

# Learning cluster-based structure to solve constraint satisfaction problems

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Published online: 7 September 2010  
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**Abstract** The hybrid search algorithm for constraint satisfaction problems described here first uses local search to detect crucial substructures and then applies that knowledge to solve the problem. This paper shows the difficulties encountered by traditional and state-of-the-art learning heuristics when these substructures are overlooked. It introduces a new algorithm, *Foretell*, to detect dense and tight substructures called *clusters* with local search. It also develops two ways to use clusters during global search: one supports variable-ordering heuristics and the other makes inferences adapted to them. Together they improve performance on both benchmark and real-world problems.

**Keywords** Cluster · Structure learning · Hybrid search · Constraint satisfaction problem

**Mathematics Subject Classification (2010)** 68T20

## 1 Introduction

A problem solver that begins with a general algorithm for search may well be surprised, or even defeated, by difficult subproblems that lurk within the larger one.

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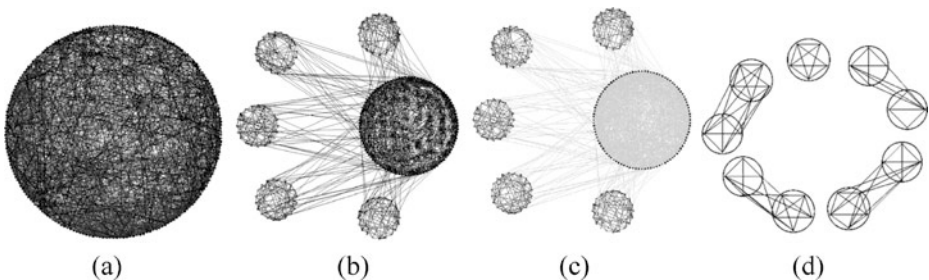
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Indeed, many modern algorithms are now designed to learn about such subproblems *during* search, as they encounter them. The thesis of this paper, however, is that it is possible, and better, to identify such subproblems *before* search, and then exploit them both to solve and to understand the problem at hand. The principle result of this paper is that it is feasible to detect difficult subproblems within a challenging constraint satisfaction problem (CSP) and then exploit a structure composed of them, both to solve the CSP effectively and to explain it to a user. On certain benchmark problems, this approach achieves as much as an order of magnitude speedup. Moreover, the learned structure provides a meaningful high-level formulation of the problem.

Search-ordering heuristics can be misled if they overlook the inherent structure of a problem. A binary CSP, for example, has a set of variables, each with a domain of values, and a set of *constraints*, each of which restricts how some pair of variables can be bound simultaneously. Such a problem is often represented as a graph where each variable is a node and each constraint is an edge between the pair of variables it restricts. When search for a CSP's solution assigns values to variables one at a time, the order in which the variables are addressed is crucial to search performance. Such search is typically supported by variable-ordering heuristics that prefer variables of high degree in the associated graph. Difficult subproblems, however, are not necessarily characterized by such variables.

What is really needed is an effective reasoning mechanism that predicts and exploits difficult subproblems. The traditional graph in Fig. 1a, for example, plots its 200 variables on a circle, and offers little insight or guidance to a solver. Figure 1b redraws Fig. 1a by extracting some variables from the central circle, and Fig. 1c darkens the more restrictive constraints. This formulation is clear only to the problem generator, however. Without the knowledge in Fig. 1c, the problem could not be solved in 30 min by a traditional heuristic. Two learning heuristics (described below) solved it in about 127 and 88 s. This paper develops algorithms that detected the graph in Fig. 1d and used it to solve the same problem, all in 3.56 s.

Although many challenging real-world problems can be modeled as CSPs, their graphs often have non-random structure that makes them difficult to solve for heuristics that ignore it. Here, a *cluster* in a CSP is a subproblem that is particularly difficult



**Fig. 1** A cluster graph predicts subproblems crucial to search. **a** An uninformative graph for a CSP places the variables on the circumference of a circle. **b** The hidden structure of the same problem, given perfect knowledge. **c** Darker edges represent tighter constraints in the same problem. **d** A cluster graph displays critical portions of the problem. This graph was detected from (a) by local search

to solve, and a *cluster graph* is a structural model that illustrates the relationships among all a CSP's discovered clusters. The next section provides background and formal definitions for constraint satisfaction problems and search for solutions to them. Section 3 describes related work on structure, while Section 4 investigates how perfect structural foreknowledge like Fig. 1c might best be used during search. Section 5 describes *Foretell*, a local search algorithm that detects clusters; Section 6 provides cluster graphs for several interesting CSPs. Section 7 explores several variable-ordering heuristics that are based on structural knowledge learned by *Foretell*. Section 8 investigates how such structural knowledge can be used in constraint inference; Section 9 provides experimental results for these algorithms on benchmark and read-world problems. The final section discusses the results and future work.

## 2 Background

Constraint satisfaction searches for a solution to a problem that satisfies a set of restrictions. Many combinatoric problems, such as scheduling, can be represented as CSPs. Because CSP solution is NP-complete [26], no general-purpose polynomial time algorithm is likely to exist. Instead, much work has been directed toward the design of algorithms with shorter average run-times. This section introduces basic CSP terminology and some relevant search methods.

### 2.1 Constraint satisfaction problems

A constraint satisfaction problem is represented by a triple  $\langle X, D, C \rangle$ , where:

- $X$  is a set of variables,  $X = \{X_1, \dots, X_n\}$
- The *domain* of variable  $X_i$  is a finite and discrete set of its possible values  $D_i \in D$
- $C$  is a set of constraints that restrict the values the variables in it *scope*  $S_i \subseteq X$  can hold simultaneously

A *constraint*  $C_i$  uses a relation  $R_i$  to indicate which tuples in the Cartesian product of the domains of the variables in  $S_i$  are acceptable. If  $C_i$  is *extensional*,  $R_i$  enumerates the tuples; if  $C_i$  is *intensional*,  $R_i$  is a boolean predicate over all the tuples. This paper deals primarily with *extensional CSPs*, all of whose constraints are extensional. (Some empirical results on *intensional CSPs*, all of whose constraints are intensional, appear in Section 9.) A *subproblem*  $\langle V, D', C' \rangle$  of a CSP  $\langle X, D, C \rangle$  is a CSP induced by a subset of variables  $V \subseteq X$ , where  $D' = \{D_i \in D \mid X_i \in V\}$  and  $C' = \{C_i \mid S_i \text{ is the scope of } C_i \text{ and } S_i \subseteq V\}$ .

The *arity* of a constraint is the *cardinality*, or size, of its scope. A *unary* constraint is defined on a single variable; a *binary* constraint, on two variables. Let binary constraint  $C_{ij}$  have scope  $(X_i, X_j)$ , where  $X_i$ 's domain is  $D_i$  and  $X_j$ 's domain is  $D_j$ . If  $C_{ij}$  excludes  $r$  value pairs from  $X_i \times X_j$ , then its *tightness*  $t(C_{ij})$  is defined as the percentage of possible value pairs  $C_{ij}$  excludes:

$$t(C_{ij}) = \frac{r}{|D_i| \times |D_j|} \quad \text{where } r \leq |D_i| \times |D_j| \quad (1)$$

Although for intensional  $C_i$  (1) could require a test on every possible tuple, some predicates (e.g., disjunctive inequalities over integer intervals) permit direct calculation of  $r$ .

This paper focuses on *binary CSPs*, which have only unary and binary constraints. A binary CSP can be represented as a *constraint graph*, where each variable is a node, each binary constraint is an edge, nodes are labeled by their domains, and each edge is labeled by the legal value pairs for that constraint. Two variables whose nodes are joined by an edge in the constraint graph are *neighbors*, and the *static degree* of a variable is the number of its neighbors. An *instantiation* of size  $k$  is a value assignment of a subset of  $k$  variables in  $X$ . If  $k < n$ , it is a *partial instantiation*; if  $k = n$ , it is a *full instantiation*. A *solution* to a CSP is a full instantiation that satisfies every constraint. A *solvable CSP* has at least one solution; an *unsolvable CSP* has no solution.

A *CSP problem class* is a set of CSPs that are categorized as similar, such as all CSPs that share the same parameters  $\langle n, k, d, t \rangle$ , where  $n$  denotes the problem's number of variables,  $k$  its maximum domain size,  $d$  its density, and  $t$  the average of the tightness of its individual constraints. The *density*  $d(P)$  of a CSP  $P$  is defined as the fraction of the  $\frac{n(n-1)}{2}$  possible pairs of variables that could be restricted:

$$d(P) = \frac{2|C|}{n(n-1)} \tag{2}$$

A class of structured problems can also be parameterized. For example, a class of artificially generated *composed* CSPs can be written as

$$\langle n, k, d, t \rangle s \langle n', k', d', t' \rangle d'' t'' \tag{3}$$

Each problem in (3) partitions its variables into subproblems, one designated as the *central component* and the others as *satellites*. There are some constraints (*links*) between satellite variables and variables in the central component, but none between variables in distinct satellites. The central component is in  $\langle n, k, d, t \rangle$ . There are  $s$  satellites (each in  $\langle n', k', d', t' \rangle$ ), and links with density  $d''$  and tightness  $t''$ . Let there be  $l$  links between the central component and the  $s$  satellites. Then the link density  $d''$  is the fraction of the  $nsn'$  possible link edges present:

$$d'' = \frac{l}{nsn'} \quad \text{where } l \leq nsn', \tag{4}$$

and the link tightness  $t''$  is the average tightness of all its links. Figure 1a is an unsolvable problem in

$$Comp = \langle 100, 10, 0.15, 0.05 \rangle 5 \langle 20, 10, 0.25, 0.50 \rangle 0.12, 0.05$$

which has one tightness uniformly within its central component and along its links, but another uniform tightness within its satellites. Figure 1c redraws it with the central component along the large circle and each satellite on a separate circle. *Structured* CSPs, including *Comp*, have characteristics that can be exploited by a specialized method to outperform a more general one. They also offer an opportunity to explore the impact and management of difficult subproblems.

