Scalable Heuristics for a Class of Chance Constrained Stochastic Programs

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We describe computational procedures for solving a wide-ranging class of stochastic programs with chance constraints where the random components of the problem are discretely distributed. Our procedures are based on a combination of Lagrangian relaxation and scenario decomposition, which we solve using a novel variant of Rockafellar and Wets’ progressive hedging algorithm. Experiments demonstrate the ability of the proposed algorithm to quickly find near-optimal solutions – where verifiable – to both difficult and very large chance constrained stochastic programs, both with and without integer decision variables. The algorithm exhibits strong scalability in terms of both CPU time required and final solution quality on large-scale instances.

Key words: Stochastic Programming, Chance Constraints, Scenario-Based Decomposition, Heuristics.


1. Introduction

Reliability considerations are always an intrinsic component of decision making under uncertainty and the design of good optimization models must necessarily include them. One way to do this is via a “penalty function” that measures the discrepancies between the decisions’ output and the potential future events, i.e., scenarios. This leads to stochastic programs with recourse where the recourse costs play the role of the penalty function (Kall and Wallace, 1994; Birge and Louveaux, 1997). Another possibility is to model non-compliance via a risk measure and optimize to minimize this risk measure; value-at-risk is a popular choice
(Holton, 2003). One could also ask explicitly for compliance by including in the model a probabilistic constraint. This leads to the formulation of a stochastic program with chance constraints, i.e., certain constraints will have to be satisfied with a high probability (Prekopa, 2003).

There are many real-world situations where chance constrained stochastic programs are an appropriate model. A commonly discussed example is the so-called 100-year disaster planning problem where a small fraction (e.g., one percent or less) of the scenarios do not have to be considered. Another example involves supply chain design, where it is typically not cost-effective to satisfy performance constraints in all realizable scenarios, as is the case in various forms of robust optimization. In general, the use of chance constraints provides modelers an opportunity to express the idea that constraints need not be satisfied or costs minimized across every conceivable eventuality.

Unfortunately, chance constrained stochastic programs are inherently difficult to solve except when they fall in a very narrow family. Stochastic programs are usually solved by relying on an approximating problem obtained via discretization of the probability space. Stochastic programs with recourse preserve convexity under discretization but, in general, that is not the case for stochastic programs with chance constraints; note however, that for problems with continuous variables, Nemirovski and Shapiro, by relying on “Bernstein approximations”, are able to build convex approximations for a relatively significant class of chance constrained programs (Nemirovski and Shapiro, 2006).

In this paper, we are concerned with efficient computational procedures for solving a wide ranging class of stochastic programs with chance constraints where the random components of the problem are discretely distributed, i.e., with finite support. We shall not be concerned if the original problem was already of this type, if it resulted from a discretization of the underlying probability distributions, or is obtained via a sampling scheme such as in the approach suggested by Luedtke and Ahmed (2008); cf. also Salinetti (1983), who deals with convergence issues, and Infanger (1993). Rather, we concern ourselves with the very important and practical problem of finding a good solution to a general chance constrained stochastic program once presented with a set of scenarios.

We formulate our decision model as a two stage stochastic program where some portion of the scenarios can be ignored. Each scenario $s$ has an associated weight or probability given as $p_s$. A formal statement of such a problem is as follows. Given a scenario index set $\mathcal{S}$ of size $|\mathcal{S}|$, a vector $x$ of $n$ first-stage decision variables, a cost vector $c$ of length $n$, and a
scalar \( 0 \leq \alpha < 1 \), find a vector \( x \in \mathbb{R}^n \) and binary vector \( d \) of length \(|S|\) to

\[
\begin{align*}
\text{minimize} & \quad c \cdot x + \sum_{s \in S} d_s (f_s \cdot y_s) \\
\text{subject to:} & \quad (x, y_s) \in Q_s, \quad \forall s \in \{S : d_s = 1\} \\
& \quad \sum_{s \in S} p_s d_s \geq (1 - \alpha) \\
& \quad d_s \in \{0, 1\}, \quad \forall s \in S
\end{align*}
\]  

(E)

where \( d_s \) represents the binary decision to enforce constraints \( Q_s \) for scenario \( s \in S \) and the \( y_s \) represent second-stage, scenario-specific decision vectors with associated cost coefficient vectors \( f_s \), which are determined given \( x \) and a particular \( s \in S \). The use of a common decision vector \( x \) for all \( s \in S \) implicitly implements the non-anticipativity constraints that avoid allowing the decisions to depend on the scenario. The \( Q_s \) summarize the problem constraints. Specific examples of (E) are provided in §5, where computational experiments concerning problems with both linear and mixed-integer constraints are described. Regardless of the nature of the constraints on \( x \), the interesting aspect of this formulation is the ability to select which scenarios are to be considered and which can be ignored.

Note that Problem (E) is non-convex due to the integrality of the \( d \) vector even if the rest of the problem happens to be convex. This significantly complicates solution procedures and inflates computational costs. This formulation is similar to one given by Ruszczyński (2002), who provided cutting planes and an exact algorithm useful for the case where the constraints are linear and \( x \) is real-valued. However, our interest here is in algorithms for more general (including non-convex) and computationally difficult forms (e.g., due to integer constraints), and for problem instances that are too large to be solved directly, and consequently requires decomposition to meet computer memory constraints. Ultimately, our goal is computational tractability, which has historically been a barrier to widespread adoption of chance constrained stochastic integer programs of the type occurring in many key application domains including logistics, transportation, supply chain management, and network design.

We conclude by observing that formulation (E) (ignoring the second-stage decision variables and costs for notational simplicity) is commonly used as a discretization of a standard chance constrained stochastic programming formulation (Ahmed and Shapiro, 2008; Ruszczyński, 2002), given as follows:

\[
\begin{align*}
\text{minimize} & \quad c \cdot x \\
\text{subject to:} & \quad \Pr\{G(x, \xi) \leq 0\} \geq (1 - \alpha)
\end{align*}
\]

where \( \xi \) is a random vector from a given probability distribution, \( G(x, \xi) \) represents the constraints associated with the random vector \( \xi \) (i.e., a scenario), and \( \alpha \) is the “risk” parameter,
i.e., the proportion of scenarios that we are willing to ignore.

The remainder of this paper is organized as follows. In §2, we discuss specific relaxations and decompositions of Problem (E). In §3, we introduce simple candidate algorithms based on scenario decomposition to solve instances of Problem (E). A more sophisticated algorithm is then introduced in §4. Computational experiments comparing the various algorithmic alternatives are presented in §5. We conclude by summarizing our contributions in §6.

