATES assumed FALSE
            by the PROCESS are TRUE.
    IF no such page is found, FAULT,
        setting the newly read page's
        PREDICATES to that of the PROCESS
    ELSE return a pointer to PAGE FI.
```

and

```
WRITE_PAGE:
    Obtain the PREDICATES of the calling PROCESS.
    Examine the PAGE TABLE for a PAGE
        such that No PREDICATES assumed FALSE
            by the PROCESS are TRUE.
    IF such a PAGE is found
        IF No PREDICATES assumed TRUE
            by the PROCESS are FALSE, update the
            PAGE and return a pointer to it.
        ELSE
            Make a new copy of PAGE.
        FI.
    ELSE
        Allocate a new PAGE.
    FI.
    Set the PAGE's PREDICATES to that of
        the PROCESS.
    Update the PAGE
    Return a pointer to the PAGE.
```

To see how this works, consider the following example, with Parent Process $\mathbf{P}$ having spawned Alternates $\mathbf{A}_{1}$ and $\mathbf{A}_{2}$. The predicate associated with $\mathbf{A}_{1}$ is $\mathbf{A}_{1}$ and $\neg \mathbf{A}_{2}$, which we'll represent as 10 , and $\neg \mathbf{A}_{1}$ and $\mathbf{A}_{2}$, which we'll represent as 01 . The state previous to the alternates is then 00 , and 11 can't happen. The state 00 implies FAILURE after the alternates have been spawned, as it implies
$\neg \mathbf{A}_{1}$ and $\neg \mathbf{A}_{2}$. Let the initial state of the page map ( $\mathrm{P}^{\prime} s$ ) be:

| Virtual <br> Page\# | Predi- <br> cates | Real <br> Page\# |
| :---: | :---: | :---: |
| 0 | 00 | 19 |
| 1 | 00 | 12 |
| 3 | 00 | 3 |

Assume further that $\mathbf{A}_{1}$ has the following operations: READ 0, READ 1, WRITE 3, READ 0. and that $A_{2}$ has READ 0, WRITE 0, WRITE 0, READ 3. The resulting state of the page map will be:

| Virtual <br> Page\# | Predi- <br> cates | Real <br> Page\# |
| :---: | :---: | :---: |
| 0 | 00 | 19 |
| 1 | 00 | 12 |
| 3 | 00 | 3 |
| 0 | 01 | 37 |
| 3 | 10 | 8 |

The predicates are lists of process identifiers, some of which the sending process depends on completing successfully and others on which the sending process depends on not completing successfully. Thus, these are even simpler and easier to manage then the predicates described by Eswaran, et al.[Eswaran 1976a] The advantage of this representation over predication of data objects is that we can update the value of these elements as processes change status (e.g., running, blocked), with the idea that processes change status much less frequently than they make memory references to objects. Note that since the processes are mutually exclusive, there is no problem sharing the other objects on a page.

These lists are constructed in two ways. First, the predicates of a "child" process consist of those of the "parent:" this allows for nesting and potentially complex dependencies. Second, when the "parent" spawns each of its alternative "children," each of
the children additionally assumes that it will complete successfully, and that its siblings will not ${ }^{6}$. The state management strategy is "copy-on-write" [Bobrow1972a] with page map inheritance from the parent, thus it is easily implemented within the context of a system which provides such features, e.g., Mach [Young 1987a], and benefits from existing hardware support, e.g., for the $W^{\circledR}{ }^{\circledR} 32101$ Memory Management Unit [AT\&T1986a]. The software-implemented predicates are used in the process control and message transmission activities to maximize sharing. Updated and newly-written pages are predicated by virtue of their residence in a per-process descriptor table.

### 3.4. Interprocess Communication

### 3.4.1. Messages

A message from $P_{m}$ to $P_{j}$ has the following three part structure:

1) A sending predicate, encapsulating the assumptions under which the sender, $P_{m}$ sends the message.
2) The data comprising the message contents.
3) Some control information, e.g., sender id, destination id, etc.

Each process in a multiprocessing (e.g., timesharing, multiprocessor, or distributed) system has a unique identifier. used to identify the process both within the system (e.g.. for scheduling and resource allocation), and further, for interaction with other processes. This unique identifier can be constructed by concatenating several quantities at the time of process creation if local process ids are not unique (as in a distributed system), but processor names are:

```
<processor name, local process id, timestamp>
```

The timestamp is included to mitigate against re-use of local process ids. In the singleprocessor case. a local process id must be unique at any given time in order for the process to be named in an unambiguous fashion. However, as soon as the process terminates, the name, e.g., a table address or small integer, can be re-used. The local

[^3]picture of time may not hold across processors, and thus we force uniqueness with a timestamp.

### 3.4.2. Multiple Worlds

An idea from science fiction, inspired by Dewitt's [DeWitt1973a] multiple worlds notion, is appropriate here. The problem with interprocess communication stems from the fact that a given alternative may or may not be successful. In the case where it is successful, its execution results are available to the calling process. Where it is not successful, its results and any side-effects it may have generated must not be observable. These include side-effects due to interprocess communication.

The specialized side effects resulting from interprocess communication are controlled by a message layer which insures that any receiver of a message makes the same assumptions about the state of the world as the sender. Since a given sender may fail and make its assumptions invalid (as well as irrelevant) two copies of the receiver are created when the sent message causes the receiver's state to change; one copy is in the sender's "world,' the other is not.

In any case, the message subsystem must be aware of the decisions made, so that the copies are made once, at the first message receipt, not at each message receipt. Thus, as far as the receiving process is concerned, there are two worlds: one where the sender exists, and one where it doesn't. Depending upon whether the sending process completes successfully, one "world" may be realized.

The message system, the virtual addressing mechanism, and the process management mechanism are linked in the following way. When a receiving process accepts a message, its predicates ( $\mathbf{R}$ ) are checked against those attached to the message ( $\mathbf{S}$ ). If the assumptions that the receiver makes about the "state of the world," as encapsulated in the predicates, agree with those of the sender (e.g., $\mathbf{S} \subseteq \mathbf{R}$ ), the message is immediately accepted. If the receiver's predicates conflict ( $p \in \mathrm{~S}$ and $\neg p \in \mathrm{R}$ ), the message is ignored. and if the receiver must make further assumptions to accept the message ( $p \in \mathrm{~S}$ and $p \notin \mathbf{R}$ ), two copies of the receiver are created. One copy is created with the predicates set to the previous values with complete ( $S)^{7}$ : the other is set up with its predicates as

[^4]before, except that complete ( S ) is negated. ${ }^{8}$ This is shown in a revision of a previous figure:


Figure 7: Use of predicates
This is easy given the representation as two lists (i.e., "must complete" and "can't complete') of process identifiers. When the sending process succeeds or fails, one receiver must be eliminated to maintain a consistent "state of the world;'" at this point the additional assumptions which receipt of the message caused will become TRUE, and they can be eliminated from the lists.

To illustrate the idea, consider a group of communicating processes composed of P , Q. and S. P has children $\mathrm{a}, \mathrm{b}$. and $\mathrm{c}: \mathrm{Q}$ has children $\mathrm{d}, \mathrm{e}, \mathrm{f}$. This is illustrated by figure 8.

[^5]

Figure 8: Communicating processes
Suppose both $b$ and $f$ wish to communicate with $S$. We use the predicates to create multiple copies of $S$, with which the alternatives communicate. This is illustrated in figure 9 .


Figure 9: Communicating processes after message receipt
There are several implications to this scheme:

1. There is clearly a potential for a combinatorial explosion in the number of processes that exist, and amount of state which must be copied. Since the process creation is based on message receipt. it is "lazy'.
2. Commit is very fast, as it can be accomplished using only pointer (page descriptor) manipulation. For example, $\neg b f$ can be committed by simply updating the page descriptor table associated with S.
3. Messages must be duplicated, as all valid receivers must get the message.
4. The method is optimistic. Rather than locking a resource based on a predicate, the assumption embodied in the predicate is sent with messages and used to create new "worlds."

While a process has predicates which are unsatisfied, it is restricted from causing
observable side-effects, and thus cannot interface with sources.
This behavior is similar to that required of transactions. Transactions [Gray1978a, Lampson1981a, Traiger1982a, Gray 1981a] are a structuring concept for operations: transactions are required to be atomic with respect to any observer. This atomicity property means that a transaction either executes in its entirety or does not execute. Thus, any side-effects of a transaction which is in progress (i.e., not complete) must not be visible, since the transaction might fail. The method described here might be viewed as a set of competing transactions, at most one of which will complete, or commit. The competing transactions must not only be isolated from external computations, but they must be isolated from each other as well.

### 3.5. Discussion

Upon receipt of a message, the predicates associated with a message (by virtue of it being sent by some process with assumptions about its existence and the existence of other processes) are checked against those of the receiver. Remember, the predicates associated with a process encapsulate the assumptions it has made about the world in order to continue executing. It has to be done this way to avoid blocking or waiting for the predicated condition (typically successful termination) to become true. The idea is that for some process these assumptions will be right. In order that the interaction with other processes leave a consistent 'world view,' we are forced to perform some special actions upon message transmission between two processes, e.g., one alternative and some server process. The idea is that the message layer, as illustrated above, is involved in process management as well. When a process receives a message which causes it to update its state (read-only messages don't change the receiver's world view, and so they can be ignored by our mechanisms but are of course used by the receiver). it must be careful. The care is taken in examining the assumptions which must be made by the receiver to continue executing. If a sending process has made an assumption which the receiver has not made, the receiver has two choices. The receiver can share the assumption with the sender, and reply accordingly. Or, the receiver can reject the assumption, and hence the message and implied updates of its state. If the sending process completes successfully, then its updates become visible when the predicates are removed from its state and from the receivers, but not before.

One issue which should be discussed is locks. If one of the alternatives successfully obtains a lock on a data object, then other alternatives will be blocked and the concurrent execution will not be transparent. Two scenarios are possible where a lock is involved. If the lock is acquired before the alternative block is entered, a copy of the state indicating successful lock acquisition will be made when the alternatives are spawned. Copies of the locked object will be made as the object is changed, and one of the set of changes will be available after the synchronization point is reached. The other scenario is lock acquisition by an alternative. In order for our mechanism to work in preserving the transparency, the lock must be accessed through another process. Then, the predicated message mechanism can be used in lock acquisition, so that two "worlds" exist, one where the lock has been acquired, and one where it has not. Otherwise, there are a variety of situations from which inconsistency or deadlock could arise.

The opportunity for a performance increase arises when the methods require different amounts of execution time.

This is illustrated in the timeline diagram of Figure 10:


Figure 10: Alternates vs. Time
It's clear that Alternate A3 is the fastest. If this had been predictable a priori, the
programmer probably wouldn't have used alternatives. We'd like to take advantage of the fact that A3 executes most quickly by selecting it and deleting the other alternatives; this is done by "sibling elimination." This is done because any computation done after A3 has terminated (the point in time indicated by the dotted line) is wasted computation.

In the next chapter, we evaluate implementation strategies, and the overhead associated with several execution environments. The careful performance measurement methodology is of particular interest, since this need be done only once for a particular execution environment. Several application areas are discussed. and experimental results demonstrate that speedup is possible with the competitive scheme on problems where no cooperative solution is possible.

## 4. Implementation, Applications and Experiments

"...the sole test of the validity of any idea is experiment." [Feynman1963a]

### 4.1. Measurement of Overhead Costs

It is informative to examine measured values of possible contributors to $\tau$ (overhead). Earlier, we described these as the cost of creating alternates, the cost of maintaining the alternates, and the cost of sibling elimination.

We have developed analysis techniques and software to evaluate the sources of overhead. The analysis techniques were applied on two workstations, the AT\&T $3 \mathrm{~B} 2 / 310^{\mathrm{TM}}$ and the Hewlett-Packard HP9000/350 ${ }^{\text {TM }}$

We begin by determining the relationships between the amount of memory in the parent's data segment, the fraction of this memory which is written by the child, and the improvement in execution time due to "copy-on-write." Since the implementation of "copy-on-write" is straightforward with modern [AT\&T1986a] memory management units (MMUs), our results for these workstations are readily generalized to other workstations.

The results show that the size of the parent's allocated memory has little direct effect on performance, because only page table entries are copied during the fork() operations. The execution time is most influenced by the amount of memory that must be copied, which can be determined from the product of memory allocated and the fraction of memory written. Thus, the worst case occurs when large address space programs update much of their memory.

To observe what occurs in practice, we measured two programs that have what are currently considered large address spaces. These programs, which we believe to be representative of the sorts of programs which use large amounts of system resources, updated less than half of the memory in their data segments.

### 4.2. Copy-on-write

The UNIX ${ }^{\circledR}$ fork() operation creates a copy of the calling process which is differentiated from its creator by the return value of fork(). The two processes have separate address spaces. Traditionally, UNIX systems copied the contents of the caller's address space to create the new process. Since the porion of the address space containing executable code was read-only, copying was not needed and an incremented reference count and text table entry sufficed [Ritchie1978a|. Clearly, the fork() operation can be expensive in system resources. Thus, some attempts were made to take advantage of special cases. An example is the 4.2 BSD [Joy1982a] vfork() call, which does not make a copy of the address space for the new process but instead allows it to share the address space with its creator. The creator is not runnable until the new process has replaced its image via an $\operatorname{exec}()$ operation. The $\operatorname{exec}()$ operation replaces the caller's image with an image derived from the contents of the named executable file. It is common for the operation which immediately follows a fork() operation (after some descriptor manipulation) to be an $\operatorname{exec}()$ operation. In particular, this happens frequently in the shell [Bourne1978a], which is the main user interface to UNIX. Thus, vfork(), in not copying, avoids unneeded work. However, the shared, not copied, address spaces force the programmer to be very aware of the differences between fork() and vfork().

Another approach is to alter the implementation of fork() to take advantage of favorable circumstances such as the shell's usage. This change should be made transparent to the application. The alteration is done with a so-called "copy-on-write" fork(), where portions of addressable memory are shared until they are changed. Similar memory management is done in TENEX [Bobrowl972a] and more recently, Mach [Young1987a]. Each process has a page table which maps its virtual addresses to physical addresses: when the fork() operation is performed, the new process has a new page table created in which each entry is marked with a "copy-on-write" flag; this is also done for the caller's address space. When the contents of memory are to be updated, the flag is checked. If it is set, a new page is allocated, the data from the old page copied. the update is made on the new page. and the "copy-on-write' flag is cleared for the new page. Thus, unexpected changes to shared state do not occur, as independent copies are created "on demand." This is effective in the special case of the shell, where almost no copying has to be done before an exec() replaces the address space. A thorough
description of the mechanism as implemented in UnIX is given by Bach [Bach1986a].

### 4.2.1. Motivation

In section 4.3, we present results from a paper [Smith1989a] where we discuss an implementation of a mechanism to fork() a process on a remote workstation; the major cost in execution time is incurred by data copying. Thus, we were interested in reducing the amount of copying, especially that which takes place over a communications channel. One strategy which we devised (assuming either homogeneous software configurations on the workstations or NFS-available [Sandberg1985a] binaries) was to have program images available on the remote system and send only the changes [Maguire,Jr.1988a] which have been made to the address space, i.e., those which would be copied by a "copy-on-write" scheme. To understand the engineering tradeoffs, we examined the local case in some detail.

The arguments presented for "copy-on-write" have so far been qualitative; we felt that detailed quantitative data were necessary.

### 4.2.2. Data Acquisition

There are two parameters of interest, i.e., the size of the storage to be "copied" in the new process and the fraction (between 0.0 and 1.0 ) of memory references which are writes. The number of times each parameter was exercised was also made variable, to remove various small-sample artifacts that can occur. Such artifacts are illustrated by the plots for small sample sizes in the copy-on-write fork() measurements. The desired data were gathered with the C program presented as Appendix I, do fork.c. A script was written in order to drive the do fork() program with various values: the values used for the measurements described here were gathered with this shell script:

```
if [ ! -f do_fork ]
then
    echo "Making do_fork."
    make do_fork
Ei
if [ : -f do_fork ]
then
    echo "No do_fork. Exiting."
    exit 1
fi
echo "size do_fork:"
size do_fork
fo: forks in 0 l 3 10 32 100 316 1000
do
    Eor heap_size in 0 1000 3162 10000 31622 100000 316228
    do
        for write_frac in 0.0 0.1 0.3 0.5 0.7 0.91.0
        do
            echo "time do_fork Sforks Sheap_size $write_frac"
            time do_fork $forks Sheap_size Sw:ite_frac
        done
    done
done
```

The script first ensures that an executable do fork binary is available, attempting to make one if not. Once do fork is available, it is invoked in the innermost of three nested loops, which vary its parameters controlling the number of fork() operations to be executed, the size of the heap to allocate, and the fraction of the allocated heap which is to be written to. Before each invocation, a message is written with echo, stating what the invocation parameters of do fork are.

Data sets for analysis by S [Becker1984a] are then created using the shell script, by, e.g. for the $3 B 2$.

```
seript 2>8: |
    grep "nreal" |
    cut -f2 |\
    awk '{ i=index(50,m); m=subscr($0,1,i-1);\
    s=scibst=(50,i+1, length($0)-i-1); s=60*m+s;\
    print s)' > real.jB2
```

and reading the list of numbers into an $S$ vector. The following data sets were extracted from the script output:
number: The number of times an invocation of do fork was to create a child process. The values $0,1,3,10,32,100,316$ and 1000 ( 0 plus powers of $\operatorname{sqrt}(10)$ were selected to make both order of magnitude induced effects (as we are changing by orders of magnitude) and implementation artifacts (because we start at small values, e.g., 0 and 1) visible.
mem: The number of bytes allocated to the process's heap, via malloc(). The values $0 ; 1,000 ; 3,162 ; 10,000 ; 31,622 ; 100,000$ and 316,228 were chosen for both artifact and order of magnitude visibility, as discussed previously; the extra factor of 1000 (over the values of number) is to compensate for the page size, since otherwise it would require (for a 2 K page ${ }^{9}$ ) 8 values before we accessed a page other than the first one. Clearly, there is no practical difference between 316,228 and 310 K : it is merely aesthetically appealing to use the correct digits.
frac: $\quad$ The fraction of memory which is to be written (we write one byte per page in order that the memory access loop not contribute to the response time beyond causing faults). The interesting boundary values of 0.0 and 1.0 were chosen, as well as the values $0.1,0.3,0.5,0.7$, and 0.9 , which were chosen for their coverage of the input domain.
real: The real time, in seconds, printed by an invocation of "time do_fork" with the parameters as set in the other vectors.
user: Likewise for user time.
sys: Likewise for system time.

### 4.2.3. Data Analysis

Given the data discussed in the previous section, we wish to analyze the data in order that we can qualitatively discuss the effects of "copy-on-write" page management on response time. One difficulty is that by our experimental design, the measured response time is a function of not one, but three quantities, number, mem, and frac. There are

[^6]two obvious hypotheses which we can propose for our analysis to refute or verify. First, that the response time increases as the size of the data segment increases, for a fixed fraction of write references. Second, that the response time increases as the fraction of write references increases, for a fixed data segment size. Figure 11 shows


Figure 11: Effect of fraction of memory written

$$
\text { (number }=1000 \text { ) }
$$

mem plotted on the $x$ axis against real on the $y$ axis for an AT\&T 3B2/310 with 2 megabytes of memory (of which 1.2 megabytes are available to user processes), a 30 megabyte hard disk, and running UNIX System V, Release 3.0, Version 2. All times are given in units of seconds. We have fixed the value of number to be 1000 to remove artifacts. The dependent variable, plotted vertically, is the real time. in seconds. The independent
variable, the size of the data memory in bytes, is on the horizontal axis. Regression lines are drawn through the plotted points corresponding to frac values of $0.1,0.5$, and 0.9. These regression lines have equations $y=1.709 e-4 \cdot x+31.4, y=7.670 e-4 \cdot x+30.5$, and $y=1.349 e-3 \cdot x+30.7$ for the respective frac values. Thus, with these equations, we could estimate that a process with a 1 megabyte data segment which writes into half of that segment would take about 800 ( $797.5=7.67 e-4 \cdot 1.0 e 6+30.5$ ) seconds of real time to perform 1000 fork() operations. The lines fit the plotted points well, indicating that the relationship is close to linear.

The same data are plotted for a Hewlett-Packard HP9000/350 with 8 megabytes of main memory and an HP7945 70 megabyte hard disk, running HP-UX ${ }^{\text {TM }} 6.0$ (same units. restrictions, and axis markings) in Figure 12.


Figure 12: Effect of write fraction (HP-UX)

$$
\text { (number }=1000 \text { ) }
$$

The equations for the lines with frac set to $0.1,0.5$, and 0.9 are $y=2.952 e-5 \cdot x+12.7$. $y=1.264 e-4 \cdot x+12.4$, and $y=2.124 e-4 \cdot x+12.2$, respectively. The effect of the faster processor in the HP9000/350) is clear from the extent of the $y$ axis in this figure versus that of the previous one. The important parameter in comparing processor speeds under this workload is memory-copying speed. To measure this, we wrote a short $C$ program which took the number of bytes to copy as an argument, the relevant fragment of which is:

```
p = malloc( size );
Ere_page( p ):
clock = times( 6cbl);
memcpy( p, g, size );
clock = :imes( &tb2 ) - clock;
```

For size set to 316,228 and a page size of 2 K bytes ( 4 K on the HP9000) we measured 0.40 seconds of real time, 0.39 seconds of user time, and 0.00 seconds of system time on the $3 B 2 / 310$. The values were 0.06 seconds of real time, 0.06 seconds of user time, and 0.00 seconds of system time on the HP9000/350. These values held true through several trials, and show that for memory-copying the HP is about (to the limited accuracy of the measurements) 6.7 times faster than the 3 B 2 . They also provide an upper bound on the memory copy rate which can be used to evaluate overhead incurred by page management operations. For the HP, we get $5 \mathrm{M}(5,270.467=316,228 / 0.06)$ bytes per second, or about $1,300(1,286=5,270,467 / 4,096) 4 \mathrm{~K}$ pages ${ }^{10}$ per second. For the 3 B 2 , we get 0.8 M ( $810,841=316.228 / 0.39$ ) bytes per second or about $400(396=810,841 / 2,048) 2 \mathrm{~K}$ pages per second.

We can use the regression lines we have presented for further analysis. The $y$ intercept (about 31 seconds for the 3B2/310 and 12 seconds for the HP9000/350) should represent the time required for 1000 forks which allocate 0 bytes of memory; examination of the script output confirms that this figure is accurate. Since do fork is written to be compact (no standard I/O, etc.) this should accurately indicate the cost of performing a fork when divided by the number of operations performed. Thus, using the computed intercepts we have given for number set to 1000 , the average $3 \mathrm{~B} 2 / 310$ fork requires about $31(=(31.4+30.5+30.7) /(3 \cdot 1000))$ milliseconds of real time. For a fixed number of fork() operations the $y$-intercept is not nearly as interesting as the slope of the line. We should note that in reality, the function is not a line, as the quantization of bytes into page size quantities forces a staircase function. However, for purposes of analysis we can assume that a linear function exists. The slope of the line for some known value of frac gives the relationship between changes in real caused by changes in mem. Hence, we can use the slope of the regression line to estimate the rate at which page faults are

[^7]serviced. Mem•frac gives a fixed amount of memory, with which we use the equation of the regression line to compute a real time estimate. Then, the observed page fault service rate can be computed with the simple formula $\frac{\text { mem•frac }}{\text { real time }}$. The slope of the line can be used to compute the service rate directly, for a known value of frac and number; this rate is given by $\frac{\text { number•frac }}{\text { slope }}$, which calculates a value in units of bytes per second. For the 3B2, these values are 585,$138 ; 651,890$; and 667,161 ( 286,319 , and 3262 K pages/second, respectively) for the three values of frac plotted. The corresponding values for the HP9000 are $3.387,534 ; 3,955,696$; and $4,237,288$ bytes per second ( 827. 965 , and $1,0344 \mathrm{~K}$ pages/second ${ }^{11}$, respectively). Using the best observed page fault service rates for each processor, we calculate the ratio of the page fault service rate and the time for memory-copying, which is $0.823(=667,161 / 810,841)$ for the 3 B 2 , and 0.804 $(=4,237.288 / 5.270 .466)$ for the HP. Values for the ratio can range between 0 and 1 ; the best case is a value near 1 , as this indicates that the virtual memory management incurs little overhead. We can estimate this overhead using the information we have. Using $\tau()$ to measure time, we know that
\[

$$
\begin{aligned}
\tau(\text { fork })= & \tau(\text { copy one page }) \cdot \text { frac } \# \text { pages }+ \\
& \tau(\text { overhead for page table entry }) \nexists \text { pages }+ \\
& \tau(\text { overhead to create new process })
\end{aligned}
$$
\]

Now, $\tau$ (copy one page) is really a function of the hardware components (e.g., bus, processor, memory) comprising a system, and we've shown how it can be gathered with a small auxiliary program. But from our numbers and analysis we can get $\tau$ (overhead for page table entry) and $\tau$ (overhead to create new process). Thus, for any given fork operation, the time required is completely parameterized by $\tau$ (copy one page), $\tau$ (overhead for page table entry), $\tau$ (overhead to create new process), frac, and mem (\# pages). The key is that the first three are determined by the system characteristics, and they can thus be precomputed; the application-dependent influences are completely encapsulated in the latter two parameters. Thus, an application can be characterized on a given system by its size in pages and the fraction of those pages which are written to.

[^8]
### 4.2.4. Relationships

The shapes of the plots we generated are similar for both processors. The HP9000 plots' time values (the y axis) are scaled differently because the HP9000 is significantly faster than the 3 B 2 . We'll use the 3 B 2 to illustrate the analysis in the remaining figures.

One failing of the $x-y$ plots is that there are two independent variables. The perspective plot shows that the real time increases as a product of mem and frac; the maximum value is 505.82 seconds, for number at 1,000 ; mem at 316,228 ; and frac at 1.0 . Figure 13 shows a 3-D perspective plot of real ( $z$-axis), mem ( $x$-axis), and frac ( $y$-axis). The point $(316,228 ; 1.0 ; 505.82)$ is the furthest, highest point on the graph. Thus mem is increasing from our left to our right (it's an exponential curve as the data values are chosen to increase exponentially), and frac is increasing from our right to our left, moving away from us. Figure 13 would then be overlaid cross-sections taken from the perspective plot by intersecting a series of $y$-planes with it. Of course, not all the data of figure 13 are available due to hidden line elimination.


Figure 13: Perspective plot, mem vs. frac vs. real
We've limited the data shown in Figure 13 to that gathered with number set to 1000 . This was done after analysis of the raw data showed two things which limited the value of the data gathered for a small number of fork operations. First, there was little opportunity for the data to become evident against the overhead of executing the parent program. This could, of course, have been removed by calling times() from inside do fork, but given our strong preference for the shell as a measurement apparatus, this was not done. Second, the timing data were apparently overwhelmed by other sampling noise. such as that caused by various background processes and network daemons (although the processors used for these tests were otherwise idle). These other processes were not shut down due to the effect on our working environment.

If we plot a 3-D perspective plot with parameters as before, except that number is 1, we get Figure 14, which demonstrates what sort of artifacts, or "noise" can arise due to inadequate sample size.


Figure 14: Perspective plot, mem vs. frac vs. real

The deduction one can make from examining this plot is that there is no obvious relationship between the input values and the response time output. The errors and spurious values have dominated the measurements to the extent that visual tools such as graphs are no longer useful. It is doubtful whether any tools are useful under these circumstances, and the lesson is clear: the analysis must be aware of the sources of error. and the measurements must be made in such a way as to minimize these sources. In our case, the minimization was achieved by using an adequate sample size.

The question might be raised as to why real time is used, not sys. Philosophically, the real time is what is most relevant to an observer. Scientifically, analysis shows that for number large, real is less than 20 percent greater than sys, and that they are closely correlated. This is illustrated in figure 15, where the x axis has values of number, and the $y$ axis is the value $\frac{\text { real-sys }}{\text { sys }}$.


Figure 15: Relation between real and sys
The plot shows that the relationship between real and sys is not good for small values of number; they differ by almost a factor of 20 . However. things improve as number gets larger: a detailed graph is provided in figure 16 by restricting number to values of 100 or more. It's clear from this illustration that for number at 1000 , sys and real are
reasonably good approximations of each other.


Figure 16: Relation between real and sys
Incidentally, we should note that for small values of number (e.g., 1), sys is subject to the same noise problem that real suffers from; this is easily observed with another perspective plot, which we will not present due to space considerations.

### 4.2.5. Write Fraction for Real Programs

In the last section, we saw that the factors mem and frac influenced the real time requirements of our test program. The biggest savings for the "copy-on-write" scheme would come from programs with large address spaces which updated a small fraction of their
data before exiting or exec()ing a new binary. As discussed before, this works well for the shell, but the shell typically uses little of its data segment. While it may expand the address space as necessary to store new variables or metacharacter expansions, this does not account for many pages. We thus sought programs with large address spaces, to see what effect the "copy-on-write" scheme would have.

As they had the largest address spaces (of programs in common use in our department), we set out to take some measurements of the memory utilization of two symbolic interpreters. We chose 4.2BSD's [BSD1982a] Franz Lisp (Opus 38.92), as it is widely available. Another less detailed set of measurements was taken using the GNU Emacs [Stallman1986a] LISP interpreter, which is also widely available. These measurements were taken on a DEC ${ }^{\text {TM }}$ VAX-11/750, because both pieces of software were available there (Franz Lisp is not available on our HPs and 3B2s, although GNU Emacs is). Since we are measuring data segment utilization, and the machines discussed in this paper all have 32 bit architectures, the measurement results should be portable. This is particularly true because we use relative measures, such as the fraction of the data segment which has changed. While a particular architecture may have a less efficient representation of the data, this should not change the fraction of the data altered by the program significantly.

### 4.2.5.1. Franz Lisp

Our first exercise was choosing a computationally intensive process so that we could gather some statistics on the sort of processes which one would want to improve the performance of [Leland1986a] : that is, those that consume many resources. Experience with an ABSTRIPS [Sacerdotil974a] implementation led us to use this system to gather statistics. ABSTRIPS is a "planning'" system which works by constructing increasingly detailed series of actions at decreasing levels ("criticality levels") of abstraction. There are primitives defined (in predicate logic) for each level of abstraction: as the levels are traversed, we gradually "flesh out" the details of a plan for achieving the goal. ABSTRIPS relies heavily on the use of a theorem prover: hence, it is representative of much current AI computation. An example of its output is given in Figure 17.

