

Branching Matters: Alternative Branching in Graphplan

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Abstract

Graphplan can be understood as a heuristic search planner that performs an IDA* regression search with a heuristic function encoded in the plan graph. In this paper we study two alternatives to Graphplan where the IDA* search algorithm and the heuristic encoded in the plan graph are maintained but the branching scheme is changed: rather than constructing plans from the tail, commitments are allowed anywhere in the plan. These commitments force certain actions *in or out* of certain time steps. While the regression search allows Graphplan to build the plan graph only once, in the new branching scheme the plan graph must be computed anew for each state. This re-computation is expensive but is often compensated by a reduced number of states in the search. We show through a number of experiments that the resulting planner scales up much better than the original Graphplan in domains involving a higher degree of parallelism, and compares well to state-of-the-art domain-independent optimal parallel planners.

Introduction

Heuristic search has been shown to be an effective approach in AI Planning. In this approach, plans are searched for with the help of a heuristic function that is extracted automatically from the problem encoding (McDermott 1999; Bonet & Geffner 2001). Most current domain-independent planners make use of heuristic estimators of some sort, and even planners such as Graphplan, that have not been designed from this perspective, can be understood in those terms (Bonet & Geffner 1999).

Current heuristic search planners, however, are strong for optimal and non-optimal sequential planning (Haslum & Geffner 2000; Hoffmann & Nebel 2001) but weaker for parallel planning. In optimal parallel planning, the task is to compute a plan that involves a minimum number of time steps where in each time step many actions can be done in parallel. Optimal parallel planning is the basic type of planning supported by current Graphplan and SAT planners (Blum & Furst 1995; Kautz & Selman 1999).

An important obstacle faced by current heuristic search planners in the parallel setting is the *high branching factor*. Indeed, in heuristic regression based planners such as

Graphplan and HSP^r* (Haslum & Geffner 2000), the number of sets of concurrent actions that need to be considered in each state s grows *exponentially* with the degree of parallelism: one set for each combination of supports for the atoms in s . It is thus not surprising that the best current optimal parallel planners are based on either SAT or CSP formulations where branching is accomplished by trying the different values of selected boolean variables, and hence where the branching factor is always two (Kautz & Selman 1999; Rintanen 1998; Baiocchi, Marcugini, & Milani 2000; Do & Kambhampati 2000). Of course, a small branching factor by itself is not sufficient for good performance, and indeed, these SAT and CSP approaches supplement the *branching strategy* with effective *pruning mechanisms* based on constraint propagation and consistency checking.

In this work, we deal with the problem of optimal parallel planning from the perspective of heuristic search. However, rather than searching for plans either forward from the initial state or backward from the goal, we consider a branching scheme that is closer to the one found in SAT and CSP approaches: we pick an action variable a and a time point i , and branch making the action a at time i either true or false. States σ in the search are thus no longer sets of *atoms* but sets of *commitments*. Yet, as before, we use a heuristic function $h(\sigma)$ to provide a lower bound on the cost of achieving the goals given the commitments in σ , and prune σ from the search when $h(\sigma)$ exceeds the IDA* bound.

More precisely, we define a *branch and bound* planner, that we call BBG (Branch-and-Bound-Graphplan), on top of the IPP implementation (Koehler *et al.* 1997) of Graphplan. As argued in (Bonet & Geffner 1999) and (Haslum & Geffner 2000), Graphplan can be understood as a heuristic search planner that searches for parallel plans from the goal using the IDA* search strategy and a heuristic that is implicitly encoded in the plan graph. BBG retains from Graphplan the same IDA* search strategy and the same plan graph heuristic function, but changes the branching strategy allowing commitments anywhere in the plan to be made. In the resulting search space, the heuristic is obtained by *recomputing the plan graph in each node*. While this re-computation is expensive, we show that it is often compensated by a reduced number of states in the search. Indeed, the resulting planner scales up much better than the original Graphplan in domains involving a higher degree of parallelism where it

compares well against state-of-the-art domain independent parallel planners such as Blackbox (Kautz & Selman 1999) using Chaff (Moskewicz *et al.* 2001).

The notion of branching in AI planning has been often analyzed implicitly in terms of the *space* searched during plan construction. Namely, linear planners (including progression and regression planners) are said to search in the space of *states*, while non-linear planners are said to search in the space of *plans* (e.g., (Weld 1994)). This is a useful distinction, yet as discussed in more detail in (Geffner 2001), focusing explicitly on the notion of branching sheds light also on the relation between linear and non-linear planning on the one hand, and SAT/CSP formulations and other well-known combinatorial optimization problems on the other. For example, the best codes for the Traveling Salesman Problem do not branch by considering all possible next cities - as would correspond to a linear planner - but by selecting an edge in the graph, and forcing this edge *in* or *out* of the solution (Balas & Toth 1985). Something similar occurs in the Job Shop Problem (Carlier & Pinson 1989).

Non-directional branching schemes in the Graphplan setting have been introduced and analyzed in (Rintanen 1998) and (Baiocchi, Marcugini, & Milani 2000) among other proposals (see also (Kautz & Selman 1999) and (Do & Kambhampati 2000) for SAT and CSP translations). What is particular to the approach that we pursue in this paper is that we introduce an alternative branching scheme in Graphplan while *retaining Graphplan's IDA* search algorithm and plan graph lower bounds*. This allows us to draw conclusions about the different branching schemes and not only about the resulting planners. Indeed, while the above proposals combine new branching schemes with new pruning mechanisms (based on some form of constraint propagation), our proposal combines a new branching scheme with a pruning mechanism directly inherited from Graphplan. The result is that differences in performance are not the result of carefully crafted propagation rules but are exclusively the result of the different branching schemes. The importance of controlled experimentation in the evaluation and analysis of algorithms is discussed in (Hooker 1996).

The paper is organized as follows. The next section outlines the Graphplan algorithm and its interpretation as a heuristic search planner. We then include two sections describing the new branching schemes and the branching selection criteria. Then, after providing an overview of the implementation, we evaluate the approach empirically and discuss the results. A short appendix provides additional details on the implementation.