2. Relaxation and Decomposition

Relaxation and decomposition are standard computational techniques for addressing situations involving both very large-scale and non-convex (or both) problems. First, we consider a Lagrangian relaxation of Problem (E), as follows:

\[
\begin{align*}
\text{minimize} & \quad c \cdot x + \sum_{s \in S} d_s p_s (f_s \cdot y_s) - \lambda \left( \sum_{s \in S} p_s d_s - (1 - \alpha) \right) \\
\text{subject to:} & \quad (x, y_s) \in Q_s, \quad \forall s \in \{S : d_s = 1\} \\
& \quad d_s \in \{0, 1\}, \quad \forall s \in S
\end{align*}
\]

For any fixed \( \lambda \geq 0 \), the optimal objective function value of Problem (L) is a lower bound on (E) as one would expect for the Lagrangian relaxation.

We further wish to exploit scenario decomposition, which in turn facilitates decomposition of \( x \) and \( d \). The decomposition of Problem (E) by scenarios (i.e., temporarily ignoring the coupling constraint) results in a problem of obtaining a solution \( x_s \) for each scenario \( s \in S \), as follows:

\[
\begin{align*}
\text{minimize} & \quad c \cdot x_s + f_s \cdot y_s \\
\text{subject to:} & \quad (x_s, y_s) \in Q_s
\end{align*}
\]

When we decompose (E) by scenarios we obtain a set of problems that are, even collectively, significantly easier to solve than the full Problem (E). Depending on the structure of the \( Q_s \), the subproblems can be tractable when the full problem is computationally out-of-reach. This situation is illustrated in our computational experiments described in §5.

For our purposes, a key aspect of scenario-based decomposition of Problem (L) is that \( d \) appears only in the coupling constraint. Thus, once the \( x_s \) variables for the scenario subproblems have been optimized, the optimal assignment of \( d_s \) variables for Problem (L) is
Let $x_s$ be the optimal value for a scenario subproblem of (L), i.e.,

$$\begin{align*}
& \text{minimize} \quad \tilde{c}_s - \lambda d_s |d_s \in \{0, 1\} \\
& \text{subject to:} \quad (x_s, y_s) \in Q_s \\
& \text{where:} \quad \tilde{c}_s = cx_s + f_s y_s
\end{align*}$$

This particular formulation borrows notation from Lulli and Sen (2004), who also address the problem of scenario selection for stochastic mixed-integer programs. Note that the optimal value of $x_s$ is independent of the value of $d_s$ for this single scenario problem. Given $x_s$ and $y_s$, there are two choices for $d_s$: if the optimal values would contribute a negative term in the objective of Problem (L), then $d_s = 1$; otherwise it is optimal to set $d_s$ to zero for that scenario. This observation is formalized as follows:

**Remark 1** When non-anticipativity in Problem (L) is relaxed, if $c \cdot x_s + f_s \cdot y_s \leq \lambda$, then $d_s = 1$ is optimal, otherwise $d_s = 0$ is optimal.

In order to exploit this decomposition, we need to deal with the fact that the decision vector $x$ cannot depend on $s$. If we could obtain an optimal or nearly optimal $x^*$ such that for all $s \in S$, $x_s = x^*$, we could immediately set the $d$ values and terminate the search. We accomplish this using algorithms described next in §3. Alternatively, when all scenarios have the same probability, then a very simple greedy algorithm is obtained by setting $d_s = 1$ for the $(1 - \alpha)|S|$ scenarios for which $c \cdot x_s + f_s \cdot y_s$ is lowest. This greedy algorithm is used as a computational baseline in §5.

### 3. Applications of Progressive Hedging for Scenario Selection

The progressive hedging algorithm proposed by Rockafellar and Wets (1991) provides a mechanism for combining scenario sub-problem solutions and enforcing non-anticipativity. Progressive hedging (PH) is sometimes referred to as a horizontal decomposition method because it decomposes the problem by scenarios rather than by time stages. The algorithm obtains solutions $x_s$ for each scenario $s$, and uses them to construct a unified solution.

The algorithm is particularly appropriate when methods exist to solve a deterministic version of the problem, but when multiple scenarios are introduced, solutions are not obtainable by directly solving the full problem due to either memory or time limitations. The
algorithm offers the additional advantage that it extends immediately to more than two time
stages and can be used heuristically for problems that are formulated with integer constraints
and other non-convexities. There are a couple of ways to gain insight or to motivate the
algorithm. One way is to think of it as blending the solutions for each scenario to form a
solution where decisions that cannot depend on the scenario do not. A more sophisticated
way is to think of non-anticipativity as a constraint of the form \( x_s = \bar{x} \) for all \( s \); then one
sees the algorithm as computing successive approximations to a sub-gradient multiplier for
this constraint.

In this section, we outline two ways to use PH to address the solution of Problem (E). The
first approach serves as a straightforward introduction to the PH algorithm, and can
solve Problem (E) when the \( d \) vector is given. This algorithm is used effectively as a post-
processor in §5. The second approach is an implementation for solving Problem (L) which
can then be embedded in a search for the smallest \( \lambda \) that results in \( \sum_{s \in S} p_s d_s \geq (1 - \alpha) \).
This algorithm is used as a comparative baseline in §5.2. A more sophisticated algorithm is
presented in §4 that simultaneously embeds the search for \( \lambda \) and \( d \) in the progressive hedging
algorithm.

3.1. PH for Problem (E) Given \( d \)

Given a particular \( d \) vector, PH for the solution of Problem (E) reduces to the basic PH
algorithm, which takes a perturbation vector \( \rho > 0 \) of length \( n \) and a convergence tolerance
\( \epsilon \) as input parameters (Rockafellar and Wets, 1991). Pseudo-code for PH in this context is
given as follows:

1. \( k = 0 \)

2. For all \( s \in S \), \( x_s^{(k)} = \arg\min_{x,y} (c \cdot x + f_s \cdot y_s) : (x, y_s) \in Q_s \)

3. \( \bar{x}^k = (\sum_{s \in S} p_s d_s x_s^{(k)}) / \sum_{s \in S} p_s d_s \)

4. For all \( s \in S \), \( w_s^{(k)} = \rho(x_s^{(k)} - \bar{x}^{(k)}) \)

5. \( k = k + 1 \)

6. For all \( s \in S \), \( x_s^{(k)} = \arg\min_{x,y} (c \cdot x + w_s^{(k-1)} x + \rho/2 \| x - \bar{x}^{(k-1)} \| ^2 + f_s \cdot y_s) : (x, y_s) \in Q_s \)

7. \( \bar{x}^{(k)} = (\sum_{s \in S} p_s d_s x_s^{(k)}) / \sum_{s \in S} p_s d_s \)
8. For all \( s \in S \), \( w_s^{(k)} = w_s^{(k-1)} + \rho \left( x_s^{(k)} - \bar{x}^{(k)} \right) \)