## 2.2 Search for a solution to a CSP

*Search* for a solution to a CSP moves through the space of its possible instantiations. Global search traverses the space of partial and full instantiations, while local search is restricted to the space of full instantiations.

### 2.2.1 Global search

*Global search* begins with an *empty instantiation*, where no variable is assigned a value. Global search traverses the search space systematically, assigning a value to one unbound variable at a time. Global search is complete because it is always able to find every solution. Thus, failure by global search to find a solution proves that the problem is unsolvable. A problem is labeled “solved” in this paper if global search finds a solution or proves that none exists.

During global search, from the perspective of a given node, an instantiated variable is called a *past variable* and an unbound variable is called a *future variable*. The *dynamic degree* of a variable is the number of its neighbors that are future variables. An instantiation for a future variable is *consistent* if it does not violate any constraint with a past variable. An inconsistent instantiation violates one or more constraints. Because global search traverses a search space systematically to find a solution, pruning the search space could accelerate this process. *Inference* attempts such pruning; it propagates the effect of an instantiation on the domains of the future variables. If a domain becomes empty (a *wipeout*), search *backtracks*, that is, it retracts the instantiation of the current variable and restores the domains that had been reduced by that instantiation. This paper uses *chronological backtracking*, which revisits each value assignment in order of recency. Figure 2 provides pseudocode for global search.

Two well-known inference methods are forward checking and arc consistency. Immediately after the instantiation of a variable  $x$ , *forward checking* removes all inconsistent values from the domains of all future variables that are neighbors of  $x$  [19]. Arc consistency is a potentially more powerful inference method. Along a binary constraint  $C_{ij}$  between variables  $X_i$  and  $X_j$  in a CSP, value  $a \in D_i$  is *supported* by  $b \in D_j$  if and only if  $x_i = a$  and  $x_j = b$  together satisfy  $C_{ij}$ .  $C_{ij}$  is *arc consistent* if and only if every value in  $D_i$  has some support in  $D_j$  and every value in  $D_j$  has some support in  $D_i$ . If every constraint in a CSP is arc consistent, then the CSP is arc consistent. *MAC-3* is an inference method that maintains a problem’s arc consistency during search. Immediately after the instantiation of variable  $x$ , *MAC-3* enqueues the edges from  $x$  to all its future variable neighbors, and then checks each element

**Algorithm 1:** Pseudocode for CSP global search

Until all variables have values that satisfy all constraints OR  
 some variable at the root has an empty domain  
**Select** a variable  
**Assign** a value to this variable  
**Infer** the impact of this assignment ; *\*inference\**  
 If a wipeout occurs, **backtrack**

**Fig. 2** Pseudocode for global search

of the queue for domain reduction [34]. Whenever this process reduces the domain of any future variable  $z$ , MAC-3 enqueues every constraint between  $z$  and its future variable neighbors, and iterates until its queue is empty.

Proper choice of the next variable to instantiate can significantly improve search performance. *Variable-ordering* heuristics may provide crucial advice for global search. A variable-ordering heuristic usually follows the fail-first principle which states “To succeed, try first where you are most likely to fail” [19]. *MinDom* seeks to minimize search tree size by reducing the branch factor; it prefers variables with small *dynamic domains*, the values remaining after inference. *MaxDeg* focuses on variables with many constraints; it prefers variables with high dynamic degree. *MinDomDeg* combines the two: it prefers variables that minimize their ratio of dynamic domain size to dynamic degree. *MinDomDeg* is an effective off-the-shelf variable-ordering heuristic, but can be outperformed by other heuristics that learn during search.

Weighted degree is a conflict-directed variable-ordering heuristic [2]. It associates each constraint with a weight, initialized to 1. During search, whenever a wipeout is encountered, the weight of the constraint that causes this wipeout is incremented by 1. The weighted degree of a variable  $X_i$  is the sum of the weights of all constraints between  $X_i$  and its unbound neighbors. *MaxWdeg* is a variable-ordering heuristic that selects a variable with the highest weighted degree. *MaxWdeg* is adaptive; it gradually guides search toward constraints that cause wipeouts. Another adaptive heuristic, *MinDomWdeg*, minimizes the ratio of dynamic domain size to weighted degree.

### 2.2.2 Local search

*Local search* begins with a full instantiation. It moves from one full instantiation to another until some full instantiation is a solution. The *distance* between two full instantiations is the number of different variable assignments. The *k-neighborhood* of a full instantiation  $s$  includes  $s$  and all the full instantiations within distance  $k$  of  $s$ . Any full instantiation in  $s$ 's neighborhood is  $s$ 's *neighbor*. In local search, each move is determined by a decision based on only *local knowledge*, the information inherent in the neighborhood of the current full instantiation being investigated. Figure 3 provides pseudocode for local search.

Within a neighborhood, local search uses an evaluation function to determine which neighbor is the most improved full instantiation. This function maps full instantiations onto the real numbers, and maps solutions onto a global maximum. Given such an evaluation function, local search chooses a neighbor whose value is a maximum within the neighborhood and is larger than that of the current full

#### Algorithm 2: Pseudocode for CSP local search

```

current-solution ← Generate-Initial-Full-Instantiation( )
Until current-solution violates no constraint
  best-neighbor ← Find-best-neighbor(current-solution)
  if best-neighbor is better than current-solution
    then current-solution ← best-neighbor
  else Escape-local-optimum( )

```

**Fig. 3** Pseudocode for local search

instantiation. Given an evaluation function  $f$ , a search space  $S$  and a neighborhood relation  $N \subseteq S \times S$ , a full instantiation  $s$  is a *local optimum* if and only if for all  $s' \in N(s)$ ,  $f(s') \leq f(s)$  and  $s$  is not a solution.

Local search normally requires less space than systematic search because it remembers few visited states. It often finds CSP solutions much faster than systematic search [30] and may be more effective on some real-time problems (e.g., game playing). Local search can become trapped at a local optimum, however, and some points in the search space may be inaccessible from the start state. One way to escape from local optima is to change neighborhoods during search.

### 2.2.3 Variable neighborhood search

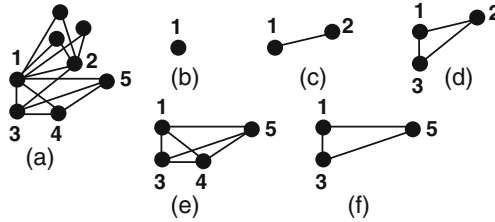
The local optima of one neighborhood are not necessarily also local optima in another. The larger the neighborhood a full instantiation checks, the better the chance to improve the current full instantiation and the less likely it is to get trapped in local optima. Variable Neighborhood Search (VNS) uses multiple neighborhoods of increasing size [18]. Figure 4 provides pseudocode for VNS, a non-deterministic search through  $k$  neighborhoods.

VNS succeeds on a wide range of combinatorial and optimization problems [17]. It works outward from an initial subset of variables (Fig. 4, line 1) in a relatively small neighborhood in a graph through  $k$  pre-specified, increasingly large neighborhoods (lines 2–3). Each neighborhood restricts the current options; as VNS iterates, each new neighborhood provides a larger search space. Within a neighborhood, Variable Neighborhood Descent (VND) is a local search algorithm that tries to improve the current subset (*best-yet*) according to a metric, *score*. A better local optimum resets *best-yet* and returns to the first neighborhood (lines 6–9); otherwise search proceeds to the next neighborhood (lines 10–11). *Shaking* (line 5) shifts search within the current neighborhood and randomizes the current *best-yet* to explore different portions of the search space. As *index* increases, the neighborhoods become larger, so that the shaken version of *best-yet* becomes less similar to *best-yet* itself. The user-specified stopping condition (line 4) is either elapsed time or movement through some number of increasingly large neighborhoods without improvement. The initial subset, the score metric, and the local search routine vary with the application.

VND greedily extends *best-yet* from *neighborhood*. Once its greedy steps are exhausted, VND repeatedly interchanges one element of its current subset for two

Algorithm 3: VNS on $k$ neighborhoods	
1	$best\text{-}yet \leftarrow \text{initial-subset}$
2	$index \leftarrow 1$
3	$neighborhood \leftarrow neighborhood(index)$
4	<b>until</b> <i>stopping condition</i> or $index = k$
5	<b>unless</b> $index = 1$ , $best\text{-}yet \leftarrow \text{shake}(best\text{-}yet, index)$
6	$local\text{-}optimum \leftarrow \text{local-search}(best\text{-}yet, neighborhood)$ ; *VND*
7	<b>if</b> $\text{score}(local\text{-}optimum) > \text{score}(best\text{-}yet)$
8	<b>then</b> $best\text{-}yet \leftarrow local\text{-}optimum$
9	$index \leftarrow 1$
10	<b>else</b> $index \leftarrow index + 1$
11	$neighborhood \leftarrow neighborhood(index)$

**Fig. 4** Pseudocode for VNS



**Fig. 5** Selected VND steps to find a maximum clique in graph (a). **b** A starting vertex. **c**, **d** Greedy steps add vertices adjacent to every selected vertex, one at a time. **e** A swap replaces vertex 2 with vertices 4 and 5. **f** VNS for  $index = 1$  shakes out one randomly selected vertex. See the text for further details

elements in *neighborhood* and breaks ties greedily. In the search for a maximum clique, for example, VND swaps out some variable  $v$  in *best-yet* for two adjacent variables that are not neighbors of  $v$  and were not in *best-yet*, but are neighbors of all the other variables already in *best-yet*, and breaks ties with maximum variable degree. An alternative produced by VND replaces *best-yet* only if it outscores it.