```
Eranz Lisp, Opus 38.92
-> [load abstrips.lsp]
\tau
-> criticality level: 4
skeletor plan : ((goal c))
criticality level: 3
skeleton plan : ({get-slippers d)
(give-slippers DOG ME) (goal c))
criticality level: 2
skelecon plan : ((gothrudoor c b DOG)
(gothrudoor b a DOG)
(gothrucoor a d pOG)
(get-slippers d)
(gothrucoor d a DOG)
(gotirudoor a b DOG)
(gothrudoor b c DOG)
(give-slippers DOG ME) (goal c))
criticality level: !
skeleton plan : ((gothrucoor c b DOG)
(gothrudoor b a DOG)
(pushopen a d)
(gothrucoor a d DOG)
(get-slippers d)
(gothrudoor d a DOG)
(gothrucioor a b DOG)
(gothrudoor b e DOG)
(give-slippers DOG ME)
(goal c))
((gothrvioor c b DOG)
(gotirudoo: b a DOG)
(pushoper a d)
(gotn=udoo: a a DOG)
(get-slippers d)
(gothrudoor d a DOG)
(gothrudeor a b DOG)
(gothrudcor b c DOG)
(give-slippers DOG ME) (goal c))
->
```

Figure 17: ABSTRIPS Output

The problem in our example was to have a dog fetch your slippers from another room. This problem takes about 15 minutes to plan on a VAX-11/750; the implementation makes heavy use of recursion and maintains several large lists.

The size of the Franz executable (from the UNIX size command) is 139,264 (text) $+511,488$ (data). The data on memory usage was obtained by using the UNIX system's ability to create a core dump of a process's address space; since the text segment is read-only, only the data and stack segments are dumped. Sending the SIGQUIT signal to a process causes a core dump; this was done at the following points in the execution of
the ABSTRIPS planner.

1. When the LISP interpreter was started. This gives us a baseline value, with no program loaded and no code executed. The core dump occupied 528,384 bytes.
2. Immediately after ABSTRIPS was loaded. This tells us how much of the address space change is due to storage of the ABSTRIPS program. The core dump occupied 556,032 bytes; a bytewise comparison with the previous dump showed that 56,937 bytes had changed.
3. Immediately after ABSTRIPS execution is terminated. This tells us how much of the address space has changed during execution. The core dump occupied 613,376 bytes, and differed from the previous dump at 77,910 bytes. The difference between this dump and the first dump was a total of 123.942 bytes changed. No garbage collection was announced.

An important issue is the locality of reference; our measurement programs for the "copy-on-write" fork performance showed that we could write every page by writing one byte on each page. The byte comparison routine delivers addresses where it found differences between two files; the difference in bytes could then be measured by piping the output to "wc -1;" if we divide each address by the pagesize ( 512 on the VAX) and pass the results to "uniq । wc -1 ," we can find the number of pages that have changed: in this case 270 of the $1.198(=613,376 / 512)$ pages changed, for a write fraction of 0.23 .

### 4.2.5.2. GNU Emacs

GNU Emacs provides a facility to dump the currently executing image into an executable file. When this file is executed, the state of the Emacs interpreter is restored to the state it had when the (dump-emacs) was invoked. We took the following measurements on the VAX/l1-750 ${ }^{12}$. The size of the GNU Emacs editor we measured was 437,248 (text) +208.896 (data), determined with size. We sought an example program

[^9]which had the sort of behavior (computation-oriented) that we desired. We based our desire for computationally-intensive examples on the observation that as heavy resource users, these programs would demonstrate the greatest effects from an optimization. GNU Emacs provides a library of LISP code; one routine provides a graphic solution of the classic "Towers of Hanoi'" problem. We ran the GNU LISP interpreter on the following input:

```
(dump-emacs "pre-hanol" "/usr/local/emacs")
(hanol 10)
(dump-emacs "post-hanoi" "/usr/local/emacs")
```

(As might be expected, this requires patience at 9600 bits per second!) The interpreter emitted several messages to the effect that it was performing garbage collection.

At the completion of the computation, we performed a bytewise comparison on the two dump files:

