

An Integrated Solver for Optimization Problems

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One of the central trends in the optimization community over the past several years has been the steady improvement of general-purpose solvers. A logical next step in this evolution is to combine mixed-integer linear programming, constraint programming, and global optimization in a single system. Recent research in the area of integrated problem solving suggests that the right combination of different technologies can simplify modeling and speed up computation substantially. Nevertheless, integration often requires special-purpose coding, which is time consuming and error prone. We present a general-purpose solver, SIMPL, that allows its user to replicate (and sometimes improve on) the results of custom implementations with concise models written in a high-level language. We apply SIMPL to production planning, product configuration, machine scheduling, and truss structure design problems on which customized integrated methods have shown significant computational advantage. We obtain results that either match or surpass the original codes at a fraction of the implementation effort.

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1. Introduction

One of the central trends in the optimization community over the past several years has been the steady improvement of general-purpose solvers. Such mixed-integer solvers as CPLEX (ILOG 2007) and XPRESS-MP (Guéret et al. 2002) have become significantly more effective and robust, and similar advancements have occurred in continuous global optimization (BARON, Tawarmalani and Sahinidis 2004; LGO, Pintér 2005) and constraint programming (CHIP, Dincbas et al. 1988; ILOG Solver, ILOG 2003). These developments promote the use of optimization and constraint-solving technology because they spare practitioners the inconvenience of acquiring and learning different software for every application.

A logical next step in this evolution is to combine mixed-integer linear programming (MILP), constraint programming (CP), and global optimization in a single system. This not only brings together a wide range of methods under one roof, but it allows users to reap the advantages of integrated problem solving. Recent research in this area shows that the right combination of different technologies can simplify modeling and speed up computation substantially. Commercial-grade solvers are already moving in this direction, as witnessed by the ECLⁱPS^e solver (Rodošek

et al. 1999), OPL Studio (Van Hentenryck et al. 1999), and the Mosel language (Colombani and Heipcke 2002, 2004), all of which combine mathematical programming and constraint programming techniques to a greater or lesser extent.

Our objective is not to demonstrate, in a single paper, that integrated methods can provide a superior alternative to state-of-the-art standalone MILP or CP solvers. This must be accomplished by an entire community of researchers examining a wide variety of problems. There is, in fact, a growing literature that documents the advantages of integrated methods. Table 1 presents a sampling of these results, and Hooker (2007) cites many additional examples. Our contribution is to show, both theoretically and empirically, that it is possible to unify most of what is known about integrated methods under one modeling and solution paradigm, without sacrificing performance.

Integration is most effective when techniques interleave at a micro level. To achieve this in current systems, however, one must often write special-purpose code, which slows research and discourages application. We have attempted to address this situation by designing an architecture that achieves low-level integration of solution techniques with a high-level modeling language. The ultimate goal is to build an integrated solver that can be used as conveniently

Table 1. Sampling of computational results for integrated methods.

Source	Type of problem/method	Speedup
Loose integration of CP and MILP		
Hajian et al. (1996)	British Airways fleet assignment. CP solver provides starting feasible solution for MILP.	Twice as fast as MILP, 4 times faster than CP.
CP plus relaxations similar to those used in MILP		
Focacci et al. (1999)	Lesson timetabling. Reduced-cost variable fixing using an assignment problem relaxation.	2 to 50 times faster than CP.
Refalo (1999)	Piecewise-linear costs. Method similar to that described in §6.	2 to 200 times faster than MILP. Solved two instances that MILP could not solve.
Hooker and Osorio (1999)	Boat party scheduling, flow shop scheduling. Logic processing plus linear relaxation.	Solved 10-boat instance in 5 min that MILP could not solve in 12 hours. Solved flow shop instances 4 times faster than MILP.
Thorsteinsson and Ottosson (2001)	Product configuration. Method similar to that described in §7.	30 to 40 times faster than MILP (which was faster than CP).
Sellmann and Fahle (2001)	Automatic digital recording. CP plus Lagrangean relaxation.	1 to 10 times faster than MILP (which was faster than CP).
Van Hoeve (2003)	Stable set problems. CP plus semidefinite programming relaxation.	Significantly better suboptimal solutions than CP in fraction of the time.
Bollapragada et al. (2001)	Nonlinear structural design. Logic processing plus convex relaxation.	Up to 600 times faster than MILP. Solved 2 problems in <6 min that MILP could not solve in 20 hours.
Beck and Refalo (2003)	Scheduling with earliness and tardiness costs.	Solved 67 of 90 instances, while CP solved only 12.
CP-based branch and price		
Easton et al. (2002)	Traveling tournament scheduling.	First to solve 8-team instance.
Yunes et al. (2005)	Urban transit crew management.	Solved problems with 210 trips, while traditional branch and price could accommodate only 120 trips.
Benders-based integration of CP and MILP		
Jain and Grossmann (2001)	Min-cost planning and disjunctive scheduling. MILP master problem, CP subproblem (§8).	20 to 1,000 times faster than CP, MILP.
Thorsteinsson (2001)	Jain and Grossmann problems. Branch and check.	Additional factor of 10 over Jain and Grossmann (2001).
Timpe (2002)	Polypropylene batch scheduling at BASF. MILP master, CP subproblem.	Solved previously insoluble problem in 10 min.
Benoist et al. (2002)	Call center scheduling. CP master, LP subproblem.	Solved twice as many instances as traditional Benders.
Hooker (2004)	Min-cost and min-makespan planning and cumulative scheduling. MILP master, CP subproblem.	100 to 1,000 times faster than CP, MILP. Solved significantly larger instances.
Hooker (2005b)	Min-tardiness planning and cumulative scheduling. MILP master, CP subproblem.	10 to >1,000 times faster than CP, MILP when minimizing number of late jobs; ~10 times faster when minimizing total tardiness, much better solutions when suboptimal.
Rasmussen and Trick (2005)	Sports scheduling to minimize number of consecutive home or away games.	Speedup of several orders of magnitude compared to previous state of the art.

as current mixed-integer, global, and constraint solvers. We have implemented our approach in a system called SIMPL, which can be read as a permuted acronym for Modeling Language for Integrated Problem Solving.

SIMPL is based on two principles: algorithmic unification and constraint-based control. *Algorithmic unification* begins with the premise that integration should occur at a fundamental and conceptual level, rather than postponed to

the software design stage. Optimization methods and their hybrids should be viewed, to the extent possible, as special cases of a single solution method that can be adjusted to exploit the structure of a given problem. We address this goal with a *search-infer-and-relax* algorithmic framework, coupled with *constraint-based control* in the modeling language. The search-infer-and-relax scheme encompasses a wide variety of methods, including branch-and-cut methods for integer programming, branch-and-infer methods for constraint programming, popular methods for continuous global optimization, nogood-based methods such as Benders decomposition and dynamic backtracking, and even heuristic methods such as local search and greedy randomized adaptive search procedures (GRASPs).

Constraint-based control allows the design of the model itself to tell the solver how to combine techniques so as to exploit problem structure. Highly structured subsets of constraints are written as metaconstraints, which are similar to “global constraints” in constraint programming. Syntactically, a metaconstraint is written much as linear or global constraints are written, but it is accompanied by parameters that specify how the constraint is to be implemented during the solution process (see §5 for details). For example, a metaconstraint may specify how it is to be relaxed, how it will filter domains, and/or how the search procedure will branch when the constraint is violated. When such parameters are omitted, a prespecified default behavior is used. The relaxation, inference, and branching techniques are devised for each constraint’s particular structure. For example, a metaconstraint may be associated with a tight polyhedral relaxation from the integer programming literature and/or an effective domain filter from constraint programming. Because constraints also control the search, if a branching method is explicitly indicated for a metaconstraint, the search will branch accordingly.

The selection of metaconstraints to formulate the problem determines how the solver combines algorithmic ideas to solve the problem. This means that SIMPL deliberately sacrifices independence of model and method: The model must be formulated with the solution method in mind. However, we believe that successful combinatorial optimization leaves no alternative. This is evident in both integer programming and constraint programming because in either case one must carefully write the formulation to obtain tight relaxations, effective propagation, or intelligent branching. We attempt to make a virtue of necessity by explicitly providing the resources to shape the algorithm through a high-level modeling process.

We focus here on branch-and-cut, branch-and-infer, generalized Benders, and a subclass of global optimization methods because these have been implemented so far in SIMPL. The system architecture is designed, however, for extension to general global optimization, general nogood-based methods, and heuristic methods.

A key contribution of this paper is to demonstrate that a general-purpose solver and modeling system can achieve

the computational advantages of integrated methods while preserving much of the convenience of existing commercial solvers. We use SIMPL to model and solve four classes of problems that have been successfully solved by custom implementations of integrated approaches. We find that a properly engineered high-level modeling language and solver can match and even exceed the performance of handcrafted implementations. We presented the basic ideas of SIMPL’s architecture, without computational results, in Aron et al. (2004). The present paper provides more detailed descriptions of SIMPL’s syntax and semantics and demonstrates its performance empirically on a collection of problem instances.

After a brief survey of previous work, we review the advantages of integrated problem solving and present the search-infer-and-relax framework. We then summarize the syntax and semantics of SIMPL models. Following this, we describe how to model and solve the four problem classes just mentioned in an integrative mode. A production planning problem with semicontinuous piecewise-linear costs illustrates metaconstraints and the interaction of inference and relaxation. A product configuration problem illustrates variable indices and how further inference can be derived from the solution of a relaxation. Finally, a machine-scheduling problem, followed by network design and truss structure design problems, show how Benders decomposition and global optimization methods, respectively, fit into our framework. For the truss structure design problem, in particular, we also demonstrate the use of an effective quasi-relaxation technique that can be applied to a general class of global optimization problems (Hooker 2005c). In each case, we exhibit the SIMPL model and present computational results in online Appendix C. We conclude with suggestions for further development. An electronic companion to this paper is available as part of the online version that can be found at <http://or.journal.informs.org/>.

2. Previous Work

Comprehensive surveys of hybrid methods that combine CP and MILP are provided by Hooker (2000, 2002, 2006, 2007), and tutorial articles may be found in Milano (2003).

Various elements of the search-infer-and-relax framework presented here were proposed by Hooker (1994, 1997, 2000, 2003), Bockmayr and Kasper (1998), Hooker and Osorio (1999), and Hooker et al. (2000). An extension to dynamic backtracking and heuristic methods is given in Hooker (2005a). The present paper builds on this framework and introduces the idea of constraint-based control, which is key to SIMPL’s architecture. A preliminary description of the architecture appears in a conference paper (Aron et al. 2004), without computational results. The present paper develops these ideas further and demonstrates that SIMPL can reproduce the computational advantages of integrated methods with much less implementation effort.

Existing hybrid solvers include ECLⁱPS^e, OPL Studio, Mosel, and SCIP. ECLⁱPS^e is a Prolog-based constraint logic programming system that provides an interface with linear and MILP solvers (Rodošek et al. 1999, Cheadle et al. 2003, Ajili and Wallace 2003). The CP solver in ECLⁱPS^e communicates tightened bounds to the MILP solver, whereas the MILP solver detects infeasibility and provides a bound on the objective function that is used by the CP solver. The optimal solution of the linear constraints in the problem can be used as a search heuristic.

OPL Studio provides an integrated modeling language that expresses both MILP and CP constraints (Van Hentenryck et al. 1999). It sends the problem to a CP or MILP solver, depending on the nature of constraints. A script language allows one to write algorithms that call the CP and MILP solvers repeatedly.

Mosel is both a modeling and programming language that interfaces with various solvers, including MILP and CP solvers (Colombani and Heipcke 2002, 2004). SCIP is a callable library that gives the user control of a solution process that can involve both CP and MILP solvers (Achterberg et al. 2008).