Figure 5 is a sample of steps that might occur during a call to VND during search for a maximum clique in a simple graph. The initial subset is a vertex that is a neighbor of every vertex in the graph, and the local search metric is subset size. VND adds one vertex adjacent to every vertex in the growing subgraph. (This is the greedy step; ties are broken on maximum degree in the original graph.) When greedy steps are no longer possible, local search swaps out one vertex for a pair of adjacent vertices that are also adjacent to every other vertex in the subgraph, as in Fig. 5e. Eventually neither greedy steps nor swaps can be found. Then the subgraph is returned to VNS, scored, stored if it is the best so far, and then shaken before local search resumes.

Local search is incomplete because it does not search systematically. Thus, it cannot prove a CSP is unsolvable, as real-world problems often are. The work reported here first exploits local search to consider the  $O(2^n)$  set of possible subproblems in a CSP, and then guides global search with the outcome of local search to solve the problem.

### 3 Related work

The heavily constrained, highly interactive subproblems our algorithms identify and exploit in a CSP are called *clusters*. “Cluster” has been used elsewhere to describe aggregations of data, portions of a solution space [22, 29], or relatively isolated, dense areas in a graph [38]. Other work on clusters as subproblems assumed an acyclic metastructure and addressed two classes of artificial problems, half the size of those studied here, and offered no structural description or explanation [31].

A cluster graph identifies dense, tight subproblems before search begins. A cluster graph groups together selected variables and their edges. Variables and constraints not explicit in a cluster graph have nonetheless influenced its formation (via pressure, described in the next section). Thus a cluster graph captures a kind



of *fail-first* metastructure that anticipates and confronts difficulties. This approach differs, therefore, from methods that relax, remove, or soften constraints. The fail first principle underlies many traditional variable-ordering heuristics intended to speed global search [1, 14, 37].

With respect to a given search algorithm, the *backdoor* of a CSP is a set of variables that, once assigned values, make the remainder of search trivial [33]. A backdoor is typically less than 30% of the variables, but its identification before search is NP-complete. Recent work suggested that both static and dynamic properties should be considered during search for a backdoor [9]. The formation of a cluster graph prior to search considers both static (initial) shape and potential (dynamic) changes in domain size. A cluster graph would, ideally, contain the backdoor, but no claim is made here that it does so. Unlike [20, 21], cluster-based explanations are available whether or not a problem has a solution.

An abstraction can be used to simplify a problem temporarily (e.g., [35]). Its solution is then gradually revised to accept additional problem detail, until the revision solves the original problem. Because a cluster graph is applied with the traditional graph, rather than as a replacement for it, no re-resolution is necessary.

Variable-ordering heuristics respond to structure detected in the constraint graph. A CSP whose graph is an arc-consistent tree can be solved without backtracking [12, 13]. SAT problems generated with unsatisfiable large cyclic cores have stumped many proficient SAT solvers [20]. To reduce a cyclic CSP to a tree, a solver could first identify and then address some heuristic approximation of the (NP-hard) minimal cycle cutset [6]. Cycle-ridden problems like those addressed here, however, have cycle cutsets far too large to provide effective guidance. Most related work on elaborate structural features that might facilitate search (including trees [27], acyclic graphs [7], tree decomposition [5, 8, 36], hinges [16], and other complex structures [15, 39, 40]) ignores the tightness of individual constraints and is primarily theoretical, or it incurs considerable computational overhead unjustifiable on easy problems.

#### 4 On the exploitation of structural foreknowledge

This section explores the power of foreknowledge about difficult subproblems to guide search. The approaches it tests are not ultimately allowable as variable-ordering heuristics. Rather they gauge how well knowledge about structure supports search, and how best to use that knowledge. Results appear in Table 1. Note that all experiments reported in this paper were run in *ACE* (the Adaptive Constraint Engine), a test-bed for CSP solution [10]. Because *ACE* is a research tool that gathers extensive data, it is highly informative but not honed for speed. Performance is therefore reported here both as elapsed CPU time in seconds and as number of nodes in the search tree. All cited differences are statistically significant at the 95% confidence level under a one-tailed *t*-test.

Standard variable ordering heuristics did poorly on *Comp* problems. Because variables in the central component have much higher degrees, *MinDomDeg* was immediately drawn to the central component and solved only two of 50 problems within the time limit. Because links in *Comp* are so few and loose, wipeouts began fairly deep in the search tree, after at least 36 variables had been bound. Retraction only led *MinDomDeg* to repair its partial instantiation of the central component,

**Table 1** On 50 *Comp* problems, mean and standard deviation for nodes and CPU seconds, including time to find clusters. Search heuristics appear above the line. Search methods with perfect knowledge (below the line) are not legitimate heuristics because they apply structural foreknowledge available only to the problem generator, not the search engine. *Until-11* is therefore only a target

Heuristic	Time, $\mu$ ( $\sigma$ )		Nodes, $\mu$ ( $\sigma$ )	
<i>MinDomDeg</i>	1,728.157	(355.877)	285,751.970	(61,368.701)
<i>MaxWdeg</i>	123.000	(128.580)	20,817.640	(22,954.165)
<i>MinDomWdeg</i>	83.580	(38.964)	12,519.360	(5,811.370)
<i>Satellite</i>	No problem solved			
<i>Stay</i>	2.848	(3.584)	511.922	(416.345)
<i>Until-11</i>	1.612	(1.866)	398.776	(244.112)

while the true difficulties lay elsewhere, in the satellites. Since weighted degrees are equal to actual degrees at the beginning of search, both *MaxWdeg* and *MinDomWdeg* initially suffered from the same attraction to the central component. After enough experience within the satellites, however, they eventually recovered and solved all the problems. Although *MinDomDeg* cannot solve the problem in Fig. 1 within 30 min, the two learning heuristics eventually recover, with the solution time reported in Section 1. Learning lacks the foresight clusters are intended to provide.

Now consider how heuristics might exploit foreknowledge about the problem. Assume one was given the structure shown in Fig. 1c, and believed that the satellites contained the backdoor. In that case, preference for satellite variables should speed search. Rather than discard traditional variable-ordering heuristics, however, each approach investigated here makes satellites a priority and then breaks ties with *MinDomDeg*. Each approach was given 30 min to solve each problem.

The next experiments seek to exploit perfect structural foreknowledge. The variable-ordering heuristic *satellite* examines whether mere presence in a satellite is sufficient to warrant prioritization. On a *Comp* problem, this approach binds all 100 satellite variables first, in a random order, and then uses *MinDomDeg* on the central component. On a single run, *satellite* never solved any problem within 30 min. (Given its lack of promise, this is the only randomized heuristic that was tested only once. All other non-deterministic experiments here report on an average of 10 runs.)

The variable-ordering heuristic *stay* addresses entire satellites first, one at a time in a random order, before it selects any variable from the central component. *Stay* selects a satellite at random, binds all its variables, and then proceeds to another randomly chosen satellite. Within a satellite and within the central component, *stay* breaks ties with *MinDomDeg*. Guided by the satellites, *stay* with *MinDomDeg* yields dramatically improved results over the traditional heuristics; it averages less than 3 s per problem with a 96% smaller search tree than *MinDomWdeg*.

Given that noteworthy improvement, and the fact that the satellites may only estimate the backdoor of a *Comp* problem, the next approach binds only some of the variables in each satellite. If MAC-3 is in use, for example, there would appear to be little point in “finishing” a satellite once it is reduced to only a pair of variables (with at most a single edge between them); *until-2* selects a different satellite at that point. The generalization of this approach, *until-i*, instantiates variables within a randomly chosen satellite until all but  $i$  variables are bound, and then moves on to another randomly chosen satellite. (*Stay* is equivalent to *until-0*.) Within a selected satellite

and later, within the central component and any “leftover” satellite variables, *until-i* also uses *MinDomDeg*. We tested a range of values:  $i = 2, 3, \dots, 15$ .

Surprisingly, search need not stay long in a given satellite. For *Comp*, where the satellites are of size 20, the clear winner was *until-11*, that is, search can address as few as 45% of the variables in a satellite before safely moving on to the next one. In contrast, the variable-ordering heuristic *satellite-i*, which randomly chooses satellite variables that are not among the last  $i$  future variables in their satellite, performed poorly. (Data omitted.) As  $i$  increases, *satellite-i* behaves more like *MinDomDeg* alone does. (Section 10.2 considers why  $i = 11$  was so successful.)

Clearly, structural foreknowledge is critical to search performance: known satellites speed the solution of *Comp* problems when search addresses them one at a time. The next section describes how knowledge about such dense, tight substructures can be detected automatically, prior to search.

