Abstract of “A unifying framework for modeling and solving optimization problems” by Ionuț D. Aron, Ph.D., Brown University, May, 2007

The idea of a unifying framework for optimization has been around for about a decade, but very few attempts have been made at providing a description of the major components and their interaction in such a framework. A notable exception is the recent book by Hooker, which builds on the earlier work of Chandru and Hooker, Hooker, and Aron, Hooker and Yunes to provide a consistent view of optimization techniques as they are used in mathematical and constraint logic programming. The framework we describe here has evolved from a joint work with Hooker and Yunes, and it provides the basic blocks for a modeling language of metaconstraints, as well as the mechanisms that a generic metasolver needs in order to solve a problem according to the specification given by the metaconstraints. The thesis on which this framework is based is that problem solving is a combination of two fundamental steps, search and inference, whose efficient utilization requires modeling and logical deduction across theory lines. Such deduction is made possible by equipping metaconstraints with reformulation rules for multiple theories and using these rules throughout the search to draw inference from each of those theories.
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A unifying framework for modeling and solving optimization problems

by

Ionuț D. Aron

A dissertation submitted in partial fulfillment of the requirements for the Degree of Doctor of Philosophy in the Department of Computer Science at Brown University

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This dissertation by Ionuț D. Aron is accepted in its present form by
the Department of Computer Science as satisfying the dissertation requirement
for the degree of Doctor of Philosophy.

Date ______________________

Pascal Van Hentenryck, Director
Brown University,
Department of Computer Science

Recommended to the Graduate Council

Date ______________________

John N. Hooker, Reader
Carnegie Mellon University,
Tepper School of Business

Date ______________________

Meinolf Sellmann, Reader
Brown University,
Department of Computer Science

Approved by the Graduate Council

Date ______________________

Sheila Bonde
Dean of the Graduate School
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A unifying framework for modeling and solving optimization problems

Ionuț D. Aron

Brown University

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Chapter 1

Introduction

Whenever we are faced with a new problem, we attempt to identify existing theories, or perhaps create new ones that would provide us with (1) a language to express the problem and (2) axioms, inference rules and theorems to reason about it. There may be more than one suitable theory, so part of the problem solving process is to also decide which one is best equipped (perhaps has more advanced theorems, and therefore might require the smallest number of extra inference steps) to solve the problem at hand as efficiently and conveniently as possible. For instance, there are problems to which a geometric reasoning is best, and thus we have geometry, with a language, axioms and theorems as a tool for solving such problems. Other problems are better suited for an algebraic reasoning, and for them we use algebra. There are problems where set theory is best, and so on. It is also not uncommon to see elements of two or more theories being used in solving a problem. This is particularly true for optimization problems. Even though we have specific theories for certain classes of problems, such as linear programming, pure and mixed integer linear programming, nonlinear programming and constraint programming, the literature is rich in examples in which elements of these theories are combined in various ways in order to obtain solutions to difficult problems. The particular combination depends on the problem and even though it could be done in arbitrary ways, certain patterns are used more frequently than others.
A unifying framework

The scope of this thesis is to provide a unifying framework under which these common patterns can be employed in a systematic manner in order to solve computational optimization problems. This framework is based on the following thesis:

*The problem solving process, in optimization in particular, is a combination of two fundamental steps, search and logical inference, whose efficient utilization requires modeling and deduction across theory lines.*

The search step is used to advance towards a proof (solution) and it generally involves making uninformed decisions. Logical inference, or simply *inference*, plays a dual role, by detecting infeasible search directions and, when possible, preventing the search from even trying directions which can be proved wrong a priori.

1.1 Theories: language and logic

A scientific theory is a system of sentences which are accepted as true and which may be called *laws or statements*. In mathematics, these statements follow one another in a definite order according to certain principles (syntax) and are accompanied by considerations intended to establish their validity (semantics). More precisely, a theory consists of a linguistical part (*language*) and a logical part (*logic*). The language includes elementary *symbols* (constants, variables and logical connectives), as well as more complex constructions such as *terms, formulas and sentences*, all described by a grammar (*syntax*) and having a well defined meaning (*semantics*). The logical part allows us to manipulate elements of the language in order to establish new laws. It consists of *logical axioms and inference rules*. In order to *solve* a given problem with a particular theory, we must first state it using the linguistical elements of the theory and then apply the logical elements to construct a proof that the formed statement is either true or false. In the context of optimization, the language provides the tools we need to create a *model* of the problem, while the logic gives us the tools to use during the solution process.
Searching for a proof: variable elimination

Among the symbols occurring in mathematical models (as well as in theorems and proofs) we distinguish **constants** and **variables**. For instance, in arithmetic, which is the theory concerned with the investigation of the general properties of numbers, relations between numbers and operations on numbers, we encounter such constants as "number", "zero"/"0", "one"/"1", "sum"/"+", and so on. Constants have a well determined meaning, which remains unchanged throughout the course of proofs. Variables, on the other hand, do not possess any meaning by themselves. Thus, while questions like "Is zero an even number?" can always be answered in the affirmative or negative (the answer may be true or false, but at any rate it is meaningful), a question concerning a variable "x", for example the question "Is x an even number?", cannot be answered meaningfully. So even though they are syntactically correct with respect to the linguistic part, constructions like "x is an even number" do not actually express a definite assertion and can neither be confirmed or refuted. We only obtain a sentence whose truth value can be established within the theory if we somehow manage to remove the variable "x" from the statement and replace it with symbols whose meaning is well defined.

One way in which we can achieve this is to directly replace "x" with a constant denoting a definite number; thus, for instance, if "x" is replaced by the symbol "1", the result is a false statement whereas a true statement arises on replacing "x" by "2" or "10". A more desirable (but not always possible) way is to apply symbolic manipulation to the statement in order to obtain an equivalent one without variables. For example, in order to decide whether the statement "∃x ∈ R : ax² + bx + c = 0" is true, we know that it is sufficient to examine the equivalent statement "b² - 4ac ≥ 0", which can be obtained from the original through symbolic manipulation (elimination of variables, or, more precisely, of quantifiers). For simple problems, like the one above, this type of variable elimination (through symbolic manipulation) is possible, but for more complex problems, we need to rely on the first method, that of directly replacing variables with symbols. This method is known in the optimization and artificial intelligence communities as **search**. It typically involves two decisions: which variable to eliminate at a given stage,
and what constant to replace it with.\footnote{For some problems it may actually be more effective to first restrict the number of available options for a given variable, without replacing it immediately with any of them.} Since there are no general rules for either of these decisions that would guarantee a short proof, search needs to rely on guidance from the logical part of the theory at hand. This guidance comes in the form of inference and allows the search to avoid looking at all possible variable-constant substitutions.

**Using inference to guide the search**

In a given theory, logical axioms are universally true formulas and inference rules are elementary steps of the logical reasoning. For example, in propositional calculus, logical reasoning is driven by rules like *modus ponens* ("from $A \rightarrow B$ and $A$, infer $B$") and *modus tollens* ("from $A \rightarrow B$ and not $B$, infer not $A$"). A theorem is either an axiom or an immediate consequence of other theorems (by application of some inference rule). A proof is a sequence of theorems such that each theorem is either an axiom or an immediate consequence of the theorems before it. The process through which we find the immediate consequences of the existing theorems is called inference.

Inference assists the solution process in two important, distinct ways. One one hand, the search should be prevented from attempting directions which can be proved \textit{a priori} to lead to false statements. When such a priori proofs of infeasibility are not possible, the inference step is called upon after the search picks a direction and commits to it, since the effects of that decision may be easier to evaluate \textit{a posteriori}. This time, inference plays a corrective role: it checks whether the most recent decision has created a false statement and if so it informs the search component that the decision must be undone.

**Inference from multiple theories**

While the choice of a language to express the problem at hand is very important, there is no reason why the inference step should be based on a single theory. Indeed, if we manage to properly express the current state of the proof (search) in multiple languages, we could use each of the corresponding theories in order to decide on the validity of the proof so far and on the best way to proceed.
1.2 Towards a unified method: previous work

A particularly interesting possibility is to use both logic and mathematical modeling and attempt to draw inference from both theories. Early ideas hinting at the benefits of combining logic and mathematical programming can be traced back to the late 1960s, in the work of Hammer and Rudeanu[40] and later in the work of Jeroslow[54] on logic-based decision support. Jeroslow’s mixed background and interests, initially in logic and later in polyhedral theory, played a key role in his pathbreaking work towards the application of mathematical programming techniques in artificial intelligence and viceversa. He was the first to articulate the vision that we could create more effective systems, by exploiting the potential of polyhedral techniques to assist in logic based approaches and, conversely, the potential for results in applied logic to be relevant in mixed integer programming.

The research community followed suit and in the 1990s we witnessed an abundance of research on the path of integration of logic methods with optimization, much of which was captured in two books by Chandru and Hooker[28], Hooker[17], a paper collection by Milano[65] and most recently another, more crystallized book by Hooker[50]. Not only did the research community realize that Jeroslow’s vision was of great practical interest, but it became apparent that logic and optimization methods essentially share the same ingredients. The names of these ingredients vary from author to author, such as branch and infer in Bockmayr and Kasper[23], branch and bound in Land and Doig[60], branch and cut in Caprara and Fischetti[25], branch and relax, branch and price in Barnhart et al[17], branch and prune in Van Hentenryck[73] and Vu et al[74], augment and branch and cut in Letchford and Lodi[61], search and infer and relax in Hooker[48], Aron, Hooker and Yuness[6], Hooker[50], etc. At their core, however, these methods employ a common solution process, which relies on two basic components: (1) one driving the search (branching in most of them), and (2) one\(^2\) restricting the search (cutting planes, relaxations, bounding techniques, pruning techniques, variable pricing and so on).

\(^2\)Some of them break up this component into various tasks, such as cut-and-price, or infer-and-relax.
1.2.1 General purpose solvers

The first general method based on a combination of search and inference is due to Land and Doig, who proposed what became known as the branch-and-bound method for solving integer programs, as an alternative to the apparently more sophisticated cutting plane method of Gomory. This was a method restricted to mixed integer programming at the time, but when looked at carefully, it can be easily seen as being a particular case of a more general method. In fact, many of the existing techniques enumerated above, and not just branch and bound, are particular cases of a more general scheme, as pointed out by Hooker. This recognition has led to several attempts to bring these ideas to life and prove that unified systems can be as useful and powerful as various specialized methods when implemented in a carefully engineered computer software. A majority of these attempts focused on particular applications and were therefore typically restricted to them. They include applications such as matching in Focacci et al., traveling salesman in Focacci et al or scheduling in Refalo and Maravelias and Grossmann. A few, however, were geared towards producing general purpose solvers, by focusing on the common aspects of the methods. These include the branch and cut framework ABACUS, the hybrid CP-IP solver ECLiPSe, the IP-CP solver SCIP and the CP-IP-NLP metasolver SIMPL.

1.2.2 Modeling languages

In order for the integration to be effective, general purpose solvers need to rely on an expressive enough language, which can provide the modeler with sufficient opportunities to reveal problem structure to the underlying inference engine. The most successful existing implementations of integrated methods are accompanied by fairly rich languages, some of which even provide mechanisms for specifying how the search should be conducted, not only how inference should be drawn. Such languages are called procedural, because the modeler can provide, to some extent, a procedure, or recipe on how the problem is to be solved. They are clearly more powerful, but require more expertise from the modeler, than the declarative languages, which only allow the specification of the problem, not of the search procedure.
Among the most well known modeling languages for optimization we count, for instance, OPL\textsuperscript{[44]}, which provides the modeling layer for the ILOG optimization suite, namely the mixed integer programming solver ILOG CPLEX\textsuperscript{[52]} and the constraint programming solver ILOG SOLVER\textsuperscript{[51]}. A similar language, MOSEL\textsuperscript{[20]}, comes with the Dash Optimization suite, again a combination of several separate solvers: one for mixed integer programming, XPRESS-OPTIMIZER\textsuperscript{[31]}, one for constraint programming, XPRESS-KALIS\textsuperscript{[32]}, as well as two more recent solvers for stochastic and nonlinear programming. Several other languages have been created in the 1990s for various special purpose solvers. Examples include HELIOS\textsuperscript{[45]}, NEWTON\textsuperscript{[40]} and NUMERICA\textsuperscript{[43]}, which were tailored for global optimization with nonlinear constraints using interval analysis, AMPL\textsuperscript{[36]}, GAMS\textsuperscript{[24]} and AIMMS\textsuperscript{[22]} for mathematical programming and PLAM\textsuperscript{[18]} for algebraic modeling of integer programs and constraint logic programs.

Several of these languages support ideas related to the integration of optimization methods, some to a larger extent than others. Even though it would be possible to create hybrid models in these languages, this task is not always simple, since in fact none of them was created with the stated goal of bringing together multiple methods under one roof, at least not at the level where inference and search interact. The modeling aspect of optimization is an ongoing research issue and it is reasonable to expect that languages of the future would pay significantly more attention to the integration process, especially that we can already see such integrated methods being used with increasing frequency.

1.3 Organization

The material in this thesis is organized as follows. Chapter 2 introduces the notion of reformulation and establishes its connection with logical inference. Chapter 3 then presents an overview of integer programming and constraint programming, with an in-depth discussion of the pervasive disjunctive constraints and techniques for reformulating them as linear inequalities and using the derived inequalities for inference. Chapter 4 describes a unifying modeling and solution framework that can immediately incorporate linear programming, integer programming, disjunctive programming and constraint programming.
The key concept of this framework is that of metaconstraints, which can be equipped with reformulation rules so that inference is derived using multiple theories at each step of the process. The modeling concepts and solution process of this framework are illustrated using three application examples: production planning, product configuration and machine scheduling. Finally, Chapter 5 presents a more in-depth discussion of a robust optimization problem, which emphasizes the advantages of modeling using higher level metaconstraints. The results shown at the end of this chapter compare the performance of a constraint programming approach to that of a pure mixed integer approach. They show that the additional inference obtained by using graph theoretical arguments at each step of the solution process is instrumental in obtaining results for larger problem instances, as well as getting significantly faster results on small instances.
Chapter 2

Inference via reformulation

It is often the case that in order to prove that a given statement is false, it may be useful to restate it in a different language and use different types of arguments. Such a reformulation, which is in essence a syntactic translation, cannot be done in arbitrary ways. It must be done such that the resulting statement is always an implication, if not an equivalence, of the original. The relaxation techniques used in optimization are perfect examples of syntactic translations which result in such an implication. If the implication is then found to be false, we can infer by application of the modus tollens rule that the original statement is false too.

2.1 The importance of reformulation

Reformulation is a valuable tool in problem solving, whose role cannot be understated. To illustrate its power, let us look at an instructive example, the pigeonhole problem, adapted from McAloon et al\textsuperscript{[64]}. Further examples of inference via reformulation will be discussed later in the context of disjunctive programming, where we look at two important reformulation techniques: conjunction elimination from systems of linear inequalities (known as the derivation of surrogates) and disjunction elimination from systems of linear inequalities (known as the derivation of disjunctive cuts). In the pigeonhole problem, we have \( m \) pigeons and must place them in \( n \) holes such that only one pigeon is in any one hole. The problem can be easily modeled using the language of propositional logic, and
we could attempt to solve it within that theory, but as it will become clear, the solution process (proof) can be substantially shortened if we use reformulation.

**A propositional logic model**

Let \( p_{ij} \) be a variable with possible values \{true, false\} that indicates whether the \( i^{th} \) pigeon is in the \( j^{th} \) hole. Then

\[
\bigwedge_i \bigvee_j p_{ij}
\]

(2.1)

states that "every pigeon has to be placed in (at least) one hole", and

\[
\bigwedge_j \bigwedge_i \left( p_{ij} \rightarrow \bigwedge_{k<i} \neg p_{kj} \right)
\]

(2.2)

states that "any hole can contain at most one pigeon". Note that we are not able to state the condition "at most one pigeon can be in hole \( j \)", using a single formula for each \( j \), since propositional logic does not have a connective for this.

If we now step back and read the description of the problem again, we can immediately see that the problem admits a solution if and only if \( m \leq n \). We know this because we subconsciously use some additional arguments, namely counting arguments, which are actually not part of the propositional logic theory. In particular, we use knowledge about the "sum" (+) and "greater than" (>) operators: we add up the pigeons, then the holes, then we compare the resulting values and decide that the problem is unsolvable if there are more pigeons than holes. However, without these two operators, classical theorem provers may find it very hard to prove that no solution exists in the case when \( m > n \).

The shortest resolution proof that detects the unsatisfiability of this problem involves a number of steps that is exponential in \( n \), as shown in Haken\[39\].

Since it appears that counting arguments can be quite effective in detecting the infeasibility of this problem, let us attempt to translate this propositional model into a theory which allows such arguments. Integer 0-1 programming is a good candidate, because we have a set of simple rules which can translate from proposition logic syntax to 0-1 integer programming syntax (see, for example, Hammer and Rudeanu\[40\]).
A 0-1 integer programming reformulation

Let $x_{ij}$ be a variable with possible values $\{0, 1\}$, such that $x_{ij} = 1$ iff $p_{ij} = true$ and $x_{ij} = 0$ iff $p_{ij} = false$. Using these variable translation rules, it is easy to verify that the propositional logic condition

$$\bigvee_j p_{ij} \quad (2.3)$$

implies the 0-1 integer programming condition

$$\sum_j x_{ij} \geq 1 \quad (2.4)$$

Thus, we can deduce the following:

$$\bigwedge_i \bigvee_j p_{ij} \rightarrow \bigwedge_i \sum_j x_{ij} \geq 1 \rightarrow \left( \sum_j x_{ij} \geq 1, \forall i \right) \quad (2.5)$$

which essentially translates the statement that "every pigeon has to be placed in (at least) one hole" from propositional logic to 0-1 integer programming.

In order to translate the second condition, let us first rewrite it as a conjunction of disjunctions, so that we can apply the same translation rule as above. This is known as writing the formula in conjunctive normal form (CNF). To obtain this form, we use the following equivalences, which are easily obtained starting from the axioms of propositional calculus and applying the modus ponens inference rule:

$$(A \rightarrow B) \leftrightarrow (\neg A \lor B) \quad (2.6)$$

and

$$A \lor (B \land C) \leftrightarrow (A \land B) \lor (A \land C) \quad (2.7)$$

Now we can write

$$\bigwedge_i \bigwedge_{j \neq k} \left( p_{ij} \rightarrow \bigwedge_{k < i} \neg p_{kj} \right) \rightarrow \bigwedge_i \bigwedge_{j \neq k} \left( \neg p_{ij} \lor \bigwedge_{k < i} \neg p_{kj} \right) \rightarrow \bigwedge_i \bigwedge_{j < k} \left( \neg p_{ij} \lor \neg p_{kj} \right)$$

The variable mapping we established above allows us to say that $\neg p_{ij}$ is true iff $1 - x_{ij} = 1$, so we can now rewrite the condition that any hole can contain at most one pigeon using
A linear programming reformulation

In this form, the problem still remains quite difficult to solve for integer programming systems, even for small values of $m > n$. However, we are now able to express the condition that at most one pigeon can be in hole $j$ quite naturally with a single formula for each $j$:

$$
\sum_i x_{ij} \leq 1, \forall j
$$

which we can use in place of the above $O(nm^2)$ inequalities. Together with the condition that each pigeon be placed in at least one hole,

$$
\sum_j x_{ij} \geq 1, \forall i
$$

and with a further reformulation of the $x$ variables into continuous $y$ variables, we obtain the linear program

$$
\sum_i y_{ij} \leq 1, \forall j
$$

$$
\sum_j y_{ij} \geq 1, \forall i
$$

whose feasible region is the empty set if $m > n$.

Thus, the fact that the problem does not have any solutions for $m > n$ can be established without performing any search by working with an appropriate theory: in this case, the theory of continuous linear programming.

2.2 A special case: relaxation

In general, reformulation does not have to be done for the entire problem, and even when it is done, individual constraints could be reformulated in different theories. However, it is often the case that in practice all (or a subset of) the constraints are reformulated in the same target language, leading to what is usually referred to as a relaxation of the
original problem. This, for example, is the way integer programming solvers approach the inference step of the solution process. The use of discrete variables introduces discontinuities and nonconvexity, which makes it considerably more difficult to reason about them than it would be to reason about continuous variables. Therefore the original constraints (including the integrality constraints) are reformulated into linear inequalities (the integrality constraints \( x \in D_x \subseteq \mathbb{Z} \) are reformulated into the conjunction of linear inequalities \( \inf\{D_x\} \leq x \land x \leq \sup\{D_x\} \)) and then the simpler linear programming theory can be used to derive inference. Such inference comes in various forms, but serves the two basic roles we mentioned earlier: (a) refutation of the statement represented by the current search state, or (b) guidance for restricting and continuing the search.

**Using relaxation to detect infeasibility**

A continuous view of the problem can provide an easy proof that the search has gone wrong, if it detects that its own feasible region is empty. Secondly, if the region is not empty, nothing can be said about the feasibility of original problem, but due to the special properties of the continuous relaxation, we might be able to say something about its optimality (which could also be regarded as feasibility). Thus, if the original problem includes an objective function in its statement, the continuous relaxation could still help us decide that there is no point in continuing with this search direction, if we can show that there could be no solution inside the continuous relaxation which provides a better value than whatever best solution we may have found previously. This conclusion of infeasibility can be reached in two ways: either the best solution of the continuous relaxation is already worse than our best discrete solution, or we could show that it would become worse no matter what search decisions we make from here. The first is easy to check, by simply evaluating the objective function on the most extreme point of the continuous feasible region. The second involves evaluating the changes in the objective function by plugging in some of the values for the original variables which the search has not yet explored. Values which would make the objective worse than the best discrete solution so far can be safely discarded. A similar technique is used in constraint programming, where it is known as reduced-cost based filtering (see Focacci et al.\[34\]).
Using relaxation to avoid infeasibility

Note here the second role played by the continuous view of the problem, that of prevention, or guidance: even if we are unable to refute the current state, because perhaps we cannot rule out all remaining values using the objective function, we can still safely remove some of them, essentially reducing the number of (bad) choices the search would have to try from this point.

An additional (and particularly useful) preventive inference technique based on a continuous view of the problem involves cutting planes. These are linear inequalities meant to help the bounding process through which we attempt to detect infeasible (sub-optimal) search states, as indicated before. Cutting planes can be viewed as being of two types, depending on how they are inferred: (1) either directly from the problem, or (2) from an implication of the problem (which could be the continuous linear relaxation). In the first case, they are called structural cuts (or sometimes polyhedral cuts), because they are implied by the structure of the problem and depend neither on the language in which the problem is stated, nor on the search state. These cuts are direct linear implications of the original problem. The well-known comb inequalities for the traveling salesman problem (see Applegate et al. [3]) are a perfect example of such inequalities. The second type of cuts are inferred from the continuous view of the problem, by focusing on removing a tentative solution if it fails to satisfy all the constraints of the original problem. They are derived at some stage during the search, using two basic inference rules: (1) non-negative combinations of linear inequalities (this is where the continuous view plays a role) and (2) integer rounding (this is where the original integrality constraints play a role). As such, these cuts are indirect linear implications of the problem and are valid only during subsequent stages of the search (although they can be made valid everywhere through a process called lifting). Examples of such cuts include Gomory cuts (see Gomory [38]), mixed integer rounding cuts (see Marchand et al. [63]), intersection cuts (see Balas [9]) and disjunctive cuts (see Balas [10]). We will have a chance to look at this type of cutting planes in more detail in Section 3.1.3 when we discuss inference based on a special subclass of disjunctive cuts, namely lift-and-project cuts (see Balas [7]).
Chapter 3

Optimization theories

In this chapter we review two commonly used optimization theories: integer programming (IP) and constraint programming (CP). They constitute the starting point for the unifying framework presented in Chapter 4, since they share many features in terms of solution method, but at the same time complement each other in terms of inference capabilities. Even though the scope of the framework we propose is in no way limited to these two theories, we restrict the discussion to them for the following reasons: (1) they are both well studied and understood, (2) they have been successfully applied to a wide range of practical applications, (3) they are among the most frequently used theories in various hybrid solutions proposed in the last decade and (3) our implementation of this framework is currently limited\footnote{This limitation refers to available and tested functionality.} to these two theories.

