

PySP: Modeling and Solving Stochastic Programs in Python

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Abstract Although stochastic programming is a powerful tool for modeling decision-making under uncertainty, various impediments have historically prevented its widespread use. One key factor involves the ability of non-specialists to easily express stochastic programming problems as extensions of deterministic models, which are often formulated first. A second key factor relates to the difficulty of solving stochastic programming models, particularly the general mixed-integer, multi-stage case. Intricate, configurable, and parallel decomposition strategies are frequently required to achieve tractable run-times. We simultaneously address both of these factors in our PySP software package, which is part of the COIN-OR CoopR open-source Python project for optimization. To formulate a stochastic program in PySP, the user specifies both the deterministic base model and the scenario tree with associated uncertain parameters in the Pyomo open-source algebraic modeling language. Given these two models, PySP provides two paths for solution of the corresponding stochastic program. The first alternative involves writing the extensive form and invoking a standard deterministic (mixed-integer) solver. For more complex stochastic programs, we provide an implementation of Rockafellar and Wets' Progressive Hedging algorithm. Our particular focus is on the use of Progressive Hedging as an effective heuristic for approximating

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general multi-stage, mixed-integer stochastic programs. By leveraging the combination of a high-level programming language (Python) and the embedding of the base deterministic model in that language (Pyomo), we are able to provide completely generic and highly configurable solver implementations. PySP has been used by a number of research groups, including our own, to rapidly prototype and solve difficult stochastic programming problems.

1 Introduction

The modeling of uncertainty is widely recognized as an 必需 integral component in most real-world decision problems. Typically, uncertainty is associated with the problem input parameters, e.g., consumer demands or construction times. In cases where parameter uncertainty is independent of the decisions, stochastic programming is an appropriate and widely studied mathematical framework to express and solve uncertain decision problems [Birge and Louveaux, 1997, Wallace and Ziemba, 2005, Kall and Mayer, 2005a, Shapiro et al., 2009]. However, stochastic programming has not yet seen widespread, routine use in industrial applications – despite the significant benefit such techniques can confer over deterministic mathematical programming models.

The growing practical importance of stochastic programming is underscored by the recent proposals for and additions of capabilities in many commercial algebraic modeling languages [AIMMS, Valente et al., 2009, Xpress-Mosel, 2010, Maximal Software, 2010].

Over the past decade, two key impediments have been informally recognized as central in inhibiting the widespread industrial use of stochastic programming. First, modeling systems for mathematical programming have only recently begun to incorporate extensions for specifying stochastic programs. Without an integrated and accessible modeling capability, practitioners are forced to implement custom techniques for specifying stochastic programs. Second, stochastic programs are often extremely difficult to solve – especially in contrast to their deterministic mixed-integer counterparts. There exists no analog to CPLEX [CPLEX, 2010], Gurobi [GUROBI, 2010], Maximal [Maximal Software, 2010], or XpressMP [XpressMP, 2010] for stochastic programming, principally because the algorithmic technology is still under active investigation and development, particularly in the multi-stage and mixed-integer cases.

Some commercial vendors have recently introduced modeling capabilities for stochastic programming, e.g., LINDO Systems [LINDO, 2010], FICO [XpressMP, 2010], and AIMMS [AIMMS]. On the open-source front, the options are even more limited. FLOPC++ (part of COIN-OR) [FLOPCPP, 2010] provides an algebraic modeling environment in C++ that allows for specification of stochastic linear programs. APLEpy provides similar functionality in a Python programming language environment. In either case, the available modeling extensions have not yet seen widespread adoption.

The landscape for solvers (open-source or otherwise) devoted to generic stochastic programs is extremely sparse. Those modeling packages that do provide stochastic programming facilities with few exceptions rely on the translation of problem into the *extensive form* – a deterministic mathematical programming representation of the stochastic program in which all scenarios are explicitly represented and solved simultaneously. The extensive form can then be supplied as input to a standard (mixed-integer) linear solver. Unfortunately, direct solution of the extensive form is unrealistic in all but the simplest cases. Some combination of either the number of scenarios, the number of decision stages, or the presence of discrete decision variables typically leads to extensive

forms that are either too difficult to solve or exhaust available system memory. Iterative decomposition strategies such as the **L-shaped method** [Slyke and Wets, 1969] or **Progressive Hedging** [Rockafellar and Wets, 1991] directly address both of these scalability issues, but introduce fundamental parameters and algorithmic challenges. Other approaches include **coordinated branch-and-cut procedures** [Alonso-Ayuso et al., 2003]. In general, the solution of difficult stochastic programs requires both experimentation with and customization of alternative algorithmic paradigms – necessitating the need for generic and configurable solvers.

In this paper, we describe an open-source software package – PySP – that begins to address the issue of the availability of generic and customizable stochastic programming solvers. At the same time, we describe modeling capabilities for expressing stochastic programs. Beyond the obvious need to somehow express a problem instance to a solver, we identify a fundamental characteristic of the modeling language that is in our opinion necessary to achieve the objective of generic and customizable stochastic programming solvers. In particular, the modeling layer must provide mechanisms for accessing components via direct introspection [Python, 2010b], such that solvers can generically and programmatically access and manipulate model components. Further, from a user standpoint, the modeling layer should differentiate between abstract models and model instances – following best practices from the deterministic mathematical modeling community [Fourer et al., 1990].