### 5 Cluster detection with *Foretell*

Intuitively, *Foretell*, the cluster finder described here, assembles sets of tightly related variables whose domains are likely to reduce during search. *Foretell* was inspired by VNS’ state-of-the-art speed and accuracy on the DIMACS maximum clique problems [18]. A *clique* is a maximally dense graph, that is, one with all possible edges between its variables. Intuitively, a *near clique* is a clique with a few missing edges. (A more formal definition appears in Section 8.) *Foretell* searches for subproblems that are tight near cliques, where the *tightness* of a subproblem  $P' = \langle V, D', C' \rangle$  is the ratio of the product of the subproblem’s size and density to the sum of the relevant domain products of its constraints;

$$tightness(P') = \frac{|V|d(P')}{\sum_{C_{ij} \in C'} (|D_i| |D_j|)} \tag{5}$$

Note, for example, the missing edges in the clusters of Fig. 1d.

*Foretell* adapts VNS to detect substructures that are clusters. It relies on the *pressure* on a variable  $v$ , the probability that, given all the constraints upon it, when one of  $v$ ’s neighbors is assigned a value, at least one value will be excluded from  $v$ ’s domain. *Foretell* uses pressure (instead of degree, as maximum clique search did in Section 2.2.3) to greedily select variables. Precise calculation of the series that defines pressure is computationally expensive. Instead, we devised an algorithm to approximate quickly the first term in that series, corrected to avoid bias in favor of variables with high degrees or large domains. For variable  $V_i$  with domain size  $|D_i|$  neighbors  $N_i$  and constraint with tightness  $t_{ik}$  between  $V_i$  and  $V_k \in N_i$ , the approximate pressure on  $V_i$ , given the constraints on it, is

$$p(V_i) = \frac{1}{\text{degree}(V_i)} \sum_{V_k \in N_i} \frac{\left( \frac{(|D_i| - 1) \cdot |D_k|}{(1 - t_{ik}) |D_i| \cdot |D_k|} \right)}{\left( \frac{|D_i| \cdot |D_k|}{(1 - t_{ik}) |D_i| \cdot |D_k|} \right)} \tag{6}$$

A cluster’s *score* is the ratio of the product of its number of variables and density to its average edge tightness. No cluster is returned unless it has at least 3 variables.

To find multiple clusters in a problem, *Foretell* finds a first cluster, removes those variables and all constraints that include them in their scopes, and then iterates to find the next cluster among the remaining variables and their constraints. Ties unbroken by maximum pressure are broken by maximum degree, and then, if need be, at random. Clusters are typically (but not always) detected in decreasing size order. Because this is local search, some variation is expected from one pass to the next. The maximum neighborhood index was taken from the original work on maximum cliques: the minimum of 10 and the current cluster size [18].

## 6 Cluster structure detected in CSPs

To automatically generate a cluster graph, *Foretell* identifies clusters in a CSP and then adds any constraints whose scope is fully contained among the cluster variables. Applied to a broad range of benchmark problems taken from [23], this approach produces cluster graphs that make explicit a variety of structure among the clusters themselves. Figure 6 was produced with DrawCSP [24], a visualization tool that displays the original constraint graph or identified cluster graph of binary CSPs.

The first category of structure is a set of isolated clusters, or pairs of clusters, in the cluster graphs for composed problems. The cluster graph in Fig. 6a for a *Comp* problem is suggestive of its tightest edges. Because satellites, with density 0.25, are far from cliques, quite often more than one cluster lies in the same satellite. Thus some clusters are linked. Figure 6b shows a similar structure for another composed CSP, designated 25–10–20 in [23] or, in our notation,

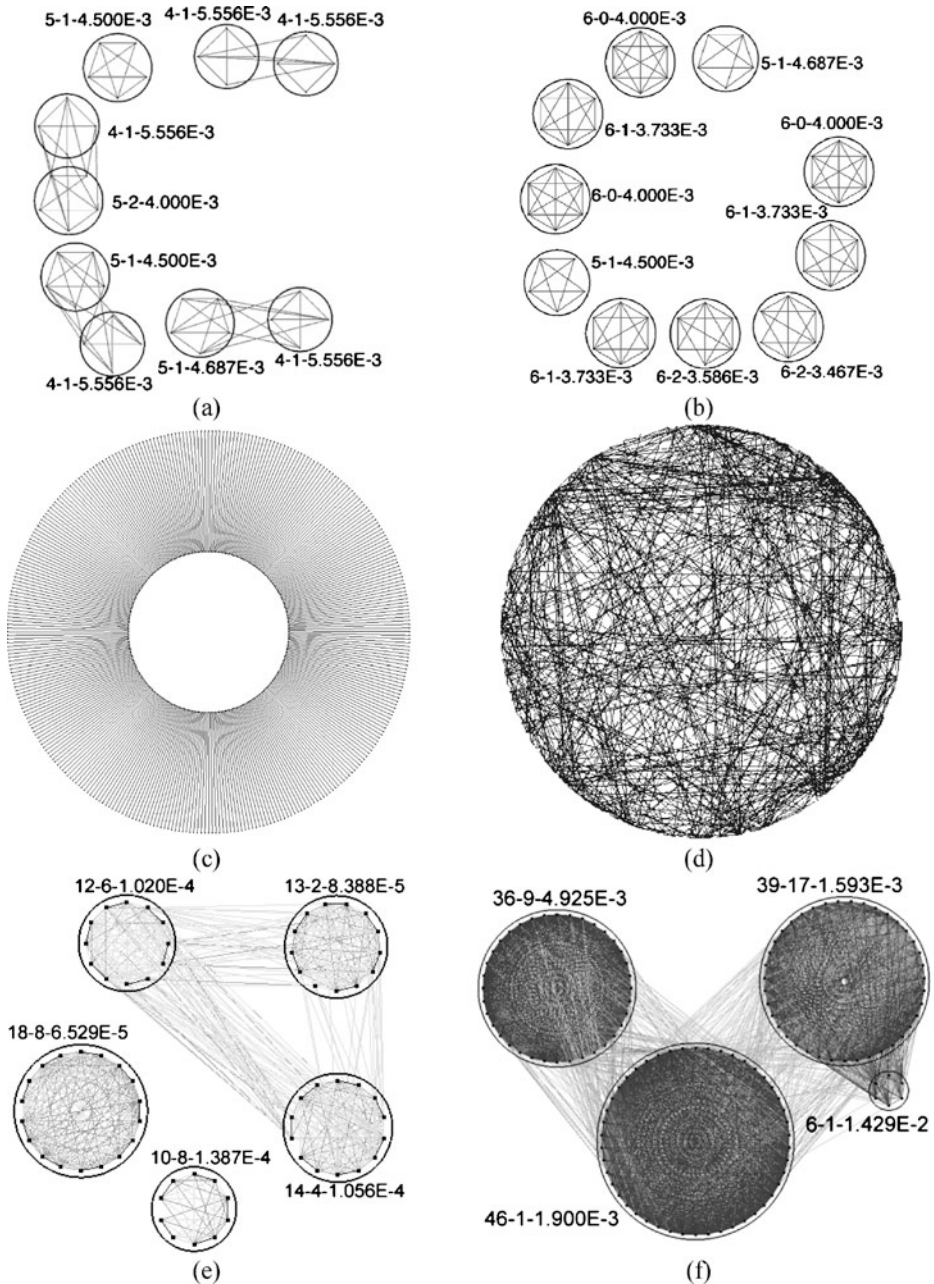
$$\langle 25, 10, 0.667, 0.15 \rangle 10 \langle 8, 10, 0.786, 0.5 \rangle 0.01, 0.05$$

The larger satellite density (0.786) in 25–10–20 encourages the formation of somewhat larger clusters, but typically leaves too few edges to form two clusters in one satellite. Thus its clusters are isolated from one another.

Clusters are not always composed from only the tightest edges. RLFAP here is scene 11 of the radio link frequency problems [3]. Its many constraints vary dramatically in tightness. Figure 6c shows only the tightest constraints, with the 680 variables on two concentric circles. This is a bipartite graph with 340 constraints each of which has tightness greater than 0.9. The cluster graph in Fig. 6e is considerably more informative. *Foretell* found the same 4 clusters, of size 18, 13, 12 and 10, on every run (Not every run detected the cluster of size 14). These 53 variables are the crux of the RLFAP scene 11 problem, the part that makes its rapid solution possible. Only 26, out of 340, of the tightest edges appear in the cluster graph at all. We also tested driverlog problems from a planning competition [23]. The one in Fig. 6d displays a more closely coupled cluster graph in Fig. 6f. Other driverlog problems also display a path-like structure in their cluster graphs.