3. Advantages of Integrated Problem Solving

One obvious advantage of integrated problem solving is its potential for reducing computation time. Table 1 presents a sampling of some of the better computational results reported in the literature, divided into four groups: (a) Early efforts at integration coupled solvers rather loosely but obtained some speedup nonetheless. (b) More recent hybrid approaches combine CP or logic processing with various types of relaxations used in MILP, and they yield more substantial speedups. (c) Several investigators have used CP for column generation in a branch-and-price MILP algorithm, as independently proposed by Junker et al. (1999) and Yunes et al. (1999). (d) Some of the largest computational improvements to date have been obtained by using generalizations of Benders decomposition to unite solution methods, as proposed by Hooker (2000) and implemented for CP/MILP by Jain and Grossmann (2001). MILP is most often applied to the master problem and CP to the subproblem.

We solve four of the problem classes in Table 1 with SIMPL: piecewise-linear costs (Refalo 1999; Ottosson et al. 1999, 2002), product configuration (Thorsteinsson and Ottosson 2001), planning and scheduling (Jain and Grossmann 2001), and truss structure design (Bollapragada et al. 2001). We selected these problems because the reported results are among the most impressive, and because they illustrate a variety of solution approaches.

Our experiments show that SIMPL reproduces or exceeds the reported advantage of integrated methods over the state of the art at that time. This level of performance can now be obtained with much less effort than invested by

the original authors. All of the problems in Table 1 can in principle be implemented in a search-infer-and-relax framework, although SIMPL in its current form is not equipped for all of them.

Aside from computational advantages, an integrated approach provides a richer modeling environment that can result in simpler models and less debugging. The full repertoire of global constraints used in CP are potentially available, as well as nonlinear expressions used in continuous global optimization. Frequent use of metaconstraints not only simplifies the model, but allows the solver to exploit problem structure.

This implies a different style of modeling than is customary in mathematical programming, which writes all constraints using a few primitive terms (equations, inequalities, and some algebraic expressions). Effective integrated modeling draws from a large library of metaconstraints and presupposes that the user has some familiarity with this library. For example, the library may contain a constraint that defines piecewise-linear costs, a constraint that requires flow balance in a network, a constraint that prevents scheduled jobs from overlapping, and so forth. Each constraint is written with parameters that specify the shape of the function, the structure of the network, or the processing times of the jobs. When sitting down to formulate a model, the user would browse the library for constraints that appear to relate to the problem at hand.

Integrated modeling therefore places on the user the burden of identifying problem structure, but in so doing it takes full advantage of the human capacity for pattern recognition. Users identify highly structured subsets of constraints, which allows the solver to apply the best-known analysis of these structures to solve the problem efficiently. In addition, only certain metaconstraints tend to occur in a given problem domain. This means that only a relevant portion of the library must be presented to a practitioner in that domain.

4. The Basic Algorithm

The search-infer-and-relax algorithm can be summarized as follows:

Search. The search proceeds by solving problem *restrictions*, each of which is obtained by adding constraints to the problem. The motivation is that restrictions may be easier to solve than the original problem. For example, a branch-and-bound method for MILP enumerates restrictions that correspond to nodes of the search tree. Branch-and-infer methods in CP do the same. Benders decomposition enumerates restrictions in the form of subproblems (slave problems). If the search is exhaustive, the best feasible solution of a restriction is optimal in the original problem. A search is exhaustive when the restrictions have feasible sets whose union is the feasible set of the original problem.

Infer. Very often the search can be accelerated by inferring valid constraints from the current problem restriction, which are added to the constraint set. MILP methods

infer constraints in the form of cutting planes. CP methods infer smaller variable domains from individual constraints. (A variable’s domain is the set of values it can take.) Classical Benders methods infer valid cuts from the subproblem by solving its dual.

One can often exploit problem structure by designing specialized inference methods for certain metaconstraints or highly structured subsets of constraints. Thus, MILP generates specialized cutting planes for certain types of inequality sets (e.g., flow cuts). CP applies specialized domain reduction or “filtering” algorithms to such commonly used global constraints as *all-different*, *element*, and *cumulative*. Benders cuts commonly exploit the structure of the subproblem.

Inference methods that are applied only to subsets of constraints often miss implications of the entire constraint set, but this can be partially remedied by *constraint propagation*, a fundamental technique of CP. For example, domains that are reduced by a filtering algorithm for one constraint can serve as the starting point for the domain reduction algorithm applied to the next constraint, and so forth. Thus, the results of processing one constraint are “propagated” to the next constraint.

Relax. It is often useful to solve a relaxation of the current problem restriction, particularly when the restriction is too hard to solve. The relaxation can provide a bound on the optimal value, perhaps a solution that happens to be feasible in the original problem, and guidance for generating the next problem restriction.

In MILP, one typically solves linear programming or Lagrangian relaxations to obtain bounds or solutions that may happen to be integer. The solution of the relaxation also helps to direct the search, as, for example, when one branches on a fractional variable. In Benders decomposition, the master problem is the relaxation. Its solution provides a bound on the optimum and determines the next restriction (subproblem) to be solved.

Like inference, relaxation is very useful for exploiting problem structure. For example, if the model identifies a highly structured subset of inequality constraints (by treating them as a single metaconstraint), the solver can generate a linear relaxation for them that contains specialized cutting planes. This allows one to exploit structure that is missed even by current MILP solvers. In addition, such global constraints as *all-different*, *element*, and *cumulative* can be given specialized linear relaxations (Hooker 2007, Williams and Yan 2001, Yan and Hooker 1999).

5. The Syntax and Semantics of SIMPL

An optimization model in SIMPL is comprised of four main parts: declarations of constants and problem data, the objective function, declaration of metaconstraints, and search specification. The first two parts are straightforward in the sense that they look very much like their counterparts in other modeling languages such as AMPL (Fourer et al.

2003), GAMS (Brooke et al. 1988), Mosel (Colombani and Heipcke 2002, 2004), and OPL (Van Hentenryck et al. 1999). Hence, this section concentrates on explaining the syntax of the metaconstraints and search specification, as well as the semantics of the model. Complete modeling examples are given in online Appendix C.

5.1. Multiple Problem Relaxations

Each iteration in the solution of an optimization problem P examines a *restriction* N of P . In a tree search, for example, N is the problem restriction at the current node of the tree. Because solving N can be hard, we usually solve a *relaxation* N_r of N , or possibly several relaxations.

In an integrated CP-MILP modeling system, linear constraints are posted to a linear programming (LP) solver, normally along with linear relaxations of some of the other constraints. Constraints suitable for CP processing, perhaps including some linear constraints, are posted to a CP solver as well. Extending this idea to other kinds of relaxations is straightforward.

5.2. Constraints and Constraint Relaxations

In SIMPL, each constraint is associated with one or more *constraint relaxations*. To *post* a constraint means to add its constraint relaxations to the appropriate problem relaxations. For example, both the LP and the CP relaxations of a linear constraint are equivalent to the constraint itself. The CP relaxation of the *element* constraint is equivalent to the original constraint, but its LP relaxation can be the convex hull formulation of its set of feasible solutions (Hooker 2000).

In branch-and-bound search, problem relaxations are solved at each node of the enumeration tree. In principle, the relaxations could be generated from scratch at every node because they are a function of the variable domains. Nonetheless, it is more efficient to regenerate only relaxations that change significantly at each node. We therefore distinguish *static* constraint relaxations, which change very little (in structure) when the domains of its variables change (e.g., relaxations of linear constraints are equal to themselves, perhaps with some variables removed due to fixing), from *volatile* relaxations, which change radically when variable domains change (e.g., linear relaxations of global constraints). When updating the relaxations at a new node in the search tree, only the volatile constraint relaxations are regenerated. Regeneration is never necessary to create valid relaxations, but it strengthens the relaxation bounds.

A metaconstraint in SIMPL receives a declaration of the form

```
<name> means {
    <constraint-list>
    relaxation = { <relaxation-list> }
    inference = { <inference-list> }
}
```

where the underlined words are reserved words of the modeling language. <name> is an arbitrary name given to the metaconstraint, whose purpose will be explained in §A.1 of the online supplement. <constraint-list> is a list of one or more constraints that constitute the metaconstraint. For example, this list could be a single global or linear constraint, a collection of linear constraints, or a collection of logical propositions. Finally, there are two other statements: *relaxation*, which is mandatory, and *inference*, which is optional. The <relaxation-list> contains a list of problem relaxations to which the constraint relaxations of <constraint-list> should be posted. An example of such a list is {lp, cp}, which indicates a linear programming relaxation and a constraint programming relaxation. The <inference-list> contains a list of types of inference that <constraint-list> should perform. For example, if <constraint-list> is a global constraint such as *cumulative*, we could invoke a particular inference (or filtering) algorithm for that constraint by writing *inference* = {edge-finding3}, which indicates an $O(n^3)$ edge-finding filtering algorithm (Baptiste et al. 2001). In the absence of the inference statement, either a default inference mechanism will be used or no inference will be performed. The actual behavior will depend on the types of constraints listed in <constraint-list>.

5.3. Specifying the Search

Syntactically, the search section of a SIMPL model is declared as follows:

```
SEARCH
  type = { <search-type> }
  branching = { <branching-list> }
  inference = { <general-inference-list> }
}
```

where the underlined words are reserved words of the modeling language. <search-type> indicates the type of search to be performed. For example, the key words bb, benders, bp, and ls mean branch and bound, logic-based Benders decomposition, branch and price, and local search, respectively. Sometimes, an argument can be added to the search type to specify a particular node selection strategy. For example, bb:bestbound means we want to do branch and bound with best-bound node selection.

The <branching-list> is a comma-separated list of terms in which each term assumes the following form: <name>:<selection-module>:<branching-module>. <name> is the name of a metaconstraint, as described in §5.2. <selection-module> and <branching-module> represent, respectively, the selection module and branching module to be used when branching on the constraint named <name>. Constraints are checked for violation (for branching purposes) in the order they appear in <branching-list>, from left to right. When <constraint-list> in the metaconstraint <name> contains more than one constraint, <selection-module> will specify the order in

which to check those constraints for violation. Current possibilities are most (most violated first), least (least violated first), and first (first violation in the list). Once a violated constraint c is selected, the optional argument *branching-module* specifies how to branch on c . For example, let us say our model has two metaconstraints named $c1$ and $c2$. The branching statement could look like *branching* = { $c1$:most, $c2$:first:sos1 }. This means we first check the constraints in $c1$ for violation and branch on the most violated of those, if any, according to some default criterion. If the solution of the current relaxation satisfies $c1$, we scan the constraints in $c2$ for violation in the order they are listed (because of *first*). If one is found violated, we use *sos1* branching. In case a variable name is used in the branching list instead of a metaconstraint name, this means we want to use the *indomain* constraints of that set of variables for branching. The indomain constraint of a variable is the constraint that specifies the possible values of that variable in the variable declaration section. Examples of such constraints are: $x[1..10]$ in $\{0, 1\}$ for binary variables, and $y[1..5]$ in $[0..10, 12..15]$ for real-valued variables with holes in their domains.

The <general-inference-list> is an optional list of inferences that should be performed in addition to the inferences specified inside each individual metaconstraint (see §5.2). If, for example, we want to generate lift-and-project cuts (Balas et al. 1993) (recall that cutting planes are a form of inference) and also perform reduced-cost fixing on our x variables, the inference statement in the search section of the model would look like *inference* = { lift-and-project, x :redcost }.

Due to space limitations, further details on SIMPL's implementation of search and inference are given in online Appendix A.1.

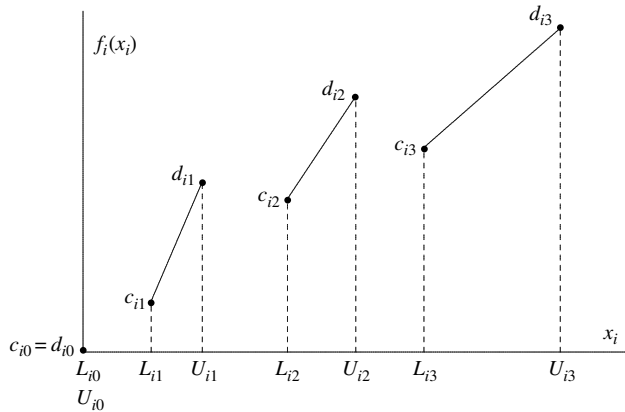
6. Example: Piecewise-Linear Functions

A simple production planning example with piecewise-linear functions illustrates integrated modeling as well as the search-infer-and-relax process. The approach taken here is similar to that of Refalo (1999) and Ottosson et al. (1999, 2002).