To express a stochastic program in PySP, the user specifies both the **deterministic base model** and **the scenario tree model with associated uncertain parameters** in the Pyomo open-source algebraic modeling language [Hart et al., 2010]. This separation of deterministic and stochastic problem components is similar to the mechanism proposed in SMPS [Birge et al., 1987, Gassmann and Schweitzer, 2001]. Pyomo is a Python-based modeling language, and provides the ability to model both abstract problems and concrete problem instances. The embedding of Pyomo in Python enables model introspection and the construction of generic solvers.

Given the deterministic and scenario tree models, PySP provides two paths for the solution of the corresponding stochastic program. The first alternative involves writing the extensive form and invoking a deterministic (mixed-integer) linear solver. For more complex stochastic programs, we provide a generic implementation of Rockafellar and Wets’ Progressive Hedging algorithm [Rockafellar and Wets, 1991]. Our particular focus is on the use of Progressive Hedging as an effective heuristic for approximating general multi-stage, mixed-integer programs. By leveraging the combination of a high-level programming language (Python) and the embedding of the base deterministic model in that language (Pyomo), we are able to provide completely generic and highly configurable solver implementations. Karabuk and Grant [2007] describe the benefits of Python for such model-building and solving in more detail.

The remainder of this paper is organized as follows. We begin in Section 2 with a brief overview of stochastic programming, focusing on multi-stage notation and expression of the scenario tree. In Section 3, we describe the PySP approach to modeling a stochastic program, illustrated using a well-known introductory model. PySP capabilities for writing and solving the extensive form are described in Section 4. In Section 5, we compare and contrast PySP with other open-source software packages for modeling and solving stochastic programs. Our generic implementation of Progressive Hedging is described in Section 6, while Section 7 details mechanisms for customizing the behavior of PH. By basing PySP on Coopr and Python, we are able to provide straightforward mechanisms to support distributed parallel solves in the context of PH and other de-

composition algorithms; these facilities are detailed in Section 8. Finally, we conclude in Section 9, with a brief discussion of the use of PySP on a number of active research projects.

2 Stochastic Programming: Definition and Notations

PySP is designed to express and solve stochastic programming problems, which we now briefly introduce. More comprehensive introductions to both the theoretical foundations and the range of potential applications can be found in [Birge and Louveaux, 1997], [Shapiro et al., 2009], and [Wallace and Ziemba, 2005].

We concern ourselves with stochastic optimization problems where uncertain parameters (data) can be represented by a set of scenarios \mathcal{S} , each of which specifies both (1) a full set of random variable realizations and (2) a corresponding probability of occurrence. The random variables in question specify the evolution of uncertain parameters over time. We index the scenario set by s and refer to the probability of occurrence of s (or, more accurately, a realization “near” scenario s) as $\Pr(s)$. Let the number of scenarios be given by $|\mathcal{S}|$. The source of these scenarios does not concern us in this paper, although we observe that they are frequently obtained via simulation or formed from expert opinions. We assume that the decision process of interest consists of a sequence of discrete time stages, the set of which is denoted T . We index T by t , and denote the number of time stages by $|T|$.

Although PySP can support some types of non-linear constraints and objectives, we develop the notation primarily for the linear case in the interest of simplicity. For each scenario s and time stage t , $t \in \{1, \dots, |T|\}$, we are given a row vector $c(s, t)$ of length $n(t)$, a $m(t) \times n(t)$ matrix $A(s, t)$, and a column vector $b(s, t)$ of length $m(t)$. Let $N(t)$ be the index set $\{1, \dots, n(t)\}$ and $M(t)$ be the index set $\{1, \dots, m(t)\}$. For notational convenience, let $A(s)$ denote $(A(s, 1), \dots, A(s, |T|))$ and let $b(s)$ denote $(b(s, 1), \dots, b(s, |T|))$.

The decision variables in a stochastic program consist of a set of $n(t)$ vectors $x(t)$; one vector for each scenario $s \in \mathcal{S}$. Let $X(s)$ be $(x(s, 1), \dots, x(s, |T|))$. We will use X as shorthand for the entire solution system of x vectors, i.e., $X = x(1, 1), \dots, x(|\mathcal{S}|, |T|)$.

If we were prescient enough to know which scenario $s \in \mathcal{S}$ would be ultimately realized, our optimization objective would be to minimize

$$f_s(X(s)) \equiv \sum_{t \in T} \sum_{i \in N(t)} [c_i(s, t) x_i(s, t)] \quad (\text{P}_s)$$

subject to the constraint

$$X \in \Omega_s.$$

We use Ω_s as an abstract notation to express all constraints for scenario s , including requirements that some decision vector elements are discrete or more general requirements such as

$$A(s)X(s) \geq b(s).$$

The notation $A(s)X(s)$ is used to capture the usual sorts of single period and inter-period linking constraints that one typically finds in multi-stage mathematical programming formulations.

We must obtain solutions that do not require foreknowledge and that will be feasible independent of which scenario is ultimately realized. In particular, lacking prescience,

only solutions that are implementable are practically useful. Solutions that are not admissible, on the other hand, may have some value because while some constraints may represent laws of physics, others may be violated slightly without serious consequence.