Graphplan and Heuristic Search

We provide a brief summary of Graphplan, see (Blum & Furst 1995) for details. We use the letters I and G to denote the initial and goal situations of a STRIPS planning task; actions a are the usual triples $pre(a)$, $add(a)$, $del(a)$, and for each proposition p , we assume the presence of a 'dummy' action $a = NOOP(p)$ with $pre(a) = add(a) = \{p\}$ and $del(a) = \{\}$. Graphplan builds a *plan graph* which is a layered graph consisting of *fact layers* and *action layers*. Fact layer 0 is the initial state I , and action layer 0 is the set

of actions (including NO-OPs) whose preconditions are in I . Fact layer i , for $i > 0$, contains the add effects of actions in layer $i - 1$, and action layer i , for $i > 0$, contains the actions whose preconditions are in fact layer i without a mutex. Mutex relations are defined as follows. A pair of actions in action layer 0 is *mutex* if they *interfere*, i.e., if one action deletes a precondition or add effect of the other (a non-interfering pair of actions is called *compatible*). A pair of facts at layer $i > 0$ is mutex if there is no non-mutex pair of actions at layer $i - 1$ achieving the facts. A pair of actions at a layer $i > 0$ is mutex if the actions interfere, or if they have *competing needs*, i.e., mutex preconditions. A set of facts is *reachable* at layer i if all these facts are contained in layer i and no pair of them is mutex. From the first layer where G is reachable, Graphplan starts a regression search, where a set of (sub)goals s at layer i yields a new set of subgoals s' at layer $i - 1$ for each set of pairwise non-mutex actions (possibly including NO-OPs) that achieve s . In that case, s' is the union of those actions preconditions. The search terminates successfully when the initial layer is reached, or negatively when all options have been explored unsuccessfully. In the latter case the graph is extended by another layer, and the search continues. When the search beneath a subgoal set s at layer i fails, s is *memoized*, i.e., it is remembered as unsolvable at layer i .

As argued originally in (Bonet & Geffner 1999), and further elaborated in (Haslum & Geffner 2000), Graphplan can be understood as a heuristic search planner with the following features:

1. **Branching:** States are sets of atoms (subgoals), and the children s' of a state s are obtained by parallel regression with NO-OPs as described above. The initial state in the search is G and the goal states s are such that $s \subseteq I$.
2. **Bounds:** The plan graph encodes an *admissible heuristic* h_G for parallel planning, estimating the parallel cost (number of time steps) for reaching s from I , where $h_G(s) = i$ iff i is the index of the first fact layer where s is reachable and not memoized (memoizations are updates on the heuristic function; see 4).
3. **Search Algorithm:** The search algorithm is an enhanced version of IDA* (Korf 1985), where the sum of the accumulated cost $g(s)$ (distance from G), and the estimated cost $h_G(s)$ (distance estimate to I) is used to prune states s whose cost exceeds the current bound (the index of the current top graph layer). Actually Graphplan never generates such nodes. The algorithm takes advantage of additional information stored in the plan graph to convert the search into a more efficient 'solution extraction' procedure. For example, compatible but mutex (due to competing needs) pairs of actions are not used for regressing a state; the resulting state would indeed have a cost exceeding the IDA* bound.
4. **Memoization:** Memoizations are updates on the heuristic function h_G . If a state s is expanded and the search beneath it fails, then the optimal cost $h^*(s)$ must be strictly greater than $h_G(s)$ (otherwise a solution through s would have been found given the same bound), and hence the estimated cost $h_G(s)$ can be increased. This

is a common extension in modern IDA* implementations which refer to the technique as *transposition tables*; e.g., (Sen & Bagchi 1989; Reinfeld & Marsland 1994; Junghanns & Schaeffer 1999; Haslum & Geffner 2000).

The dramatic improvement in performance that Graphplan brought to AI planning in the middle 90's can be traced from this perspective to the use of an informative *heuristic function*. Indeed, previous partial order planners, unlike Graphplan, perform *blind* search (using a different branching scheme).

Branching

The search for optimal solutions in planning, and more generally, in combinatorial optimization, is the result of two main operations: *branching* and *pruning*.¹ Branching refers to the scheme used for generating the child nodes in the search, while pruning refers to the criterion used for eliminating some of them from consideration. In Graphplan, *branching* is done by parallel regression, and *pruning* is done when the lower bound $f(s) = g(s) + h_G(s)$ exceeds the current bound. SAT and CSP approaches rely on the same search strategy, IDA*, but branch and prune in a different way: *branching* is done by trying the different values of a selected variable, and *pruning* is done when the choices, along with the problem constraints and the bound condition, yield an inconsistency by constraint propagation. Branching and pruning, however, are largely two orthogonal operations, and in this paper we aim to explore alternative branching schemes to Graphplan's parallel regression, *leaving its pruning technique, and in particular its plan graph heuristic estimator, untouched*.

Branching on Actions

In the first branching scheme we consider, *branching on actions*, an action a and time d (the decision layer) smaller than or equal to the current bound are selected, and a *binary split* is generated: one in which a is made true at time d , the other where it is made false. A state σ in this search is a set of choices or commitments of this type. For example in the blocks world, a state σ will contain a set of moves to be done at various time points, and a set of moves *not* to be done at possibly other time points.

We let $h^*(\sigma)$ refer to the cost of the best (parallel) plan that is compatible with the commitments contained in σ . Since computing the exact cost is intractable ($h^*(\emptyset)$ is the length of an optimal parallel plan), we compute a lower bound $h_G(\sigma) \leq h^*(\sigma)$. This lower bound (admissible heuristic) is obtained by building a plan graph as in Graphplan *but* respecting the commitments made in σ ; namely, if $a[d] = \mathbf{true} \in \sigma$, then action a must be in action layer d of the plan graph for σ and all actions mutex with a at d must be excluded from that layer. Similarly, if $a[d] = \mathbf{false} \in \sigma$, action a is excluded from layer d in the plan graph for σ .

¹This applies to linear-space search algorithms like IDA* and DFS Branch and Bound, but not to exponential-space algorithms like A* that, in any case, are not feasible for large problems. See (Korf 1993).

As for Graphplan, $h_G(\sigma)$ is given by the index of the first layer in the plan graph for σ that contains the goals without a mutex. The initial state $\sigma = \sigma_0$ is the empty set of commitments, and thus $h_G(\sigma_0)$ is computed from exactly the same plan graph that is constructed by Graphplan. For other states, the plan graph is constructed incrementally from the parent graph by retracting one or more actions in the corresponding layer, and propagating this retraction forward. A state σ in this search is *pruned* when $h_G(\sigma)$ exceeds the current bound or when the plan graph construction procedure cannot comply with the commitments made (the latter happens when a new commitment at layer d makes the preconditions of an action a previously committed to a layer i , $i > d$, either unachievable or mutex). The terminal states in this search are those for which the plan graph contains no 'flaws', and hence, where a valid plan can be extracted without backtracking. This is explained more precisely below, when the selection criteria for choosing the action a and time d for branching are presented.