```
$ ls -l post-hanoi pre-hanoi
-rwxr-xr-x l jms pha ( 851968 Oc= 27 08:25 pos:-nano:
-rwxr-xr-x 1 jms phd 737280 Oct \(2616: 01\) pre-hanoi
```

which showed that 183,312 bytes had changed, which for the computed data segment size of $414.720(=851,968-437,248)$ is slightly less than thirty-five percent of the dump: that is, almost the same percentage we had observed with Franz Lisp and ABSTRIPS. Several times during the computation and in the dump-emacs function the garbage collector was run. Thus the amount which appears to have changed may include parts which did not change but were relocated and thus appear to have changed. It also compacted storage which appeared to be changed (since newly-allocated storage is considered changed from the previous non-allocated storage). The important point is that these changes would be seen by a page-management mechanism in either case.

### 4.2.6. Conclusions about copy-on-write

"Copy-on-write" paging strategies for address space inheritance have been shown to be effective in reducing the real time required to perform UNIX fork() operations. This qualitative assessment is based on the quantitative data we gathered and analyzed. For large processes, the time required is proportional to the fraction of write references, so
that a child process which updates half $(0.5)$ of its address space will spend half the time doing copying that a child process which updates all (1.0) of its address space will. For a pair of interpreters with large address spaces, we showed that the portion of the address space changed from process startup until process termination was small, typically less than 0.5. These measurements concur with those of Zayas [Zayas 1987a], who measured program behavior in an Accent environment, and confirm the desirable properties observed of a similar scheme for fast state transfers to remote systems in the V [Theimer 1985a] system ${ }^{13}$.

Thus, if these interpreters or programs which behave similarly were to fork() child processes which executed tasks similar to those described, a reduction of 50 percent or more of the system time devoted to copying data might be achieved. This confirms that the scheme for remote fork() using 'lazy' copying has considerable merit.

This reduction in copying also reduces the amount of swap space required, reduces the amount of time spent swapping, increases the number of processes which can be run without paging, and decreases the cost of context switches (where the cost of paging out the written pages and the paging in of pages which are only read and have not been modified is included). Thus the advantages of the text table are extended to unmodified pages (or viewed another way UNIX gains via "copy-on-write" the ability to eliminate the text table and improved fork() performance). With respect to these page management strategies, note that TENEX [Bobrow1972a] had these advantages ten years earlier and needed neither a distinguished text table nor the confusion of two varieties of fork().

The cost figures we present should be representative of a shared memory configuration of equivalent processor technology. The fact that we have provided a methodology for gathering such measurements ensures that the techniques are portable, even if the measurements themselves are not.

[^10]
### 4.3. Remote fork()

There is more overhead associated with the distributed case, due to the increased costs of copying. We describe a method of implementing a distributed fork() operation in Smith and Ioannidis [Smith1989a]. A process successfully executing a fork() operation generates two copies of its address space; these are often distinguished as parent and child by the return value of the fork() call. If the child process continues its execution with the containing address space located on a processor different from the parent process, we have achieved a "remote fork."

By distinguishing between the state saving activity and the state transfer activity, we were able to measure and refine the performance of each activity independently. Once the design and initial implementation were complete, we analyzed the performance, and reimplemented pieces of the system (on several different machine architectures) to improve the response time. This improvement was dramatic: from about 7 seconds of real time on the HP9000 and the 3B2, to less than 1 second on the HPs and Suns using NFS. The major savings came from reducing the execution time devoted to copying state information from point to point.

### 4.3.1. Further process migration ideas

As the major cost of process migration is copying [Zayas1987a], attempts have been made to reduce this overhead. More sophisticated migration schemes, using "ondemand" state management techniques have been constructed [Theimer1985a]. Most programs exhibit locality of reference; in particular symbolic computations which use large amounts of system resources [Smith1988a]. These computations are representative of those that present the greatest opportunity [Leland1986a] for applying process migration to load-balancing across multiple processors. In early computer systems, the notion of a relocatable (position-independent) module of executable code improved the degree of multiprocessing. Relocatable code allowed multiprocessing systems to exploit available memory more effectively. This exploitation increased the degree of multiprogramming, achieving an increase in throughput. Likewise, relocatable processes give operating systems the capability to exploit the availability of multiple processors. This exploitation can result in improved throughput, improved response time, or both. What makes a process relocatable is a description of its state which can be used to continue the
computation elsewhere. This description is often achieved via a copy, thus forcing the expensive copying operations. Measured in execution time, the cost of copying can be reduced by faster networks or data transfer software. Our implementation of rfork() shows that this approach can be effective. However, for any data transfer scheme there is a limit on its performance. This limit is imposed by the combination of the transfer hardware and the software used to access it. Also, although the mapping between data volume and execution time may increase in steps rather than smoothly, more data implies more time. Thus, for a limited data transfer rate, we should seek methods of reducing data copying.

Several ideas suggest themselves:

1. Be lazy. This describes the approaches of Zayas and Theimer; they took advantage of locality to reduce copying or its effect on program execution speeds. "Demandpaging" has illustrated the effectiveness of lazy copying in computer systems.
2. Encode Symbolically. The state can be encoded symbolically. Symbolic encoding can be achieved by use of interpreters or compiled interpretative languages. When the execution of a process is halted, the program and the state of the interpreter are re-represented in a symbolic form. This symbolic form is then passed to another interpreter, which can restart the program from the "symbolic checkpoint." This representation may also be more compact than the running program. Note that such machine- independent representations offer the only hope for true "heterogeneous" process migration.
3. Compress. Another possibility for re-representation is compression, where the encoded state is produced by applying a data compression algorithm [Lelewer 1987a] to the saved state. When the process is restarted, the state is uncompressed and used to recreate the running process.

Schemes 2 and 3 pay a computational cost in encoding and decoding. Thus, there is a performance tradeoff between the cost of copying and the cost of encoding. For straightforward compression techniques, the cost of encoding can be calculated using the size of the input state. For "symbolic checkpoints." cost estimates are less predicatable from the size of the process, but can easily be made using the symbol table which is necessary for the encoding to take place. Copying costs can be estimated using the techniques we have described in this thesis, so that decisions can be made by a dynamic migration
manager. These decisions, about whether to migrate or not, can help in load-balancing.

### 4.4. Disk Response Time

Referenced pages will not always be available in memory, thus disk access (or network access, to be discussed later) may be needed for a reference to be satisfied. One difficulty with simulation, or with "toy" implementations is that the quantitative data necessary for accurate response time evaluation are not used in the simulation. To make our results more accessible to practitioners, we have used data on disk response times [Johnson 1987a] which was gathered on a running UNIX system, operating the Digital Equipment Corporation ${ }^{\text {TM }}$ (DEC) RA81 drives whose characteristics are summarized in the table,

| DEC RA81 Disk Drive Characteristics |  |
| :--- | :--- |
| Cylinders | 2516 |
| Transfer Rate | 2.2 megabytes/second |
| Rotational Speed | 3600 rpm |
| Average Rotational Latency | 8.33 milliseconds |
| Head Switch Latency | 6 milliseconds |
| Average Seek | 28 milliseconds |
| One Cylinder Seek | 7 milliseconds |
| Maximum Seek | 50 milliseconds |

attached to a DEC UDA50 controller. Under the load conditions described by Johnson, Smith, and Wilson as "normal," the average number of ticks per response was 2.1, thus the average response time using a tick of $1 / 60$ second was $\$ 2.1 \operatorname{cdot}(1 / 60) \$$. or 35 milliseconds. From their data analysis, it can be seen that this number is constant across drives, interfaces, transfer sizes, and transfer start time. It varies between reads and writes, with writes taking longer, and it varies with the location on the drive. Writes take longer because they are bunched together at times when UNIX flushes its buffer cache,
thereby lengthening the time spent in the drive's request queue. The response time varies with the location on the drive due to hot-spots found at the $i$-lists of file systems found on the drive. Both of these variations are due to the nature of the UNIX file system, and thus are not significant for our discussion.

### 4.5. Network Response Time

To gather information on the response time for network page requests, the program ("netrand.c") was written. Since it is short, a source listing is provided as Appendix II. The experiment was to operate the program on two files, one on local disk, and the other on an NFS-mounted file system. The system employed for testing was a HewlettPackard HP9000/350 with 8 megabytes of main memory and an HP7945 70 megabyte hard disk, running HP-UX ${ }^{\text {TM }} 6.0$.

The UNIX buffer cache mechanism was frustrated by copying a large ( 2 megabyte) file previous to running the tests. Three executions of the test were run on each file; for the purposes of benchmarking we ran the program with an argument of 400 blocks. Since the randomization frustrates the buffer cache mechanism to some degree, the buffer cache has a significant effect by the third execution. The results of running the program on the local file were $10.68,10.24$, and 9.74 seconds of elapsed time for the three runs. For the remote file (the server is a machine of the same type, accessed over a lightly-loaded 10Mbit Ethernet) the results were $15.42,15.02$, and 13.32 seconds of elapsed time for the three runs. The improvement in performance seen as the runs progressed is due to the success of the UNIX buffer cache in retaining recently-read blocks. The network access seems to indicate a penalty of about a factor of 1.5 for network access of pages, which is encouraging. The time per page (using the worst case, where the cache is flushed) gives us a per-page time of $\frac{15.42}{400}$ seconds, or about 39 milliseconds per page. This works out to about 26 pages per second for 4 K pages. This cost compares unfavorably with the cost of a local page copy, which is about 1 millisecond on this machine.

### 4.6. Sibling Elimination

A remaining source of overhead is the cost of eliminating unwanted computations; we have speculated, selected, and now we must eliminate. What will this cost us in time? A program, do_elim.c, is presented in Appendix XII. It was constructed to address the various factors which might influence the time involved in eliminating processes. While the details of the construction are evident in the program, some discussion of the goals and philosophy is in order. The goal was to give us some idea of the cost in execution time attributable to sibling elimination which would not be paid in the case where the fastest alternative was selected "at random." This helps us to estimate (overhead).

The philosophy was to use a highly-tuned existing operating system, in this case Hewlett-Packard's HP-UX 6.0 implementation of UNIX System V. The idea behind that is that we can experiment with various tunable parameters of interest, while taking account of details which a simulation might otherwise ignore. In addition, the numbers should be close, certainly less than an order of magnitude away, from values gathered in an implementation. We gathered data for real, user, and system times, obtained from the UNIX times() system call. The UNIX timing facility is not particularly accurate, but we applied techniques which should, in the average case, remove much of the error content from the data.

The basic design of the experiment was to create some number of processes, have them each do something after spawning, and then eliminate them. Only the elimination portion of the experiment is bracketed with timing requests. Elimination is done by means of the UNIX kill() system call, which sends a small (less than a byte) amount of data from a sender to a receiver in the form of a signal. The signal causes the receiver to execute some signal-handling action, which can include terminating, ignoring the signal, or handling the signal in some specialized fashion. The processes in our experiment were set up in a fashion that ensured process termination upon receipt of a signal. Bach [Bach1986a] provides discussion of these mechanisms in such detail that further detail here is unnecessary.

The factors we chose to examine were:
Groups: UNIX provides a facility to send a signal to (1) a single process, or (2) a group of processes. selectable by virtue of either the owner's user id or the process group. Use of the process group signaling facility allowed us to emulate the
effects of a multicast message-sending facility, so that we could evaluate the effects of point-to-point elimination versus multicast.

Files: Among the items of system-maintained state associated with a UNIX process are a number of open files and their associated data structures. Upon exiting, these files must be closed, which involves deallocating them, flushing some buffers, etc. Thus, varying the open files should vary the system state maintained by a process which is to be deleted.

Size: As we saw the effect of virtual memory copying on response time earlier in the thesis, it may have some impact on process deletion, in particular the deallocation of allocated pages and page descriptors. Varying the size gives the best estimate of the impact on response time, as we saw earlier.

Work: The spawned subprocesses either loop sleep()ing, or they alternate between sleeping and iterating in an empty for() loop. The latter case tries to estimate the effect of the UNIX process scheduling policy on processes; in particular, if the wait for a working process would be longer than the wait for a sleeping process.

Asynch: After signaling the spawned subprocesses with kill(). we can either wait for them to complete inside the timing block, or we can exit the timing block, to emulate a situation where we don't wait for the processes to die, but just signal that they should die, and then continue. The waiting case is synchronous, and the other is asynchronous. With a large degree of parallelism present, asynchronous elimination seemed a better policy, as the hardware parallelism could be employed to speed up the elimination.

Dirty: As with the size of allocated memory, we earlier discovered that the write fraction had a determining effect on response time when coupled with memory

Procs: This varies the number of spawned sub-processes. The idea is to see how the increase in sibling elimination costs varies with both the policy employed for elimination, and the number of processes the policy is applied to.

The measurement program, do_elim.c, was run using the shell script in Appendix XII. The numbers were extracted, and entered into an $S$ [Becker1984a] dataset for analysis. The statistics system was used as a tool for data analysis, in the sense that various parameters were related to the real and sys times they exhibited, to determine the importance
of their effect on the response time. The next two sections present the data in graphical form, with a short discussion and explanation following each figure. The first of these sections presents the real time measured, and the second section shows the sys time measured. The point of real time is that it's the best measure of response time; sys time tells us how much effort the system is exerting on our behalf. In response to well-known problems with the granularity of the UNIX clock scheme, we ran the creation- timed deletion- cleanup loop 100 times (by setting REP_COUNT to 100 in do_elim.c ) for each variety of do_elim invocation. This has two effects. First, it gives us enough magnitude in the data so that comparisons can be made between the different test inputs. Second, the timing errors in such a loop can be lesser or greater than the correct time. With repetitions, the idea is that the errors will in effect cancel. so that the repetitions will in the average case purify the data, much like oversampling techniques in audio. Of course, in the worst case, correlated or systematic errors, there is the possibility that the results will be all noise. This possibility is refuted later.

### 4.6.1. Real Time

To measure the effects of the different variables on the execution time ( real time. in UNIX jargon) we plotted the real time values as a function of the variable values. The most useful technique we have found for representing this data is the boxplot; it provides much more information than an $\mathbf{X}-\mathbf{Y}$ plot for this type of data. For a given $\mathbf{x}$ value, the box defines the middle 50 percent of the data, the horizontal line inside the box is the median, and the bar at the end of the dashed line marks the nearest value not beyond some standard range (in this case, $1.5 \cdot$ (inter-quartile range) ) from the quartiles. Points outside these ranges ("outliers") are shown individually. Details of boxplot presentation can be found in Chambers, et al. [Chambers1983a]

The first plot, figure 18 , indicates that the use of the process group mechanism reduces the execution time devoted to sibling elimination. While the median improvement is not great, the larger top half of the left hand box for not killing by groups indicates that for the slowest half of the times, group signaling had a more significant effect.


Figure 18: Effect of signaling process groups on real time (For 100 repetitions)

This is an intuitive result. since broadcast should be significantly cheaper in wall clock time than a serial sending of the "messages."

Figure 19 shows the effect on real time of the number of open files: it appears that the amount of system state which must be changed when these are closed on process termination is not significant in its effect on sibling elimination times.


Figure 19: Effect of number of open files on real time (For 100 repetitions)

Figure 20 shows the effect of performing the elimination synchronously (Asynch=0) versus asynchronously (Asynch=1). There appears to be no difference in the real time required. This is surprising. as intuitively, not waiting for something to happen should be faster than waiting for something to happen. There are two possibilities: (1) the effect is too insignificant to be discerned against the measurement error (a real possibility, given the granularity of the UNIX clock facility) or (2) the experimental apparatus is flawed. We address this counterintuitive result later in the thesis.


Figure 20: Effect of Asynchronous elimination on real time (For 100 repetitions)

Figure 21 indicates that the effect of dirtying pages in the child ("dirtying" is done by the child, as it would be in truly concurrent execution) is negligible when measured in real time.


Figure 21: Effect of Dirtying child pages on real time (For 100 repetitions)

Figure 22 indicates a slight effect on real time caused by the process behavior of the spawned children. The idea here was to see if the signals would take longer to propagate if the multiprocessing involved processing and not merely waiting for receipt of a signal. As one might imagine, it takes longer to eliminate busy children than it does sleeping children.


Figure 22: Effect of child work/sleep activity on real time (For 100 repetitions)

Figure 23 indicates almost no effect contributed by the process size. There is a slight difference, most evident when the least ( $\mathrm{size}=0$ ) is compared with the greatest ( size $=100.000$ ) where we see a slight increase in the max, median, and top 50 percent. The increase is so slight that it would be difficult to attribute a significant time cost to process size in sibling elimination.


Figure 23: Effect of child process size on real time (For 100 repetitions)

Figure 24 shows the effect the number of child processes (which are spawned and then eliminated) has on the measured real time. This plot indicates that the number of child processes has a significant effect on the required real time for sibling elimination, and that the cost increases in proportion to the number of children. The sample values for Procs formed an exponentially increasing sequence, giving rise to the shape of the curve drawn through the medians or the curve through the top "whiskers." These drawn curves differ slightly in their shape; 8 child processes is a jump in the value of the top whisker, where the curve through the medians is smoother. This discontinuity was not consistently reproducible. For example, real time and system time should be well-
correlated on these measurements, and yet a comparison of the graphs does not show the slight discontinuity in the curve connecting the top whiskers. Therefore it is probably measurement error; see the section below on errors in the measurements for a detailed discussion of the sources of such errors.


Figure 24: Effect of number of child processes on real time (For 100 repetitions)

Thus, the data we have gathered indicate that there are two significant variables in the real time devoted to sibling elimination. These are (1) the use of process groups in communicating message information through signals (roughly equivalent to a multicast), and (2) the number of processes which are spawned and eliminated. Surprisingly, there was little effect seen from the use of asynchronous messaging. To refine our estimates of
measurement errors, we decided that plots of the system time required for each of the variables we analyzed the real time for were necessary. These are presented in the next section.

### 4.6.2. System Time

As in the last section, the time required for a multicast sibling elimination is reduced significantly when compared to serial message-sending. In light of the intuition and previous measurement results, this helps to confirm that the results are significant (and not noise) if the experimental apparatus is correct.


Figure 25: Effect of signaling process groups on sys time (For 100 repetitions)

As before, the number of open files was examined as a contributor to the measured time values. There does not seem to be any contribution to the system time, as is evidenced by the plotted data of figure 26.


Figure 26: Effect of number of open files on system time (For 100 repetitions)

Figure 27 shows the effect on system time of making the elimination process asynchronous. As in the analysis for real time, the results are counterintuitive; that is, there seems to be no difference. This seems to make more sense in the system time setting, as the system must do approximately the same work in each case; the exception might be in table searches which could be done on the basis of process group membership, rather than multiple searches for single process identifiers; however, the effect of this would probably be lost in the noise. Likewise, there is an increased number of system calls: their contribution appears to be minimal according to the plot.


Figure 27: Effect of Asynchronous elimination on system time (For 100 repetitions)

Figure 28 shows the effect of the child process's page updating behavior on the system time; as before, there is no observable difference.


Figure 28: Effect of Dirtying child pages on sys time (For 100 repetitions)

Figure 29 shows the effect of the child process's computational behavior on the system time; there is no observable difference.


Figure 29: Effect of child work/sleep activity on system time (For 100 repetitions)

Figure 30 shows the effect of the child process's allocated memory size on the system time; there is no observable difference, not even the slight effect we observed previously for real time. This may be due to the accounting scheme used for faulting pages; if the page fault time is charged to some kernel "process' the extra costs associated with page management (including deletion of dirty pages) may not be charged to the proper entity.


Figure 30: Effect of child process size on system time (For 100 repetitions)

Figure 31 illustrates the effect the number of processes has on the system time. The increase is once again proportional to the number of processes; this increases our confidence in the behavior of the observed real time. It's interesting to observe that there is also a correlation between the number of processes and the dispersion of the system time values measured: much more so than our observations for real time. We have no explanation for this other than clock inaccuracy and the contribution of other variables. For example, the contribution of data from both values of a bimodally distributed variable such as Groups may cause an apparent dispersion. Also, since the system time values are always smaller than real time values (on a uniprocessor) the effect of clock granularity on
measurement accuracy will be more pronounced. Clock granularity as a source of error in measurement is discussed below.


Figure 31: Effect of number of child processes on system time (For 100 repetitions)

Thus, our observations of the system time correlate well with our observations of the real time. What this does is to confirm that where the data shows an influence, there exists an influence. The non-intuitive values for asynchronous elimination drove us to further examination of the data and the apparatus.

Since there was a clear correlation between the number of processes spawned and eliminated and the time required for both timing measurements, it seemed worthwhile to examine the most expensive case separately, to remove extraneous influences form the
data. In the next section, we examine the influences of the variables for the special case of 16 spawned and eliminated child processes.

### 4.6.3. Real Time, 16 Procs only

As in the less restricted set of measurements, the use of process groups seems to provide a significant improvement in real time performance. Figure 32 illustrates the magnitude of this improvement.


Figure 32: Effect of signaling process groups on real time. 16 procs (For 100 repetitions)

Figure 33 indicates a small contribution from the number of open files; this makes sense, as for the larger number of processes involved, the system has significantly more state to
manage. However, the contribution still appears slight: it is slight enough to be accounted for by errors, as discussed below as "Sources of Errors in Measurements."


Figure 33: Effect of number of open files on real time, 16 procs (For 100 repetitions)

Surprisingly, asynchronous elimination still does not have any effect on the real time, as illustrated in Figure 34. These results inspire a certain amount of suspicion, since the synchronous version must unfailingly wait() for each of 16 children to complete, while the asynchronous case (at least according to the code) returns without wait() -ing: collection is done outside of the timing loop.


Figure 34: Effect of Asynchronous elimination on real time, 16 procs (For 100 repetitions)

Figure 35 indicates that dirtying pages has a slight effect on the real time: we doubt that the effect is significant.


Figure 35: Effect of Dirtying child pages on real time, 16 procs (For 100 repetitions)

Child process computational activity indicates a minor influence on real time, as shown in figure 36. Work=1, that is. the process is computing rather than sleeping, seems to cause a greater amount of dispersion as well as a slight increase in the median value: little significance can be attached to the observation.


Figure 36: Effect of child work/sleep activity on real time, 16 procs (For 100 repetitions)

Process size has a greater effect on the measured real time in the 16 process case than we observed for all the values of Proc lumped together. This effect is demonstrated by figure 37. The fact that the effect begins to be significant only on the larger sizes, not linearly or proportionally, makes us suspicious that the cause is some system artifact such as a limitation on the system memory size forcing a swapout of one or more of the processes under test.


Figure 37: Effect of child process size on real time, 16 procs
(For 100 repetitions)
What we can infer from the data for the special case of 16 spawned and terminated subprocesses is that the observed influences on real time are pretty much the same. The effect of process groups is more dramatic, and there began to be an influence exerted by the size of the processes. There was also an observed, but likely to be insignificant, increase in costs associated with the number of open files at elimination time. What the 16 process case should illuminate is the increasing effects of various variables with a change of scale.

We also thought it provident to examine the case of 16 processors for system time as well, to isolate any influences which might not otherwise be visible. This data is
examined in the next section.

### 4.6.4. System Time, 16 Procs only

As before, the Group signaling mechanism seems to have a significant effect on the observed performance, in this case with respect to system time. While the real time impact is about a factor of 1.3 , the system time impact is about a factor of 3 ; comparison with the earlier graphs shows that clearly, the effect of this variable increases with the number of child processes.


Figure 38: Effect of signaling process groups on sys time, 16 procs (For 100 repetitions)

Open files, shown in figure 39 , have little effect on the system time for sixteen processes:
this suggests that the results observed for real time with 16 processes may be spurious.


Figure 39: Effect of number of open files on system time, 16 procs (For 100 repetitions)

Asynchronous elimination had no effect, as is illustrated in figure 40 ; this is interesting only in that it correlates well with the previous observations.


Figure 40: Effect of asynchronous elimination on system time, 16 procs (For 100 repetitions)

Dirtying had no effect on the system time, as figure 41 illustrates.


Figure 41: Effect of dirtying child pages on system time, 16 procs (For 100 repetitions)

Processing behavior of the spawned child processes had no effect on the system time, as figure 42 illustrates.


Figure 42: Effect of child work/sleep on system time, 16 procs (For 100 repetitions)

And finally, process size had no effect on the system time, as figure 43 illustrates.


Figure 43: Effect of child process size on system time, 16 procs
(For 100 repetitions)

After reviewing the results of the last four sections, it seems clear that such factors as the number of open files, subprocess actions in executing instructions and dirtying pages, and for the most part, the sizes of the child processes have little effect on time required for sibling elimination.

The time increases with the number of processes, and this was used to see what factors change with scale; there were really no significant changes. What was surprising, however, was the major impact of process group signaling. and the non-existent impact of asynchronous execution. After some further examination of the code, we changed the code so that the do_elim process forked a copy of itself just previous to the sibling
elimination phase. In the synchronous case, the termination of this process was waited for. The asynchronous case returned without waiting. All children and grandchildren are eliminated by a post timing-loop cleanup routine.

This had a dramatic effect on the timing results, as is shown in the next section. We believe that the relationship between Groups and Asynch was due to a subtlety in the UNIX process scheduling which defeated the intent of our measurement apparatus.

### 4.6.5. Correction for process scheduling

To scale the graphs properly, all real time measurements longer than twenty seconds were removed from the graphs. This only removed outliers from the full graphs, and allows an enlargement of the relevant detail. The first graph, figure 44, of Groups against real time, shows that the effects of Groups have gone away.


Figure 44: Effect of group signaling on real time, corrected
(For 100 repetitions)

The expected change in real time for the Asynchronous case occurred, as is shown in figure 45: note that a factor of two difference is observed.


Figure 45: Effect of Asynchronous elimination, corrected (For 100 repetitions)

Groups was examined for the case Procs $=16$ to see if scale changed anything. It did not, as figure 46 illustrates.


Figure 46: Effect of process groups on real time, corrected, 16 procs (For 100 repetitions)

Finally, the effect of asynchronous elimination shows a factor of more than two improvement in the real time case with 16 processes, scaling as expected. This is illustrated by figure 47.


Figure 47: Effect of asynchronous elimination on real time, corrected, 16 procs (For 100 repetitions)


Figure 48: Effect of group elimination on system time, 16 procs (For 100 repetitions)


Figure 49: Effect of asynchronous elimination on system time, 16 procs
(For 100 repetitions)

### 4.7. Possible Sources of Error in Measurements

There are two places where errors can affect the measurements. One is the measurement apparatus, in this case the UNIX timing facility. The other is in the experimental apparatus which gathers the raw data from the measurement when experiments are performed. In this discussion, the UNIX clock facility was used for all measurements. and is a common weakness or strength of the experiments.

### 4.7.1. UNIX Clock Facility

The UNIX clock facility is implemented [Bach1986a] as follows:

1. The underlying architecture provides a "timer" facility. The timer is typically implemented as a storage location into which a number is written. This number's value represents hardware "clock ticks'; the value is decremented by the machine clock on each of its cycles. After the hardware clock has "ticked" down to zero, a timer interrupt is generated.
2. At the interrupt vector for the clock is the address of a timer routine. The timer routine first increments a counter, the "software clock." The timer routine then examines a linked list of queued tasks (the "c-list"), decrementing the value of the first task's timer. The tasks are queued in the order they should be executed; each task's "timer" is stored as the difference between it and the preceding task on the list. Storing times this way reduces searching to update timer values; the relative value is set upon list insertion. Examples of tasks are (1) checking for characters on a teletype line and (2) sending an alarm(l) signal to a sleeping process.
3. The state of the system when interrupted determines which of a set of auxiliary clock timers is incremented, e.g., USER, SYS, WAITING, or IDLE. Tasks which have non-positive timer values are executed
4. The value stored in the hardware timer is determined by selecting a HZ value for the software clock, such as 60 . For this value, software clock ticks are to occur every $\frac{1}{60}$ second, so if there are $T$ hardware timer units per second, $\frac{T}{60}$ is loaded into the hardware clock

Running processes are preempted and rescheduled at the "soft'" clock interrupt if they have not voluntarily released the processor. To reduce the overhead of such a timer facility. it can either be made fast. or called infrequently. The UNIX time facility was created for use on small slow machines, so that the second approach was applied after the first was inadequate. Thus, the clock granularity is large relative to the execution speeds of current processors. For example, the workstations we used for our measurements execute about 2 or 3 million instructions per second. So, in $\frac{1}{60}$ second, about 40,000 instructions can be executed, which is sufficient for several significant system events. e.g., context switches, to take place. Most system events, e.g., the execution of a system
call, require less than one "tick," so that timing a single system call will often show that the call took no time at all. Yet these events, for example spawning a new process with the fork() system call, are precisely what we are interested in measuring. Thus, the clock timer seems to be an insurmountable error source; the difficulties it causes in performance measurement have been noticed before [Johnson 1987a]

There are two techniques which can be used to gather useful data in spite of the large error caused by the clock granularity:

1. One trial consisting of repeated experiments from which the mean value is returned.
2. Repeated trials.

The mean of repeated samples is calculated by performing a series of measurements and timing the entirety. The value is divided by the number of experiments to get the mean. The difficulty with this technique in computer system measurement strategies is that architects have constructed system features to perform faster under this behavior, e.g., caches. Thus, the measurements may not truly be indicative of the cost of performing a single operation. The "random block number" approach of netrand.c, given as Appendix II. attempts to circumvent these caching strategies to give a truer reading for the network page access time.

Repeating trials of the experiment indicates that the experiments are reproducible. Reproducibility, a fundamental requirement of science, reduces the possibility of some artifact such as a network overload temporarily perturbing the measurements. Such an artifact would not be present in the isolated system under study. These techniques were applied in the development of our experimental apparatus, namely the programs in Appendices I, II, XI and XII.

### 4.7.2. Experimental Apparatus

We discussed the effect of repetitions on the reproducibility of experiments in the discussion of "copy-on-write" fork performance. It was clear that a small number of repetitions produced data that was not reliable, as the poor qualities of the clock for fine grained timing measurements displayed themselves. Repeating the inner loop and deducing an aggregate value produces results that are consistent with prediction. The prediction in this case is not sophisticated: it's mainly that "copy-on-write" has the most effect when little is copied. The value of this 'prediction'" is that it gives us a quick check on
the results, and it's clear that small numbers of repetitions do not adhere to the model. Our study of copy-on-write fork performance gave a graphic illustration of the effect of repetition. We also demonstrated relations between different types of time measurements (which should be closely related) which change as a function of the number of repetitions.

Repeated trials test the validity of the experiment. For example, if the experiment is to repeat some action 100 times and derive a mean execution time, repetitions of this experiment should produce approximately the same mean execution time, under the assumption of a reasonable variance. If they do not, then the figure for the mean execution time has no experimental validity. In science, since other researchers cannot reliably reproduce the number under the same conditions, it has no value. All experiments reported in the measurements section were repeated at least twice in order to ensure some measure of validity. Other validation comes from repetitions with adjusted variables. Predictions coming from models of system behavior will give estimates of the sensitivity of the experiment to various external factors. e.g., memory size or machine speed. If, for environmental factors to which the experiments are insensitive, the experimental results appear to be duplicates, the results have validity.

To test the results of the "sibling elimination" measurements, which seemed to be only grossly reproducible, we tried another strategy. The problem with the clock timer is that its granularity is constant across systems of widely varying performance. Modern processors are sufficiently fast that events can escape detection by the timer. On a slower processor, the clock appears to be of finer granularity. A subset (for example, 1, 4, and 8 child processes) of the do_elim experiments were re-run using an AT\&T 3B2/310, which as we have mentioned earlier, is significantly slower (by about a factor of 7 ) than the HP9000. The curves generated were consistent with the measurements we presented in this chapter. The consistency of the curves implies that the effort taken to limit the effects of errors in the measurement apparatus were successful. The validity of our conclusions is reinforced by the relationships maintained between parameters. These stay the same even as the magnitudes change due to the slower processing unit.

### 4.8. Discussion

We have discussed use of alternatives and shown how execution of the alternatives in a concurrent fashion can lead to performance increases. Further. we have parameterized characteristics of the alternatives and the implementation overhead so that the effectiveness of the technique can be evaluated. Many of the contributors to $\tau$ (overhead) have been identified and measured, so that the performance improvement can be evaluated on the basis of the alternative's characteristics, using these characteristics to calculate $\tau$ (overhead).

### 4.9. Applications

> "In science the primary duty of ideas is to be useful and interesting even more than to be 'true' ." |Trotter1941a|

What properties must we have, other than minimal implementation overhead, for the concurrent execution method we describe to be useful? The analysis of Chapter 2 and the measurements earlier in this chapter suggest several:

1. A large portion of the shared state is read-only.
2. There is some state shared between the alternatives which each may update. With no shared state, no work is necessary for transparent concurrent execution.
3. There are execution time differences between the alternatives, due to data dependencies, use of heuristic methods, or other influences.

Application areas for our design are described in the following sections.

### 4.9.1. Distributed Execution of Recovery Blocks

The Recovery Block [Horning1974a, Randell1975a] is a method for writing software which is tolerant of mistakes in its own logic, from which failures can arise. The idea is simple. It is assumed that the software in question has been written to some specification. Several alternative versions of the software are written, according to the specification. A boolean "acceptance test," which checks the results of the software is developed along with the software, using the specification. The acceptance test, which either succeeds or fails, will be refined once some experience with the software is developed.

The alternatives and the acceptance test are gathered into an ALGOL-like block
construct, where the alternatives are typically ordered on the basis of observed or estimated characteristics such as reliability and execution speed.

When the acceptance test succeeds, the results (including all state changes) of the alternative which passed the test are made available. When the acceptance test fails, the state of the program is "rolled back" to the state the program had before the block was entered, and the next alternative is tried. If the last alternative in the sequence results in a failed acceptance test, the block as a whole fails.

The scheme is conceptually similar to the "standby spare" technique used in hardware. $N$ alternate methods of passing an acceptance test are provided. The first such method is referred to as the primary; they have typically been rank-ordered by some metric, e.g., observed performance. Assuming that the acceptance test performs perfectly, the recovery block method fails on inputs where all methods fail the acceptance test. Note that the acceptance test is application-specific; Hecht [Hecht1979a] provides a detailed discussion of the forms such acceptance tests might take. Cha, et al. [Cha1987a] have shown that self-checks (a generalization of the acceptance tests used by recovery blocks ) can be effective in finding faults. However, there is difficulty both in the writing of the self-checks and their placement within the program structure. They also note a great variation in the ability to write effective self-checks, and the efficacy of combining code-based checks with specification-based checks compared to specification-based alone.

The recovery block is different in behavior than the "Alternative Block'" we proposed as a sequential model in the Introduction. First, rather than having one guard per body, the Recovery Block possesses one guard to which all the alternatives are passed. Second, the guard is applied after the body is executed, rather than before. However, neither of these are problems for our design, as (1) the computation can be viewed as part of the guard, with the body consisting solely of updates to external variables, or (2) the blocks can be viewed as self-checking entities where the guard is always enabled for scheduling of the computation, but which may fail due to self-checks.

The changes to the program's state space are equivalent to some execution which selected exactly one alternative at each Recovery Block encountered in the execution. This the nondeterministic selection which we discussed in the introduction. The RB language, developed by Smith and Maguire [Smith 1988c] is designed for the distributed
execution of recovery blocks.
Since Recovery Block alternates may attempt to update shared state, e.g., database files or external variables, our mechanism for preventing observation of a sibling's actions is necessary, and the "copy-on-write" memory management reduces the amount of state which must be maintained. One special problem which arises with the parallel execution of Recovery Block alternates is the fact that the method is designed to cope with failures, so that we must do more work in order not to add new failure modes. Two issues in particular are important. First, we may copy all the state rather than copying as necessary, in order that the state not become inaccessible ${ }^{14}$ and so cause a failure. Second. the synchronization must not introduce a single point of failure. This is remedied by the use of majority consensus. as discussed above, to achieve a fault-tolerant 0-1 semaphore for use in synchronization. A combination of existing techniques [Schneider1983a, Schneider 1982a] may be needed for the distributed case.

### 4.9.2. Polynomial Root-finding

The Jenkins-Traub [Jenkins1972a, Jenkins1970a] algorithm for finding roots of polynomial equations is a highly robust, rapid, and portable method for solving for the zeros of a polynomial over the complex numbers. Those interested in the internal workings of the algorithm should consult Ralston [Ralston1978a] for details. The property of the algorithm which makes it attractive as an application for our method is that it is adaptive, in the following sense: as it begins the search for a zero in the complex plane, the algorithm chooses an angle with which it attempts to approach the root. After iterating towards the root, the algorithm tests for convergence. If the iteration is not converging, the algorithm retries with another angle, which in the published version (ACM Algorithm 419: 'Zeros of a Complex Polynomial') is a fixed rotation in the plane. The application of the method for concurrent execution is in the choice of the angle of approach. Several such angles are selected, and alternative root-finders are spawned with the angle set. If an

[^11]alternative algorithm does not converge, it "fails" by exiting; successful alternatives are selected using our "fastest first" synchronization scheme; our performance analysis of the speedup applies here as well. The minimum time is the CPU time it took for the first rootfinder to discover all the zeros of the polynomial. Failures are excluded. since they did not discover all the zeros. The maximum time is the CPU time required for the last rootfinder to discover all the zeros of the polynomial, or fail. Failures are included, since we would have to wait for them if we didn't know whether the rootfinder was converging. The average time is the mean of all execution times, including failures. These times are determined on a uniprocessor, since each angle choice will be executing on such a machine in the experimental work.

The granularity of finding a single zero may be too fine to dominate the overhead involved, but alternatives can easily be spawned which retry using different rotations or different starting angles for the entire set of zeros in this case.

As a test, we re-coded the published Fortran program for the Jenkins-Traub method in "C." This is provided as Appendix XIII. A variety of test cases were run against the algorithm, using a driver which varied the initial angle for the sequence of shifts. The published algorithm had used 94 degrees as an angle; we parameterized cpoly() with an angle argument which allowed different choices to be tried. The driver program, cvaryangle.c, is given as Appendix XIV. The compiled program's address space had a write fraction of less than 0.25 due to the large portion comprised of program text.

The test cases were generated using r2p.c which is given as Appendix XV. The idea is to generate a distribution of roots, which are used as input to $r 2 p . r 2 p$ constructs a monic polynomial having these roots, and outputs its coefficients into a form usable by cvaryangle.c.

### 4.9.2.1. Example

Consider the following set of points, which are to be the roots of a polynomial:

$$
\begin{aligned}
& -1.1+0.1 \mathrm{i} \\
& -1.1+0.1 \mathrm{i} \\
& -1.0+3.1 \mathrm{i} \\
& -1.2+0.334 \mathrm{i} \\
& -1.3+1.276576 \mathrm{i} \\
& -0.4+0.026 \mathrm{i} \\
& 1.5+1.1 \mathrm{i} \\
& 0.3+1.0 \mathrm{i} \\
& 0.4+1.0 \mathrm{i} \\
& 0.5+1.0 \mathrm{i} \\
& -1.6+0.0 \mathrm{i} \\
& -0.7+0.76 \mathrm{i} \\
& -1.8+1.9 \mathrm{i} \\
& -0.9+0.0 \mathrm{i} \\
& 0.1+0.042 \mathrm{i} \\
& 0.1+0.042 \mathrm{i} \\
& -1.0+3.1 \mathrm{i} \\
& -1.0+3.111 \mathrm{i} \\
& 0.2+0.5 \mathrm{i} \\
& 0.7+0.3 \mathrm{i} \\
& 0.4+0.7 \mathrm{i}
\end{aligned}
$$

The roots are dispersed through the top half of the complex plane:


When processed by $r 2 p$, they yield ${ }^{15}$ the monic 21 st degree complex polynomial $z^{21}+$ $(8.9-19.491576 i) \cdot z^{20}+$
$(-1274.642064+381.680467 i) \cdot z^{18}+$ $(16752.870876+10043.831986 i) \cdot z^{16}+$ $(-23799.211445-106982.985377 i) \cdot z^{14}+$ $(-152123.766242+236186.810121 i) \cdot z^{12}+$ $(342742.413912-83786.997252 i) \cdot z^{10}+$ $(-195080.181222-100059.575279 i) \cdot z^{8}+$

$$
\begin{array}{r}
(-140.045104-161.369678 i) \cdot z^{19}+ \\
(-743.821643+5810.837248 i) \cdot z^{17}+ \\
(41763.432806-29981.366628 i) \cdot z^{15}+ \\
(-189267.174496-34407.932181 i) \cdot z^{13}+ \\
(198013.830239+280742.352179 i) \cdot z^{11}+ \\
(37460.901644-301901.877085 i) \cdot z^{9}+ \\
(-94732.08529+90223.058904 i) \cdot z^{7}+
\end{array}
$$

${ }^{15}$ R2p.c has lost much precision; the real coefficients of the polynomial are different from those it computes. This approximation is not bad, $\mathrm{P}(-0.9+0.0 \mathrm{i})$ is $0.8 \mathrm{E}-7$, which is not far from zero. The polynomial $\mathrm{P}(0$ for which $\mathrm{P}(-0.9+0.0 \mathrm{i})$ is $-0.26 \mathrm{E}-26$ has vastly different cocfficients. For example the coefficient of $z^{0}$ is $-1088.35+607.0 \mathrm{i}$ [Chang1989a]. However, this precision is not necessary to prove our point about parallel processing!

| $(27282.343436+57440.498715 i) \cdot z^{6}+$ | $(24271.198674-3724.963179 i) \cdot z^{5}+$ |
| :--- | ---: |
| $(682.559499-7237.864131 i) \cdot z^{4}+$ | $(-1528.494152-397.912167 i) \cdot z^{3}+$ |
| $(-63.349391+237.887291 i) \cdot z^{2}+(27.878345+0.897986 i) \cdot z^{1}+(-0.80232-1.101871 i)$. |  |

When the program of Appendix XIV, cvaryangle.c, is run on this input, the following output is produced on a Hewlett-Packard HP9000/350 with 8 megabytes of main memory and an HP7945 70 megabyte hard disk, running HP-UX 6.0:
nprocs: l, max: 1044, min: 0, avg: 1044, fails: 1 nprocs: 2, max: 1015, min: 473, $\exists \mathrm{vg}: 744$, fails: 1 nprocs: 3, max: 1733, min: 1733, avg: 1246, fails: 2 nprocs: 4, max: 1604, min: 474, avg: 1047, fails: 1 nprocs: 5, max: 1531, min: 814, avg: 1117, fails: 3 nprocs: 6, max: 1734, min: 473, avg: 938, fails: 2 nprocs: 7, max: 1330, min: 791, avg: 1023, fails: 2 nprocs: 8, max: 1571, min: 461, avg: 1037, fails: 1 nprocs: 9, max: 1706, min: 1080, avg: 1206, fails: 3 nprocs: 10, max: 1877, min: 459, avg: 1076, fails: 3 nprocs: 11, max: 1896, min: 965, avg: 1324, fails: 1 nprocs: 12, max: 1758, min: 472, avg: 1127, fails: 3 nprocs: 13, max: 3307, min: 775, avg: 1437, fails: 4 nprocs: 14, max: 1885, min: 790, avg: 1275, fails: 5 nprocs: 15, max: 1748, min: 293, avg: 1053, fails: 6 nprocs: 16, max: 2312, min: 476, avg: 1194, fails: 2 nprocs: 17, max: 1937, min: 656, avg: 1159, fails: 5 nprocs: 18, max: 1816, min: 504, avg: 1066, fails: 4 nprocs: 19, max: 2085, min: 738, avg: 1223, fails: 2 nprocs: 20, max: 2381, min: 474, avg: 1238, fails: 3 nprocs: 21, max: 2100, min: 619, avg: 1235, fails: 8 nprocs: 22, max: 1887, min: 380, avg: 1135, fails: 4 nprocs: 23, max: 2088, min: 643, avg: 1344, fails: 3 nprocs: 24, max: 2056, min: 472, avg: 1147, fails: 4 nprocs: 25, max: 2317, min: 699, avg: 1154, fails: 8 nprocs: 26, max: 2003, min: 566, avg: 1249, fails: 6 nprocs: 27, max: 1902, min: 590, avg: 1167, fails: 5 nprocs: 28, max: 1930, min: 545, avg: 1149, fails: 8 nprocs: 29, max: 2346, min: 332, avg: 1297, fails: 7 nprocs: 30, max: 1948, min: 312, avg: 1110, fails: 6 nprocs: 31, max: 1950, min: 657, avg: 1201, fails: 5 nprocs: 32, max: 2273, min: 471, avg: 1183, fails: 7 nprocs: 33, max: 2529, min: 505, avg: 1227, fails: 5 nprocs: 34, max: 2378, min: 490, avg: 1330, fails: 4 nprocs: 35, max: 1774, min: 420, avg: 1104, fails: 7 nprocs: 36, max: 1758, min: 471, avg: 1088, fails: 7 nprocs: 37, max: 2245, min: 299, avg: 1224, fails: 8 nprocs: 38, max: 2556, min: 568, avg: 1222, fails: 7 nprocs: 39, max: 1867, min: 680, avg: 1140, fails: 4 nprocs: 40, max: 2317, min: 473, avg: 1255, fails: 5 nprocs: 41, max: 2165, min: 552, avg: 1252, fails: 11 nprocs: 42, max: 2703, min: 621, avg: 1382, fails: 7 nprocs: 43, max: 2232, min: 515, avg: 1268, fails: 13 nprocs: 44, max: 2140, min: 425, avg: 1199, fails: 4 nprocs: 45, max: 2171, min: 312, avg: 1225, fails: 8 nprocs: 46, max: 1911, min: 429, avg: 1159, fails: 7 nprocs: 47, max: 2511, min: 439, avg: 1265, fails: 5 nprocs: 48, max: 2271, min: 471, avg: 1229, fails: 9 nprocs: 49, max: 2698, min: 476, avg: 1211, fails: 6 nprocs: 50, max: 3216, min: 543, avg: 1231, fails: 9

The timing numbers are given in units of clock ticks; there are 60 clock ticks per second
on this machine. The relations are graphed in figure 50:


Figure 50: Varying angle versus execution time

## Legend:

"*" - Minimum execution time
"@" - Maximum execution time
"-_" - Average Execution time
"Failures" - Number of choices causing failure
Times are in CPU clock tick units.
For a number of processors $N$, the ensemble of angles $<3.0+\frac{0 \cdot 360}{N}, 3.0+\frac{1 \cdot 360}{N}, \ldots, 3.0+\frac{(N-1) \cdot 360}{N}>$ is tried in parallel. The time for failures
is included in the averages, but the minimum time is always derived from successful executions. The number of failures is calculated by using a counter, which is output. These results illustrate two important facts:

1. A speedup is possible using only a few variations. For example, comparing the average case with 4 processors to the best case gives a speedup of about 2.2. The speedup in the two-processor case is 1.6 . Of course, these measurements ignore overhead. More importantly from the point of view of a numerical analyst, the scheme tolerates failures.
2. Certain choices of angle result in failures. This is strikingly illustrated for 1 processor, and is due to, among other things, the condition of this polynomial. Picking the first successful execution would work for any number of processors greater than 2 in this experiment.

### 4.9.2.2. Parallel Execution

The Jenkins-Traub algorithm displayed significant variance on this input and others. Since we had a working program available, we decided to measure the performance of our scheme on a multiprocessor system to see how much overhead arises in the complete, end-to-end algorithm. Copies of the Jenkins-Traub algorithm were run using the parallel version of cvaryangle.c given as Appendix XVI. Cmach.c was run on several multiprocessor systems. For an otherwise idle two processor Ardent Titan, the first six lines of $c j t$ produced the results:

| procs | $\max$ | $\min$ | avg | fails | par |
| :---: | :---: | :---: | :---: | :---: | :---: |
| 1 | 4.01 | 4.01 | 4.01 | 0 | 4.37 |
| 2 | 4.49 | 4.07 | 4.28 | 0 | 4.25 |
| 3 | 4.45 | 2.03 | 3.50 | 0 | 4.74 |
| 4 | 4.48 | 1.37 | 3.31 | 0 | 5.19 |
| 5 | 4.27 | 2.36 | 3.35 | 2 | 8.61 |
| 6 | 4.50 | 2.02 | 3.65 | 0 | 7.03 |

The column labeled "par" is the parallel execution time measured with cmach. All times
below the double lines were with processor contention, since the number of processes was greater than the number of available processors.

Thus, the execution time overhead of creating two processes and running them concurrently can be computed as $4.25-4.07 \mathrm{sec}$., or about .18 sec . But the average time was 4.28 sec , so even with the additional overhead, the parallel execution finished first.

While encouraging, the numbers suggested that more processors might yield more impressive results. On an experimental [Garcia1989a] multiprocessor, cmach was run on the degree 16 polynomial $z^{16}+\quad(19.800000-3.200000 i) \cdot z^{15}+$ $(176.930000-58.940000 i) \cdot z^{14}+$ $(940.246000-494.724000 i) \cdot z^{13}+$ $(3284.874700-2495.183000 i) \cdot z^{12}+\quad(7821.151500-8375.778560 i) \cdot z^{11}+$ $(12521.112583-19491.045162 i) \cdot z^{10}+\quad(12045.424807-31405.685381 i) \cdot z^{9}+$ $(2785.185034-32740.983162 i) \cdot z^{8}+\quad(-10378.708048-15172.462694 i) \cdot z^{7}+$ $(-16870.986162+13554.317038 i) \cdot z^{6}+\quad(-12430.788642+33571.308326 i) \cdot z^{5}+$ $(-3673.995257+33389.404113 i) \cdot z^{4}+\quad(1360.752190+20180.264386 i) \cdot z^{3}+$ $(1689.471245+7674.191752 i) \cdot z^{2}+\quad(622.122192+1700.277923 i) \cdot z^{1}+$ ( $86.407858+168.238421 i$ ). An eight-processor configuration running MACH produced the results illustrated in the following table:

| procs | $\max$ | $\min$ | avg | fails | par |
| :---: | :---: | :---: | :---: | :---: | :---: |
| 1 | 12.1 | 12.1 | 12.1 | 0 | 12.8 |
| 2 | 13.1 | 6.2 | 9.6 | 0 | 6.3 |
| 3 | 22.9 | 13.1 | 17.1 | 1 | 12.3 |
| 4 | 13.4 | 6.23 | 10.7 | 0 | 6.4 |
| 5 | 19.9 | 5.8 | 12.2 | 0 | 6.3 |
| 6 | 23.0 | 6.2 | 13.8 | 1 | 11.4 |
| 7 | 16.9 | 5.9 | 10.1 | 0 | 7.4 |

When plotted yields:
$\mathrm{M} \quad \mathrm{M}$
$20-\quad$ M
Time A
M
(Secs)
A


Figure 51: Performance of polynomial root-finder
The legend is quite simple: $\mathbf{M}$ designates the maximum time of an ensemble of angles, $\mathbf{m}$ designates the minimum time, A designates the mean time (excluding failures), and $\|$ is the execution time, including all overheads, required for parallel execution. The overhead can be calculated from the gap between $m$ and $\mathbf{A}$ values in the plot. The speedup can be seen by comparing A to $\|$. This legend is used for the plots of Appendix III.

For these tests, a single process was run before the testing started, in order to prepage the code into the multiprocessor from the host; this single process took 17.2 seconds. This was done because the ACE is attached to an IBM PC/RT host, from which code is loaded into the ACE address space before execution. This initial load would introduce a spurious execution time increase, and must be removed from the results. A variety of polynomials were tested; this was neither the best nor worst performance observed. A number of other test sets and performance tables are given in Appendix III. Figure 52 summarizes the speedups seen on these examples, which were chosen based on
the advice given in several papers on testing rootfinders. The summary statistics clearly reflect the limited dispersion available, as illustrated in the graph.

We first calculate the speedup for a given number of processors on each polynomial by computing the ratio of the average time to the observed time for parallel execution. The mean of these speedups is then computed, giving the value in the table.

| procs | speedup |
| :---: | :---: |
| 1 | 0.99 |
| 2 | 1.16 |
| 3 | 1.23 |
| 4 | 1.28 |
| 5 | 1.26 |
| 6 | 1.11 |
| 7 | 1.10 |

So, for example, the method produced, on average, a 28 percent speedup when 4 processors were utilized. This is a real speedup, since all processing overhead is accounted for in the observed execution times.

The research contribution of this method of concurrently executing the alternative versions of the Jenkins-Traub algorithm is significant. Certain component operations of the algorithm can be vectorized, so that processors such as a Cray [Russell1978a] or CDC 6600 [Thornton1970a] CPU could extract some parallelism from the execution. However, this parallelism is available to speed up all the alternatives, since they differ only by an initial angle and not in their executable code, which is shared. Other than exploitation of vectorizable code, or perhaps fine-grained dataflow properties, the algorithm is inherently sequential, due to the use of polynomial deflation [Ralston1978a] when a root is found. Yet, we have shown a method by which significant speedups can be achieved through use of multiple processors, and we have demonstrated those speedups in practice.

### 4.9.3. Other Applications of the Technique

We discuss other applications of the technique. These are drawn from varying areas of computer science.

### 4.9.3.1. OR-parallelism in Prolog

Logic programming is a new method of harnessing computers to solve problems. The major idea is that the "logic program'" declares what must be true, and the logic programming system finds a way of making this true. This results in a solution to the problem. Logic programming is discussed by Kowalski [Kowalski1979a]. Logic programming's attraction to researchers is primarily because of the freedom from constraints on "how" the "what" of the program is to be achieved. Thus, schemes for short-cuts and parallel execution abound. Much of this research activity has had as its focus the Prolog language, to which we will restrict our attention.

The Prolog [Clocksin1984a] programming language is based on predicate logic [Kowalski1979a], using "Horn clauses'" [Rich1983a] to describe data and interrelationships. Many normal operations are subsumed by the unification algorithm by which Pro$\log$ attempts to satisfy predicates: variables are bound during the unification process to values which caused the predicates to become true. Thus equal ( $\mathrm{X}, \mathrm{elrod}$ ) will cause the variable X to take on the value elrod, as this binding is the only one which allows the predicate equal () to be satisfied.

Progress is achieved with a goal-oriented predicate-satisfaction algorithm; a database of predicate values and rules is used to construct a set of dependency relations; toplevel goals are decomposed into sub-goals using the relations between the rules, objects, and predicates. For example, testing equality of lists implies that their elements are equal: testing element-wise equality may then give a list of sub-goals. This gives rise to a possibility for parallel execution, however the granularity of such parallelism seems inappropriate. More appropriate is rule-level parallelism, which is centered on two types, AND-parallelism and OR-parallelism. The idea with AND-parallelism is that if we have a situation where goals $A$ and $B$ must be simultaneously satisfied, we can pursue the satisfaction of $A$ and $B$ in parallel, and deal with the simultaneity later. This, however, has proven possible but difficult. due to the requirement of consistent variable bindings in A and B . The situation is similar for OR-parallelism: this is more interesting to us,
since it maps closely to our problem of attempting alternatives in parallel.

### 4.9.3.1.1. Tutorial Example

We'll start with a database of "facts," which are essentially ground literals of first-order predicate logic. The example we'll construct will be useful in understanding the Prolog execution engine, so that we can examine what opportunities exist for parallel execution. The knowledge base consists of the following facts:

```
father( charles, jonathan ).
father( frederick, charles ).
father( raymond, jacqueline ).
father( john, gertrude ).
father( charles, steven ).
mother( jacqueline, jonathan ).
mother( gertrude, jacqueline ).
mother( julia, charles ).
mother( mary, gertrude ).
father( frederick, elizabeth ).
father( frederick, frederickjr ).
father( frederick, patricia ).
```

First, let's consider some queries in the sequential execution of a Prolog interpreter. Those desiring a more detailed tutorial can consult Clocksin and Mellish's [Clocksin1984a] book. The clause

```
grandfather( X, Z ) :-
    father( X, Y ), father( Y, Z ).
```

defines a predicate which defines grandfatherhood in terms of two related fatherhoods. The interpretation of the comma is that it combines the clauses separated by it in such a way that all of them must be true. The " $:$ :-" can read so that the right-hand side implies the left hand side. Thus, both father ( X, Y ) and father ( Y, Z ) must be
true for grandfather ( $\mathrm{X}, \mathrm{Z}$ ) to be true, and $Y$ must take on the same value in both clauses. Consider the Prolog query:
?grandfather ( G, jonathan ).
G=[frederick]
What has happened here is that the Prolog interpreter has taken the query and decomposed it into the two sub-queries. It first tries to match father ( $X, Y$ ), and the first match (assuming the literals are organized as shown) in the database will be the literal father( charles, jonathan ). Thus, $X$ is instantiated as charles and $Y$ is instantiated as jonathan. The second clause must then match father ( jonathan, jonathan), which it does not. Since the match failed, the algorithm backtracks to the last choice point. and the search for a value of father ( $\mathrm{X}, \mathrm{Y}$ ) continues, and $X$ and $Y$ are instantiated as frederick and charles, respectively. Since $Y$ has now been "bound" to charles, the database is examined for a match for father ( charles, jonathan ) which it finds as the first entry. The interpreter then sets $G$ to frederick and prints the value of $G$. The interpreter also waits for an indication that the search is to be continued, in case there are more solutions to the query ${ }^{16}$.

Now, we have ignored the possibility of a maternal grandfather, which can be specified with the additional clause

```
grandfather( X, Z ) :-
```

    father ( X, Y ), mother ( Y, Z ).
    The existence of this clause changes the interpreter's control flow in the following way, as a result of there being two alternative methods of demonstrating grandfatherhood. Consider the query:

```
?grandfather( raymond, jonathan ).
```

T
There is no pair of values which satisfies the first clause, so it would fail. However, there remains a second clause which can be tried, and father ( raymond, Y ),

[^12]mother ( $Y$, jonathan ) is satisfied by $Y=j a c q u e l i n e$, and the query is satisfied.

The first query offers an opportunity for parallelism in the database search. Elements satisfying the first sub-query can be matched against elements satisfying the second; this merging process produces a result which can satisfy the original query. If the conjunction of clauses does not share variables, then the merging is easier. Management of the general case of these queries is difficult, since the sub-queries may have caused shared variables to exhibit side-effects, i.e., "bindings." However, in having two methods for "proving" grandfather ( raymond, jonathan ) we have introduced a new opportunity for parallel execution. The query is true if either top-level clause results in an answer. Thus, we can attempt to process both of these queries in parallel, since a successful sub-query in either case can cause the top-level query to be satisfied.

One major problem, as pointed out by Warren in his survey [Warren1987a] (ignoring side effects) is the problem of "multiple binding environments." What this means is that as the clauses execute in parallel, they bind values to variables, e.g., jacqueline to $Y$. Now. different alternatives may bind different values to the variables at different times, and hence interfere with each other, in a way that would not be possible in the single threaded stack-oriented interpretation style we described. How this problem can be dealt with is a question which has inspired many researchers, and engendered many solutions. A good, but somewhat dated, survey is provided by Wise [Wise 1986a] who surveys work in parallel logic programming for comparison with his EPILOG system. Some of these are described in the next section.

### 4.9.3.1.2. Existing Solutions

The basic problem is to prevent the clauses from interfering with each other, through side-effects, during their execution. One solution which has been examined is called committed-choice non-determinism. Clauses are partitioned into guards and bodies, so that they take the form:

```
head :- guard, body.
```

The guards of all clauses selected by the matching algorithm are evaluated in parallel. From among the clauses whose guards are true, a body is selected for execution. The
choice of this clause is the committed-choice. In order that they can be executed in this fashion, the guards are often [Clark 1987a] constrained to be read-only, i.e., side-effect free. For example, static analysis by the PARLOG [Clark1987b] compiler insures that the clauses are read-only, before execution. The GHC [Ueda1987a] scheduling mechanism blocks processes which update shared state.

Another approach is maintain the language features of Prolog, and introduce the parallelism in a transparent fashion. This is the approach of the systems Warren describes in his survey. They use a combination of interpreter-controlled variable management and interpreter-controlled process scheduling to prevent updates which would result in an inconsistent state. Processes are typically modeled as independent threads of control working in a common address space. Thus, we could look at the forked() sub-processes as inheriting the state of their parent process; classical resolution ${ }^{17}$ would copy all the state at each sub-node in a computation tree. A "node" is this tree is a list of goals; for example the root node's goal is satisfaction of the initial query. The computation consists of a series of decompositions of the initial query into clauses which are satisfied, left to right in a traditional Prolog system. This results in a depthfirst search of the tree; search termination is achieved when a goal is "resolved," or matched with some entry in the knowledge base. As each sub-node is executed, it consists of a copy of its parent node, with the left-most clause instantiated to whatever knowledge base entry matched it. Aside from the execution cost, the main additional overhead here is the copying of the parent's goals. This burden of copying would occur at each branch point, where a fork would occur, in a parallel execution. Aside from this burden, Warren points out that the approach does offer the attractive properties of (1) the processors can work independently on physically separate data, and (2) in all other respects, the implementation would be the same as a sequential implementation, and therefore would be able to take advantage of any optimizations, etc., that might apply.

A simple modification, and an optimization, is the "Naive Model," where rather than copying state, binding lists (<name,value> pairs) for variables are associated with each node in the tree. The sub-node instantiates the elements of the binding list; no physical copy need be made of an unchanged variable. To look up a variable's value, a

[^13]search through the binding list must be made, as bindings form a LIFO queue. The major drawback is that the size (and hence the search time) of the binding list is not bounded. The 'Naive Model" is close to the way sequential Prolog maintains its trail stack, used for backtracking from a failed goal.

To address this problem of unbounded lookup time for a binding list, the "binding array" was proposed. The idea is that each processor has a binding array in local store; there is one node per processor. The array is instantiated when a goal is selected for resolution by a processor: it is a constant cost lookup, so that the "binding list's major problem is eliminated. A clever implementation uses the binding array data structure to "shadow' the contents of the binding list, as goals are executed. When a processor begins to execute a new goal, the binding array must be set up, so that task switching now gains more overhead; this can be considerable, depending on how many bindings must be changed.

The "Argonne Model" uses hash table lookup to cut down the cost of searching the binding array by a constant factor. There is in addition the idea of "favored'' bindings. which are local to the processor, and "unfavored," which are non-local in the sense that they are bindings that are "favored" by some other processor. Non-shared variables are denoted "private." Only "unfavored" bindings must be looked up in the hash table, although hash tables are created for each arc of the OR-parallel tree, and are kept in a linked "chain." Figure 4 of Disz, et al's paper [Disz1987a] is particularly illustrative. "Private" and "favored" variables have constant times for binding (i.e. assigning) values to variables. "Unfavored" bindings are more costly, as they involve table accesses and updates. If most shared variables are "favored,' this scheme is effective. However, as Warren points out in his survey [Warren1987a], the fraction of variable references which are unfavored is high, i.e., 0.2->0.5. Disz, et al. [Disz1987a] were similarly disappointed by the performance of the scheme on benchmarks which they reported. A "context-switch" (where the processor suspends execution of the currently executing process, and resumes the execution of some process) is expected to be cheap compared to the context-switch cost for the "binding arrays" scheme, as it involves only establishing pointers to the proper hash chain, not copying. For the "unfavored" bindings, the lookup cost, while reduced by a large constant due to the hashing technique, can still be (potentially) unbounded, as it remains proportional to the number of unfavored bindings.

The "Manchester-Argonne" model improves upon the Argonne model by applying the observation that the Argonne model's hash chains are not necessary unless sharing occurs. Thus, in the optimized model, allocation is postponed until a processor wants to share an 'arc;'" at this point the number of relevant bindings is known and assuming a reasonable 'randomizer" the hash table technique can yield a constant-bounded time lookup, and Warren argues that it will, with scheduling and merging techniques which he describes.

Finally, the "Argonne-SRI' model uses the "binding array" technique of the "SRI' model combined with the favored/unfavored distinction between shared variables from the "Argonne" model. The private "binding array" is used for unfavored bindings only; "favored" bindings are marked with some flag indicating their status; this flag is associated with the value. (Note that the space for this flag must be available in all value cells. If the 'space"' devoted to this is greater than the fraction of shared variables, the straight "binding-array" may be more space-efficient.) Thus, if many bindings are "unfavored." this technique will cause an extra access (of the binding array) for each of them, but the time will be constant. Warren's opinion of the merit of the various schemes is summarized as:

## Argonne<Manchester-Argonne Abstract<Naive< ${ }_{S R I<A r g o n n e-S R I ~}^{\text {S }}$

Now, all these schemes make the following assumptions:

1. The Prolog implementation (compiler/interpreter) must manage its own memory.
2. If multi-processing, memory is shared.
3. The Prolog implementation is responsible for task management.
4. Variable-binding is considered to be the major problem. The implication of this is that other concerns, e.g., side-effects and IPC, are ignored or given short shrift.
5. Adherence to sequential Prolog syntax and semantics must be maintained. This is a more modern version of the "dusty-deck" arguments associated with any changes in an existing language.

Unfortunately, these assumptions are not always met, or in some cases, reasonable. We will examine the assumptions point-by-point.

1. Unless a special-purpose Prolog machine is built, and in addition is economically viable, the cost/utility/performance of virtual memory will dictate its inclusion in general-purpose computers. Thus, in reality, an abstract 'memory reference" must go through two memory management schemes: Prolog's, and the hardware's virtual addressing support (this ignores other "transparent" architectural features such as caches).
An interesting question which has arisen in the study of block sizes (see Deitel [Deitel1984a] for a discussion) for page-oriented virtual memory systems is the penalty paid for fetching data in blocks which may or may not be well-correlated with the problem characteristics. For example, if accesses are done on memory-word sized units which are widely separated, each word access may require several thousand instructions to handle the associated page fault; the same problem arises when choosing a cache line size. While this difficulty can be reduced, if not eliminated, by problem-specific approaches such as overlays, these are clumsy to manage and are not transparent to the programmer. In any case, most programs exhibit locality of reference; LISP and Prolog programs are no exception.
2. While shared-memory architectures are common and provide a nice abstraction for the programmer, distributed computing is different with respect to the latency and bandwidth of communication versus the latency and bandwidth of local memory referencing [Smith 1986a], therefore the economics of communication become much more interesting, even if the communication is buried beneath a shared memory abstraction LAbramson1985a. Delp1988a. Cheriton1986a, Li1986a, Li1986b, Stout1983a, Poplawski1987a]. For example, locality of reference becomes much more important because its impact on performance is so much greater. Some attention has been paid to the issue of Prolog memory referencing behavior in the literature. Tick's [Tick1988a] thesis provides a wealth of detail, but assumes a specific model for the abstract machine. Nguyen [Nguyen1988a] uses a variety of benchmarks for a Prolog running on an existing system, some of which are discussed later in this section. Ross [Ross1986a] discusses detailed measurements of virtual memory access behavior for a Prolog system, and makes several suggestions for improved performance.

While attractive in terms of reducing unnecessary references, indirection and pointer-based <name, value> retrieval look less attractive when the number of remote
references is calculated. Crammond [Crammond1985a] looks at three methods of creating an execution environment for parallel execution of disjunctions. He provides some analysis of mechanisms designed for efficient reference of shared data, in particular the update of shared data. He studies three algorithms, which he calls "Directory," "Hash Window," and "Variable Importation." 'Directory" and "Hash Window" do bindings (i.e., variable instantiations) on demand, via pointers, while "Variable Importation" imports everything it may need, at higher initial overhead, but taking more advantage of locality once it performed the importation. Crammond points out in his study that copy-based schemes ("variable importation') for multiple binding environments, such as Lindstrom's [Lindstrom1984a, Lindstrom1984b, Tinker1987al begin to look more attractive under such circumstances, since their performance is competitive with the other schemes in any case.
3. This is not a bad assumption in intent, as the Prolog implementation's knowledge of semantics and scheduling heuristics (which are not applicable to more generalpurpose operating system process schedulers) may make it a much more effective scheduling tool. However, if interaction with the scheduler of a general-purpose operating system is necessary, as we argued it would be without Prolog machines, then heuristics will be applied based on CPU utilization rather than execution-time performance, (they may be different, as we saw earlier in our measurements of overhead) and due to load-balancing schemes using process migration [Ferguson1988a], not easily predictable. Communication between the scheduling component of the general-purpose operating system and the Prolog implementation thus seems like an effective strategy in these circumstances.
4. One major problem which the committed-choice non-determinism logic programming languages [Shapiro1986a] address is that of side-effects. How is an "allsolutions" requirement dealt with if each solution writes a payroll check? Butler, et al. [Butler1988a] point out how these issues damped their initial enthusiasm [Ciepelewski1985a] for the "Dusty Deck" solutions, and led them to include a "commit'" operator in their language [Hausman 1987a, Lusk1988a]. While it could be argued that the programmer must be aware of such possibilities and adjust for them, this seems wrong, as experience has shown that good design of support mechanisms enhances programmer productivity, and myriad subtleties and
restrictions reduce it. "Committed-choice" semantics address the issue simply: they guarantee that only one result will be produced by the construct, and do not specify the selection criterion in order to allow themselves full use of the degree of freedom provided by nondeterminism.
5. Cost-benefit analysis usually dictates whether changes can reasonably be made, if they are technically justifiable. When a large amount of software assuming a certain semantics is written, this large installed base and its assumptions represents a huge cost which must be weighed against improvements derived from the change. For example, the performance benefits of pipelining in the IBM 360 Model 91 were considered significant enough so that the architectural specification of the 360 for that particular machine was changed to include an imprecise interrupt. Prolog has the advantage in that it is newer, and the installed base is much smaller, and that a small change can yield such benefits, especially when considered in the context of Point \#4. In addition, the "guarded command" structure has strong intuitive appeal, as it has appeared independently in several settings [Dijkstra1976a, Randell1975a, Ledgard1981a].

### 4.9.3.1.3. Discussion

What our method does is copy using virtual memory support provided by the operating system, and since we choose only one alternative, no merging is necessary. Since there are no extra (beyond whatever is required for sequential execution) pointer chains to traverse on variable references, memory access is fast. Use of the method requires changing the Prolog interpreter to detect and exploit OR-parallelism. How aggressively available parallelism is exploited is a function of the overhead associated with maintaining a process. However, once this is known (we show how to measure the overhead in Sections 4.2-7 of this thesis), the proper granularity can be used as a factor in the decomposition process.

We see the following advantages to our approach:

1. Virtual memory support in some form is going to be available on general purpose computers, both because it is useful, and because it makes a great deal of economic sense. While Prolog machines are desirable, as were the LISP machines, it's not clear that the performance advantages which accrue from special-purpose
architectures can overcome the steady increase in performance seen by general purpose computing engines.
2. Our arguments for performance were completely general, and our example applications have been drawn from a large spectrum of computing tasks. Thus, the mechanism itself is useful outside of Prolog, and thus should be made available to other applications.
3. The workstations our measurements were made on are representative of the processing components which comprise many of the commercially- produced multiprocessor engines. For examples, see the Sequent [Beck1985a] or Encore [Encore1985a] multiprocessor machines.
4. The problem with copying is that it's costly. But, (1) as we've shown in this thesis, the cost can be parameterized (2) the cost can be reduced through optimizations such as copy-on-write (3) once the copy is made, it takes full advantage of any available per-processor cache [Smith1982a]. Following a pointer chain or indirection through a hash table will then cost more after the first reference. Analysis of the page-fault cost amortized over subsequent references versus the cost of following chains (much of which may be cached as well) on each reference should be done, although the analysis of Crammond, as discussed above, points to the fact that one implementation of a copying scheme (not page-level) did not perform badly compared to indirectionbased schemes.
5. When considering a distributed system, as opposed to the bus-connected shared memory multiprocessor which most studies have assumed, the cost of remote references is increased to the point where copying looks increasing attractive, as the copy which is made serves as a cache.
6. The support of Prolog OR-parallelism entails certain overheads even in interpreterbased memory management schemes. While the more advanced schemes described above tend to take a more demand-based approach to copying, there is overhead associated with adding these features which is incurred by all Prolog programs running using the augmented interpreter, e.g., extra tables, extra pointers, or restricted syntax. Our method takes advantage of services already available under general-purpose operating systems, and a Prolog program which does not use the features runs no more slowly than the same Prolog program running on a system without the features.

It must be noted that the interpreter-controlled approaches which use their own tables to handle multiple binding environments will be portable, while our virtual memory based scheme will require interaction with an operating system, which may have an adverse impact on portability. However, as we've argued, and as others have observed, solving the multiple binding environments problem is not equivalent to managing all side-effects. In summary, we've offered an interesting and sometimes advantageous scheme for executing Prolog programs in parallel. In addition, since we have quantified where the advantage will occur, measurements of program execution time combined with our overhead figures from previous sections of this chapter will give us a good estimate for the technique's effectiveness.

### 4.9.3.1.4. Measurements of Published Prolog Programs

To test the efficacy of our technique, we measured some programs from a standard Prolog reference, Clocksin \& Mellish [Clocksin1984a]. The programs are different algorithms for sorting, applied to various data sets. The sources for the sorting programs are given as Appendix IV. The programs were applied to three data sets, those of Appendices V, VI, and VIII. The data of Appendix VII was used to indicate the growth rate of the execution time with the data set size for a naive "generate-and-test" sorting strategy. The results of sorting the large data set of Appendix V are shown in the following table.

| Sort <br> Name | CPU <br> Sec. |
| :---: | :---: |
| qsort <br> quisortx $\dagger$ | 24.4 |
| quisort | 4.283 |
| insort | 6097.0 |
| busort $\ddagger$ | 65.9 |
| sort | $\infty$ |

These results show an incredible amount of variation, although the strategy would prefer
$\dagger$ Failed, no explanation.
$\ddagger$ Failed, stack overflow
qsort on this size of list. The naive sort took 2811.8 seconds of CPU time for 9 elements (X38) which is indicative of why it has so much trouble. If the "random selection' picked this algorithm for a large list, execution time would be infinite for all practical purposes. Smaller lists exhibit more variation. For example the small ordered list benchmarks are shown in the next table.

| Sort <br> Name | CPU <br> Sec. |
| :---: | :---: |
| qsort | 0.267 |
| quisortx | 0.267 |
| quisort | 0.233 |
| insort | 0.100 |
| busort | 0.083 |
| sort | 0.133 |

Ordered lists were chosen because the implementations of quicksort are known to have trouble with them, and for this task it is obvious. The win on a small "random' list is not so dramatic, although there is some variance:

| Sort <br> Name | CPU <br> Sec. |
| :---: | :---: |
| qsort | 0.117 |
| quisortx | 0.133 |
| quisort | 0.117 |
| insort | 0.117 |
| busort | 0.633 |
| sort $^{*}$ | $\infty$ |

These simple tasks used a significant amount of CPU time, and the cases that exhibited

[^14]usable variance used little memory. For the larger lists, more memory was used, implying more copying. This is due to the deeper recursion.
Nguyen and Despain [Nguyen1988a] have compiled a series of tables containing memory access statistics for Prolog programs applied to standard benchmarks. The statistics indicate that much of the traffic is to stack data segments (Stack+Trail) ( $\min =14.6 \%$ for cmatch_strA , $\max =77.9 \%$ for cdeep_bakA, mean=46.1\% ); the rest is to the heap (mean $=23.4 \%$ ) and H2. H2 is used for control of parallel execution. Other statistics breaking the accesses down into reads and writes show that on average, the benchmarks write to the heap about 7.7 percent of the time. What these results indicate is that for almost all the benchmarks, there was more write traffic to the stack(s) than to the heap. This is important to our method, since stack writes will occur after the alt_spawn (), and hence will be local to whatever processor the process is executing on, rather than causing the extra traffic caused by the copy which must be made when the heap is written to. Writes to the heap are the major source of copying in the address space model used by our method. Thus, the copy-on-write technique can save by a factor of $\frac{100}{7.7}$ or about 13 in the amount of state which must be copied. Thus, we expect the combination of "copy-on-write" and "fastest-first" OR-parallel execution to produce a significant speedup for many programs.

### 4.9.3.2. Polyalgorithms

Polyalgorithms [Rice1968a, Symes1971a] have been suggested as a method for encapsulating a numerical analyst's knowledge into a system for solving numerical problems. The basic idea is that several methods are combined [Rice 1969a] along with information about the circumstances under which a method is likely to be successful. As different methods are tried and fail. information about the problem is built up until either there are no successful solutions, or a solution method succeeds (for example, discovering multiple zeros in a failing root-finder may be useful to the next solution method).

Our scheme could be used by creating artificial "alternatives" with the available solution methods. Each "alternative" trys a different solution method 'first,' to create alternative versions of the polyalgorithm. "Fastest first" scheduling could improve the response time properties of a system such as NAPSS [Rice1973a], especially since the performance of the system was perceived to be a problem [Rice1988a].

We should note that Traub [Traub1964a] mentioned such an idea (exploring multiple solution methods) as a direction for future work in his book on iterative methods.

### 4.9.3.3. Simulation

Simulation tasks are attractive applications for speculative methods, as some success has already been demonstrated by Jefferson [Jefferson1985a, Jefferson 1987a], in his "Virtual Time" scheme, which uses rollback-recovery to mask unsuccessful speculations. Note that we delete unsuccessful speculation by deleting the processes which contain its results. Consequently, we never roll back.

## 5. Related Work

Exploring alternatives in parallel is far from a new idea; hardware engineers looked to it as a way of maintaining pipeline ${ }^{18}$ utilization in some high-speed computers, most notably the IBM 360 Model 91 [Anderson1967a]. Their approach was to prefetch components of both possible branch paths until either the results of the conditional execution are available (in which case the correct stream can be chosen and the other discarded) or an irreversible side effect (such as instruction execution) would occur. One possible approach is to prefetch components of both possible branch paths until either the results of the conditional execution are available (in which case the correct stream can be chosen and the other discarded) or an irreversible side effect (such as instruction execution) would occur. This approach is analyzed by Riseman and Foster [Riseman 1972a], who conclude that speedups are possible if the correct branch is taken; their results are from a set of 7 programs run on the CDC 3600 computer. In the architecture setting, there is the problem that hardware is needed for each execution path and all paths must be explored, leading to a combinatorial explosion which affects even small values of the number of conditionals. Our management of side effects lets us go further.

Bernstein [Bernstein1980a] points out two problems with Hoare's CSP proposal. The first is that a programmer may want to have some method for choosing between alternatives with open guards; he solves this with priorities. The second is that CSP guards cannot contain output expressions; Bernstein solves this by use of the fact that the I/O primitives name a specific process: thus, guards specifying that process can be delayed until the specified process is ready. This is essentially a scheduling trick.

Version control systems such as SCCS [Rochkind1975a] and RCS [Tichy 1985a] use the idea of deltas to store multiple versions of data. More related to our predicates is the idea used in the PEDIT [Kruskal1984a] parametric line editor. Associated with each line

[^15]Difficulty is caused by conditional statements which can take one of two paths, thus inhibiting pre-fetching of instruction components. There are several approaches to resolving this difficulty; Lilja's [Lilja1988a] survey discusses the strategies.
of text is a set of parameters, hence the name parametric editor. These parameters are state variables, e.g. SYSTEM=UNIX, VERSION=SySV, et cetera. The line is selected for display if the mask set in the view of the file matches the settings of the state variables; thus, the viewer of a source program in a particular environment might see the source without the obscuring effect of various conditional compilation directives. Each setting of the state variables gives a distinct version, but in practice most of the text is shared between the versions.

Our method uses predicates to detect conflicts, but delays their resolution as long as is possible. Thus, it is optimistic in the sense that each timeline assumes that it will succeed. At each point where this success may come into question, it generates a predicate. These predicated processes are similar to the possibilities and dependencies discussed by Reed [Reed1978a] in his thesis: however, his NAMOS system was further from realization than the methods described here.

The notion of multiple alternatives is orthogonal to the transaction concept; if we view an alternative "block" as effecting a transaction on the system state, the specification is a description of how to accomplish the transaction reliably. It could also be viewed as a set of "competing' transactions, at most one of which will take effect.

One significant feature of our use of predicates is that there is as little waiting as possible in the system: each process which could only execute under a set of assumptions simply makes that set of assumptions, until some conflict with the correctness policies results. In other settings, such methods are called optimistic [Kung1981a, Strom1987a] because they assume that delay-causing or failure-causing conditions happen infrequently. Thus, normal operation is made cheap, at the expense of more expensive handling when the assumption is wrong. In our setting, the operant optimistic assumption is that the executing alternative is the one which will complete successfully. Thus, the predicates indicate that a process assumes that it will complete successfully; rather than waiting, it continues under that assumption. Strom and Yemini's [Strom1985a] dependency vectors behave much like our predicates.

Distribution of computation across several nodes offers attractive possibilities for both reliability and performance. Cooper [Cooper1985a] discusses the use of replicated distributed programs to take advantage of this potential. Cooper's CIRCUS [Cooper1984al system transparently replicates computations across several nodes in
order to increase reliability. Goldberg [Goldberg 1987a] has also discussed process replication, with a focus more on performance than fault tolerance. Replication is different than the problem we have examined, mainly because we cannot count on all the concurrent alternatives exhibiting the same behavior, e.g., reading and writing. For example, when managing I/O for replicated computations, only one read operation can be performed, and its results buffered for subsequent readers of the same data. Thus, idempotency of some source state can be forced through buffering.

Transparent replication can easily be combined with the use of parallel execution of several alternatives for increases in performance, reliability, or both.

## 6. Conclusions

This thesis has carefully examined problems for which there are multiple alternative solutions. When all such solutions are equally acceptable, the strategy of executing all alternative solutions concurrently has some benefits. In particular, this thesis has examined selecting the first successful computation as an approach to improve response time. We described scheduling strategies and memory management policies to ensure consistency and correctness. To restrain the growth in state required for concurrent execution of "Multiple Worlds" we suggested the use of "copy-on-write" storage managed in units of pages. External interactions are controlled by (1) an extension of the page management scheme to include slower storage, i.e.. files; and (2) an extension of the page management scheme to interprocess communication, where interacting processes are also 'copy-on-write.'

The results of this thesis are dependent on achieving an improvement in response time. We first identified the overhead of concurrent execution using the algorithms of Chapter 3. Using this model of overhead, we provide a measure of the performance improvement ( $P I$ ) which can be used to compare execution strategies. We use the measure of performance improvement to identify opportunities for response time improvement. The best situation (discussed formally in Chapter 2) for the approach presented here is one where:

- Alternatives require a significant amount of computation time, as encapsulated in $\tau\left(C_{\text {mean }}, \vec{x}\right)$.
- Each alternative changes a small amount of the state of the calling process, thus reducing the penalty $\tau$ (overhead).
- There is enough difference between the execution times of the alternatives that choosing the fastest and killing the others is worth the overhead of spawning the copies and deleting the slower siblings.

We made the observation that the speedup is dependent on both the execution time devoted to overhead and the dispersion exhibited by the execution times of the alternatives. The implication is that if overhead can be understood and controlled, there is an opportunity for speedup roughly proportional to the dispersion; thus superlinear speedups, where the execution time is less than $O\left(\frac{1}{N}\right)$ for $N$ processors, are possible under this
scenario.
Since the potential for speedup is sensitive to overheads, we examined these in Chapter 4. Copying is the major overhead in the creation and maintenance of concurrent executions. There had been no previous work examining the efficacy of "copy-on-write;' our results indicate that the technique is extremely effective in practice. The thesis provides several useful measurement tools to gather execution time data. These tools are used to adapt the measurements we reported here to new computers; thus, with a few measurements the domain for performance improvement using this method can be delimited. We have implemented a system which performs a 'remote fork,' which is similar to process migration with the exception that two processes exist when the operation is successful, rather than one migrated process. The measurements confirmed that the major cost in overhead was copying, and further, offered empirical proof that child processes could be "spawned" in a distributed execution environment. Response time is affected by the response times of several system components; we examined two subcomponents, the disk and the network, in Chapter 4. The final overhead was "sibling elimination,' which we modeled on a multiprocessing timesharing system. This should represent the worst case execution, and the results were encouraging, as they show the elimination to be remarkably cheap, and insensitive to process behavior. Thus, they are predictable and can be used in deciding whether to apply the concurrent execution scheme.

After identifying the opportunities with analytic work and measurements, some applications were. Distributed Execution of Recovery Blocks and parallel execution of logic programming languages are well-matched to the semantics of our execution scheme, and both can benefit from the higher performance. In each case, there are some restrictions imposed by the setting. Parallel implementation of logic programming languages provides an appropriate environment, because the computation is data-driven, and thus the execution time and control flow can vary greatly with the input. The way in which unification operates (as a 'sophisticated pattern matcher') leads to an overwhelming preponderance of read references made to page-managed memory: while a high percentage of references are writes, these are mainly to the stack, and thus locality should be high; stack 'growth'" can be handled locally, reducing copying. Many logic programs have a great degree of parallelism, so that appropriate opportunities must be identified with respect to the overheads implied by our scheme. In particular, coarsegrained parallelism is better to exploit than fine-grained parallelism. Major advantages
of the "Multiple Worlds'" scheme are that (1) it deals with side-effects other than variable binding, and (2) it can run efficiently on general-purpose hardware.

Distributed execution of recovery block alternates uses the "fastest-first" behavior in an attempt to find a rapid failure-free path through the computation. The restriction imposed by recovery blocks is that they are designed for fault-tolerance. Thus, there may be further requirements beyond fast execution time; by using our RB language and suggested modifications to the concurrent execution scheme which increases the amount of state available in a system to facilitate recovery.

Numerical Analysis is rich with examples for which the choice of solution method or the parameters of a particular solution method are free variables. The Jenkins-Traub polynomial root finding scheme was executed in parallel on multiprocessors; both speedup and fault-tolerance were observed across a selection of polynomials. The distribution of execution times did not show sufficient dispersion for a linear speedup, but the results were extremely promising. In particular, the overhead of the prototype '"Multiple Worlds" execution scheme was very small for this problem.

We also briefly mentioned applications to simulation. These example applications are from vastly different areas of computer science, and illustrate the general utility of the scheme. Throughput is clearly traded away for improved response time in our scheme, since the number of processors doing useful work (work that will eventually be used) at any given time will be $\leq 1$. This method may be particularly useful in real-time systems, where the bias is clearly towards response time, and the sibling elimination can be carried out asynchronously with respect to result delivery.

A question which remains is the effectiveness of exploring alternatives in parallel (which is speculative computation using serial algorithms) versus other methods for utilizing available processors. Now, for a problem which is easily divided into subproblems (although this division may not be complete, and thus the sequential portion will be subject to the "Amdahl limit"), the parallel execution will most likely show an improved response time proportional to the number of processors, which means that our method will improve performance when

$$
\frac{\tau\left(C_{\text {avg }}, \vec{x}\right)}{\tau\left(C_{\text {best }}, \vec{x}\right)}>\frac{\tau\left(C_{\text {avg }}, \vec{x}\right)}{N}
$$

for $N$ processors.
Does this happen? It's clear that it can happen, through choice of a particularly poor algorithm to parallelize. To focus the discussion, let's look at an example.

Problem: Given two sorted lists, L 1 and L2, find an element in their intersection.
Operators: There are two operators available for the problem solution. Operator element $(L, i)$ returns the $i$-th element of $L$. or an indication of failure if $i$ is greater than size $(\mathrm{L})$. A call to element $(\mathrm{L}, \mathrm{i})$ takes 1 unit of time. Operator $\operatorname{search}(L, x)$ is a boolean operator which returns success when $x$ is an element of L, and failure otherwise. Since L1 and L2 are sorted lists, for a list of size N , search $(\mathrm{L}, \mathrm{x})$ requires time $O\left(\log _{2}(N)\right)$ : to ease calculation we can assume the calculation takes exactly $\log _{10}(N)$.

Algorithm $F$ For each $x$ in $L 1$, search $L 2$ for $x$.
Algorithm B For each $x$ in L2, search $L 1$ for $x$.
If there are two processors, the data parallel algorithm will apply Algorithm F (or Algorithm B ; it's clear that it doesn't matter) to two half-lists. The two alternatives, Algorithm F and Algorithm B , are run in parallel. The following table compares the performance under various size assumptions for the lists:

| size(L1) | 10 | 100 | 100 |
| :---: | ---: | ---: | ---: |
| size(L2) | 100 | 100 | 10 |
| Parallel | 10 | 100 | 50 |
| F | 20 | 200 | 100 |
| B | 100 | 200 | 20 |

Of course, the operators are rather constrained; for example it's clear that if testing for the size of the lists was easy, then the smallest list would always be used for the linear search.

There are several issues which concern implementation on existing processors, or processors similar to these. First, it might be argued that memory contention caused by several algorithms accessing the same data simultaneously will cause shared memory multiprocessors or like architectures to be slowed. We expect that this will not be true in
practice. There are three reasons:

- There is typically a cache memory per processor in such a system, to allow the existing memory technology to support more processors.
- Once the first reference is made which forces a copy of the storage, the storage will be local and accessed through the local cache.
- When the instruction stream requires memory access on a cache miss, there will be less temporal locality exhibited by the distinct instruction streams. Thus, the likelihood of undesirable interactions between the degree of parallelism and such memory properties as the cache size, the memory interleaving factor, and the cache line size. are reduced. Whatever locality is exhibited by the algorithms can be taken advantage of in the local caches, while there will not be problems if the parallel processing is slightly out of lockstep, causing repeated 'miss-fill-replace-miss' sequences to occur.

These caching arguments can hold in a distributed setting as well. For example, rfork() used UNIX systems connected by a network file system. When shared memory is accessed over a network, each child process will be a "client," and there will be one "server," the parent or spawning process. As the children begin executing, state will be copied to their processors as necessary. If the parent has accessed the instructions or data, the state is likely to be in a buffer cache in main memory: such caches are useful in maintaining performance [Nelson 1988a] in distributed systems. Main memory caches of disk objects are most often managed in LRU fashion, so that temporal locality will result in common requests being serviced by the cache. The essence of the cache argument is that if the clients access different data, there will be less contention. If they access the same data, and there is more contention, fast caches will resolve the performance problem. This effect will hold true in networks, where main memory serves to cache slower portions of the storage hierarchy.

Thus, we expect that the contention is at most only slightly worse for the MISD case than the SIMD case. The point of this is that if there is an opportunity for superlinear speedup in the execution times of the alternatives, we will not be prevented from exploiting it by some architectural difficulty. This means that applying N processors to a problem can give superlinear speedup if the execution time variance is sufficient, i.e.,

$$
\tau\left(C_{\text {best }}, \stackrel{\rightharpoonup}{x}\right)<\frac{\tau\left(C_{\text {mean }}, \vec{x}\right)}{N}
$$

and that the execution time overhead of this approach is comparable to SIMD parallel approaches. The speedup is a property of the algorithms involved and the availability of processing units. If there are fewer processors than algorithms available, the "speedup" must be adjusted to reflect multiprocessing or delays if scheduling is non-preemptive. If there are exactly as many processors as needed, the speedup follows our analysis. If there are more processors than algorithms, there will be unused processors which some other form of parallel execution might be able to take advantage of. However, in several applications that we examined, the number of available tasks can be controlled dynamically. OR-parallelism in Prolog can keep the number of tasks proportional to the number of processors by using the Prolog interpreter to schedule sub-computations. The Jenkins-Traub algorithm's non-deterministic component allows for an arbitrary number of alternatives to be created.

## 7. Directions for Future Work

Future work can proceed from this thesis in several significant directions. In the theoretical domain, the relationship between the distribution of execution times and the speedup should be analyzed. For example, the properties of an execution time distribution necessary for a given speedup could be very useful; experimental work would determine how such a distribution might be generated.

Algorithmic work and analyses should compare optimistic and pessimistic strategies, e.g., by predicting abort costs, the number of aborts, and estimate delays due to waiting on locks. The synchronization scheme should be compared to other commit/synchronization mechanisms. Some measure of cost, such as the number of messages, could be used to refine and optimize the protocols.

Page-oriented management of binding environments for Prolog programs should be compared to the variable-oriented schemes which are currently used to maintain multiple binding environments.

The technique does not preclude having an alternative identify itself with a state variable. Systems could be constructed which apply the concurrent execution of alternatives to an appropriately chosen subset of the data in order to see which was fastest. Several trials could be done. The results could then be used to "predict'" which alternative would be fastest on the complete data set. Thus, the system could "learn' from samples to favor alternatives which perform better.

Finally, a complete system for concurrent execution of alternatives should be built. This thesis identifies the opportunities and provides enough of a roadmap so that the implementation should be straightforward; source code for the RB processor and fork() are already available. The completed system should provide a chance to develop better execution strategies and further, refine the boundaries of the domain for which performance improvements are possible.

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## 9. Appendix I: do fork.c