We refer to a solution that satisfies constraints for all scenarios as *admissible*. We refer to a solution vector as *implementable* if for all pairs of scenario s and s' that are indistinguishable up to time t , $x_i(s, t') = x_i(s', t')$ for all $1 \leq t' \leq t$ and each i in each $N(t)$. We refer to the set of all implementable solutions as $\mathcal{N}_{\mathcal{S}}$ for a given set of scenarios, \mathcal{S} .

To achieve admissible and implementable solutions, the expected value minimization problem then becomes:

$$\min \sum_{s \in \mathcal{S}} [\Pr(s) f(s; X(s))] \quad (\text{P})$$

subject to

$$\begin{aligned} X &\in \Omega_s \\ X &\in \mathcal{N}_{\mathcal{S}}. \end{aligned}$$

Formulation (P) is known as a stochastic mathematical program. If all decision variables are continuous, we refer to the problem simply as a stochastic program. If some of the decision variables are discrete, we refer to the problem as a stochastic mixed-integer program.

In practice, the parameter uncertainty in stochastic programs is often encoded via a *scenario tree*, in which a node specifies the parameter values $b(s, t)$, $c(s, t)$, and $A(s, t)$ for all $t \in T$ and $s, s' \in \mathcal{S}$ such that s and s' are indistinguishable up to time t . Scenario trees are discussed in more detail in Section 6.

3 Modeling in PySP

Modern algebraic modeling languages (AMLs) such as AMPL [AMPL, 2010] and GAMS [GAMS, 2010] provide powerful mechanisms for expressing and solving deterministic mathematical programs. AMLs allow non-specialists to easily formulate and solve mathematical programming models, avoiding the need for a deep understanding of the underlying algorithmic technologies; we desire to retain the same capabilities for stochastic mathematical programming models. Many AMLs differentiate between the abstract, symbolic model and a concrete instance of the model, i.e., a model instantiated with data. The advantages of this approach are widely known [Fourer et al., 1990, p.35]. Thus, a design objective is to retain this differentiation in our approach to modeling stochastic mathematical programs in PySP. Finally, our ultimate objective is to develop and maintain software under an open-source distribution license. Consequently, PySP itself must be based on an open-source AML.

These design requirements have led us to select Pyomo [Hart et al., 2010] as the AML on which to base PySP. Pyomo is an open-source AML developed and maintained by Sandia National Laboratories (co-developed by authors of this paper), and is distributed as part of the COIN-OR initiative [COIN-OR, 2010]. Pyomo is written in the Python high-level programming language [Python, 2010a], which possesses several features that enable the development of generic solvers. AML alternatives to Pyomo

(many of which are also written in Python) do exist, but the discussion of the pros and cons are beyond the present scope; we defer to [Hart et al., 2010] for such arguments.

In this section, we discuss the use of Pyomo to formulate and express stochastic programs in PySP. As a motivating example, we consider the well-known Birge and Louveaux [Birge and Louveaux, 1997] “farmer” stochastic program. Mirroring several other approaches to modeling stochastic programs (e.g., see [Thénié et al., 2007]), we require the specification of two related components: the deterministic base model and the scenario tree. In Section 3.1 we discuss the specification of the deterministic reference model and associated data; Section 3.2 details the model and data underlying PySP scenario tree specification. The mechanisms for specifying uncertain parameter data are discussed in Section 3.3. Finally, we briefly discuss the programmatic compilation of a scenario tree specification into objects appropriate for use by solvers in Section 3.4.

3.1 The Deterministic Reference Model

The starting point for developing a stochastic programming model in PySP is the specification of an abstract *reference* model, which describes the deterministic multi-stage problem for an representative scenario. The reference model does not make use of, or describe, any information relating to parameter uncertainty or the scenario tree. Typically, it is simply the model that would be used in single-scenario analysis, i.e., the model that is commonly developed before stochastic aspects of an optimization problem are considered. PySP requires that the reference model – specified in Pyomo – is contained in a file named **ReferenceModel.py**. As an illustrative example, the complete reference model for Birge and Louveaux’s farmer problem is shown Figure 1. When looking at this figure, it is useful to remember that backslash is the python line continuation character.

For details concerning the syntax and use of the Pyomo language, we defer to [Hart et al., 2010]. Here, we simply observe that a detailed knowledge of Python is *not* necessary to develop a reference model in Pyomo; users are often unaware that a Pyomo model specifies executable Python code, or that they are using a class library. Relative to the AMPL formulation of the farmer problem, the Pyomo formulation is somewhat more verbose – primarily due to its embedding in an high-level programming language.

While the reference model is independent of any stochastic components of the problem, PySP does require that the objective cost component for each decision stage of the stochastic program be assigned to a distinct variable or variable index. In the farmer reference model, we simply label the first and second stage cost variables as *FirstStageCost* and *SecondStageCost*, respectively. The corresponding values are computed via the constraints *ComputeFirstStageCost* and *ComputeSecondStageCost*. We initially imposed the requirement concerning specification of per-stage cost variables (which are not a common feature in other stochastic programming software packages) primarily to facilitate various aspects of solution reporting. However, the convention has additionally proved very useful in implementation of various PySP solvers.