Branching on Mutexes

In the second branching scheme that we consider, *branching on mutexes*, a pair of actions a_1 and a_2 that are mutex at time d are selected, and a *three way split* is generated: one in which a_1 is made true at d , one in which a_2 is made true at d , and finally, one in which a_1 and a_2 are both false at d . As before, if a state σ contains a commitment of the form $a[d] = \mathbf{true}$, then the plan graph for σ will have all actions that are mutex with a at d excluded. The heuristic function, pruning, and initial state are the same as those for branching on actions.

The choice of the two branching schemes above is somewhat arbitrary: we could as well have done branching not on actions but on facts or both (as in SAT). In principle, any partition of the space of plans that could be reflected in the plan graph construction would do. Ours is thus an initial and not an exhaustive exploration of alternative branching schemes in the Graphplan setting.

Branching Selection Criteria

We refer to the choice of the action (mutex) and time step on which to branch as the branch point selection. Like the similar variable-selection heuristic criterion in constraint-based search, the branch point selection criterion does not affect the soundness or completeness of the branching scheme, but affects the size of the search tree and thus is critical for performance. After some limited empirical exploration, we settled on two goal-oriented criteria that select branching points in order to remove *flaws* in a *relaxed plan*; a state σ for which the relaxed plan contains no flaw is a *terminal state* from which a valid parallel plan can be extracted backtrack free. The flaw identification and repair loop in BBG is reminiscent of a similar loop in partial-order planning (Weld 1994); yet the nature of the flaws and repairs, as well as the pruning criteria, are different.

A relaxed plan in a state σ is defined as in the FF planner (Hoffmann & Nebel 2001): as a solution extracted from the plan graph built in σ ignoring mutex relations between

actions. Thus, construction of a relaxed plan from the plan graph in σ is backtrack free and fast. While there may be an exponential number of relaxed plans in σ , we care only about the first relaxed plan found, that we call the relaxed plan for σ . Since we would like this relaxed plan to have as few flaws as possible, we extract it by ordering the atoms and their supporters in each layer so that 1) atoms with smaller number of supporters are regressed first, and 2) no-ops are chosen for support if possible, otherwise, a regular action is chosen greedily, minimizing the number of mutex relations with supporters already selected.

The relaxed plan, like a regular parallel plan, is a set of action occurrences (a, i) where a is an action and i is a time point. Unlike regular plans, however, the relaxed plan explicitly represents the occurrence of NO-OP actions.

A *flaw* in the relaxed plan is a triplet (a, a', i) where (a, i) and (a', i) are two action occurrences in the relaxed plan such that a and a' are mutex at i .

The first selection criterion for choosing branching points in both branching schemes focuses on flaws that occur *latest* in the relaxed plan, i.e. the flaws (a, a', d) with maximal time index d . In both branching schemes, an arbitrary latest flaw (a, a', d) is chosen (actually, the first flaw found in the relaxed plan extraction procedure). When branching on mutexes this flaw forms the new branching point; when branching on actions one of (a, d) or (a', d) is chosen arbitrarily (the action with the lower internal index). We have tried other simple static selection criteria (earliest flaws, middle flaws, etc) but the experiments didn't show improvements, and in certain cases showed a clear deterioration. The focus on the 'latest flaws' does not mean that we are back to a regression search as these are the latest flaws in the relaxed plan and not in the plan graph itself. Criteria for choosing branching points that do not make use of the relaxed plan do not appear to do as well.

We consider also a dynamic selection criterion based on estimating the *criticality* of action occurrences in the relaxed plan. The *criticality measures* $c(a, i)$ are computed in a single backward pass over the plan graph and aim to approximate the relative number $c^*(a, i)$ of relaxed plans in which a occurs at time i , a number that ranges from $1/NR$ to 1, where NR is the number of relaxed plans.² Ideally, we would like to choose a branching point that removes highly critical actions in all its branches. For that, let b be a branching point, and let b_1, \dots, b_n , refer to the commitments made in each of its branches ($n = 2$ when branching on actions and $n = 3$ when branching on mutexes). We define the criticality of a commitment b_j , $c(b_j)$, as the measure $c(a, i)$ of

²Criticality measures are computed for both facts (p, i) and actions (a, i) as follows. The goal facts g are assigned a measure $c(g, m) = 1$ where m is the current IDA* search bound. The process then proceeds backwards from graph layer m to graph layer 0. If a fact f at layer i has achievers $a_1 \dots a_n$ at layer $i - 1$ then it propagates a measure $c(f, i)/n$ into each of these actions. The measure $c(a, i)$ of an action a at layer i is then the max measure it gets from the facts at layer $i + 1$. The action propagates in turn its own measure $c(a, i)$ into its preconditions at i . The measure $c(f, i)$ of a fact at layer $i < m$ is the max measure it gets from the actions a at layer i .

the most critical action-time pair a, i that is forced out by b_j ,³ and the criticality of a branching point b , $c(b)$, as the product $c(b_1) \cdot \dots \cdot c(b_n)$.⁴ When branching on mutexes, we select for branching the most critical relaxed plan flaw $b = (a, a', d)$, and when branching on actions, we select for branching the most critical branching point $b = (a, d)$ that occurs in a relaxed plan flaw (a, a', d) . Ties, as before, are broken arbitrarily.

Implementation Overview

Since the plan graph must be recomputed in every single search state, an incremental implementation of this re-computation is crucial for performance. Our implementation builds on the ideas introduced in STAN (Long & Fox 1999), following their realization in IPP (due to some minor changes in the re-implementation our graph building procedure is in fact somewhat more efficient than IPP's in most of the cases we tried). Here, we provide a brief overview over the implementation; see the appendix for further details.

The plan graph is represented as a single layer of facts and actions, where only the layer-dependent information is updated along the time steps. When initially building the graph, all the optimizations described in (Long & Fox 1999) are applied in order to avoid unnecessary re-computations of the same piece of information (such computations are surprisingly plentiful in the original Graphplan). Information that must be accessed frequently, such as mutex information, is kept in bit vectors.