9. \( g^{(k)} = \frac{1}{\sum_{s \in S} p_s d_s} \sum_{s \in S} p_s d_s \| \frac{x_s^{(k)} - \bar{x}^{(k)}}{\bar{x}^{(k)}} \| \)

10. If \( g^{(k)} > \epsilon \), then go to step 5. Otherwise, terminate.

In the pseudocode, \( x_s^{(k)} \) denotes the value of \( x_s \) for scenario \( s \in S \) at iteration \( k \) of PH, while \( \bar{x}^{(k)} \) denotes the corresponding average \( x_s^{(k)} \) over all scenarios \( s \) with \( d_s = 1 \). The \( w_s^{(k)} \) are PH-specific, per-scenario vectors of length \( n \), and serve as the mechanism through which the \( x_s^{(k)} \) are eventually forced to agreement. PH is terminated once the scenario sub-problems are sufficiently homogeneous, as quantified by \( g^{(k)} \) and thresholded by \( \epsilon \). For each scenario \( s \in S \), the quantity \( g^{(k)} \) captures the difference between the selected (as determined by \( d_s \)) scenario solutions \( x_s^{(k)} \) and the average \( \bar{x}^{(k)} \), normalized in turn by \( \bar{x}^{(k)} \) to control for disparate variable scales; in a slight abuse of notation, we interpret the vector division operator as an element-wise operator. If Problem (E) is convex given a fixed \( d \), PH is guaranteed to locate optimal solutions given appropriate values of \( \rho \) and \( \epsilon \). Otherwise, the use of PH is as a heuristic, with the objective of quickly locating high-quality solutions.

3.2. PH for Problem (L) Given \( \lambda \)

Remark 1 enables modification of the PH algorithm given above in §3.1 by adding the logic:

If \( c \cdot x_s^{(k)} + f_s \cdot y_s \leq \lambda \) then \( d_s = 1 \) else \( d_s = 0 \)

to Steps 2 and 6. The result is that for a given \( \lambda \), a straightforward PH algorithm for Problem (L) can be stated as follows, again taking \( \rho > 0 \) and \( \epsilon \) as input parameters:

1. \( k = 0 \)

2. For all \( s \in S \), \( x_s^{(k)} = \text{argmin}_{x, y_s} (c \cdot x + f_s \cdot y_s) : (x, y_s) \in Q_s \)

   If \( (c \cdot x_s^{(k)} + f_s \cdot y_s) \leq \lambda \) then \( d_s = 1 \) else \( d_s = 0 \)

3. \( \bar{x}^k = \frac{\sum_{s \in S} p_s d_s x_s^{(k)}}{\sum_{s \in S} p_s d_s} \)

4. For all \( s \in S \), \( w_s^{(k)} = \rho (x_s^{(k)} - \bar{x}^{(k)}) \)

5. \( k = k + 1 \)
6. For all $s \in S$, 
\[
x_s^{(k)} = \text{argmin}_{x, y_s} \left( (c \cdot x + w_s^{(k-1)} x + \rho/2 \|x - \bar{x}^{(k-1)}\|^2 + f_s \cdot y_s) \right)
\]
\[
: (x, y_s) \in \mathcal{Q}_s
\]

If $(c \cdot x_s^{(k)} + f_s \cdot y_s) \leq \lambda$ then $d_s = 1$ else $d_s = 0$

7. $\bar{x}^{(k)} = \sum_{s \in S} p_s d_s x_s^{(k)} / \sum_{s \in S} p_s d_s$

8. For all $s \in S$, $w_s^{(k)} = w_s^{(k-1)} + \rho \left( x_s^{(k)} - \bar{x}^{(k)} \right)$

9. $g^{(k)} = \frac{1}{\sum_{s \in S} p_s d_s} \sum_{s \in S} p_s d_s \left\| \frac{x_s^{(k)} - \bar{x}^{(k)}}{x_s^{(k)}} \right\|$

10. If $g^{(k)} > \epsilon$, then go to step 5. Otherwise, terminate.

In Steps 2 and 6, the search for optimal $x$ and $d$ values are separated exploiting Remark 1. The intent behind the $g^{(k)}$ metric is identical to that discussed in §3.1. As a practical matter, Steps 3 and 7 must include a test for the possibility that $\sum_{s \in S} p_s d_s = 0$; in this case the algorithm terminates and reports that $\lambda$ is too small. Due to non-convexity, PH for the solution of Problem (L) is again a heuristic solution technique.

4. Progressive Hedging for Simultaneous Determination of $\lambda$, $d$, and $x$

In order to simultaneously determine $\lambda$, $d$, and $x$, we use progressive hedging to find values of $x_s$ that progressively satisfy the non-anticipativity requirement. In the process, we determine a minimal $\lambda$ that would result in $\sum_{s \in S} p_s d_s \geq (1 - \alpha)$ and set the $d_s$ values accordingly. In order to improve the stability of the $\bar{x}$ estimates (and therefore the $w$ estimates), we relax the strict binary constraints on the $d_s$ variables, and instead progressively bias the values of $d_s$ toward either 0 or 1 as the algorithm converges, as discussed in §4.1. The details of the enhanced PH algorithm are provided in §4.2.

4.1. Techniques for Biasing the $d_s$ Variables

In order to improve the stability of the PH algorithm in early iterations, we relax the binary restrictions on the $d_s$ variables. As the PH algorithm progresses the $d_s$ are gradually forced to become discrete. An effective and mathematically motivated mechanism to accomplish this behavior is via an augmentation function. We compose a piecewise step function with a sequence of mollifiers that serve to smooth the step augmentation functions; cf. (Ermoliev
Figure 1: A mollifier function, $\phi(\Delta; x)$, based on the Beta density function with parameters $a = 7$ and $b = 1.75$ when $\Delta = 1$.

et al., 1995) for a related approach when dealing with the minimization of discontinuous functions. Our “limiting” step function, with argument $\tau = c \cdot x_s + f_s \cdot y_s$, is given by:

$$
\psi(\tau, \lambda, \lambda_{\text{max}}) = \begin{cases} 
1 & \text{when } 0 \leq \tau \leq \lambda, \\
0 & \text{for } \lambda < \tau \leq \lambda_{\text{max}}
\end{cases}
$$

where $\lambda_{\text{max}}$ represents an upper bound on the cost $\tau$. Our collection of mollifiers are Beta density functions defined on the interval $[0, \Delta]$ with $\Delta \searrow 0$ as the PH algorithm converges:

$$
\phi(\Delta; x) = \begin{cases} 
\frac{1}{B(7, 1.75)} (x/\Delta)^6 (1 - x/\Delta)^{0.75} & \text{when } x \in [0, \Delta], \\
0 & \text{everywhere else}
\end{cases}
$$

where the Beta function $B(a, b) = \int_0^1 x^{a-1}(1 - x)^{b-1} dx$. In order to control the transition from a smooth representation of $d_s$ to a discrete one as PH converges, we use $\Delta = g^{(k)}/g^0$, i.e., the PH convergence gap. A graphical depiction of our particular parameterization $B(7, 1.75)$ is provided in Figure 1, for $\Delta = 1$.