```
#include <errno.h>
#include <sys/param.h>
#ifndef NBPC
#define PAGE_SIZE 2048
#else
#define PAGE_SIZE NBPC
#endif
#ifndef NULL
#define NULL 0
#endif
main( argc, argv )
int argc;
char *argv[];
l
    int count = 0, heap_size = 0, pid, status;
    double atof(), write_fraction = 0.0,
        write_count = 0.0, write_size = 0.0;
    register char *ptr;
    char *malloc();
    extern int errno;
    if( argc > 1 )
    {
        count = atoi( argv[l] );
        if( argc > 2 )
        |
            heap_size = atoi( argv[2] );
                if( ( ptr = malloc( heap_size ) )
                    == (char *) NULL )
                        error( "Insufficient memory available. Exiting.\n" );
                if( argc > 3)
                l
                write_fraction = atof( argv[3] );
                if( write_fraction < 0.0 || write_fraction > 1.0)
                        error( "0.0 <= writes <= 1.0; Exiting.\n" );
                        write_size = write_fraction * (double) heap_size;
                }
        }
    }
    while( count > 0 )
    l
        switch( (pid = fork()) )
        |
        case -1: /* failed. If EAGAIN, wait. */
            if( errno == EAGAIN )
                wait( &status );
            break;
```

```
            case 0: /* child. make refs if needed, and exit */
                    while( write_count < write_size )
                        l
                    *ptr = ' ';
                    ptr = &ptr[PAGE_SIZE];
                    write_count += (double) PAGE_SIZE;
                    }
                    exit( 0):
            default:
                    count -= 1;
            l
        }
    exit( 0 );
}
error( string )
char *string;
l
    write( 2, string, strlen( string ) );
    exit(1);
}
```


## 10. Appendix II: netrand.c

```
#include <sys/types.h>
#include <sys/param.h>
#include <sys/times.h>
long times();
struct tms tbuf;
#include <stdio.h>
main( argc, argv )
int argc;
char *argv[];
l
    long start, diff;
    int count;
    FILE *fp;
    if( argc< 2 )
    l
        fprintf( stderr, "number required.\n" );
        exit( 1 );
    }
    count = atoi( argv[1] );
    if( count <= 0 )
        exit( 1 );
    if( argc > 2 )
    I
        fp = fopen( argv[2], "r" );
        if( fp == (FILE *) NULL )
        |
            perror( argv[1] );
            exit( l );
        }
    }
    else
        fp = stdin;
    start = times( &tbuf );
    random_read( fp, count );
    diff = times( &tbuf ) - start;
    printf( "elapsed time, %g seconds.\n", (float) diff / (float) HZ );
    exit( 0 );
|
random_read( fp, count )
FILE *fp;
int count;
|
```

```
    int i, fd, block_no, bytes;
    char page[NBPG_M320];
#ifdef DEBUG
    FILE *lfp;
    lfp = fopen( "/tmp/RANDBLOCKS", "w" );
    if( lfp == (FILE *) NULL )
        lfp = stdout;
#endif
        fd = fileno( fp );
        for( i = 0, block_no = random( times(&tbuf ) ) % count;
        i < count;
        block_no = random( 0 ) % count, i = i + 1 )
    {
        lseek( fd, block_no * sizeof(page), 0 );
        bytes = read( fd, page, sizeof(page));
#ifdef DEBUG
            fprintf( lfp, "block_no: %d, bytes read: %d\n",
                block_no, bytes );
    }
    fclose( lfp);
#else
    }
#endif
    return;
}
int
random( start )
int start;
{
    static int last_val;
    if( start != 0 )
        last_val = start;
    last_val = (17*last_val+123) % 65521;
    while( last_val < 0 )
        last_val += 65521;
    return( last_val );
}
```


## 11. Appendix III: Further Jenkins-Traub executions

### 11.1. Polynomial \#1




### 11.2. Polynomial \#2

Degree 43 Polynomial:

$$
\begin{aligned}
& z^{43}+ \\
& (1.0 e-0.140045104+1.0 e-0.161369678 i) \cdot z^{41}+ \\
& (1.0 e-0.1274642064+1.0 e-0.381680467 i) \cdot z^{40}+ \\
& (1.0 e-0.743821643+1.0 e-0.5810837248 i) \cdot z^{39}+ \\
& (1.0 e-0.16752870876+1.0 e-0.10043831986 i) \cdot z^{38}+ \\
& (1.0 e 1.41763432806+1.0 e 1.29981366628 i) \cdot z^{37}+ \\
& (1.0 e 1.23799211445+1.0 e-0.106982985377 i) \cdot z^{36}+ \\
& (1.0 e-0.189267174496+1.0 e-0.34407932181 i) \cdot z^{35}+ \\
& (1.0 e-0.152123766242+1.0 e 1.236186810121 i) \cdot z^{34}+
\end{aligned}
$$

$(1.0 e-0.198013830239+1.0 e 1.280742352179 i) \cdot z^{33}+$ $(1.0 e-0.342742413912+1.0 e 1.83786997252 i) \cdot z^{32}+$ $(1.0 e-0.37460901644+1.0 e-0.301901877085 i) \cdot z^{31}+$ $(1.0 e-0.195080181222+1.0 e-0.100059575279 i) \cdot z^{30}+$ $(1.0 e-0.94732085290+1.0 e-0.90223058904 i) \cdot z^{29}+$ $(1.0 e 1.27282343436+1.0 e-0.57440498715 i) \cdot z^{28}+$ $(1.0 e 1.24271198674+1.0 e-0.3724963179 i) \cdot z^{27}+$ ( $1.0 e 1.682559499+1.0 e-0.7237864131 i) \cdot z^{26}+$ $(1.0 e-0.1528494152+1.0 e-0.397912167 i) \cdot z^{25}+$ $(1.0 e 1.63349391+1.0 e 1.237887291 i) \cdot z^{24}+\quad(1.0 e 1.27878345+1.0 e 1.0897986 i) \cdot z^{23}+$ $(1.0 e 1.0802320+1.0 e-0.1101871 i) \cdot z^{22}+\quad(1.0 e-0.10+1.0 e 1.0 i) \cdot z^{21}+$ $(1.0 e 1.890+1.0 e-0.19491576 i) \cdot z^{20}+\quad(1.0 e-0.140045104+1.0 e-0.161369678 i) \cdot z^{19}+$ $(1.0 e-0.1274642064+1.0 e-0.381680467 i) \cdot z^{18}+$ $(1.0 e-0.743821643+1.0 e-0.5810837248 i) \cdot z^{17}+$ $(1.0 e-0.16752870876+1.0 e-0.10043831986 i) \cdot z^{16}+$ $(1.0 e 1.41763432806+1.0 e 1.29981366628 i) \cdot z^{15}+$ $(1.0 e 1.23799211445+1.0 e-0.106982985377 i) \cdot z^{14}+$ $(1.0 e-0.189267174496+1.0 e-0.34407932181 i) \cdot z^{13}+$ $(1.0 e-0.152123766242+1.0 e 1.236186810121 i) \cdot z^{12}+$ $(1.0 e-0.198013830239+1.0 e 1.280742352179 i) \cdot z^{11}+$ $(1.0 e-0.342742413912+1.0 e 1.83786997252 i) \cdot z^{10}+$ $(1.0 e-0.37460901644+1.0 e-0.301901877085 i) z^{9}+$ $(1.0 e-0.195080181222+1.0 e-0.100059575279 i) \cdot z^{8}+$ $(1.0 e-0.94732085290+1.0 e-0.90223058904 i) \cdot z^{7}+$ $(1.0 e 1.27282343436+1.0 e-0.57440498715 i) \cdot z^{6}+$ $(1.0 e 1.24271198674+1.0 e-0.3724963179 i) \cdot z^{5}+$ (1.0e $1.682559499+1.0 e-0.7237864131 i) \cdot z^{4}+$ $(1.0 e-0.1528494152+1.0 e-0.397912167 i) \cdot z^{3}+$ $(1.0 e 1.63349391+1.0 e 1.237887291 i) \cdot z^{2}+\quad(1.0 e 1.27878345+1.0 e 1.0897986 i) \cdot z^{1}+$ (1.0e $1.0802320+1.0 e-0.1101871 i)$


### 11.3. Polynomial \#3

Degree 16 Polynomial: $z^{16}+(199.80+33.20 i) \cdot z^{15}+(1766.930+588.940 i) \cdot z^{14}+$ $(9400.2460+4944.7240 i) \cdot z^{13}+\quad(32844.87470+24955.1830 i) \cdot z^{12}+$ $(78211.15150+83755.778560 i) \cdot z^{11}+$ $(120455.424807+314055.685381 i) \cdot z^{9}+$ $(103788.708048+151722.462694 i) \cdot z^{7}+$ $(124300.788642+335711.308326 i) \cdot z^{5}+$ $(13600.752190+201800.264386 i) \cdot z^{3}+$ $(125211.112583+194911.045162 i) \cdot z^{10}+$ $(27855.185034+327400.983162 i) \cdot z^{8}+$ $(168700.986162+135544.317038 i) \cdot z^{6}+$ $(36733.995257+333899.404113 i) \cdot z^{4}+$ $(16899.471245+76744.191752 i) \cdot z^{2}+$ $(6222.122192+17000.277923 i) \cdot z^{1}+(866.4078 .58+1688.238421 i)$


### 11.4. Polynomial \#4

| Degree | Polynomial | $z^{43}+$ | $(1.8900000-0.19491576 i) \cdot z^{42}+$ |
| :--- | ---: | ---: | ---: |
| $(-0.140045104-0.161369678 i) \cdot z^{41}+$ | $(-0.1274642064-0.381680467 i) \cdot z^{40}+$ |  |  |
| $(-0.743821643-0.5810837248 i) \cdot z^{39}+$ | $(-0.16752870876-0.10043831986 i) \cdot z^{38}+$ |  |  |
| $(1.41763432806+1.29981366628 i) \cdot z^{37}+$ | $(1.23799211445-0.106982985377 i) \cdot z^{36}+$ |  |  |
| $(-0.189267174496-0.34407932181 i) \cdot z^{35}+$ | $(-0.152123766242+1.236186810121 i) \cdot z^{34}+$ |  |  |
| $(-0.198013830239+1.280742352179 i) \cdot z^{33}+$ | $(-0.342742413912+1.83786997252 i) \cdot z^{32}+$ |  |  |
| $(-0.37460901644-0.301901877085 i) \cdot z^{31}+$ | $(-0.195080181222-0.100059575279 i) \cdot z^{30}+$ |  |  |
| $\left(-0.9473208529(-0.902230589() 4 i) \cdot z^{29}+\right.$ | $(1.27282343436-0.57440498715 i) \cdot z^{28}+$ |  |  |
| $(1.24271198674-0.3724963179 i) \cdot z^{27}+$ | $(1.682559499-0.7237864131 i) \cdot z^{26}+$ |  |  |
| $(-0.1528494152-0.397912167 i) \cdot z^{25}+$ | $(1.63349391+1.237887291 i) \cdot z^{24}+$ |  |  |
| $(1.27878345+1.0897986 i) \cdot z^{23}+$ | $(1.0802320-0.1101871 i) \cdot z^{22}+$ | $(-0.10+1.0 i) \cdot z^{21}+$ |  |


| $(1.890-0.19491576 i) \cdot z^{20}+$ | $(-0.140045104-0.161369678 i) \cdot z^{19}+$ |
| :--- | ---: |
| $(-0.1274642064-0.381680467 i) \cdot z^{18}+$ | $(-0.743821643-0.5810837248 i) \cdot z^{17}+$ |
| $(-0.16752870876-0.10043831986 i) \cdot z^{16}+$ | $(1.41763432806+1.29981366628 i) \cdot z^{15}+$ |
| $(1.23799211445-0.106982985377 i) \cdot z^{14}+$ | $(-0.189267174496-0.34407932181 i) \cdot z^{13}+$ |
| $(-0.152123766242+1.236186810121 i) \cdot z^{12}+$ | $(-0.198013830239+1.280742352179 i) \cdot z^{11}+$ |
| $(-0.342742413912+1.83786997252 i) \cdot z^{10}+$ | $(-0.37460901644-0.301901877085 i) \cdot z^{9}+$ |
| $(-0.195080181222-0.100059575279 i) \cdot z^{8}+$ | $(-0.94732085290-0.90223058904 i) \cdot z^{7}+$ |
| $(1.27282343436-0.57440498715 i) \cdot z^{6}+$ | $(1.24271198674-0.3724963179 i) \cdot z^{5}+$ |
| $(1.682559499-0.7237864131 i) \cdot z^{4}+$ | $(-0.1528494152-0.397912167 i) \cdot z^{3}+$ |
| $(1.63349391+1.237887291 i) \cdot z^{2}+$ | $(1.27878345+1.0897986 i) \cdot z^{1}+$ |
| $(1.0802320-0.1101871 i)$ |  |

20- M


## Time

$(\operatorname{Sec} \$)^{-}$

$$
5-
$$



### 11.5. Polynomial \#5

Degree 16 Polynomial: $z^{16}+(1998.0+332.0 i) \cdot z^{16}+(17669.30+5889.40 i) \cdot z^{14}+$ $(94002.460+49447.240 i) \cdot z^{13}+\quad(328448.7470+249551.830 i) \cdot z^{12}+$ $(782111.5150+837557.78560 i) \cdot z^{11}+\quad(1252111.12583+1949110.45162 i) \cdot z^{10}+$ $(1204554.24807+3140556.85381 i) \cdot z^{9}+$ $(1037887.08048+1517224.62694 i) \cdot z^{7}+$ $(278551.85034+3274009.83162 i) \cdot z^{8}+$ $(1243007.88642+3357113.08326 i) \cdot z^{5}+$ $(136007.52190+2018002.64386 i) \cdot z^{3}+$ $(1687009.86162+1355443.17038 i) \cdot z^{6}+$ $(367339.95257+3338994.04113 i) \cdot z^{4}+$ $(62221.22192+170002.77923 i) \cdot z^{1}+(8664.07858+16882.38421 i)$

## 20 -

|  | M |  | M |  | M |
| :---: | :---: | :---: | :---: | :---: | :---: |
| $15-\mathbb{W}$ | A İ m | M |  |  |  |

A
$\underset{10}{\text { Time }}$ (Secs)

A
II

Hh m
5 -


### 11.6. Polynomial \#6

Degree 21 Polynomial: $z^{21}+(8.90+19.491576 i) \cdot z^{20}+(140.045104+161.369678 i) \cdot z^{19}+$ $(1274.642064+381.680467 i) \cdot z^{18}+$ $(16752.870876+10043.831986 i) \cdot z^{16}+$ $(23799.211445+106982.985377 i) \cdot z^{14}+$ $(152123.766242+236186.810121 i) \cdot z^{12}+$ $(342742.413912+83786.997252 i) \cdot z^{10}+$ $(195080.181222+100059.575279 i) \cdot z^{8}+$ $(27282.343436+57440.498715 i) \cdot z^{6}+$ $(682.559499+7237.864131 i) \cdot z^{4}+$ $(743.821643+5810.837248 i) \cdot z^{17}+$ $(41763.432806+29981.366628 i) \cdot z^{15}+$ $(189267.174496+34407.932181 i) \cdot z^{13}+$ $(198013.830239+280742.352179 i) \cdot z^{11}+$ $(37460.901644+301901.877085 i) \cdot z^{9}+$ $(94732.085290+90223.058904 i) \cdot z^{7}+$ $(24271.198674+3724.963179 i) \cdot z^{5}+$ $(63.349391+237.887291 i) \cdot z^{2}+(27.878345+0.897986 i) \cdot z^{1}+(0.802320+1.101871 i)$

$$
20 \text { - }
$$

|  | M | M | M | M | M | M |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 15 | A |  |  |  |  |  |


|  | m | A | A | A |  |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- |
| Time <br> (Secs) |  |  |  | A |  |  |
|  |  | m |  | $\\|$ | m | $\\|$ | th

II
m
m
m
5 -


### 11.7. Polynomial \#7





### 11.8. Polynomial \#8

Degree 43 Polynomial:

$$
\begin{array}{lr}
z^{43}+ & (0.890+0.19491576 i) \cdot z^{42}+ \\
(0.1274642064+0.381680467 i) \cdot z^{40}+ & (0.140045104+0.161369678 i) \cdot z^{41}+ \\
(0.16752870876+0.10043831986 i) \cdot z^{38}+ & (0.743821643+0.5810837248 i) \cdot z^{39}+ \\
(0.23799211445+0.106982985377 i) \cdot z^{36}+ & (0.189267174496+0.34407932181 i) \cdot z^{35}+ \\
(0.152123766242+0.236186810121 i) \cdot z^{34}+ & (0.198013830239+0.280742352179 i) \cdot z^{33}+ \\
(0.342742413912+0.83786997252 i) \cdot z^{32}+ & (0.37460901644+0.301901877085 i) \cdot z^{31}+ \\
(0.195080181222+0.100059575279 i) \cdot z^{30}+ & (0.94732085290+0.90223058904 i) \cdot z^{29}+ \\
(0.27282343436+0.57440498715 i) \cdot z^{28}+ & (0.24271198674+0.3724963179 i) \cdot z^{27}+ \\
(0.682559499+0.7237864131 i) \cdot z^{26}+ & (0.1528494152+0.397912167 i) \cdot z^{25}+
\end{array}
$$


#### Abstract

$(0.63349391+0.237887291 i) \cdot z^{24}+$ $(0.27878345+0.0897986 i) \cdot z^{23}+$ $(0.0802320+0.1101871 i) \cdot z^{22}+$ $(0.10+0.0 i) \cdot z^{21}+$ $(0.890+0.19491576 i) \cdot z^{20}+$ $(0.140045104+0.161369678 i) \cdot z^{19}+$ $(0.743821643+0.5810837248 i) \cdot z^{17}+$ $(0.41763432806+0.29981366628 i) \cdot z^{15}+$ $(0.189267174496+0.34407932181 i) \cdot z^{13}+$ $(0.152123766242+0.236186810121 i) \cdot z^{12}+$ $(0.198013830239+0.280742352179 i) \cdot z^{11}+$ $(0.342742413912+0.83786997252 i) \cdot z^{10}+$ $(0.37460901644+0.301901877085 i) \cdot z^{9}+$ $(0.195080181222+0.100059575279 i) \cdot z^{8}+$ $(0.94732085290+0.90223058904 i) \cdot z^{7}+$ $(0.24271198674+0.3724963179 i) \cdot z^{5}+$ $(0.1528494152+0.397912167 i) \cdot z^{3}+$ $(0.27878345+0.0897986 i) \cdot z^{1}+(0.0802320+0.1101871 i)$


| $M$ | $M$ | $M$ | $M$ | $M$ | $M$ |
| :--- | :--- | :--- | :--- | :--- | :--- |

A
20 -

Time

## m

(Secs)
th

m
A
\|

$10-$

A A II II
m
m


## 12. Appendix IV: Source, Prolog Sorts