When a new constraint is introduced at a layer d , the graph is updated by removing actions at layer d and propagating forward the changes. In general, we have a layer i , a set *ops* of actions to be removed at i , and a set *opmutexes* of action mutexes to be introduced at i . These changes are propagated to the next fact layer, i.e., the actions *ops* are removed from the possible achievers for the respective facts at layer $i + 1$, and the mutexes *opmutexes* are introduced (all these operations are done implicitly by updating bit vectors and setting flags). In consequence, facts can become unachievable or mutex. We thus get a set *fts* of facts to be removed at $i + 1$, and a set *ftmutexes* of fact mutexes to be introduced at $i + 1$. Propagating these changes to action layer $i + 1$ amounts to removing actions which need a fact in *fts*, or a fact pair in *ftmutexes* as precondition, and introducing mutexes between actions which now have competing needs due to *ftmutexes*. So we get a list *ops* of actions to be removed at $i + 1$, and a set *opmutexes* of action mutexes to be introduced at $i + 1$. We are now back in the situation we started from, so the algorithmic loop is closed. The loop terminates when the current deadline layer is reached. If the goals are not reachable in the deadline layer then it is known that the

³More precisely, if the commitment b_j in a state σ is $a[i] = \text{true}$, then $c(b_j)$ is defined as $\max_{a'} c(a', i)$ over all actions a' that are mutex with a at time i in the plan graph corresponding to σ . Similarly, if the commitment b_j is $a[i] = \text{false}$ for one or more actions a , then $c(b_j)$ is defined as $\max_{a'} c(a', i)$ over all such actions a .

⁴Similar product aggregation functions are used in the popular MOM variable selection heuristic in SAT.

heuristic value is above the deadline and thus the node is pruned. All changes made during the graph update are kept in memory so they can easily be retracted upon backtracking.

Results

The new branching schemes aim to improve the performance over highly-parallel domains where the branching factor in Graphplan and similar parallel heuristic search planners grows exponentially (see (Haslum & Geffner 2000) for a discussion). We have thus compared BBG with IPP over two different suits of domains: one containing domains that are highly parallel, the other containing domains that are mostly serial. The results, as we will see, show a significant improvement in the former, and little, if any improvement (and in certain cases a deterioration) in the latter. We also evaluate how close the change in the branching scheme takes the 'new Graphplan', BBG, in relation to state of the art domain-independent parallel planners such as Blackbox using Chaff. At the end of this section we include such a comparison.

All experiments were conducted on a notebook using a Pentium-III processor running at 850 MHz with 256 MBytes main memory. The source code and testing examples are available from the authors.

Parallel Domains

The first parallel domain we consider is the classical *Logistics* domain where packages must be transported between locations by means of trucks (within cities) and aeroplanes (between different cities). The parallelism lies in that moving actions for different vehicles, and loading / unloading actions for different packages, do not interfere with each other.

We obtained a suit of random *Logistics* instances using the generator available from the FF homepage.⁵ We generated instances scaling in 5 different sizes, 10 instances per size. The instances all had 4 cities with 2 locations each, 1 truck per city, and 2 aeroplanes. Size was scaled by increasing the number of packages to be transported from 7 to 15 in steps of 2 (we chose these specific size parameters after some experimentation in order to make the resulting instances challenging but feasible for the planners in question; in comparison, the well-known *Logistics* instance "log-c", e.g., corresponds exactly to our smallest instance set). See the data in Table 1.

The six planners we tested are the following: BBG-BA-* is BBG branching on actions, BBG-BM-* is BBG branching on mutexes; BBG-*-L uses the latest flaw selection, while BBG-*-C uses the criticality flaw selection. IPP, in turn, is the IPP 4.0 implementation available from the IPP homepage;⁶ IPP-N is the same implementation but with memoization turned off — we include this data as comparing it to BBG, in which no transposition tables are used either, yields a more immediate picture of what the effects of our new branching techniques are.

planner	Logistics				
	size 1	size 2	size 3	size 4	size 5
BBG-BA-L	1.00	1.00	0.90	1.00	0.70
BBG-BM-L	1.00	1.00	0.90	0.70	0.70
BBG-BA-C	1.00	1.00	1.00	1.00	0.80
BBG-BM-C	1.00	1.00	1.00	1.00	0.80
IPP	1.00	0.80	0.20	0.00	0.10
IPP-N	0.90	0.40	0.20	0.00	0.00

Table 1: Fraction of solved instances in the *Logistics* domain. See explanation in the text.

Each planner was given a time bound of 500 seconds and a memory bound of 256 MBytes. Table 1 shows the fractions of the 10 instances per size that were solved within these bounds by the respective planners. Clearly, all BBG configurations scale up much better than the IPP configurations. We also measured the search time spent and the number of search states encountered by the planners. We do not include this data in the tables for two reasons. First, the fractions of solved instances are often sufficient to rank the planners in the experiments. Second, time averages in the presence of a significant number of unsolved instances are a problem (should unsolved instances be ignored? should they be weighted by the cut-off time/space? ...). For these reasons, we report only some of these data in the text, when directly comparing pairs of planners where the time and states averages are taken over the instances that are solved by both planners (ignoring the rest).

In the *Logistics* suit, all BBG configurations behave roughly similar with a slight advantage for BBG-BM-C. On the instances solved by both BBG-BM-C and IPP, the average search times per size are 0.19, 0.45, 0.34, -, 0.31 seconds for BBG-BM-C, and 1.72, 117.16, 18.16, -, 447.11 seconds for IPP (in the second largest size class, no instance is solved by both planners). These measurements are pure search time, i.e., they do *not* include the initial planning graph building phase which, as said before, is somewhat more efficient in BBG due to minor implementation differences. The average numbers of search states visited are 24, 77, 28, -, 22 for BBG-BM-C, and 5786, 104818, 33646, -, 239888 for IPP. The search states we count in BBG are the ones in which a commitment has been made successfully (without exceeding the current bound on the heuristic function), while in IPP the states we count are the successfully created subgoal sets (the subgoal sets produced after successful backchaining with non-mutex actions that are not memoized). Clearly, the reduction in the number of states in the search is dramatic in this case (several orders of magnitude), while the reduction in search time is significant, although not as large, due to the additional overhead incurred by BBG in the computation of the heuristic. In comparison to IPP-N, the data are about an order of magnitude more drastic: comparing BBG-BM-C to IPP-N, the search time values are 0.19, 0.22, 0.33, -, - for BBG-BM-C and 2.13, 27.97, 296.03, -, - for IPP-N. The search state numbers are 23, 20, 28, -, - respectively 37529, 387862, 1998143, -, -. We have made similar observations — i.e., that the data in

⁵At <http://www.informatik.uni-freiburg.de/~hoffmann/ff.html>.