Given the family of mollifier functions parameterized on $\Delta$, the corresponding augmentation functions are obtained via the convolution:

$$
m(\Delta, \lambda_{\text{max}}; x, \lambda) = \int_0^\Delta \psi(x - z, \lambda, \lambda_{\text{max}}) \phi(\Delta; z) \, dz, \quad x \in [0, \lambda_{\text{max}}].
$$

A graphical depiction of the augmentation function based on a Beta mollifier is provided in Figure 2.
4.2. Progressive Hedging for Scenario Selection

Given the family of augmentation functions $m(\cdot)$, a PH algorithm for scenario selection can be stated as follows:

1. $k = 0$ and $\Delta = 1$.

2. For all $s \in S$, $x_s^{(k)} = \arg\min_{x, y_s} (c \cdot x + f_s \cdot y_s) : (x, y_s) \in Q_s$

3. Assign an upper bound on the cost function to $\lambda_{\text{max}}$

4. $(\lambda, d) = \arg\min_{\lambda \leq \lambda_{\text{max}}} \lambda$ such that for all $s \in S$,

   $$d_s = m(\Delta, \lambda_{\text{max}}; c \cdot x_s^{(k)} + f_s \cdot y_s, \lambda) \quad \text{and} \quad \sum_{s \in S} p_s d_s \geq (1 - \alpha)$$

5. $\bar{x}^{(k)} = \sum_{s \in S} p_s d_s x_s^{(k)} / \sum_{s \in S} p_s d_s$

6. For all $s \in S$, $w_s^{(k)} = \rho(x_s^{(k)} - \bar{x}^{(k)})$

7. $g^{(k)} = \frac{1}{\sum_{s \in S} p_s d_s} \sum_{s \in S} p_s d_s \frac{|x_s^{(k)} - \bar{x}^{(k)}|}{x_s^{(k)}}$

8. $k = k + 1$

9. For all $s \in S$, $x_s^{(k)} = \arg\min_{x, y_s} (c \cdot x + w_s^{(k-1)} x + \rho / 2 \|x - \bar{x}^{(k-1)}\|^2 + f_s \cdot y_s) : (x, y_s) \in Q_s$

Figure 2: The Beta mollifier-based augmentation function $m(\Delta, \lambda_{\text{max}}; x, \lambda)$ given $\Delta = 1$, $\lambda_{\text{max}} = 3$, and $\lambda = 1$; $x$ is given on the horizontal axis.
10. \((\lambda, d) = \arg\min_{\lambda \leq \lambda_{\text{max}}} \lambda \) such that for all \(s \in S\),
\[
d_s = m(\Delta, \lambda_{\text{max}}; c \cdot x_s^{(k)} + f_s \cdot y_s, \lambda) \quad \text{and} \quad \sum_{s \in S} p_s d_s \geq (1 - \alpha)
\]

11. \(\bar{x}^{(k)} = \sum_{s \in S} p_s d_s^{(k)} x_s^{(k)} / \sum_{s \in S} p_s d_s\)

12. For all \(s \in S\), \(w_s^{(k)} = w_s^{(k-1)} + \rho \left(x_s^{(k)} - \bar{x}^{(k)}\right)\)

13. \(g^{(k)} = \frac{1}{\sum_{s \in S} p_s d_s} \sum_{s \in S} p_s d_s \|\frac{x_s^{(k)} - \bar{x}^{(k)}}{\bar{x}^{(k)}}\|\) and \(\Delta = g^{(k)} / g^{(0)}\)

14. If \(g^{(k)} > \epsilon\), then go to step 8. Otherwise, terminate.

We denote the above algorithm simply by SSPH. Recall that \(n\) is the number of first-stage decision variables in Problem (E). Steps 4 and 10 determine the value of \(\lambda\) that will result in the desired sum over the \(d\) vector. As can be seen from Figure 2, the values of \(d_s\) are monotonic in \(\lambda\) so the minimization can be done easily.

In Steps 4 and 10, the condition for achieving the target number of scenarios in which constraints are satisfied is based on non-binary (relaxed) \(d_s\). As SSPH converges, certain \(d_s\) are driven toward 0, and in practice once a \(d_s\) is “sufficiently” close to 0, it is highly unlikely that the same \(d_s\) will equal 1 in a fully converged solution. Further, the SSPH algorithm can exhibit “thrashing” behavior (i.e., spending large numbers of iterations attempting to eliminate very minor scenario solution differences) with \(d_s\) near 0, as these values lead to instability in the \(\bar{x}^{(k)}\) and \(g^{(k)}\). As a result, we impose \(d_s = 0\) once \(d_s\) drops below some threshold \(\gamma\) in both Steps 4 and 10 of the SSPH algorithm. For the experiments described in §5, we set \(\gamma = 0.10\) based on limited computational exploration.

In many situations, the solution of the extensive form (E) is computationally efficient given a binary \(d\) vector. In particular, this is generally the case for the network flow test problems described below in §5.1.1 and §5.1.2. In such cases, it is possible to prematurely terminate SSPH after Step 10 once (1) a binary \(d\) vector is obtained (following truncation relative to the threshold \(\gamma\)) and (2) \(\sum_{s \in S} p_s d_s = (1 - \alpha)\). The latter equality condition is required to prevent inclusion of excess scenarios (leading to cost inflation) and must be appropriately modified if equality is mathematically unattainable (e.g., via some achievable lower bound). The resulting \(d\) are then fixed in the extensive form (E), which is solved, for example, using a commercial MIP solver. We refer to this technique as a “quick exit” strategy. The baseline SSPH, which determines both \(d\) and the decision variable vectors \(x\) and \(y_s\), is referred to as the “full” (convergence) strategy.
5. Computational Experiments

We now examine the performance of the proposed PH algorithms for scenario selection in stochastic programming, considering two test cases: a small-scale, yet difficult “laboratory” problem and a large-scale, real-world problem. The advantage of the laboratory problem is that performance can be assessed relative to both established algorithms in the literature and extensive form solutions obtained via commercial mixed-integer programming solvers. In contrast, the extensive form of the real-world problem is not solvable with commercial solvers, and competing approaches have yet to be introduced. However, the size of the real-world problem instances serves to demonstrate the scalability and applicability of the proposed algorithms.