To create a concrete instance from the abstract reference model, a Pyomo data file must also be specified. The data can correspond to an arbitrary scenario, and must completely specify all parameters in the abstract reference model. The reference data file must be named **ReferenceModel.dat**. An example data file corresponding to the

```

from coopr.pyomo import *
model=Model()

# Parameters
model.CROPS=Set()
model.TOTALACREAGE=Param(within=PositiveReals)
model.PriceQuota=Param(model.CROPS, within=PositiveReals)
model.SubQuotaSellingPrice=Param(model.CROPS, within=PositiveReals)
model.SuperQuotaSellingPrice=Param(model.CROPS)
model.CattleFeedRequirement=Param(model.CROPS, \
    within=NonNegativeReals)
model.PurchasePrice=Param(model.CROPS, within=PositiveReals)
model.PlantingCostPerAcre=Param(model.CROPS, within=PositiveReals)
model.Yield=Param(model.CROPS, within=NonNegativeReals)

# Variables
model.DevotedAcreage=Var(model.CROPS, \
    bounds=(0.0, model.TOTALACREAGE))
model.QuantitySubQuotaSold=Var(model.CROPS, bounds=(0.0, None))
model.QuantitySuperQuotaSold=Var(model.CROPS, bounds=(0.0, None))
model.QuantityPurchased=Var(model.CROPS, bounds=(0.0, None))
model.FirstStageCost=Var()
model.SecondStageCost=Var()

# Constraints
def total_acreage_rule(model):
    return summation(model.DevotedAcreage) <= model.TOTALACREAGE
model.ConstrainTotalAcreage=Constraint(rule=total_acreage_rule)

def cattle_feed_rule(i, model):
    return model.CattleFeedRequirement[i] <= \
        (model.Yield[i] * model.DevotedAcreage[i]) + \
        model.QuantityPurchased[i] - \
        model.QuantitySubQuotaSold[i] - \
        model.QuantitySuperQuotaSold[i]
model.EnforceCattleFeedRequirement=Constraint(model.CROPS, \
    rule=cattle_feed_rule)

def limit_amount_sold_rule(i, model):
    return model.QuantitySubQuotaSold[i] + \
        model.QuantitySuperQuotaSold[i] <= \
        (model.Yield[i] * model.DevotedAcreage[i])
model.LimitAmountSold=Constraint(model.CROPS, \
    rule=limit_amount_sold_rule)

def enforce_quotas_rule(i, model):
    return (0.0, model.QuantitySubQuotaSold[i], model.PriceQuota[i])
model.EnforceQuotas=Constraint(model.CROPS, \
    rule=enforce_quotas_rule)

# Stage-specific cost computations
def first_stage_cost_rule(model):
    return model.FirstStageCost == \
        summation(model.PlantingCostPerAcre, model.DevotedAcreage)
model.ComputeFirstStageCost=Constraint(rule=first_stage_cost_rule)

def second_stage_cost_rule(model):
    expr=summation(model.PurchasePrice, model.QuantityPurchased)
    expr -= summation(model.SubQuotaSellingPrice, \
        model.QuantitySubQuotaSold)
    expr -= summation(model.SuperQuotaSellingPrice, \
        model.QuantitySuperQuotaSold)
    return (model.SecondStageCost - expr) == 0.0
model.ComputeSecondStageCost=Constraint(rule=second_stage_cost_rule)

# Objective
def total_cost_rule(model):
    return (model.FirstStageCost + model.SecondStageCost)
model.Total_Cost_Objective=Objective(rule=total_cost_rule, \
    sense=minimize)

```

Fig. 1 ReferenceModel.py: The deterministic reference Pyomo model for Birge and Louveaux's farmer problem.

```

set CROPS := WHEAT CORN SUGAR_BEETS ;

param TOTALACREAGE := 500 ;

param PriceQuota := WHEAT 100000 CORN 100000 SUGAR_BEETS 6000 ;

param SubQuotaSellingPrice := WHEAT 170 CORN 150 SUGAR_BEETS 36 ;

param SuperQuotaSellingPrice := WHEAT 0 CORN 0 SUGAR_BEETS 10 ;

param CattleFeedRequirement := WHEAT 200 CORN 240 SUGAR_BEETS 0 ;

param PurchasePrice := WHEAT 238 CORN 210 SUGAR_BEETS 100000 ;

param PlantingCostPerAcre := WHEAT 150 CORN 230 SUGAR_BEETS 260 ;

param Yield := WHEAT 3.0 CORN 3.6 SUGAR_BEETS 24 ;

```

Fig. 2 ReferenceModel.dat: The Pyomo reference model data for Birge and Louveaux’s farmer problem.

farmer reference model is shown in Figure 2. Although Pyomo supports various data file formats, the example illustrates the use of a file that uses Pyomo data commands – the most commonly used data file format in Pyomo. Pyomo data commands include data commands for set and parameter data that are consistent with AMPL’s data commands. Our adoption of this convention minimizes the effort required to translate deterministic reference models expressed in AMPL into Pyomo.