3.1 Integer and disjunctive programming

Integer programming is linear programming with the additional constraints that some of the variables must take integer (discrete) values. For this reason, integer programs are typically much more difficult to solve than pure linear programs. It is interesting to point out here that it was unknown for a long time whether even solving a pure linear program can be done efficiently. The original proof proposed by DANTZIG\textsuperscript{[30]} in 1947, based on linear
algebraic arguments, required an exponential number of steps in the worst case, as shown by Klee and Minty\cite{Klee1972}. Three decades later, however, the Soviet mathematician Leonid Khachyan\cite{Khachyan1980} was able to show that linear programming does admit a polynomial length proof, but for that he had to use arguments from mathematical analysis, as opposed to just linear algebra. No such proof has been found for integer programming, and it is widely believed that none exists.

Disjunctive programming goes one step further from integer programming, by allowing in the language an additional type of constraints, namely disjunctive (linear) constraints. These constraints naturally capture the idea of choice, which is inherent to all combinatorial optimization problems. Integer programs, pure and mixed, and a host of other non-convex programming problems can be stated more naturally as linear programs with logical conditions, more precisely statements about linear inequalities involving the operators “or”. For a good introduction to disjunctive programming see Balas\cite{Balas1998}.

**Modeling and solution process**

Integer programming solvers typically solve a linear program at every node of the search tree in order to obtain bounds on the objective function (and sometimes with the hope that the resulting solution is also feasible with respect to the integrality constraints). There are no integer programming solvers that deal directly with disjunctive constraints at the modeling level. Instead, these constraints are either enforced implicitly, through a branching mechanism, or reformulated (relaxed) into linear constraints that integer solvers can deal with. In the remaining of this section, we focus on this reformulation and present one type of linear inequalities (cuts) that can be derived from disjunctions and can be used as an inference tool in an integrated solver. The algorithm discussed here is an extension of the one created by Balas and Perregaard\cite{Balas1995}, and it is the latest development in a field of research that originated in the work on intersection cuts (also known as convexity cuts) by Young\cite{Young1985}, Balas\cite{Balas1982}, Glover\cite{Glover1988} and Owen\cite{Owen1988}.
3.1.1 Disjunctive linear constraints

Consider the family of inequalities \( \{ a^i x \geq b^i \}_{i \in I} \), with \( x \in \mathbb{R}_+ \), \( a^i \in \mathbb{R} \), \( b^i \in \mathbb{R} \). A linear disjunction (or simply, disjunction) is a proposition which states that "at least one of the inequalities in this family is satisfied" and has the form:

\[
\bigvee_{i \in I} a^i x \geq b^i
\]  

(3.1)

Note that when \( b^i > 0 \), which is always the case when we derive cuts from a (feasible basic solution in a) simplex tableau, disjunction (3.1) is clearly equivalent with one in which we multiply each disjunct with the inverse of its right hand side:

\[
\bigvee_{i \in I} a^i x \geq b^i \iff \bigvee_{i \in I} \frac{a^i}{b^i} x \geq 1
\]  

(3.2)

It is often convenient to represent a disjunction in the form (3.2) whenever \( b^i > 0 \), because it allows us to compare the coefficients (and hence the strength) of inferred inequalities one by one without having to worry about the right hand side. We say that disjunctions in this form, as well as the inequalities implied by them, are normalized. We will later see that it is also desirable, not just convenient, to represent disjunctions in normalized form.

More generally, we can define disjunctions for systems (conjunctions) of linear inequalities

\[
\bigvee_{i \in I} A^i x \geq b^i
\]  

(3.3)

where \( A \) is an \( m \times n \) matrix, \( b^i \in \mathbb{R}^m \), \( x \in \mathbb{R}^n \), with the important special case \( m = 1 \)

\[
\bigvee_{i \in I} a^i x \geq b^i
\]  

(3.4)

with \( a^i \in \mathbb{R}^n \). Note also that the normalization argument used in (3.2) can be easily generalized to apply to these cases as well.

Disjunctive programming is closely related to linear programming, since a sentence involving the disjunctive connective "\( \lor \)" can be replaced by an implication using only linear connectives. This is a reformulation process, and it can help detecting when a disjunctive model is infeasible.
3.1.2 Inference via reformulation

We now discuss the basic translation rules that allow us to convert a disjunction into a linear inequality. By conversion, it is understood that we want to derive an inequality such that all solutions that satisfy the original disjunction, also satisfy that inequality. In other words, the derived inequality should be an *implication* of the disjunction, and as such, if it turns out to be a false statement, then we can say the same about the disjunction.

**Linear implications that eliminate disjunction**

The following basic translation rules provide the main instrument used to derive a class of linear inequalities known as *disjunctive cuts* (see BALAS[11, 12]). These inequalities are reformulations of the original disjunction, into a language from which the "∨" connective has been eliminated, and as such, they can be reasoned about using the simpler theory of linear programming.

**Proposition 3.1.1.** If the disjunction \( \bigvee_{i \in I} a^i x \geq b^i \) is true, then the following linear inequality must hold:

\[
\max_i \{a^i\} x \geq \min_i \{b^i\} \tag{3.5}
\]

Similarly, if \( \bigvee_{i \in I} a^i x \leq b^i \) is true, then

\[
\min_i \{a^i\} x \leq \max_i \{b^i\} \tag{3.6}
\]

is also true.

These rules can be easily generalized to the case when \( a \) and \( x \) are \( n \)-dimensional, as well as to disjunctions of systems of linear inequalities, such as (3.3) and (3.4).

**Proposition 3.1.2.** Let \( A^i \) be an \( m \times n \) matrix, \( a^i \in \mathbb{R}^n \), \( b^i \in \mathbb{R}^m \), for any \( i \in I \), and \( x \in \mathbb{R}^n_+ \). Then the following implications hold:

\[
\bigvee_{i \in I} a^i x \geq b^i \rightarrow \sum_{j=1}^n \max_i \{a^i_j\} x_j \geq \min_i \{b^i\} \tag{3.7}
\]

\[
\bigvee_{i \in I} a^i x \leq b^i \rightarrow \sum_{j=1}^n \min_i \{a^i_j\} x_j \leq \max_i \{b^i\} \tag{3.8}
\]
\[ \bigvee_{i \in I} A^i x \geq b^i \rightarrow \sum_{j=1}^{n} \max \{ A^i_j \} x_j \geq \min \{ b^i \} \quad (3.9) \]

\[ \bigvee_{i \in I} A^i x \leq b^i \rightarrow \sum_{j=1}^{n} \min \{ A^i_j \} x_j \leq \max \{ b^i \} \quad (3.10) \]

where \( A^i_j \) represents the \( j^{th} \) column of matrix \( A^i \) and \( v \) (respectively \( v \)) represents the largest (respectively smallest) element of vector \( v \).

The reformulation rules in Proposition 3.1.2 are of central importance in disjunctive (and therefore integer) programming, because they can be used to derive a large family of cutting planes, including ones for mixed integer programming, such as mixed integer Gomory cuts and mixed integer rounding (MIR) cuts (see Nemhauser and Wolsey\textsuperscript{[66]} and Marchand and Wolsey\textsuperscript{[63]}), where the Chvatal-Gomory rounding procedure (see Schrijver\textsuperscript{[70]}) fails. The consequents of these implications are called disjunctive cuts (see Balas\textsuperscript{[11]}).

**Linear implications that eliminate conjunction**

The other reformulation rule which plays a key role in integer and linear programming is the following:

**Proposition 3.1.3.** Let \( A \) be an \( m \times n \) matrix, \( b \in \mathbb{R}^m \) and \( x \in \mathbb{R}^n_+ \). Then the following implication holds:

\[ (Ax \geq b \land u \geq 0) \rightarrow uAx \geq ub \quad (3.11) \]

This property allows us to summarize an entire system (conjunction) of inequalities with a single inequality, which is much easier to check against infeasibility. The above implication is clearly a reformulation, since it removes from the language the conjunction connective "\( \land \)" , and therefore allows us to use a simpler theory in order to detect if the original system is inconsistent. The resulting inequality, \( uAx \geq ub \), is called a surrogate. It is used to derive so-called valid inequalities for the linear programming relaxation of an integer program.

**Definition 3.1.4 (Valid inequality).** An inequality \( \alpha x \geq \beta \) is said to be valid for a set \( S \subseteq \mathbb{R}^n \) if it is satisfied by all the points \( x \in S \).
Clearly, all surrogates $uAx \geq ub$ are valid for the set $P = \{ x \in \mathbb{R}_+^n : Ax \geq b \}$. The requirement that $u \geq 0$ ensures that the sense of the inequalities is preserved. Obviously, this requirement can be relaxed to $u_i \in \mathbb{R}$ if the $i^{th}$ inequality of the system $Ax \geq b$ is in fact an equality (because a negative multiplier would not affect its sense).

**Stronger inference via a combination of these rules**

If we now look back at the linear inequalities obtained through Propositions 3.1.1 and 3.1.2, it is not difficult to see that they are not the strongest inferences we can make from a disjunction of linear inequalities. Indeed, let us take a very simple example, to illustrate what can be done in order to obtain a stronger inequality from such a disjunction. Consider the disjunction:

$$4x \geq 3 \lor x \geq 1$$

(3.12)

If we simply apply Proposition 3.1.1 to this disjunction, we obtain the valid inequality $4x \geq 1$. However, we can do better. Notice that if we multiply the second disjunct with 3, we obtain an equivalent disjunction:

$$4x \geq 3 \lor 3x \geq 3$$

(3.13)

But the inequality we can now infer from this disjunction by applying Proposition 3.1.1 is much stronger: $4x \geq 3$. This is so because it has the same left hand side, but a tighter right hand side (that is, there are fewer values of $x$ that satisfy it). We say that this inequality *dominates* the one we had obtained before.

**Definition 3.1.5 (Dominating inequality).** An inequality $\alpha x \geq \beta$ is said to *dominate* another inequality $\alpha' x \geq \beta'$ if $\alpha \leq \alpha'$ and $\beta \geq \beta'$.

What did actually happen in the transformation from (3.12) to (3.13)? We managed to infer a tighter valid inequality by:

1. collecting together all the inequalities of the disjuncts into a system (*conjunction*) of inequalities:

$$\begin{pmatrix}
4x \geq 3 \\
x \geq 1
\end{pmatrix}$$
2. then carefully choosing a non-negative multiplier \( \mathbf{u} = (u_1, u_2) = (1, 3) \) for this system such that to preserve (or decrease) the maximum coefficient of \( x \) in the left hand side, and to increase (or preserve) the minimum value on the right hand side:

\[
\begin{pmatrix}
4x & \geq & 3 \\
\times 1 \\
3x & \geq & 3
\end{pmatrix}
\]

3. and finally applying disjunction elimination to the resulting inequalities:

\[
(4x \geq 3 \lor 3x \geq 3) \rightarrow 4x \geq 3
\]

Note that when we derived the weaker \( 4x \geq 1 \) from the original disjunction, we in fact used a default multiplier, namely \( \mathbf{u} = (u_1, u_2) = (1, 1) \). The new multiplier is certainly better, and we can tell that because it reduced the difference between the left hand side and the right hand side of the resulting inequality. The fact that we were able to increase the right hand side means that the original inequality, although valid, was not maximal.

**Definition 3.1.6 (Maximal inequality).** A maximal valid inequality is one which is not dominated by any other valid inequality.

Such inequalities are also known as face defining, or supporting inequalities. Our goal is to devise a technique to find valid inequalities that are as strong as possible (maximal).

**Remark 3.1.7.** We said before that it is often convenient to normalize the disjunction when the right hand side is strictly positive. It should be clear by now that it is not just convenient. Note that the effect of this normalization is the same as what we have done with the multiplier \( \mathbf{u} \) in order to strengthen the inferred inequality. Indeed, had we first normalized the disjunction, we would have obtained:

\[
\frac{4}{3} x \geq 1 \lor x \geq 1
\]

which now leads directly (by Proposition 3.1.2) to the stronger valid inequality (3.14). So it is always a good idea to normalize disjunctions that satisfy this condition.
Connection with linear programming duality

You might have recognized already the principle at work here (and perhaps the connection with linear programming duality). Indeed, in linear programming, we try to find a non-negative multiplier \( u \) for a system of inequalities \( Ax \geq b \) which transforms it into an implied inequality \( uAx \geq ub \), such that we keep \( uA \) below some predetermined objective vector \( c \) (i.e. \( uA \leq c \)) and at the same time increase the right hand side \( ub \) by as much as possible. We can find such a multiplier \( u \) by solving a linear programming dual of the original LP:

\[
\begin{array}{c|c}
\text{LP} & \text{Dual} \\
\hline
\min & \max u \\
\text{s.t} & \text{s.t} \\
Ax & uA \leq c \\
x & u \geq 0 \\
\end{array}
\]

There is a difference, however, with finding the multiplier \( u \) that strengthens an inequality inferred from a disjunction of single linear inequalities \( a'x \geq b' \) (collected into a system of inequalities \( Ax \geq b \)). The difference is that we are not computing the dot products \( uA_j \) and \( ub \), as we do in the dual linear program, but instead we take the maximum and minimum of the element-wise products \( u \odot A \) and \( u \odot b \), where:

\[
u \odot A = \begin{pmatrix}
u_1 a_{11} & u_1 a_{12} & \ldots & u_1 a_{1n} \\
u_2 a_{21} & u_2 a_{22} & \ldots & u_2 a_{2n} \\
\vdots & \vdots & \ddots & \vdots \\
u_m a_{m1} & u_m a_{m2} & \ldots & u_m a_{mn}
\end{pmatrix}
\]

\[
u \odot b = \begin{pmatrix}
u_1 b_1 \\
u_2 b_2 \\
\vdots \\
u_m b_m
\end{pmatrix}
\]

So if we denote

\[
\alpha = u \odot A = \left( \max_i \{ u_ia_{i1} \}, \max_i \{ u_ia_{i2} \}, \ldots, \max_i \{ u_ia_{in} \} \right) \in \mathbb{R}^n
\]

\[
\beta = u \odot b = \min_i \{ ub_i \} \in \mathbb{R}
\]

then finding the strongest inferred inequality \( \alpha x \geq \beta \) amounts to solving the following problem:

\[
\max \beta \\
\text{s.t} \quad \alpha \leq c \\
\quad \quad u \geq 0
\]
where the unknown is the multiplier $u$ and where the bounding vector $c = \mathbf{1} \odot \mathbf{A}$ contains the coefficients of the inequality obtained using the default multiplier $u = 1$.

**Finding valid inequalities for a disjunction of conjunctions**

It is easy to verify that an inequality $\alpha \mathbf{x} \geq \beta$ is valid for the set $P = \{ \mathbf{x} \in \mathbb{R}^n_+ : \mathbf{A} \mathbf{x} \geq \mathbf{b} \}$ if and only if it is dominated by a surrogate of $\mathbf{A} \mathbf{x} \geq \mathbf{b}$. Furthermore, if we have a disjunction of such sets, written as

$$\bigvee_{i \in I} \mathbf{x} \in P^i \iff \bigvee_{i \in I} \mathbf{A}^i \mathbf{x} \geq \mathbf{b}^i$$

(3.16)

a theorem by Balas\cite{11} says that $\alpha \mathbf{x} \geq \beta$ is valid for (3.16) if and only if it is dominated by a surrogate $\mathbf{u}^i \mathbf{A}^i \mathbf{x} \geq \mathbf{u}^i \mathbf{b}^i$ of each of the disjuncts $i \in I$. In other words, if and only if it is valid for each of the sets $P^i = \{ \mathbf{x} \in \mathbb{R}^n_+ : \mathbf{A}^i \mathbf{x} \geq \mathbf{b}^i \}$. So, in order to find a valid inequality for (3.16), it is enough to find a set of multipliers $\mathbf{u}^i$ with this property, and then use Proposition 3.1.2. We obtain the following valid inequality:

$$\sum_j \max_{i \in I} \{ \mathbf{u}^i \mathbf{A}^i_j \} x_j \geq \min_{i \in I} \{ \mathbf{u}^i \mathbf{b}^i \}$$

(3.17)

To find these multipliers, we can set up a simple linear program and require that they define inequalities $\mathbf{u}^i \mathbf{A}^i \mathbf{x} \geq \mathbf{u}^i \mathbf{b}^i$ which dominate some $\alpha \mathbf{x} \geq \beta$:

$$\begin{align*}
\text{max} & \quad 0 \\
\text{s.t} & \quad \alpha \geq \mathbf{u}^i \mathbf{A}^i \quad \forall i \in I \\
& \quad \beta \leq \mathbf{u}^i \mathbf{b}^i \quad \forall i \in I \\
& \quad \mathbf{u}^i \geq 0 \quad \forall i \in I \\
& \quad \alpha, \beta \in \mathbb{R}^n
\end{align*}$$

(3.18)

The unknowns in this system are $\mathbf{u}^i$, $\alpha$ and $\beta$. The first two inequalities simply state that the surrogates $\mathbf{u}^i \mathbf{A}^i \mathbf{x} \geq \mathbf{u}^i \mathbf{b}^i$, for all $i \in I$, dominate $\alpha \mathbf{x} \geq \beta$, which implies that $\alpha \mathbf{x} \geq \beta$ will be valid for (3.16). Solving this system for $\mathbf{u}$ ($\alpha$ and $\beta$ being unconstrained in sign, can be eliminated), we obtain the coefficients of valid inequalities of the form $\alpha \mathbf{x} \geq \beta$ that obviously satisfy:

$$\alpha_j = \max_{i \in I} \{ \mathbf{u}^i \mathbf{A}^i_j \}$$
and

$$\beta = \min_{i \in I} \{ u^i b^i \}$$

In other words, we have obtained the same inequality as we would (if we knew $u^i$) using the reformulation rules in Proposition 3.1.2, namely (3.17).

**Finding stronger inequalities**

We obviously wrote (3.18) such that it finds one set of multipliers $u^i$ in order to obtain a valid inequality. We saw before that it may be possible to find stronger inequalities, by finding *better* multipliers. In order to do this, note first that we can rewrite the system (3.18), by moving all the variables on the left hand side, in the following form:

$$\begin{align*}
\max & \quad 0 \\
\text{s.t} & \quad \alpha - u^i A^i \geq 0 \quad \forall i \in I \\
& \quad -\beta + u^i b^i \geq 0 \quad \forall i \in I \\
& \quad u^i \geq 0 \quad \forall i \in I \\
& \quad \alpha, \beta \in \mathbb{R}^n
\end{align*}$$

(3.19)

which makes it apparent that this is a polyhedral cone. This observation becomes useful if we want to optimize a more interesting function than $\max 0$, one which could involve $\alpha$ and $\beta$.\(^2\) It tells us that we would have to truncate this cone with some bounding inequality (or inequalities) in order to prevent $\alpha$ and $\beta$ from scaling up indefinitely. Evidently, the choice of the bounding constraints used to truncate the cone could have an impact on the type (and quality) of solutions we obtain, so it is important to pay attention to this aspect.

*Balas et al.*\(^14\) consider adding one of three possible bounding constraints (which they call normalizations) to the system (3.19) in order to guarantee that optimal solutions exist when the objective function involves the unknowns $\alpha$ and $\beta$ (i.e. the coefficients of the valid inequality). Their bounding constraints focused on one of these variables only, and they were: (i) $|\beta| \leq 1$, or (ii) $|\alpha_j| \leq 1, \forall j$, or (iii) $\sum_j |\alpha_j| \leq 1$. Clearly, by imposing such constraints, we limit the class of inequalities that we can obtain with (3.19), but at least

\(^2\)Therefore it could help us find better valid inequalities $\alpha x \geq \beta$, in some sense.
we hope that the system has a (finite, nontrivial) solution. In the computational study of the original lift-and-project method by Balas et al.\textsuperscript{[15]}, which generated cuts from 0-1 disjunctions on a single variable (i.e. $x_k \leq 0 \lor x_k \geq 1$ for some integer constrained $x_k$), the bounding constraint $|\beta| \leq 1$ was the best (in terms of cut quality) among the three. Unfortunately, the problem with this constraint is that $\alpha$ can still be unbounded. Indeed, to find a cut which is (maximally) violated at some point $\tilde{x}$, we would use an objective like $\min \alpha \tilde{x} - \beta$. But if there exists a valid cut of the form $\alpha' x \geq 0$ with $\alpha' \tilde{x} < 0$, then the objective is clearly unbounded. The other two bounding constraints pose different problems. Even though they both achieve the goal of bounding $\alpha$, they either produce cuts of lesser quality (as Balas et al.\textsuperscript{[15]} found for $|\alpha_j| \leq 1$), or they are computationally more demanding, by increasing the size of the linear program (as it is the case with $\sum_j |\alpha_j| \leq 1$, which in order to be used inside a linear solver requires that we split the unrestricted variables $\alpha$ into two positive variables $\alpha = \alpha^+ - \alpha^-$).

A better bounding constraint was proposed by Ceria and Pataki\textsuperscript{[26]}:

$$\sum_{i \in I} \sum_j u^i_j \leq 1 \quad (3.20)$$

Because the multipliers $u^i$ are non-negative, and because $\alpha$ and $\beta$ are obtained as linear combinations of $u^i$, this constraint is sufficient to bound all the variables in (3.19). This and the previous three bounding constraints are discussed in detail by Ceria and Soares\textsuperscript{[27]}. They also give some nice interpretations of the dual formulation of (3.19) when the different constraints are used.

**Generating the deepest cuts**

The problem of finding better valid inequalities arises when we have a certain reference point $\bar{x} \in \mathbb{R}^n$ which does not satisfy disjunction (3.16). In this case, we want to derive an inequality $\alpha x \geq \beta$ which is valid for the disjunction, but is maximally violated by this point. To find this inequality, we can use the linear program (3.19), to which we add the bounding constraint (3.20) and the objective function $\min \alpha \bar{x} - \beta$. We obtain the
following linear program, which Balas et al. call *cut generating linear program*:

\[
\begin{align*}
\min & \quad \alpha x - \beta \\
\text{s.t} & \quad \alpha - u^i A^i \geq 0 \quad \forall i \in I \\
& \quad -\beta + u^i b^i \geq 0 \quad \forall i \in I \\
& \quad u^i \geq 0 \quad \forall i \in I \\
& \quad \sum_i u^i \cdot 1 \leq 1 \\
& \quad \alpha, \beta \in \mathbb{R}^n
\end{align*}
\] (CGLP)

Solving this system yields an inequality \( \alpha x \geq \beta \), with

\[
\alpha_j = \max_{i \in I} \{ u^i A^i_j \} \quad \text{and} \quad \beta = \min_{i \in I} \{ u^i b^i \}
\] (3.21)

where \( A^i_j \) is the \( j^{th} \) column of \( A \). This inequality (cut) maximizes the amount \( \beta - \alpha x \) by which \( x \) is cut off (separated from) the solutions of the original disjunctive program.