```
/*
    * Naive sort
    * Clocksin & Mellish, P. 155
    */
sort(L1,L2) :- permutation(L1,L2), sorted(L2), !.
permutation(L,[H|T]) :-
    append(V,[H|U],L),
    append(V,U,W),
    permutation(W,T).
permutation([],[]).
sorted(L) :- sorted(0,L).
sorted(_,[]).
sorted(N,[H|T]) :- N =< H, sorted(H,T).
/*
    * Insertion sort
    * Clocksin & Mellish, p. }15
    */
insort([],[]).
insort([X|L],M) :- insort(L,N), insortx(X,N,M).
insortx(X,[A|L],[A|M]) :- A =< X, !, insortx(X,L,M).
insortx(X,L,[X|L]).
/*
    * Bubble sort
    * Clocksin & Mellish, p. 156
    */
busort(L,S) :-
    append(X,[A,B|Y],L),
    B < A,
    append(X,[B,A|Y],M),
    busort (M,S).
busort(L,I).
/*
    * Quicksort #1 from
    * Clocksin & Mellish, p. }15
    */
split(H,[A|X],[A|Y],Z) :- A =< H, split(H,X,Y,Z).
split(H,[A|X],Y,[A|Z]) :- A > H, split(H,X,Y,Z).
split(_,[],[],[]).
quisort( [],[] ).
quisort( [H|T], S) :-
    split(H,T,A,B),
    quisort(A,Al),
```

```
    quisort(B,B1),
    append(Al, [H|Bl], S).
/*
    * Quicksort #2 from
    * Clocksin & Mellish, p.157
    */
quisortx([H|T],S,X) :-
    split(H,T,A,B),
    quisortx(A,S,[H|Y]),
    quisortx(B,Y,X).
quisortx([],X,X).
/*
    * Quicksort
    * from DEC-10 library
    */
qsort([X|L],RO,R) :-
    partition(L,X,L1,L2),
    qsort(L2,R0,R1),
    qsort(L1,[X|R1],R).
qsort([],R,R).
partition([X|L],Y,[X|L1],L2) :- X =< Y, !,
    partition(L,Y,L1,L2).
partition([X|L],Y,L1,[X|L2]) :- X > Y, !,
    partition(L,Y,L1,L2).
partition([],_,[],[]).
/* define append(). */
append([], L, L).
append( [H|T], L, [H|V] ) :- append(T,L,V).
```


## 13. Appendix V: Timings, Prolog Sorts, Large Lists

## statistics.

qsort $([4,4,8,6,7,9,1,6,6,1,1,9,0,4,3,3,3,4,7,9,4,9,5,9,4,7,0,3,1,1$, $7,0,7,5,9,2,1,9,5,1,2,9,5,7,0,7,7,1,0,8,7,3,7,3,3,3,1,7,9,3,1,4,3,3$, $1,1,0,9,2,3,5,3,1,4,0,5,7,1,9,4,1,6,3,6,5,2,4,0,9,0,7,5,1,8,1,1,8,2$, $8,5,0,4,9,5,5,7,1,8,3,6,1,5,9,0,0,5,7,0,9,9,1,7,9,1,9,1,4,9,7,3,3,7$, $3,5,2,7,8,9,1,4,3,6,4,0,2,0,4,6,3,8,4,2,6,0,2,7,2,6,3,7,5,4,8,7,5,2$, $7,6,4,3,4,1,4,9,7,5,6,9,2,0,2,0,2,0,0,0,7,9,3,4,6,2,5,5,6,6,6,6,1,5$, $1,6,6,5,1,5,2,5,8,2,9,1,9,3,9,3,1,3,4,3,3,9,1,2,2,2,2,6,1,4,6,2,1,2$, $4,2,0,1,4,1,5,3,3,6,5,0,3,7,2,8,6,8,3,3,6,6,2,9,2,1,1,8,6,2,9,0,4,2$, $3,5,4,7,7,8,9,6,6,3,7,8,3,7,8,5,4,6,7,8,9,3,0,1,2,6,3,8,0,4,1,0,8,2$, $1,4,3,7,8,5,4,8,7,3,9,8,8,6,6,4,8,7,9,9,2,3,4,1,1,7,9,9,4,8,5,0,4,3$, $3,8,6,6,7,7,8,1,2,5,5,9,4,5,4,1,3,4,7,2,4,6,5,2,4,6,2,3,1,1,9,4,8,8$, $1,4,0,1,3,1,6,0,7,1,6,2,8,4,2,7,2,8,1,7,1,3,0,4,2,2,4,7,8,6,9,1,8,5$, $6,3,1,2,0,0,1,9,2,3,3,3,6,9,8,9,6,6,9,3,3,5,4,4,3,6,1,6,2,9,3,9,1,3$, $1,1,0,4,1,7,3,0,9,5,6,5,0,1,6,9,4,6,6,2,7,5,4,5,5,8,8,7,5,6,4,4,3,4$, $5,1,9,1,2,6,9,2,7,9,6,0,0,6,9,3,5,5,1,8,0,9,2,7,1,9,5,6,4,5,0,2,6,4$, $2,9,4,0,9,2,8,5,7,4,1,0,8,2,8,3,5,3,5,1,1,8,8,2,7,5],[], X 10)$.
statistics.
quisortx $([4,4,8,6,7,9,1,6,6,1,1,9,0,4,3,3,3,4,7,9,4,9,5,9,4,7,0,3,1,1$, $7,0,7,5,9,2,1,9,5,1,2,9,5,7,0,7,7,1,0,8,7,3,7,3,3,3,1,7,9,3,1,4,3,3$, $1,1,0,9,2,3,5,3,1,4,0,5,7,1,9,4,1,6,3,6,5,2,4,0,9,0,7,5,1,8,1,1,8,2$, $8,5,0,4,9,5,5,7,1,8,3,6,1,5,9,0,0,5,7,0,9,9,1,7,9,1,9,1,4,9,7,3,3,7$, $3,5,2,7,8,9,1,4,3,6,4,0,2,0,4,6,3,8,4,2,6,0,2,7,2,6,3,7,5,4,8,7,5,2$, $7,6,4,3,4,1,4,9,7,5,6,9,2,0,2,0,2,0,0,0,7,9,3,4,6,2,5,5,6,6,6,6,1,5$, $1,6,6,5,1,5,2,5,8,2,9,1,9,3,9,3,1,3,4,3,3,9,1,2,2,2,2,6,1,4,6,2,1,2$, $4,2,0,1,4,1,5,3,3,6,5,0,3,7,2,8,6,8,3,3,6,6,2,9,2,1,1,8,6,2,9,0,4,2$, $3,5,4,7,7,8,9,6,6,3,7,8,3,7,8,5,4,6,7,8,9,3,0,1,2,6,3,8,0,4,1,0,8,2$, $1,4,3,7,8,5,4,8,7,3,9,8,8,6,6,4,8,7,9,9,2,3,4,1,1,7,9,9,4,8,5,0,4,3$, $3,8,6,6,7,7,8,1,2,5,5,9,4,5,4,1,3,4,7,2,4,6,5,2,4,6,2,3,1,1,9,4,8,8$, $1,4,0,1,3,1,6,0,7,1,6,2,8,4,2,7,2,8,1,7,1,3,0,4,2,2,4,7,8,6,9,1,8,5$, $6,3,1,2,0,0,1,9,2,3,3,3,6,9,8,9,6,6,9,3,3,5,4,4,3,6,1,6,2,9,3,9,1,3$, $1,1,0,4,1,7,3,0,9,5,6,5,0,1,6,9,4,6,6,2,7,5,4,5,5,8,8,7,5,6,4,4,3,4$, $5,1,9,1,2,6,9,2,7,9,6,0,0,6,9,3,5,5,1,8,0,9,2,7,1,9,5,6,4,5,0,2,6,4$, $2,9,4,0,9,2,8,5,7,4,1,0,8,2,8,3,5,3,5,1,1,8,8,2,7,5]$, [], X11).
statistics.
quisort $([4,4,8,6,7,9,1,6,6,1,1,9,0,4,3,3,3,4,7,9,4,9,5,9,4,7,0,3,1,1$, $7,0,7,5,9,2,1,9,5,1,2,9,5,7,0,7,7,1,0,8,7,3,7,3,3,3,1,7,9,3,1,4,3,3$, $1,1,0,9,2,3,5,3,1,4,0,5,7,1,9,4,1,6,3,6,5,2,4,0,9,0,7,5,1,8,1,1,8,2$, $8,5,0,4,9,5,5,7,1,8,3,6,1,5,9,0,0,5,7,0,9,9,1,7,9,1,9,1,4,9,7,3,3,7$, $3,5,2,7,8,9,1,4,3,6,4,0,2,0,4,6,3,8,4,2,6,0,2,7,2,6,3,7,5,4,8,7,5,2$, $7,6,4,3,4,1,4,9,7,5,6,9,2,0,2,0,2,0,0,0,7,9,3,4,6,2,5,5,6,6,6,6,1,5$, $1,6,6,5,1,5,2,5,8,2,9,1,9,3,9,3,1,3,4,3,3,9,1,2,2,2,2,6,1,4,6,2,1,2$, $4,2,0,1,4,1,5,3,3,6,5,0,3,7,2,8,6,8,3,3,6,6,2,9,2,1,1,8,6,2,9,0,4,2$, $3,5,4,7,7,8,9,6,6,3,7,8,3,7,8,5,4,6,7,8,9,3,0,1,2,6,3,8,0,4,1,0,8,2$, $1,4,3,7,8,5,4,8,7,3,9,8,8,6,6,4,8,7,9,9,2,3,4,1,1,7,9,9,4,8,5,0,4,3$, $3,8,6,6,7,7,8,1,2,5,5,9,4,5,4,1,3,4,7,2,4,6,5,2,4,6,2,3,1,1,9,4,8,8$,
$1,4,0,1,3,1,6,0,7,1,6,2,8,4,2,7,2,8,1,7,1,3,0,4,2,2,4,7,8,6,9,1,8,5$, $6,3,1,2,0,0,1,9,2,3,3,3,6,9,8,9,6,6,9,3,3,5,4,4,3,6,1,6,2,9,3,9,1,3$, $1,1,0,4,1,7,3,0,9,5,6,5,0,1,6,9,4,6,6,2,7,5,4,5,5,8,8,7,5,6,4,4,3,4$, $5,1,9,1,2,6,9,2,7,9,6,0,0,6,9,3,5,5,1,8,0,9,2,7,1,9,5,6,4,5,0,2,6,4$, $2,9,4,0,9,2,8,5,7,4,1,0,8,2,8,3,5,3,5,1,1,8,8,2,7,5], \mathrm{X} 12)$.
statistics.
insort ( $[4,4,8,6,7,9,1,6,6,1,1,9,0,4,3,3,3,4,7,9,4,9,5,9,4,7,0,3,1,1$, $7,0,7,5,9,2,1,9,5,1,2,9,5,7,0,7,7,1,0,8,7,3,7,3,3,3,1,7,9,3,1,4,3,3$, $1,1,0,9,2,3,5,3,1,4,0,5,7,1,9,4,1,6,3,6,5,2,4,0,9,0,7,5,1,8,1,1,8,2$, $8,5,0,4,9,5,5,7,1,8,3,6,1,5,9,0,0,5,7,0,9,9,1,7,9,1,9,1,4,9,7,3,3,7$, $3,5,2,7,8,9,1,4,3,6,4,0,2,0,4,6,3,8,4,2,6,0,2,7,2,6,3,7,5,4,8,7,5,2$, $7,6,4,3,4,1,4,9,7,5,6,9,2,0,2,0,2,0,0,0,7,9,3,4,6,2,5,5,6,6,6,6,1,5$, $1,6,6,5,1,5,2,5,8,2,9,1,9,3,9,3,1,3,4,3,3,9,1,2,2,2,2,6,1,4,6,2,1,2$, $4,2,0,1,4,1,5,3,3,6,5,0,3,7,2,8,6,8,3,3,6,6,2,9,2,1,1,8,6,2,9,0,4,2$, $3,5,4,7,7,8,9,6,6,3,7,8,3,7,8,5,4,6,7,8,9,3,0,1,2,6,3,8,0,4,1,0,8,2$, $1,4,3,7,8,5,4,8,7,3,9,8,8,6,6,4,8,7,9,9,2,3,4,1,1,7,9,9,4,8,5,0,4,3$, $3,8,6,6,7,7,8,1,2,5,5,9,4,5,4,1,3,4,7,2,4,6,5,2,4,6,2,3,1,1,9,4,8,8$, $1,4,0,1,3,1,6,0,7,1,6,2,8,4,2,7,2,8,1,7,1,3,0,4,2,2,4,7,8,6,9,1,8,5$, $6,3,1,2,0,0,1,9,2,3,3,3,6,9,8,9,6,6,9,3,3,5,4,4,3,6,1,6,2,9,3,9,1,3$, $1,1,0,4,1,7,3,0,9,5,6,5,0,1,6,9,4,6,6,2,7,5,4,5,5,8,8,7,5,6,4,4,3,4$, $5,1,9,1,2,6,9,2,7,9,6,0,0,6,9,3,5,5,1,8,0,9,2,7,1,9,5,6,4,5,0,2,6,4$, $2,9,4,0,9,2,8,5,7,4,1,0,8,2,8,3,5,3,5,1,1,8,8,2,7,5], \mathrm{X} 13)$.

## statistics.

busort ( $[4,4,8,6,7,9,1,6,6,1,1,9,0,4,3,3,3,4,7,9,4,9,5,9,4,7,0,3,1,1$, $7,0,7,5,9,2,1,9,5,1,2,9,5,7,0,7,7,1,0,8,7,3,7,3,3,3,1,7,9,3,1,4,3,3$, $1,1,0,9,2,3,5,3,1,4,0,5,7,1,9,4,1,6,3,6,5,2,4,0,9,0,7,5,1,8,1,1,8,2$, $8,5,0,4,9,5,5,7,1,8,3,6,1,5,9,0,0,5,7,0,9,9,1,7,9,1,9,1,4,9,7,3,3,7$, $3,5,2,7,8,9,1,4,3,6,4,0,2,0,4,6,3,8,4,2,6,0,2,7,2,6,3,7,5,4,8,7,5,2$, $7,6,4,3,4,1,4,9,7,5,6,9,2,0,2,0,2,0,0,0,7,9,3,4,6,2,5,5,6,6,6,6,1,5$, $1,6,6,5,1,5,2,5,8,2,9,1,9,3,9,3,1,3,4,3,3,9,1,2,2,2,2,6,1,4,6,2,1,2$, $4,2,0,1,4,1,5,3,3,6,5,0,3,7,2,8,6,8,3,3,6,6,2,9,2,1,1,8,6,2,9,0,4,2$, $3,5,4,7,7,8,9,6,6,3,7,8,3,7,8,5,4,6,7,8,9,3,0,1,2,6,3,8,0,4,1,0,8,2$, $1,4,3,7,8,5,4,8,7,3,9,8,8,6,6,4,8,7,9,9,2,3,4,1,1,7,9,9,4,8,5,0,4,3$, $3,8,6,6,7,7,8,1,2,5,5,9,4,5,4,1,3,4,7,2,4,6,5,2,4,6,2,3,1,1,9,4,8,8$, $1,4,0,1,3,1,6,0,7,1,6,2,8,4,2,7,2,8,1,7,1,3,0,4,2,2,4,7,8,6,9,1,8,5$, $6,3,1,2,0,0,1,9,2,3,3,3,6,9,8,9,6,6,9,3,3,5,4,4,3,6,1,6,2,9,3,9,1,3$, $1,1,0,4,1,7,3,0,9,5,6,5,0,1,6,9,4,6,6,2,7,5,4,5,5,8,8,7,5,6,4,4,3,4$, $5,1,9,1,2,6,9,2,7,9,6,0,0,6,9,3,5,5,1,8,0,9,2,7,1,9,5,6,4,5,0,2,6,4$, $2,9,4,0,9,2,8,5,7,4,1,0,8,2,8,3,5,3,5,1,1,8,8,2,7,5], \mathrm{X} 14)$.

## statistics.

/*

* can't be done; would take forever!!!
sort $([4,4,8,6,7,9,1,6,6,1,1,9,0,4,3,3,3,4,7,9,4,9,5,9,4,7,0,3,1,1$, $7,0,7,5,9,2,1,9,5,1,2,9,5,7,0,7,7,1,0,8,7,3,7,3,3,3,1,7,9,3,1,4,3,3$, $1,1,0,9,2,3,5,3,1,4,0,5,7,1,9,4,1,6,3,6,5,2,4,0,9,0,7,5,1,8,1,1,8,2$, $8,5,0,4,9,5,5,7,1,8,3,6,1,5,9,0,0,5,7,0,9,9,1,7,9,1,9,1,4,9,7,3,3,7$, $3,5,2,7,8,9,1,4,3,6,4,0,2,0,4,6,3,8,4,2,6,0,2,7,2,6,3,7,5,4,8,7,5,2$, $7,6,4,3,4,1,4,9,7,5,6,9,2,0,2,0,2,0,0,0,7,9,3,4,6,2,5,5,6,6,6,6,1,5$,

[^16]
## 14. Appendix VI: Sort Performance, Small Lists

## statistics.

qsort ( $[4,4,8,6,7,9,1,6,6,1,1,9,0],[], \times 30)$.
statistics.
quisortx $([4,4,8,6,7,9,1,6,6,1,1,9,0], \mathrm{X} 31,[])$.
statistics.
quisort ( $[4,4,8,6,7,9,1,6,6,1,1,9,0], \times 32$ ).
statistics.
insort ( $[4,4,8,6,7,9,1,6,6,1,1,9,0], \mathrm{x} 33$ ).
statistics.
busort ( $[4,4,8,6,7,9,1,6,6,1,1,9,0], \times 34)$.
statistics.
sort $([4,4,8,6,7,9,1,6,6,1,1,9,0], X 35)$.
statistics.

## 15. Appendix VII: Naive Sort Performance

```
statistics.
sort( [4], X30).
statistics.
sort( [4,4], X31).
statistics.
sort( [4,4,8], X32).
statistics.
sort( [4,4,8,6], X33).
statistics.
sort( [4,4,8,6,7], X34).
statistics.
sort( [4,4,8,6,7,9], X35).
statistics.
sort( [4,4,8,6,7,9,1], X36).
statistics.
sort( [4,4,8,6,7,9,1,6], X37).
statistics.
sort( [4,4,8,6,7,9,1,6,6], x38).
statistics.
sort( [4,4,8,6,7,9,1,6,6,1], X39).
statistics.
```


## 16. Appendix VIII: Performance on Small, Ordered Lists

```
statistics.
qsort( [0,1,2,3,4,5,6,7,8,9,10,11,12], [], x30).
statistics.
quisortx( [0,1,2,3,4,5,6,7,8,9,10,11,12], X31, []).
statistics.
quisort( [0,1,2,3,4,5,6,7,8,9,10,11,12], x32).
statistics.
insort( [0,1,2,3,4,5,6,7,8,9,10,11,12], x33).
statistics
busort( [0,1,2,3,4,5,6,7,8,9,10,11,12], X34).
statistics
sort( [0,1,2,3,4,5,6,7,8,9,10,11,12], X35).
statistics
```


## 17. Appendix IX: Program to estimate memory speeds

```
/*
    * program to estimate memory speeds.
    * One argument: size of memory to {mem|b)copy, once.
    * Strategy:
    * We page it in via the first lcop.
    * The loop is set up to page, and not utilize the CPU's data cache.
    * Then, the time is obtained, we do the copy (in place) and,
    * the resulting time is calculated.
    */
#include <stdio.h>
#include <sys/types.h>
#include <sys/param.h>
#include <sys/times.h>
#include <errno.h>
#include <memory.h>
#ifndef NBPC
#define PAGE_SIZE 2048
#else
#define PAGE_SIZE NBPC
#endif
#ifndef NULL
#define NULl 0
#endif
main( argc, argv )
int argc;
char *argv[];
l
    int size;
    char *pl, *p2, *malloc();
    struct tms tbl, tb2;
    long clock, times();
    if( argc <= 1)
        error( "usage: mem_speed size\n" );
    size = atoi( argv[l] );
    if( (pl = malloc( size ) ) == (char *) NULL )
        error( "mem_speed: can't allocate memory.\n" );
    for( p2 = p1; p2 < &pl[size]; F2 = &p2[PAGE_SIZE] )
        *p2 = ' ';
    p2 = pl;
    clock = times( &tbl );
```

```
    memcpy( p2, p1, size );
    clock = times( &tb2 ) - clock;
    printf( "Real: %.2f, User: %.2f, System: %.2f\n",
        (1.0* (double)clock)/(1.0* (double)HZ),
        (1.0*(tb2.tms_utime-tb1.tms_utime))/(1.0*HZ),
        (1.0*(tb2.tms_stime-tbl.tms_stime))/(1.0*HZ) );
    exit(0);
l
error( string )
char *string;
l
    fprintf( stderr, "%s", string );
    exit( 1 );
}
```


## 18. Appendix X: Lower bound affects dispersion

```
#include <stdio.h>
#include <math.h>
#include <sys/types.h>
#include <sys/times.h>
#include <sys/time.h>
#define MILLION 1000000
#define TRUE 1
#define FALSE 0
#define MAX_SIZE 100000
#define CCOST 5 /* cost of comparison */
int S[MAX_SIZE],
    Ticks[MAX_SIZE],
    P[MAX_SIZE],
    COPy[MAX_SIZE],
    SSize = -1, PSize = -1;
int Perm = FALSE;
int Least = FALSE;
/*
    * Program to test performance of various quicksorting variations,
    * but mostly the application of randomness to partition element
    * selection.
*
* Options:
* -p: generate a permutation to sort rather than a random (might
* contain dupiicates) list.
* -s: specifies size of permutation or list to generate. Limited
* to value of symbol MAX_SIZE.
* -c: count of attempts to sort the list.
* -l: use element of list with smallest index as the pivot rather
* than selecting partition at random: see mysort(), below.
*
*/
main( argc, argv )
int argc;
char *argv[];
l
    double total, scaler, dSize, floor(), drand48();
    register int i, j, k;
    int min, max;
    long clock;
    struct timeval tvi;
    struct timezone tz;
    while( argc > 1 )
    l
        if( argv[argc-1][0] != '-' )
                usage():
```

```
    switch( argv[argc-1][1] )
    |
    case 's':
        PSize = atoi( &argv[argc-1][2] );
        if( PSize < 1 || PSize > MAX_SIZE )
            usage():
        break;
    case 'c':
        SSize = atoi( &argv[argc-1][2] );
        if( SSize < l || SSize > MAX_SIZE )
            usage();
        break:
    case 'p': /* use random permutation rather than
                * random list
                */
        Perm = TRUE;
        break;
    case 'l': /* use element of smallest index in sort
            * rather than a random element
            */
        Least = TRUE;
        break;
    default:
        usage();
    |
    argc -= 1;
}
if( PSize == -1 )
    PSize = MAX_SIZE;
if( SSize == - ) 
    SSize = MAX_SIZE;
dSize = (double) PSize;
gettimeofday( &tvl, &tz );
clock = MILLION*tvi.tv_sec+tvi.tv_usec;
srand48( clock );
if( Perm == TRUE )
1
    for( i = 0; i < PSize; i += l )
        P[i] = i;
    for( i = 0; i < PSize; i += l )
    {
        k = (int) floor( drand48()*dSize );
        j = P[i]; P[i] = P[k]; P[k] = j;
    }
}
```

```
    else
    l
        for( i = 0; i < PSize; i += 1)
        P[i] = (int) floor( drand48()*dSize );
    }
    for( i = 0; i < SSize; i += 1 )
    |
        for( j = 0; j < PSize; j += 1)
            Copy[j] = P[j];
        gettimeofday( &tvl, &tz );
        clock = MILLION*tvl.tv_sec+tv1.tv_usec;
        Ticks[i] = clock;
        mysort( Copy, 0, PSize-1 );
        gettimeofday( &tvl, &tz );
        clock = MILLION*tvl.tv_sec+tvi.tv_usec;
        Ticks[i] = clock - Ticks[i];
    l
    /* dump statistics */
    scaler = 1.0/((double) MILLION);
    min = max = Ticks[0];
    /* we pre-scale Ticks to prevent overflow */
    total = scaler*Ticks[0];
    for( i = 1; i < SSize; i += 1 )
    l
        if( Ticks[i] < min )
            min = Ticks[i];
        if( Ticks[i] > max )
            max = Ticks[i];
        total += scaler*Ticks[i];
    }
    printf( "min: %g sec, max: %g sec, avg: %g sec.\n",
        scaler* (double) min,
        scaler* (double) max,
        total / (double) SSize );
    exit( 0 );
}
usage()
|
    fprintf( stderr, "Usage: rand_qsort [-ssize] [-ccount] [-1] [-p]\n" );
    fprintf( stderr, "size & count <= %d\n", MAX_SIZE );
    exit(1 );
J
/* implementation of quicksort which randomizes.
    * There are some special tricks applied here which makes the
```