3.2 The Scenario Tree

Given a deterministic reference model, the second step in developing a stochastic program in PySP involves specification of the scenario tree structure and associated parameter data. A PySP scenario tree specification supplies all information concerning the time stages, the mapping of decision variables to time stages, how various scenarios are temporally related to one another (i.e., scenario tree nodes and their inter-relationships), and the probabilities of various scenarios. As discussed below, the scenario tree does not directly specify uncertain parameter values; rather, it specifies references to data files containing such data.

As with the abstract reference model, the abstract scenario tree model in PySP is expressed in Pyomo. However, the contents of the scenario tree model – called **ScenarioStructure.py** and shown in Figure 3 for reference – are fixed. The model is built into and distributed with PySP; the user does not edit this file. Instead, the user must supply values for each of the parameters specified in the scenario tree model. Finally, we observe that the scenario tree model is simply a data collection that is specified using Pyomo parameter and set objects, i.e., a very restricted form of a Pyomo model (lacking variables, constraints, and an objective).

The precise semantics for each of the parameters (or sets) indicated in Figure 3 are as follows:

- **Stages:** An ordered set containing the names (specified as arbitrary strings) of the time stages. The order corresponds to the time order of the stages.


```

from coopr.pyomo import *
scenario_tree_model=Model()

scenario_tree_model.Stages=Set(ordered=True)
scenario_tree_model.Nodes=Set()

scenario_tree_model.NodeStage=Param(scenario_tree_model.Nodes, \
                                     within=scenario_tree_model.Stages)
scenario_tree_model.Children=Set(scenario_tree_model.Nodes, \
                                 within=scenario_tree_model.Nodes, \
                                 ordered=True)
scenario_tree_model.ConditionalProbability= \
    Param(scenario_tree_model.Nodes)

scenario_tree_model.Scenarios=Set(ordered=True)
scenario_tree_model.ScenarioLeafNode= \
    Param(scenario_tree_model.Scenarios, \
          within=scenario_tree_model.Nodes)

scenario_tree_model.StageVariables=Set(scenario_tree_model.Stages)
scenario_tree_model.StageCostVariable= \
    Param(scenario_tree_model.Stages)

scenario_tree_model.ScenarioBasedData=Param(within=Boolean, \
                                             default=True)

```

Fig. 3 ScenarioStructure.py: The PySP model for specifying the structure of the scenario tree.

- **Nodes:** A set of the names (specified as arbitrary strings) of the nodes in the scenario tree.
- **NodeStage:** An indexed parameter mapping node names to stage names. Each node in the scenario tree must be assigned to a specific stage.
- **Children:** An indexed set mapping node names to sets of node names. For each non-leaf node in the scenario tree, a set of child nodes must be specified. This set implicitly defines the overall branching structure of the scenario tree. Using this set, the parent nodes are computed internally to PySP. There can only be one node in the scenario tree with no parents, i.e., the tree must be singly rooted.
- **ConditionalProbability:** An indexed, real-valued parameter mapping node names to their conditional probability, relative to their parent node. The conditional probability of the root node must be equal to 1, and for any node with children, the conditional probabilities of the children must sum to 1. Numeric values must be contained within the interval $[0, 1]$.
- **Scenarios:** An ordered set containing the names (specified as arbitrary strings) of the scenarios. These names are used for two purposes: reporting and data specification (see Section 3.3). The ordering is provided as a convenience for the user, to organize reporting output.
- **ScenarioLeafNode:** An indexed parameter mapping scenario names to their leaf node name. Facilitates linkage of the scenario to their composite nodes in the scenario tree.

```

set Stages := FirstStage SecondStage ;

set Nodes := RootNode
             BelowAverageNode
             AverageNode
             AboveAverageNode ;

param NodeStage := RootNode      FirstStage
                   BelowAverageNode SecondStage
                   AverageNode    SecondStage
                   AboveAverageNode SecondStage ;

set Children[RootNode] := BelowAverageNode
                          AverageNode
                          AboveAverageNode ;

param ConditionalProbability := RootNode      1.0
                               BelowAverageNode 0.33333333
                               AverageNode    0.33333334
                               AboveAverageNode 0.33333333 ;

set Scenarios := BelowAverageScenario
                  AverageScenario
                  AboveAverageScenario ;

param ScenarioLeafNode := BelowAverageScenario BelowAverageNode
                          AverageScenario      AverageNode
                          AboveAverageScenario AboveAverageNode ;

set StageVariables[FirstStage] := DevotedAcreage[*] ;
set StageVariables[SecondStage] := QuantitySubQuotaSold[*]
                                   QuantitySuperQuotaSold[*]
                                   QuantityPurchased[*] ;

param StageCostVariable := FirstStage FirstStageCost
                           SecondStage SecondStageCost ;

```

Fig. 4 The PySP **ScenarioStructure.dat** data file for specifying the scenario tree for the farmer problem.

- **StageVariables:** An indexed set mapping stage names to sets of variable names in the reference model. The sets of variables indicate those variables that are associated with the given stage. Implicitly defines the non-anticipativity constraints that should be imposed when generating and/or solving the PySP model.
- **ScenarioBasedData:** A Boolean parameter specifying how the instances for each scenario are to be constructed. The semantics for this parameter are detailed below in Section 3.3.

Data to instantiate these parameters and sets must be provided by a user in a file named **ScenarioStructure.dat**. The scenario tree structure specification for the farmer problem is shown in Figure 4, specified using the AMPL data file format.

