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A SIMPLE PREPROCESSING SCHEME TO EXTRACT AND BALANCE IMPLICIT PARALLELISM IN THE CONCURRENT MATCH OF PRODUCTION RULES.

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A Simple Preprocessing Scheme to Extract and Balance Implicit Parallelism in the Concurrent Match of Production Rules¹

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Abstract

In this brief paper we report a simple scheme to extract implicit parallelism in the low-level match phase of the parallel execution of production system programs. The essence of the approach is to replicate rules while introducing new constraints within each copy to restrict each individual rule to match a potentially smaller set of data elements. Speed up is achieved by matching each copy of a rule in parallel. Variations of this approach may be applicable to logic-based programming systems, such as PROLOG, executed in a parallel environment. Indeed, sequential implementations of OPS-style production systems based on the Rete match algorithm may enjoy performance advantages as well. This scheme may be implemented by a simple preprocessing stage which requires no modification to the underlying match algorithms.

1 Introduction

We have previously reported a number of parallel algorithms to accelerate the execution of characteristically different *Production System* (PS) programs [Stolfo 1984]. The simplest, called the *Full Distribution Algorithm*, is based on allocating each rule, as well as the *Working Memory* (WM) elements relevant to its left-hand side, to a single processing element (PE) of a large-scale, fine-grain multiprocessor, such as the DADO machine. In essence, the original PS is converted into a large number of "one-rule" PS's each processed concurrently.

For some PS programs, however, the potential speed-up of the match phase for the Full Distribution Algorithm is not nearly as great as might be expected. In programs such as R1, where few rules may potentially match newly asserted WM elements on each cycle, few PE's may perform useful work, while in programs such as ACE that work with large databases, local requirements for WM elements may exceed the capacity of some PE's. In other cases, certain anomalous rules may require more processing on average than other rules, thus producing "hot spots" of sequential execution in a distributed environment.

In this brief paper, we report a simple scheme to extract implicit parallelism in the low-level match phase of an individual rule which has the potential to improve greatly the performance of a number of different parallel PS algorithms while mitigating the problem of PE memory overflow and reducing the effects of "hot-spot" rules. The essence of the approach is to replicate anomalous rules and to introduce constraints within the copies which restrict them to match smaller, disjoint portions of the set of potentially relevant WM elements. This scheme is particularly advantageous for the Full Distribution Algorithm since it is the simplest of the entire set of reported algorithms and requires the least inter-PE communication. Before describing the load balancing scheme, we detail the operation of a production

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system program. We also describe the DADO machine and two parallel algorithms for production system execution.

2 Production Systems

In general, a Production System [Newell 1973, Davis and King 1977] is defined by a set of rules, or productions, which form the Production Memory(PM), together with a database of assertions, called the Working Memory(WM). Each production consists of a conjunction of pattern elements, called the left-hand side (LHS) of the rule, along with a set of actions called the right-hand side (RHS). The RHS specifies information that is to be added to (asserted) or removed from WM when the LHS successfully matches against the contents of WM. An example production, borrowed from the blocks world, is illustrated in figure 1 in the syntax style of OPS5 [Forgy 1981].

Figure 1: An Example Production.

(p clear-block (Goal `name Clear-top-of Block `status ON) (Physical-object `name <x> `type Block) (On-top-of `bottom-object <x> `top-object <y>) (Physical-object `name <y> `type Block) -> (delete 3) (assert (On-top-of `bottom-object Table `top-object <y>))).

If the goal is to clear the top of a block, and there is a block (x) covered by something (y) which is also a block, then remove the fact that y is on x from WM and assert that y is on top of the table.

In operation, the production system repeatedly executes the following cycle of operations:

- 1. Match: For each rule, determine whether the LHS matches the current environment of WM. All matching instances of the rules are collected in the conflict set of rules.
- 2. Select: Choose exactly one of the matching rules according to some predefined criterion.
- 3. Act: Add to or delete from WM all assertions specified in the RHS of the selected rule or perform some operation.