## 7 Exploitation of structural knowledge to guide search

This section seeks to exploit clusters detected automatically by *Foretell*, much the way foreknown satellites were used in Section 4. *Foretell* never found a cluster larger



**Fig. 6** CSPs display different structures. Each cluster is drawn within a circle; tighter edges are darker. Cluster labels are number of variables, number of additional edges needed to make it a clique, and *Foretell* score. **a** Cluster graph for a *Comp* problem. **b** Cluster graph for a composed 25-10-20 problem. **c** Edges with tightness > 0.9 in RLFAP scene 11 and **d** in the driverlogw-08c problem. **e** Cluster graph for RLFAP scene 11 and **f** for the driverlogw-08c problem

than 6 variables in a *Comp* problem; instead it found multiple (disjoint) clusters in individual satellites, clusters that covered satellites only partially. The primary question then becomes how best to exploit clusters. Is it, for example, better to shift from one cluster to another during search, or to solve them one at a time? And if one at a time, in what order should the clusters be considered? Perhaps one would address the cluster that at the moment is the tightest. The true *dynamic tightness of a cluster*  $c$  on variables  $V_c$  is the ratio of the number of tuples that satisfy its unbound variables under the current partial instantiation to the product of their dynamic domain sizes:

$$\text{dynamic-tightness}(c) = \frac{|\text{satisfying-assignments}(c)|}{\prod_{v \in V_c} \text{original-domain}(v)} \quad (7)$$

(7) is too expensive to calculate repeatedly, as is a dynamic version of pressure in (6). Instead, the dynamic tightness of a cluster  $c$  is estimated here as the ratio of the product of the current domain sizes of those variables to the product of their original domain sizes:

$$\text{estimated-tightness}(c) = \frac{\prod_{v \in V_c} |\text{dynamic-domain}(v)|}{\prod_{v \in V_c} |\text{original-domain}(v)|} \quad (8)$$

The variable-ordering heuristic *tight* selects a variable from the (estimated) dynamically tightest cluster. Search guided by *tight*, however, could shift from one cluster to another, and therefore from one satellite to another in *Comp*, the way the poorly-performing *satellite* did. The improvement produced by *stay* in Table 1 therefore inspired heuristics that treat one cluster at a time. *Concentrate* chooses a cluster at random, selects variables from it until all of them are bound, and then selects the next cluster at random. In contrast, *focus* selects the (estimated) dynamically tightest cluster, selects variables from it using a traditional variable-ordering heuristic (e.g., *MinDomWdeg*) until all of them are bound, and then uses estimated dynamic tightness to select the next cluster. Note that *tight*, *concentrate* and *focus* only select clusters, not variables. A traditional variable-ordering heuristic is required to select variables within the current cluster of interest. *Concentrate-i* and *focus-i* are analogous to *until-i* in Section 4; they instantiate within a cluster until all but  $i$  of its variables have been bound. In all these heuristics, if clusters have the same maximum tightness, ties are broken by maximum dynamic cluster size, the number of unbound variables in a cluster.

In the experiments in this section, each heuristic had 30 min to solve each *Comp* problem. Data for all non-deterministic algorithms, including those involving clusters, is reported as an average across 10 runs. For the heuristics that use cluster detection, time  $e$  is allocated to VNS per cluster. Thus a problem in which  $s$  clusters were detected could require up to  $se$  time. (Because elapsed time is tested only at the end of a loop iteration, it is possible to slightly exceed  $se$  in practice.) The total VNS time required to find clusters is included in all search time data.

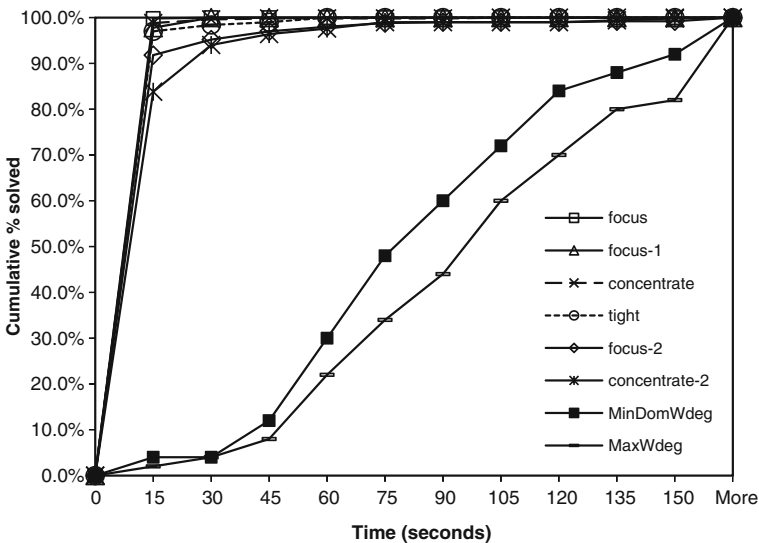
On *Comp* problems, *Foretell* finds clusters that form a structure remarkably like foreknowledge. It found between 6 and 19 clusters per problem, all of sizes 3 to 6. It found at least one cluster in every satellite in every problem on every run. A typical

**Table 2** Cluster-guided search speeds traditional heuristics on *Comp* by more than an order of magnitude. Average and standard deviation are shown for nodes and time in CPU seconds, including time for cluster detection. Data above the line is repeated from Table 1. Except for *MinDomDeg*, every method solved every problem. *Focus* is statistically significantly better (in bold) than all the heuristics tested. *Until-11* is a target, not a legitimate heuristic; it applies foreknowledge about structure available only to the problem generator, not to the search engine

Heuristic	Time, $\mu$ ( $\sigma$ )		Nodes, $\mu$ ( $\sigma$ )	
<i>MinDomDeg</i>	1,728.157	(355.877)	285,751.970	(61,368.701)
<i>MaxWdeg</i>	123.000	(128.580)	20,817.640	(22,954.165)
<i>MinDomWdeg</i>	83.580	(38.964)	12,519.360	(5,811.370)
<i>Until-11</i>	1.612	(1.866)	398.776	(244.112)
<i>Tight</i>	4.705	(6.252)	505.296	(718.029)
<i>Concentrate</i>	5.461	(5.628)	836.434	(876.539)
<b><i>Focus</i></b>	<b>4.311</b>	(2.411)	<b>497.964</b>	(324.327)
<i>Focus-1</i>	5.267	(3.215)	516.406	(425.739)
<i>Focus-2</i>	8.713	(22.442)	1,371.338	(2,765.681)

result appears in Fig. 1d. With  $e = 0.2$  s per cluster, VNS search time averaged 2.111 s per problem, 49% of the total time.

Clusters guide search in *Comp* effectively, as shown below the line in Table 2. *Concentrate*'s weaker performance clearly indicates that the order in which clusters are addressed is important. Unlike *stay*, however, *focus* appears to need to finish a cluster to produce its best performance. Essentially, by  $i = 3$ , both *concentrate-i* and *focus-i* deteriorate to *MinDomWdeg*. (Data omitted.) A full graphic comparison (Fig. 7) indicates that even *focus-2* and *concentrate-2* solve most *Comp* problems far more quickly than the learning heuristics *MaxWdeg* and *MinDomWdeg* do alone. Also, *concentrate* solves more *Comp* problems than *tight* in 45 s or less. One



**Fig. 7** Cumulative percentage of 50 *Comp* problems solved. Solvers were allocated 30 min per problem. Because it solved only two of the problems, *MinDomDeg* was omitted

possible explanation is that in these CSPs *Foretell's* clusters are all of roughly equal importance, so that *concentrate* benefits from a refusal to shift from one cluster to another. As more clusters are solved, however, it becomes important to instantiate within tight clusters, so that *tight* would then have an advantage. *Focus* combines the best of both approaches.

Given the vagaries of local search, one cannot expect VNS to produce an adequate set of clusters every time. Rather than allot substantial time to VNS (which should ultimately find adequate clusters that way), we used *MinDomWdeg* to select individual variables within a cluster. *MinDomWdeg* is slightly slower than *MinDomDeg* at selecting variables inside a cluster, but it also provides backup if *Foretell's* local search is simply “unlucky.” Learning is there to help, although it is rarely necessary.

### 8 Clusters and inference

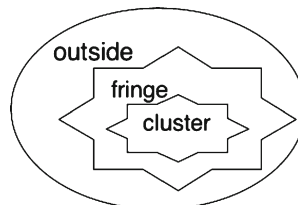
A cluster graph also provides information that can be harnessed to guide inference. Inference methods can be characterized along a spectrum by the effort they expend. More inference does not always result in more domain reduction—it is often faster to risk and retract mistakes than to anticipate them. Inference after every assignment, as in Algorithm 1, is called *consistency maintenance*. FC and MAC-3 (as described in Section 2.2.1) are commonly used to maintain consistency. FC enforces a lower level of consistency by only propagating the effect of a variable assignment to its neighbors. MAC-3 does considerably more work to propagate that effect to the entire problem. *ACR-k* takes a stance between FC and MAC-3 [10]. It begins with the same initial queue as MAC-3, but subsequently enqueues only constraints on variables whose dynamic domains lose at least *k%* of their values. (The R is for “response.”) Intuitively, higher values for *k* make ACR lazier.

With the structural knowledge detected by *Foretell*, it becomes possible to fine tune consistency enforcement. *Cluster-based inference* considers where other variables lie with respect to the clusters. Each cluster *C* in problem *P* delineates a *fringe* (variables in  $P - C$  within *width* edges of some variable in *C*), and an *outside* ( $P - C - fringe(C)$ ), as shown in Fig. 8. The question then becomes how to select propagation methods for the cluster, the fringe, and the outside.