⁶At <http://www.informatik.uni-freiburg.de/~koehler/ipp.html>.

direct comparison to IPP-N are around an order of magnitude more drastic than in direct comparison to IPP — in all the other parallel domains below; we do not report all this data as it seems redundant.

Our second parallel domain is a variant of the *Blocksworld* where blocks can be moved in a move-A-from-B-to-C fashion, thus enabling parallel moves of different blocks to different destinations. Our testing suit contains random instances (generated with the generator available from the FF homepage) with 6 to 14 blocks, increasing in steps of 2, 10 instances per size. See the data in Table 2.

<i>Blocksworld</i>					
planner	size 1	size 2	size 3	size 4	size 5
BBG-BA-L	1.00	1.00	1.00	1.00	0.90
BBG-BM-L	1.00	1.00	1.00	0.90	0.90
BBG-BA-C	1.00	1.00	1.00	0.60	0.70
BBG-BM-C	1.00	0.70	0.60	0.40	0.30
IPP	1.00	1.00	1.00	0.80	0.60
IPP-N	1.00	1.00	1.00	0.70	0.60

Table 2: Fraction of solved instances in the *Blocksworld* domain. See explanation in the text.

The tested planners, the time and memory bounds in the experiment, and our form of data presentation are the same as explained above for the *Logistics* experiment. As can be seen from Table 2, in the *Blocksworld*, the branching scheme and branching point selection criterion used make a big difference. BBG configurations with latest flaw selection (-L) solve more instances than IPP and are more efficient. The average search times on those instances solved by both BBG-BA-L and IPP are 0.09, 0.20, 0.29, 1.47, 1.53 for BBG-BA-L and 0.00, 0.01, 0.05, 1.44, 7.12 for IPP. The average number of states visited are 5, 31, 17, 73, 52, and 6, 60, 80, 408, 3060 respectively. The reduction in the number of states is smaller in this domain but still significant. On the other hand, the time reduction almost vanishes in this case, due to the larger overhead per node in BBG (once again, note that these figures are averages over the instances that IPP found relatively easy, ignoring the instances solved by BBG but not by IPP). As in *Logistics*, the data in comparison to IPP-N is an order of magnitude more drastic.

Our third test is over the *Satellite* domain as used in the 3rd International Planning Competition (IPC-3) (Fox & Long 2002). A number of satellites must take images of phenomena in space, which involves, among other things, turning the satellites into the right direction, and calibrating the instruments. Individual satellites can act in parallel. Our testing suit contains random instances (generated with the official competition generator) with 5 satellites each and 6 to 10 images to be taken, 10 instances per size (we will later also present results for random *Satellite* instances with only 1 satellite, where there is less parallelism). See the data in Table 3.

The difference in performance that branching makes in this case is very significant (see Table 3). All instances where more than 7 images must be taken are solved by BBG while none is solved by IPP. The smallest instances, on the

<i>Satellite</i> — parallel suit					
planner	size 1	size 2	size 3	size 4	size 5
BBG-BA-L	1.00	1.00	1.00	1.00	1.00
BBG-BM-L	1.00	1.00	1.00	1.00	1.00
BBG-BA-C	1.00	1.00	1.00	1.00	1.00
BBG-BM-C	1.00	1.00	1.00	1.00	1.00
IPP	1.00	1.00	0.00	0.00	0.00
IPP-N	1.00	1.00	0.00	0.00	0.00

Table 3: Fraction of solved instances in the *Satellite* domain, (more) parallel random suit. See explanation in the text.

other hand, are simple and are solved by IPP with no search (BBG solves them as well). The branching scheme and branching point selection make a lot of difference in this case, with advantages for branching on actions (-BA) and latest flaw selection (-L): the average number of states is 9, 110, 1821, 3557, 4567 for BBG-BA-L, and 8, 190, 12789, 58568, 100817 for BBG-BM-C.

The fourth domain is *Depots*, which also originates from the IPC-3 suit. It is a combination of *Logistics* and *Blocksworld* where objects (“crates”) must be transported in trucks and then be arranged in stacks using a number of robot arms, “hoists”. The robot arms can act independently of each other. In all our random examples (generated with the official competition generator) there are 2 locations, and 2 trucks. One can use 7 hoists to arrange 6 to 10 crates, 10 instances per size (we will later also present results for random *Depots* instances with only 2 hoists, where there is less parallelism). See the data in Table 4.

<i>Depots</i> — parallel suit					
planner	size 1	size 2	size 3	size 4	size 5
BBG-BA-L	1.00	1.00	1.00	1.00	0.70
BBG-BM-L	1.00	1.00	1.00	1.00	0.60
BBG-BA-C	1.00	0.60	0.30	0.20	0.20
BBG-BM-C	1.00	0.60	0.30	0.00	0.10
IPP	0.80	1.00	1.00	1.00	0.10
IPP-N	0.80	1.00	0.90	0.80	0.10

Table 4: Fraction of solved instances in the *Depots* domain, (more) parallel random suit. See explanation in the text.

In Table 4, we see again that BBG configurations that use latest flaw selection (-L) perform better. Comparing BBG-BA-L to IPP we obtain average search times 0.81, 1.22, 6.24, 5.06, 1.40 for BBG-BA-L and 0.39, 0.59, 19.07, 78.69, 11.08 for IPP (the single largest instance solved by IPP thus appears to be relatively easy). The number of search states is 235, 199, 435, 662, 66 for BBG-BA-L and 1076, 860, 15432, 21719, 24422 for IPP.

The last two domains we consider are *Mystery* and *Mprime*, both introduced by Drew McDermott in the 1st International Planning Competition. These are *Logistics* variations where trucks consume one fuel unit per move. The fuel units are associated with the locations (rather than with the trucks), and moving away from a location decrements the amount of fuel available at that location. In *Mprime* there

is an additional operator to transfer fuel between locations. To obtain instances that were within the range of solvability of the planners at hand but that still featured a significant amount of potential parallelism we had to restrict the number of locations to 2. Our random instances (generated with the generators available from the FF homepage) all have 3 trucks (truck transport capacity is limited to a random number between 1 and 3 objects), and scale from 6 to 10 objects to be transported. See the data in Table 5.