We observe that PySP provides a simple “slicing” syntax to specify subsets of indexed variables. In particular, the “*” character is used to match all values in a particular dimension of an indexed parameter. In more complex examples, variables are typically indexed by time stage. In these cases, the slice syntax allows for very concise

specification of the stage-to-variable mapping. Finally, we observe that PySP makes no assumptions regarding the linkage between time stages and variable index structure. In particular, the time stage need not explicitly be referenced within a variable's index set. While this is often the case in multi-stage formulations, the convention is not universal, e.g., as in the case of the farmer problem.

3.3 Scenario Parameter Specification

Data files specifying the deterministic and stochastic parameters for each of the scenarios in a PySP model can be specified in one of two ways. The simplest approach is “scenario-based”, in which a single data file containing a complete parameter specification is provided for each scenario. In this case, the file naming convention is as follows: If the scenario is named *ScenarioX*, then the corresponding data file for the scenario must be named **ScenarioX.dat**. This approach is often expedient – especially if the scenario data are generated via simulation, as is often the case in practice. However, there is necessarily redundancy in the encoding. Depending on the problem size and number of scenarios, this redundancy may become excessive in terms of disk storage and access. Scenario-based data specification is the default behavior in PySP, as indicated by the default value of the `ScenarioBasedData` parameter in Figure 3. We note that the listing in Figure 2 is an example of a scenario-based data specification.

Node-based parameter specification is provided as an alternative to the default scenario-based approach, principally to eliminate storage redundancy. With node-based specification, parameter data specific to each *node* in the scenario tree is specified in a distinct data file. The naming convention is as follows: If the node is named *NodeX*, then the corresponding data file for the node must be named **NodeX.dat**. To create a scenario instance, data for all nodes associated with a scenario are accessed (via the `ScenarioLeafNode` parameter in the scenario tree specification and the computed parent node linkages). Node-based parameter encoding eliminates redundancy, although typically at the expense of a slightly more complex instance generation process. To enable node-based scenario initialization, a user needs to simply add the following line to **ScenarioStructure.dat**:

```
param ScenarioBasedData := False ;
```

In the case of the farmer problem, all parameters except for *Yield* are identical across scenarios. Consequently, these parameters can be placed in a file named **RootNode.dat**. Then, files containing scenario-specific *Yield* parameter values are specified for each second-stage leaf node (in files named **AboveAverageNode.dat**, **AverageNode.dat**, and **BelowAverageNode.dat**).

3.4 Compilation of the Scenario Tree Model

The PySP scenario tree structure model is a declarative entity, merely specifying the data associated with a scenario tree. PySP internally uses the information contained in this model to construct a `ScenarioTree` object, which in turn is composed of `ScenarioTreeNode`, `Stage`, and `Scenario` objects. In aggregate, these Python objects allow

programmatic navigation, query, manipulation, and reporting of the scenario tree structure. While hidden from the typical user, these objects are crucial in the processes of generating the extensive form (Section 4) and generic solvers (Section 6).

4 Generating and Solving the Extensive Form

Given a stochastic program encoded in accordance with the PySP conventions described in Section 3, the next immediate issue of concern is its solution. The most straightforward method to solve a stochastic program involves generating the *extensive form* (also known as the *deterministic equivalent*) and then invoking a standard deterministic (mixed-integer) programming solver, e.g., CPLEX. The extensive form given as problem (P) in Section 2 completely specifies all scenarios and the coupling non-anticipativity constraints at each node in the scenario tree.

In many cases, particularly when small numbers of scenarios are involved or the decision variables are all continuous, the extensive form can be effectively solved with off-the-shelf solvers [Parija et al., 2004]. Further, although decomposition techniques may ultimately be needed for large, more complex models, the extensive form is usually the first attempted method to solve a stochastic program.

In this section, we describe the use and design of facilities in PySP for generating and solving the extensive form. Section 4.1 describes a user script for generating and solving the extensive form; an overview of the implementation of this script is then provided in Section 4.2.

4.1 The **runef** Script

PySP provides an easy-to-use script – **runef** – to both generate and solve the extensive form of a stochastic program. We now briefly describe the primary command-line options for this script; note that all options begin with a double dash prefix:

```
--help
Display all command-line options, with brief descriptions, and exits.

--verbose
Display verbose output to the standard output stream, above and beyond the usual
status output. Disabled by default.

--model-directory=MODEL_DIRECTORY
Specifies the directory in which the reference model (ReferenceModel.py) is
stored. Defaults to “.”, the current working directory.

--instance-directory=INSTANCE_DIRECTORY
Specifies the directory in which all reference model and scenario model data files
are stored. Defaults to “.”, the current working directory.

--output-file=OUTPUT_FILE
Specifies the name of the LP format output file to which the extensive form is
written. Defaults to “efout.lp”.

--solve
Directs the script to solve the extensive form after writing it. Disabled by default.

--solver=SOLVER_TYPE
Specifies the type of solver for solving the extensive form, if a solve is requested.
Defaults to “cplex”.
```

`--solver-options=SOLVER_OPTIONS`

Specifies solver options in keyword-value pair format, if a solve is requested.