**Making the cuts valid for the original problem**

If separating inequalities such as (3.21) are generated at nodes other than the root of a search tree, they subsume the decisions that search has made so far, by assigning values to components of \( x \). Those components \( x_j \) of \( x \) that have been fixed no longer participate actively in the selection of the dual multipliers \( u^i \) in (CGLP), since their corresponding columns from \( A^i_j \) have been moved (added as a constant factor) to the right hand side \( b^i \). Thus, the fact that \( \alpha x \geq \beta \) is dominated by the surrogates \( u^i A^i x \geq u^i b^i \) (which makes it a valid inequality) is likely to change when we undo those assignments (during backtracking). So these inequalities are valid only in the subtree where the assignments hold, and cannot be used in other parts of the search space.

In principle, however, it is possible to make them valid for the whole search space, by calculating appropriate values for the coefficients of the fixed variables. This process is called *lifting*. In general, this is a difficult task, which may require the solution of an integer program for every missing coefficient. Fortunately, the cuts we are looking at here have an important advantage: we can use the multipliers \( u \) obtained (along with the cut coefficients \( \alpha \) and \( \beta \)) by solving (CGLP) to calculate, by closed form expressions, the objective coefficients \( \alpha_j \) whose corresponding variable \( x_j \) has been fixed by search. The
expressions for these coefficients are given by:

\[ \alpha_j = \max_{i \in I} \{ u^i A^i_j \} \]  

(3.22)

as shown by BALAS ET AL\(^{[14, 15]}\).

### 3.1.3 Lift and project as an inference technique

We now have the background necessary to discuss a more effective algorithm for generating disjunctive cuts. We focus here on lift and project cuts (see BALAS ET AL\(^{[14]}\)), which are a special class of disjunctive cuts, derived from a two term disjunction of the form \((x_k \leq 0 \lor x_k \geq 1)\) imposed to the set \(P = \{ x \in \mathbb{R}_+^n : Ax \geq b \}\). In other words, they are valid inequalities for the set of solutions of:

\[
\begin{align*}
Ax & \geq b \\
x & \geq 0 \\
x_k & \leq 0 \lor x_k \geq 1
\end{align*}
\]

(3.23)

or, equivalently,

\[
\begin{align*}
\left( \begin{array}{c}
Ax \\
x \\
-x_k
\end{array} \right) \geq \begin{array}{c} b \\
x \geq 0 \\
x_k \geq 1
\end{array}
\end{align*}
\]

(3.24)

for some \(k \in \{ 1, \ldots, n \}\) such that \(0 < \bar{x}_k < 1\), where \(\bar{x}_k\) is the value of variable \(x_k\) in the current solution of the LP relaxation. We want to impose the (nonlinear and non-convex) condition that \(x_k \in \{ 0, 1 \}\). But since integer solvers cannot handle it in the model, we have to obtain a linear implication of this condition, one which is hopefully strong enough that it removes (at least) the current reference point \(\bar{x}\) from further consideration.

**Unstrengthened lift and project cutting planes**

Note that disjunction (3.24) has the same form as the general disjunction (3.16), which means that we can find valid inequalities implied by (3.24) using the linear program defined in (3.19). In other words, we find a non-negative multiplier for each of the disjuncts, require that \(\alpha x \geq \beta\) be dominated by the resulting surrogates, and then check to see whether
the current solution \( \bar{x} \) of \( P \) violates \( \alpha x \geq \beta \). If it does (and it should, because we also took into account the two inequalities that forbid the current value of \( \bar{x}_k \)), that means we have a cut already, and we can try to find an even better one (one which \( \bar{x} \) violates “more”) by trying a new multiplier. This, in essence, is the same procedure as the one described in Balas et al.\(^\text{[15]}\). They call the resulting system, together with the bounding constraint (3.20) and the objective function \( \min \alpha \bar{x} - \beta \) a cut generating linear program for variable \( x_k \) (CGLP\(_k\)). For each binary variable that happens to have a fractional value in the current LP solution, they solve one such system and add the resulting inequality to the problem formulation.

The cut generating linear program

Let us illustrate the process of constructing the cut generating LP and finding the right multipliers for the two disjuncts. To simplify notation, we can include the non-negativity constraints \( x \geq 0 \) together with the rest, \( Ax \geq b \), to obtain a new system \( A'x \geq b' \), where \( A' = \left( \frac{A}{I} \right) \) and \( b' = \left( \frac{b}{0} \right) \). Now we can write the disjunction (3.24) as

\[
\begin{pmatrix}
A'x & \geq & b' \\
-x_k & \geq & 0
\end{pmatrix}
\lor
\begin{pmatrix}
A'x & \geq & b' \\
x_k & \geq & 1
\end{pmatrix}
\] (3.25)

In fact, let us make life even easier and simply drop the notation \( A' \) and \( b' \). We will use the original notation, \( A \) and \( b \), but implicitly assume that \( Ax \geq b \) includes the non-negativity constraints \( x \geq 0 \). So (3.25) becomes simply:

\[
\begin{pmatrix}
Ax & \geq & b \\
-x_k & \geq & 0
\end{pmatrix}
\lor
\begin{pmatrix}
Ax & \geq & b \\
x_k & \geq & 1
\end{pmatrix}
\] (3.26)

To find an inequality \( \alpha x \geq \beta \) which is implied by this disjunction, and at the same time is maximally violated by the current solution \( \bar{x} \), we need to find some corresponding non-negative multipliers \((u, v) \in \mathbb{R}^m \times \mathbb{R}\) for the first disjunct

\[
u \times \begin{pmatrix}
Ax & \geq & b \\
-x_k & \geq & 0
\end{pmatrix} \Leftrightarrow \begin{pmatrix}
u Ax & \geq & ub \\
u x_k & \geq & 0
\end{pmatrix}
\] (3.27)
and \((v, v) \in \mathbb{R}^m \times \mathbb{R}\) for the second disjunct

\[
\begin{align*}
\mathbf{v} \times \left( \begin{array}{c} A \mathbf{x} \geq \mathbf{b} \\ x_k \geq 1 \end{array} \right) & \iff \left( \begin{array}{c} \mathbf{vA} \mathbf{x} \geq \mathbf{v} \mathbf{b} \\ v x_k \geq v \end{array} \right) \\
\end{align*}
\]

such that the resulting surrogates dominate \(\alpha \mathbf{x} \geq \beta\). In other words, we need to solve the following linear program:

\[
\begin{align*}
\min \quad & \alpha \mathbf{x} - \beta \\
\text{s.t.} \quad & \alpha - u A + v e_k \geq 0 \\
& -\beta + ub \geq 0 \\
& \alpha - v A - v e_k \geq 0 \\
& -\beta + vb + v \geq 0 \\
& u \geq 0 \\
& u \cdot 1 + v \cdot 1 + u + v \leq 1 \\
& \alpha, \beta \in \mathbb{R}^n \\
\end{align*}
\]

where \(e_k\) is the \(k^{th}\) unit vector. Note that in order to guarantee that an optimal solution to \((\text{CGLP}_k)\) exists, we have also added the bounding constraint \(\sum_i (u_i + v_i) + u + v \leq 1\), which is the same as (3.20) proposed by Ceria and Pataki [26].

As we have already seen in (3.21), solving this system yields a cut \(\alpha \mathbf{x} \geq \beta\), with

\[
\alpha_j = \begin{cases} 
\max \{ u A_j , \ v A_j \} & j \neq k \\
\max \{ u A_j - u , \ v A_j + v \} & j = k 
\end{cases}
\]

and

\[
\beta = \min \{ u b , v b + v \}
\]

where \(A_j\) is the \(j^{th}\) column of matrix \(A\).

**Strengthening cuts using integrality conditions**

We have so far focused on strengthening the derived inequality by finding better multipliers \(u\) using the linear program (CGLP). However, a further strengthening is possible, by exploiting the integrality conditions on variables. For instance, inequality (3.21) can be
further strengthened using this technique, as shown by Balas and Jeroslw[16]. This can be done by replacing each coefficient $\alpha_j$ with

$$\alpha'_j = \min \{ uA_j + u_0[m_j], vA_j - v_0[m_j] \}$$

(3.29)

where

$$m_j = \frac{vA_j - uA_j}{u_0 + v_0}$$

Note that this strengthening can be applied to more general disjunctions (i.e. not necessarily two term disjunctions, or involving just 0-1 variables). However, in that case, the expression for $m_j$ is no longer a closed form, but has to be computed procedurally, in $O(p)$, where $p$ is the number of terms in the disjunction. For details, see Balas and Jeroslw[16].

**Lift and project cuts without lifting and projection**

Since the size of the cut generating linear program can be quite large relative to the size of the original disjuncts, solving it may prove too expensive for practical purposes. In fact, the experiments of Balas et al[15] showed that the most efficient way to generate cuts using (CGLP$_k$) is to stop short of solving it to optimality. This was achieved by ignoring the multipliers (variables of (CGLP$_k$)) that correspond to constraints of $Ax \geq b$ not tight at the optimum $x$ (except for the lower and upper bounding constraints on the 0-1 variables). The cuts were then lifted to the full $x$ space using the closed form expression (3.22).

However, even when cuts are generated in a subspace as described above, this method still remains computationally challenging, due to the large size of (CGLP$_k$). Recent work by Balas and Perregaard[8] showed that it is in fact possible to compute the coefficients of $\alpha$ and $\beta$ without actually solving the linear program (CGLP$_k$). The implication of this result is that we no longer need to lift the problem, by formulating it as a linear program, into the much higher dimensional space of $(u, v, u, v, \alpha, \beta) \in \mathbb{R}^m \times \mathbb{R}^m \times \mathbb{R} \times \mathbb{R} \times \mathbb{R}^n \times \mathbb{R}^n$ and then project it back onto the $x \in \mathbb{R}^n$ space, but rather obtain the same inequalities directly in the $x$ space. By performing certain pivots in the original simplex tableau, we can essentially simulate one or more pivots in the higher dimensional (CGLP$_k$). In other
words, we can determine the reduced costs of the dual multipliers $u$ and $v$ in $(CGLP_k)$ by simply looking at a closed formula based on the current simplex tableau in the $x$ space. Therefore, we can simply start with the original (optimal) LP solution $\bar{x}$ and perform a sequence of pivots while monitoring what happens to these reduced costs. As long as there is at least one negative reduced cost, this means that we could further improve the objective function $\min \alpha \bar{x} - \beta$ of $(CGLP_k)$, which in turn means that we can improve the distance by which the inequality cuts off $\bar{x}$. Due to this correspondence, the cut obtained in the end is exactly the cut we would have obtained by actually doing the pivoting in (i.e. solving to optimality) the larger $(CGLP_k)$. This procedure is illustrated in Figure 3.1. More detailed steps of the algorithm can be found in Section A.1.

\begin{algorithm}
\textbf{Lift and project} $(k)$

1. $\text{optimal} \leftarrow \text{false}$ /* will be set to true when we have an optimal lift-and-project cut from row $k$ */

2. \textbf{while} (not optimal) \textbf{do}

3. \quad /* Are there negative reduced costs corresponding to the dual multipliers in $CGLP_k$? */

4. \quad if ($i^* \leftarrow \text{find cut improving pivot row} (k)$) then

5. \quad \quad $j^* \leftarrow \text{find best pivot column} (k, i^*)$

6. \quad \quad \text{change basis} $(i^*, j^*)$

7. \quad \quad else

8. \quad \quad \quad $\text{optimal} \leftarrow \text{true}$ /* we are done, the cut is optimal */

9. \quad \textbf{end if}

10. \textbf{end while}

11. \quad $(\alpha, \beta) \leftarrow \text{create intersection cut} (\tilde{A}, k)$ /* $(\alpha, \beta)$ is now a lift-and-project cut */

12. \quad $\text{strengthen cut} (\tilde{A}, k, \alpha)$ /* apply integrality strengthening */

\textbf{return} $(\alpha, \beta)$

\end{algorithm}

Figure 3.1: Generating lift-and-project cuts in the lower dimensional space

\section*{Two term simple disjunctions}

Let us start our discussion of this algorithm by looking at the simple disjunction which is the one used to derive lift and project cuts, namely $(x_k \leq 0 \lor x_k \geq 1)$. This belongs to a very useful special class of disjunctions, namely two term simple disjunctions. They have the general form:

$$ax \leq a_0 \lor bx \geq b_0$$

(3.30)
where \(a \in \mathbb{R}^n\), \(b \in \mathbb{R}^n\) and \(x \in \mathbb{R}^n_+\). When the two hyperplanes are parallel (that is, \(a = b\) and \(b_0 = a_0 + \epsilon\), where \(\epsilon > 0\)), these disjunctions are called *split disjunctions*. An important special case of split disjunctions is when \(a, b, a_0\) and \(b_0\) are integer and \(\epsilon = 1\). This case is particularly interesting because there are no integer solutions between the hyperplanes, so by applying them to an integer program we lose no feasible points.

**Split disjunctions and branching**

Branching on an integer constrained variable \(x_k\), whose value \(\bar{x}_k\) in the solution of the LP relaxation \(P = \{ x \in \mathbb{R}^n_+ : Ax \geq b \}\) is fractional, is equivalent to adding the split disjunction:

\[
    x_k \leq \lfloor \bar{x}_k \rfloor \lor x_k \geq \lfloor \bar{x}_k \rfloor + 1
\]

(3.31)
to the original formulation of the problem. In a typical branch and bound algorithm, however, the disjunction cannot be added explicitly, and it is instead enforced implicitly by creating two branches to be explored independently, one for each term of the disjunction. Since the language of integer programming does not include the \(\lor\) logical connective, if we want to take advantage of its semantics (other than enforcing it through branching), we have to derive an implication of this disjunction that takes the form of a linear inequality.

Note that (3.31) does not exclude any integer solution that is feasible for the original problem. Indeed, its job is limited to preventing the fractional value \(x_k = \bar{x}_k\), and in fact any other fractional value \(\lfloor \bar{x}_k \rfloor < x_k < \lfloor \bar{x}_k \rfloor + 1\), from ever being considered again. This means that any implication we derive directly from this disjunction will be valid for the convex hull of integer points that satisfy the original problem.

**Intersection cuts from the current simplex tableau**

A good point to start is the current LP solution, which is readily available in the simplex tableau. Let us extract the expression of \(x_k\) from the simplex tableau corresponding to the current solution \(\bar{x}\)

\[
    x_k = \bar{a}_{k0} - \sum_{j \in J} \bar{a}_{kj} s_j
\]

(3.32)
and plug it in (3.31). We obtain the following equivalent disjunction, which enforces
\[
\tilde{a}_{k0} - \sum_{j \in J} \tilde{a}_{kj} s_j \in \mathbb{Z}: \\
\tilde{a}_{k0} - \sum_{j \in J} \tilde{a}_{kj} s_j \leq \lfloor \tilde{x}_k \rfloor \lor \tilde{a}_{k0} - \sum_{j \in J} \tilde{a}_{kj} s_j \geq \lceil \tilde{x}_k \rceil + 1
\]  

(3.33)

We can now derive an inequality which is implied by this disjunction, by first changing the sign of the first disjunct, replacing \(\tilde{a}_{k0} - \lfloor \tilde{x}_k \rfloor\) with \(f_{k0}\) to simplify the notation, and then using the inference rule (3.7):

\[
\begin{align*}
\sum_{j \in J} \tilde{a}_{kj} s_j & \geq f_{k0} \\
\sum_{j \in J} -\tilde{a}_{kj} s_j & \geq 1 - f_{k0} \\
\sum_{j \in J} |\tilde{a}_{kj}| s_j & \geq \min\{f_{k0}, 1 - f_{k0}\}
\end{align*}
\]

(3.34)

Note that the current LP solution \(\tilde{x}\) indeed violates this inequality, because the nonbasic variables \(s_j\) are all zero, while \(0 < f_{k0} < 1\) (because \(\tilde{a}_{k0} = \tilde{x}_k\) is the value of \(x_k\) in the current solution and \(\tilde{x}_k\) is fractional). Since we were careful to construct (3.34) such that it remained valid for the convex hull of integer feasible points, we have essentially obtained a cut (a hyperplane that separates the current infeasible solution \(\tilde{x}\) from the set of feasible points). Because we derived it by intersecting the disjunction (3.31) with the polyhedral cone obtained by relaxing \(P\) (i.e. dropping all the nonbasic inequalities), this cut is called an intersection cut (see Balas[9]), and its coefficients can be computed in \(O(n)\) from the \(k^{th}\) row of the simplex tableau:

\[
\begin{align*}
\alpha_j &= |\tilde{a}_{kj}| \quad \forall j \in J \\
\beta &= \min\{f_{k0}, 1 - f_{k0}\}
\end{align*}
\]

(3.35)

Since \(f_{k0} > 0\), we can use Remark 3.1.7 to normalize the disjunction before taking the cut in (3.34), and we obtain the following (generally) stronger cut:

\[
\begin{align*}
\alpha_j &= \max\left\{\frac{\tilde{a}_{kj}}{f_{k0}}, -\frac{\tilde{a}_{kj}}{1 - f_{k0}}\right\} \quad \forall j \in J \\
\beta &= 1
\end{align*}
\]

(3.36)

Note that this is not necessarily the only (or the best) normalization. For example, one that could be (computationally) more advantageous is obtained by multiplying the first
inequality with $1 - f_{k0}$ and the second with $f_{k0}$. We get:

$$\begin{align*}
\alpha_j &= \max \{ \tilde{a}_{kj}(1 - f_{k0}), -\tilde{a}_{kj}f_{k0} \} \quad \forall j \in J \\
\beta &= f_{k0}(1 - f_{k0})
\end{align*}$$

(3.37)

**Expressing the cut in terms of the original variables**

There is one more step we need to complete in order to be able to add this cut to the original model as an inferred inequality. We need to make sure that we have the cut expressed in terms of the original model variables. As it is now, $\alpha s_j \geq \beta$, we have no guarantee that the variables $s_j$ appearing in the expression of (3.34) are all part of the original problem description (they could also be slack or surplus variables added by the simplex algorithm when it relaxed the problem to eliminate the $\leq$ and $\geq$ relational operators and transform it into a system of equations). So we now need to run Gaussian elimination again, this time to eliminate these extra variables (i.e. express them in terms of the originals). To perform this substitution of variables, note that originally, for each inequality $ax \leq b$ ($ax \geq b$), the simplex algorithm added a slack (surplus) variable $s$ to transform it into the equality $ax + s = b$ ($ax - s = b$). So, for those nonbasic variables $s_j$ that are not part of the original problem, we can simply replace $s_j$ with the equation of the original row to which they were added as slack (surplus). That is, $s_j = b^i - a^i x$ ($s_j = a^i x - b^i$) for some row $i$. This way we obtain a cut $\alpha' x \geq \beta'$ expressed solely in terms of the original variables and we can safely add it to the model as an inequality implied by the original ones (and, of course, by the decisions we have made so far).

**Special case: cuts from disjunctions on binary variables**

Binary variables are a very useful modeling device in integer programming. For such variables $x_k \in \{0, 1\}$, we have that $[\tilde{x}_k] = 0$ in (3.34), so the intersection cut obtained by applying the disjunction $x_k \leq 0 \lor x_k \geq 1$ has the same coefficients as (3.35), but a simpler right hand side: $\beta = \min\{\tilde{a}_{k0}, 1 - \tilde{a}_{k0}\}$. When derived at the end of the pivoting procedure shown in Figure 3.1, this cut is a deepest *lift and project* cut (as shown by BALAS AND PERREGAARD[8]).
Strengthened lift and project cuts

The precise correspondence between the bases of the lower dimensional simplex tableau and those of the corresponding CGLP tableau enables us to find deepest lift and project cutting planes without going through the trouble of explicitly setting up and solving (CGLP_k) for each binary variable x_k which has a fractional value in the optimal LP solution. When and optimal solution to (CGLP_k) is found, through either procedure, we know that the cut can no longer be improved by finding better multipliers u and v (in other words, by adding multiples of inequalities in Ax ≥ b to either term of the simple disjunction x_k ≤ ⌊x̅_k⌋ ∨ x_k ≥ ⌊x̅_k⌋ + 1). However, a further strengthening is possible by using integrality conditions on the remaining variables, as shown in Section 3.1.3. This, in fact, is the way to obtain mixed integer Gomory cuts from the optimal simplex tableau: we first derive an intersection cut from the tableau, without performing any additional pivots, as shown in (3.35) (or the stronger version (3.36)) and then apply integrality strengthening to it. On the other hand, the procedure outlined by Balas and Perregaard [8] for finding lift and project cuts, performs first a sequence of pivots and at the very end applies integrality strengthening to the resulting cut. This leaves open the following question: would it not be possible to obtain a better cut by mixing the pivots with integrality strengthening steps? This is the question we address in what follows.

Disjunctive modularization

Balas [13] calls the strengthening procedure (3.29) disjunctive modularization. In order to implement it, we can modify the procedure of Balas and Perregaard [8] in the following way. Instead of pivoting all the way to optimality of (CGLP_k), and then applying disjunctive modularization to the resulting cut, we will apply it to the row of the simplex tableau corresponding to x_k after each pivot. This way, we effectively combine the two cut strengthening methods: surrogates (represented by pivots) and integrality (represented by disjunctive modularization (3.29)).

Cut generation procedure

The idea of the procedure is the following. We start from the optimal solution x of
the LP relaxation (i.e. a vertex of the relaxed feasible region), derive an intersection cut from each fractional component $\bar{x}_k$ of $\bar{x}$ and try to improve that cut by pivoting on the original LP tableau, using rules for pivot selection derived from the properties of $\text{(CGLP}_k\text{)}. The original procedure of Balas and Perregaard\textsuperscript{[8]} generates lift and project cuts by first applying several rounds of strengthening by combination (i.e. pivots) and a final strengthening by integer rounding. We now use a modified version in which we apply the two strengthening methods in an interleaved manner: we apply integer rounding immediately after each pivot. Moreover, the intermediate (strengthened) cut is added to the model, which means that subsequent pivots (i.e. strengthening by combination) will take it into account. The process is repeated as long as there are negative reduced costs for the variables $u$ and $v$ of $\text{(CGLP}_k\text{)}$.

Note that our goal is not to maintain feasibility and improve the objective function of the LP (which is essentially the goal of pivoting in the traditional simplex method), but to obtain a better cut, by stepping around (and possibly outside) the feasible region of LP. Therefore, our pivots will quite likely take us to points in space where one or more constraints of the original LP don’t hold. For this reason, we need to save the optimal basis of the LP before starting generating cuts and restore it when we are done with cut generation, so that the integer programming solver does not get confused when trying to make search decisions.

**Experimental results**

We ran experiments the evaluate the effectiveness of interleaving the two cut strengthening methods. For these experiments, we used the latest version of the MIPLIB library of benchmark problems, MIPLIB 2003. Table A.1 shows the characteristics of the problems used: the number of rows and columns in $A$, the number of binary components and the number of integer (non-binary) components of $x$. All results reported here were obtained using XPRESS-MP version 15.3. In order to evaluate the impact of the resulting lift and project cuts in the cleanest possible way, we turned off the presolve and cut generation functions of XPRESS. For details on both the algorithm and on the results of these experiments, please see Appendix A.1.
3.2 Constraint programming

The theory of constraint programming (CP) represents a significant departure from the philosophy of its predecessors. We are no longer restricted to a closed language, as it was the case with linear, mixed integer linear and disjunctive programming, but we can now freely define new operators (constraints), with their own semantics and possibly new inference rules. The most obvious such operator would be $\neq$, which now allows us for the first time to reason about linear disequations (if you remember our discussion of linear and mixed integer linear programming, such reasoning is not possible within the framework of those theories). There are already several hundred such operators, some more frequently encountered than others. The interested reader can find more information about these operators (global constraints in the CP community) in the catalog maintained by Beldiceanu\textsuperscript{[20]}.