Mystery					
planner	size 1	size 2	size 3	size 4	size 5
BBG-BA-L	1.00	1.00	1.00	0.40	0.40
BBG-BM-L	1.00	1.00	1.00	0.40	0.40
BBG-BA-C	1.00	1.00	1.00	0.60	0.50
BBG-BM-C	1.00	1.00	1.00	0.50	0.50
IPP	1.00	1.00	1.00	0.40	0.40
IPP-N	1.00	0.90	1.00	0.40	0.20

Mprime					
planner	size 1	size 2	size 3	size 4	size 5
BBG-BA-L	0.80	1.00	0.60	1.00	0.50
BBG-BM-L	0.80	1.00	0.50	1.00	0.50
BBG-BA-C	1.00	1.00	1.00	1.00	0.90
BBG-BM-C	1.00	1.00	0.90	1.00	0.70
IPP	1.00	1.00	0.80	1.00	0.50
IPP-N	0.70	1.00	0.50	0.90	0.50

Table 5: Fraction of solved instances in the *Mystery* and *Mprime* domains. See explanation in the text.

In these domains the planners seem to perform more evenly, with a small edge for the BBG configurations with criticality flaw selection (-C), which solve more instances than the other planners. The *Mystery* search times compared to IPP are 1.24, 5.55, 9.40, 7.77, 28.09 for BBG-BA-C and 6.09, 16.31, 17.29, 13.78, 140.40 for IPP; the number of states are 824, 4757, 5729, 5141, 14418 for BBG-BA-C and 17159, 63019, 71707, 55638, 468469 for IPP. Like in *Blocksworld*, the reduction in the number of states is quite significant while the reduction in search time is moderate. Something similar occurs in *Mprime*.

Serial Domains

To test the hypothesis that the new branching schemes are suitable in particular for parallel domains where the branching factor of the regression search is very high, we ran experiments in domains with less parallelism. To exclude, as far as possible, effects that arise from structural differences between planning domains, we used serialized versions of (four of) the domains described above. In *Logistics* and *Blocksworld*, we introduced artificial preconditions and effects that force actions to be applied sequentially;⁷ in *Satellite* and *Depots* we generated random suits with less parallelism. The *Logistics* and *Blocksworld* data are displayed in Table 6.

⁷One copy of the original operators require a new fact f ; they delete f , and add f' ; a second copy of the operators require f' , delete f' , and add f . This way all actions interfere with each other while the domain semantics remain the same.

Logistics-serialized					
planner	size 1	size 2	size 3	size 4	size 5
BBG-BA-L	1.00	1.00	1.00	0.80	0.70
BBG-BM-L	1.00	1.00	1.00	0.80	0.50
BBG-BA-C	1.00	1.00	1.00	0.90	0.70
BBG-BM-C	1.00	1.00	1.00	0.80	0.70
IPP	1.00	1.00	1.00	1.00	1.00
IPP-N	1.00	1.00	1.00	0.80	0.70

Blocksworld-serialized					
planner	size 1	size 2	size 3	size 4	size 5
BBG-BA-L	1.00	0.90	0.90	0.40	0.10
BBG-BM-L	1.00	1.00	0.90	0.30	0.10
BBG-BA-C	1.00	0.50	0.60	0.10	0.00
BBG-BM-C	1.00	0.30	0.60	0.00	0.00
IPP	1.00	1.00	1.00	0.90	0.60
IPP-N	1.00	1.00	1.00	0.40	0.10

Table 6: Fraction of solved instances in *Logistics-serialized* and *Blocksworld-serialized*. See explanation in the text.

The serialized *Logistics* instances become harder for all the planners considered (probably due to the low quality of the plan graph lower bound for such a domain). We thus had to consider simpler instances with a single city with two locations and one truck. The number of objects scales from 5 to 9, and as before, we have 10 random instances per size. The best BBG planner is BBG-BA-C, which in this case, is not as good as IPP. The runtimes obtained are 0.01, 0.02, 0.06, 0.18, 0.39 for IPP and 4.15, 0.51, 10.68, 68.18, 18.39 for BBG-BA-C, while the search states are 218, 284, 854, 1957, and 3350 for IPP, and 11305, 890, 19323, 99084, and 19846 for BBG-BA-C. In this case, the new branching schemes yields a search that visits more states and takes more time. The comparison to IPP without memoization (IPP-N) is better for BBG: the runtimes are 1.57, 0.25, 3.39, 7.11, 26.18 for IPP-N and 4.15, 0.51, 10.68, 16.39, 18.29 for BBG-BA-C, while the states visited are 101676, 12467, 147355, 278770, 670861 for IPP-N and 11305, 890, 19323, 25912, 19846 for BBG-BA-C. This suggests that in the serialized form of *Logistics* the new branching schemes reduce the search space, but not as much as memoization does; also, we see that the search space reduction in comparison to plain regression, i.e., IPP-N, is not sufficient to make up for the more costly computation.

In our serialized *Blocksworld* suit, we have instances with 8 to 12 blocks, 10 instances per size. The conclusions that can be drawn are similar to those for *Logistics-serialized*: IPP performs best, and IPP-N performs close to the best BBG planner, BBG-BA-L.

The serial *Satellite* and *Depots* suits differ from the parallel suits in that there are only 1 satellite and 2 hoists respectively. The resulting data are displayed in Table 7.

In the serial *Satellite* suit, memoization is again a more effective enhancement to the basic Graphplan search than the new branching schemes: the runtimes are 0.07, 0.25, 0.36, 1.11 for IPP, and 0.16, 0.90, 10.79, 10.51, 204.00 for the best BBG planner in this case, BBG-BA-L. Likewise, the states visited are 223, 752, 2164, 2397, 6194 for IPP and 264, 2107, 18698, 12875, 183273 for BBG-BA-L (the BBG-

Satellite — serial suit					
planner	size 1	size 2	size 3	size 4	size 5
BBG-BA-L	1.00	1.00	1.00	1.00	1.00
BBG-BM-L	1.00	1.00	1.00	1.00	1.00
BBG-BA-C	1.00	1.00	0.80	1.00	0.00
BBG-BM-C	1.00	1.00	0.00	0.00	0.00
IPP	1.00	1.00	1.00	1.00	1.00
IPP-N	1.00	1.00	0.00	0.00	0.00
Depots — serial suit					
BBG-BA-L	1.00	1.00	1.00	1.00	1.00
BBG-BM-L	1.00	1.00	1.00	1.00	0.90
BBG-BA-C	1.00	1.00	0.70	0.50	0.20
BBG-BM-C	1.00	1.00	0.80	0.50	0.20
IPP	1.00	1.00	1.00	1.00	1.00
IPP-N	1.00	1.00	0.90	0.70	0.30

Table 7: Fraction of solved instances in *Satellite*, (more) serial random suit, and *Depots*, (more) serial random suit. See explanation in the text.