During the selection phase of production system execution, a typical interpreter provides conflict resolution strategies based on the recency of matched data in WM, as well as syntactic discrimination. Rules matching data elements that were more recently inserted in WM are preferred, with ties decided in favor of rules that are more specific (i.e., have more constants) than others.

In its current form, R1, an expert system for Vax computer configuration, contains approximately 2500 rules operating on a WM containing several hundred data items, describing a partially configured VAX. Running on a DEC VAX 11/780 computer and implemented in OPS5, a highly efficient production system language, R1 executes from 2 to 600 production system cycles per minute with an average performance of 10 cycles per second. Configuring an entire VAX system requires a considerable amount of computing time on a moderately large and expensive computer. The performance of such systems will quickly worsen as experts are designed with not only one to two thousand rules, but perhaps with *tens of thousands* of rules. Indeed, several such large-scale systems are currently under development at various research centers. Statistics are difficult to calculate in the absence of specific empirical data, but it is conceivable that such large systems may require an unacceptable amount of computing time for a medium size conventional computer to execute a single cycle of production system execution! Thus, we consider the design and implementation of a specialized production system machine to warrant serious attention by parallel architects and VLSI designers.

3 DADO

Simply stated, the goal of the DADO machine project is the design and implementation of a cost effective high performance rule processor, based on large-scale parallel processing, capable of rapidly executing a production system cycle for very large rule bases. The essence of our approach is to execute a very large number of pattern matching operations on concurrent hardware, thus substantially accelerating the match phase. Our goals also include increased performance via multiple rule application executed in parallel, which will not be discussed in the present paper.

DADO [Stolfo and Miranker 1984] is a fine-grain, parallel machine where processing and memory are extensively intermingled. A full-scale version of the machine would comprise a large set of processing elements (PE's) (on the order of thousands), each containing its own processor, a small amount (16K bytes, in the current prototype design) of local random access memory, and a specialized I/O switch. The PE's are interconnected to form a complete binary tree.

Within the DADO machine, each PE is capable of executing in either of two modes under the control of run-time software. In the first, which we will call SIMD mode (for single instruction stream, multiple data stream), the PE executes instructions broadcast by some ancestor PE within the tree. In the second, which will be referred to as MIMD mode (for multiple instruction stream, multiple data stream), each PE executes instructions stored in its own local RAM, independently of the other PE's. A single conventional coprocessor, adjacent to the root of the DADO tree, controls the operation of the entire ensemble of PE's.

When a DADO PE enters MIMD mode, its logical state is changed in such a way as to effectively "disconnect" it and its descendants from all higher-level PE's in the tree. In particular, a PE in MIMD mode does not receive any instructions that might be placed on the tree-structured communication bus by one of its ancestors. Such a PE may, however, broadcast instructions to be executed by its own descendants, providing all of these descendants have themselves been switched to SIMD mode. The DADO machine can thus be configured in such a way that an arbitrary internal node in the tree acts as the root of a tree-structured SIMD device in which all PE's execute a single instruction (on different data) at a given point in time. This flexible architectural design supports *multiple-SIMD* execution (MSIMD). Thus, the machine may be logically divided into distinct partitions, each executing a distinct task, and is the primary source of DADO's speed in executing a large number of primitive pattern matching operations concurrently.

A small (15 processor) prototype of the machine, constructed at Columbia University from components supplied by Intel Corporation, has been operational since April 1983. Based on our experiences with

constructing this small prototype, we have embarked on the task of implementing a larger DADO2 prototype comprising 1023 commercially available processors, which should be completed in the summer of 1985. (A 15 node single-board version of DADO2 is presently operational.) We believe that this larger experimental device will provide us with the vehicle for evaluating the performance, as well as the hardware design, of a full-scale version of DADO implemented entirely with custom VLSI circuits.