3.2.1 Modeling and solution process

The fact that constraint programming allows a higher degree of modeling flexibility makes it an attractive alternative to linear and integer programming. However, before a new operator can be added to the language, it must be equipped with specialized inference algorithms in order to be useful in deriving proofs. Such inference algorithms are commonly referred to as filtering algorithms, or domain reduction algorithms in the CP community. For many operators, there may exist multiple such algorithms, with varying complexity and performance guarantees. For instance, some algorithms guarantee completeness (that is, after they are applied, no values are left in the domains of their variables which do not belong to some feasible solution). This property is known as hyper-arc consistency. Others do not provide such guarantees, but execute much faster, and are therefore preferable under certain circumstances.

Constraint solvers employ a seemingly different approach than integer solvers, in the sense that at every step of the process, each constraint is responsible for drawing inference in isolation. In integer programming, inference is typically obtained from a collection of constraints and from the objective function, considered together. When no more inference
is possible, the search process is invoked. This time, the process is similar to the one used in integer programming: they both share a divide and conquer type of strategy. The CP community, however, was far more active in this area than the IP community, and so there are many more variants of divide and conquer search encountered in CP than in IP, such as limited discrepancy search (see Harvey and Gingsberg\cite{75}) and first-fail search (see Haralick and Elliott\cite{42}).

One important drawback of constraint programming, when compared with integer programming, is that it cannot generate useful bounds on the value of an objective function, if one is present. In integer programming, the simplicity of constraints allows the solver to automatically generate a convex reformulation of the problem (by linearizing the integrality constraints), which can then be used to compute a bound on the value of the objective. CP constraints do not offer this flexibility, and this is the main reason why CP-based methods have not been as successful for optimization problems as IP-based methods. However, when dealing with pure feasibility questions, CP solvers are typically much more effective than IP solvers, since they have access to more information about the nature of problem (and therefore its solutions) than pure IP models can provide.

This contrast of capabilities, together with the fact that parts of their solution processes overlap, makes the combination of the two techniques desirable. This is the starting point of our discussion of a unifying framework, in which these two (and other) methods can work together, at each step of the solution process, in order to take advantage of their complementary and common strenghts.
Chapter 4

A unifying framework

The idea of a unifying framework for optimization has been around for over a decade, but very few attempts have been made at providing a description of its major components and their interaction. A notable exception is the recent book of Hooker\cite{50}, which builds on some of the earlier work of Chandru and Hooker\cite{28}, Hooker\cite{48} and Aron, Hooker and Yunes\cite{6} to provide a consistent view of optimization techniques as they are used in mathematical and constraint logic programming. The framework we describe here has evolved from a joint work with Hooker and Yunes\cite{6}, and it provides the basic blocks for a modeling language of metaconstraints, as well as the mechanisms that a generic metasolver needs in order to solve a problem according to the specification given by the metaconstraints. The solution process employed by this framework is an interleaving of two fundamental steps, search and inference, each of which is reflected at the modeling level in the language of metaconstraints. Inference can be drawn using multiple techniques from (a) individual constraints (similar to domain reduction in constraint programming or bound preprocessing in integer programming) or (b) groups of constraints (similar to relaxation solving in mathematical programming). Search is performed using constraint selection rules and search direction selection rules, which can be statically specified by the modeler and dynamically prioritized by the inference component during the solution process.
4.1 Modeling

A key decision in optimization is the choice of a modeling language. Modeling must be done at a high enough level so that the problem statement appears clear to the modeler and at the same time provides sufficient structural information to the underlying solver. We saw earlier that in order to efficiently prove that the pigeonhole problem does not admit solutions if there are more pigeons than holes, we needed to state the problem in a language whose semantics offered powerful enough tools for such inference. We needed to communicate to the solver that it should use counting arguments. In this case, we reformulated the problem from a statement in propositional logic to one in 0-1 integer programming, and later to a continuous linear programming problem. It happened that in this case the translation was straightforward. The difficulty, however, is that we cannot efficiently represent any arbitrary problem in propositional logic. So we need a more expressive language, but which still allows us to easily reformulate problems into perhaps continuous linear programming, or even other paradigms.

Representability

One language which could easily provide such immediate reformulation rules is the language of mixed integer programming. However, mixed integer programming itself is not expressive enough to allow one to efficiently state all the problems one might need to solve. In fact, Jeroslow\textsuperscript{[55]} showed that the expressiveness of mixed integer programming is surprisingly limited, by proving that a set $S \subset \mathbb{R}^n$ is mixed integer representable if and only if it is a finite union of polyhedra whose set of recession directions coincide. In other words, even simple statements such as $”x$ is different from 3” ($x \neq 3$) have no representation in mixed integer programming. Constraint programming, on the other hand, provides an open language, in which problem characteristics can be encoded into new constraints, as necessary. Thus, an unrestricted combination of theories can be used when deriving inference, which is a very desirable property. This is the model we want for a language of metaconstraints, since it gives us the flexibility to add specialized inference algorithms to these constraints, or create new constraints when the existing ones are not expressive enough.
4.1.1 Metaconstraints

Informally speaking, a metaconstraint is a relation. It operates on the variables of a problem and produces a true or false answer depending on whether there exists a replacement of the variables with constants which turns the relation into a true statement. A metaconstraint can itself be viewed as an independent problem and considered in isolation. This is an important observation, because it tells us that even if we have a very complex problem, consisting of a large set of metaconstraints, proving that the problem represents a false statement (that is, it has no solution) could be as easy as proving that any one of these metaconstraints, or subproblems, is false (infeasible). This in fact is the approach taken in constraint programming. The solution process there is based solely on this type of infeasibility proof: the specialized inference algorithm(s) of each constraint are queried in isolation, and if one of them responds with false, the problem itself is false. What constraint programming notably lacks is the second type of inference, which is mostly used in mathematical programming, namely inference via reformulation. Besides providing additional tools for inference which are not available in the original language, reformulation has the great advantage that, if constraints are reformulated into a common language, then we could draw inference based on all at once, rather than querying each of them separately, as it is the case in constraint programming. This way we obtain a more global view of the problem, which may not necessarily be available in the original language. This global view is actually what’s behind Hooker’s choice to distinguish between inference and relaxation, since the relaxation is precisely a collection of constraints that have been reformulated into the same language.

Multiple reformulations

It should be clear by now that it is useful to equip metaconstraints with translation rules, preferably rules that translate into some common language. In fact, not only can we associate with a metaconstraint one set of translation rules for each given target language, but we can create alternative translations of that metaconstraint into the same language. To see why this is advantageous, let us examine the following example.
Suppose that we want to reformulate the following metaconstraint
\[ \sum_{i=1}^{n} |x_i| \leq 1 \] into an integer program. As it is stated, this constraint cannot be used in an integer programming solver, because the "absolute value" (|x|) is not part of the language of integer programming. One plausible reformulation would be to enumerate all possible cases of signs. For example, for \( n = 2 \), the constraint
\[ |x_1| + |x_2| \leq 1 \]
would be reformulated as the system of four inequalities
\[
\begin{align*}
-x_1 - x_2 & \leq 1 \\
-x_1 + x_2 & \leq 1 \\
x_1 - x_2 & \leq 1 \\
x_1 + x_2 & \leq 1
\end{align*}
\] representing all possible cases. Clearly, this is not a desirable representation for practical purposes, as it requires all the \( 2^n \) inequalities in the general case (see Jeroslow[54]).

However, an alternative reformulation is possible, with a much more compact representation. If we define a new variable \( y_i \geq 0 \), then we can write a reformulation of \( C_1 \) which is linear in size:
\[
\begin{align*}
\sum_{i=1}^{n} y_i & \leq 1 \\
x_i & \leq y_i \quad \forall i \\
-x_i & \leq y_i \quad \forall i
\end{align*}
\] Note that this new model is much smaller in size than \( C_1 \rightarrow R_1 \), even though it actually requires more variables.\(^1\)

**Constraint syntax**
We can now define a new metaconstraint based on \( C_1 \) which knows how to reformulate itself in the language of integer programming (and therefore automatically into linear programming).

\(^1\)So it is not always a good idea to stick to models with fewer variables!
In doing this, we implicitly assume that the implication $C_1 \rightarrow R_2$ is known to the language, via a keyword which links to the special translation procedure outlined above for the reformulation of the metaconstraint ($C_1$) into ($C_1 \rightarrow R_2$). This procedure is stored in a module, which does not need to be seen or used directly by the modeler. Upon reading the model, the metasolver brings in the module and uses it to obtain the system described by ($C_1 \rightarrow R_2$).

Note that there is nothing to prevent us from reformulating the constraint in multiple ways. For instance, we could declare it as in Figure 4.2. However, this does not make much sense and would not be useful at all in this case, for two reasons. One, we saw that the first reformulation we came up with, namely ($C_1 \rightarrow R_1$), had an exponential number of constraints. Secondly, even if it had a more reasonable size, it would still not be of much use since both ($C_1 \rightarrow R_1$) and ($C_1 \rightarrow R_2$) translate the constraint into the same language, which is somewhat redundant. The idea is to be able to obtain reformulations in a variety of languages, and not multiple reformulations into the same language. However, of course, having more than one reformulation into a given language allows us to choose the one better suited to the problem at hand.

---

$$C_1 \{$$
\[ \text{sum i of } | x[i] | \leq 1 \]
\[ \text{infer using } C_1 \rightarrow R_2 \]
$$\}$$

Figure 4.1: Modeling: example of a metaconstraint with reformulation rules

$$C_1 \{$$
\[ \text{sum i of } | x[i] | \leq 1 \]
\[ \text{infer using } C_1 \rightarrow R_1, C_1 \rightarrow R_2 \]
$$\}$$

Figure 4.2: Modeling: metaconstraint with multiple reformulation rules
4.2 Solution process

We have argued so far that the solution process in optimization is a combination of two fundamental steps: search and inference. These two steps are interleaved and rely on each other in order to advance towards a solution of the problem or prove that none exists. The advancing process is done by the search component, while the verification of feasibility is handled by the inference part. Figure 4.3 gives an outline of this solution process.

```
Algorithm solve (P)
  I ← ∅  /* I is the set of inferences made by the inference component */
  D ← ∅  /* D is the set of decisions made by the search component */
  while (∃ unexplored (variable, constant) substitutions) do
    I ← infer (P, I, D)
    if (false) then
      /* The inference component decided that we made some wrong decisions along the way, so we must backtrack */
      (D, I) ← backtrack (P, I, D)
    else
      if (P is TRUE) then
        /* All variables have been eliminated and we have determined that P is true (consistent) */
        return true
      else
        /* We are not done yet, but no further inference is possible, so we'll have to make some new decisions */
        D ← search (P, I, D)
      end if
    end if
  end while
  return false  /* All possible eliminations were tried and none turns P into a true statement */
```

Figure 4.3: General algorithm to solve an optimization problem P

The algorithm starts with the statement of $P$ and searches over all possible ways to eliminate variables from $P$. The search is kept in check by the inference module, which is called at every stage in order to (a) detect wrong decisions made earlier, and (b) help the search avoid new ones. The state of the algorithm is kept in two sets, $I$ and $D$. The set $I$ contains all the inferences made by the inference module up to this stage. Similarly, $D$ is the set of all decisions made by the search module up to now. Note that new elements are added to $I$ only by the inference module, while $D$ is augmented only by the search
module. However, whenever a dead end is detected during inference (that is, the inference module decides that the combination of the original statement $P$ with $I$ and $D$ leads to a false statement), both sets are modified by the backtrack module. This module has the role of reversing some of the decisions made by the search module, and with them the inferences from $I$ that were made on the basis of these decisions. The algorithm ends when all possible decisions have been either explored by the search or ruled out by the inference module. The decision that $P$ is true is made when at least one elimination was found which leads to a true statement and either we exhausted all other eliminations or inference tells us that there are no others which could provide a better conclusion (in the case of optimization, no other eliminations will provide a solution with a better value).

Note that there is no reference to an objective function. This is because the only role of the objective function in optimization is to help the inference process, by using bounding arguments, and so all references to such a function are captured inside the inference module.

### 4.2.1 Search

Search becomes useful when inference alone cannot decide on the feasibility or infeasibility of the problem. In general, search takes the form of variable elimination, in the sense that it attempts to remove variables from the problem statement by replacing them with constants from their corresponding domains. However, other ways to conduct search are possible, and it may be the case than sometimes a better decision is not to eliminate a variable, but instead to just restrict the number of possible replacements we can make for that variable. This is because sometimes inference is able to determine infeasibility even if the variable is still in the problem, based only on its restricted domain.

The typical search process is mostly unguided. Inference may try to suggest ways for the search to proceed, but this is in general not well understood and poorly used in most existing solvers. In integer programming, the way search proceeds in most solvers is by looking at the solution of the relaxation, which in general has little or nothing to do with what we are in fact looking for. There are examples in which such a solution is arbitrarily far from any feasible integer solutions, and using it as a guide for the search is
indeed a very poor decision. The integer solver will attempt to search so that it corrects a problem it sees in the solution of the relaxation. It does so by choosing some variable \( x_k \) which does not have an integer value in the relaxation and creating a disjunction of the type \( x_k \leq \lfloor \tilde{x}_k \rfloor \lor x_k \geq \lceil \tilde{x}_k \rceil \), where \( \tilde{x}_k \) is the value of \( x_k \) in the solution found by the continuous solver. This is a very weak way of searching, since there is no guarantee that either (1) \( x_k \) will receive an integer value, or (b) any of the variables which happened to be integral in the current relaxation solution will remain so.

**Search rules**

A better way to search is to exploit knowledge about the problem itself, and the best handle we have into the problem structure is the collection of metaconstraints. Let us illustrate this with a simple example (which happens to be useful also in the context of integer solvers). Consider the linear constraint

\[
x_1 + x_2 + x_3 \leq 1
\]

where \( x_i \in \{ 0, 1 \}, \forall i \) and suppose that one of the variables was assigned a fractional (non-integer) solution by the continuous solver during the inference process (remember that the solver attempts to bound the current problem using the objective function, and in the process it finds the most extreme point of the polyhedron - that point is the solution we are using here). One option is to create two search directions (branches), using the disjunction we mentioned above, in order to "avoid" this particular fractional solution in the future. A wiser choice would be to avoid all fractional solutions in the future, not just this one. We can do so by creating four search directions, each of which eliminates all three variables at once:

- direction 1: \( x_1 \leftarrow 0, \ x_2 \leftarrow 0, \ x_3 \leftarrow 0 \)
- direction 2: \( x_1 \leftarrow 1, \ x_2 \leftarrow 0, \ x_3 \leftarrow 0 \) (\( C_2 :: S_1 \))
- direction 3: \( x_1 \leftarrow 0, \ x_2 \leftarrow 1, \ x_3 \leftarrow 0 \)
- direction 4: \( x_1 \leftarrow 0, \ x_2 \leftarrow 0, \ x_3 \leftarrow 1 \)

This way of searching covers all feasible cases, and avoids potential fractional (and therefore infeasible) ones. It could therefore lead to a much shorter proof, although this generally depends, of course, on the rest of the problem.
$C_2 \{ \\
\quad \text{sum } i \text{ of } x[i] \leq 1 \\
\quad \text{search using } C_2 :: S_1 \\
\} \\
$

Figure 4.4: Modeling: example of a metaconstraint with search rules

So we could also equip this metaconstraint with a search rule, which we can later refer to when we use it in the statement of a problem. The resulting metaconstraint is shown in Figure 4.4. Just as it was the case with inference rules, the language is now aware of the keyword $C_2 :: S_1$, and it uses it to make search decisions based on this metaconstraint.

Note that this way of specifying search rules for individual constraints does not prevent us from using the already existing strategy from integer programming, namely adding the disjunction $x_k \leq \lfloor \bar{x}_k \rfloor \lor x_k \geq \lceil \bar{x}_k \rceil$, when the value $\bar{x}_k$ of $x_k$ is fractional in the continuous linear relaxation. Indeed, we can use a metaconstraint called $\text{integer}(x_k)$, whose search rule produces:

\begin{align*}
\text{direction 1: } & x_k \leq \lfloor \bar{x}_k \rfloor \\
\text{direction 2: } & x_k \geq \lceil \bar{x}_k \rceil \\
& (\text{int :: frac})
\end{align*}

The metaconstraint could then be declared as in Figure 4.5.

$\text{integer} \{ \\
\quad x \text{ is int} \\
\quad \text{search using int::frac} \\
\} \\
$

Figure 4.5: Modeling: example of a metaconstraint enforcing integrality of variables

**Rule selection**

Since a model is a collection of metaconstraints, and each metaconstraint has its own search rule, we must be able to decide which rule to use at a given point during the search. Obviously we can only use one, but which one we choose can have dramatic effects on the length of the search. The rule selection is performed dynamically by the underlying
metasolver. As pointed out earlier, the main task of inference is to come up with a proof of infeasibility. If it fails, there are still some positive side effects of the inference process, such as reduced variable domains, cutting planes, new bounds on the objective function, and so on. This additional information can be used in order to come up with recommendations for the search. One possible strategy, which is also the one we employ in our implementation (see Aron et al. [6]), is to evaluate how close to infeasibility each metaconstraint is (given the new information) and then select the one which is closest to infeasibility (or feasibility) in order to decide the next search direction. Once such a constraint is selected, the particular search decisions, and their order, are left to its search rule. In order to allow the modeler some flexibility in choosing what the inference module

```plaintext
search {
    create using least::\{C_2, C_3\}, most::\{C_1, C_4\}, first::all
    explore using best::bound::dive
}
```

Figure 4.6: Modeling: example of search module specification

should recommend, one must either allow procedural elements into the language, or at least provide some declarative means of choosing among the most common alternatives. In Aron et al. [6], we opt for the latter, since the goal is to provide a framework that is powerful enough while still easy to grasp and use. The statement in Figure 4.6 says that search should first consider the rules of constraints $C_2$ and $C_3$, and use the search rule of the least feasible of the two. If the two are feasible, then search will consider the next group of constraints, $C_1$ and $C_4$, and use the one which is closest to feasibility, as indicated by the keyword most (feasible). If both these constraints are feasible as well, then it selects the first infeasible (first) from the remaining constraints (all) to continue the search. If no infeasible constraint exists, a solution (and perhaps a new incumbent) has been found, so the search can now focus on some unexplored direction, as discussed next.

**Direction selection**
As it is clear from the previous examples, each search rule will generally add more than one direction to the search. Since only one such direction can be explored at a time, the rest are kept in a pool. This pool can be prioritized in various ways, which are indicated by keywords like `best::bound::dive`, or `best::bound::explore` (see Figure 4.6) representing commonly used pool management strategies. The search module always picks the direction which is at the top of this pool. Internally, the pool is stored as a multiple key priority queue, to allow ordering based on several characteristics, including search depth (number of decisions made from the original problem statement), objective bound, degree of infeasibility, etc. The main search procedure takes a very simple form, as shown in Figure 4.7, since most of the search related decisions are left to the constraints. This is the procedure being called in line 11 of Figure 4.3.

```
Algorithm search (P, I, D)
/* Find the one constraint which the inference engine has ranked first as most promising for search */
1 foreach (constraint group G_i of P) do
2   if (∃c ∈ G_i | c is infeasible according to I) then
3     select the first such constraint c and break
4   end if
5 end for

/* Generate new search directions based on the search rule of the selected constraint */
6 D ← D ∪ c.search_rule()

/* D now reflects the current state of the search, plus the new alternatives generated by the search rule of c */
7 return D
```

Figure 4.7: Search algorithm based on metaconstraint search rules

### 4.2.2 Inference

The inference module is called upon before and after a search decision is made. The role of this module is to decide whether the current state of the search is infeasible. While attempting to prove infeasibility, the module can use a variety of tools and techniques, as instructed by the metaconstraints. It may happen, and it is often the case, that inference
cannot (yet) conclude that the problem is infeasible, so further search is required. Even
when this is the case, important information becomes available as a result of attempting to
prove infeasibility, which can be used to further limit the number of possibilities available
to the search module.

Using multiple reformulations
As we have seen, each metaconstraint is equipped with special inference rules, all of which
essentially reinterpret the constraint in some theory and attempt to prove its infeasibility
using that theory. For instance, symbolic constraints such as \texttt{alldifferent}(x) can be re-
formulated as graph theoretical problems, and solved using theorems from graph theory
(see \texttt{R\textsc{\textregistered}EGIN}[69]). Others have better translations into \textit{continuous linear programming}, and
many can even be reformulated efficiently in multiple languages. We could, for instance,
draw inference from the \texttt{alldifferent}(x) constraint both by using the graph theoretical re-
formulation, as well as by reformulating it as a linear programming problem, although the
linear reformulation is not nearly as useful in practice for this constraint. However, it is
an instructive exercise.

The \texttt{alldifferent}(x) constraint states that the components of an n-dimensional vector
of discrete variables $x \in \{1, \ldots, m\}^n$ must be pairwise different. In other words, no two
$x_i$ and $x_j$ with $i \neq j$ can have the same value. We could start a reformulation process
by first stating the constraint using a simpler relation, \"\neq\" in place of the more complex
\"alldifferent\". Thus, we can write

$$x_i \neq x_j, \forall 1 \leq i < j \leq n$$

which is semantically equivalent to the original constraint. Since the \"\neq\" relation is not
part of the language of linear programming, we must replace it with inequalities. An
equivalent statement involving inequalities would be

$$x_i \neq x_j \leftrightarrow x_i < x_j \lor x_i > x_j$$

but this is still not representable as a linear program (neither \"<\" nor \"\lor\" are part of
that language). Fortunately, since $x_i$ and $x_j$ can only take discrete values, we can escape
the "<" relation and replace it with "≤", which is allowed in linear programming:

\[ x_i < x_j \lor x_i > x_j \leftrightarrow x_i \leq x_j - 1 \lor x_i \geq x_j + 1 \]

We still need to eliminate the "either-or" disjunction relation "\lor". One way to do this is to introduce additional variables into the model, one for each disjunct, to play the role of selectors for which disjunct is true. Let \( z_{ij} \in \{0, 1\} \) be such a variable to indicate which of the two inequalities \( x_i \leq x_j - 1 \) and \( x_i \geq x_j + 1 \) holds true. We can now write a linear system which is implied by the original constraint:

\[
\begin{align*}
    x_i - x_j + 1 & \leq mz_{ij} & \forall 1 \leq i < j \leq n \\
    x_j - x_i + 1 & \leq m(1 - z_{ij}) & \forall 1 \leq i < j \leq n
\end{align*}
\]

(\textit{aldiff} \rightarrow \textit{linear})

Whenever \( z_{ij} = 0 \), the system is equivalent to \( x_i \leq x_j - 1 \) (the second inequality becomes true since no two numbers in the set \( \{1, \ldots, m\} \) can be \( m \) units apart), and when \( z_{ij} = 1 \), it is equivalent with \( x_i \geq x_j + 1 \) (the first inequality becomes true). Thus,

```plaintext
alldifferent {
    aldiff(x)
    infer using linear::default, specialized::default
}
```

Figure 4.8: Modeling: alldifferent with reformulation in multiple languages

we can declare a metaconstraint \textit{alldifferent} using two different inference (reformulation) rules, as shown in Figure 4.8. The first reformulation (default linear) points to a procedure which creates \textit{aldiff} \rightarrow \textit{linear}. The second reformulation (specialized) of \textit{alldifferent(x)}, into a graph theoretical problem, converts the statement of alldifferent into a \textit{bipartite matching} problem. On one side of the graph are the variables \( x_i \), on the other side their possible values. Then alldifferent is infeasible if there is no matching that covers all variables. To determine that no such matching exists, the algorithm proposed by RÉGIN\textsuperscript{[69]} attempts to remove all edges from the graph that are not part of a maximum cardinality matching, using a theorem due to BERGE\textsuperscript{[21]}. This algorithm is what is invoked by the framework upon encountering the \texttt{specialized::default} rule in the definition of the alldifferent metaconstraint in Figure 4.8.
There is an interesting distinction to be made here between the two reformulations of the \textit{alldifferent}(x) metaconstraint, which in fact is behind Hooker\textsuperscript{[50]}'s view of relaxation and inference as being different. The graph theoretical reformulation, \textit{alldiff} $\rightarrow$ \textit{regin}, is a \textit{specialized} one, meaning that it was not intended to be used in combination with reformulations of other metaconstraints into graph theory (although it could, in theory, but nobody has looked in that direction so far). The other reformulation, into continuous linear programming, \textit{alldiff} $\rightarrow$ \textit{linear}, can in fact be used \textit{in conjunction with} reformulations of other metaconstraints as linear programs, in order to strengthen the inference by taking more aspects of the problem into account at the same time. Hooker's definition of a relaxation is based precisely on this property of the continuous reformulation: namely that it can collect implications from more than one constraint and attempt to prove infeasibility based on \textit{all} of them at once. In fact, the definition he uses starts from an alternative, equivalent viewpoint: it regards a relaxation as being obtained from the original problem by \textit{dropping} some of the original constraints form the problem statement. Regardless of the viewpoint, there is a clear advantage in using relaxations (as \textit{collections} of constraint implications), since in many cases it may be difficult to prove infeasibility by looking at constraints in isolation, but once two or more constraints are considered together, infeasibility becomes much easier to detect.