BA-L search space size is still a vast improvement over basic regression search: IPP-N searches 54812 and 9116650 states on average in the smallest two groups of instances). In the serial *Depots* suit, the best BBG planner BBG-BA-L is still better than IPP with memoization, but not as much as in the parallel suit: the runtimes are 0.00, 0.01, 0.47, 5.94, 49.46 for IPP and 0.12, 0.17, 0.87, 2.10, 28.86 for BBG-BA-L; the search state data are 39, 113, 1382, 9768, 59587 for IPP and 50, 50, 418, 488, 7793 for BBG-BA-L. As in the parallel domains, IPP-N is an order-of-magnitude slower than IPP.

We have also collected some data to make sense of the differences in performance that result from the different branching schemes. To make precise the degree of parallelism in a planning task and its effect on the branching factor during Graphplan-style backchaining, we measured the following two parameters: the *normalized potential parallelism* — averaged over all actions a , and standing for the number of actions that a does not interfere with, over the total number of actions; and *IPP branching factor* — averaged over all non-leaf nodes n in IPP’s search space, and standing for the number of subgoal sets produced by successful backchaining from n (including memoized subgoal sets). The resulting data is shown in Table 8.

The data in Table 8 reflect averages over all instances in the respective sets; dashes in the entries for the IPP branching factor, appear thus in cases where IPP did not show all the corresponding instances. Clearly, in *Logistics*, *Blocksworld*, *Satellite*, and *Depots* the parallel suits contain a higher degree of potential parallelism; and except for the *Satellite* domain, where the only parallel instances solved by IPP get solved without any search at all, the IPP branching factor in the parallel suits is higher. This supports our hypothesis about the effect of parallelism on Graphplan’s backward search, and helps to explain the performance results obtained. Making inter-domain comparisons, on the other hand, is more difficult; there is certainly a correlation between the degree of parallelism and the relative behavior of IPP and BBG, yet exact form of the correlation appears

domain	size 1	size 2	size 3	size 4	size 5
average normalized potential parallelism					
<i>Logistics</i>	0.96	0.97	0.97	0.98	0.98
<i>Logistics-ser.</i>	0.00	0.00	0.00	0.00	0.00
<i>Blocksworld</i>	0.42	0.52	0.60	0.65	0.69
<i>Blocksworld-ser.</i>	0.00	0.00	0.00	0.00	0.00
<i>Satellite-par.</i>	0.93	0.94	0.94	0.95	0.95
<i>Satellite-ser.</i>	0.68	0.71	0.73	0.76	0.77
<i>Depots-par.</i>	0.68	0.69	0.70	0.71	0.72
<i>Depots-ser.</i>	0.47	0.47	0.48	0.49	0.49
<i>Mystery</i>	0.74	0.74	0.74	0.75	0.75
<i>Mprime</i>	0.74	0.74	0.75	0.76	0.76
average IPP branching factor					
<i>Logistics</i>	59.53	176.26	167.98	-	133.61
<i>Logistics-ser.</i>	2.52	2.50	3.59	3.79	5.56
<i>Blocksworld</i>	1.18	4.35	6.71	7.91	46.46
<i>Blocksworld-ser.</i>	3.65	4.89	4.28	6.68	5.32
<i>Satellite-par.</i>	1.00	1.17	-	-	-
<i>Satellite-ser.</i>	9.85	12.32	14.39	15.90	17.88
<i>Depots-par.</i>	13.14	24.78	55.07	84.71	54.79
<i>Depots-ser.</i>	2.06	4.57	10.17	13.59	17.44
<i>Mystery</i>	66.08	30.65	74.45	107.47	131.87
<i>Mprime</i>	47.50	48.51	50.68	46.22	67.28

Table 8: Parallelism and regression branching factor data across all domains. See explanation in the text.

to depend on the individual domains (see the discussion below).

Comparison with the State-of-the-art

In order to assess how far the new branching schemes take Graphplan in relation to state-of-the-art parallel planners, we report some results comparing one of the options in BBG, BBG-BA-L (i.e., BBG with branching on actions schemes and latest flaw selection criterion) against Blackbox using Chaff. Considering the success of Chaff in the SAT community, we figure that Blackbox using Chaff is one of the strongest SAT-based optimal parallel planners around. We ran Blackbox on the instances considered in the previous section. The resulting data are displayed in Table 9.

In the *Logistics* (both serial and parallel) as well as the *Mystery* and *Mprime* suits, Blackbox is superior to BBG-BA-L. In the *Satellite* suits the planner performances are similar modulo an average search time factor of 1.45 (in the parallel suit) and 1.69 (in the serial suit) in favor of Blackbox (recall that we always report search times, and in Blackbox, this means the time spent in Chaff). In the *Blocksworld* and *Depots* suits BBG-BA-L is superior to Blackbox in our experiment. Note that the behavior of Blackbox in these latter 4 suits is rather poor. This is because over the larger instances, the planner often ran out of memory. How much of this phenomenon is due to the Graphplan implementation in Blackbox, to the size of the CNFs generated, or to the clause learning mechanism in Chaff, it is hard to say. In those examples that are solved by both planners, BBG-BA-L is superior to Blackbox in search time by an average factor of 7.25 (note that Chaff is a highly optimized implementation