The DADO I/O switch, which is implemented in semi-custom gate array technology and incorporated within the 1023 processing element version of the machine, has been designed to support rapid global communication. In addition, a specialized combinational circuit incorporated within the I/O switch will allow for the very rapid selection of a single distinguished PE from a set of candidate PE's in the tree, a process we call *maz-resolving*. Currently, the 15 PE version of DADO performs these operations in firmware embodied in its off-the-shelf components.

4 Parallel Execution of Production Systems

On first glance it appears that each phase of the production system cycle is suitable for direct execution on parallel hardware, with the greatest opportunity for a speed-up in the match phase. This requires a partitioning of PM and WM among the available processors: some subset of processors would store and process the LHS of rules, while another possibly intersecting subset of processors would store and process WM elements. Thus, we envisage a set of processors concurrently executing pattern matching tests for a number of rules assigned to them. Similarly, once a conflict set of rules is formed, high-speed selection can be implemented in parallel as a logarithmic time algebraic operation. Finally, the RHS of a rule can be processed by a parallel update of WM. We summarize this approach by the abstract algorithm illustrated in figure 2.

Figure 2: Abstract Production System Algorithm.

- 1. Assign some subset of rules to a set of (distinct) processors.
- 2. Assign some subset of WM elements to a set of processors (possibly distinct from those in step 1).
- 3. Repeat until no rule is active:
 - a. Broadcast an instruction to all processors storing rules to begin the match phase, resulting in the formation of a local conflict set of matching instances.
 - b. Considering each maximally rated instance within each processor, compute the maximally rated rule within the entire system. Report its instantiated RHS.
 - c. Broadcast the changes to WM reported in step 3.b to all processors, which update their local WM accordingly. end Repeat;

This very simple view of the parallel implementation of the production system cycle forms the basis of our parallel algorithms previously reported and currently being implemented. For pedagogical reasons we outline two of these algorithms to illustrate the flexibility of the DADO design and to explicate our load balancing scheme.

4.1 Algorithm 1: Full Distribution of PM

In this case, a very small number of distinct production rules are distributed to each of the DADO PE's, as well as all WM elements relevant to the rules in question, i.e., only those data elements which match some pattern in the LHS of the rules. Algorithm 1 alternates the entire DADO tree between MIMD and SIMD modes of operation. The match phase is implemented as an MIMD process, whereas selection and act execute as SIMD operations.

In simplest terms, each PE executes the match phase for its own small production system. One such production system is allowed to "fire" a rule, however, which is communicated to all other PE's. The algorithm is illustrated in figure 3.

Figure 3: Full Distribution of Production Memory.

- 1. Initialize: Distribute a simple rule matcher to each PE. Distribute a few distinct rules to each PE. Set CHANGES to initial WM elements.
- 2. Repeat the following:
- 3. Act: For each WM-change in CHANGES do:
 - a. Broadcast WM-change (add or delete a specific WM element) to all PE's.
 - b. Broadcast a command to match locally. [Each PE operates independently in MIMD mode and modifies its local WM. If this is a deletion, it checks its local conflict set and removes rule instances as appropriate. If this is an addition, it matches its set of rules and modifies its local conflict set accordingly].
 - c. end do;
- 4. Find local maxima: Broadcast an instruction to each PE to rate its local matching instances according to some predefined criteria (conflict resolution strategy).
- 5. Select: Using the high-speed max-RESOLVE circuit of DADO2, identify a single rule for execution from among all PE's with active rules.
- 6. Instantiate: Report the instantiated RHS actions. Set CHANGES to the reported WMchanges.
- 7. end Repeat;

4.2 Algorithm 2: Original DADO Algorithm

The original DADO algorithm makes direct use of the machine's ability to execute in both MIMD and SIMD modes of operation at the same point in time. The machine is logically divided into three conceptually distinct components: a *PM-level*, an *upper tree* and a number of *WM-subtrees* (see figure 4). The PM-level consists of MIMD-mode PE's executing the match phase at one appropriately chosen level of the tree. A number of distinct rules are stored in each PM-level PE. The WM-subtrees rooted by the PM-level PE's consist of a number of SIMD mode PE's collectively operating as a hardware contentaddressable memory. WM elements relevant to the rules stored at the PM-level root PE are fully distributed throughout the WM-subtree. The upper tree consists of SIMD mode PE's lying above the PM-level, which implement synchronization and selection operations. Concurrency is achieved between PM-level PE's as well as in accessing PE's of the WM-subtrees. The algorithm is illustrated in figure 5.