`--output-solver-log`

Specifies that the output of the solver is to be echoed to the standard output stream. Disabled by default. Useful to ascertain status for extensive forms with long solve times.

For example, to write and solve the farmer problem (provided with the Coop installation, in the directory `coopr/examples/pysp/farmer`), the user simply executes:

```
runef --model-directory=models \\
      --instance-directory=scenariodata \\
      --solve
```

The double forward-slash characters are simply continuation characters, used (here and elsewhere in this article) to restrict the width of the example inputs and outputs.

Following solver execution, the resulting solution is loaded and displayed. The solution output is split into two distinct components: variable values and stage/scenario costs. For the farmer example, the per-node variable values are given as:

Tree Nodes:

```
Name=RootNode
Stage=FirstStage
Variables:
    DevotedAcreage [CORN]=80.0
    DevotedAcreage [SUGAR_BEETS]=250.0
    DevotedAcreage [WHEAT]=170.0

Name=AboveAverageNode
Stage=SecondStage
Variables:
    QuantitySubQuotaSold [CORN]=48.0
    QuantitySubQuotaSold [SUGAR_BEETS]=6000.0
    QuantitySubQuotaSold [WHEAT]=310.0

Name=AverageNode
Stage=SecondStage
Variables:
    QuantitySubQuotaSold [SUGAR_BEETS]=5000.0
    QuantitySubQuotaSold [WHEAT]=225.0

Name=BelowAverageNode
Stage=SecondStage
Variables:
    QuantitySubQuotaSold [SUGAR_BEETS]=4000.0
    QuantitySubQuotaSold [WHEAT]=140.0
    QuantityPurchased [CORN]=48.0
```

Similarly, the per-node stage cost and per-scenario overall costs are given as follows:

Tree Nodes :

```

Name=RootNode
Stage=FirstStage
Expected node cost=-108390.0000

Name=AboveAverageNode
Stage=SecondStage
Expected node cost=-275900.0000

Name=AverageNode
Stage=SecondStage
Expected node cost=-218250.0000

Name=BelowAverageNode
Stage=SecondStage
Expected node cost=-157720.0000

```

Scenarios :

```

Name=AboveAverageScenario
Stage=FirstStage      Cost=108900.0000
Stage=SecondStage     Cost=-275900.0000
Total scenario cost=-167000.0000

Name=AverageScenario
Stage=FirstStage      Cost=108900.0000
Stage=SecondStage     Cost=-218250.0000
Total scenario cost=-109350.0000

Name=BelowAverageScenario
Stage=FirstStage      Cost=108900.0000
Stage=SecondStage     Cost=-157720.0000
Total scenario cost=-48820.0000

```

Currently, the extensive form is output for solution in the CPLEX LP file format. In practice, this has not been a limitation, as nearly all commercial and open-source solvers support this format.

Various other command-line options are available in the **runef** script, including those related to performance profiling and Python garbage collection. Further, the **runef** script is capable of writing and solving the extensive form augmented with a weighted Conditional Value-at-Risk term in the objective [Schultz and Tiedemann, 2005].

We conclude by noting that the **runef** script, as with the decomposition-based solver script described in Section 6, relies on significant functionality from the Coopr Python optimization library in which PySP is embedded. This includes a wide range of

solver interfaces (both commercial and open-source), problem writers, solution readers, and distributed solvers. For further details, we refer to [Hart et al., 2010].

4.2 Under the Hood: Generating the Extensive Form

We now provide an overview of the implementation of the **runef** script. In doing so, our objectives are to (1) illustrate the use of Python to create generic writers and solvers and (2) to provide some indication of the programmatic-level functionality available in PySP.

The high-level process executed by the **runef** script to generate the extensive form in PySP is as follows:

1. Load the scenario tree data; create the corresponding instance.
2. Create the ScenarioTree object from the scenario tree Pyomo model.
3. Load the reference model; create the corresponding instance from reference data.
4. Load the scenario instance data; create the corresponding instances.
5. Create the master “binding” instance; instantiate per-node variable objects.
6. Add master-to-scenario instance equality constraints to enforce non-anticipativity.

Steps 1 and 2 simply involve the process of creating the Pyomo instance specifying all data related to the scenario tree structure, and creating the corresponding ScenarioTree object to facilitate programmatic access of scenario tree attributes. In Step 3, the core deterministic abstract model is loaded. The abstract model is then used in Step 4, in conjunction with the ScenarioTree object, to create a concrete model instance for each scenario in the stochastic program. The scenario instances are at this point – and remain so – completely independent of one another. This approach differs from that of some of the software packages described in Section 5, in which variables are instantiated for each node of the scenario tree and shared across the relevant scenarios. While our approach does introduce redundancy, the replication introduces only moderate memory overhead and confers significant practical advantages when implementing generic decomposition-based solvers, e.g., as illustrated below in Section 6. In particular, we note that scenario-based decomposition solvers gradually and incrementally enforce non-anticipativity, such that replicated variables are required.

Next, a master “binding” instance is created in Steps 5 and 6. The purpose of the master binding instance is to enforce the required non-anticipativity constraints at each node in the scenario tree. Using the ScenarioTree object, the tree is traversed and the collection of variable names (including indices, if necessary) associated with each node is identified – initially specified via the *StageVariables* attribute of the scenario tree model. The corresponding variable objects are then identified in the reference model instance, cloned, and attached to the binding instance. This step critically requires the Python capability of *introspection*: the ability, at run-time, to gather information about objects and manipulate them.