To begin, we generated classes of small, not necessarily solvable CSPs similar to the clusters *Foretell* finds. The densest possible graph is a clique. Intuitively, a near clique is a subgraph that is a few edges short of a clique. A *near clique* is defined recursively as follows:

- The clique on 3 vertices  $K_3$  is a near clique.
- Given a near clique  $NC = \langle V, E \rangle$  with missing edges  $m = \frac{|V|(|V|-1)}{2} - |E|$

**Fig. 8** Propagation regions delineated with respect to a cluster

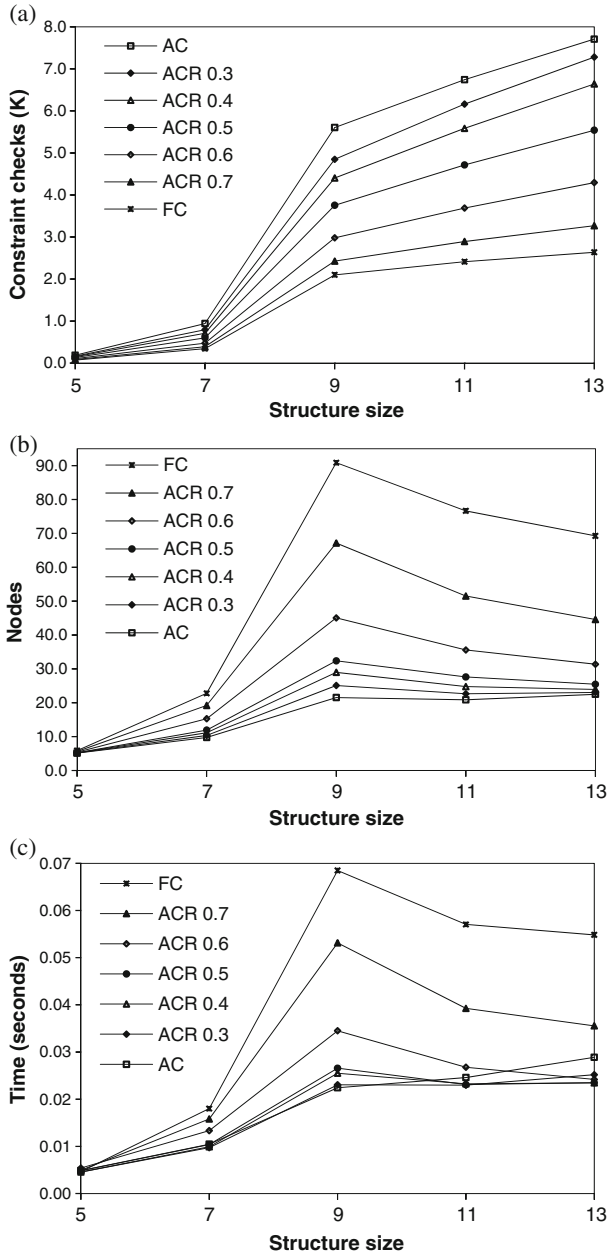




a new near clique  $NC' = \langle V \cup \{v\}, E \cup \{e_1, e_2, \dots, e_k\} \rangle$  can be constructed from  $NC$  by the introduction of a new vertex  $v$  and  $k$  new edges  $e_1, e_2, \dots, e_k$  to  $NC$  such that the increase  $\Delta m$  in the number of missing edges conforms to

$$\Delta m < \frac{|V|}{2} + \frac{m}{|V| - 1} \tag{9}$$

**Fig. 9** Number of **a** checks, **b** expanded nodes, and **c** CPU seconds required to solve cluster-like graphs of various sizes under seven different propagation methods. Lazier propagation does fewer checks, expands more nodes, and is sometimes faster



For  $|V| > 3$ , (8) requires

$$m \leq \left\lfloor \frac{|V| - 1}{2} \right\rfloor \quad (10)$$

To simulate *Foretell*'s clusters, we generated classes of CSPs that were near cliques of sizes 5 to 13 with edge tightness 0.5, and solved them with *MinDomDeg* in separate runs that maintained consistency with FC, MAC-3, or a MAC-3-like version of ACR- $k$  for  $k$  from 0.3 to 0.7. The results appear in Fig. 9. AC is warranted while clusters are of size no more than 7, but on larger simulated clusters it is statistically significantly slower than ACR-0.4. ACR-0.4 also showed low variation in performance on the simulated clusters.

The structure of a cluster graph suggests the design of cluster-based inference methods. Let  $c/w/f/o$  be a cluster-based method that propagates within a cluster with method  $c$ , within its fringe of width  $w$  with method  $f$ , and outside with method  $o$ . For *Comp*, the clusters in Fig. 6a are small; that mandates AC propagation within them. Because *Comp* clusters are often linked, even a fringe of width 1 may reach other clusters. Thus AC is a wise approach within the fringe as well. Once outside the clusters, propagation can afford to be lazier. Thus a reasonable cluster-based propagation method for *Comp* is as AC/2/AC/FC. RLFAP's cluster graph is different; the clusters are larger, so that propagation within clusters of ACR 0.4 is reasonable. The relative isolation of the four crucial clusters there suggests propagation in the fringe with ACR-0.6, but FC is expected to be safe in the rest of the graph. This produces the method ACR-0.4/1/ACR-0.6/FC. Finally, the large clusters in the driver problems are so closely connected that it may only be reasonable to attempt ACR-0.4/1/ACR-0.5/FC.

## 9 Experimental design and results

The experiments in this section compare *MinDomWdeg* with the strongest cluster-based search heuristic: *Foretell* and *focus* with *MinDomWdeg*. To control for the vagaries of local search, performance during any experiment with *Foretell* was averaged across 10 trials for each problem. On each problem, *Foretell* was given some number of milliseconds (ms.) per cluster, and identified as many clusters as it could until a call to VNS failed. Table 3 lists the number, average size, and maximum size of the clusters detected with *Foretell* prior to search.

Algorithms were tested on all the classes of composed problems on the benchmark website [23], where there are 10 problems per class. A problem described there by  $a - b - c$  denotes a central component with  $a$  variables,  $b$  satellites of eight variables each and  $c$  links. All variables have domain size 10, with constraint tightness 0.150 within the central component and 0.050 on each link. Constraint density within a satellite is always 0.786. The structure of the composed problems in these classes is deliberately obscured. Nonetheless, *Foretell*'s output matches the descriptions provided for those problems.

The upper section of Table 3 reports those results. In 30 min each, *MinDomDeg* could solve only nine of those 90 problems. That and the search tree sizes for *MinDomWdeg* suggest that these benchmarks are easier than *Comp*. On six classes,

**Table 3** At the 95% confidence level, *focus* outperforms *MinDomWdeg* on these problem classes. Order of magnitude improvements over *MinDomWdeg* are in bold. Classes above the line are composed, with central component density  $d$ , satellite tightness  $t'$ , and link density  $d''$ . Time is in CPU seconds. Data for *Foretell* includes number of clusters, average cluster size, and maximum cluster size, averaged across 10 runs. Data for *focus* is mean and standard deviation over 10 runs

	Number of instances			MinDomWdeg			Foretell's clusters			Focus time			Focus Nodes		
	$d$	$t'$	$d''$	Time	Nodes	Count	Size	Max	$\mu$	$\sigma$	$\mu$	$\sigma$	$\mu$	$\sigma$	
<b>Composed CSPs</b>															
25-10-20	0.667	0.50	0.010	2.49	670.10	10.17	5.20	5.58	0.88	0.47	192.07	149.88			
25-1-80	0.667	0.65	0.010	0.95	308.00	5.60	5.28	6.08	0.26	0.25	94.50	71.81			
75-1-80	0.216	0.65	0.133	2.32	595.20	9.09	4.86	5.90	0.37	0.17	181.40	21.69			
25-1-2	0.667	0.65	0.010	1.01	553.00	1.01	5.77	5.77	<b>0.02</b>	0.00	<b>41.40</b>	1.36			
25-1-25	0.667	0.65	0.125	0.91	465.70	2.30	5.60	5.90	<b>0.04</b>	0.02	<b>41.60</b>	1.29			
25-1-40	0.667	0.65	0.200	1.10	473.80	5.00	5.37	6.40	<b>0.07</b>	0.02	<b>41.50</b>	1.21			
75-1-2	0.216	0.65	0.003	3.33	1171.70	1.00	5.69	5.69	<b>0.04</b>	0.01	<b>91.60</b>	1.50			
75-1-25	0.216	0.65	0.042	3.29	1084.40	5.40	5.24	6.46	<b>0.15</b>	0.12	<b>91.40</b>	1.29			
75-1-40	0.216	0.65	0.067	2.97	960.90	4.60	5.29	5.80	<b>0.15</b>	0.14	<b>91.30</b>	1.28			
Comp	0.150	0.50	0.120	83.58	12519.40	11.00	4.31	5.15	<b>4.31</b>	2.41	<b>497.96</b>	324.33			
<b>Real-world CSPs</b>															
	$n$	$k$	Number of constraints												
RLFAP scene 11	680	44	4103	36.97	2810.00	5.50	12.48	18.00	13.22	0.14	980.10	1.45			
RLFAP scene 11_f10	680	34	4103	133.24	8768.00	13.00	11.95	17.80	35.46	0.06	2644.00	0.00			
driverlogw 09	650	12	17447	498.98	15987.00	12.40	17.43	30.10	179.13	17.95	5119.00	489.70			
driverlogw 08cc	408	11	9312	85.82	4880.00	4.00	31.75	46.00	41.58	0.06	2544.00	0.00			
driverlogw 08c	408	11	9312	86.11	4820.00	4.00	31.75	46.00	35.67	0.06	2289.00	0.00			
driverlogw 04	272	11	3876	6.01	751.00	18.80	9.01	23.80	3.17	0.38	350.10	51.85			
driverlogw 02	301	8	4055	16.11	1862.00	18.80	8.01	15.80	5.61	1.42	649.30	189.61			
os-taillard-4-95-0	16	182	48	28.58	303.00	5.00	3.20	4.00	6.13	0.09	95.00	0.00			
os-taillard-4-95-1	16	220	48	255.43	4721.00	5.00	3.18	3.90	142.76	2.30	1577.80	10.12			

*focus* once again provided an order of magnitude speedup. Note that, for a fixed central component size, *focus* scales about linearly with problem size.