The objective is to manufacture several products at a plant of limited capacity C so as to maximize net income. Each product must be manufactured in one of several production modes (small scale, medium scale, large scale, etc.), and only a certain range of production quantities are possible for each mode. Thus, if x_i units of product i are manufactured in mode k , $x_i \in [L_{ik}, U_{ik}]$. The net income $f_i(x_i)$ derived from making product i is linear in each interval $[L_{ik}, U_{ik}]$, with $f_i(L_{ik}) = c_{ik}$ and $f_i(U_{ik}) = d_{ik}$. Thus, f_i is a continuous or semicontinuous piecewise-linear function (Figure 1):

$$f_i(x_i) = \frac{U_{ik} - x_i}{U_{ik} - L_{ik}} c_{ik} + \frac{x_i - L_{ik}}{U_{ik} - L_{ik}} d_{ik} \quad \text{if } x_i \in [L_{ik}, U_{ik}]. \quad (1)$$

Figure 1. A semicontinuous piecewise-linear function $f_i(x_i)$.



Making none of product i corresponds to mode $k = 0$, for which $[L_{i0}, U_{i0}] = [0, 0]$.

An integer programming model for this problem introduces 0-1 variables y_{ik} to indicate the production mode of each product. The functions f_i are modeled by assigning weights λ_{ij} , μ_{ik} to the endpoints of each interval k . The model is

$$\begin{aligned} \max \quad & \sum_{ik} \lambda_{ik} c_{ik} + \mu_{ik} d_{ik} \\ \sum_i x_i & \leq C, \\ x_i = \sum_k \lambda_{ik} L_{ik} + \mu_{ik} U_{ik}, \quad \sum_k \lambda_{ik} + \mu_{ik} & = 1 \quad \text{for all } i, \quad (2) \\ \sum_k y_{ik} & = 1 \quad \text{for all } i, \\ 0 \leq \lambda_{ik} \leq y_{ik}, \quad 0 \leq \mu_{ik} \leq y_{ik}, \quad y_{ik} \in \{0, 1\} & \quad \text{for all } i, k. \end{aligned}$$

If desired, one can identify $\lambda_{i0}, \mu_{i0}, \lambda_{i1}, \mu_{i1}, \dots$ as a specially ordered set of type 2 for each product i . However, specially ordered sets are not particularly relevant here because the adjacent pair of variables $\mu_{ik}, \lambda_{i, k+1}$ are never both positive for any k .

We formulate a model that directly informs the solver of the piecewise-linear nature of the costs. For each product i , the point (x_i, u_i) must lie on one of the line segments of the piecewise-continuous cost function. If the solver is aware of this fact, it can construct a tight linear relaxation by requiring (x_i, u_i) to lie in the convex hull of these line segments, thus resulting in a substantially faster solution.

This is accomplished by equipping the modeling language with metaconstraint *piecewise* to model continuous or semicontinuous piecewise-linear functions. A single piecewise constraint represents the constraints in (b) that correspond to a given i . The model is written as

$$\begin{aligned} \max \quad & \sum_i u_i \\ \sum_i x_i & \leq C, \quad (a) \quad (3) \\ \text{piecewise}(x_i, u_i, L_i, U_i, c_i, d_i) & \quad \text{for all } i. \quad (b) \end{aligned}$$

Here, L_i is an array containing L_{i0}, L_{i1}, \dots , and similarly for U_i, c_i , and d_i . Each piecewise constraint enforces $u_i = f_i(x_i)$.

In general, the solver has a library of metaconstraints that are appropriate to common modeling situations. Typically, some constraints are written individually, as is (a) above, whereas others are collected under one or more metaconstraints to simplify the model and allow the solver to exploit problem structure. It does so by applying inference methods to each metaconstraint, relaxing it, and branching in an intelligent way when it is not satisfied. In each case, the solver exploits the peculiar structure of the constraint.

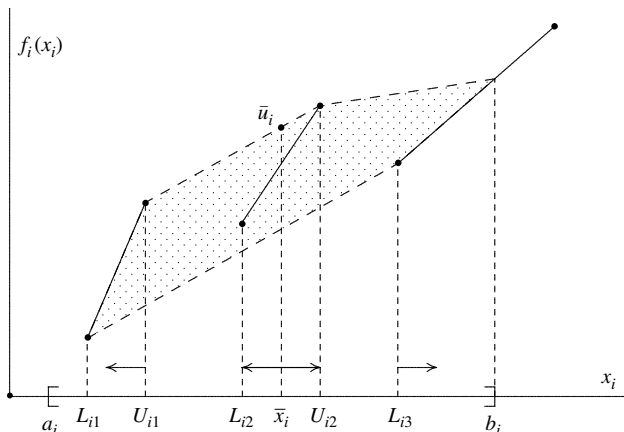
Let us suppose the solver is instructed to solve the production planning problem by branch and bound, which defines the *search* component of the algorithm. The search proceeds by enumerating restrictions of the problem, each one corresponding to a node of the search tree. At each node, the solver *infers* a domain $[a_i, b_i]$ for each variable x_i . Finally, the solver generates bounds for the branch-and-bound mechanism by solving a *relaxation* of the problem. It branches whenever a constraint is violated by the solution of the relaxation, and the nature of the branching is dictated by the constraint that is violated. If more than one constraint turns out to be violated, the solver will branch on the one with the highest priority, as specified by the user (see §C.1 in the online supplement for an example).

It is useful to examine these steps in more detail. At a given node of the search tree, the solver first applies inference methods to each constraint. Constraint (a) triggers a simple form of *interval propagation*. The upper bound b_i of each x_i 's domain is adjusted to become $\min\{b_i, C - \sum_{j \neq i} a_j\}$. Constraint (b) can also reduce the domain of x_i , as will be seen shortly. Domains reduced by one constraint can be cycled back through the other constraint for possible further reduction. As branching and propagation reduce the domains, the problem relaxation becomes progressively tighter until it is infeasible or its solution is feasible in the original problem.

The solver creates a relaxation at each node of the search tree by pooling relaxations of the various constraints. It relaxes each constraint in (b) by generating linear inequalities to describe the convex hull of the graph of each f_i , as illustrated in Figure 2. The fact that x_i is restricted to $[a_i, b_i]$ permits a tighter relaxation, as shown in the figure. Similar reasoning reduces the domain $[a_i, b_i]$ of x_i to $[L_{i1}, b_i]$. The linear constraint (a) also generates a linear relaxation, namely, itself. These relaxations, along with the domains, combine to form a linear relaxation of the entire problem:

$$\begin{aligned} \max \quad & \sum_i u_i \\ \sum_i x_i & \leq C, \quad (4) \\ \text{conv}(\text{piecewise}(x_i, u_i, L_i, U_i, c_i, d_i)) & \quad \text{for all } i, \\ a_i \leq x_i \leq b_i & \quad \text{for all } i, \end{aligned}$$

Figure 2. Convex hull relaxation (shaded area) of $f_i(x_i)$ when x_i has domain $[a_i, b_i]$.



where *conv* denotes the convex hull description just mentioned.

The solver next finds an optimal solution (\bar{x}_i, \bar{u}_i) of the relaxation (4) by calling a linear programming plug-in. This solution will necessarily satisfy (a), but it may violate (b) for some product i , for example, if \bar{x}_i is not a permissible value of x_i , or \bar{u}_i is not the correct value of $f_i(\bar{x}_i)$. The latter case is illustrated in Figure 2, where the search creates three branches by splitting the domain of x_i into three parts: $[L_{i2}, U_{i2}]$, everything below U_{i1} , and everything above L_{i3} . Note that in this instance the linear relaxation at all three branches will be exact, so that no further branching will be necessary.

The problem is therefore solved by combining ideas from three technologies: search by splitting intervals, from continuous global optimization; domain reduction, from constraint programming; and polyhedral relaxation, from integer programming.

Summary of Computational Results

SIMPL is much faster than the original hand-coded implementation of the integrated approach, which was comparable to state-of-the-art commercial MILP technology at the time (CPLEX). Although CPLEX has since improved by a factor of a thousand on the larger instances, SIMPL is about twice as fast as the most recent version of CPLEX on the original problem set (up to 100 jobs). SIMPL's advantage grows dramatically on larger instances (300 to 600 jobs). The search tree, compared that of CPLEX, is 1,000 to 8,000 times smaller, and the computation time 20 to 120 times less. For further details, see §C.1 in the online supplement.

7. Example: Variable Indices

A variable index is a versatile modeling device that is readily accommodated by a search-infer-and-relax solver. If an expression has the form u_y , where y is a variable, then y is a *variable index* or variable subscript. A simple product

configuration problem (Thorsteinsson and Ottosson 2001) illustrates how variable indices can be used in a model and processed by a solver.

The problem is to choose an optimal configuration of components for a product, such as a personal computer. For each component i , perhaps a memory chip or power supply, one must decide how many q_i to install and what type t_i to install. Only one type of each component may be used. The types correspond to different technical specifications, and each type k of component i supplies a certain amount a_{ijk} of attribute j . For example, a given type of memory chip might supply a certain amount of memory, generate a certain amount of heat, and consume a certain amount of power; in the last case, $a_{ijk} < 0$ to represent a negative supply. There are lower and upper bounds L_j, U_j on each attribute j . Thus, there may be a lower bound on total memory, an upper bound on heat generation, a lower bound of zero on net power supply, and so forth. Each unit of attribute j produced incurs a (possibly negative) penalty c_j .

A straightforward integer programming model introduces 0-1 variables x_{ik} to indicate when type k of component i is chosen. The total penalty is $\sum_j c_j v_j$, where v_j is the amount of attribute j produced. The quantity v_j is equal to $\sum_{ik} a_{ijk} q_{ik} x_{ik}$. Because this is a nonlinear expression, the variables q_i are disaggregated, so that q_{ik} becomes the number of units of type k of component i . The quantity v_j is now given by the linear expression $\sum_{ik} a_{ijk} q_{ik}$. A big- M constraint can be used to force q_{ij} to zero when $x_{ij} = 0$. The model becomes

$$\begin{aligned} \min \sum_j c_j v_j \\ v_j = \sum_{ik} a_{ijk} q_{ik}, \quad L_j \leq v_j \leq U_j \quad \text{for all } j, \\ q_{ik} \leq M_i x_{ik} \quad \text{for all } i, k, \\ \sum_k x_{ik} = 1 \quad \text{for all } i, \end{aligned} \tag{5}$$

where each x_{ij} is a 0-1 variable, each q_{ij} is integer, and M_i is an upper bound on q_i .

An integrated model uses the original notation t_i for the type of component i , without the need for 0-1 variables or disaggregation. The key is to permit t_i to appear as a subscript:

$$\begin{aligned} \min \sum_j c_j v_j \\ v_j = \sum_i q_i a_{ij t_i} \quad \text{for all } j, \end{aligned} \tag{6}$$

where the bounds L_j, U_j are reflected in the initial domain assigned to v_j .

The modeling system automatically decodes variably indexed expressions with the help of the *element* constraint, which is frequently used in CP modeling. In this case,

the variably indexed expressions occur in *indexed linear* expressions of the form

$$\sum_i q_i a_{ij t_i}, \quad (7)$$

where each q_i is an integer variable and each t_i a discrete variable. Each term $q_i a_{ij t_i}$ is automatically replaced with a new variable z_{ij} and the constraint

$$\text{element}(t_i, (q_i a_{ij1}, \dots, q_i a_{ijn}), z_{ij}) \quad (8)$$

This constraint in effect forces $z_{ij} = q_i a_{ij t_i}$. The solver can now apply a domain reduction or “filtering” algorithm to (8) and generate a relaxation for it. The filtering procedure is straightforward, if tedious, and is described in Appendix B of the online supplement.