\textbf{Relaxation as a special case}

The framework proposed here includes the notion of relaxation as a special case, since it allows for reformulations of individual constraints to \textit{also} be collected into a common implication whenever the target theory allows it. In particular, this is done automatically for continuous linear reformulations, as well as reformulations into propositional logic. For specialized reformulations such as Régis's graph theoretical view of \textit{alldifferent}, inference is drawn only in isolation from each such reformulation at a time. In fact, this is also done for general reformulations: the method first attempts to prove that at least one of the individual reformulations is infeasible and \textit{only if} no infeasibility is detected, the globalized reformulation (which could be quite large for some problems) is considered. The outline of the inference algorithm is given in Figure 4.10.
Inference from relaxations

In order to take better advantage of relaxation as a special case of reformulation, we should instruct the underlying solver on what sort of inference it must draw from such relaxations. For instance, with a collection of linear constraints, we can derive cutting planes of various types and strengths, bounds, etc. For a collection of logical clauses, we can use resolution. The default behaviour is to use all the available techniques for a given theory, but this can be further specialized by adding another section to the model, as shown in Figure 4.9. This says that for the collection of linear implications of the problem (that is, the linear relaxation) inference should use cutting plane generators, and in particular Gomory mixed integer cuts (see Gomory[38]), lift-and-project cuts (the algorithm described in Chapter 3.1.3), mixed integer rounding cuts (see Marchand et al.[63]) and flow cover cuts. Furthermore, it should only keep the best 60% of the Gomory cuts and 10% of the lift and project cuts that are generated, ranked by the distance from the solution of the linear relaxation (this limitation ensures that subsequent inference steps will not take too long, which can happen when too many such cuts are being added to the formulation).

Guiding the search

The degree of feasibility of each constraint is determined by the inference module, which also provides a ranking of the constraints according to feasibility degrees. As shown in Figure 4.6, constraints can be grouped by priority, in which case the inference module ranks constraints in each group separately, to allow for even more specialized search decisions. Using this ranking mechanism, the search module can make a deterministic decision. This decision is based on as much information as we can obtain during the inference process and therefore reflects the problem structure as revealed by the metaconstraints.
Algorithm \texttt{infer} \((P, I, D)\)

This first loop is similar to what constraint programming solvers do - run inference on each constraint in isolation. However, the procedure described here is more general, because it allows multiple reformulations of the same constraint to be used.

\begin{verbatim}
/* Try each individual constraint reformulation and see if any comes back infeasible */
1 foreach (metaconstraint \(C_i \in P\)) do 
2    foreach (reformulation rule \(C_i \rightarrow R_i\)) do 
3        if ( \(I \leftarrow \text{infeasible} \ (C_i \rightarrow R_i, I)\) ) then 
4            return false 
5        end if 
6    end for 
7 end for 

The remaining part is similar to what mathematical programming solvers do - run inference on a relaxation of the entire problem. The procedure described here is more general, because it also allows multiple reformulations of the problem to be used.

/* No individual constraint could be proved infeasible, so we construct problem reformulations */
8 foreach (reformulation rule \(C_i \rightarrow R_i\)) do 
9    if (theory \(T_{R_i}\) of \(R_i\) allows global inference) then 
10       \(P^{T_{R_i}} \leftarrow P^{T_{R_i}} \cup R_i\) /* Add the reformulation \(R_i\) of \(C_i\) to relaxation \(P^{T_{R_i}}\) */ 
11    end if 
12 end for 

/* Try each problem reformulation now and see if any comes back infeasible */
13 foreach (reformulation \(P^{T_{R_i}}\) of \(P\)) do 
14    if ( \(I \leftarrow \text{infeasible} \ (P^{T_{R_i}}, I)\) ) then 
15        return false 
16    end if 
17 end for 

/* No infeasibility detected, so we just return the inferences made along the way */
18 return \(I\)
\end{verbatim}

Figure 4.10: Inference algorithm based on metaconstraint inference rules
4.3 Example: Production planning

Let us illustrate these concepts with a practical example, namely a production planning problem. The objective is to manufacture several products on a line of limited capacity $C$ so as to maximize net income. Each product must be manufactured in one of several production scale modes (small, medium, large), or not manufactured at all, and only a certain range of production quantities are possible for each product in each mode. Thus if $x_i$ units of product $i$ are manufactured in mode $k$, $x_i \in [L_{ik}, U_{ik}]$. The income $f_i(x_i)$ obtained from making product $i$ is linear in each interval $[L_{ik}, U_{ik}]$, which means that $f_i$ is a semicontinuous piecewise linear function (Figure 4.11):

$$f_i(x_i) = \frac{1}{U_{ik} - L_{ik}} [(U_{ik} - x_i)c_{ik} + (x_i - L_{ik})d_{ik}], \quad \text{if } x_i \in [L_{ik}, U_{ik}] \quad (4.1)$$

The mode $k = 0$, for which $[L_{i0}, U_{i0}] = [0, 0]$, corresponds to the case when product $i$ is not produced at all.

![Figure 4.11: Production planning: a semicontinuous piecewise linear function $f_i(x_i)$.](image)
4.3.1 Modeling

An integer programming approach to modeling this problem requires the introduction of some indicator (0-1) variables $y_{ik}$ to select the production mode of each product. The functions $f_i$ can be modeled by assigning weights $\lambda_{ij}, \mu_{ik}$ to the endpoints of each interval $k$, and so a linear model for this problem could look like the following:

$$\max \sum_{ik} \lambda_{ik} c_{ik} + \mu_{ik} d_{ik}$$

$$\sum_i x_i \leq C$$

$$x_i = \sum_k \lambda_{ik} L_{ik} + \mu_{ik} U_{ik}, \text{ all } i$$

$$\sum_k \lambda_{ik} + \mu_{ik} = 1, \text{ all } i$$

$$0 \leq \lambda_{ik} \leq y_{ik}, \text{ all } i, k$$

$$0 \leq \mu_{ik} \leq y_{ik}, \text{ all } i, k$$

$$\sum y_{ik} = 1, \text{ all } i$$

$$y_{ik} \in \{0, 1\}, \text{ all } i, k$$

(4.2)

**CP model**

However, a perhaps more natural way to model this problem is to use conditional constraints of the form $A \Rightarrow B$, which means that whenever the antecedent $A$ is true, the consequent $B$ must be enforced. Note that this is not the same as an implication, since it does not say that whenever $B$ is not true, $A$ must also be false. In this case, it simply acts as an on-off switch for constraint $B$ based on the truth value of $A$. Furthermore, the piecewise linear cost function $f_i(x_i)$ can be coded much as it appears in (4.1), so the new model looks like:

$$\max \sum_i u_i$$

$$\sum_i x_i \leq C$$

$$x_i \in [L_{ik}, U_{ik}] \Rightarrow u_i = \frac{1}{U_{ik} - L_{ik}} [(U_{ik} - x_i)c_{ik} + (x_i - L_{ik})d_{ik}], \text{ all } i, k$$

(4.3)
Note that no discrete variables are required, and the model is quite simple. This model can now be solved by branching on the continuous variables \(x_i\) in a branch-and-bound method. In this case the domain of \(x_i\) is an interval of real numbers. The search branches on an \(x_i\) by splitting its domain into two or more smaller intervals, much as is done in continuous global solvers, so that the domains become smaller as one descends into the search tree. The solver processes the conditional constraints (b) by adding the consequent to the constraint set whenever the current domain of \(x_i\) lies totally within \([L_{ik}, U_{ik}]\).

**A model based on metaconstraints**

Although the above model is a perfectly legitimate approach to solving the problem and might appear as quite natural initially, it does not fully exploit the problem's structure. Note that for each product \(i\), the point \((x_i, u_i)\) must lie on one of the line segments defined by consequents of constraint (b). *If the solver were aware of this fact, it could 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 (potentially) faster solution.* This can be accomplished by equipping the modeling language with a metaconstraint `piecewise` which models continuous or semicontinuous piecewise linear functions. A single piecewise constraint represents the constraints in (b) that correspond to a given \(i\). The model now becomes

\[
\begin{align*}
\max & \sum_i u_i \\
\sum_i x_i & \leq C \quad \text{(a)} \\
\text{piecewise} & (u_i, x_i, L_i, U_i, c_i, d_i), \text{ all } i \quad \text{(b)}
\end{align*}
\]

Here \(L_i\) is an array containing \(L_{i0}, L_{i1}, \ldots\), and similarly for \(U_i, c_i,\) and \(d_i\). Each piecewise constraint enforces \(u_i = f_i(x_i)\). The high level representation of this model, in compilable form, is shown in Figure 4.12. The key points in the model are the inference rules on the piecewise constraint (which is linearized using the default available algorithm and produces the convex hull (4.5) of feasible values of \(x_i\) and \(u_i\) given the current state of the search), the search rules that say we should branch on the least feasible piecewise constraint (and use the algorithm “three-split” in order to create new branches) and the branch selection rule which says that we should select the branch with the best bound and then dive from
there until a leaf is reached in the search tree. The degree of infeasibility of each piecewise constraint is calculated by measuring how far the values of $x_i$ and $u_i$ obtained from the linear solver are from the closest linear segment (piece) of the function.

maximize { 
    income = sum i of u[i]
}

capacity { 
    sum i of x[i] <= C
}

piecewise { 
    _piecewise(x[i],u[i],L[i],U[i],c[i],d[i]) forall i
    infer using specialized::default, linear::default
    search using three_split
}

infer { 
    linear using gomory::40 , mir::15
}

search { 
    create using least::{ piecewise }
    explore using best::bound::dive
}

Figure 4.12: Production planning: integrated model

### 4.3.2 Solution process

Let us suppose the solver is instructed to solve the problem by best::dive (a variant of branch and bound), which is defined as a direction exploration strategy in the search component of the model in Figure 4.12. 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 (necessary for inference on the linear reformulation, as well as for ordering new restrictions in a pool as indicated best::bound::dive) by solving a linear relaxation of the problem. This
relaxation is (4.5). The solver branches whenever a constraint is violated by the solution of the relaxation. The nature of the branching is dictated by the constraint that is violated.

**Specialized inference**

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 \( \min \left\{ b_i, C - \sum_{j \neq i} a_j \right\} \). 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 its solution is feasible in the original problem.

**Inference via reformulation**

The solver creates a linear relaxation at each node of the search tree by pooling linear reformulations of the various constraints. For example, it reformulates each constraint in (b) by generating linear inequalities to describe the convex hull of the graph of each \( f_i \), as illustrated in Figure 4.13. 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 \([L_i, b_i]\). The linear constraint (a) also contributes to the relaxation, by simply sending a reformulation of itself (identity). These relaxations, along with the domains, combine to form a linear relaxation of the entire problem:

\[
\begin{align*}
\max & \sum_i u_i \\
\sum_i x_i & \leq C \\
\text{conv(piecewise}(u_i, x_i, L_i, U_i, c_i, d_i)), & \text{all } i \\
a_i & \leq x_i \leq b_i, & \text{all } i
\end{align*}
\]

(4.5)

where \( \text{conv} \) denotes the convex hull description just mentioned.

The solver next finds an optimal solution \((\bar{x}_i, \bar{u}_i)\) of the relaxation (4.5) by calling a
linear programming solver. This solution will necessarily satisfy (a), but it may violate (b) for some product \(i\), for instance 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 Fig. 4.13, 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 optimization areas: (1) search by splitting intervals, from *continuous global optimization*, (2) domain reduction, from *constraint programming* and (3) polyhedral relaxation, from *integer programming*.

![Figure 4.13: Production planning: convex hull relaxation (shaded area) of \(f_i(x_i)\).](image)

### 4.3.3 Experimental results

We ran both the pure MILP model (4.2) and the above integrated model over 10 randomly generated instances with the number of products \(n\) ranging from 5 to 50. In all instances, products have the same cost structure with five production modes. In order to break symmetry, the models also include constraints of the form \(x_i \leq x_{i+1}\) for \(i \in \{1, \ldots, n-1\}\). The number of search nodes and CPU time (in seconds) required to solve each of the ten instances.
instances to optimality are shown in Table 4.1. As the number of products increases, one

| Number of Products | IP (CPLEX 9.0) | | Integrated |
|--------------------|----------------|------------------|
|                    | Searched nodes | Time (s) | Searched nodes | Time (s) |
| 5                  | 93             | 0.03         | 13             | 0.09     |
| 10                 | 423            | 0.12         | 40             | 0.35     |
| 15                 | 1321           | 0.34         | 33             | 0.58     |
| 20                 | 4573           | 1.14         | 53             | 1.00     |
| 25                 | 5105           | 2.43         | 19             | 0.30     |
| 30                 | 4504           | 2.10         | 43             | 0.82     |
| 35                 | 6089           | 3.30         | 33             | 1.11     |
| 40                 | 7973           | 4.06         | 36             | 1.15     |
| 45                 | 23414          | 14.72        | 40             | 1.76     |
| 50                 | 18795          | 9.45         | 47             | 1.77     |

Table 4.1: Production planning: search nodes and CPU time.

can see that the number of search nodes required by a pure MILP approach can be more
than 500 times larger than the number of nodes required by the integrated approach.
This can be explained by the fact that more powerful inference is applied before creating
new nodes, and furthermore the search process uses more specific problem knoweldge (by
exploiting the properties of the piecewise constraint). A MILP solver may find it very hard
(and many times impossible) to uncover this information from a purely linear model. The
integrated approach is also superior in all (but the three smallest) instances with respect
to CPU time. The fact that CPLEX can solve the smaller IP instances faster than the
integrated solver, despite the larger number of nodes explored, can be explained if we take
into account the additional bookkeeping that goes on underneath the integrated solver at
each node, like automatically reformulating (tightening the convex hull linearization of)
the piecewise constraint. In general, the integrated solver will spend significantly more
time at a node that a MILP solver does, but the hope is that this time is well spent
and will reduce the number of nodes (and therefore the total solution time) considerably.
This transpires quite clearly from Table 4.1, where the number of nodes required by the
integrated solver does not grow nearly as fast as it does for the IP solver (in fact, a nice
feature of this model seems to be that it results in a more robust behavior across problem
instances, whereas the IP model’s performance degrades significantly with problem size).
4.4 Example: Product configuration

In the second example we consider, the problem is to choose an optimal configuration of components for a product, such as a personal computer. For each component $i$, such as 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 instance, 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, $A_{ijk} < 0$ is used 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 $c_j$.

4.4.1 Modeling

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 $\sum_{ik} A_{ijk} q_{ik}$. Since this is a nonlinear expression, the variables $q_{ik}$ 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,

$$\min \sum_j c_j v_j$$

$$v_j = \sum_{ik} A_{ijk} q_{ik}, \text{ all } j \quad (a)$$

$$L_j \leq v_j \leq U_j, \text{ all } j \quad (b)$$

$$q_{ik} \leq M_i x_{ik}, \text{ all } i, k \quad (c)$$

$$\sum_k x_{ik} = 1, \text{ all } i \quad (d)$$

(4.6)
where each \( x_{ij} \) is a 0-1 variable, each \( q_{ij} \) is integer, and \( M_i \) is an upper bound on \( q_i \). If the modeling language accommodates specially ordered sets of nonbinary variables, the variables \( x_{ij} \) can be eliminated and constraints (c) and (d) replaced by a stipulation that \( \{q_{i1}, q_{i2}, \ldots\} \) is a specially ordered set of type 1 for each \( i \).

**A model based on metaconstraints**

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:

\[
\begin{align*}
\min & \sum_j c_j v_j \\
\text{s.t.} & \quad v_j = \sum_i q_i A_{ij} t_i , \text{ all } j 
\end{align*}
\]

(4.7)

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

A variable index is a very versatile modeling device. If an expression has the form \( u_y \), where \( y \) is a variable, then \( y \) is a variable index or variable subscript. The modeling system automatically decodes variably indexed expressions with the help of the element metaconstraint, which is frequently used in constraint programming. In this case the variably indexed expression has the indexed linear form \( ua_y \), where \( u \) is an integer variable and \( y \) a general discrete variable. An expression of this form is replaced with a new variable \( z \) and an additional constraint

\[
\text{element}(y, (ua_1, \ldots, ua_n), z)
\]

(4.8)

This constraint in effect forces \( z = ua_y \). The solver can now apply a domain reduction (or filtering) algorithm to (4.8) and generate a relaxation for it.

**4.4.2 Solution process**

Filtering for (4.8) is straightforward. If \( z \)'s domain is \([z, z']\), \( y \)'s domain is \( D_y \), and \( u \)'s domain is \( \{u, u + 1, \ldots, u'\} \) at any point in the search, then the reduced domains \([z', z']\),
$D_y'$, and $\{u', \ldots, \bar{u}'\}$ are given by

$$z' = \min \{z, \min_k \{a_k q\}\}, \quad \bar{z}' = \max \{\bar{z}, \max_k \{a_k \bar{q}\}\},$$

$$D_y' = D_y \cap \{k \mid [z', \bar{z}'] \cap [a_k \bar{q}, a_k q] \neq \emptyset\}$$

$$u' = \min \{u, \min_k \{\lceil z'/a_k \rceil\}\}, \quad \bar{u}' = \max \{\bar{u}, \max_k \{\lfloor \bar{z}'/a_k \rfloor\}\}$$

**Inference via reformulation**

Since (4.8) implies a disjunction $\bigvee_{k \in D_y} (z = a_k u)$, it can be given the standard convex hull relaxation for a disjunction, which in this case simplifies to

$$z = \sum_{k \in D_y} a_k u_k, \quad u = \sum_{k \in D_y} u_k$$

where $u_k \geq 0$ are new variables. Based on this idea, the relaxation of (4.7) becomes

$$\min \sum_j v_j$$

$$v_j = \sum_i \sum_{k \in D_{t_i}} A_{ijk} q_{ik}, \quad \text{all } j$$

$$q_i = \sum_{k \in D_{ti}} q_{ik}, \quad \text{all } i$$

$$L_j \leq v_j \leq U_j, \quad \text{all } j$$

$$q_i \leq q_i \leq q_i, \quad \text{all } i$$

$$q_{ik} \geq 0, \quad \text{all } i, k$$

(4.9)

**Preventing future infeasibilities**

There is also an opportunity for *post-relaxation inference*, which in this case takes the form of reduced cost variable fixing. Suppose the best feasible solution found so far has value $z^*$, and let $\hat{z}$ be the optimal value of (4.9). If $\hat{z} + r_{ik} \geq z^*$, where $r_{ik}$ is the reduced cost of $q_{ik}$ in the solution of (4.9), then $k$ can be removed from the domain of $t_i$. In addition one can infer

$$\bar{q}_i \leq \min_{k \in D_{t_i}} \{[(z^* - \hat{z})/r_{ik}]\}, \quad \text{all } i$$

Post-relaxation 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 it is enough to branch on the domain constraints \( t_i \in D_{t_i} \). Since \( t_i \) does not appear in the linear relaxation, it does not have a determinate value until it is fixed by branching. The domain constraint \( t_i \in D_{t_i} \) is viewed as unsatisfied as long as \( t_i \) is undetermined. The search branches on \( t_i \in D_{t_i} \) by splitting \( D_{t_i} \) into two subsets. Branching continues until all the \( D_{t_i} \) are singletons, at which point the solution of the relaxation will necessarily be feasible.

4.4.3 Experimental results

For our computational experiments, we used ten of the problem instances proposed by Thorsteinsson and Ottosson\[71\], which have 26 components, up to 30 types per component, and 8 attributes. We also used the same branching strategy on \( q_i \) as they suggested: 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 descendents 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 \( q_i \geq \bar{q}_i + 1 \).

The number of search nodes and CPU time (in seconds) required to solve each of the ten

<table>
<thead>
<tr>
<th>Instance</th>
<th>IP Solver (CPLEX 9.0)</th>
<th>Integrated solver</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Nodes</td>
<td>Time (s)</td>
</tr>
<tr>
<td>1</td>
<td>1</td>
<td>0.06</td>
</tr>
<tr>
<td>2</td>
<td>1</td>
<td>0.08</td>
</tr>
<tr>
<td>3</td>
<td>184</td>
<td>0.79</td>
</tr>
<tr>
<td>4</td>
<td>1</td>
<td>0.04</td>
</tr>
<tr>
<td>5</td>
<td>723</td>
<td>4.21</td>
</tr>
<tr>
<td>6</td>
<td>1</td>
<td>0.05</td>
</tr>
<tr>
<td>7</td>
<td>111</td>
<td>0.59</td>
</tr>
<tr>
<td>8</td>
<td>20</td>
<td>0.19</td>
</tr>
<tr>
<td>9</td>
<td>20</td>
<td>0.17</td>
</tr>
<tr>
<td>10</td>
<td>1</td>
<td>0.03</td>
</tr>
</tbody>
</table>

Table 4.2: Product configuration: search nodes and CPU time.
instances to optimality are shown in Table 4.2. Although the integrated model solves considerably slower than the pure MILP model, it again appears to be more robust than the MILP model in terms of the number of search nodes required, as was also the case in the production planning example. It is worth noting that Thorsteinsson and Ottosson\cite{71} used CPLEX 7.0 to solve the MILP model (4.6) and it managed to solve only 3 out of the above 10 instances with fewer than 100,000 search nodes. The average number of search nodes explored by CPLEX 7.0 over the 3 solved instances was around 77,000.

4.5 Example: Logic-based Benders decomposition

A third example we consider involves a decomposition approach for a machine scheduling problem. 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.