Clusters are often readily detected in CSPs for real-world problems too. The lower section of Table 3 includes RLFAP, driverlog, and Taillard jobshop problems. Scene11 is the most difficult RLFAP; scene11\_f10 is a modified scene11 problem with its highest 10 radio frequencies (domain values) removed. This modification makes the problem unsolvable and considerably more difficult than the original problem. The jobshop problems (os-taillard) are intensional CSPs, whose constraints are defined by predicates. Since their predicates are disjunctive inequalities over integer intervals, ACE solves the inequalities to calculate constraint tightness, which *Foretell* requires. For cluster-based search, time includes the time used by *Foretell* to detect clusters. The difficulty of a class of problems is gauged here by the resources *MinDomWdeg* required to solve it. On all real-world CSPs, cluster-based search significantly outperforms *MinDomWdeg*, at the 95% confidence level, on both time and nodes. These results support the premise that clusters detected by *Foretell* address the hardest parts of a problem. Far fewer incorrect assignments were made under cluster-based search.

Cluster-based inference was added to cluster-guided search and tested on all the problems in Table 3. On all the classes of composed problems there was little room for improvement over cluster-guided search, and none appeared. On RLFAP, however, cluster-based inference further accelerated cluster-guided search by 4%, a statistically significant improvement. As Fig. 9 anticipated, the laziness of ACR-*k* engendered more mistakes than AC, so there was no concomitant reduction in nodes. The success of cluster-based inference on RLFAP suggests that *Foretell*'s clusters cover enough of the backdoor so that FC suffices for the "outside." (This improvement is not attributable solely to FC; FC alone is dramatically slower on this problem.) On the driverlog problems, cluster-based inference did not improve cluster-guided search. We suspect that this is because the structure of cluster graphs is markedly different.

## 10 Discussion and conclusion

Density and tightness are synergistic. Earlier work [11] tested them separately on classes of smaller, considerably easier composed problems, ones that even *MinDomDeg* could solve. A heuristic that prioritized variables by tightness roughly halved *MinDomDeg*'s search time. A heuristic that prioritized variables by density (with VNS-based near clique detection) consumed about a third of the search time. When combined in an earlier version of *Foretell*, however, density and tightness did an order of magnitude better, and produced nearly backtrack-free search trees [11]. On smaller composed problems with one or two satellites in the earlier study, clusters did not harm performance, and they sometimes improved it. While earlier work and that reported here both use the same local search mechanism to find crucial substructure, they differ in two significant ways: the key measurement used to guide local search and the problems tested. The earlier work used *tension*, the dynamic reduction in the domains of the neighbors of a variable, while this paper uses *pressure* as defined in (6), an estimate of the probability that a variable's domain will be reduced when its neighbor is assigned a value. The earlier paper experimented

only on smaller composed problems with extensional constraints generated by the authors, while this paper emphasizes real-world and benchmark problems, including far larger composed problems whose scale is closer to that of real-world problems.

### 10.1 Cluster detection with *Foretell*

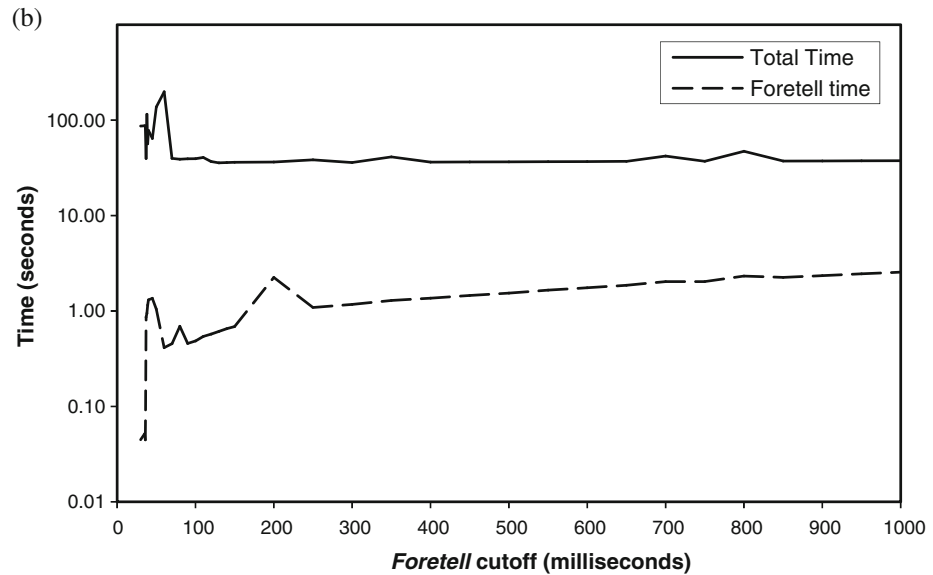
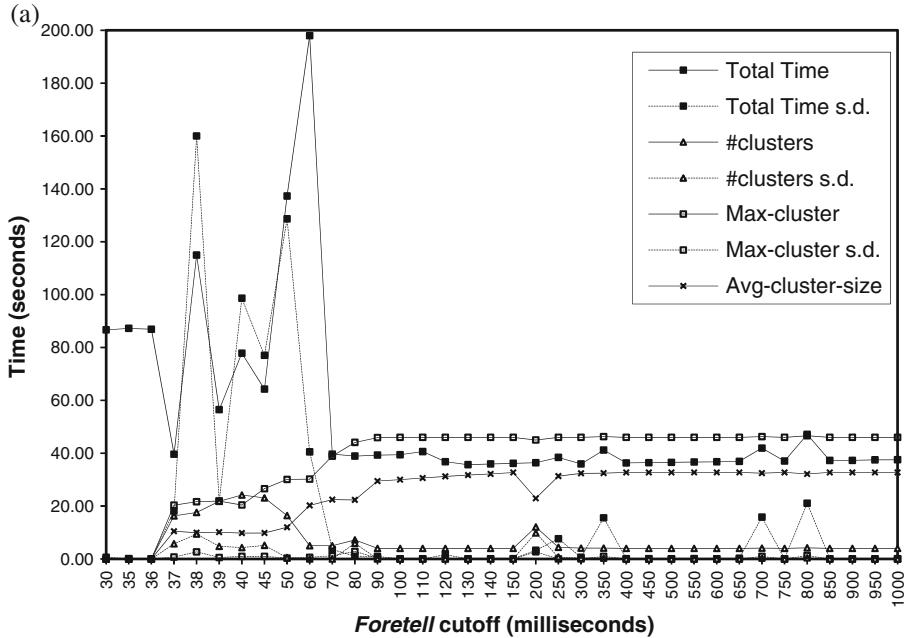
*Foretell*'s key parameter, *cutoff*, determines how much time to devote to the detection of any single cluster. *Foretell* has no prior knowledge about how many clusters lie within a problem or about how many might be necessary to solve it. We have found empirically that too few clusters provide inconsistent guidance, and that, as *cutoff* is increased, *Foretell* finds more consistent sets of clusters from one run to the next. Consider, for example, the experiments on *driverlog-08c* reported in Fig. 10, where each data point represents 10 runs with a single *cutoff* value. At 30, 35 and 36 ms. per cluster, *cutoff* was too small—*Foretell* found no clusters, and the resultant performance was effectively that of *MinDomWdeg*. We then tested small *cutoff* values further. Values between 37 and 80 ms, produced large variations in both the identified cluster graphs and the resultant search performance. *Foretell* found as many as 29 clusters on some runs at 38 and 45 ms. The best run, however, with *cutoff* = 37 ms, found 18 clusters and solved the problem in 18.819 s, about half the time reported in Table 3. While 37 ms and other *cutoff* values in that range do not perform well on average, the structure that led to the 18.819 s solution merits further study, and may ultimately motivate new heuristics able to support consistent performance at this level. Larger *cutoff* values produced more stable *Foretell* results.

As *cutoff* increases, the number of detected clusters stabilizes and decreases. Given more than 80 ms. the same largest cluster in *driverlog-08c* was found consistently. In general, there is no difference among the identified cluster graphs (Fig. 6f) and search performance beyond 80 ms. Because total time is the time *Foretell* uses to find all clusters plus the time to search for a solution, increasing *cutoff* can increase total time. As Fig. 10b shows, however, there is a wide range of cutoff values (from 37 to 1,000 ms), in which the time *Foretell* actually consumes is lower than this maximum allocation, indeed at least an order of magnitude lower than the total time, thereby assuring robust search performance.

In general, smaller *cutoff* values, although of interest for structure analysis and the design of new heuristics, are too aggressive to produce consistent performance. Larger cutoff values lead to consistent cluster graphs and benefit users who seek to solve the problem effectively. Current work addresses additional termination conditions for *Foretell* (line 4 in Algorithm 3). These include an overall VNS time limit, a Luby-like adaptive cutoff [25] for allocations on successive clusters, and a limit on the percentage of variables that may be included in either an individual cluster or the entire cluster graph.