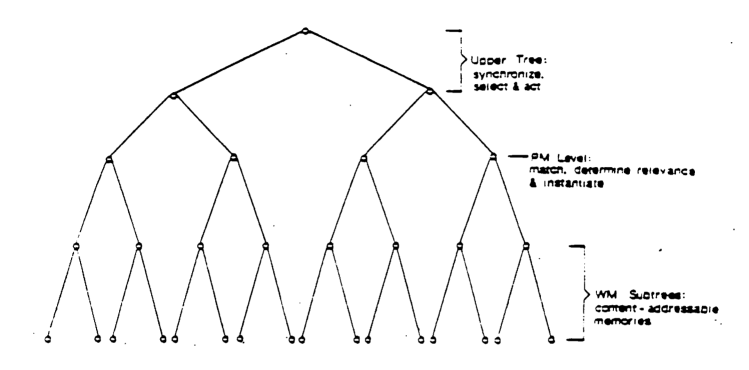


Figure 4: Functional Division of the DADO Tree.

Figure 5: Original DADO Algorithm.

- 1. Initialize: Distribute a match routine and a partitioned subset of rules to each PM-level PE. Set CHANGES to the initial WM elements.
- 2. Repeat the following:
- 3. 'Act: For each WM-change in CHANGES do;
 - a. Broadcast WM-change to the PM-level PE's and an instruction to match.
 - b. The match phase is initiated in each PM-level PE:
 - i. Each PM-level PE determines if WM-change is relevant to its local set of rules by a partial match routine. If so, its WM-subtree is updated accordingly. [If this is a deletion, an associative probe is performed on the element (relational selection) and any matching instances are deleted. If this is an addition, a free WM-subtree PE is identified, and the element is added.]
 - ii. Each pattern element of the rules stored at a PM-level PE is broadcast to the WM-subtree below for matching. Any variable bindings that occur are reported sequentially to the PM-level PE for matching of subsequent pattern elements (relational equi-join).

- iii. A local conflict set of rules is formed and stored along with a priority rating in a distributed manner within the WM-subtree.
- c. end do;

4. Upon termination of the match operation, the PM-level PE's synchronize with the upper tree.

- 5. Select: The max-RESOLVE circuit is used to identify the maximally rated conflict set instance.
- 6. Report the instantiated RHS of the winning instance to the root of DADO.
- 7. Set CHANGES to the reported action specifications.
- 8. end Repeat;

5 Copy and Constrain Rules

Measurements reported in [Gupta and Forgy 1983] show that in an average OPS5 production system, only about 32 rules are affected by changes to working memory during each production cycle. Furthermore, even if all active rules are assigned to different PE's, some PE's will take longer than others to complete the match phase. Since the select phase of the Full Distribution Algorithm cannot begin until all rules have finished matching, the total time spent in the match phase is the time taken by the slowest, not the average, rule. Simulations on OPS5 programs [Gupta 1984] indicate that because of the large variation in processing time among the affected rules, the average speed-up obtainable in the match phase from production-level parallelism is a factor of about 6. The approach we present below has the potential for transcending these limitations and increasing parallel speed-up by augmenting the number of affected rules and decreasing the variance of their processing times.