4.5.1 Modeling

An integer programming model can be written by discretizing time and letting \( x_{ijt} = 1 \) if job \( j \) starts processing at time \( t \) on machine \( i \).

\[
\min \sum_{ijt} c_{ij} x_{ijt}
\]

\[
\sum_{i,t} x_{ijt} = 1, \text{ all } i
\]

\[
\sum_{j} \sum_{t \in T_{ijt}} x_{ijt} \leq 1, \text{ all } i, t \tag{4.10}
\]

\[
x_{ijt} = 0, \text{ all } i, j, t \text{ with } d_j - p_{ij} < t < r_j \text{ or } t > N - p_{ij}
\]

\[
x_{ijt} \in \{0, 1\}
\]

where \( N \) is the number of discrete times (beginning with time 0), and

\[
T_{ijt} = \{t' | t - p_{ij} < t' \leq t\}
\]
is the set of times at which a job $j$ in progress on machine $i$ at time $t$ might start processing. There is also a continuous-time IP model suggested by Türkay and Grossmann\cite{72}, but computational testing (Hooker\cite{49}) indicates that it is much harder to solve than (4.10).

**A model based on metaconstraints**

A hybrid model can be written with the *cumulative* metaconstraint, 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 $i$ be always less than or equal to $U_i$. Here $t = (t_1, \ldots, t_n)$, $p_i = (p_{i1}, \ldots, p_{in})$, and similarly for $u_i$.

In the present instance, jobs must be 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

$$
\min \sum_j c_{y_j}
\quad r_j \leq t_j \leq d_j - p_{y_j}, \quad \forall j
\quad \text{cumulative}((t_j|y_j = i), (p_{ij}|y_j = i), e, 1), \quad \forall i
$$

(4.11)

where $e$ is a vector of ones.

**4.5.2 Solution process**

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 IP and a subproblem to be solved by CP. The assignment portion becomes the Benders master problem, which
allocates job $j$ to machine $i$ when $x_{ij} = 1$:

$$\min \sum_{ij} c_{ij} x_{ij}$$
$$\sum_i x_{ij} = 1, \text{ all } j$$

relaxation of subproblem

Benders cuts

$x_{ij} \in \{0, 1\}$

Inference

The solution $\bar{x}$ of the master problem determines the assignment of jobs to machines. Once these assignments are made, the problem (4.11) separates into a scheduling feasibility problem on each machine $i$:

$$r_j \leq t_j \leq d_j - p_{\bar{y}_j,j}, \text{ all } j$$

$$\text{cumulative}((t_j|\bar{y}_j = i), (p_{ij}|\bar{y}_j = i), e, 1)$$

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 (4.13) 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:

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

where $J_i = \{j \mid \bar{x}_{ij} = 1\}$. 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 examining the proof of infeasibility and noting which jobs actually play a role in the proof (see Hooker\cite{49}). 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 instance, in minimum makespan and minimum tardiness problems.

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:

\[
\sum_j p_{ij} x_{ij} \leq \max_j\{d_j\} - \min_j\{r_j\} \tag{4.15}
\]

A Benders method (as well as any nogood-based method) fits easily into this unifying 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

\[
\begin{align*}
\min & \sum_{ij} c_{ij} x_{ij} \quad & \text{(a)} \\
\sum_i x_{ij} &= 1, \text{ all } j & \text{(b)} \\
(x_{ij} = 1) &\iff (y_j = i), \text{ all } i, j & \text{(c)} \\
r_j &\leq t_j \leq d_j - p_{i_{y_j}}, \text{ all } j & \text{(d)} \\
\text{cumulative}((t_j|y_j = i), (p_{i_{y_j}}|y_j = i), e, 1), \text{ all } i & \text{(e)}
\end{align*}
\]

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 the master problem constraints (a) and (b) indicate that these constraints are themselves part of an IP relaxation of the problem. The relaxation parameter for (d) adds the inequalities (4.15) to the relaxation. The inference parameter for (e) specifies 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.
4.5.3 Experimental results

For our computational experiments, we used the instances proposed by Jain and Grossmann\cite{53} and we created three additional instances with more than 20 jobs. These are the last instance in Table 4.3 and the last two instances in Table 4.4. Although state-of-the-art IP solvers have considerably improved since Jain and Grossmann’s results were published, these instances are still intractable with the IP model (4.10). In addition to being orders of magnitude faster in solving the smallest problems, the integrated Benders approach can easily tackle larger instances as well. After 48 hours of CPU time and more than 5 million branch-and-bound nodes, the best solution found by the IP model to the last instance of Table 4.3 had value 176, whereas the optimal solution has value 175. When processing times are shorter the problem tends to become easier, and we report results for this case in Table 4.4. Even here, the IP model is still much worse than the integrated Benders approach as the problem size grows. For instance, after 16.9 million search nodes the IP solver could not find a single feasible solution to the instance with 22 jobs and 5 machines. The best solution it found for the last instance, after 48 hours and 4.5 million search nodes, had value 181, and the optimal value is 179.

<table>
<thead>
<tr>
<th>Jobs</th>
<th>Machines</th>
<th>IP Solver (CPLEX 9.0)</th>
<th>Integrated</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Nodes</td>
<td>Time (s)</td>
</tr>
<tr>
<td>3</td>
<td>2</td>
<td>1</td>
<td>0.00</td>
</tr>
<tr>
<td>7</td>
<td>3</td>
<td>1</td>
<td>0.02</td>
</tr>
<tr>
<td>12</td>
<td>3</td>
<td>11060</td>
<td>16.50</td>
</tr>
<tr>
<td>15</td>
<td>5</td>
<td>3674</td>
<td>14.30</td>
</tr>
<tr>
<td>20</td>
<td>5</td>
<td>159400</td>
<td>3123.34</td>
</tr>
<tr>
<td>22</td>
<td>5</td>
<td>&gt; 5.0M</td>
<td>&gt; 48h</td>
</tr>
</tbody>
</table>

Table 4.3: Parallel machine scheduling: long processing times.

<table>
<thead>
<tr>
<th>Jobs</th>
<th>Machines</th>
<th>IP Solver (CPLEX 9.0)</th>
<th>Integrated</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Nodes</td>
<td>Time (s)</td>
</tr>
<tr>
<td>3</td>
<td>2</td>
<td>1</td>
<td>0.00</td>
</tr>
<tr>
<td>7</td>
<td>3</td>
<td>1</td>
<td>0.01</td>
</tr>
<tr>
<td>12</td>
<td>3</td>
<td>4950</td>
<td>1.98</td>
</tr>
<tr>
<td>15</td>
<td>5</td>
<td>14000</td>
<td>19.80</td>
</tr>
<tr>
<td>20</td>
<td>5</td>
<td>140</td>
<td>5.73</td>
</tr>
<tr>
<td>22</td>
<td>5</td>
<td>&gt; 16.9M</td>
<td>&gt; 48h</td>
</tr>
<tr>
<td>25</td>
<td>5</td>
<td>&gt; 4.5M</td>
<td>&gt; 48h</td>
</tr>
</tbody>
</table>

Table 4.4: Parallel machine scheduling: short processing times.
Chapter 5
Applications

5.1 The robust spanning tree problem

Given an undirected graph with interval edge costs, a robust spanning tree is one whose
cost is as close as possible to that of a minimum spanning tree under any possible as-
signment of costs. This section defines the problem formally and introduces the main
concepts and notations necessary to understand the proposed solution.

Let $G = (V, E)$ be an undirected graph with $|V| = n$ nodes and $|E| = m$ edges and
an interval $[c^e, \overline{c^e}]$ for the cost of each edge $e \in E$. A scenario $s$ is a particular assignment
of a cost $c^e \in [c^e, \overline{c^e}]$ to each edge $e \in E$. We use $c^s_e$ to denote the cost of edge $e$ under
a given scenario $s$. Recall that a spanning tree for $G = (V, E)$ is a set of edges $T \subseteq E$
such that the subgraph $G' = (V, T)$ is acyclic and $\forall i \in V, \exists j \in V : (i, j) \in T$. The
cost of a spanning tree $T$ for a scenario $s$, denoted by $c^s_T$, is the sum of the costs of all
edges under scenario $s$:

$$c^s_T = \sum_{e \in T} c^s_e.$$  

A minimum spanning tree for scenario $s$, denoted by $MST^s$, is a spanning tree with
minimal cost for scenario $s$ and its cost is denoted by $c_{MST}^s$. Following YAMAN ET AL\cite{[76]}
we now define the worst case scenario for a spanning tree $T$ and the robust deviation of
$T$, two fundamental concepts for this section.
Definition 5.1.1 (Relative worst case scenario). Given a spanning tree $T$, a scenario $w(T)$ which maximizes the difference between the cost of $T$ and the cost of a minimum spanning tree under $w(T)$ is called a relative worst case scenario for $T$. More precisely, a relative worst case scenario $w(T)$ satisfies

$$w(T) \in \arg\max_{s \in S} (c^*_T - c_{MST^s})$$

(5.1)

where $S$ is the set of all possible scenarios.

Note that $\arg\max_{s \in S} f(s)$ denotes the set

$$\{ e \in S \mid f(e) = \max_{s \in S} f(s) \}.$$ 

and $\arg\min$ is defined similarly.

Definition 5.1.2 (Robust Deviation). The robust deviation of a spanning tree $T$, denoted by $\Delta_T$, is the distance between the cost of $T$ and the cost of a minimum spanning tree under the relative worst case scenario of $T$:

$$\Delta_T = c^w_{w(T)} - c_{MST^w(T)}.$$ 

(5.2)

The goal is to compute a robust spanning tree, i.e., a spanning tree whose robust deviation is minimal.

Definition 5.1.3 (Robust Spanning Tree). A (relative) robust spanning tree is a spanning tree $T^*$ whose robust deviation is minimal, i.e.,

$$T^* \in \arg\min_{T \in \Gamma_G} \max_{s \in S} (c^*_T - c_{MST^s})$$

where $\Gamma_G$ is the set of all spanning trees of $G$ and $S$ is the set of all possible scenarios. According to the definition of $w(T)$ (Eq. 5.1), this is equivalent to:

$$T^* \in \arg\min_{T \in \Gamma_G} (c^w_{w(T)} - c_{MST^w(T)}).$$

Figure 5.1 depicts these concepts graphically. The horizontal axis depicts various scenarios. For each scenario $s$, it shows the cost $c^s_T$ of $T$ under $s$ and the cost $c_{MST^s}$ of an MST under $s$. In the figure, scenario $s_5$ is a worst-case scenario, since the distance $c^s_T - c_{MST^s}$
is maximal and this distance is then the robust deviation of $T$. Such a figure can be drawn for each spanning tree and we are interested in finding the spanning tree with the smallest robust deviation. Kouvelis and Yu\cite{59} conjectured that the robust spanning tree problem with interval edges is NP-complete, and Aron and Van Hentenryck\cite{5} proved that this conjecture was true.

**Prior Work**

Yaman et al\cite{76} proposed an elegant MIP formulation for the RSTPIE problem. The formulation combines the single commodity model of the minimum spanning tree with the dual of the multicommodity model for the same problem. In addition, they introduced the concept of weak and strong edges and used them for preprocessing. They showed that preprocessing significantly enhances the performance of their MIP implementation. We now summarize their relevant results.

We first introduce the concept of weak edges, i.e., the only edges that need to be considered during the search.

**Definition 5.1.4 (Weak Tree).** A tree $T \subseteq E$ is weak if there exists at least one scenario under which $T$ is a minimum spanning tree of $G$.

**Definition 5.1.5 (Weak Edge).** An edge $e \in E$ is weak if it lies on at least one weak tree.
Strong edges are edges that are necessarily part of a robust spanning tree.

**Definition 5.1.6 (Strong Edge).** An edge \( e \in E \) is strong if it lies on a minimum spanning tree of \( G \) for all possible scenarios.

The following propositions characterize weak and strong edges in terms of the minimum spanning trees of two scenarios.

**Proposition 5.1.7.** An edge \( e \in E \) is weak if and only if there exists a minimum spanning tree using edge \( e \) when its cost is at the lowest bound and the costs of the remaining edges are at their highest bounds.

**Proposition 5.1.8.** An edge \( e \in E \) is strong if and only if there exists a minimum spanning tree using edge \( e \) when its cost is at the highest bound and the costs of the remaining edges are at their lowest bounds.

As a consequence, Yaman et al.\[76\] showed that it is possible to use a slightly modified version of Kruskal’s algorithm to find all the weak and strong edges of a given graph in \( O(m^2 \log m) \) time. In Section 5.1.1, we give an improved algorithm for finding weak edges, which runs in \( O(m \log m) \). The following two propositions capture the intuition we gave earlier on weak and strong edges.

**Proposition 5.1.9.** A relative robust spanning tree \( T \) is a weak tree. Thus an edge can be part of a relative robust spanning tree only if it is a weak edge.

**Proposition 5.1.10.** There exists a relative robust tree \( T \) such that every strong edge in the graph lies on \( T \).

The next result is also fundamental: it makes it possible to characterize precisely the worst case scenario of a spanning tree \( T \).

**Proposition 5.1.11.** The scenario in which the costs of all edges in a spanning tree \( T \) are at upper bounds and the costs of all other edges are at lower bounds is a relative worst case scenario for \( T \). In other words, \( w(T) \) is specified as

\[
c_{\text{w}(T)} = \begin{cases} 
    c_e, & e \in T \\
    \bar{c}_e, & e \in E \setminus T 
\end{cases}
\]  

(5.3)
In other words, once we select a tree $T$, we can easily find the worst-case scenario for $T$ by assigning to the edges in $T$ their upper bounds and to the edges not in $T$ their lower bounds. We use Proposition 5.1.11 in our new algorithm. In the rest of this section, we also use the notation $w(S)$ to denote the scenario where $c_e = \overline{c}_e$ if $e \in S$ and $c_e = c_e$ otherwise even when $S$ is not a tree.

**Solution process**

```
Algorithm Search ((S, R))
1  if (|S| < n − 1) then
2     ⟨S, R⟩ ← PruneInfeasible ((S, R))
3     ⟨S, R⟩ ← PruneSuboptimal ((S, R))
4     if (LB((S, R)) ≤ f*) then
5         e ← SelectEdge (E \ (S ∪ R))
6         Search ((S, R ∪ {e}))
7         Search ((S ∪ {e}, R))
8     else
9         if (∆S < f*) then
10            T* ← S
11            f* ← ∆S
12         end if
13     end if
14  end if
```

Figure 5.2: Robust spanning tree: the search algorithm

Figure 5.2 gives a high-level description of the search algorithm. A node in the search tree is called a configuration to avoid confusion with the nodes of the graph. A configuration is a pair $⟨S, R⟩$, where $S$ represents the set of selected edges and $R$ represents the set of rejected edges. The algorithm receives a configuration as input. If the configuration is not a spanning tree, the algorithm prunes infeasible and suboptimal edges. Both steps remove edges from $E \setminus (S \cup R)$ and adds them to $R$. If the lower bound of the resulting configuration is smaller than the best found solution, the algorithm selects an edge and explores two subproblems recursively. The subproblems respectively reject and select the
Notation | Definition
--- | ---
$c^s_T$ | the cost of tree $T$ under scenario $s$
$c^e_T = \sum_{e \in T} c^s_e$ | worst case scenario for edge set $T$: $w(T) = \left\{ \begin{array}{ll} c^s_e, & e \in T \\ c^e_e, & e \in E \setminus T \end{array} \right.$
$\Gamma_G$ | the set of all spanning trees of graph $G$
$T(S, R)$ | the set of spanning trees derived from $\langle S, R \rangle$
$T(S, R) = \{ T \in \Gamma_G | S \subseteq T \subseteq E \setminus R \}$
$M^s(S, R)$ | the set of spanning trees from $T(S, R)$ with minimal cost under scenario $s$
$M^s(S, R) = \text{arg-min}_{T \in T(S, R)} c^s_T$
$MST^s(S, R)$ | an arbitrary tree from $M^s(S, R)$
$\mathcal{M}^s \mathcal{M}^s(\emptyset, \emptyset)$ | an arbitrary tree from $\mathcal{M}^s(\emptyset, \emptyset)$

Table 5.1: Robust spanning tree: notations

edge. The best found solution and upper bound are updated each time a spanning tree with a smaller robust deviation is obtained.

### 5.1.1 Inference

Infeasibility pruning is relatively simple in our algorithm. It ensures that $S$ can be extended into a spanning tree and removes edges that would create cycles. We do not discuss infeasibility further and instead focus on the lower bound and suboptimality pruning. Before doing so, we introduce some additional notations. We use $T(S, R)$ to denote the set of all spanning trees that can be derived from configuration $\langle S, R \rangle$, i.e.,

$$T(S, R) = \{ T \in \Gamma_G \mid S \subseteq T \subseteq E \setminus R \}.$$ 

Given a scenario $s$, $M^s(S, R)$ denotes the set of all minimum spanning trees which are derived from configuration $\langle S, R \rangle$ under scenario $s$ and $MST^s(S, R)$ is a representative of $\mathcal{M}^s(S, R)$. For simplicity, we use $\mathcal{M}^s$ and $MST^s$ when $S = \emptyset$ and $R = \emptyset$. All relevant notations are summarized in Table 5.1 for convenience.

### The Lower Bound

We now present a lower bound to the robust deviation of any spanning tree derived from
\( \langle S, R \rangle \). In other words, we need to find a lower bound on the value of \( \Delta_T \) (as defined by Eq. 5.2), for any tree \( T \in \mathcal{T}(S, R) \). Recall that, for any such tree \( T \), the robust deviation is given by \( \Delta_T = c_T^w - c_{\text{MST}}^w(T) \). We can approximate \( \Delta_T \) by finding a lower bound to \( c_T^w \) and an upper bound to \( c_{\text{MST}}^w(T) \). Since \( S \subseteq T \subseteq S \cup L \) where \( L = E \setminus (S \cup R) \), both bounds can be obtained by considering the scenario \( w(S \cup L) \). As a consequence, we define the lower bound \( LB(\langle S, R \rangle) \) of a configuration \( \langle S, R \rangle \) as

\[
LB(\langle S, R \rangle) = c_{\text{MST}}^w(S \cup L)(S, R) - c_{\text{MST}}^w(S \cup L).
\]

The following proposition proves that \( LB(\langle S, R \rangle) \) is indeed a lower bound.

**Proposition 5.1.12.** Let \( \langle S, R \rangle \) be an arbitrary configuration and let \( L = E \setminus (S \cup R) \). Then, for all \( T \in \mathcal{T}(S, R) \), we have

\[
\Delta_T \geq c_{\text{MST}}^w(S \cup L)(S, R) - c_{\text{MST}}^w(S \cup L).
\]

**Proof:** Since \( T \in \mathcal{T}(S, R) \), it follows that \( S \subseteq T \subseteq E \setminus R = S \cup L \). Therefore,

\[
c_{\text{MST}}^w(T) \leq c_{\text{MST}}^w(S \cup L)
\]

On the other hand, by the definition of a minimum spanning tree and since \( T \subseteq S \cup L \), we have that

\[
c_{\text{MST}}^w(S \cup L)(S, R) \leq c_T^w(S \cup L) = c_T^w(T).
\]

The result follows since

\[
c_{\text{MST}}^w(S \cup L)(S, R) - c_{\text{MST}}^w(S \cup L) \leq c_T^w(T) - c_{\text{MST}}^w(T) = \Delta_T.
\]

Observe that this lower bound only requires the computation of two minimum spanning trees and hence it can be computed in \( O(m \log m) \) time. Interestingly, \( c_{\text{MST}}^w(S \cup L) \) can use any edge in the graph and is thus independent of the edges selected in \( S \) and \( R \). Of course, the scenario \( w(S \cup L) \) is not! It is easy to see that this lower bound is monotone in both arguments

\[
LB(\langle S, R \rangle) \leq LB(\langle S \cup \{e\}, R \rangle)
\]

\[
LB(\langle S, R \rangle) \leq LB(\langle S, R \cup \{e\} \rangle).
\]
Suboptimality Pruning

A fundamental component of our algorithm is suboptimality pruning, i.e., the ability to remove all non-weak edges at every configuration of the search tree. The results in YAMAN ET AL.\[76\] allow us to detect all weak edges in time \(O(m^2 \log m)\) by solving \(m\) MSTs. This cost is prohibitive in practice. We now show how to detect all weak edges by solving a single MST and performing a postprocessing step for each edge. The overall complexity is \(O(n^2 + m \log m)\), which is \(O(m \log m)\) on dense graphs.

The key idea is to characterize all weak edges in terms of a unique scenario. The characterization is based on the following results, which specify when a tree remains an MST under cost changes and the cost of an MST which must contain a specified edge.

Proposition 5.1.13. Let \(s\) be a scenario and \(T \in M^s\). Let \(e = (u, v) \notin T\) and \(f = (x, y)\) be the edge of maximal cost on the path from \(u\) to \(v\). Consider \(\hat{s}\) the scenario \(s\) where \(c^s_e\) is replaced by \(c^\hat{s}_e\), all other costs remaining the same. Then \(T \in M^{\hat{s}}\) if \(c^\hat{s}_e \geq c^\hat{s}_f\).

Proof: By contradiction. Assume that there exists a tree \(T'\) containing \(e\) such that \(c^\hat{s}_{T'} < c^\hat{s}_T\). Removing \(e\) from \(T'\) produces two connected components \(C_1\) and \(C_2\). If \(x \in C_1\) and \(y \in C_2\), then we can construct a tree

\[
T'' = T' \setminus \{e\} \cup \{f\}
\]

and we have

\[
c^\hat{s}_{T''} = c^\hat{s}_{T'} - (c^\hat{s}_e - c^\hat{s}_f) < c^\hat{s}_{T'} < c^\hat{s}_T.
\]

Since \(e \in T''\) and \(e \in T\), we have

\[
c^\hat{s}_{T''} = c^\hat{s}_T < c^\hat{s}_T = c^\hat{s}_T
\]

which contradicts the fact that \(T \in M^s\). If \(x, y \in C_1\) (resp. \(C_2\)), since there exists a cycle in the graph containing \(e\) and \(f\), there exists at least one edge \(g\) on the path from \(u\) to \(v\) in \(T\) such that \(g \in T'\) (otherwise \(T'\) would not be a tree). By hypothesis, \(c^s_g \leq c^s_f\) and hence \(c^\hat{s}_e \geq c^\hat{s}_g\). We can thus apply the same construction as in the case \(x \in C_1\) and \(y \in C_2\) with \(f\) replaced by \(g\).
Proposition 5.1.14. Let \( s \) be a scenario and \( T \) be an \( MST^s \). Let \( e = (u, v) \notin T \) and \( f = (x, y) \) be the edge of maximal cost on the path from \( u \) to \( v \). Then, \( T \setminus \{ f \} \cup \{ e \} \in \mathcal{M}^s(\{ e \}, \emptyset) \).

The proof of this result is similar to the proof of Proposition 5.1.13. We are now ready to present a new characterization of weak edges. The characterization only uses the scenario \( \bar{s} \) where all costs are at their upper bounds.

Proposition 5.1.15. Let \( \bar{s} \) be the scenario where all costs are at their upper bounds and \( T \in \mathcal{M}^\bar{s} \). An edge \( e = (u, v) \) is weak if \( e \in T \) or if an edge \( f \) of maximal cost on the path from \( u \) to \( v \) in \( T \) satisfies \( c_e \leq c_f \).

Proof: Let \( \hat{s} \) the scenario \( \bar{s} \) where \( c_{\bar{s}} e \) is replaced by \( c_{\hat{s}} e \). If \( e \in T \), then \( T \in \mathcal{M}^{\hat{s}} \) as well and hence \( e \) is weak by Proposition 5.1.7. If \( e \notin T \) and \( c_{\bar{s}} e > c_{\hat{s}} f \), then \( T \in \mathcal{M}^{\hat{s}} \) by Proposition 5.1.13. By Proposition 5.1.14, \( T \setminus \{ f \} \cup \{ e \} \in \mathcal{M}^{\hat{s}}(\{ e \}, \emptyset) \) and its cost is greater than \( c_T \) since \( c_{\bar{s}} e > c_{\hat{s}} f \). Hence \( e \) is not weak. If \( e \notin T \) and \( c_{\bar{s}} e = c_{\hat{s}} f \), then \( T \setminus \{ f \} \cup \{ e \} \) is an \( MST^{\bar{s}} \) and hence \( e \) is weak. The same holds for the case where \( e \in T \) and \( c_{\bar{s}} e < c_{\hat{s}} f \).