As for the relaxation, note that since (8) implies a disjunction $\bigvee_{k \in D_i} (z_{ij} = a_{ijk} q_i)$, it can be given the standard convex hull relaxation for a disjunction, which in this case simplifies to

$$z_{ij} = \sum_{k \in D_i} a_{ijk} q_{ik}, \quad q_i = \sum_{k \in D_i} q_{ik}, \quad (9)$$

where $q_{ik} \geq 0$ are new variables.

If there is a lower bound L on expression (7), the relaxation used by Thorsteinnsson and Ottosson (2001) can be strengthened with integer knapsack cuts (and similarly if there is an upper bound). Because

$$\begin{aligned} \sum_i q_i a_{ij t_i} &= \sum_i \sum_{k \in D_i} a_{ijk} q_{ik} \leq \sum_i \max_{k \in D_i} \{a_{ijk}\} \sum_{k \in D_i} q_{ik} \\ &= \sum_i \max_{k \in D_i} \{a_{ijk}\} q_i, \end{aligned}$$

the lower bound L on (7) yields the valid inequality

$$\sum_i \max_{k \in D_i} \{a_{ijk}\} q_i \geq L. \quad (10)$$

Because the q_i s are general integer variables, integer knapsack cuts can be generated for (10).

Based on these ideas, the automatically generated relaxation of (6) becomes

$$\begin{aligned} \min \sum_j c_j v_j \\ v_j &= \sum_i \sum_{k \in D_i} a_{ijk} q_{ik}, \quad L_j \leq v_j \leq U_j \quad \text{for all } j, \\ q_i &= \sum_{k \in D_i} q_{ik}, \quad \underline{q}_i \leq q_i \leq \bar{q}_i \quad \text{for all } i, \end{aligned} \quad (11)$$

knapsack cuts for $\sum_i \max_{k \in D_i} \{a_{ijk}\} q_i \geq L_j$ and

$$\sum_i \min_{k \in D_i} \{a_{ijk}\} q_i \leq U_j \quad \text{for all } j,$$

$q_{ik} \geq 0$ for all i, k .

There is also an opportunity for *postrelaxation inference*, which in this case takes the form of reduced-cost variable fixing. Suppose that the best feasible solution found so far has value z^* , and let \hat{z} be the optimal value of (11). If $\hat{z} + r_{ik} \geq z^*$, where r_{ik} is the reduced cost of q_{ik} in the solution of (11), then k can be removed from the domain of t_i . In addition, if $r_{ik} > 0$, one can infer

$$\bar{q}_i \leq \max_{k \in D_i} \left\{ \left\lfloor \frac{z^* - \hat{z}}{r_{ik}} \right\rfloor \right\} \quad \text{for all } i.$$

Postrelaxation inference can take other forms as well, such as the generation of separating cuts.

The problem can be solved by branch and bound. In this case, we can start by branching on the domain constraints $t_i \in D_i$. Because t_i does not appear in the linear relaxation, it does not have a determinate value until it is fixed by branching. The domain constraint $t_i \in D_i$ is viewed as unsatisfied as long as t_i is undetermined. The search branches on $t_i \in D_i$ by splitting D_i into two subsets. Branching continues until all the D_i are singletons, or until at most one q_{ik} (for $k \in D_i$) is positive for each i . At that point we check if all q_i variables are integer and branch on q if necessary.

Summary of Computational Results

The original integrated approach was orders of magnitude faster than the contemporary CPLEX solver, both in terms of search nodes and time. SIMPL is even better than the original implementation and therefore achieves our main goal. It generates a search tree that is 10 times smaller on average. However, CPLEX has again become far better at solving this problem and now requires an average of 0.15 seconds, compared to 0.40 seconds for SIMPL, because it solves most problems at the root node. One might expect a commercial implementation of SIMPL to see comparable speedups as the technology and implementation improve. For further details, see §C.2 in the online supplement.

8. Example: Logic-Based Benders Decomposition

Nogood-based methods search the solution space by generating a *nogood* each time a candidate solution is examined. The nogood is a constraint that excludes the solution just examined, and perhaps other solutions that can be no better. The next solution enumerated must satisfy the nogoods generated so far. The search is exhaustive when the nogood set becomes infeasible.

Benders decomposition is a special type of a nogood-based method in which the nogoods are Benders cuts and the master problem contains all the nogoods generated so far. In classical Benders, the subproblem (slave problem) is a linear or nonlinear programming problem, and Benders cuts are obtained by solving its dual—or in the nonlinear case by deriving Lagrange multipliers. *Logic-based*

Benders generalizes this idea to an arbitrary subproblem by solving the *inference dual* of the subproblem (Hooker and Yan 1995; Hooker 1996, 1999).

A simple planning and scheduling problem illustrates the basic idea (Hooker 2000, Jain and Grossmann 2001, Bockmayr and Pizaruk 2003). A set of n jobs must be assigned to machines, and the jobs assigned to each machine must be scheduled subject to time windows. Job j has release time r_j , deadline d_j , and processing time p_{ij} on machine i . It costs c_{ij} to process job j on machine i . It generally costs more to run a job on a faster machine. The objective is to minimize processing cost. The MILP model used by Jain and Grossmann appears in §C.3 of the online supplement.

A hybrid model can be written with the *cumulative* meta-constraint, which is widely used in constraint programming for “cumulative” scheduling, in which several jobs can run simultaneously but subject to a resource constraint and time windows. Let t_j be the time at which job j starts processing and u_{ij} the rate at which job j consumes resources when it is running on machine i . The constraint

$$\text{cumulative}(t, p_i, u_i, U_i)$$

requires that the total rate at which resources are consumed on machine i always be less than or equal to U_i . Here $t = (t_1, \dots, t_n)$, $p_i = (p_{i1}, \dots, p_{in})$, and similarly for u_i .

In the present instance, jobs must run sequentially on each machine. Thus, each job j consumes resources at the rate $u_{ij} = 1$, and the resource limit is $U_i = 1$. Thus, if y_j is the machine assigned to job j , the problem can be written as

$$\begin{aligned} \min \sum_j c_{y_j j} \\ r_j \leq t_j \leq d_j - p_{y_j j} \quad \text{for all } j, \\ \text{cumulative}((t_j | y_j = i), (p_{ij} | y_j = i), e, 1) \quad \text{for all } i, \end{aligned} \quad (12)$$

where e is a vector of ones.

This model is adequate for small problems, but solution can be dramatically accelerated by decomposing the problem into an assignment portion to be solved by MILP and a subproblem to be solved by CP. The assignment portion becomes the Benders master problem, which allocates job j to machine i when $x_{ij} = 1$:

$$\begin{aligned} \min \sum_{ij} c_{ij} x_{ij} \\ \sum_i x_{ij} = 1 \quad \text{for all } j, \\ \text{relaxation of subproblem,} \\ \text{Benders cuts,} \\ x_{ij} \in \{0, 1\}. \end{aligned} \quad (13)$$

The solution \bar{x} of the master problem determines the assignment of jobs to machines. Once these assignments

are made, problem (12) separates into a scheduling feasibility problem on each machine i :

$$\begin{aligned} r_j \leq t_j \leq d_j - p_{\bar{y}_j j} \quad \text{for all } j, \\ \text{cumulative}((t_j | \bar{y}_j = i), (p_{ij} | \bar{y}_j = i), e, 1), \end{aligned} \quad (14)$$

where $\bar{y}_j = i$ when $\bar{x}_{ij} = 1$. If there is a feasible schedule for every machine, the problem is solved. If, however, the scheduling subproblem (14) is infeasible on some machine i , a Benders cut is generated to rule out the solution \bar{x} , perhaps along with other solutions that are known to be infeasible. The Benders cuts are added to the master problem, which is re-solved to obtain another assignment \bar{x} .

The simplest sort of Benders cut for machine i rules out assigning the same set of jobs to that machine again:

$$\sum_{j \in J_i} (1 - x_{ij}) \geq 1, \quad (15)$$

where $J_i = \{j | \bar{x}_{ij} = 1\}$. A stronger cut can be obtained, however, by deriving a smaller set J_i of jobs that are actually responsible for the infeasibility. This can be done by removing elements from J_i one at a time, and re-solving the subproblem, until the scheduling problem becomes feasible (Hooker 2005a). Another approach is to examine the proof of infeasibility in the subproblem and note which jobs actually play a role in the proof (Hooker 2005b). In CP, an infeasibility proof generally takes the form of edge-finding techniques for domain reduction, perhaps along with branching. Such a proof of infeasibility can be regarded as a solution of the subproblem’s *inference dual*. (In linear programming, the inference dual is the classical linear programming dual.) Logic-based Benders cuts can also be developed for planning and scheduling problems in which the subproblem is an optimization rather than a feasibility problem. This occurs, for example, in minimum makespan and minimum tardiness problems (Hooker 2004).

It is computationally useful to strengthen the master problem with a relaxation of the subproblem. The simplest relaxation requires that the processing times of jobs assigned to machine i fit between the earliest release time and latest deadline:

$$\sum_j p_{ij} x_{ij} \leq \max_j \{d_j\} - \min_j \{r_j\}. \quad (16)$$

A Benders method (as well as any nogood-based method) fits easily into the search-infer-and-relax framework. It solves a series of problem restrictions in the form of subproblems. The search is directed by the solution of a relaxation, which in this case is the master problem. The inference stage generates Benders cuts.

The decomposition is communicated to the solver by writing the model

$$\begin{aligned} \min \sum_{ij} c_{ij} x_{ij} & \quad (a) \\ \sum_i x_{ij} = 1 \quad \text{for all } j, & \quad (b) \\ (x_{ij} = 1) \Leftrightarrow (y_j = i) \quad \text{for all } i, j, & \quad (c) \quad (17) \\ r_j \leq t_j \leq d_j - p_{y_j j} \quad \text{for all } j, & \quad (d) \\ \text{cumulative}((t_j | y_j = i), (p_{ij} | \bar{y}_j = i), e, 1) & \quad (e) \\ & \quad \text{for all } i, \end{aligned}$$

where the domain of each x_{ij} is $\{0, 1\}$. Each constraint is associated with a relaxation parameter and an inference parameter. The relaxation parameters for constraints (b) and (d) will indicate that these constraints contribute to the MILP master relaxation of the problem. Note that (d) and (e) are part of the Benders subproblem. The relaxation parameter for (e) will add the inequalities (16) to the linear relaxation. The inference parameter for (e) will specify the type of Benders cuts to be generated. When the solver is instructed to use a Benders method, it automatically adds the Benders cuts to the relaxation. For more details on how these parameters are stated, see §C.3 in the online supplement.

Summary of Computational Results

SIMPL matches or exceeds the performance of the original integrated implementation. SIMPL is also several orders of magnitude faster than the most recent version of CPLEX, and the advantage increases rapidly with problem size. It solves, in a second or less, some instances that are intractable for CPLEX. For further details, see §C.3 in the online supplement.

9. Example: Global Optimization

We solved two sets of global problems. We first solved a few bilinear problems taken from Chapter 5 of Floudas et al. (1999). Bilinear problems are an important subclass of nonconvex quadratic programming problems whose applications include pooling and blending, separation sequencing, heat exchanger network design, and multicommodity network flow problems.

The global solver BARON already implements what is in many ways an integrated approach to solving a problem of this kind because it uses filtering techniques (known in the field as range reduction techniques) as well as relaxations. We implemented a similar approach in SIMPL and obtained performance that is roughly competitive with but somewhat slower than BARON's. This suggests that SIMPL could benefit from the primal heuristics and other techniques used in BARON. Further details may be found in §C.4 of the online supplement.