```
* code more complex (and faster) than many published versions.
* In particular, if there are more than one occurrence of the
* partitioning element, they create a "center" to the array
* which must be removed in order for the partitioning to be correct.
* "Software Tools" quicksort avoids this problem by always
* choosing the last element as the partition, and pushing
* other elements up against it.
*
* options: Least == TRUE; choose least element rather than
* a random element.
*/
mysort( array, start, finish )
int array[], start, finish;
l
    int i, random, pivot, hi, lo, tmp, exchanges;
    double drand48(), floor();
    if( Least == TRUE )
        random = start;
    else
        random = start + (int)floor( drand48()*( (double)(finish-start+1) ) );
    pivot = array[random];
    10 = start; /* upper bnd, current set of values < pivot */
    hi = finish; /* lower bnd, current set of values < pivot */
    while( lo < hi )
    |
        while( (lo < hi) && (compare( array[lo], pivot ) < 0 ) )
        10 = lo+1 ;
        while( (lo < hi) && ( compare( array[hi], pivot ) > 0 ) )
        hi = hi-1 ;
        if( lo < hi )
        |
                /* this block of code insures that all of
                * the elements with value == pivot are
                * bunched and ordered correctly.
                */
                if( (compare( array[hi], pivot ) == 0 ) &&
                ( compare( array[lo], pivot ) == 0 ))
            1
                exchanges = 0;
            for( i = lo+1; i <= hi-1; i += 1)
                |
                    if( compare( array[i], pivot ) < 0 )
                    l
                        tmp = array[lo);
                        array[lo] = array[i];
                        array[i] = tmp;
```

```
                    10 += 1 ;
                    ++exchanges;
                }
                else if( compare( array[i], pivot ) > 0 )
                l
                    tmp = array[hi];
                    array[hi] = array[i];
                        array[i] = tmp;
                    hi -= 1 ;
                    ++exchanges;
                }
            |
            if( exchanges == 0 )
                        goto recurse;
            }
            else
            l
                        tmp = array[lo];
                        array[lo] = array[hi];
                        array[hi] = tmp;
            J
        }
    1
recurse:
    if( lo-start > 1 )
        mysort( array, start, lo-1 );
    if( finish-hi > 1 )
        mysort( array, hi+1, finish );
    return;
}
/*
* this makes comparison costs significant compared to
* random # generation costs.
* CCOST can be altered to change significance of
* comparison costs.
*/
int
compare( i, j )
int i, j;
{
    int waste, diff;
    double drand48(), floor();
    for( waste = 0; waste <= ccosT; waste += 1 )
        diff = (int)floor( drand48()*( (double)(waste-j) ) );
    diff = i - j;
    if(diff< < )
        return( -1 ):
```

```
    if( diff > 0 )
        return( l );
    return( 0 );
}
$for i in 1 5 10 50 100 500
> do
> rand_qsort -s500 -c{{i}
> done
min: 4.88386 sec, max: 4.88386 sec, avg: 4.88386 sec.
min: 4.60353 sec, max: 5.2892 sec, avg: 4.78954 sec.
min: 4.5822 sec, max: 5.63769 sec, avg: 4.96763 sec.
min: 4.68141 sec, max: 6.8771 sec, avg: 5.49659 sec.
min: 4.58186 sec, max: 12.6386 sec, avg: 5.52683 sec.
min: 4.45218 sec, max: 14.1808 sec, avg: 5.46629 sec.
```


## 19. Appendix XI: do_elim Script

```
if [ ! -f do_elim ]
then
    echo "Making do_elim."
    make do_elim
fi
if [ ! -f do_elim ]
then
    echo "No do_elim. Exiting."
    exit 1
fi
echo "size do_elim:"
size do_elim
if [ -f/tmp/niceit ]
then
    rm -f /tmp/niceit
    exec nice -20 script
fi
for Groups in 0 1
do
    for Files in 0 5 10 15
    do
        for Asynch in 0 1
        do
            for Work in 0 1
            do
                for Size in 0 1000 3162 10000 31622 100000
                do
                    for Dirty in 0 1
                        do
                        for Procs in 1 2 4 8 16
                        do
                                Output=`do_elim\
                        -g$(Groups)\
                        -f${Files}\
                        -s$(Size)\
                        -w${Work}\
                        -aS{Asynch}\
                        -dS{Dirty}\
                        -p${Procs:'
                                echol
                                    "do_elim\
                            -g${Groups)\
                        -f${Files}\
                        -s$(Size)\
                        -w${Work}\
                        -aS{Asynch}\
                        -d$(Dirty)\
                        -ps{Procs}:\
                            ${Output}" |\
                                sed -e 's/ */ /g'
```

$$
\begin{aligned}
& \text { done done } \\
& \text { done } \\
& \text { done } \\
& \text { done }
\end{aligned}
$$

## 20. Appendix XII: do_elim.c

```
#include <errno.h>
#include <sys/param.h>
#include <sys/types.h>
#include <sys/times.h>
#include <sys/signal.h>
#include <sys/stat.h>
#ifndef NPROC
#define NPROC 100
#endif
#ifndef NBPC
#define PAGE_SIZE 2048
#else
#define PAGE_SIZE NBPC
#endif
#ifndef NULL
#define NULL 0
#endif
#define EOS '\0'
#define EVER ;;
#define HUGE 0x10000000
#define TRUE l
#define FALSE 0
#define TMP_PREFIX "/tmp/d_eFXXXXXX"
#define REP_COUNT 100
int
    ProcIdTable[NPROC],
    Groups = 0,
    Files = 0,
    Size = 0,
    Nork = 0,
    Asynch = 0,
    Dirty = 0,
    Procs = 0;
char *TmpEileNames[NOFILE];
/*
    * do_elim:
    * To evaluate costs of sibling elimination.
    * Eliminates siblings with a SIG_INTR signal (no dump)
    *
    * flags:
    * -g[01]: 1-use process group feature of kill(). Default: 0
    * -p[n]: # of processes to spawn. Default: 0
    * -s[n]: bytes of memory to allocate. Default: 0
    * -f[n]: open files per process. Default: 0
```

```
    * -w[01]: work mix: sleep vs. sleep + busy idle loop. Default: 0
    * -a[01]: asynchronous: (kind of hard to measure!). Default: 0
    * -d[01]: dirty all pages (to defeat c-o-w management). Default: 0
    */
main( argc, argv )
int argc;
char *argv[];
{
    struct tms tbl, tb2;
    long clock, times(), real_t, user_t, sys_t;
    double scaler;
    int i;
    char *osbrk, *sbrk();
    for( ++argv, --argc; argc > 0; ++argv, --argc )
    {
        if( argv[0][0] != '-' )
            usage();
        switch( argv[0][1] )
        l
        case 'g':
            Groups = getnum( &argv[0][2], 0, 1 );
            break;
        case 'p':
            Procs = getnum( &argv[0][2], 0, NPROC );
            break;
        case 's':
            Size = getnum( &argv[0][2], 0, HUGE);
            break;
            case 'f':
                Files = getnum( &argv[0][2], 0, NOFILE );
                break;
            case 'w':
                Work = getnum( &argv[0][21, 0, 1 );
                break;
            case 'a':
                Asynch = getnum( &argv[0][2], 0, 1 );
                break;
            case 'd':
                Dirty = getnum( &argv[0][2], 0, 1 );
                break;
            default:
                usage();
                break;
            }
        |
```

```
    /* set up new process group so that our parent not killed */
    if( setpgrp() < 0)
        fail( "Can't set new process group.\n" );
    /* timing phase; do repetitions help? */
    for( i = 0, real_t = user_t = sys_t = 0.0;
        i < REP_COUNT;
        i += 1)
    I
        startup();
        clock = times( &tb1 );
        eliminate();
        clock = times( &tb2 ) - clock;
        real_t += clock;
        user_t += (tb2.tms_utime-tbl.tms_utime);
        sys_t += (tb2.tms_stime-tb1.tms_stime);
        cleanup();
    I
    scaler = 1.0/(((double) HZ));
    printf( "Real: %.3f, User: %.3f, System: %.3f\n",
    ((double) real_t)*scaler,
    ((double) user_t)*scaler,
    ((double) sys_t)*scaler );
    exit(0);
}
int
getnum( str, min, max )
char *str;
int min, max;
{
    int val;
    val = convt( str );
    if( val < min || val > max )
        usage();
    return( val ):
}
usage()
{
    fail( "Usage: do_elim [-G] [-p] [-s] [-f] [-w] [-d] [-a]\n" );
|
int
```

```
convt( str )
char *str;
{
    int i;
    for( i = 0; *str != EOS; ++str )
    {
            if( *str >= '0' && *str <= '9' )
                i = 10*i + (*str - '0' );
            else
                    return( -1 ); /* invalid char. */
    |
    return( i );
}
startup()
l
    char *ptr, *emalloc(), *temp_file();
    int i, pid;
    ptr = emalloc( Size );
    for( i = 0; i < Files; i += 1 )
    {
            TmpFileNames[i] = temp_file();
        if( creat( TmpFileNames[i], 0 ) < 0 )
        |
            perror( TmpFileNames[i] );
            fail( "Can't create temporary file.\n" );
        }
    }
    /* set up signals so message not ignored */
    signal( SIGTERM, SIG_DFL ):
    for( i = 0; i < Procs; i += 1 )
    l
        switch( (pid = fork()) )
        l
        case -1:
            fail( "Can't fork.\n" );
            break;
        case 0: /* in child. Dirty and Work if necessary */
                if( Dirty)
                write_it( ptr, Size ):
            for( EVER )
            {
                if( Work )
                        iterate( 10000);
                        sleep( l );
                |
                break;
```

```
            default:
                ProcIdTable[i] = pid;
                break;
            |
        }
    /* only the Parent should ever be able to get here */
    free( ptr );
    return;
)
iterate( count )
register int count;
{
    while( count-- )
        ;
        return;
}
eliminate()
1
    int i, status;
    if( Groups )
    1
        /* set up signals so *we* don't get eliminated */
        signal( SIGTERM, SIG_IGN );
        if( kill( 0 , SIGTERM ) < 0 )
                fail( "Group kill failed.\n" );
    }
    else
    l
        for( i = 0; i < Procs; i += 1 )
        |
            if( kill( ProcIdTable[i], SIGTERM ) < 0 )
                fail( "Kill of proc failed.\n" );
        }
    }
    if( Asynch )
        return;
    else
    {
        for( i = Procs; i < 0; i = i-1 )
        {
            if( wait( &status ) < 0 )
            l
                if( errno == ECHILD )
                    continue:
                    else
                        fail( "Wait failed, !ECHILD\n" );
            }
                if( (0xFFFF & status) := SIGTERM )
```

```
                |
                    printf( "status: 0x%y\n", status );
                    fail( "Process terminated, wrong reason.\n" );
                    }
            }
    }
    return;
1
int
cleanup()
l
    int i;
    for( i = 0; i < Files; i += 1)
    1
        unlink( TmpFileNames[i] );
        free( TmpFileNames[i] );
        close( i+3 ): /* ??? */
    1
    while( wait( &i) >= 0 )
        ;
    if( errno != ECHILD )
        fail( "Cleanup: Wait failed, != ECHILD\n" );
    return;
}
char *
strsave( str)
char *str;
l
    char *p, *emalloc();
    p = emalloc( strlen( str ) + 1 );
    strcpy( p, str );
    return( p );
}
char *
temp_file()
{
    char *s, *strsave():
    void unique_temp();
    s = strsave( TMP_PREFIX );
    unique_temp( s );
    return( s );
}
void
unique_temp( s )
```

```
char *s;
|
    static long random;
    long work:
    char *p;
    register i;
    p =&s[strlen(s )-6];
    do
    l
        random = (random + 32647 + getpid() ) * 32653;
        work = random;
        for( i=0; i<6; i++ )
        |
            p[i] = "abcdefghijklmnopqrstuvwxyz012345"[ work & 0x1F ];
            work = work >> 5;
        }
    | while( exists( s ) == TRUE );
    return;
}
int
exists( name )
char *name;
l
    struct stat sb;
    int ret;
    extern errno;
    ret = stat( name, &sb );
    if( ret < 0 )
        if( errno == ENOENT )
            return( FALSE );
    return( TRUE );
l
char *
emalloc( size )
unsigned int size;
{
    char *ptr, *malloc();
    if( (ptr = malloc( size)) == (char *) NULL )
    l
        fail( "No Memory. bailing out!\n" );
    I
    return( ptr );
}
```

```
write_it( mem, size )
char *mem;
int size;
|
    double write_count = 0.0, write_size = 0.0;
    write_size = 1.0 * (double) size;
    while( write_count < write_size )
    l
            *mem = EOS;
            mem = &mem[PAGE_SI2E];
            write_count += (double) PAGE_SIZE;
    }
    return;
}
fail( string )
char *string;
{
    write( 2, string, strlen( string ) );
    exit( 1 );
}
```


## 21. Appendix XIII: "C' version of Jenkins-Traub algorithm

```
/*
    *
    * This is the Jenkins-Traub root-finding
    * algorithm from CACM, February 1972, V 15, Number 2,
    * pages 97-99. It is listed as ''Algorithm 419:
    * Zeros of a Complex Polynomial''
    * The algorithm has been recoded in "C",
    * which necessitated some changes.
    * To activate the debugging statements,
    * compile with -DEBUG.
    *
    * the machine constants have been changed to
    * reflect the IEEE floating point standard,
    * rather than the IBM/360.
    * a minor bug in cmod() was repaired; it had
    * divided by zero if given the complex origin
    * as an argument.
    * added angle (in radians) as an argument to cpoly().
    * all changes noted in ACM corrigendum (CACM 3/74)
    * have been applied.
    * the number of fixed and variable shifts has been
    * increased so that the algorithm is more effective
    * at multiple roots.
    *
    * if you find errors, contact me, Jonathan M. Smith,
    * at 450 Computer Science, Columbia University, NY, NY 10027
    * or jms@close.cs.columbia.edu, on the ARPAnet
    *
    * remainder is straight out of listing:
    * finds the zeros of a complex polynomial.
    * opr, opi - double precision vectors of real and
    * imaginary parts of the coefficients in
    * order of decreasing powers.
    * zeror, zeroi - output double precision vectors of
    * real and imaginary parts of the zeros.
    * fail - output logical parameter, true only if
    * leading coefficient is zero or if cpoly
    * has found fewer than degree zeros.
    * the program has been written to reduce the chance of overflow
    * occurring. if it does occur, there is still a possibility that
    * the zerofinder will work provided the overflowed quantity is
    * replaced by a large number.
    */
/* includes */
#include <math.h>
#include <stdio.h>
/* definitions */
#define MAX_DEGREE 50 /* biggest polynomial it can solve */
```

```
#define TRUE 1
#define FALSE 0
#define dabs(_x) ((_x<0.0)?(-_x):(_x))
#ifndef M_SQRT2
#define M_SQRT2 1.41421356237309504880
#define M_SQRT1_2 0.70710678118654752440
#endif
/* Globals */
double smalno, sr, si, tr, ti, pvr, pvi, are, mre, eta, infin,
base, errev(), cmod(), scale(), cauchy(), sqrt(), cos(), sin(),
pr[MAX_DEGREE],
pi[MAX_DEGREE],
hr[MAX_DEGREE],
hi[MAX_DEGREE],
qpr[MAX_DEGREE],
qPi[MAX_DEGREE],
qhr[MAX_DEGREE],
qhi[MAX_DEGREE],
shr[MAX_DEGREE],
shi[MAX_DEGREE];
int nn;
cpoly(opr,opi,degree,zeror,zeroi, fail,angle)
double *opr, *opi, *zeror, *zeroi, angle;
int *fail, degree;
l
    double xx,yy,cosr,sinr,xxx,zr,zi,bnd;
        int i, conv, cnt1, cnt2, idnn2;
    mcon();
    are = eta;
    mre = 2.0*M_SQRT2*eta;
#ifdef EBUG
    printí( "mre=%e, are=%e\n", mre, are );
#endif
    xx = M_SQRT1_2; /* 0.70710678 */
    yy = -xx;
    cosr = cos( angle ); /* -.069756474 */
    sinr = sin( angle ); /* .99756405 */
    *fail = FALSE;
    nn = degree+1;
    if(opr[0] != 0.0 || opi[0] != 0.0 ) /* fail if lead coeff. zero */
    l
        /* remove any zeros at origin */
        while( !( opr[nn-1] != 0.0 || opi[nn-1] != 0.0 ) )
        {
            idnn2 = degree-nn+2;
                zeror[idnn2-1] = 0.0;
                zeroi[idnn2-1] = 0.0;
                    nn = nn - 1;
```

```
    }
}
else
|
    *fail = TRUE;
    return;
I
/* copy coefficients */
for( i = 1; i <= nn; i += l )
l
    pr[i-1] = opr[i-1];
    pi[i-1] = opi[i-1];
    shr[i-1] = cmod(pr[i-1], pi[i-1]);
}
/* scale the polynomial */
bnd = scale( shr );
if( bnd != 1.0 )
l
    for( i=1; i <= nn; i += l )
    l
            pr[i-1] = bnd*pr[i-1];
            pi[i-1] = bnd*pi[i-1];
        1
l
/* start the algorithm */
while( nn > 2 )
l
    for( i=1; i <= nn; i t= 1)
    l
        shr[i-1] = cmod(pr[i-1], pi[i-1]);
    }
    bnd = cauchy( shr, shi);
    /* 2 major passes, different sequences of shifts */
    for( cnt1 = 1; cntl <= 2; cntl += 1 )
    l
        noshft( 5 ); /* first stage, no shift */
        /* inner loop to select a shift */
        for( cnt2 = 1; cnt2 <= 9; cnt2 += 1 )
        l
                /* shift is chosen with modulus bnd
                    * and amplitude rotated by angle
                        * degrees from the previous shift
                        */
                xxx = cosr*xx-sinr*yy;
                yy = sinr*xx+cosr*yy;
                xx = xxx;
                sr = bnd*xx;
                si = bnd*yy;
```

```
#ifdef EBUG
                printf( "shift: %g+ %gi\n",
            sr,si );
#endif
    /* 2nd stage, fixed shift */
        fxshft(10*cnt2, &zr, &zi, &conv):
        if( conv )
        l
        idnn2 = degree-nn+2;
        zeror[idnn2-1] = zr;
        zeroi[idnn2-1] = zi;
#ifdef EBUG
#endif
        printf( "zero: %q+ %gi\n", zr,zi );
        nn = nn-1;
        for( i = 1; i <=nn; i += 1)
        l
            pr[i-1] = qpr[i-1];
            pi[i-1] = GPi[i-1];
        }
        goto found_root; /* necessary evil */
            }
        }
        |
        *fail = TRUE;
        return;
        found_root:
        ;
        }
        if( nn >= 1 )
            cdivid( -pr[1], -pi[1], pr[0], pi[0],
            &zeror[degree-1], &zeroi[degree-1] );
#ifcef EBUG
        printf( "nn was fd at return.\n",nn );
#endif
            return;
l
/*
    * computes the derivative polynomial as the initial h
    * polynomial and computes ll no-shift h polynomials.
    */
noshft(ll)
int ll;
I
        double xni,t1, t2, dn;
        int i, n, nml, jj, j;
        n = nn - 1;
        nml = n - l;
        dn = (double) n;
```

```
    for( i = 1; i <=n; i += 1)
    l
        xni = (double) (nn - i);
        hr[i-1] = xni*pr[i-1]/dn;
        ni[i-1] = xni*pi[i-1]/dn;
    }
    for( jj = 1; jj<= l1; jj += 1 )
    l
        if(cmod(hr[n-1],hi[n-1])> eta*10.0* cmod(pr[n-1],pi[n-1]))
        }
            cdivid(-pr[nn-1],-pi[nn-1],hr[n-1],hi[n-1],&tr,&ti);
            for( i = 1; i <= nm1; i += 1 )
            {
                        j = nn-i;
                        t1 = hr[j-2];
                        t2 = hi[j-2];
                        hr[j-1] = tr*t1-ti*t2+pr[j-1];
                        hi[j-1] = tr*t2+ti*t1+pi[j-1];
                }
                hr[0] = pr[0];
                hi[0] = pi[0];
        }
        else
        l
            for( i=1; i <= nml; i += 1)
            {
            j = nn-i;
            hr[j-1] = hr[j-2];
            hi[j-1]=ni[j-2];
            }
            hr[0] = 0.0;
            hi[0] = 0.0;
        }
    }
    return;
}
/*
    * computes }12\mathrm{ fixed-shift h polynomials and tests for
    * convergence
    * initiates a variable-shift iteration and returns with the
    * approximate zero if successful.
    * l2 - limit of fixed shift steps
    * zr,zi - approx zero if conv is .true.
    * conv logical indicating convergence of stage 3 iteration
    */
fxshft(12,zr,zi,conv)
int 12, *conv;
double *zr, *zi;
l
    double otr,oti,svsr,svsi;
    int test,pasd,bool;
    int n, i, j;
```

```
n = nn-1;
polyev(nn,sr,si,pr,pi,qpr,qpi,&pvr,&pvi); /* evaluate p at s */
test = TRUE;
pasd = FALSE;
calct(&bool ): /* lst t = -p(s)/h(s) */
/* main loop, 2nd stage step */
for( j = 1; j <=12; j += 1 )
l
    otr = tr;
    oti = ti;
    nexth( &bool ); /* next polynomial */
    calct( &bool ); /* new t */
        *zr = sr+tr;
        *zi=si+ti;
        /* convergence test */
        if( bool || (!test) || j == l2)
            continue;
        if( cmod(tr-otr,ti-oti) >= . 5* cmod(*zr,*zi)) /* weak conv. */
        {
            pasd = FALSE;
            continue;
        }
        if( ! pasd )
        {
            pasd = TRUE;
            continue;
        }
        for( i = 1; i <=n; i += 1)
        {
            shr[i-1] = hr[i-1];
            shi[i-1] = hi[i-1];
        )
        svsr = sr;
        svsi = si;
        vrshft(60,zr,zi,conv);
        if( *conv)
            return;
        test = FALSE;
        for( i = 1; i <= n; i += 1)
        |
            hr[i-1] = shr[i-1];
            hi[i-1] = shi[i-1];
        }
        sr = svsr;
        si = svsi;
        polyev(nn,sr,si,pr,pi,qpr,qpi,&pvr,&pvi);
        calct( &bool );
l
vrshft(50, zr, zi, conv);
return;
```

```
/*
    * carries out the third stage i=eration
    * 13 - limit of steps in stage 3.
    * zr,zi - on entry contains the initial iterate, if the
    * iteration converges it contains the final iterate
    * on exit.
    * conv - .true. if iteration converges
    */
vrshft(13,zr,zi,conv)
int l3, *conv;
double *zr, *zi;
{
    double mp,ms,omp,relstp,r1,r2,tp, twenerr;
    int b,bool;
    int i, j;
        *conv = FALSE;
        b = FALSE;
        - sr = *zr;
        si = *zi;
#ifdef EBUG
        printf( "shift (stage 3): %g+ %gi\n", sr,si );
#endif
    for( i=1; i<=13; i += 1)
    l
            polyev(nn,sr,si,pr,pi,qpr,qpi,&pvr,&pvi);
            mp = cmod(pvr,pvi):
            ms = cmod(sr,si);
            twenerr = 20.0*errev(qpr,qpi,ms,mp,are,mre);
#ifdef EBUG
            printf( "mp: %%g, twenerr: %%g\n", mp, twenerr );
#endif
        if(mp<= twenerr )
        {
            *conv = TRUE;
            *zr = sr;
            *zi = si;
            return;
        1
        if( i > l )
        l
            if( b || mp < omp || relstp >= .05 )
            l
                    if(mp*.1 > omp )return;
            |
            else
            |
                    tp = relstp;
                    b = TRUE;
```

```
        if( relstp < eta )
            tp = eta;
        rl = sqrt( tp );
        r2 = sr*(1.0+r1)-si*r1;
        si = sr*rl+si*(1.0+rl);
        sr = r2;
        polyev(nn,sr,si,pr,pi,qpr,qpi,&pvr,&pvi);
        for( j = 1; j <=5; j += 1 )
        {
            calct( &bool );
            nexth( &bool );
        )
        omp = infin;
        }
        }
        omp = mp;
        calce( &bool );
        nexth( &bool );
        calct( &bool );
        if( !bool)
        {
            relstp = cmod(tr,ti)/cmod(sr,si);
            sr = sr+tr;
            si = si+ti;
        J
        J
        return;
}
/*
    * computes t = -p(s)/h(s).
    * bool - logical set true if h(s) is essentially zero.
    */
calct(bool)
int *bool;
l
    double hvr,hvi;
        int n;
        n=nn-1;
        polyev(n,sr,si,hr,hi,qhr,qhi,&hvr,&hvi); /* h(s) */
        *bool = ((cmod(hvr,hvi) <= are*l0.0*\operatorname{mod}(hr[n-1],hi[n-1]))
            ? TRUE : FALSE );
        if( *bool)
        {
        tr = 0.0;
        ti=0.0;
        }
        else
        {
            cdivid(-pvr,-pvi,hvr,hvi,&tr,&ti);
```

```
        |
        return;
}
/*
    * calculates the next shifted h polynomial.
    * bool - logical, if .true. h(s) is essentially zero
    */
nexth(bool)
int *bool;
{
    double tl,t2;
    int n, nml, j;
        n = nn-1;
        nm1 = n-1;
        if( *bool)
        |
            for( j = 2; j <= n; j += l )
            {
                    hr[j-1] = qhr[j-2];
                    hi[j-1]=qhi[j-2];
                }
                hr[0] = 0.0;
                hi[0] = 0.0;
                return;
        }
        else
        {
            for( j = 2; j <=n; j += 1 )
                |
                    t1 = qhy[j-2];
                    t2 = qhi[j-2];
                    hr[j-1]= tr*t1-ti*t2+qpr[j-1];
                    hi[j-1] = tr*t2*ti*t1+qpi[j-1];
                }
                hr[0] = qpr[0];
                hi[0] = qpi[0];
        }
        return;
|
/*
    * evaluates a polynomial p at s by the horner recurrence
    * placing the partial sums in q and the computed value in pv.
    */
polyev(n,sr,si,pr,pi,qr,qi,pvr,pvi)
double *pr, *pi, *qr, *qi, sr,si, *pvr, *pvi;
int n;
{
        int i;
```

```
    double t;
    qr[0] = pr[0];
    qi[0] = pi[0];
    *pvr = qr[0];
    *pvi = qi[0];
    for( i = 2; i <= n; i += 1 )
    {
        t = (*pvr)*sr-(*pvi)*si+pr[i-1];
        *pvi = (*pvr)*si+(*pvi)*sr+pi[i-1];
        *pvr = t;
        qr[i-1] = *pvr;
        qi[i-1] = *pvi;
|
return;
}
/*
    * bounds the error in evaluating the polynomial by the horner
    * recurrence.
    * qr,qi - the partial sums
    * ms - modulus of the point
    * mp - modulus of the polynomial value
    * are, mre - error bounds on complex addition and multiplication
    */
double errev(qr,qi,ms,mp,are,mre)
double *qr, *qi, ms,mp,are,mre;
{
        int i;
        double e;
        e = cmod(qr[0],qi[0])*mre/(are+mre);
        for( i = 1; i <= nn; i t= l i
        l
        e = e*ms+cmod(qr[i-1],qi[i-1]);
        }
        return( e*(are+mre)-mp*mre );
}
/*
    * cauchy computes a lower bound on the moduli of the zeros of a
    * polynomial - pt is the modulus of the coefficients
    */
double cauchy(pt,q)
double *q, *pt;
|
        double x,xm,f,dx,df;
        int i, n;
        n = nn-1;
        pt[n] = -pt[n];
```

```
    /* upper estimate of bound */
    x = exp( (log(-pt[n]) - log(pt[0]))/( ( double )n) );
    if(pt[n-1] != 0.0 )
    l
        xm = -pt[n]/pt[n-1];
        if( xm<x ) x = xm;
    }
    for( f = 1.0; /* kluge */ f > 0.0; x = xm )
    l
        xm = x*0.1;
        f = pt[0];
        for( i = 2; i <=nn; i += 1)
        l
            f= f*xm+pt[i-1];
        }
    |
    /* Newton iteration until x converges to two decimal places */
    dx = x;
    while( dabs(dx/x) > 0.005)
    l
        q[0] = pt[0];
        for( i = 2; i <=nn; i += 1)
        {
            q[i-1] = q[i-2]*x+pt[i-1];
        }
        f = q[n];
        df = q[0];
        for( i = 2; i <= n; i += 1 )
        {
            df = df*x+q[i-1];
        }
        dx = f/df;
        x = x-dx;
    l
    return( x );
}
/*
    * returns a scale factor to multiply the coefficients of the
    * polynomial. the scaling is done to avoid overflow and to avoid
    * undetected underflow interfering with the convergence
    * criterion. the factor is a power of the base.
    * pt - modulus of the coefficients of p
    * eta,infin,smalno,base - constants describing the
    * floating point arithmetic.
    */
double scale(pt)
double *pt;
l
```