Both test cases are of a form that facilitates easy computation of an upper bound on cost, due to the absence of second-stage decision variables in the objective function. In particular, Step 3 in the SSPH algorithm is replaced by: \( \forall i \in [1..n], x_{\text{max}}(i) = \max_{s \in S} x_s^{(k)}(i) \) and \( \lambda_{\text{max}} = c \cdot x_{\text{max}}. \)

5.1. Ruszczyński’s Network Flow Model

Ruszczyński (2002) makes use of a chance constrained network flow problem to illustrate his method for solving Problem (E). We adopt and also extend this problem for use in our experiments. For the purpose of describing the formulation, we stay as close as possible to his notation, deviating only when necessary to avoid conflicts with our own. Ruszczyński’s notation allows for scenarios with different probabilities, but the example he used has equal probabilities, so we introduce the model formulation in the context of that assumption.

5.1.1. The Basic Model

We begin with a node set \( \mathcal{V} \) and a directed arc set \( \mathcal{A} \subseteq \mathcal{V} \times \mathcal{V} \). For each scenario \( s \in \mathcal{S} \) a quantity \( D_{kl}(s) \) must be shipped from node \( k \) to node \( l \) for all \( (k,l) \in \mathcal{V} \times \mathcal{V}, \ k \neq l \). The optimization objective is to determine arc capacities \( x(a), a \in \mathcal{A} \), that minimize the total cost \( \sum_{a \in \mathcal{A}} c(a)x(a) \) while enabling the shipping requirements (i.e., demands) to be met with probability \( 1 - \alpha \). The \( c(a) \) represent capacity cost parameters associated with each arc \( a \in \mathcal{A} \). Variables \( y_{kl}(a,s) \) are introduced to represent the flow from \( k \) to \( l \) passing through arc \( a \in \mathcal{A} \) in scenario \( s \in \mathcal{S} \). The problem formulation, which we denote (NF), is then given
as:

\[
\begin{align*}
\text{minimize} & \quad \sum_{a \in A} c(a)x(a) \\
\text{subject to:} & \quad \sum_{a \in A^+(\nu)} y_{kl}(a, s) - \sum_{a \in A^-(\nu)} y_{kl}(a, s) = \begin{cases} 
-D_{kl}(s) & \text{if } \nu = k \\
D_{kl}(s) & \text{if } \nu = l \\
0 & \text{otherwise}
\end{cases} \\
& \quad \forall \nu, k, l \in V, \forall s \in S \\
& \quad \sum_{s \in S} d_s \geq (1 - \alpha)|S| \\
& \quad x(a) \geq \sum_{k, l \in V} y_{kl}(a, s), \quad \forall a \in A, \forall s \in \{S : d_s = 1\} \\
& \quad x \geq 0, \ y \geq 0, \ d_s \in \{0, 1\}, \ \forall s \in S
\end{align*}
\]

where the notations $A^+(\nu)$ and $A^-(\nu)$ respectively indicate the set of arcs into and out of node $\nu$. In practice, $\alpha$ and $|S|$ are selected such that $(1 - \alpha)|S|$ is integer. The arc capacity constraint can be linearized as $x(a) \geq \sum_{k, l \in V} y_{kl}(a, s) - (1 - d_s)M$, where $M$ denotes a sufficiently large constant, e.g., the sum of all demands in scenario $s$.

The particular test case of (NF) used by Ruszczyński is a small example with 5 nodes and 7 arcs and is intended to demonstrate his method rather than to be the basis of extensive computational experiments (see Ruszczyński (2002) for a full description of the data used, including the specific network configuration and arc capacity costs). In this case all arcs are bi-directional and the capacities are assumed to be symmetric across an arc, i.e., if $a_1 = (i, j)$ and $a_2 = (j, i)$, then $x(a_1) = x(a_2)$ in any feasible solution. Similarly, the demand between each pair of distinct nodes is symmetric. The risk level $\alpha = 0.1$ was used in all experiments.

Building on this example, we generated ten random 100-scenario and five random 400-scenario (NF) instances using the procedure described in Ruszczyński (2002). In particular, the $D_{kl}(s)$ for a specific scenario $s \in S$ are given by $D_{kl}(s) = 0.1D_{s} + \epsilon_{kl}$, where $D_{s}$ represents the aggregate flow volume for scenario $s$, sampled from a normal distribution $N(30, 5)$; the $\epsilon_{kl}$ are independent normal variables with mean 0 and standard deviation 0.25.

For each instance, we compute the optimal solution of the (NF) extensive form using Ruszczyński’s exact method; we are grateful to Ruszczyński for sharing his AMPL code for solving (NF). Additionally, we allocate one day (1440 minutes) of CPU time to CPLEX 10.1 (ILOG, 2007) on these same instances. We then compare the performance of our SSPH algorithm against these baselines. In particular, we consider two variants of SSPH: one variant in which the “quick-exit” logic described in §4 is utilized, and another in which the algorithm is allowed to run to full convergence. In all SSPH runs, we define for each variable $x(a), a \in A$, the PH parameter $\rho_a = c(a)$; this particular choice was based on our prior experience with variable-dependent $\rho$ values (Watson et al., 2007). In all runs, $\epsilon = 0.01$. 

13
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<th>Instance</th>
<th>CPLEX Obj.</th>
<th>CPLEX T.</th>
<th>Ruszczyński Obj.</th>
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<th>Greedy T.</th>
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% Gap | 0.0 | 0.0 | 0.185 | 0.081 | 0.049 |
Time  | 53.75 | 4.4 | 0 | 0.15 | 3.05 |

Table 1: Performance results for various scenario selection algorithms on 100-scenario network flow problems. For each algorithm, the total cost (Obj. labeled columns) and run time (T. labeled columns) are reported. Run-time units are minutes, rounded to the nearest half minute increment. The final two rows record the average percentage above optimal solution cost (row labeled “Gap”) and the average run-time (row labeled “Time”).

Upon termination, both variants of SSPH have identified a candidate set of scenarios, as defined by the resulting $d_s$ variables. The $d_s$ are then fixed in Problem (NF), yielding a straightforward linear program which is then solved using CPLEX. Finally, for an additional baseline, we compare the performance of SSPH variants relative to that of a simple greedy approach that selects the least expensive $(1 - \alpha)|S|$ scenarios, as described in §2. As with SSPH, the obtained $d_s$ are then fixed in Problem (NF), and the resulting linear program is solved via CPLEX. The times reported for the SSPH and greedy algorithms include all overhead processing and final CPLEX linear program solve times. All execution times are rounded to the nearest half minute increment. All experiments were executed on a 2.2GHz AMD Athlon architecture running Linux, with 64GB of RAM.