Our research objective in this paper, however, is to determine whether SIMPL can perform well on problems for which hand-coded integrated methods have been found to deliver substantial speedups. For this purpose we solved the mixed discrete/nonlinear truss structure design problems to which Bollapragada et al. (2001) applied an integrated method.

Truss structure optimization is one of the fundamental problems in engineering design. The goal is to find a minimum-cost placement and sizing of truss members (bars) to support a given load. The possible cross-sectional areas of the bars come from a discrete set of values that correspond to commonly manufactured sizes. Due to the presence of nonconvex physical constraints (i.e., Hooke's law), these problems are very difficult to solve. The nonlinear formulation can be converted into an MILP model by adding 0-1 and continuous variables (Ghattas and Grossmann 1991), which allows us to compare SIMPL's performance with that of both global and MILP solvers.

The structure to be built is represented as a network of nodes and arcs in two or three dimensions. The coordinates of the nodes are given, and structural bars are joined at the nodes. The problem consists of selecting the cross-sectional area of the bar to be placed along each arc, where the area is zero if no bar is placed. The objective is to minimize cost, which in our case is the volume of metal (aluminum) used in the bars. Each node has freedom of movement in a specified number of directions (degrees of freedom). Several possible loading conditions are anticipated, each of which is represented by a force applied to a subset of the nodes and along one or more degrees of freedom. There are limits on the displacement of each node along their degrees of freedom for each loading condition. In addition, the elongation, compression, and stress on each bar must lie within given limits.

This problem can be formulated as follows:

$$\begin{aligned} \min \sum_{i=1}^I c_i h_i A_i & \\ \sum_{i=1}^I b_{ij} s_{il} = p_{jl} \quad \text{for all } j, l, & \quad (a) \\ \sum_{j=1}^J b_{ij} d_{jl} = v_{il} \quad \text{for all } i, l, & \quad (b) \\ \frac{E_i}{h_i} A_i v_{il} = s_{il} \quad \text{for all } i, l, & \quad (c) \\ v_i^L \leq v_{il} \leq v_i^U \quad \text{for all } i, l, & \quad (d) \\ d_j^L \leq d_{jl} \leq d_j^U \quad \text{for all } j, l, & \quad (e) \\ \bigvee_{k=1}^{K_i} (A_i = A_{ik}) \quad \text{for all } i, & \quad (f) \end{aligned} \quad (18)$$

where I is the number of bars, J the number of degrees of freedom (summed over all nodes), L the number of loading

conditions, and K_i the number of discrete cross-sectional areas for bar i . Also, h_i is the length of bar i , A_{ik} is the k th discrete cross-sectional area of bar i , E_i is the modulus of elasticity of bar i , p_{jl} is the force imposed by loading condition l at degree of freedom j , b_{ij} is the cosine of the angle between bar i and degree of freedom j , and c_i is the cost per unit volume of bar i (typically the weight density). Finally, σ_i^L and σ_i^U are the minimum and maximum allowable stress in bar i , v_i^L and v_i^U are the limits on elongation/contraction of bar i , and d_j^L and d_j^U are the limits on displacement for degree of freedom j .

The variables are as follows. A_i is the cross-sectional area chosen for bar i , where the absence of bar i is represented by assigning a very small value to A_i ; s_{il} is the force in bar i due to loading condition l ; σ_{il} is the stress in bar i due to loading condition l ; v_{il} is the elongation (or contraction, if negative) of bar i due to loading condition l ; and d_{jl} is the node displacement along degree of freedom j for loading condition l . Constraints (a) represent equilibrium equations that balance the external loads with the forces induced in the bars; (b) are compatibility equations that relate the displacement of the nodes with the elongation of the bars; (c) represents Hooke's law, which relates the elongation or compression of a bar to the force applied to it (note that this constraint is nonlinear); and the disjunctive constraints (f) require that each area A_i take one of the discrete values A_{ik} .

If we add 0-1 variables y_{ik} that are equal to 1 when $A_i = A_{ik}$, (f) can be replaced by two constraints, $A_i = \sum_k A_{ik}y_{ik}$ and $\sum_k y_{ik} = 1$, for each i . This transforms (18) into a mixed-integer nonlinear programming model (MINLP), which can be solved by a global optimization solver like BARON. Furthermore, if we disaggregate the v_{il} variables, we can convert (18) to an MILP model by replacing v_{il} with $\sum_k v_{ikl}$ and replacing (c) with

$$\frac{E_i}{h_i} \sum_{k=1}^{K_i} A_{ik} v_{ikl} = s_{il} \quad \text{for all } i, l.$$

A disadvantage of the MILP model is the large number of variables. Bollapragada et al. (2001) show how to obtain a much smaller relaxation of a problem that has the same optimal value as (18). The resulting *quasi-relaxation* of (18) therefore provides a valid bound on the optimal value of (18).

The quasi-relaxation technique, generalized in Hooker (2005c), applies to any constraint of the form $g(x, y) \leq 0$, where g is semihomogeneous in x and concave in y , and where $x \in \mathbb{R}^n$ and y is a scalar. The function $g(x, y)$ is semihomogeneous in x when $g(\alpha x, y) \leq \alpha g(x, y)$ for all x , y and $\alpha \in [0, 1]$ and $g(0, y) = 0$ for all y . We also suppose there are bounds $x^L \leq x \leq x^U$ and $y_L \leq y \leq y_U$, and the

objective function involves variables in x . Then, a quasi-relaxation can be obtained by replacing $g(x, y) \leq 0$ with

$$\begin{aligned} g(x^1, y_L) + g(x^2, y_U) &\leq 0, \\ \alpha x^L &\leq x^1 \leq \alpha x^U, \\ (1 - \alpha)x^L &\leq x^2 \leq (1 - \alpha)x^U, \\ x &= x^1 + x^2. \end{aligned}$$

The bilinear constraints (c) in (18) have the form $g(x, y) \leq 0$ and satisfy the conditions for quasi-relaxation. The following is therefore a quasi-relaxation of (18):

$$\begin{aligned} \min \sum_{i=1}^I c_i h_i [A_i^L y_i + A_i^U (1 - y_i)] \\ \sum_{i=1}^I b_{ij} s_{il} &= p_{jl} \quad \text{for all } j, l, \\ \sum_{j=1}^J b_{ij} d_{jl} &= v_{i0l} + v_{i1l} \quad \text{for all } i, l, \\ \frac{E_i}{h_i} (A_i^L v_{i0l} + A_i^U v_{i1l}) &= s_{il} \quad \text{for all } i, l, \\ v_i^L y_i &\leq v_{i0l} \leq v_i^U y_i, \\ v_i^L (1 - y_i) &\leq v_{i1l} \leq v_i^U (1 - y_i) \quad \text{for all } i, l, \\ d_j^L &\leq d_{jl} \leq d_j^U \quad \text{for all } j, l, \\ 0 &\leq y_i \leq 1 \quad \text{for all } i. \end{aligned} \tag{19}$$

The quasi-relaxation technique is sufficiently general to justify a metaconstraint that represents constraints of the form $g(x, y) \leq 0$ satisfying the conditions for quasi-relaxation. However, because any bilinear function satisfies the condition, we invoke the quasi-relaxation simply by adding an additional relaxation option to the bilinear constraint introduced earlier.

Model (18) can be solved by branch and bound using (19) as the relaxation at each node. If $0 < y_i < 1$ in the optimal solution of (19) and $A_i^L \neq A_i^U$, let $A_i^* = A_i^L y_i + A_i^U (1 - y_i)$. We split the domain of A_i into two parts: the cross-sectional areas strictly smaller than A_i^* , and the cross-sectional areas greater than or equal to A_i^* . We branch first on the second part because it is more likely to lead to a feasible solution. Another property of the quasi-relaxation (19) is that whenever it is feasible, there exists a feasible solution in which both v_{i0l} and v_{i1l} have the same sign. Therefore, if v_{i0l} and v_{i1l} have opposite signs in the solution of (19), we can branch by introducing *logic cuts* of the form $v_{i0l}, v_{i1l} \geq 0$ (left branch) and $v_{i0l}, v_{i1l} \leq 0$ (right branch). These logic cuts are particularly useful in the presence of displacement bounds ($d_j^L > -\infty$ and $d_j^U < \infty$), and they are checked for violation before the check on y_i described above.

Summary of Computational Results

SIMPL replicates the performance of the hand-coded integrated method for this problem, becoming superior to it as the problem size increases from 10 to 108 bars. In addition, SIMPL solves the problem instances between two and seven times faster than the most recent version of CPLEX. The advantage over BARON is substantially greater. None of the methods solves a 200-bar problem to optimality, but after 24 hours of computation, SIMPL finds a better feasible solution than the original integrated implementation, whereas neither BARON nor CPLEX finds a feasible solution. For further details, see §C.5 of the online supplement.

10. Final Comments and Conclusions

In this paper, we describe a general-purpose integrated solver for optimization problems, to which we refer by the acronym SIMPL. It incorporates the philosophy that many traditional optimization techniques can be seen as special cases of a more general method, one that iterates a three-step procedure: solving relaxations, performing logical inferences, and intelligently enumerating problem restrictions. The main objective of SIMPL is to make the computational and modeling advantages of integrated problem solving conveniently available.

We tested SIMPL's modeling and solution capabilities on five types of optimization problems. We found that SIMPL (a) reproduces or exceeds the computational advantage of custom-coded integrated algorithms on four of these problems; (b) solves three of the problem classes faster than current state of the art, one of them by orders of magnitude, even though SIMPL is still an experimental code; (c) provides these advantages with modest effort on the part of the user because the integrated models are written in a concise and natural way; and (d) accommodates a wide range of problem types.

One may argue that it is unfair to compare SIMPL with an off-the-shelf commercial solver because the latter does not contain facilities to exploit problem structure in the way that SIMPL does. However, a major advantage of an integrated solver is precisely that it can exploit structure while remaining a general-purpose solver and providing the convenience of current commercial systems. SIMPL's constraint-based approach automatically performs the tedious job of integrating solution techniques while exploiting the complementary strengths of the various technologies it combines.

Our examples suggest that a SIMPL user must be more aware of the solution algorithm than an MILP user, but again this allows the solver to benefit from the user's understanding of problem structure. We anticipate that future development of SIMPL and related systems will allow them to presuppose less knowledge on the part of the average user to solve less difficult problems, while giving experts the power to solve harder problems within the same modeling framework. In addition, we plan to increase SIMPL's

functionality by increasing its library of metaconstraints, solver types, constraint relaxations, and search strategies, with the goal of accommodating the full spectrum of problems described in Table 1.

Those interested in reproducing our results can download a demo version of SIMPL from <http://moya.bus.miami.edu/~tallys/simpl.php>. The package includes all the problem instances used in our experiments.

11. Electronic Companion

An electronic companion to this paper is available as part of the online version that can be found at <http://or.journal.informs.org/>.

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A SIMPL's Implementation of Search and Inference

This section explains how SIMPL internally implements a model's search specification and its inference parameters.

A.1 Search

The main search loop inside SIMPL is implemented as shown below.

```
procedure Search(A)
  If  $A \neq \emptyset$  and stopping criteria not met
     $N := A.\text{getNextNode}()$ 
     $N.\text{explore}()$ 
     $A.\text{addNodes}(N.\text{generateRestrictions}())$ 
  Search(A)
```

Here, N is again the current problem restriction, and A is the current list of restrictions waiting to be processed. Depending on how A , N and their subroutines are defined, we can have different types of search, as mentioned in Section 1. The routine $N.\text{explore}()$ implements the infer-relax sequence. The routine $N.\text{generateRestrictions}()$ creates new restrictions, and $A.\text{addNodes}()$ adds the restrictions to A . Routine $A.\text{getNextNode}()$ implements a mechanism for selecting the next restriction, such as depth-first, breadth-first, or best-bound.