```
    double hi, lo, max, min, x, sc, l;
    int i;
    /* find largest and smallest moduli of coefficients */
    hi = sqrt(infin);
    lo = smalno/eta;
    max = 0.0;
    min = infin;
    for( i = 1; i <= nn; i += l )
    {
        x = pt[i-1];
        if( }x>>max 
            max = x;
        if( x != 0.0 && x < min )
        min = x;
    }
    /* only necessary with very large or very small components */
    if( min >= lo && max <= hi )
        return( 1.0 );
    x = lo/min;
    if( x > 1.0 )
    {
        sc = x;
        if( infin/sc > max )
            sc = 1.0;
    }
    else
    |
        sc = 1.0/(sqrt(max)*sqrt(min));
    |
    l = ceil(log(sc)/log(base));
    return( pow(base,1) );
}
/* complex division, avoiding overflow */
cdivid(ar,ai,br,bi,cr,ci)
double ar,ai,br,bi,*cr,*ci;
l
    double r,d;
    if(br != 0.0 || bi != 0.0 )
    {
        if( dabs(br) >= dabs(bi))
        |
                r = bi/br;
                d = br+r*bi;
                *cr = (ar+ai*r)/d;
                *ci = (ai-ar*r)/d;
                return;
            }
            elsel
```

```
                r = br/bi;
                    d = bi+r*br;
                    *cr = (ar*r+ai)/d;
                    *ci = (ai*r-ar)/d;
                    return;
    }
    }
    else /* division by zero, c = infinity */
    l
        mcon():
        *cr = infin;
        *ci = infin;
        return;
    }
}
/* modulus of a complex number, avoiding overflow */
double cmod(r,i)
double r,i;
l
    double ar, ai;
    ar = dabs(r);
    ai = dabs(i);
    if( ar > ai )
        return(ar*sqrt(1.0+(ai/ar)*(ai/ar)));
    else
    l
        if( ar< ai )
                        return( ai*sqrt(1.0+(ar/ai)*(ar/ai)));
            else
                return( ar*M_SQRT2 );
    }
}
    * mcon provides machine constants used in various parts of the
    * program. the user may either set them directly or use the
    * statements below to compute them. the meaning of the four
* constants are -
* eta the maximum relative representation error
* which can be described as the smallest positive
* floating-point number such that 1.0d0 + eta is
* greater than 1.0d0
* infiny the largest floating point number
* smalno the smallest positive floating-point number
* base the base of the floating-point number system used
* let t be the number of base-digits in each floating-point
* number(double precision). then eta is either . 5* b**(1-t)
* or b**(1-t) depending on whether rounding or truncation
* is used
* let m}\mathrm{ be the largest exponent and n the smallest exponent
```

```
    * in the number system. then infiny is (1-base**(-t))*base**m
    * and smalno is base**n.
    * the values for base,t,m,n below correspond to the ibm/360.
    *
    * *NOTE* revised for IEEE floating point
    *
    */
mcon()
{
    base = 2.0;
#ifdef vax
    eta = pow(base, -54.0);
    infin = pow(base,126.0);
    smalno = 1.0/infin;
#else
    eta = pow(base,(1.0-52.0));
    infin = pow(base,1023.0);
    smalno = 1.0/infin;
#endif
#ifdef EBUG
    printf( "eta=%e, 1.0+eta=%e, infin=%e, smalno=%e\n",
        eta, 1.0+eta, infin, smalno );
#endif
        return;
]
```


## 22. Appendix XIV: cvaryangle.c

```
#include <stdio.h>
#include <sys/param.h>
#include <sys/types.h>
#include <sys/times.h>
#define MAX_DEGREE 50
/*
    * batch interface program for j-t method,
    * found in this directory in file "cjt.c"
    */
main( argc, argv, env )
char *argv[], *env[];
{
    long clock, times(), min, avg, max, user, nprocs;
    struct tms tbl, tb2;
    int i, degree, flflag, count, fail_count;
        double init, coeffr[MAX_DEGREE], coeffi[MAX_DEGREE],
        resulr[MAX_DEGREE], resuli[MAX_DEGREE], offset;
        scanf( "%d", &degree );
#ifdef EBUG
        fprintf( stderr, "degree: %d\n", degree );
#endif
    if( degree > MAX_DEGREE-1 || degree < 0)
    1
        fprintf( stderr, "0<= degree <= %d", MAX_DEGREE-1 );
        exit( l );
    }
    for( i = 0; i <= degree; i += 1)
    {
        scanf( " %lf, %lf", &coeffr[i], &coeffi[i]);
    }
```

\#ifdef EBUG
for ( $i=0$; $i$ < degree; $i+=1$ )
!
fprintf( stderr, "\%g, \%g\n", coeffr[i], coeffi[i]);
\}
\#endif
/*
* apply the method for a sequence of angles
* 1
for ( nprocs=1; nprocs $<=50$; nprocs $i=1$ )
1
fail_count $=$ count $=\min =\max =a v g=0$;
offset $=$ (double) $360.0 /$ nprocs;
for ( init=3.0;
init <= 360.0; init $+=$ offset )

```
    {
        count += 1;
        clock = times( &tb1 );
        cpoly( coeffr, coeffi, degree, resulr, resuli, &flflag,
        (init/180.0)*3.14159265358979323846 );
    clock = times( &tb2 ) - clock;
    user = tb2.tms_utime - tbl.tms_utime;
    if( flflag )
    |
#ifdef FAIL
        fprintf( stderr,
            "cpoly() failed on input. Exiting.\n" );
                exit( l );
#else
#endif
                fail_count += 1;
    }
#ifdef VERBOSE
    printf( "\nangle: %g, time %d ticks, zeros at:\n",
                init, user );
    for( i=0; i <= degree-1; i += 1)
    i
        printf( "各e, ofe\n", resulr[i], resuli[i] );
    }
#endif
    if(min == 0 && !flflag )
        min = user;
    if( user < min && !flflag )
        min = user;
    if( user > max )
        max = user;
    avg += user;
1
avg = avg / count;
printf( "nprocs: %d, max: %d, min: %d, avg: %d, fails: %d\n",
    nprocs, max, min, avg, fail_count );
}
exit( 0 );
I
```


## 23. Appendix XV: r2p.c

```
#include <stdio.h>
#define MAX_DEGREE 50
#define EOL '\n'
#define BUFSIZE 512
#define EOS '\0'
struct complex (
    double r;
    double i;
};
typedef struct complex COMPLEX;
COMPLEX tmp[MAX_DEGREE],
    roots[MAX_DEGREE],
    coeff[MAX_DEGREE];
main( argc, argv, env )
int argc;
char *argv[], *env[];
l
    FILE *fp;
    if( argc > 2 )
        usage();
    if( argc == 2 )
    {
        fp = fopen( argv[l], "r" );
        if( fp == (FILE *) NULL )
        l
            perror( argv[1] );
                    exit( 1 );
            1
        }
        else
            fp = stdin;
    load_roots( fp );
    close( fp );
    compute_poly();
    print poly();
    exit( 0 );
}
usage()
|
    fprintf( stderr, "Usage: r2p [filename]\n" );
    exit( 1 );
}
```

```
print_poly()
|
    int deg;
    register int i;
    deg = degree( coeff );
    printf( "%d\n", deg );
    for( i = 0; i <= deg; i += 1)
    {
        printf( "%f, %f\n", coeff[i].r, coeff[i].i );
    )
    return;
}
load_roots( file )
EILE *file;
{
    int i, getline();
    char line[BUFSIZE], *line_ptr, *get_float();
    for( i = 0;
            getline( line, BUFSIZE ) > 0;
            i += 1 )
    1
            line_ptr = line;
            line_ptr = get_float( line_ptr, &roots[i].r );
            get_float( line_ptr, &roots[i].i );
#ifdef EBUG
            printf( "load_roots: got %g+%gi\n",
                        roots[i].r, roots[i].i );
#endif
    }
    return;
}
compute_poly()
{
    int i;
    int deg;
    deg = degree( roots );
    coeff[1].r = -roots[0].r;
    coeff[1].i = -roots[0].i;
    coeff[0].r = 1.0;
    coeff[0].i = 0.0;
    for( i = 1; i <= deg; i += 1)
    {
```

```
            roots[i].r = -roots[i].r;
            roots[i].i = -roots[i].i;
                do_root( &roots[i] );
    I
    return;
1
do_root( root )
COMPLEX *root;
|
    register int i;
    int deg;
    COMPLEX c;
    deg = degree( coeff );
    cprod( root, &coeff[deg], &c );
    tmp[deg+1].r = c.r;
    tmp[deg+1].i = c.i;
    tmp[0].r = 1.0;
    tmp[0].i = 0.0;
    for( i = 1; i <= deg; i += 1 )
    {
            cprod( root, &coeff[i-1], &c );
            csum( &coeff[i], &c, &tmp[i] );
        }
        for( i = 0; i <= deg+1; i += 1 )
            coeff[i] = tmp[i]; /* needs structure assign. */
        return;
}
cprod( a, b, c )
COMPLEX *a, *b, *C;
I
    c->r = a->r*b->r - a->i*b->i;
    c->i = a->i*b->r + a->r*b->i;
    return;
}
csum( a, b, c )
COMPLEX *a, *b, *c;
|
    c->r = a->r + b->r;
    c->i = a->i +b->i;
    return;
}
int
degree( poly )
COMPLEX *poly;
|
```

```
    register int i;
    int deg:
    for( i = 0; i < MAX_DEGREE; i += l )
        if( poly[i].r !=0.0 || poly[i].i != 0.0 )
                deg = i;
#ifdef EBUG
    printf( "degree: returning %d\n", deg );
#endif
    return( deg );
}
/*
    * safe "gets()"
    * leaves terminating newline
    * always returns null-terminated string.
    */
int
getline( buf, size )
char *buf;
int size;
l
    int c, counter;
    if( buf == (char *) NULL )
            return( 0 );
        for( counter = 0; counter < size-1; )
        |
            c = getchar();
            if(c == EOF )
                        break;
            buf[counter++] = c;
            if( c == EOL )
                    break;
        |
        buf[counter] = EOS;
        return( counter );
}
char *
get_float( str, dub )
char *str;
double *dub;
l
    char copy[BUFSIZE], *ptr;
    if( str == (char *) NULL )
            return( (char *) NULL );
        while( *str != EOS )
```

```
    1
        if( *str >= '0' && *str <= 'g' )
        break;
        if( (*str == '+' || *str == '-' || *str == '.') &&
            (*(str+1) >= '0' && *(str+1) <= '9' ) )
            break;
        ++str;
    }
    ptr = copy;
    while( *str != EOS )
    l
        if( (*str < '0' || *str > '9' )
            && *str != '+'
            && *str != '_'
            && *str != '.'
            && *str != 'e' )
        1
            *ptr = EOS;
#ifdef EBUG
            printf( "copy was: ofs\n", copy );
#endif
            sscanf( copy, "%lf", dub );
            return( str );
        }
        *ptr = *str;
        ++ptr;
        ++str;
]
return( str );
1
```


## 24. Appendix XVI: cmach.c

```
#include <stdio.h>
#include <sys/param.h>
#include <sys/types.h>
#include <sys/times.h>
#include <sys/signal.h>
#include <sys/time.h>
#define MILLION 1000000
#define MAX_DEGREE 50
#define FILE_NAME_LEN 128
char sema_file[FILE_NAME_LEN];
/*
    * "parallel" batch interface program for j-t method,
    * found in this directory in file "cjt.c"
    */
main( argc, argv, env )
char *argv[], *env[];
l
    int catcher();
    long clock, user, nprocs;
    struct tms tbl, tb2;
    int i, fd, degree, flflag, count, pid, ppid, status;
    double init, coeffr[MAX_DEGREE], coeffi[MAX_DEGREE],
    resulr[MAX_DEGREE], resuli[MAX_DEGREE], offset;
#ifdef SYSTEM_FIVE
    long times():
#else
    struct timeval tvl, tv2;
    struct timezone tz;
#endif
    scanf( "%d", &degree ):
    if( degree > MAX_DEGREE-1 || degree < 0 )
    |
        fprintf( stderr, "0<= degree <= %od", MAX_DEGREE-1 );
        exit( 1 );
    }
    for( i = 0; i <= degree; i += 1 )
    {
        scanf( " &lf, %lf", &coeffr[i], &coeffi[i] );
    }
    /*
        * apply the method for a sequence of angles
        */
    pid = fork();
```

```
    switch( pid )
    {
    case -1:
        fprintf( stderr, "Can't fork; exiting!\n"):
        exit( l );
    case 0:
        break;
    default:
        while ( wait( &status ) >= 0 )
        ;
        _exit( 1 ):
    l
    for( nprocs=1; nprocs <= 6; nprocs += 1)
    l
#ifdef SYSTEM_FIVE
    clock = times( &tbl );
#else
    gettimeofday( &tv1, &tz );
#endif
    setpgrp();
    ppid = getpid();
    get_sema();
    signal( SIGTERM, catcher );
    offset = (double) 360.0/mprocs;
    for( init=3.0;
        init <= 360.0; init += offset )
    l
            pid = fork();
            switch( pid)
            l
            case -1:
                    fprintf( stderr, "Can't fork, exiting!\n" );
                    exit( l );
                        case 0:
                            cpoly( coeffr, coeffi, degree,
                    resulr, resuli, &flflag,
                    (init/180.0)*3.14159265358979323846 ):
                    if(flflag )
                l
                    _exit( 1 );
                        l
                            lock_sema();
                            signal( SIGTERM, SIG_IGN );
                kill(0, SIGTERM );
                break;
                        default:
                        break;
```

```
    }
        if( pid == 0)
        break;
    }
    if( pid != 0 )
    l
        while( wait( &status ) > 0 )
        ;
        free_sema();
    }
#ifdef SYSTEM_FIVE
    clock = times( &tb2 ) - clock;
    printf( "nprocs: %d, time: %od ticks\n", nprocs, clock );
#else
    gettimeofday( &tv2, &tz );
    /* adjust as per manual page */
    if( tvl.tv_usec > tv2.tv_usec )
    l
        tv2.tv_usec += MILLION;
        tv2.tv_sec -= 1;
        }
        clock = MILLION*(tv2.tv_sec-tv1.tv_sec)+
            (tv2.tv_usec-tv1.tv_usec);
        printf( "nprocs: %d, time: %d usec\n", nprocs, clock );
#endif
        |
        exit( 0 ):
}
#ifdef HPUX
#include <sys/types.h>
#include <sys/ipc.h>
#include <sys/sem.h>
int Semid;
struct sembuf Sops[1];
ushort Sinit[1];
get_sema()
{
    key_t k;
    int ret;
    sprintf( sema_file, "/tmp/CMACH%d", getpid() );
    k = ftok( sema_file, 'A' );
    ret = semget ( k, 1, IPC_CREAT | 0600 );
    if( ret < 0 )
    l
        fprintf( stderr, "get of semaphore failed. Exiting.\n");
        exit( 1 );
        |
        Semid = ret;
```

```
    Sinit[0] = 1;
    if( semctl( Semid, 0, SETALL, Sinit ) < 0 )
    |
        fprintf( stderr, "set of semaphore failed. Exiting.\n");
        exit(1 );
    }
1
lock_sema()
{
        Sops[0].sem_num = 0;
        Sops[0].sem_op = - 1;
        Sops[0].sem_flg= IPC_NOWAIT;
        if( semop( Semid, Sops, 1 ) < 0)
            _exit( 1 );
        else
        return;
}
free_sema()
1
        semctl( Semid, 0, IPC_RMID, Sinit );
        return;
}
#else
get_sema()
{
    int fd;
        sprintf( sema_file, "/tmp/CMACH%d", getpid() );
        if( (fd=creat(sema_file,0)) < 0)
        {
```



```
                sema file );
            exit( l );
        }
        else
            close( fd );
        return;
}
free_sema()
[
    lock_sema();
}
lock_sema()
{
```

```
    if( unlink( sema_file ) == - 1 )
    l
        exit( 0 );
    }
    return;
l
#endif
int catcher( sig )
l
        if( sig == SIGTERM )
        |
            signal( SIGTERM, catcher );
#ifdef EBUG
            fprintf( stderr, "pid %d, caught SIGTERM\n", getpid() );
        exit( l );
        |
        else
        return;
}
```


## 25. Appendix XVII: Extremal exploitation of randomness

```
rand_search.c:
#include <stdio.h>
#include <math.h>
#include <sys/types.h>
#include <sys/times.h>
#include <sys/time.h>
#define MILLION 1000000
#define TRUE l
#define FALSE 0
#ifdef EBUG
#define PERM_FILE "/tmp/Permutation"
#endif
#define MAX_SIZE 100000
int S[MAX_SI2E],
    Ticks[MAX_SIZE],
    P[MAX_SIZE]
    SSize = -1, PSize = -1;
main( argc, argv )
int argc;
char *argv[];
l
    double total, scaler, dSize, floor(), drand48();
    register int i, j, k;
    int min, max;
    long clock;
        struct timeval tv1;
        struct timezone tz;
        while( argc > 1 )
        l
            if( argv[argc-l][0] != '-' )
                usage();
            switch( argv[argc-1][I] )
            l
            case 's':
                                PSize = atoi( &argv[argc-l][2] );
                                if( PSize < 1 || PSize > MAX_SIZE )
                                    usage();
                                break;
            case 'c':
                        SSize = atoi( &argv[argc-1][2] );
                        if( SSize < l || SSize > MAX_SIZE )
                            usage();
                                break;
```

```
        default:
            usage();
        }
        argc -= 1;
    }
    if( PSize == -1 )
    PSize = MAX_SIZE;
if( SSize == -1 )
    SSize = MAX_SI2E;
dSize = (double) PSize;
gettimeofday( &tv1, &tz );
clock = MILLION*tv1.tv_sec+tvl.tv_usec;
srand48( clock );
for( i = 0; i < PSize; i += l )
    P[i] = i;
/* this might have to be done better */
for( i = 0; i < PSize; i += l )
l
    k = (int) floor( drand48()*dSize );
    j = P[i]; P[i] = P[k]; P[k] = j;
}
#ifdef EBUG
    l
        FILE *fP;
        fp = fopen( PERM_FILE, "w" );
        if( fp == (EILE ` *) NULL )
        1
            fprintf( stderr,"Can't open %s for writing. Exiting\n",
                    PERM FILE );
                    exit( l );
            }
            fprintf( fp, "Elements: sd\n", PSize );
            for( i = 0; i < PSize; i += 1 )
                    fprintf( fp, "%d%c", P[i], (igl0)?',':'\n' );
        fclose( fp ):
    }
#endif
k = (int) floor( drand48()*dSize );
for( i = 0; i < SSize; i += 1)
l
        S[i] = (int) floor( drand48()*dSize );
        gettimeofday( &tv1, &tz );
        clock = MILIION*tvl.tv_sec+tvl.tv_usec;
```

```
        Ticks[i] = clock;
        if( P[find_index( k, S[i] )] != k )
            fprintf( stderr, "find_index failed. continuing.ln" );
        gettimeofday( &tvl, &tz );
        clock = MILLION*tvl.tv_sec+tvl.tv_usec;
        Ticks[i] = clock - Ticks[i];
    }
    /* dump statistics */
    scaler = 1.0/((double) MILLION);
    min = max = Ticks[0];
    /* we pre-scale Ticks to prevent overflow */
    total = scaler*Ticks[0];
    for( i = 1; i < SSize; i += 1 )
    i
        if( Ticks[i] < min )
            min = Ticks[i];
        if( Ticks[i] > max )
            max = Ticks[i];
        total += scaler*Ticks[i];
    }
    printf( "min: %g sec, max: %g sec, avg: %g sec.\n",
        scaler* (double) min,
        scaler* (double) max,
        total / (double) SSize );
    exit( 0 ):
1
int
find_index( element, start )
int element, start;
1
    int i, offset, halfSize;
    halfSize = PSize/2;
    for( offset = 0; offset <= halfSize; offset += 1 )
    l
        i = start - offset;
            if( i < 0)
                i += PSize;
            if( equal( P[i], element ) )
                return( i );
            i = start + offset;
            if( i >= PSize )
                i -= PSize;
            if( equal( P[i], element ) )
                return( i );
            I
```

```
    return( -1 );
}
int
equal( i, j )
int i, j;
|
    if( i == j )
        return( 1 );
    return( 0 );
}
usage()
{
    fprintf( stderr, "Usage: rand_search [-ssize] [-ccount]\n" );
    fprintf( stderr, "size & count <= %d\n", MAX_SIZE );
    exit( 1 );
}
script:
for i in 1 2 5 10 20 50 100 200 500 1000 5000 10000 50000 100000
do
    rand_search -c${i}
done
results:
min: 0.361704 sec, max: 0.361704 sec, avg: 0.361704 sec.
min: 0.038728 sec, max: 0.158708 sec, avg: 0.098718 sec.
min: 0.019244 sec, max: 0.606028 sec, avg: 0.385311 sec.
min: 0.022124 sec, max: 0.68992 sec, avg: 0.249112 sec.
min: 0.020452 sec, max: 0.698092 sec, avg: 0.456766 sec.
min: 0.031444 sec, max: 0.70512 sec, avg: 0.330535 sec.
min: 0.00248 sec, max: 0.71294 sec, avg: 0.330925 sec.
min: 0.00034 sec, max: 0.739772 sec, avg: 0.3693 sec.
min: 0.002532 sec, max: 1.18184 sec, avg: 0.383545 sec.
min: 0.001124 sec, max: 0.710796 sec, avg: 0.35026 sec.
min: 0.00024 sec, max: 2.53285 sec, avg: 0.356609 sec.
min: 0.000164 sec, max: 2.85144 sec, avg: 0.367076 sec.
min: 0.000176 sec, max: 8.0722 sec, avg: 0.371924 sec.
min: 0.00012 sec, max: 2.39245 sec, avg: 0.358988 sec.
```


## 26. Biography

Jonathan Michael Smith was born on October 10, 1959 in Taunton, Massachusetts, where he attended elementary and secondary school. He received his Bachelor of Arts degree in Mathematics, Magna Cum Laude, from Boston College, Chestnut Hill, Massachusetts in May. 1981. Jonathan's senior Honors thesis in number theory was on "Full Period Primes," which are prime numbers whose unitary fraction has the longest possible repeating decimal expansion.

After graduating, Jonathan was employed by Bell Telephone Laboratories. From September 1982 until May 1983, he was supported in his studies towards a Master of Science in Computer Science in the Laboratories' One Year on Campus graduate studies program. Columbia University granted the M.S. degree in May of 1983. On January 1 of 1984. Jonathan became an employee of Bell Communications Research. Inc. as a result of the AT\&T divestiture. He returned to Columbia University in September of 1984 in order to pursue a Doctor of Philosophy degree in Computer Science.


[^0]:    ${ }^{1}$ These are observations noted by Nelson [Nelson 1987a] in a videotaped lecture.

[^1]:    ${ }^{2}$ As reported in [Smith 1988a] we observed a write fraction, which describes the fraction of memory copied by "copy-on-write" mechanisms, to be between 0.2 and 0.5 . Thus 0.5 is reasonable, since the major overhead we observed was copying.

[^2]:    ${ }^{3}$ This statement "know little" must be qualified. On the one hand, the contents of a large personnel data base may be available before we begin our computation, so that we could "know" how long a particular formulation of a query will take. On the other hand, it seems computationally infeasible to (1) pre-compute this information or (2) store per-datum, or static aggregate, annotations of which method to choose. Also, execution time information from previous executions may have liulle predictive value, or may be sensitive to environmental change.
    ${ }^{4}$ For example, the single bit history of page-referencing behavior maintained with a "dirty" bit in a virtual memory management system.

[^3]:    ${ }^{6}$ Thus, so-called "sibling rivalry" is taken to its extreme in this design! The failure altemative assumes that none of the siblings will complete.

[^4]:    ${ }^{7}$ Thus implying all the sender's predicates.

[^5]:    ${ }^{8}$ Thus implying rejection of the sender's predicates without creating a logical impossibility. Assuming the negation of all of S 's predicates might imply that two mutually exclusive processes must complete.

[^6]:    ${ }^{9}$ HP-UX on our HP9000/350 systems uses a 4 K pagesize. With our instrumentation (e.g., do_fork.c) the difference is relevant only when the offset of a particular byte in the last page accessed causes an extra 2 K bytes of memory to be paged in. It may be more relevant to applications.

[^7]:    ${ }^{10}$ For comparison purposes, this would be about $2,600(2,573=5,270,467 / 2,048) 2 \mathrm{~K}$ pages per second.

[^8]:    ${ }^{11}$ For comparison, 1,655; 1,932; and 2,069 2 K pages/second.

[^9]:    ${ }^{12}$ The results for GNU Emacs were checked on the workstations, and they are consistent. For the "Towers of Hanoi" problem discussed below, the fraction of the data altered by the program was 0.30 on the 3 B 2 and 0.48 on the HP9000. Much of the difference is due to what features the runnable Emacs is pre-loaded with; the VAX executable has large amounts of pre-loaded information, which is read-only.

[^10]:    ${ }^{13}$ See the description in Smith's survey paper [Smith 1988b].

[^11]:    ${ }^{14}$ Evidence [Gray 1988a] from commercial systems [Bartleu1978a, Borg 1983a, Bartlett1981a] indicates that reliability is increased significantly with two copies of the state, as maintained in a "process pair." The increase in reliability by adding a third process is infinitesimal, and is achieved at a significant cost in storage and synchronization. Thus, if the location of storage copies is well-known, it may not be necessary to have more than two copies in the system; demand-copying can be used for any copies beyond the two necessary for reliability.

[^12]:    ${ }^{16}$ This presentation is more concerned with the variables and control mechanisms than with the full unification theorem proving scheme; thus our portrayal of the process as "pattern-matching" is sufficient.

[^13]:    ${ }^{17}$ Warren's survey provides a much more in-depth discussion of what he calls the "Abstract Model of classical resolution;'" sec also the discussion of resolution in Nilsson [Nilsson 1980a].

[^14]:    After 21,500 minutes ( $=358$ hours, or more than 14 days) of CPU time (HP9000/350) this process had not printed a result. I gave up.

[^15]:    ${ }^{18}$ A pipeline in a computer architecture is a logic implementation based on the principle of an assembly line. That is, there are a series of specialized functional units which solve a portion of the problem; these can be made rapid, and ideally all the functional units can be kept busy at the same time. Analogously, the ideal case is realized when everything is predetermined. As Henry Ford said,
    "You can have any color you want, as long as it's black."

[^16]:    $1,6,6,5,1,5,2,5,8,2,9,1,9,3,9,3,1,3,4,3,3,9,1,2,2,2,2,6,1,4,6,2,1,2$, $4,2,0,1,4,1,5,3,3,6,5,0,3,7,2,8,6,8,3,3,6,6,2,9,2,1,1,8,6,2,9,0,4,2$, $3,5,4,7,7,8,9,6,6,3,7,8,3,7,8,5,4,6,7,8,9,3,0,1,2,6,3,8,0,4,1,0,8,2$, $1,4,3,7,8,5,4,8,7,3,9,8,8,6,6,4,8,7,9,9,2,3,4,1,1,7,9,9,4,8,5,0,4,3$, $3,8,6,6,7,7,8,1,2,5,5,9,4,5,4,1,3,4,7,2,4,6,5,2,4,6,2,3,1,1,9,4,8,8$, $1,4,0,1,3,1,6,0,7,1,6,2,8,4,2,7,2,8,1,7,1,3,0,4,2,2,4,7,8,6,9,1,8,5$, $6,3,1,2,0,0,1,9,2,3,3,3,6,9,8,9,6,6,9,3,3,5,4,4,3,6,1,6,2,9,3,9,1,3$, $1,1,0,4,1,7,3,0,9,5,6,5,0,1,6,9,4,6,6,2,7,5,4,5,5,8,8,7,5,6,4,4,3,4$, $5,1,9,1,2,6,9,2,7,9,6,0,0,6,9,3,5,5,1,8,0,9,2,7,1,9,5,6,4,5,0,2,6,4$, $2,9,4,0,9,2,8,5,7,4,1,0,8,2,8,3,5,3,5,1,1,8,8,2,7,5]$, X15).
    *

    * statistics.
    */