domain	planner	size 1	size 2	size 3	size 4	size 5
<i>Logistics</i>	BBG	1.00	1.00	0.90	1.00	0.70
<i>Logistics</i>	BBOX	1.00	1.00	1.00	1.00	1.00
<i>Blocksworld</i>	BBG	1.00	1.00	1.00	1.00	0.90
<i>Blocksworld</i>	BBOX	1.00	1.00	0.70	0.30	0.00
<i>Satellite-par.</i>	BBG	1.00	1.00	1.00	1.00	1.00
<i>Satellite-par.</i>	BBOX	1.00	1.00	1.00	1.00	1.00
<i>Depots-par.</i>	BBG	1.00	1.00	1.00	1.00	0.70
<i>Depots-par.</i>	BBOX	0.80	0.50	0.40	0.10	0.00
<i>Mystery</i>	BBG	1.00	1.00	1.00	0.40	0.40
<i>Mystery</i>	BBOX	1.00	1.00	1.00	0.70	0.70
<i>Mprime</i>	BBG	0.80	1.00	0.60	1.00	0.50
<i>Mprime</i>	BBOX	1.00	1.00	1.00	1.00	1.00
<i>Logistics-ser.</i>	BBG	1.00	1.00	1.00	0.80	0.70
<i>Logistics-ser.</i>	BBOX	1.00	1.00	1.00	0.90	1.00
<i>Blocksworld-ser.</i>	BBG	1.00	0.90	0.90	0.40	0.10
<i>Blocksworld-ser.</i>	BBOX	0.40	0.00	0.00	0.00	0.00
<i>Satellite-ser.</i>	BBG	1.00	1.00	1.00	1.00	1.00
<i>Satellite-ser.</i>	BBOX	1.00	1.00	1.00	1.00	1.00
<i>Depots-ser.</i>	BBG	1.00	1.00	1.00	1.00	1.00
<i>Depots-ser.</i>	BBOX	1.00	1.00	1.00	0.80	0.20

Table 9: Fraction of solved instances, compared between BBG-BA-L and Blackbox using Chaff. See explanation in the text.

so this factor is due to *search*, not programming, advantages in BBG).

Discussion

We have introduced two non-directional branching schemes in Graphplan and showed that they significantly improve performance over a number of parallel domains. These results have been achieved by a *pure* modification to the branching scheme, leaving the IDA* search algorithm and the plan graph lower bounds in Graphplan untouched. Non-directional branching schemes in the Graphplan setting have been introduced and analyzed in (Rintanen 1998) and (Baioletti, Marcugini, & Milani 2000), while (Kautz & Selman 1999) and (Do & Kambhampati 2000) propose suitable SAT and CSP translations. These proposals, however, not only modify the branching scheme in Graphplan, but also its pruning mechanism, which is formulated as some form of constraint propagation and consistency checking. Our proposal, on the other hand, combines a new branching scheme with the plan graph based pruning mechanism that comes from the original Graphplan. The result is that the differences in performance follow exclusively from the different branching schemes, and not from the propagation rules. Indeed, some of these propagation rules can be added to the resulting planner, leading to further speed ups (e.g. rules that exclude actions at layer $d - 1$ that delete the precondition of an action committed at layer d , etc.).⁸ Interestingly,

⁸A preliminary implementation of two of these rules resulted actually in speed ups by factors ranging between 2 and 5 over the different domains.

even without such improvements, we have shown that the performance of the resulting planner over a wide range of domains, approaches the performance of an state-of-the-art optimal parallel planner such as Blackbox with Chaff. The importance of controlled experimentation in the empirical study of algorithms is discussed at length in (Hooker 1996).

A question raised by this work concerns the conditions that make one branching scheme more suitable than another given the same lower bound technique (such as plan graph lower bounds). In principle, it would seem that the degree of parallelism, and the branching factor in the regression search, are the two key factors. Indeed, the experiments we have presented illustrate that the higher these factors, the more effective the non-directional branching schemes become in relation to the regression search in Graphplan. Yet these results are not uniform over all domains; e.g., domains such as *Mystery* and *Mprime* involve a high degree of parallelism and a high branching factor, and yet, over these domains, the non-directional schemes do not improve much over the directional (regression-based) scheme.

On the algorithmic side, other branching schemes and branching point selection heuristics need to be studied in this setting, as they appear to be crucial for performance.

Appendix: Implementation Details

Our implementation follows the ideas in STAN (Long & Fox 1999). The plan graph is represented as a single layer of facts and actions, where only the layer-dependent information is updated along the time steps. Information that is frequently accessed is kept in bit vectors. This is the case for precondition, add, and delete lists, for the sets of facts which are mutex with an action precondition at a layer, for the sets of actions which are mutex with all achievers for a fact at a layer, and for the sets of actions that can achieve a fact at a layer. Full details about the computations are in (Long & Fox 1999). As an illustration, when $f-M$ are the bits for the actions that are mutex with all achievers of a fact f , and $f'-A$ are the bits for the actions that achieve f' , then f is mutex with f' when there is no bit b such that $f'-A[b] = 1$ and $f-M[b] = 0$.

When a new constraint is introduced at a layer d , the graph is updated by removing a set of actions at layer d , and propagating the changes. In general, we have a layer i , a set *ops* of actions to be removed at i , and a set *opmutexes* of action mutexes to be introduced at i . These changes are propagated to the next fact layer as follows. First the bits corresponding to *ops* are set to 0 in the bit vectors $f-A$ that represent the achievers of the respective facts f , and the facts whose bit vectors become empty are removed — of course, in this and the following removal operations, the facts and actions are not physically removed from the graph but only marked as deleted. Simultaneously, we build a list *chfts* of facts whose achievers have changed. Then all facts that are achieved by an action in *opmutexes* are included into *chfts*. For each fact f in *chfts*, we thereafter recompute $f-M$, i.e., the bit vector representing the actions that are mutex with all achievers of f . Finally for all pairs f and f' of facts such that at least one of them is in *chfts* it is checked (by comparing the appropriate bit vectors, $f-M$ and $f'-A$)

whether they are now mutex; if so, the respective bit vectors are updated. All propagated changes are kept in memory, and retracted upon backtracking. The result of the propagation is a set of facts *fts* to be removed at $i + 1$ (i.e., the next fact layer), and a set *ftmutexes* of fact mutexes to be introduced at $i + 1$. Propagating these changes to action layer $i + 1$ is simple. First remove all actions that have a fact in *fts* as precondition. Then proceed over all pairs o and o' of actions such that a pair $f \in pre(o)$ and $f' \in pre(o')$ is in *ftmutexes*: if $o = o'$ then remove o ; else, make o and o' mutex. Again, the changes are remembered for later retraction, and the output is a list *ops* of actions to be removed at $i + 1$, and a set *opmutexes* of action mutexes to be introduced at $i + 1$. From here, we continue as described above until the current deadline layer is reached.

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