The scheme is best introduced by means of a simple example. Consider the stylized rule

 $P_1 (C_1 C_2 ... C_n \rightarrow A_1 ... A_m)$, where the $C_i (i=1,...,n)$ are condition elements, and the A_i are actions. If we interpret WM_1 (the set of working memory elements relevant to P_1 's left-hand side) as a large relation, or set of tuples, then we may view each condition element C_i as a relational selection. The set of instantiations (matches) of P_1 is the equijoin of the relations R_i selected by the condition elements C_i subject to the restriction that variables be consistently bound across all conditions. The local memory requirements and execution time to match P_1 are thus bounded by (and indeed may achieve) the size of the full Cartesian product of the individual relations R_i .

Suppose for concreteness that C_2 is a relational selection of a large number of physical objects, represented by the OPS5-style pattern:

(PHYSICAL-OBJECT 'Name $\langle x \rangle$ 'Color $\langle y \rangle$ 'Shape $\langle z \rangle$), and that the domain of the Color attribute of the relation WM_1 is {RED, GREEN}, that is, that physical objects are either RED or GREEN. To speed up the match of rule P_1 , we split the set of working memory elements associated with P_1 and set two PE's to the concurrent tasks of matching constrained versions of P_1 . We thus construct two new condition elements:

 C'_{2} (PHYSICAL-OBJECT 'Name <x> 'Color RED 'Shape <z>)

 C''_2 (PHYSICAL-OBJECT 'Name <x> 'Color GREEN 'Shape <z>),

two new rules:

 $\begin{array}{c} \mathsf{P'}_1 \; (\mathsf{C}_1 \; \mathsf{C'}_2 \; \ldots \; \mathsf{C}_n \; -> \; \mathsf{A}_1 \; \ldots \; \mathsf{A}_m) \\ \mathsf{P''}_1 \; (\mathsf{C}_1 \; \mathsf{C''}_2 \; \ldots \; \mathsf{C}_n \; -> \; \mathsf{A}_1 \; \ldots \; \mathsf{A}_m). \end{array}$

and two new working memories:

 $WM'_1 = \{ w \text{ in } WM_1 : Color(w) = RED \text{ if } w \text{ is a PHYSICAL-OBJECT} \}$

 $WM''_1 = \{ w \text{ in } WM_1: Color(w) = GREEN \text{ if } w \text{ is a PHYSICAL-OBJECT} \}$

and assign them to distinct PE's, PE'₁ and PE''₁. P'₁ and P''₁ may clearly be matched in parallel, and the set of instantiations of P₁ is exactly the disjoint union of the instantiations of P'₁ and P''₁.

In the best case, half the tuples selected by the original condition C_2 of rule P_1 reside in each of PE'_1 and PE''_1 , and the processing time required to match P_1 decreases by half, since the two new rules can be matched in parallel. Local processing requirements and storage of WM elements for each rule decrease significantly as well. In the worst case, of course, all the tuples selected by C_2 of rule P_1 reside in one of the two PE's, PE'_1 and PE''_1 , and partitioning buys nothing. If more PE's are available, the scheme can be applied repeatedly, producing many copies of rules, each constrained to match a smaller range of distinct WM elements. Thus, with many PE's available, it should be possible to reduce the inter-PE variation in processing times, balancing the execution load over the entire system and increasing overall performance dramatically.

If a small finite domain of attribute values is not known a priori (as in the above example with RED and GREEN objects), two variations on the technique are possible. The first is *hash partitioning*. Suppose that in the above example, the domain of the Color field is not {RED, GREEN}, but is some (possibly infinite) domain D. If there is an easily computable function

f: $D \to \{1,...,k\},\$

we can split WM₁ into k partitions

 $WM_1(j) = \{ w \text{ in } WM_1: f(Color(w)) = j \text{ if } w \text{ is a PHYSICAL-OBJECT} \}, (j=1,...,k), \}$

and assign each of the k partitions to a separate PE along with a suitably constrained version of the rule P_1 . In the best case, again, the processing time for the match phase of P_1 is divided by k, the number of partitions. Thus, our scheme is similar in many respects to hash partitioning tuples in a single relational query executed iteratively on relations exceeding the size of main memory. However, in our case hash partitioning is applied in parallel to a large number of concurrent queries operating on a large number of small relations.