We first consider the results for the 100-scenario instances, which are summarized in Table 1. Consistent with the statements made in (Ruszczyński, 2002), Ruszczyński’s method significantly outperforms CPLEX, obtaining optimal solutions in roughly an order of magnitude less time. This is despite the advances in CPLEX solver technology since (Ruszczyński, 2002) appeared. The baseline greedy scenario selection approach performs remarkably well, achieving solutions on average only 0.185% above optimal in a few seconds of run-time; in half of the instances, the greedy solution is optimal. This result is likely due to the sampling
<table>
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| % Better | 0.0       | 4.441   | 3.744            | 3.872         | 4.220       |
| % Gap    | 4.72      | 0.0     | 0.73             | 0.60          | 0.22        |
| Time     | 1440      | 3323    | 0                | 1.5           | 13.5        |

| Table 2: Performance results for various scenario selection algorithms on 400-scenario network flow problems. For each algorithm, the total cost (Obj. labeled columns) and run time (T. labeled columns) are reported. Run-time units are minutes, rounded to the nearest half minute increment. The final three rows record the average percentage quality improvement over the CPLEX solution (row labeled “% Better”), the average percentage above optimal solution cost (row labeled “Gap”), and the average run-time (row labeled “Time”). |

procedure used to construct the problem instances; the $\epsilon_{kl}$ are sampled from a relatively tight distribution, such that there is no significant overlap in demands $D_{kl}(s)$ across scenarios $s \in \mathcal{S}$. On average, SSPH with the quick exit strategy enabled yields improvements over greedy scenario selection with only slight increases in run-time. SSPH with full convergence requires relatively longer run-times, but they remain lower than those obtained by Ruszczyński’s method; solutions were only 0.0488% above optimal on average, and optimal solutions were identified in eight instances.

Overall, the heuristic scenario selection algorithms were able to obtain very high-quality solutions in minimal run-times, relative to the extensive form solves via CPLEX. Clearly, Ruszczyński’s exact method is the preferred algorithm for these particular small instances. Rather, the results reported in Table 1 serve as a preliminary demonstration of the potential effectiveness of SSPH.

Next, we consider the results for the 400-scenario instances, which are summarized in Table 2. Here, we limited the run-time of CPLEX on each instance to 1440 minutes, i.e., 1 day. In no case did CPLEX prove optimality, and the optimality gaps at termination were significant. Despite the apparent simplicity of the (NF) problem formulation, obtaining high-quality solutions for moderate numbers of scenarios appears problematic. This is confirmed by the results we obtained via Ruszczyński’s method; within the allocated 1440 minutes of run-time, the method was only able to complete roughly 13-14 iterations on any given
problem instance, with some individual iterations requiring greater than half the allocated time. In no case was the algorithm able to identify a solution within the time limit. We did eventually allow runs of Ruszczyński’s method to complete, which required several days of computation; the results are reported in Table 2. These extended-duration runs allow us to obtain optimal solutions, which we use to assess the absolute performance of our heuristics. In contrast, CPLEX was unable to identify optimal solutions on any instance given a week of run-time.

Surprisingly, even greedy scenario selection obtains solutions over 3.7% better than those achieved by CPLEX, in less than 10 seconds for all instances. SSPH with the quick exit strategy yields slight improvements over greedy (to nearly 4% better than CPLEX) in reasonable run-times (≈ 1.5 minutes on average). Executing SSPH to full convergence results in further improvements, albeit at the expense of increased (but still reasonable) run-times. In absolute terms, the SSPH algorithm with full convergence is able to locate very near-optimal solutions in orders-of-magnitude less time than Ruszczyński’s method, and is able to locate higher quality solutions than CPLEX, again in a fraction of the run-time.

Overall, both SSPH algorithm variants (quick-exit and full convergence) yield improvements relative to the greedy algorithm, while still executing in reasonable run-times. In contrast to both exact methods, SSPH is conclusively more scalable; even at 400 scenarios on a 5-node instance, the exact methods are encountering serious performance issues.

5.1.2. An Extended Model With Arc Budgets

To further investigate the scalability of SSPH relative to the direct solution of the extensive form via CPLEX, we consider a slightly more complicated version of Problem (NF). To increase computational complexity, we add a per-scenario budget for the total number of arcs that can be used in a solution. For each scenario \( s \in S \) and arc \( a \in A \), we introduce a binary variable \( b(a, s) \). In each scenario \( s \in S \), an arc \( a \in A \) can be made active, subject to a total budget on the number of active arcs \( B(s) \). Given the \( M \) constant defined in Section 5.1.1, we add the following constraints to the (NF) problem formulation:

\[
\begin{align*}
y_{kl}(a, s) &\leq M \cdot b(a, s), \quad \forall k, l \in V, \forall a \in A, \forall s \in S \\
\sum_{a \in A} b(a, s) &\leq B(s), \quad \forall s \in S \\
b(a, s) &\in \{0, 1\}, \quad \forall a \in A, \forall s \in S
\end{align*}
\]
Table 3: Performance results for various scenario selection algorithms on 100-scenario budget network flow problems. For each algorithm, the total cost (Obj. labeled columns) and run-time (T. labeled columns) are reported. Run-time units are minutes, rounded to the nearest half minute increment. The final two rows record the average objective percentage improvement over the CPLEX solution (row labeled “% Better”) and the average run-time (row labeled “Time”).

To maintain consistency with the arc symmetry budget described in §5.1.1, we additionally impose a corresponding budget symmetry constraint, i.e., if \( a_1 = (i,j) \) and \( a_2 = (j,i) \) for two arcs \( a_1 \) and \( a_2 \), then \( b(a_1, s) = b(a_2, s) \) must hold for all \( s \in S \), i.e., “opening” an arc for use in one direction opens the arc in the opposite direction. We denote the resulting formulation by (BNF), which is an acronym for Budget Network Flow.