<table>
<thead>
<tr>
<th>Algorithm MaxCost ((u, v))</th>
</tr>
</thead>
<tbody>
<tr>
<td>1  if ((u = v)) then</td>
</tr>
<tr>
<td>2    return 0</td>
</tr>
<tr>
<td>3  else</td>
</tr>
<tr>
<td>4    if ((\text{level}(u) &lt; \text{level}(v))) then</td>
</tr>
<tr>
<td>5      return (\max{ \text{cost}(v, p(v)) , \text{MaxCost} (u, p(v)) } )</td>
</tr>
<tr>
<td>6    else</td>
</tr>
<tr>
<td>7      if ((\text{level}(u) &gt; \text{level}(v))) then</td>
</tr>
<tr>
<td>8      return (\max{ \text{cost}(u, p(u)) , \text{MaxCost} (p(u), v) } )</td>
</tr>
<tr>
<td>9      else</td>
</tr>
<tr>
<td>10     maxedge (\leftarrow) (\max{ \text{cost}(u, p(u)) , \text{cost}(v, p(v)) } )</td>
</tr>
<tr>
<td>11     return (\max{ \text{maxedge} , \text{MaxCost} (p(u), p(v)) } )</td>
</tr>
<tr>
<td>12     end if</td>
</tr>
<tr>
<td>13  end if</td>
</tr>
<tr>
<td>14 end if</td>
</tr>
</tbody>
</table>

Figure 5.3: Robust spanning tree: finding the largest cost of an edge

We now describe how to use Proposition 5.1.15 to obtain an \( O(m \log m) \) algorithm on dense graphs. The key idea is to compute \( MST^{\bar{s}} \), i.e., the MST when all costs are at
their upper bounds. All edges in this MST are weak. In addition, for each \( e = (u, v) \) not in the MST, we compute the largest cost of any edge on the path from \( u \) to \( v \). This can be done easily by upgrading Prim’s algorithm slightly to associate a level and a parent to each vertex. When an edge \( e = (u, v) \) with \( u \in T \) and \( v \notin T \) is added to \( T \) in Prim’s algorithm, we set

\[
\text{level}(v) = \text{level}(u) + 1;
\]
\[
\text{parent}(v) = u;
\]

These modifications do not affect the complexity of the algorithm. It now suffices to apply the algorithm depicted in Figure 5.3 to compute the cost of the maximal edge and to apply Proposition 5.1.15. The algorithm in Figure 5.3 simply follows the paths from \( u \) and \( v \) to their common ancestor, computing the cost of the maximal edge on the way. Since algorithm \( \text{MaxCost} \) takes \( O(n) \) in the worst case, the overall complexity becomes \( O(m \log m + mn) \). However, it is easy to reduce this complexity to \( O(m \log m + n^2) \) by amortizing the computation of the maximal costs. It suffices to cache the results of \( \text{MaxCost} \) in an \( n \times n \) matrix \( M \). For each edge \( e = (u, v) \in E \setminus T \) we first examine entry \( M[u, v] \) and call \( \text{MaxCost}(u, v) \) only if this entry is uninitialized. Since there are at most \( n^2 \) entries and each call to \( \text{MaxCost}(u, v) \) costs \( O(1) \) per entry it fills, the overall complexity becomes \( O(m \log m + n^2) \). We proved the following theorem.

**Theorem 5.1.16.** All weak edges of a graph can be computed in \( O(m \log m + n^2) \) time.

### 5.1.2 Search

It is well-known that a good branching heuristic may improve performance significantly. We now show a branching heuristic adapting the first-fail principle (see Haralick and Elliott[41]) to robust optimization. The key idea is to explore the most uncertain edges first, i.e., to branch on an edge \( e \) with the maximal difference \( c_e - \bar{c}_e \). Indeed, rejecting \( e \) (i.e., adding \( e \) to \( R \)) allows \( \text{MST}^{w(S \cup L)} \) to select \( e \) at a low cost, possibly giving a large deviation. Hence this branch is likely to fail early. However, this is only important if \( \text{MST}^{w(S \cup L)} \) is likely to select \( e \) which may not necessarily the case if \( \bar{c}_e \) is large compared to
other “similar” edges. Hence, it seems appropriate to select first an edge $e \in \text{MST}^w(S \cup L)$ whose difference $\overline{c_e} - \underline{c_e}$ is maximal. This justifies the following branching strategy.

**Definition 5.1.17 (Branching Strategy).** Let $\langle S, R \rangle$ be a configuration, $L^* = L \cap \text{MST}^w(S \cup L)$, and $L^- = L \setminus \text{MST}^w(S \cup L)$. The branching strategy selects an edge $s$ defined as follows:

$$s = \begin{cases} 
\arg\max_{e \in L^*} \overline{c_e} - \underline{c_e} & \text{if } L^* \neq \emptyset \\
\arg\max_{e \in L^-} \overline{c_e} - \underline{c_e} & \text{otherwise.}
\end{cases}$$

### 5.1.3 Experimental results

We now report experimental results comparing the constraint satisfaction and the MIP approaches. The comparison uses the instances in *Yaman et al.*\cite{76}, as well as some new instances which capture additional structure arising in practical applications.

The experimental setting of *Yaman et al.*\cite{76} uses complete graphs with $n$ vertices and six classes of problems. Three of the six classes use tight intervals for edge costs,
while the other three allowed larger differences between the lower and upper bounds. The edge intervals were chosen as follows. The values of \( c_e \) and \( \overline{c_e} \) are uniformly distributed in the intervals:

- **class 1:** \( c_e \in [0, 10] \), \( \overline{c_e} \in (c_e, 10] \)
- **class 2:** \( c_e \in [0, 15] \), \( \overline{c_e} \in (c_e, 15] \)
- **class 3:** \( c_e \in [0, 20] \), \( \overline{c_e} \in (c_e, 20] \)
- **class 4:** \( c_e \in [0, 10] \), \( \overline{c_e} \in (c_e, 20] \)
- **class 5:** \( c_e \in [0, 15] \), \( \overline{c_e} \in (c_e, 30] \)
- **class 6:** \( c_e \in [0, 20] \), \( \overline{c_e} \in (c_e, 40] \)

Note that the size of the search space to explore is \( O(2^{300}) \) for a complete graph of 25 nodes. Of course, our constraint satisfaction algorithm will only explore a small fraction of that space. In addition to these six classes, we also generate instances (Classes 7 and 8) whose cost structure is not the same for all the edges. Class 7 contains instances which represent a two-level network. The lower level consists of clusters of 5 nodes whose edges are generated according to Class 1 above. The upper level links the clusters and these edges have higher costs, i.e., Class 1 costs shifted by a constant which is larger than the Class 1 edges. This captures the fact that, in networks, there are often various types of edges with different costs. See Figure 5.4 for an instance of Class 7 with 35 nodes. Class 8 contains instances which are similar to Class 7, except that the upper-level layer is organized as a binary tree. See Figure 5.5 for an instance of Class 8 with 35 nodes. In
Table 5.2: Robust spanning tree: average CPU time

<table>
<thead>
<tr>
<th>Edges(Nodes)</th>
<th>Algo.</th>
<th>Class 1</th>
<th>Class 2</th>
<th>Class 3</th>
<th>Class 4</th>
<th>Class 5</th>
<th>Class 6</th>
<th>Class 7</th>
<th>Class 8</th>
</tr>
</thead>
<tbody>
<tr>
<td>45(10)</td>
<td>CSR</td>
<td>0.12</td>
<td>0.07</td>
<td>0.08</td>
<td>0.08</td>
<td>0.06</td>
<td>0.07</td>
<td>0.06</td>
<td>0.06</td>
</tr>
<tr>
<td></td>
<td>CSF</td>
<td>0.14</td>
<td>0.10</td>
<td>0.12</td>
<td>0.12</td>
<td>0.09</td>
<td>0.11</td>
<td>0.07</td>
<td>0.07</td>
</tr>
<tr>
<td></td>
<td>MIP</td>
<td>6.59</td>
<td>3.95</td>
<td>4.10</td>
<td>4.38</td>
<td>4.68</td>
<td>3.83</td>
<td>2.29</td>
<td>2.02</td>
</tr>
<tr>
<td>105(15)</td>
<td>CSR</td>
<td>1.82</td>
<td>1.96</td>
<td>1.09</td>
<td>0.36</td>
<td>0.86</td>
<td>0.33</td>
<td>1.76</td>
<td>1.15</td>
</tr>
<tr>
<td></td>
<td>CSF</td>
<td>2.53</td>
<td>3.06</td>
<td>2.15</td>
<td>0.86</td>
<td>2.01</td>
<td>0.77</td>
<td>1.83</td>
<td>1.35</td>
</tr>
<tr>
<td></td>
<td>MIP</td>
<td>245.19</td>
<td>184.14</td>
<td>136.88</td>
<td>109.30</td>
<td>91.33</td>
<td>87.62</td>
<td>730.20</td>
<td>88.66</td>
</tr>
<tr>
<td>190(20)</td>
<td>CSR</td>
<td>39.72</td>
<td>33.80</td>
<td>8.91</td>
<td>3.43</td>
<td>2.09</td>
<td>3.12</td>
<td>7.37</td>
<td>9.51</td>
</tr>
<tr>
<td></td>
<td>CSF</td>
<td>74.86</td>
<td>61.08</td>
<td>12.66</td>
<td>7.56</td>
<td>4.69</td>
<td>7.05</td>
<td>5.28</td>
<td>6.77</td>
</tr>
<tr>
<td></td>
<td>MIP</td>
<td>8620.66</td>
<td>5517.75</td>
<td>14385.55</td>
<td>3344.95</td>
<td>12862.29</td>
<td>22855.82</td>
<td>18547.27</td>
<td>1399.48</td>
</tr>
<tr>
<td>300(25)</td>
<td>CSR</td>
<td>121.85</td>
<td>181.57</td>
<td>68.42</td>
<td>20.41</td>
<td>12.23</td>
<td>13.26</td>
<td>91.89</td>
<td>61.71</td>
</tr>
<tr>
<td></td>
<td>CSF</td>
<td>244.18</td>
<td>272.63</td>
<td>145.11</td>
<td>50.37</td>
<td>30.55</td>
<td>32.41</td>
<td>97.94</td>
<td>36.21</td>
</tr>
<tr>
<td>435(30)</td>
<td>CSR</td>
<td>926.65</td>
<td>415.07</td>
<td>942.38</td>
<td>133.90</td>
<td>63.85</td>
<td>177.47</td>
<td>1719.61</td>
<td>721.28</td>
</tr>
<tr>
<td></td>
<td>CSF</td>
<td>2100.43</td>
<td>909.15</td>
<td>1811.88</td>
<td>359.72</td>
<td>167.58</td>
<td>418.41</td>
<td>804.78</td>
<td>284.39</td>
</tr>
<tr>
<td></td>
<td>MIP</td>
<td>10711.37</td>
<td>11906.01</td>
<td>4183.22</td>
<td>1100.88</td>
<td>548.01</td>
<td>1115.83</td>
<td>14585.73</td>
<td>2149.58</td>
</tr>
<tr>
<td>595(35)</td>
<td>CSR</td>
<td>4639.55</td>
<td>5095.71</td>
<td>2304.36</td>
<td>383.34</td>
<td>188.36</td>
<td>419.56</td>
<td>32511.77</td>
<td>6356.78</td>
</tr>
<tr>
<td></td>
<td>CSF</td>
<td>10771.37</td>
<td>11906.01</td>
<td>4183.22</td>
<td>1100.88</td>
<td>548.01</td>
<td>1115.83</td>
<td>14585.73</td>
<td>2149.58</td>
</tr>
<tr>
<td></td>
<td>MIP</td>
<td>27206.38</td>
<td>16388.12</td>
<td>15059.39</td>
<td>1103.50</td>
<td>1122.57</td>
<td>1071.62</td>
<td>57309.23</td>
<td>33390.67</td>
</tr>
<tr>
<td>780(40)</td>
<td>CSR</td>
<td>29421.70</td>
<td>34666.84</td>
<td>22084.00</td>
<td>3456.84</td>
<td>3241.04</td>
<td>3031.55</td>
<td>28432.91</td>
<td>11339.59</td>
</tr>
</tbody>
</table>

In general, classes 7 and 8 are significantly harder than classes 1 to 6, since preprocessing is less effective than in Classes 1 to 6 because of the additional structure. Observe also that these instances are sparser for the same number of nodes.

Table 5.2 compares the efficiency of the two approaches. It reports the computation times in CPU seconds of the MIP implementations, the constraint satisfaction algorithm with suboptimality pruning at the root node only (CSR), and the constraint satisfaction algorithm with suboptimality pruning at every node (CSF). The times are given on a Sun Sparc 440Mhz processor and the average is computed over 10 instances for each class and each size. We used CPLEX 6.51 for solving the MIP, after preprocessing of the weak and strong edges. Note that the constraint programming approach produces dramatic speedups over the pure integer programming approach. On Class 1, CSR runs about 134 times faster than the MIP on graphs with 15 nodes and about 217 times faster on graphs with 20 nodes. On Classes 7 and 8, the speed-ups are even more impressive. On graphs with 20 nodes for classes 7 and 8, CSR runs about 3500 and 200 times faster than the MIP. (Recall that these graphs are not complete). The MIP times are not given for graphs with more than 20 nodes, since they cannot be obtained in reasonable time. The results indicate that the constraint satisfaction approach is also better at exploiting the network structure, which is likely to be fundamental in practice. Observe also that the constraint
satisfaction approach is able to tackle much large instances in reasonable times.

Figure 5.6: Robust spanning tree: results using CP on class 1.

Figure 5.7: Robust spanning tree: results using CP on class 7.

Figures 5.6, 5.7, and 5.8 plot the execution times of the two constraint satisfaction algorithms. Observe also that applying suboptimality pruning at every node is not cost-effective on Classes 1 to 6. This is due to the fact that graphs are complete and costs are uniformly distributed in these instances. Classes 7 and 8, which add a simple additional cost structure, clearly indicate that suboptimality pruning becomes increasingly important when the network is more heterogeneous. The benefits of suboptimality pruning clearly appears on large graphs for class 8, where CSF is about three times as fast in the average.
In fact, CSF is significantly faster (e.g., 10 times faster) than CSR on some instances, while the two algorithms are similar on others. It is important to mention that systematic suboptimality pruning is useless without Theorem 5.1.16. Indeed, the pruning benefit is often offset by the high pruning cost otherwise.

Overall, it is clear that the constraint satisfaction approach is much more effective on these problems than the MIP approach. It produces extremely dramatic speed-ups and substantially enlarge the class of instances that are amenable to effective solutions. The constraint satisfaction approach is able to solve large-scale problems (over 1,000 edges). Since networks are often organized in hierarchies, it should scale up nicely to larger real-life instances. There is still considerable room for improvement in the implementation, since incremental MST algorithms (see Rauch et al.[68]) and enhanced feasibility pruning may decrease runtime significantly.
Bibliography


Appendix A

A.1 Lift and project cut generator: algorithm details

The procedure for generating lift and project cuts, as outlined in the current document, deals only with binary variables. That is we only generate disjunctive cuts for fractional binary variables. The method can be extended to handle non-binary (general integer constrained) variables, but this is beyond the scope of this work. However, with this extension in mind, we shall refer to these variables as constrained variables, rather than just binary. Wherever the assumption that a variable is binary is made in the pseudocode, we say so in a comment. Since the procedure involves pivoting, which starts from an optimal simplex tableau and at each step ends up with a new tableau, we adopt the following notation: all objects that are related to the initial (optimal) solution of the LP relaxation are denoted as $\bar{x}$. The objects related to the subsequent (current) LP tableau (whether optimal or not, feasible or not) are denoted with $\bar{x}$. This is a slight departure from the notation used by Balas and Perregaard\cite{Balas1982}, in the sense that they use $\bar{a}_{ij}$ to refer to the coefficient for nonbasic variable $j$ in row $i$ of the current tableau. We use $\bar{a}_{ij}$ instead (and reserve $\bar{a}_{ij}$, as well as $\bar{x}$ and $\bar{B}$ for the case when we talk about the optimal LP tableau). We also use the term slack for both slack and surplus variables, but whenever the distinction is necessary, we point it out.
Algorithm \texttt{pivot\_cuts} ($LP$, \texttt{rounds}, \texttt{max\_pivots}, \texttt{strengthen})

\begin{tabular}{|c|c|}
\hline
\texttt{rounds} & number of times we solve the LP and generate cuts for each fractional variable of the optimal solution \\
\texttt{max\_pivots} & maximum number of pivots allowed in order to strengthen each cut \\
\texttt{strengthen} & if true, we apply disjunctive modularization (i.e. integrality strengthening) to row \texttt{k} after each pivot \\
\texttt{\tilde{B}} & the current basis (which, sometimes, may be infeasible) \\
\texttt{(\tilde{x}, \tilde{s})} & the current solution (structural and slack) \\
\hline
\end{tabular}

1. \textbf{for} ($1 \leq t \leq \texttt{rounds}$) \textbf{do} \\
   /* Solve LP and let $\pi$ be the optimal solution */ \\
2. \hspace{1em} ($\pi, \pi_0, \tilde{B}, \tilde{A}$) $\leftarrow$ \texttt{optimal\_solution} ($LP$) \\
   /* Generate optimal lift-and-project cuts for all fractional components of $\pi$ */ \\
   /* \texttt{NOTE!} Since the new cuts could make the primal tableau infeasible, we must use dual-simplex from this point */ \\
3. \hspace{1em} \textbf{foreach} (structural, constrained variable $k : \pi_k$ is fractional) \textbf{do} \\
4. \hspace{2em} ($\pi, \pi_0$) $\leftarrow$ \texttt{create\_lift\_and\_project\_cut} ($k, \pi, \pi_0, \tilde{B}, \tilde{A}, \texttt{strengthen}$) \\
   /* add the L&P cut derived from $x_k \leq 0 \lor x_k \geq 1$ to the LP relaxation */ \\
5. \hspace{2em} \texttt{add\_cut} ($\pi, \pi_0, LP$) \\
6. \hspace{1em} \textbf{end for} \\
7. \textbf{end for}

Figure A.1: Algorithm for generating cuts via pivoting
Algorithm `create_lift_and_project_cut (k, π, π, B, A, strengthen)`

```plaintext
    1. optimal ← false /* will be set to true when we have an optimal lift-and-project cut from row k */
    2. degeneracy ← false /* will be set to true if we need to perturb row k to avoid degeneracy in CGLP */
    3. (B, A) ← (B, A) /* we start from the optimal LP tableau and pivot until we obtain the optimal cut from row k */
    4. if (strengthen) then
        /* Apply disjunctive modularization to row k */
        5. (φ_j, φ_0) ← create_strengthened_row (A, J, k) /* add the new row to the tableau */
        6. A ← add_LP_tableau_row (A, φ_j, φ_0)
        WARNING! The index k has changed as a result of appending a new modularized row (we now have a new variable, y_k, on which we will work instead of x_k).
        7. k ← index of the newly added row /* this new row becomes our cut generation row! */
    8. end if
    9. J ← { indices of all non-basic variables in the current solution }
    10. pivots ← 0 /* We use this to limit the number of pivots (which will lead to a suboptimal cut) */
    11. while (not optimal and pivots < max_pivots) do /* the cut is not yet an optimal lift-and-project cut */
        /* Can we strengthen the cut (i.e. improve the solution of CGLP) by pivoting in the current LP tableau? */
        12. if (i* → find_cut_improving_pivot_row (A, B, J, k)) then
            13. j* ← find_best_pivot_column (A, B, J, i*, strengthen)
            14. (A, B, J) ← change_basis (A, B, J, i*, j*, strengthen)
            15. pivots ← pivots + 1
        16. else /* in practice, this branch can be skipped, since the cut obtained without it has roughly the same strength */
            17. if (row k has no 0 entries) then
                18. optimal ← true /* we are done, the cut is optimal */
            19. else /* we must perturb row k such that the partition (M_1, M_2) (i.e. the basis in CGLP) becomes unique */
                20. degeneracy ← true
                21. A ← perturb_row (A, k)
            22. end if
            23. end if
        24. end while
    25. if (degeneracy) then
        26. A ← remove_perturbation_from_row (A, k)
    27. end if
    28. (π, π_0) ← create_intersection_cut (A, J, k) /* (π, π_0) is now a L&P cut */
    29. π ← strengthen_cut (A, J, k, π) /* (π, π_0) is now a strengthened L&P (MIG) cut */
    30. return (π, π_0)
```

Figure A.2: Algorithm for generating lift-and-project cuts
Procedure `find_cut_improving_pivot_row` ($\tilde{A}, \tilde{B}, J, k$)

/* choose a row $i$, some multiple of which will be added to row $k$ */
1 foreach (row $i \neq k$ of $\tilde{A}$) do
   /* First, assume $x_i$ will go to its lower bound (i.e. replace $x_i' = UB_i - x_i$) */
   2 `adjustTableauRow` ($A, i, UB$)
   3 $\tau_{u_i} \leftarrow \text{compute CGLP reduced cost} (\tilde{A}, \tilde{B}, J, k, x, \bar{x}, i, 1)$
      /* Should be computed only if $r_{u_i} \geq 0$ */
   4 $\tau_{v_i} \leftarrow \text{compute CGLP reduced cost} (\tilde{A}, \tilde{B}, J, k, x, \bar{x}, i, 2)$
      /* Now, assume $x_i$ will go to its lower bound (i.e. replace $x_i' = x_i - LB_i$) */
   5 `adjustTableauRow` ($A, i, LB$)
   6 $\tau_{u_i} \leftarrow \text{compute CGLP reduced cost} (\tilde{A}, \tilde{B}, J, k, x, \bar{x}, i, 1)$
      /* Should be computed only if $r_{u_i} \geq 0$ */
   7 $\tau_{v_i} \leftarrow \text{compute CGLP reduced cost} (\tilde{A}, \tilde{B}, J, k, x, \bar{x}, i, 2)$
   8 if ($r_{u_i} < 0 \lor r_{v_i} < 0$) then
      /* solution of CGLP (i.e. lift-and-project cut) can be improved */
      9 return $i$
      /* Pivoting on row $i$ in LP (i.e. pivoting $x_i$ out) corresponds to pivoting one of $u_i$ or $v_i$ into the basis of CGLP */
   10 end if
11 end for
/* No pivoting row found! */
12 return 0

Figure A.3: Finding a pivoting row (i.e. variable to be removed from $\tilde{B}$)
Procedure \textit{find\_best\_pivot\_column} ($\tilde{A}, \tilde{B}, J, k, i, strengthen$)

We want to choose a column $j$ to decide the sign and magnitude of the multiplier with which row $i$ will be added to row $k$:

$$j^*_i = \min_{j \in J} \left\{ \min_{j \in J} \gamma_j, \min_{j \in J} \frac{\gamma_j}{\delta_{ij}} \right\}$$