The second variation of the basic technique applies when the domain is a totally ordered set: we can simply split it into disjoint subranges. Continuing with the above example, suppose that working memory elements have the form

(PHYSICAL-OBJECT `Name $\langle x \rangle$ `Reflected-Wavelength $\langle y \rangle$ `Shape $\langle z \rangle$), and that a set of values

 $v_{\min} = v_0, v_1, \dots, v_k = v_{\max}$

of Reflected-Wavelength are given. We can again split working memory, this time into

 $WM_1(j) = \{ w \text{ in } WM_1 :$

Reflected-Wavelength(w) $>= v_{i-1}$ and Reflected-Wavelength(w) $< v_i$

if w is a PHYSICAL-OBJECT}, (j=1,...,k),

and assign each to a separate PE together with constrained versions of the original rule P_1 . This disjointsubranges scheme is a special case of hash partitioning that bypasses the explicit computation of a hashing function.

Rule partitioning can be incorporated into both the Full Distribution Algorithm and the original DADO algorithm without modifying either one's implementation. The copy and constrain idea would make more PE's perform useful match computation in the full distribution case simply by copying rules. In the original DADO algorithm, the scheme would copy rules to different PM-level PE's; in this case, the WMsubtrees would of course store disjoint subsets of the original subset of WM held in a single prepartitioning WM-subtree.

8 Implementation Outline

The scheme outlined can be implemented by a simple preprocessor supplied with information on how to partition the domains of working memory attributes. These pragmas, or hints, can take the form of explicit values or ranges provided by the programmer (derived from knowledge of the problem or from previous executions of the production system program), or can include hashing functions. The preprocessor's role is simply to generate new productions incorporating the value, hashing function, or subrange tests, one for each value, function value, or subrange supplied.

OPS-style productions are easily modified by adding partitioning information to *literalize* declarations. The declarations for PHYSICAL-OBJECT in our first example (when a small finite domain of attribute values is known) might be

> (literalize PHYSICAL-OBJECT Name Color (symbol RED GREEN) Shape)

To specify a hashing function defined on the Color field with range $\{1,...,k\}$, we could write

(literalize PHYSICAL-OBJECT

Name Color (hash hash-function-name k) Shape)

Specifying k subranges of a totally ordered domain is just as easy:

(literalize PHYSICAL-OBJECT

Name Reflected-Wavelength (range $v_1 \dots v_{k-1}$) Shape)

The preprocessor should split each rule containing non-constant tests on the partitioned fields into the appropriate set of more specialized rules.

This approach has the potential of greatly improving the performance of a variety of PS programs including those with thousands of rules and hundreds of WM elements, as for example R1, and conversely those with hundreds of rules and thousands of WM elements, as for example ACE. Other coarse-grained approaches to the parallel execution of PS programs may make effective use of this scheme as well. The Rete match algorithm, for example, would thus compile additional match nodes for the introduced partitioning constraints which would effectively reduce the size of the Beta memories of the original condition element. The resultant sequential search of the Alpha and Beta memories to compute partial match results would thus be quicker since fewer data elements would be compared with each other.

Indeed, logic-based programming systems may also make use of similar approaches to load balance execution by introducing copied and constrained clauses. In a PROLOG environment, the idea may be used as follows: If in a rule

a:- b,c,d

it is known via a pragma that goal b dominates the computation for satisfying goal a, then rewrite the rule as two rules:

a:-b',c,d a:-b'',c,d

where b' and b'' are copied versions of goal b with constraints on the first order terms to unify with a smaller and distinct set of terms that would match b. Since goals b' and b'' would be executed in parallel in an OR-parallel environment, a speed up is achievable. One approach to figuring out the constraints for goals b' and b'' is to execute the PROLOG program symbolically in order to identify the range of unit literals that terminate the goal tree emanating from the posting of goal b. The range of first order literals so identified may then be partitioned by suitably constraining the first order terms of literal b.

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