In tree search, N is the problem restriction that corresponds to the current node, and A is the set of open nodes. In local search, N is the restriction that defines the current neighborhood, and A is the singleton containing the restriction that defines the next neighborhood to be searched. In Benders decomposition, N is the current subproblem and A is the singleton containing the next subproblem to be solved. In the case of Benders, the role of $N.\text{explore}()$ is to infer Benders cuts from the current subproblem, add them to the master problem, and solve the master problem. $N.\text{generateRestrictions}()$ uses the solution of the master problem to create the next subproblem. In the sequel, we restrict our attention to branch-and-bound search.

A.2 Node Exploration

The behavior of $N.\text{explore}()$ for a branch-and-bound type of search is

1. Pre-relaxation inference
2. Repeat
3. Solve relaxations
4. Post-relaxation inference
5. Until (no changes) or (iteration limit)

The inference procedures in Steps 1 and 4 extract information from each relaxation in order to accelerate the search, as explained in Section A.4 below. The loop continues to execute as desired until the domains reach a fixed point.

A.3 Branching

SIMPL implements a tree search by branching on constraints. This scheme is considerably more powerful and generic than branching on variables alone. If branching is needed, it is because some constraint of the problem is violated, and that constraint should “know” how to branch as a result. This knowledge is embedded in the *branching module* associated with the constraint. For example, if a variable $x \in \{0, 1\}$ has a fractional value in the current LP, its indomain constraint I_x is violated. The branching module of I_x outputs the constraints $x \in \{0\}$ and $x \in \{1\}$, meaning that two subproblems should be created by the inclusion of those two new constraints. Traditional branching on a variable x can therefore be interpreted as a special case of branching on a constraint. In general, a branching module returns a sequence of constraint *sets* C_1, \dots, C_k . Each C_i defines a subproblem at a successor node when it is merged with the current problem. There is no restriction on the type of constraints appearing in C_i .

Clearly, there may be more than one constraint violated by the solution of the current set of problem relaxations. A *selection module* is responsible for selecting, from a given set of constraints, the one on which to branch next. Some possible criteria for selection are picking the first constraint found to be violated or the one with the largest degree of violation.

A.4 Inference

We now take a closer look at the inference steps of the node exploration loop in Section A.2. In step 1 (pre-relaxation inference), one may have domain reductions or the generation of new implied constraints (Hooker and Osorio 1999), which may have been triggered by the latest branching decisions. If the model includes a set of propositional logic formulas, this step can also execute some form of resolution algorithm to infer new resolvents. In step 4 (post-relaxation inference), other types of inference may take place, such as fixing variables by reduced cost or the generation of cutting planes. After that, it is possible to implement some kind of primal heuristic or to try extending the current solution to a feasible solution in a more formal way, as advocated in Sect. 9.1.3 of Hooker (2000).

Since post-relaxation domain reductions are associated with particular relaxations, the reduced domains that result are likely to differ across relaxations. Therefore, at the end of the inference steps, a synchronization step must be executed to propagate domain reductions across different relaxations. This is done in the algorithm below.

1. $V := \emptyset$
2. For each problem relaxation r
3. $V_r :=$ variables with changed domains in r
4. $V := V \cup V_r$
5. For each $v \in V_r$
6. $D_v := D_v \cap D_v^r$
7. For each $v \in V$
8. Post constraint $v \in D_v$

In step 6, D_v^r denotes the domain of v inside relaxation r , and D_v works as a temporary domain for variable v , where changes are centralized. The initial value of D_v is the current domain of variable v . By implementing the changes in the domains via the addition of indomain constraints (step 8), those changes will be transparently undone when the search moves to a different part of the enumeration tree. Similarly, those changes are guaranteed to be redone if the search returns to descendants of the current node at a later stage.

B Filtering for the Element Constraint

For a given j , filtering for (8) is straightforward. If z_{ij} 's domain is $[z_{ij}, \bar{z}_{ij}]$, t_i 's domain is D_{t_i} , and q_i 's domain is $\{q_i, q_i + 1, \dots, \bar{q}_i\}$ at any point in the search, then the reduced domains $[z'_{ij}, \bar{z}'_{ij}]$, D'_{t_i} , and $\{q'_i, \dots, \bar{q}'_i\}$ are given by

$$\begin{aligned}
 z'_{ij} &= \max \left\{ z_{ij}, \min_k \left\{ a_{ijk} q_i, a_{ijk} \bar{q}_i \right\} \right\}, & \bar{z}'_{ij} &= \min \left\{ \bar{z}_{ij}, \max_k \left\{ a_{ijk} q_i, a_{ijk} \bar{q}_i \right\} \right\}, \\
 D'_{t_i} &= D_{t_i} \cap \left\{ k \mid [z'_{ij}, \bar{z}'_{ij}] \cap \left[\min\{a_{ijk} q_i, a_{ijk} \bar{q}_i\}, \max\{a_{ijk} q_i, a_{ijk} \bar{q}_i\} \right] \neq \emptyset \right\} \\
 q'_i &= \max \left\{ q_i, \min_k \left\{ \left\lceil \frac{z'_{ij}}{a_{ijk}} \right\rceil, \left\lfloor \frac{\bar{z}'_{ij}}{a_{ijk}} \right\rfloor \right\} \right\}, & \bar{q}'_i &= \min \left\{ \bar{q}_i, \max_k \left\{ \left\lfloor \frac{z'_{ij}}{a_{ijk}} \right\rfloor, \left\lceil \frac{\bar{z}'_{ij}}{a_{ijk}} \right\rceil \right\} \right\}
 \end{aligned}$$

C Models and Computational Experiments

We formulate integrated models for the five examples described in sections 6 through 9 of the main paper. We solve them with SIMPL version 0.08.22 and compare its performance to the reported performance of previously implemented integrated methods. When applicable, we also formulate and solve the corresponding MILP and global optimization models with CPLEX 11.0.0 (CPLEX 11, for short) and BARON version 7.2.5, respectively. We also use CPLEX 9.0.0 (CPLEX 9, for short) for comparison purposes in some of our experiments.

We report both the number of search nodes and the computation time. Since SIMPL is still a research code, the node count may be a better indication of performance at the present stage of development. The amount of time SIMPL spends per node can be reduced by optimizing the code, whereas the node count is more a function of the underlying algorithm. Even though the focus of this paper is not to show that integrated approaches can outperform traditional optimization approaches, we note that SIMPL requires less time, as well as fewer nodes, than current technology on three of the five problem classes studied here.

Unless indicated otherwise, all the experiments reported in this section have been run on a Pentium 4, 3.7 GHz with 4GB of RAM, running Ubuntu 8.04 with Linux kernel 2.6.24-19. We used CPLEX 11 as the LP solver and ECLⁱPS^e 5.8.103 as the CP solver in SIMPL. For simplicity, when showing the SIMPL code of each model we omit the data and variable declaration statements.

C.1 Production Planning

The SIMPL code that corresponds to the integrated model (3) of the production planning problem is shown below.

```

01. OBJECTIVE
02.   maximize sum i of u[i]
03. CONSTRAINTS
04.   capacity means {
05.     sum i of x[i] <= C
06.     relaxation = { lp, cp } }
07.   piecewisectr means {
08.     piecewise(x[i],u[i],L[i],U[i],c[i],d[i]) forall i
09.     relaxation = { lp, cp } }
10. SEARCH
11.   type = { bb:bestdive }
12.   branching = { piecewisectr:most }

```

Lines 06 and 09 of the above code tell SIMPL that those constraints should be posted to both the linear programming (**lp**) and constraint programming (**cp**) relaxations/solvers. Recall that the linear programming relaxation of the i^{th} piecewise constraint is the collection of inequalities on x_i and u_i that define the convex hull of their feasible values in the current state of the search. Line 11 indicates that we use branch-and-bound (**bb**), select the current active node with the best lower bound, and dive from it until we reach a leaf node (keyword **bestdive**). Finally, in line 12 we say that the branching strategy is to branch on the piecewise constraint with the largest degree of violation (keyword **most**). The amount of violation is calculated by measuring how far the LP relaxation values of x_i and u_i are from the closest linear piece of the function. That is, we measure the rectilinear distance between the point (x_i, u_i) and the current convex hull of piecewise.

We ran the integrated model over 28 randomly generated instances with the number of products n ranging from 5 to 600. Similar problems were solved by Ottosson, Thorsteinsson and Hooker (1999, 2002), but their models had a few additional constraints and their piecewise linear functions were continuous. We did not use their original instances because we wanted to experiment with a more challenging version of the problem (including discontinuous functions and more products). The performance of their implementation was comparable to the best MILP results at that time, whereas our implementation is superior to the MILP approach. In all instances, products have the same cost structure with five production modes. For the purpose of reducing symmetry, the model also includes constraints of the form $x_i \leq x_{i+1}$ for all $i \in \{1, \dots, n-1\}$.

The number of search nodes and CPU time (in seconds) required to solve the instances to optimality are shown in Table 2. For reference, we also include results obtained with the pure MILP model (2) with the above symmetry breaking constraints. As the number of products increases, the number of search nodes required by the integrated approach can be 1000 to 8000 times smaller

than the number of nodes required by the MILP approach. This problem used to be considerably more challenging for previous versions of CPLEX, as shown by the columns under "MILP (CPLEX 9)". Nevertheless, it is also possible to find larger instances that are challenging for CPLEX 11. If we change the LP solver in SIMPL from CPLEX 11 to CPLEX 9, the numbers under the SIMPL columns in Table 2 do not change significantly. This indicates that the integrated models solved with SIMPL are more robust than the MILP models.

C.2 Product Configuration

The product configuration problem of Section 7 can be coded in SIMPL as follows.

```

01. OBJECTIVE
02.   minimize sum j of c[j]*v[j]
03. CONSTRAINTS
04.   usage means {
05.     v[j] = sum i of q[i]*a[i][j][t[i]] forall j
06.     relaxation = { lp, cp }
07.     inference = { knapsack } }
08.   quantities means {
09.     q[1] >= 1 => q[2] = 0
10.     relaxation = { lp, cp } }
11.   types means {
12.     t[1] = 1 => t[2] in {1,2}
13.     t[3] = 1 => (t[4] in {1,3} and t[5] in {1,3,4,6} and t[6] = 3 and
                    t[7] in {1,2})
14.     relaxation = { lp, cp } }
15. SEARCH
16.   type = { bb:bestdive }
17.   branching = { quantities, t:most, q:least:triple, types:most }
18.   inference = { q:redcost }

```

For our computational experiments, we used the ten hardest problem instances proposed by Thorsteinsson and Ottosson (2001), which have 26 components, up to 30 types per component, and 8 attributes. According to Thorsteinsson and Ottosson (2001), these instances were generated to "closely resemble" real-world instances. In addition, there are a few extra logical constraints on the q_i and t_i variables, which are implemented in lines 08 through 14 above. These constraints are also added to the MILP model (5).

For the `usage` constraints, note the variable subscript notation in line 05 and the statement that tells it to infer integer knapsack cuts (as described in Section 7) in line 07 (this would be the default behavior anyway). We define our branching strategy in line 17 as follows: we first try to branch on the q -implication (`quantities`), then on the indomain constraints of the t variables (most violated first), followed by the indomain constraints on the q variables (least violated first), and finally on the implications on the t variables (`types`, most violated first). The indomain constraint of a t_i variable is violated if its domain is not a singleton and two or more of the corresponding q_{ik} variables have a positive value in the LP relaxation (see Section 7 for the relationship between t_i and q_{ik}). When the element constraint (8) is relaxed, the q_{ik} variables are created and variable

q_i is marked as a variable that decomposes into those q_{ik} variables (see the convex hull relaxation (9) in Section 7). In addition, variable t_i is marked as the indexing variable responsible for that decomposition and it saves a pointer to q_i and the list of q_{ik} variables. Hence, when the indomain constraint of t_i is checked for violation as described above, it knows which q_{ik} variables to look at. The keyword `triple` that appears after `q:least` in line 17 indicates that we branch on q_i as suggested by Thorsteinsson and Ottosson (2001): let \bar{q}_i be the closest integer to the fractional value of q_i in the current solution of the LP relaxation; we create up to three descendants of the current node by adding each of the following constraints in turn (if possible): $q_i = \bar{q}_i$, $q_i \leq \bar{q}_i - 1$ and $q_i \geq \bar{q}_i + 1$. Finally, the post-relaxation inference using reduced costs is turned on for the q variables in line 18.