We mirror the experimental methodology described in §5.1.1 to generate random instances of Problem (BNF). The \( B(s) \) for each problem instance and scenario \( s \in S \) are independently sampled as follows: with probability 0.8, \( B(s) = 12 \); otherwise, \( B(s) = 10 \). Due to the arc symmetry constraints, this sampling implies that either 5 or 6 of the total 7 (symmetric) arcs may be activated. We again generate ten and five 100 and 400 scenario instances, respectively. With the exceptions noted below, we replicate the experimental methodology described in §5.1.1. In the case of Problem (BNF), our sole comparative baseline is CPLEX for solving the extensive form. SSPH parameters are identical to those described for the experiments presented in §5.1.1. Upon termination of both the greedy and SSPH heuristics, the \( d_s \) are appropriately fixed in Problem (BNF) and the corresponding mixed-integer program is solved with CPLEX. Further, the CPLEX solve is warm-started by assigning the second-stage \( b(a, s) \) variables to the heuristically obtained values; this assignment yields
moderate acceleration of CPLEX run-times. For 100-scenario instances, the resulting MIPs solve in reasonable run-times to optimality. This is not the case for 400-scenario instances; as a result, we terminate CPLEX once the first feasible incumbent is identified. All reported greedy and SSPH run-times include the cost of this final MIP solve. In practice, the necessity of this final solve could be mitigated by selecting a smaller value for the PH convergence parameter $\epsilon$.

First, we consider the results for the 100-scenario (BNF) instances, as reported in Table 3. In contrast to the (NF) results, CPLEX is unable to identify optimal solutions within the allocated run-time budget of 1440 minutes; at termination, the optimality gap is again significant (ranging from just under 2% to over 10%). We assess the performance of our heuristic scenario selection algorithms relative to the CPLEX baseline. The greedy algorithm obtains better solutions than CPLEX on average (a slight improvement of 0.184%), in substantially less run-time ($\approx 71$ minutes), although CPLEX does outperform the greedy algorithm on half of the instances. Nearly all of the run-time associated with the greedy algorithm is attributable to the cost of solving the mixed-integer program resulting from the fixing of the $d_s$ scenario selection variables. These models can be very difficult, in contrast to the linear programs resulting from the fixing of the $d_s$ in the (NF) formulation. Further, there is no possibility of “warm-starting” the fixed-$d_s$ CPLEX solve due to the lack of converged $x(a)$. SSPH with the quick exit strategy enabled yields further improvements in solution quality (0.787% better than CPLEX). On average, we observe an increased run-time relative to greedy, but this is due to the specific results for instance 10; for this instance, the initial incumbent solution yielded an optimality gap of 0.02%, and CPLEX failed to prove optimality within the 1440 minute time allocation. In general, SSPH (under both convergence criteria) often counterintuitively requires less run-time than the greedy algorithm. This is due to the ability of SSPH to warm-start MIP solves across PH iterations, as a solution to a scenario $s \in S$ obtained in PH iteration $k$ remains feasible (but typically sub-optimal with respect to cost) in PH iteration $k + 1$. Finally, the fully converged SSPH algorithm obtains the best overall performance (a 0.96% improvement over CPLEX), in roughly an hour of run-time on average.

Finally, we consider the results for the 400-scenario (BNF) instances, as reported in Table 4. Relative to the results reported in Table 3, the run-times are proportionally lower due to our limiting the run-time of the final MIP solve (using fixed $d_s$ values) in the 400-scenario case. The results exhibit similar patterns to those obtained for the (NF) and 100-
Table 4: Performance results for various scenario selection algorithms on 400-scenario budget network flow problems. For each algorithm, the total cost (Obj. labeled columns) and run-time (T. labeled columns) are reported. Run-times are reported in minutes, rounded to the nearest half minute. The final two rows record the average objective percentage improvement over the CPLEX solution (row labeled “% Better”) and the average run-time (row labeled “Time”).

5.2. An Aircraft Sustainability Planning Model

Our work on scenario selection heuristics for stochastic programming was originally motivated by a real-world application involving the allocation of spare parts and part repair-related resources to a large aircraft maintenance operation. The objective in allowing a certain proportion of infeasible scenarios in this application is to facilitate investigation into the risk-reward trade-off: the maintenance contractor may accept a certain probability $\alpha$ of failing to meet target performance criteria (e.g., the average percentage of time that aircraft in the fleet are available to fly) in exchange for a reduced capital expenditure in the deployment of the associated supply and repair chains. Monetary penalties are imposed if performance targets are not achieved.

This aircraft sustainability problem can be formulated as a stochastic mixed-integer program with the following characteristics. Given a cost vector $c \geq 0$ of length $n$, a scenario set $\mathcal{S}$ of size $|\mathcal{S}|$, a scalar $0 < \alpha \leq 1$, matrices $A(s) \geq 0$ each of which is of dimension $m$ by $n$, and $b(s)$ vectors each of length $m$ for all $s \in \mathcal{S}$, find a vector $x$ of length $n$ and vector $d$ of...
minimize \quad c \cdot x \quad \text{(ASP)}
subject to: \quad A(s)x \geq b(s)d_s, \quad \forall s \in S
\quad \sum_{s \in S} d_s \geq (1 - \alpha)|S|
\quad d_s \in \{0, 1\}, \quad \forall s \in S
\quad x \in (\mathbb{Z}^+)^n

In this formulation, which we denote (ASP), the \( A(s) \) and \( b(s) \) associated with a given scenario \( s \in S \) represent constraints that model a discrete event simulation (see (Savage et al., 2005)) of the sustainability operation, subject to a given realization of aircraft part failures. We assume that the scenario realization probabilities \( p_s \) are uniformly distributed. The variables \( x \) encode inventory policy parameters, resource levels, and time-indexed state tracking variables. The latter represent a dominant fraction of the total number of variables, which reaches over a million for the largest instances we consider below. For any scenario \( s \in S \), the number of constraints reaches over a million, with the corresponding number of non-zeros in \( A(s) \) reaching 10 million. Details of the (ASP) formulation, a scenario solver, and an enhanced PH algorithm (for models without chance constraints) are described by Watson et al. (2007). In particular, mechanisms for solving scenario subproblems heuristically, setting and varying the PH parameter \( \rho \), accelerating convergence, and economically terminating PH are introduced.

Clearly, the defining characteristic of Problem (ASP) is its size, which allows us to investigate the run-time scalability of our scenario selection heuristics. On the other hand, the scale and novelty of the formulation prevents us from obtaining a comparative performance baseline; CPLEX requires tens of gigabytes of RAM to store the extensive form of our largest problem instances, and solution of even the linear relaxation of Problem (ASP) can require several days of run-time. The problem is representative of industrial scales, for which solutions must be obtained despite the lack of a provably optimal comparative baseline. We compare the following algorithms:

- **Greedy**: Solve all scenario subproblems independently, select the resulting \((1 - \alpha)|S|\) lowest-cost scenarios (setting \( d_s = 1 \) for the lowest-cost scenarios and \( d_s = 0 \) otherwise),
and use PH (as described in §3.1) to solve the related stochastic mixed-integer program consisting strictly of scenarios \( s \in S \) with \( d_s = 1 \).

- **SSPH**: Use the SSPH algorithm described in §4 to simultaneously determine \( \lambda \), a scenario vector \( d \), and a solution vector \( x \). The “full” convergence form of SSPH is used, due to the computational difficulty of the extensive form (E) even when \( d \) is given.