1. $m^* \leftarrow 0, j^*_i \leftarrow 0$ /* $j^*_i = 0$ means that column $j^*_i$ is undefined */
2. \textbf{foreach} $(j \in J \mid \frac{\delta_{ik}}{\delta_{ij}} < \gamma_j = -\frac{\delta_{jk}}{\delta_{ij}} < \frac{1}{\delta_{ij}})$ \textbf{do}
3. \hspace{1em} $m_j \leftarrow \text{CGLP\_objective\_function} (A, J, i, j, \gamma, \pi, \bar{\pi}, strengthen)$
4. \hspace{1em} \textbf{if} $(m_j < m^*)$ \textbf{then}
5. \hspace{2em} $m^* \leftarrow m_j, \ j^*_i \leftarrow j$
6. \hspace{1em} \textbf{end if}
7. \textbf{end for}
8. \textbf{return} $j^*_i$

Figure A.4: Finding the best pivoting column in the current basis $\tilde{B}$

Procedure \textit{change\_basis} ($\tilde{A}, \tilde{B}, J, i, l, strengthen$)

1. $(\tilde{A}, \tilde{B}, J) \leftarrow \text{pivot} (\tilde{A}, \tilde{B}, J, i, l)$
2. \textbf{if} (strengthen) \textbf{then}
   /* Apply disjunctive modularization to row $k$ */
3. \hspace{1em} $(\phi_J, \phi_0) \leftarrow \text{create\_strengthened\_row} (\tilde{A}, J, k)$ /* Replace row $k$ with the newly computed row */
4. \hspace{1em} $\tilde{A} \leftarrow \text{remove\_LP\_tableau\_row} (\tilde{A}, k)$
5. \hspace{1em} $\tilde{A} \leftarrow \text{add\_LP\_tableau\_row} (\tilde{A}, \phi_J, \phi_0)$ /* add the new row to the tableau */
6. \hspace{1em} $k \leftarrow \text{index of the newly added row}$ /* this new row becomes our cut generation row */
7. \textbf{end if}
8. \textbf{return} $(k, \tilde{A}, \tilde{B}, J)$

Figure A.5: Changing the current basis $\tilde{B}$ in the LP tableau
Procedure CGLP_objective \((\tilde{A}, J, i, l, \gamma, \pi, \tau, \mathsf{strengthen})\)

If we pivot on row \(i\) and column \(l\) (i.e. \(\gamma = -\frac{\alpha_{kl}}{\mu_{kl}}\)) then:

New row \(k\):
\[
\begin{align*}
\tilde{a}_{k,l} &\leftarrow \gamma \\
\tilde{a}_{k,j} &\leftarrow \mathsf{new\_row\_coefficient}\left(\tilde{a}_{k,j}, \tilde{a}_{i,j}, \gamma\right), \text{ for } j \not\in J \setminus \{l\} \\
\tilde{a}_{k,0} &\leftarrow \mathsf{new\_row\_coefficient}\left(\tilde{a}_{k,0}, \tilde{a}_{i,0}, \gamma\right)
\end{align*}
\]

New strengthened row \(k\) (after modularization):
\[
\begin{align*}
\tilde{a}_{k,l} &\leftarrow \mathsf{strengthened\_row\_coefficient}\left(\gamma, \tilde{a}_{k,0}, \text{typeof}(\{i\}, i)\right) \\
\tilde{a}_{k,j} &\leftarrow \mathsf{strengthened\_row\_coefficient}\left(\tilde{a}_{k,j}, \tilde{a}_{i,0}, \text{typeof}(J \setminus \{l\}, j)\right), \text{ for } j \not\in J \setminus \{l\} \\
\tilde{a}_{k,0} &\leftarrow \tilde{a}_{i,0}
\end{align*}
\]

**IMPORTANT!** Note that we apply disjunctive modularization to \(\tilde{a}_{k,j}\) (i.e. the new row), NOT to \(a_{k,j}\) (i.e. the row before pivoting). This means that when we compute the CGLP objective to determine which column to pivot on, we must BOTH simulate the pivot (i.e. compute \(a_{k,j}^\gamma\)) and apply disjunctive modularization to the new (simulated) row! Of course, the second step is only necessary when we use disjunctive modularization.

New simple disjunctive cut from new row \(k\):
\[
\begin{align*}
\pi_1 &\leftarrow \mathsf{intersection\_cut\_coefficient}\left(\gamma, \tilde{a}_{k,0}\right) \\
\pi_2 &\leftarrow \mathsf{intersection\_cut\_coefficient}\left(\tilde{a}_{k,j}, \tilde{a}_{i,0}\right), \text{ for } j \not\in J \setminus \{l\} \\
\pi_0 &\leftarrow \mathsf{intersection\_cut\_rhs}\left(\tilde{a}_{k,0}\right)
\end{align*}
\]

New strengthened intersection cut from new row \(k\):
\[
\begin{align*}
\pi_1 &\leftarrow \mathsf{strengthened\_intersection\_cut\_coefficient}\left(\gamma, \tilde{a}_{k,0}, \text{typeof}(\{i\}, i)\right) \\
\pi_2 &\leftarrow \mathsf{strengthened\_intersection\_cut\_coefficient}\left(\tilde{a}_{k,j}, \tilde{a}_{i,0}, \text{typeof}(J \setminus \{l\}, j)\right), \text{ for } j \not\in J \setminus \{l\} \\
\pi_0 &\leftarrow \mathsf{intersection\_cut\_rhs}\left(\tilde{a}_{k,0}\right)
\end{align*}
\]

Objective function of \((\text{CGLP})_{k}\) corresponding to the cut \(\pi_1^\gamma x_i + \pi_2^\gamma x_j \geq \pi_0^\gamma\):
\[
\begin{align*}
\mathcal{I}_{\text{CGLP}} &= \frac{-\pi_0^\gamma + \sum_{j \not\in (J \setminus \{l\}) \cup \{i\}} \pi_2^\gamma x_j}{1 + \sum_{j \not\in (J \setminus \{l\}) \cup \{i\}} |\tilde{a}_{k,j}|}
\end{align*}
\]

**IMPORTANT!** Note that using \(|\gamma|\) in the denominator as in [8] is misleading (and not always correct)!! Thus, the formula:
\[
\frac{\gamma_i x_i + \gamma_j x_j - \pi_0^\gamma}{1 + |\gamma| + \sum_{j \not\in (J \setminus \{l\}) \cup \{i\}} |\tilde{a}_{k,j}|}
\]

is not correct if we use disjunctive modularization, because \(\gamma\) stands for the value of \(\tilde{a}_{k,j}^\gamma\), which, when we use disjunctive modularization is no longer just \(\gamma\)! The correct formula is shown below:

\[
\begin{align*}
1 \quad \text{return} \quad \frac{-\pi_0^\gamma + \pi_f^\gamma x_i + \pi_f^\gamma x_j}{1 + |k_j| + \sum_{j \not\in J'} |\tilde{a}_{k,j}|}, \text{ where } J' = J \setminus \{l\}
\end{align*}
\]

Figure A.6: Computing the CGLP objective given by a prospective pivot \((\gamma)\)
**Procedure** intersection_cut_coefficient $(\alpha, \beta)$

1. return $\max \{ \alpha (1 - \beta), -\alpha \beta \} = -\alpha \beta + \max \{ \alpha, 0 \}$

Figure A.7: Computing coefficients for intersection cuts

**Procedure** strengthened_intersection_cut_coefficient $(\alpha, \beta, \text{vartype})$

/* Strengthen the coefficients that correspond to constrained (i.e. structural) non-basic variables */
/* using the integrality conditions on these variables */

1. return

\[
\begin{cases}
\min \{ (\alpha - \lfloor \alpha \rfloor) (1 - \beta), (\lceil \alpha \rceil - \alpha) \beta \} & \text{vartype is constrained} \\
intersection_cut_coefficient (\alpha, \beta) & \text{vartype is continuous}
\end{cases}
\]

Figure A.8: Computing strengthened coefficients for intersection cuts

**Procedure** new_row_coefficient ($\tilde{a}_{kj}, \tilde{a}_{ij}, \gamma, \text{strengthen}, \text{vartype}$)

If we pivot on row $i$ and column $l$ (i.e. $\gamma = -\tilde{a}_{kl}/a_{ij}$) then:

- New row $k$:
  - $\tilde{a}_{ki}^\gamma = \gamma$
  - $\tilde{a}_{kj}^\gamma = \text{new_row_coefficient} (\tilde{a}_{kj}, \tilde{a}_{ij}, \gamma)$, for $j \in J \setminus \{ l \}$
  - $\tilde{a}_{k0}^\gamma = \text{new_row_coefficient} (\tilde{a}_{k0}, \tilde{a}_{i0}, \gamma)$

- New strengthened row $k$ (after modularization):
  - $\tilde{a}_{ki}^\gamma = \text{strengthened_row_coefficient} (\gamma, \tilde{a}_{ki}^\gamma, \text{typeof} (\{ i \}, i))$
  - $\tilde{a}_{kj}^\gamma = \text{strengthened_row_coefficient} (\tilde{a}_{kj}^\gamma, \tilde{a}_{k0}^\gamma, \text{typeof} (J \setminus \{ l \}, j))$, for $j \in J \setminus \{ l \}$
  - $\tilde{a}_{k0}^\gamma = \tilde{a}_{k0}^\gamma$

1. $\tilde{a}_{kj}^\gamma \leftarrow \tilde{a}_{kj} + \gamma a_{ij}$
2. if (strengthen) then
3.  $\tilde{a}_{kj}^\gamma \leftarrow \text{strengthened_row_coefficient} (\tilde{a}_{kj}^\gamma, \tilde{a}_{k0}^\gamma, \text{vartype})$ /* Strengthen the new coefficient */
4. end if
5. return $\tilde{a}_{kj}^\gamma$

Figure A.9: Computing the coefficient of a row after a prospective pivot ($\gamma$)
Procedure **pivot** \((\tilde{A}, \tilde{B}, J, i, l)\)

If we pivot on row \(i\) and column \(l\) (i.e. \(\gamma = -\frac{\tilde{a}_{kl}}{\tilde{a}_{il}}\)) then:

New row \(k\):
- \(\tilde{a}_{kj}^\gamma \leftarrow \tilde{a}_{kj} + \gamma \tilde{a}_{ij}\)
- \(\tilde{a}_{k0}^\gamma \leftarrow \tilde{a}_{k0} + \gamma \tilde{a}_{i0}\)

```plaintext
1 foreach (row \(k \neq i\)) do
   /* row \(k = row k + \gamma \text{ row } i*/
   2 \(\gamma \leftarrow -\frac{\tilde{a}_{kl}}{\tilde{a}_{il}}\)
   3 foreach \((j \in J \cup \{ l \})\) do
      4 \(\tilde{a}_{kj}^\gamma \leftarrow \tilde{a}_{kj} + \gamma \tilde{a}_{ij}\)
   5 end for
   6 \(\tilde{a}_{k0}^\gamma \leftarrow \tilde{a}_{k0} + \gamma \tilde{a}_{i0}\)
   7 end for
8 return \(\tilde{B}\)
```

Figure A.10: Pivoting in the \(\tilde{B}\) tableau
Procedure **strengthened_row_coefficient** \((\alpha, \beta, \text{vartype})\)

/* Strengthen the coefficients that correspond to constrained (i.e. structural) non-basic variables */
/* using the integrality conditions on these variables */

\[
\begin{cases}
\alpha - \lfloor \alpha \rfloor & \text{vartype is constrained and } \sigma \leq \beta, \text{ where } \sigma = \alpha - \lfloor \alpha \rfloor \\
\alpha - \lceil \alpha \rceil & \text{vartype is constrained and } \sigma > \beta, \text{ where } \sigma = \alpha - \lfloor \alpha \rfloor \\
\alpha & \text{vartype is continuous}
\end{cases}
\]

1 return

---

Figure A.11: Computing row coefficients resulting from disjunctive modularization

---

Procedure **compute_CGLP_reduced_cost** \((\tilde{A}, \tilde{B}, J, k, \pi, s, i, \text{selector})\)

/* selector = 1 means we compute the reduce cost for \(u_i\) */

1 \(M_i^1 \leftarrow \{ j \in J \mid \tilde{a}_{kj} < 0 \lor (\tilde{a}_{kj} = 0 \land \tilde{a}_{ij} > 0) \} \) if selector = 1

2 \(M_i^2 \leftarrow J \setminus M_i^1\)

REMARK: \(J\) corresponds to a basis in the original LP, while the partition \((M_i^1, M_i^2)\) corresponds to a basis in CGLP. Note that when there are no 0 entries in row \(k\), the partition \((M_i^1, M_i^2)\) is unique (i.e. same for all rows \(i\))!

IMPORTANT! The notation \(\pi_j\) does NOT (necessarily) mean that \(j\) indexes a slack variable!!! Because \(x_j = s_j = 0\) (from \(x \geq 0\)), Balas and Perregaard unified the notation by including under \(\pi_j\) the structural nonbasic variables \((\pi_{N \cap J})\) too! This is why we pass the whole \(\pi\) to this function (instead of just \(\pi_k\), which seems to be the only component of \(\pi\) needed)

3 \(\sigma \leftarrow \sum_{j \in M_i^2} \tilde{a}_{kj} s_j - \tilde{a}_{i0}(1 - \pi_k) \div \sum_{j \in J} |\tilde{a}_{kj}|, \quad \rho \leftarrow \sum_{j \in M_i^2} \tilde{a}_{ij} - \sum_{j \in M_i^1} \tilde{a}_{ij}\)

4 \(r \leftarrow \begin{cases} \sigma (+\rho - 1) - \sum_{j \in M_i^2} \tilde{a}_{ij} \tilde{s}_j + \tilde{a}_{i0}(1 - \pi_k) & \text{if selector = 1} \\
\sigma (-\rho - 1) - \sum_{j \in M_i^1} \tilde{a}_{ij} \tilde{s}_j + \tilde{a}_{i0}\pi_k & \text{otherwise}\end{cases}\)

5 return \(r\)

---

Figure A.12: Compute CGLP reduced cost
Procedure `typeof (J, j)`

1. return

| constrained | $j \in J$ and $x_j$ is constrained (structural only) |
| continuous  | $j \in J$ and $x_j$ is continuous (structural and slack/surplus) |
| ERROR       | $j \notin J$ |

Figure A.13: Determining the type of a variable

Procedure `intersection_cut_rhs (β)`

1. return $β(1 - β)$

Figure A.14: Computing rhs for intersection cuts

Procedure `create_intersection_cut (\tilde{A}, J, k)`

1. $\pi_0 \leftarrow \text{intersection_cut_rhs} (\tilde{a}_{k0})$ /* $\pi_0 \leftarrow \tilde{a}_{k0}(1 - \tilde{a}_{k0}) */$
2. $\pi_{j \in J} \leftarrow \text{intersection_cut_coefficient} (\tilde{a}_{kj}, \tilde{a}_{k0})$ /* $\pi_j \leftarrow \max \{ \tilde{a}_{kj}(1 - \tilde{a}_{k0}), -\tilde{a}_{kj}\tilde{a}_{k0} \} */$

3. return $(\pi, \pi_0)$

Figure A.15: Intersection cut

Procedure `create_strengthened_row (\tilde{A}, J, k)`

1. $\phi_0 \leftarrow \tilde{a}_{k0}$
2. $\phi_{j \in J} \leftarrow \text{strengthened_row_coefficient} (\tilde{a}_{kj}, \tilde{a}_{k0}, \text{typeof} (J, j))$
3. return $(\phi, \phi_0)$

Figure A.16: Disjunctive modularization (integrality based strengthening)
Procedure \texttt{create\_mixed\_integer\_gomory\_cut} ($\tilde{A}, J, k$)

1. $(\pi, \pi_0) \leftarrow \texttt{create\_intersection\_cut} (\tilde{A}, J, k)$
2. $\pi \leftarrow \texttt{strengthen\_cut} (\tilde{A}, J, k, \pi)$ \quad \COMMENT{Strengthen the cut using the integrality conditions on $x_{j \in J \setminus \{k\}}$}
3. return $(\pi, \pi_0)$

Figure A.17: Mixed integer Gomory cut

Procedure \texttt{strengthen\_cut} ($\tilde{A}, J, k, \pi$)

/* Strengthen the coefficients that correspond to constrained (i.e. structural) non-basic variables */
/* using the integrality conditions on these variables */
1. $\pi_{j \in J \cap \{1, \ldots, p\}} \leftarrow \texttt{strengthened\_intersection\_cut\_coefficient} (\tilde{a}_{k_j}, \tilde{a}_{k_0}, \texttt{typeof} (J, j))$
2. return $\pi$

Figure A.18: Cut strengthening using integrality conditions

Procedure \texttt{perturb\_row} ($\tilde{A}, \tilde{B}, k$)

1. $\tilde{a}_{kj} \leftarrow \epsilon^t, \forall \ j : \tilde{a}_{kj} = 0, t = 1, 2, \ldots$
2. return $\tilde{A}$

Figure A.19: Perturb a row of the $\tilde{B}$ tableau

Procedure \texttt{remove\_perturbation\_from\_row} ($\tilde{A}, \tilde{B}, k$)

1. $\tilde{a}_{kj} \leftarrow 0, \forall \ j : \tilde{a}_{kj} = \epsilon^t, t = 1, 2, \ldots$
2. return $\tilde{A}$

Figure A.20: Remove perturbation from a row of the $\tilde{B}$ tableau
**Procedure** XPRESS\_Get\_Tableau\_Row \((i)\)

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<thead>
<tr>
<th>Var</th>
<th>Size</th>
<th>Meaning</th>
</tr>
</thead>
<tbody>
<tr>
<td>(y)</td>
<td>rows</td>
<td>temporary array to store (c_i B^{-1})</td>
</tr>
<tr>
<td>(z)</td>
<td>rows + cols + 1</td>
<td>array in which we compute row (i)'s coefficients (and right and side)</td>
</tr>
<tr>
<td>(A^j)</td>
<td>rows</td>
<td>temporary array in which we retrieve column (j)</td>
</tr>
<tr>
<td>(b)</td>
<td>rows</td>
<td>temporary array in which we retrieve the right hand side</td>
</tr>
</tbody>
</table>

1. \(y \leftarrow c_i\) /* Set up \(y = c_i\) to pick row \(i\) */
2. \(y \leftarrow \text{XPRESS\_btran} (y)\) /* \(y = c_i B^{-1}\) */

3. **foreach** \((0 \leq j \leq \text{rows} - 1)\) **do**
   /* \(z = yA\) for each slack column of \(A\) */
   4. \(z_j \leftarrow y_j\)
   5. **end for**

6. **foreach** \((\text{rows} \leq j \leq \text{cols} - 1)\) **do**
   /* \(z_{\text{rows} + j} = yA^j\) for each structural column \(j\) of \(A\) */
   7. \(A^j \leftarrow \text{XPRESS\_getcols} (j)\) /* Retrieve column \(j\) from XPRESS */
   8. \(z_{\text{rows} + j} \leftarrow yA^j\)
   9. **end for**

10. \(b \leftarrow \text{XPRESS\_getrhs} ()\) /* Retrieve RHS from XPRESS */
11. \(z_{\text{rows} + \text{cols}} \leftarrow yb\)

**Important!** We now need to bring all nonbasic variables at a lower/upper bound of zero, because all formulas in the paper assume this fact!

12. \(z \leftarrow \text{XPRESS\_adjust\_row} (z)\)
13. **return** \(z\)

Figure A.21: Retrieving the tableau from XPRESS
Procedure XPRESS_Get_var_bound (j)

1 \[ v_j \left\{ \begin{array}{ll}
UB_j & , \ j \text{ is structural, at its upper bound} \\
LB_j & , \ j \text{ is structural, at its lower bound} \\
RANGE_{row(j)} & , \ j \text{ is slack, at its upper bound} \\
0 & , \ j \text{ is slack, at its lower bound}
\end{array} \right. \]

2 return \( v_j \)

Figure A.22: Function which returns the current bound of variable \( x_j \)

Procedure XPRESS_adjust_row (row)

\[
/* \text{row has rows + cols elements, the last one being the RHS} */
1 \text{foreach } (0 \leq j \leq \text{rows + cols - 1} | j \in J) \text{ do} \\
\text{/* bring each non-basic variable at a zero bound */}
2 \quad \text{bound}_j \leftarrow \text{XPRESS_Get_var_bound} (j) \\
3 \quad \text{if } (\text{var } j \text{ is at UB}) \text{ then} \\
4 \quad \quad \text{row}_j \leftarrow -\text{row}_j \quad \quad \quad /* \text{Var is at UB, replace with } x_j' = UB_j - x_j */
5 \quad \text{else} \\
6 \quad \quad \text{row}_{rows+cols} \leftarrow \text{row}_{rows+cols} + \text{row}_j \text{bound}_j \quad /* \text{adjust rhs to reflect the change of variable} */
7 \quad \text{end if} \\
8 \text{end for} \\
9 \text{return row}
\]

Figure A.23: Adjusting a row to reflect that all nonbasics are at a bound of zero
We want to compute:

\[ B^{-1}b - \sum_{j \in U} B^{-1}v_j A^j = B^{-1} \left( b - \sum_{j \in U} v_j A^j \right) \]

where \( U \) is the set of nonbasic variables at one of their bounds and \( v_j \) is the value of the bound at which variable \( j \) is (clearly, this only makes sense when variable \( j \) is at a bound (upper or lower) and that bound is non-zero.

```
1 b ← XPRESSgetrhs () /* Retrieve RHS from XPRESS */
2 /* Compensate for non-basic variables with non-zero values */
3 U ← \{ j ∈ J | variable (or slack) j is at a (non-zero) bound \}
4 foreach (j ∈ U) do /* look at all structural nonbasic variables at non-zero bounds */
5   bound_j ← XPRESS_get_var_bound (j) /* This test is here for efficiency, since a zero value would leave things unchanged! */
6     if (bound_j ≠ 0) then
7       if (j is structural) then
8         A^j ← XPRESS_getcols (j) /* Retrieve column j from XPRESS */
9         b ← b − bound_j A^j /* Note that this is a vector operation! */
10        end if
11      else
12        /* j is a slack, there is no column to retrieve */
13        Note! There is no need (or possibility) to retrieve column j - it’s the identity column with 1 in position i, where i is the row to which slack j corresponds. In fact, this column cannot even be retrieved from XPRESS - it is not stored explicitly (since we are dealing with columns of the original matrix, in which slacks don’t exist)!
14      end if
15    end if
16  end for
17  /* We now have b − \sum_{j ∈ U} A^j, so we multiply this with B^{-1} */
18  b ← XPRESS_ftran (b) /* rhs = B^{-1} \left( b - \sum_{j ∈ U} bound_j A^j \right) */
19 return b
```

Figure A.24: Computing the right hand side from XPRESS
Experimental results on cut modularization

The first experiment shows the effect of one round of cuts, with modularization applied only at the end of the pivoting sequence. In the second experiment, we apply modularization after each pivot. Both experiments were run with a limit of 50 cuts, and a maximum number of 20 pivots per cut. Tables A.2 (experiment 1) and A.4 (experiment 2) show the number of lift and project cuts generated for each problem, how many of them were actually optimal (i.e. we reached the CGLP minimum before exhausting the limit on the number of pivots), how many pivots we executed on average per cut, the density of the resulting cut (the number of non-zero coefficients in $\alpha$), as well as the average and total cut generation time, in CPU seconds. Finally, the amount of gap closed by cuts at the root node is shown in Tables A.3 and A.5.
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<th>Problem name</th>
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Table A.1: Lift and project: properties of the problems in MIPLIB 2003
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<th>Problem name</th>
<th>L&amp;P cuts</th>
<th>Optimal L&amp;P cuts</th>
<th>Avg pivots per cut</th>
<th>Avg cut density</th>
<th>Avg cut time</th>
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Table A.4: Lift and project: properties of modularized cuts
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<th>Percentage of gap closed by cuts</th>
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Table A.5: Lift and project: gap closed by modularized cuts