The number of search nodes and CPU time (in seconds) required to solve each of the ten instances to optimality are shown in Table 3. This problem has become easy for recent versions of CPLEX, but Thorsteinsson and Ottosson’s (2001) original implementation was orders of magnitude better than the CPLEX of the time, both in terms of number of nodes and time. SIMPL’s search tree is, on average, about 10 times smaller than Thorsteinsson and Ottosson (2001) search tree on the same problem instances. We therefore achieved our goal of replicating or improving on the performance of the original hand-coded integrated method. When we compare SIMPL’s performance to that of the most recent version of CPLEX, it ranges from comparable to an order of magnitude slower. This is perhaps because SIMPL’s current implementation does not include strong cutting plane algorithms, which make a difference at the root node for this problem. The commercialization of MILP solvers has resulted in dramatic speedups, due to the resources invested in software engineering. One might similarly expect commercialization of integrated solvers to result in significant improvements. Computational comparisons between a research implementation of integrated methods and a commercial implementation of MILP should be viewed with this in mind.

C.3 Parallel Machine Scheduling

We first describe the MILP model used by Jain and Grossmann (2001). Let the binary variable x_{ij} be one if job j is assigned to machine i , and let the binary variable $y_{jj'}$ be one if both j and j' are assigned to the same machine and j finishes before j' starts. In addition, let t_j be the time at

which job j starts. The MILP model is as follows.

$$\min \sum_{ij} c_{ij} x_{ij}$$

$$r_j \leq t_j \leq d_j - \sum_i p_{ij} x_{ij}, \text{ all } j \quad (\text{a})$$

$$\sum_i x_{ij} = 1, \text{ all } j \quad (\text{b})$$

$$y_{jj'} + y_{j'j} \leq 1, \text{ all } j' > j \quad (\text{c})$$

$$y_{jj'} + y_{j'j} + x_{ij} + x_{i'j'} \leq 2, \text{ all } j' > j, i' \neq i \quad (\text{d})$$

$$y_{jj'} + y_{j'j} \geq x_{ij} + x_{i'j'} - 1, \text{ all } j' > j, i \quad (\text{e})$$

$$t_{j'} \geq t_j + \sum_i p_{ij} x_{ij} - \mathcal{U}(1 - y_{jj'}), \text{ all } j' \neq j \quad (\text{f})$$

$$\sum_j p_{ij} x_{ij} \leq \max_j \{d_j\} - \min_j \{r_j\}, \text{ all } i \quad (\text{g})$$

$$x_{ij} \in \{0, 1\}, y_{jj'} \in \{0, 1\} \text{ for all } j' \neq j$$

Constraint (a) defines lower and upper bounds on the start time of job j , and (b) makes sure every job is assigned to some machine. Constraints (c) and (d) are logical cuts which significantly reduce solution time. Constraint (e) defines a logical relationship between the assignment (x) and sequencing (y) variables, and (f) ensures start times are consistent with the value of the sequencing variables y ($\mathcal{U} = \sum_j \max_i \{p_{ij}\}$). Finally, (g) are valid cuts that tighten the linear relaxation of the problem. There is also a continuous-time MILP model suggested by Türkay and Grossmann (1996), but computational testing indicates that it is much harder to solve than the above (Hooker 2004).

We now describe the SIMPL code to implement the Benders decomposition approach of Section 8 (model (17)) when the master problem is given by (13), with (14) as the subproblem, and (15) as the Benders cuts.

```

01. OBJECTIVE
02.   min sum i,j of c[i][j] * x[i][j];
03. CONSTRAINTS
04.   assign means {
05.     sum i of x[i][j] = 1 forall j;
06.     relaxation = { ip:master } }
07.   xy means {
08.     x[i][j] = 1 <=> y[j] = i forall i, j;
09.     relaxation = { cp } }
10.   tbounds means {
11.     r[j] <= t[j] forall j;
12.     t[j] <= d[j] - p[y[j]][j] forall j;
13.     relaxation = { ip:master, cp } }
14.   machinecap means {

```

```

15.    cumulative({ t[j], p[i][j], 1 } forall j | x[i][j] = 1, 1) forall i;
16.    relaxation = { cp:subproblem, ip:master }
17.    inference = { feasibility } }
18. SEARCH
19.    type = { benders }

```

The keywords `ip:master` that appear in the relaxation statements in lines 06, 13 and 16 indicate that those constraints are to be relaxed into an Integer Programming relaxation, which will constitute the master problem. Constraints that have the word `cp` in their relaxation statements will be posted to a CP relaxation and are common to all Benders subproblems. This is true for the `xy` and `tbounds` constraints. For the `machinecap` constraints, the keywords `cp:subproblem` in line 16, together with the `forall i` statement in line 15, indicate that, for each i , there will be a different CP subproblem containing the corresponding cumulative constraint, in addition to the common constraints mentioned above. Finally, we tell SIMPL that the cumulative constraints should generate the feasibility-type Benders cuts (15) in line 17. Hence, when a subproblem turns out to be infeasible, its cumulative constraint, which is aware of the jobs it was assigned to handle, has all the information it needs to infer a cut that looks like (15).

For our computational experiments, we used the instances proposed by Jain and Grossmann (2001) and, additionally, we followed their procedure to create three new instances with more than 20 jobs. These are the last instance in Table 4 and the last two instances in Table 5. As was the case in Section C.2, Jain and Grossmann (2001)'s instances were also generated to resemble real-world instances.

Jain and Grossmann's original implementation only needed a small number of master iterations and a few seconds to solve these instances. The results we obtain with SIMPL match this performance. Although state-of-the-art MILP solvers have considerably improved since Jain and Grossmann's results were published, the largest instances are still intractable with their pure MILP model. In addition to being orders of magnitude faster in solving the smallest problems, the integrated Benders approach can easily tackle larger instances as well. As noted by Jain and Grossmann (2001), when processing times are shorter the problem tends to become easier, and we report the results for shorter processing times in Table 5. Even in this case, the MILP model is still much worse than the integrated Benders approach as the problem size grows. For instance, after more than 27 million search nodes and a time limit of 48 hours of CPU time, the MILP solver had found an integer solution with value 156 to the problem with 22 jobs and 5 machines, whereas the optimal solution has value 155. As for the largest problem (25 jobs and 5 machines), the MILP solver ran out of memory after 19 hours of CPU time and over 5 million search nodes, having found a solution of value 182 (the optimal value is 179). It is worth mentioning that the MILP formulations of these two problems only have 594 and 750 variables, respectively.

C.4 Pooling, Distillation and Heat Exchanger Networks

We consider the following bilinear model from Floudas et al. (1999), which is often used for pooling, distillation, and heat exchanger network problems:

$$\begin{aligned}
 \min \quad & x^T A_0 y + c_0^T x + d_0^T y \\
 & x^T A_i y + c_i^T x + d_i^T y \leq b_i, \quad i = 1, \dots, p \\
 & x^T A_i y + c_i^T x + d_i^T y = b_i, \quad i = p + 1, \dots, p + q \\
 & x \in \mathbb{R}^n, \quad y \in \mathbb{R}^m
 \end{aligned} \tag{1}$$

where x and y are n - and m -dimensional variable vectors, respectively.

SIMPL replaces a nonlinear term of the form $x_i y_j$ with a new variable z_{ij} and a metaconstraint $\text{bilinear}(x_i, y_j, z_{ij})$. This constraint enforces $z_{ij} = x_i y_j$ and creates relaxations that are automatically updated. The CP relaxation is the constraint itself, because CP solvers can propagate bilinear constraints. A well-known linear relaxation is

$$\begin{aligned}
 L_j x_i + L_i y_j - L_i L_j &\leq z_{ij} \leq L_j x_i + U_i y_j - U_i L_j \\
 U_j x_i + U_i y_j - U_i U_j &\leq z_{ij} \leq U_j x_i + L_i y_j - L_i U_j
 \end{aligned} \tag{2}$$

where $[L_i, U_i]$ and $[L_j, U_j]$ are the current bounds on x_i and y_j , respectively. The search branches on $\text{bilinear}(x_i, y_j, z_{ij})$ if $x_i y_j \neq z_{ij}$ when the variables are replaced by their values in the solution of the current relaxation. Branching splits the domains of x_i and/or y_j , depending on a number of conditions.

To demonstrate SIMPL's ability to solve the bilinear global optimization problems, we selected the 6 bilinear problems from chapter 5 of Floudas et al. (1999) that BARON was able to solve in less than 24 hours. Problem names correspond to section numbers in that chapter. Problems 5.2.2.1, 5.2.2.2, 5.2.2.3, and 5.2.4 are pooling problems; problem 5.3.2 is a distillation problem; and problem 5.4.2 is a heat exchanger network problem. For illustration purposes, we describe a SIMPL model that could be used to model problem 5.2.2.1 (case 1 of section 5.2.2 in Floudas et al. 1999). Models for the other problems would be very similar to this one.

```

01. OBJECTIVE
02.   max 9*x + 15*y - 6*A - 16*B - 10*(Cx + Cy);
03. CONSTRAINTS
04.   flow means {
05.     Px + Py - A - B = 0;
06.     x - Px - Cx = 0;
07.     y - Py - Cy = 0;
08.     relaxation = { lp, cp } }
09.   pooling means {
10.     p*Px + 2*Cx - 2.5*x <= 0;
11.     p*Py + 2*Cy - 1.5*y <= 0;
12.     p*Px + p*Py - 3*A - B = 0;
13.     relaxation = { lp, cp } }
14. SEARCH

```

```

15.  type = { bb:bestdive }
16.  branching = { pooling:most }
17.  inference = { redcost }

```

If a non-linear programming solver is linked to SIMPL, line 13 could be changed to `relaxation = { lp, cp, nlp }`. Because the CP solver can handle non-linear constraints directly, they are posted to the CP relaxation as such, and ECLiPSe takes care of doing the proper bounds propagation (using BARON’s terminology, this type of inference would be called *feasibility-based range reduction*). Internally, the constraint in line 10 would be transformed into $ZpPx + 2*Cx - 2.5*x \leq 0$ and `bilinear(p,Px,ZpPx)`; the constraint in line 11 would be transformed into $ZpPy + 2*Cy - 1.5*y \leq 0$ and `bilinear(p,Py,ZpPy)`; and the constraint in line 12 would be transformed into the linear constraint $ZpPx + ZpPy - 3*A - B = 0$, because its bilinear terms have appeared before and there is no need for additional `bilinear` constraints. Branching is done on the most violated of the bilinear metaconstraints, where the violation is measured by $|x_i y_j - z_{ij}|$. Line 17 tells SIMPL to perform domain reduction based on reduced costs for all variables (using BARON’s terminology, this type of inference would be called *optimality-based range reduction*).

Computational results appear in Table 6. For comparison purposes, we solved these problems with both SIMPL and BARON version 7.2.5 (Tawarmalani and Sahinidis 2004). Because we ran BARON on another machine (an IBM workstation with two 3.2 GHz Intel Xeon processors and 2.5 GB of RAM), the times reported in the BARON column of Table 6 cannot be directly compared with the times reported for SIMPL. Even though BARON is a much more mature, stable and advanced global optimization solver than SIMPL currently is, the results of Table 6 help to demonstrate that SIMPL’s framework can also accommodate global optimization problems. As noted in Section 9, what BARON does when solving a global optimization problem, known as *Branch and Reduce*, can be interpreted as a special case of SIMPL’s search-infer-relax paradigm.