- **SSPH+**: First, execute SSPH as above to obtain a \( d \) vector. Then, form a stochastic mixed-integer program consisting strictly of scenarios \( s \in S \) where \( d_s = 1 \). Finally, use PH (as described in §3.1) to solve the resulting program.

In the case of Greedy and SSPH+, the PH algorithm *without* scenario selection is being used in a “touch-up” role, analogous to our use of CPLEX in conjunction with scenario selection heuristics as described in §5.1. As discussed above, the computational difficulty and scale of Problem (ASP) prevents exact solution of the extensive form even when the scenario selection constraint is relaxed. In contrast to our experiments with Problems (NF) and (BNF), we do not use the quick-exit SSPH strategy; accelerator techniques designed to exploit the general structure of Problem (ASP) mitigate the need for such a strategy. We defer to Watson et al. (2007) for a detailed description of these techniques.

We execute each heuristic algorithm on each of three synthetic (ASP) test instances, of varying size. For all trials, we arbitrarily use \( \alpha = 0.2 \). The computational platform is identical to that described above in §5.1. For each instance and algorithm combination, we record the overall run-time and the cost of the best solution obtained. The time and cost units are respectively minutes and USD. As previously noted, only a small fraction of the variables in the \( x \) vector require blending by progressive hedging to enforce non-anticipativity; the exact number ranges from 144 to 528 for the smallest and largest instances, respectively. All instances consist of 30 scenarios, which is deemed sufficient by the end-user given multi-year planning horizons and comparatively frequent aircraft part failures.

The obtained solution costs are reported in Table 5. The instances are characterized by the first three columns of each row and the remaining columns record the solution costs for each algorithm. For reference, we also provide two additional columns of results. The column labeled “All \( d = 1 \)” records the solution cost obtained by solving Problem (ASP) with \( \alpha = 0 \) using PH. The column labeled “\( \lambda \) Search” records the solution cost obtained by embedding
Table 5: Costs of solutions obtained using various scenario selection heuristics (see text), on a range of aircraft sustainability planning problem instances. The \( n \) and \( m \) values respectively represent the number of variables and constraints in the extensive form of the (ASP) stochastic mixed-integer program.

| \( n \) | \( m \) | \( |\mathcal{S}| \) | All \( d = 1 \) | Greedy | \( \lambda \) Search | SSPH | SSPH+ |
|---|---|---|---|---|---|---|---|
| 5M | 4M | 30 | 60,264,200 | 58,330,300 | 58,076,200 | 57,878,700 | 57,287,650 |
| 11M | 8M | 30 | 133,909,900 | 126,140,650 | 125,291,550 | 126,021,300 | 125,448,300 |
| 22M | 16M | 30 | 359,249,300 | 345,419,550 | 346,937,100 | 345,000,400 | 344,800,320 |

Table 6: Algorithm run-times in minutes on a 2.2GHz AMD Athlon Linux workstation, for the results reported in Table 5.

| \( n \) | \( m \) | \( |\mathcal{S}| \) | All \( d = 1 \) | Greedy | \( \lambda \) Search | SSPH | SSPH+ |
|---|---|---|---|---|---|---|---|
| 5M | 4M | 30 | 94 | 64 | 669 | 85 | 85+58 |
| 11M | 8M | 30 | 365 | 304 | 3949 | 377 | 377+386 |
| 22M | 16M | 30 | 3715 | 3470 | 28069 | 3356 | 3356+3287 |

the PH algorithm for a given \( \lambda \) (as given in §3.2) in a binary search for an optimal \( \lambda \). The initial lower and upper bounds \( \lambda_l \) and \( \lambda_u \) are respectively given as (1) the minimum solution cost for an individual scenario identified in PH iteration 0 and (2) the cost of the solution obtained by taking an element-wise maximum of the \( x_s \) vectors obtained for all scenarios in PH iteration 0. The binary search is terminated once \((\lambda_h - \lambda_l)/\lambda_l \cdot 100 \leq 0.01\).

Necessarily, allowing infeasibility in a proportion \( \alpha = 0.2 \) of scenarios yields significant reductions in overall solution cost. While no clear “winner” exists among the scenario selection heuristics, we observe that SSPH+ dominates SSPH, which in turn dominates the greedy algorithm. SSPH dominates \( \lambda \) search on all but the medium-sized instance, while \( \lambda \) search underperforms the greedy algorithm on the largest instance. Although we cannot draw general conclusions given the small number of test instances, the preliminary evidence suggests that SSPH and SSPH+ consistently provides high-quality solutions.

Next, we consider the relative run-times of the various algorithms, as reported in Table 6. The computational difficulty of Problem (ASP) is illustrated by the absolute run-times for the baseline PH algorithm with \( d_s = 1 \) for all \( s \in \mathcal{S} \) (§3.1). The key observation is that SSPH requires no more time to compute solutions than the baseline PH algorithm, i.e., the additional complexity associated with determination of the \( d \) vector does not translate into increased computational costs. The run-time cost of the greedy algorithm is dominated by the PH post-processing run; the analogous post-processing in SSPH+ acts to roughly double
the base SSPH run-time, with the benefit of lower-cost solutions. Finally, while PH-based \( \lambda \) search (§3.2) can locate solutions competitive with SSPH and SSPH+, the run-time cost is prohibitive; typically 10–15 outer loop iterations are required to achieve convergence.

6. Discussion and Conclusions

We have described computational procedures based on Lagrangian relaxation and scenario decomposition to heuristically solve a wide ranging class of stochastic programs with chance constraints where the random components of the problem are discretely distributed, with or without integer variable constraints. Given their well-known difficulty even for small instances, an effective and scalable method for such problems is an important addition to the suite of tools available for optimization under uncertainty.

Computational experiments demonstrate the ability of the proposed heuristics to find near-optimal solutions to small, yet very difficult laboratory test examples, in orders of magnitude less run-time than exact algorithms. On larger instances of laboratory test examples, the heuristics yield higher-quality solutions than commercial solvers, again in orders-of-magnitude lower run-times. We also report tests on a very large real-world example, which demonstrate that scenario selection analysis can be performed with modest additional computational effort relative to solving the corresponding stochastic programs in which the chance constraints are relaxed.

The use of chance constraints provides a modeler the opportunity to express the idea that constraints need not be satisfied or costs minimized across every conceivable eventuality. Efficient and scalable heuristics for scenario selection allow end-users to quickly explore risk-reward trade-offs. Further, such heuristics may be effectively leveraged to significantly accelerate exact algorithms, by providing initial near-optimal incumbents.

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