### 10.2 Why *focus* works

Variable-ordering heuristics usually do not consider persistence in a “geographic area” of a problem. Nonetheless, that was clearly *satellite*'s mistake—even with foreknowledge about *Comp*, it *satellite hopped*, that is, it failed to address enough variables in the same satellite consecutively. *Stay* forbade satellite hopping and resulted in a considerable improvement. Analogously, *cluster hopping* occurs when



**Fig. 10** Average results of 10 runs on driverlogw-08c. **a** The relationships between *cutoff*, the time allocated to find one cluster, and total time, number of detected clusters, maximum cluster size and average cluster size. Standard deviations are shown for total time, number of clusters and maximum cluster size. Total time includes both time to find all clusters and search time to solution. The x-axis is not proportional to the values of *cutoff*. **b** The relationship between *Foretell* time and total time

a heuristic fails to address enough variables in the same cluster consecutively. Because constraints within a cluster are selected for above average tightness, once any variable in a cluster has been bound, propagation is likely to reduce the domains of the other variables in that cluster. As a result, variables in a partially-instantiated cluster are more likely to have smaller domain sizes and make their cluster even more attractive to cluster-guided search. *Tight* was permitted to cluster hop, while *focus* and *concentrate* both explicitly forbade it. *Focus* does better, however, because it uses knowledge about clusters to select one.

Luckily, remaining in a cluster in *Comp* is likely to encourage remaining in any additional clusters within the same satellite. In problems that are not composed, any other region with sufficiently dense and/or tight connections to a cluster should also have the domains of its variables reduced when those of the related cluster are instantiated. In this way, cluster-guided search results in a sequence of decisions that persist in a particularly constrained region of the graph.

A subproblem reduced to an arc-consistent tree (which always has at least one solution) would make it safe to continue on to another subproblem. Binding  $w$  variables in a subproblem of size  $s$  with density  $d$ , leaves a tree only if

$$d \binom{s-w}{2} \leq s-w-1, \text{ that is, } w \geq s - \frac{2}{d} \quad (11)$$

(Of course, this only makes a tree possible, not certain.) For *Comp* satellites,  $s = 20$  and  $d = 0.25$ , so that there is no possibility of a tree unless  $w \geq 12$ , that is, we have bound 12 variables and 8 remain (*until-8*). Our empirical results, however, show that *until-11* minimizes both time and nodes (using a one-tail  $t$ -test at the 95% confidence level). This ability to leave behind a (necessarily) cyclic subgraph is probably attributable to propagation. The occasional retraction back to a “finished” satellite proved less costly than binding a few more variables in the current satellite before moving on to the next one. Because clusters in *Comp* average  $s = 4.309$ , however,  $w \geq 0.309$ , that is, only *focus-0* is safe, which is exactly what our results indicate.

### 10.3 Clusters and search

The performance of perfect foreknowledge on *Comp*, as embodied by *until-11*, is the gold standard. *Until-11* knows a superset of the backdoor and exploits it. Inspection indicates that the backdoor is probably no more than 35 variables for a *Comp* problem. The last retraction on a *Comp* problem under *focus* was at a node where an average of 15.588 variables had been bound, with a maximum of 62.

The structural knowledge learned provides an explanation of where the difficulties lie in a CSP. Figure 6a focuses attention on the satellites, but the solution with *focus* is even more descriptive: it searched within at most three satellites before it reported the insolubility of Fig. 1. To demonstrate the insolubility of the *Comp* problem in Fig. 1, *focus* bound only 12 variables (out of 200) drawn from three clusters found by *Foretell*. Those three clusters, two of size 5 and one of size 4, provide a concise and more satisfying explanation than either a search tree rooted at a single node or a collection of edge weights.

RLFAP and the driverlog problems demonstrate that a problem need not have satellites to have clusters. On small-world problems, for example, almost every variable is quickly shown to lie in some cluster. Having clusters, however, does not justify directing computational resources to *Foretell*. On easy problems, it is faster to use *MinDomWdeg* or even *MinDomdeg*. Clusters are not detected dynamically, during search, because *Foretell* does not find clusters in order of either tightness or size. To identify a good starting point, *focus* must therefore choose among a set of clusters. This static but predictive perspective serves search well.

The two driverlog problems differ in the tuples they allow, but they have the same primary structure and the same tightness on their constraints. On every run, *Foretell* found the identical sparse secondary structure in the two problems, that is, exactly the same clusters among the 408 variables: three large nodes (two single-cluster nodes and one two-cluster node) with two edges. That *focus* improves search on them both confirms its ability to manage structure dynamically.

Both composed problems and real-world problems have non-random structure and varying tightness. Traditional heuristics like *MinDomDeg* perform poorly on composed problems because of the difference in degree and tightness within the satellites and the central component. Composed problems provide an elegant, if artificial, argument for the need to consider tightness during search. Compared to real-world problems, however, composed problems offer a known structure that allows us to study (and aspire to) performance given perfect structural knowledge. (This is the point of the study of *Comp* in Section 4.) Moreover, composed problems' simpler structure makes it easier to monitor the entire search process and to understand the differences among various search regimens. What was learned from *Comp* applies to other artificial structured classes too, as shown in Section 9. While composed problems are built to confound traditional heuristics in a particular way, real-world problems are merely difficult. Not surprisingly, the performance improvements on real-world problems (below the line in Table 3) are noteworthy, but less dramatic. The insights provided by the structure detected by *Foretell*, however, could prove meaningful to a user. For example, people who know RLFAP Scene 11 well may be interested in *Foretell*'s detection of just five clusters, which include only 67 variables out of 680. Three of those clusters form a triangle in the cluster graph.

As stated earlier in this section, *Foretell* offers no advantage when a problem is too easy or is unstructured. We therefore chose representative benchmark instances that could benefit from *Foretell*, including a class of seven driverlog problems and the 11 scenes of the RLFAP problems. Problems shown in the lower part of Table 3 represent the performance improvements *Foretell* achieved on these two classes of real-world CSPs. (There was no significant difference on the easier problems.)

Search with *Foretell* is a two-stage process: First *Foretell* seeks clusters and then clusters guide search for a solution. We separated structure detection from structure guidance because, during search, retractions that back out of a cluster could demand frequent calls to find new clusters. This overhead could be expensive and reduce search performance. Thus *Foretell* is called only before search, and the structural knowledge learned by *Foretell* remains static during the subsequent search for a solution. This strategy appears to be effective for the problems we have tested.

Early departure from clusters during search, as with *focus-i* in Section 7 and Table 2, has been shown to be less effective for *Comp* than *focus*. The difference



between *until-i* and *focus-i* is the structural knowledge on which they rely. *Until-i* uses perfect knowledge, which covers the entire satellite, but *focus-i* only considers the cluster that partially covers the underlying satellite. Because of this partial coverage of satellites by clusters, *focus* needs all the structural knowledge it can muster to achieve its largest performance improvement. Topics of current work include how early departure from a cluster affects search on real-world instances, and the relation between subproblem coverage by clusters (where perfect structural knowledge about subproblems is available) and performance.

No solver, human or machine, has an efficient way to “see” Fig. 1c perfectly without knowledge about the problem generator. Generally, a cluster graph is a prediction of significant structure where global search is likely to fail. A user confronted with an unsolvable real-world problem could use clusters to reconsider the problem’s specifications, or at least to understand why a problem is difficult to solve or has no solution at all. Thus clusters can be used not only to focus attention during search but also to provide insight into the nature of the problem in a user-friendly representation.

Methods to detect tight, dense subproblems must not only be incisive, they must also scale. Every real-world problem that we tested (i.e., all the RLFAP and driver problems) contained clusters that *Foretell* found fairly quickly. The *Foretell cutoff* for RLFAP scene11, which has roughly three times as many variables as a *Comp* problem, is about three times as large as the *Foretell cutoff* for *Comp*. This suggests that *Foretell* scales linearly.

Several enhancements are currently under development. Not every problem needs *Foretell*. The “right” clusters are not necessarily many, but incisive, so *Foretell* could partition less than the entire problem. Other problems have more dense structures and may therefore respond better to other cluster-based orderings. Finally, *focus* might attend more closely to learned weights before all the cluster variables have been bound.

Cluster-based propagation is still under development. Because cluster sizes vary in real-world problems, ACR seems a wise choice unless the problem has uniformly small clusters. Cluster-based propagation should be further tailored to the metastructure of the cluster graph, including cluster size, domain size, variance in internal edge tightness, and the number and tightness of inter-cluster edges.

Other work introduces the concept of *impact*, the influence of search space reduction, for every variable [32], and uses a matrix-based representation to visualize pair-wise impacts between variables [4]. On artificial instances with tighter subproblems, this visualization shows such structure. It would be interesting to compare the structure learned by *Foretell* before search and the impact structure learned during search.

Although the experiments described here are on binary CSPs, we see no obvious impediment to adapting this approach for non-binary constraints. As long as there is some estimate of the tightness of a constraint, it is possible to estimate the pressure on a variable and to detect sets of mutually-constrained variables by local search. The swap and score functions would require only some modification. Real-valued domains present a different challenge, one we believe surmountable through the methods planned for large domains. There are also problems in which *Foretell* cannot find any clusters at all. Other structures, such as lengthy cycles, can create search difficulty without local density [28]. Something similar may be operative in these problems.

As observed earlier, ACE is not honed for speed. Nonetheless, the concomitant reductions in checks and nodes searched suggest that clusters will accelerate other, more agile solvers as well. For an easy problem, no clusters are necessary, and any reasonable amount of time spent on cluster detection will have no noteworthy impact. For more challenging problems, however, cluster-guided search substantially accelerated search with off-the-shelf heuristics on problems with sparse secondary structure. Given their acuity and explanatory ability, clusters are a worthwhile preprocessing step.

**Acknowledgements** This work was supported in part by the National Science Foundation under awards IIS-0811437 and IIS-0739122. ACE is a joint project with Eugene Freuder and Richard Wallace of the Cork Constraint Computation Centre. Thanks go to Pierre Hansen for helpful discussions on VNS.

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