SIMPL’s implementation of global optimization still has a lot of room for improvement. In addition to better memory management, these improvements include the use of more powerful inference mechanisms (e.g. using lagrangian multipliers), support for other types of non-linear constraints and, most importantly, strong pre-processing techniques and a local search mechanism that help find good solutions early in the search, such as what BARON does. The next step is to link a standalone non-linear solver to SIMPL, such as KNITRO (Byrd, Nocedal, and Waltz 2006) or MINOS (Murtagh and Saunders 1983), which would increase the range of global optimization problems that it can solve.

C.5 Truss Structure Design

We now describe the SIMPL model for (18). Note that constraints (d), (e) and (f) are part of the variable declarations.

```

01. OBJECTIVE
02.  maximize sum i of c[i]*h[i]*A[i]
03. CONSTRAINTS
04.  equilibrium means {
05.    sum i of b[i,j]*s[i,1] = p[j,1] forall j,1
06.    relaxation = { lp } }
07.  compatibility means {

```

```

08.    sum j of b[i,j]*d[j,1] = v[i,1] forall i,1
09.    relaxation = { lp } }
10.    hooke means {
11.    E[i]/h[i]*A[i]*v[i,1] = s[i,1] forall i
12.    relaxation = { lp:quasi } }
13. SEARCH
14.    type = { bb:bestdive }
15.    branching = { hooke:first:quasicut, A:splitup }

```

This model contains a few novel constructs. Internally, each constraint in line 11 will be transformed into two constraints: $E[i]/h[i]*z[i,1] = s[i,1]$ and $\text{bilinear}(A[i],v[i,1],z[i,1])$. The `lp:quasi` statement creates a quasi-relaxation of the `bilinear` constraint and sends it to the LP solver. This relaxation is volatile (i.e. it is updated whenever the lower and/or upper bounds on A_i change) and consists of

$$\begin{aligned}
v_{i\ell} &= v_{i0\ell} + v_{i1\ell} \\
A_i &= A_i^L y_i + A_i^U (1 - y_i) \\
z_{i\ell} &= A_i^L v_{i0\ell} + A_i^U v_{i1\ell} \\
v_i^L y_i &\leq v_{i0\ell} \leq v_i^U y_i \\
v_i^L (1 - y_i) &\leq v_{i1\ell} \leq v_i^U (1 - y_i) \\
y_i &\in [0, 1]
\end{aligned}$$

We first branch by looking for the first violation of the logic cuts (`hooke:first:quasicut`), and then by choosing the first A_i that violates its indomain constraint. Following a recommendation by Bollapragada, Ghattas and Hooker (2001), we turn off logic cuts when the problem has no displacement bounds (i.e. when $d_j^L = -\infty$ and $d_j^U = \infty$). When no logic cuts are found/enabled, branching is performed on the A_i variables. The domain of the chosen A_i is split at the current fractional value and we branch first on the upper (right) piece (`splitup`) (see Section 9). The problem also includes the concept of equivalence classes (or linking groups), which are subsets of bars that are supposed to have the same cross-sectional area. The choice of the A_i variable on which to branch can be prioritized to scan the bars in non-increasing order of linking group size. To do this, we can modify the `A:splitup` statement in line 15 to `A:most(1,LinkSize):splitup`. Here, `LinkSize` is a vector of numbers (with the same dimension as the `A` vector). The number 1 indicates that the values in this vector are to be used as the first sorting criterion which, as a consequence, makes the violation amount the second sorting criterion. This statement generalizes to a larger number of criteria if we were to write, for example, `A:most(1,C1,2,C2,4,C4)`. In this last example, the `A` vector will be sorted with `C1` as the first criterion, `C2` as the second criterion, the original violation as the third criterion (because the number 3 was omitted), and `C4` as the fourth criterion, in a lexicographical fashion.

For our computational experiments we used 12 instances from the literature (the same ones used by Bollapragada, Ghattas and Hooker 2001). Eleven of them come from Venkayya (1971), and one of them from Cai and Theirauf (1993). Table 7 shows the number of search nodes and CPU time (in seconds) required to solve these problems with each of the following four approaches: solving the MILP version of model (18) with CPLEX 11; solving the MINLP version of model (18) with

BARON; using the original implementation of Bollapragada, Ghattas and Hooker (2001) (referred to as BGH); and replicating the BGH approach with SIMPL. As in Section C.4, BARON was run on an IBM workstation with two 3.2 GHz Intel Xeon processors and 2.5 GB of RAM. The other three algorithms were run on the same machine as the other experiments in this paper. All runs had a time limit of 24 CPU hours.

As the problem size increases, the standard global optimization approach (BARON column) does not scale well and time becomes a factor. The MILP model solved by CPLEX tends to get too large as the number of bars increases, and the solution time grows more quickly than in the integrated approaches. It is worth noting that CPLEX found a solution of value 5096.99 for instance 3, whereas BARON, BGH and SIMPL all found a solution of value 5156.64 (if we use CPLEX version 9 instead of 11, the “optimal” solution it returns for instance 3 has value 5037.4). Both BGH and SIMPL behave very similarly (as expected), with SIMPL having some advantage on the larger instances. Surprisingly, even though SIMPL’s underlying code has considerably more overhead than the BGH code (which was solely written for the purpose of solving the truss design problem), SIMPL is only about two and a half times slower than BGH (on average, over the 12 instances) in terms of number of nodes processed per second.

For the 200-bar instance, in particular, neither BARON nor CPLEX were able to find a single feasible solution within 24 hours of CPU time. Both BGH and SIMPL found a feasible (but not provably optimal) solution in less than 24 hours, and the solution found by SIMPL (32700.25) was slightly better than the one found by BGH (32747.58). Both codes processed about 1 node per second in this particular instance.

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Table 2: Production planning: search nodes and CPU time.

Number of Products	MILP (CPLEX 9)		SIMPL (CPLEX 9)		MILP (CPLEX 11)		SIMPL (CPLEX 11)	
	Nodes	Time (s)	Nodes	Time (s)	Nodes	Time (s)	Nodes	Time (s)
5	93	0.03	16	0.02	45	0.03	20	0.02
10	423	0.12	50	0.05	198	0.08	41	0.06
15	1,321	0.34	36	0.08	265	0.19	51	0.10
20	4,573	1.14	67	0.15	652	0.64	38	0.14
25	5,105	2.43	44	0.18	660	0.78	47	0.20
30	4,504	2.10	79	0.39	375	0.44	61	0.34
35	6,089	3.30	34	0.28	550	0.79	34	0.29
40	7,973	4.06	68	0.44	592	1.29	69	0.44
45	23,414	14.72	39	0.36	746	1.44	34	0.41
50	18,795	9.45	44	0.42	677	2.39	45	0.43
55	46,349	34.60	48	0.52	679	2.14	40	0.50
60	99,606	55.07	141	1.33	767	1.78	99	1.14
65	43,759	39.85	92	1.53	784	2.86	101	1.42
70	103,470	98.77	41	0.74	899	2.56	54	0.82
75	76,540	84.54	106	2.10	698	3.61	84	1.79
80	78,479	60.82	47	1.05	865	4.55	59	1.25
85	143,457	185.11	197	2.94	1043	6.07	146	2.91
90	315,456	421.50	66	1.60	1043	6.22	69	1.69
95	345,724	503.95	164	3.36	1174	6.69	104	1.98
100	2,247,538	4,458.65	103	2.03	1229	4.40	137	2.75
300a	10,164	81.74	85	25.88	101,756	375.63	73	19.13
300b	43,242	700.78	111	41.09	128,333	372.39	58	18.74
400a	18,234	101.30	67	22.80	135,641	563.56	41	22.89
400b	44,941	221.80	58	26.55	160,972	659.96	72	31.77
500a	4,019	117.45	340	299.17	113,162	824.11	90	69.30
500b	10,862	307.60	295	64.64	186,952	1,068.97	81	64.21
600a	363,740	3,514.83	457	161.18	646,997	4,509.11	74	39.18
600b	7,732	213.59	85	39.61	1,297,071	9,416.08	214	130.55

Table 3: Product configuration: search nodes and CPU time.

Instance	MILP (CPLEX 9)		SIMPL (CPLEX 9)		MILP (CPLEX 11)		SIMPL (CPLEX 11)	
	Nodes	Time (s)	Nodes	Time (s)	Nodes	Time (s)	Nodes	Time (s)
1	1	0.06	94	0.81	1	0.07	56	0.49
2	1	0.08	32	0.27	1	0.10	32	0.25
3	184	0.79	152	1.43	31	0.68	186	1.67
4	1	0.04	22	0.19	1	0.02	28	0.24
5	723	4.21	24	0.24	1	0.12	32	0.33
6	1	0.05	18	0.19	1	0.07	9	0.09
7	111	0.59	32	0.25	10	0.18	35	0.30
8	20	0.19	39	0.34	1	0.10	32	0.25
9	20	0.17	22	0.17	1	0.05	28	0.22
10	1	0.03	17	0.16	1	0.12	14	0.13

Table 4: Parallel machine scheduling: long processing times.

Jobs	Machines	MILP (CPLEX 11)		SIMPL Benders		
		Nodes	Time (s)	Iterations	Cuts	Time (s)
3	2	1	0.00	2	1	0.00
7	3	1	0.05	13	16	0.12
12	3	3,351	6.56	26	35	0.73
15	5	2,779	8.84	20	29	0.83
20	5	33,321	882.31	43	82	5.44
22	5	352,309	10,563.15	69	98	9.60

Table 5: Parallel machine scheduling: short processing times. An asterisk * means “out of memory”.

Jobs	Machines	MILP (CPLEX 11)		SIMPL Benders		
		Nodes	Time (s)	Iterations	Cuts	Time (s)
3	2	1	0.01	1	0	0.00
7	3	1	0.02	1	0	0.00
12	3	499	0.98	1	0	0.01
15	5	529	2.63	2	1	0.06
20	5	250,047	396.19	6	5	0.28
22	5	> 27.5M	> 48h	9	12	0.42
25	5	> 5.4M	> 19h*	17	21	1.09

Table 6: Bilinear global optimization: search nodes and CPU time. BARON and SIMPL were run on different machines.

Problem	Variables	Constraints	BARON		SIMPL	
			Nodes	Time (s)	Nodes	Time (s)
5.2.2.1	9	6	1	0.03	23	0.14
5.2.2.2	9	6	26	0.05	19	0.05
5.2.2.3	9	6	7	0.03	6	0.02
5.2.4	7	6	19	0.06	179	0.64
5.3.2	24	18	19	0.13	53	0.78
5.4.2	8	6	39	0.23	313	2.96

Table 7: Truss structure design: number of search nodes and CPU time (in seconds). Only CPLEX, BGH and SIMPL were run on the same machine. *CPLEX's solution to instance 3 is apparently incorrect. †Instance 9 was run in SIMPL with depth-first search (like BGH).

Problem	#Bars	BARON		CPLEX 11		BGH		SIMPL	
		Nodes	Time	Nodes	Time	Nodes	Time	Nodes	Time
1A	10	263	5.26	390	0.40	95	0.03	83	0.08
1B	10	175	3.83	106	0.26	81	0.02	73	0.07
1C	10	479	8.12	702	0.83	521	0.16	533	0.49
1D	10	518	8.76	1,320	1.17	719	0.22	726	0.63
2	10	449	24.28	2,977	4.86	841	0.64	1,028	1.84
3	10	11,354	327.19	403,683*	146.14*	517,255	144.67	94,269	64.75
4	10	34,662	2,067.43	678,471	1,086.72	1,088,955	600.09	508,816	650.71
5	25	3,190	3,301.62	3,739	43.65	11,351	44.09	2,401	20.23
6	72	291	3,375.93	1,962	207.81	665	33.01	489	27.93
7	90	782	21,610.86	2,376	576.46	1,889	130.86	826	92.47
8	108	<i>no sol.</i>	> 24h	14,966	3,208.38	9,809	1,996.87	8,485	1,719.99
9	200	<i>no sol.</i>	> 24h	<i>no sol.</i>	> 24h	<i>feasible</i>	> 24h	<i>feasible</i>	> 24h
						<i>cost =</i>	32747.58	<i>cost =</i>	32700.25†