To illustrate how introspection is used to develop generic algorithms, consider again the farmer example from Section 3. By specifying the line

```
set StageVariables[FirstStage] := DevotedAcreage[*] ;
```

the user is communicating the requirement to impose non-anticipativity constraints on (all indices of) the first stage variable *DevotedAcreage*. The variable is specified simply as a string, which can be programmatically split into the corresponding root name and index template. Using the root name, Python can query (via the `getattr` built-in function) the reference model for the corresponding variable, validate that it exists, and if so, return the corresponding variable object. This loose coupling between the user data and algorithm code is facilitated by this simple, yet powerful, introspection mechanism.

The final primary step in the **runef** script involves construction of constraints to enforce non-anticipativity between the newly created variables in the master binding instance and the variables in the relevant scenario instances. This process again relies on introspection to achieve a generic implementation.

Overall, the core functionality of the **runef** script is expressed in approximately 700 lines of Python code – including all white-space and comments. This includes both the code for creating the relevant Pyomo instances, generating the master binding instance via the processed described above, and controlling the output of the LP file.

Finally, we observe that despite our explicit approach to writing the extensive form through the introduction of master variables and non-anticipativity constraints, we find that the impact on run-time is at worst negligible. The presolvers in commercial packages such as CPLEX, Gurobi, or XpressMP (and those available with some open-source solvers) are able to quickly identify and eliminate most of the redundant variables and constraints.

5 Related Proposals and Software Packages

We now briefly survey prior and on-going efforts to develop software packages supporting the specification and solution of stochastic programs, with the objective of placing the capabilities of PySP in this broader context. Numerous extensions to existing AMLs to support the specification of stochastic programs have been proposed in the literature; Gassmann and Ireland [1996] is an early example. Similarly, various solver interfaces have been proposed, with the dominant mechanism being the direct solution of the extensive form. Here, we primarily focus on specification and solver efforts associated with open-source and academic initiatives, which generally share the same distribution goals, user community targets, and design objectives (e.g., experimental, generic, and configurable solvers) as PySP.

StAMPL Fourer and Lopes [2009] describe an extension to AMPL, called StAMPL, whose goal is to simplify the modeling process associated with stochastic program specification. One key objective of StAMPL is to explicitly avoid the use of scenario and stage indices when specifying the core algebraic model, separating the specification of the stochastic process from the underlying deterministic optimization model. The authors describe a preprocessor that translates a StAMPL problem description into the fully indexed AMPL model, which in turn is written in SMPS format for solution. In contrast to StAMPL, PySP provides a straightforward mechanism to specify a stochastic program, and does not strive to advance the state-of-the-art in modeling. Rather, our primary focus is on developing generic and configurable solvers, and discussed in Sections 4, 6, and 7.

STRUMS STRUMS is a system for performing and managing decomposition and relaxation strategies in stochastic programming [Fourer and Lopes, 2006]. Input problems are specified in the SMPS format, and the package provides mechanisms for writing the extensive form, performing basic and nested Benders decomposition (i.e., the L-shaped method), and implementing Lagrangian relaxation; only stochastic linear programs are considered. The design objective of STRUM – to provide mechanisms facilitating automatic problem decomposition – is consistent with the design of PySP. However, PySP currently provides mechanisms for scenario-based decomposition, in contrast to stage-oriented decomposition. This emphasis is due primarily to our interest in mixed-integer stochastic programming. In contrast to STRUMS, PySP is integrated with an AML.

DET2STO Thénier et al. [2007] describe an extension of AMPL to support the specification of stochastic programs, noting that (at the time the effort was initiated) no AMLs were available with stochastic programming support. In particular, they provide a script – called DET2STO, available from <http://apps.ordecys.com/det2sto> – taking an augmented AMPL model as input and generating the extensive form via an SMPS output file. The research focus is on the automated generation of the extensive form, with the authors noting: “We recall here that, while it is relatively easy to describe the two base components - the underlying deterministic model and the stochastic process - it is tedious to define the contingent variables and constraints and build the deterministic equivalent” [Thénier et al., 2007, p.35]. While subtle modeling differences do exist between DET2STO and PySP (e.g., in the way scenario-based and transition-based representations are processed), they provide identical functionality in terms of ability to model stochastic programs and generate the extensive form.

SMI Part of COIN-OR, the Stochastic Modeling Interface (SMI) [SMI, 2010] provides a set of C++ classes to (1) either to programmatically create a stochastic program or to load a stochastic program specified in SMPS, and (2) to write the extensive form of the resulting program. SMI provides no solvers, instead focusing on generation of the extensive form for solution by external solvers. Connections to FLOPC++ [FLOPCPP, 2010] do exist, providing a mechanism for problem description via an AML. While providing a subset of PySP functionality, the need to express models in a compiled, technically sophisticated programming language (C++) is a significant drawback for many users.

APLEpy Karabuk [2008] describes the design of classes and methods to implement stochastic programming extensions to his Python-based APLEpy [Karabuk, 2005] environment for mathematical programming, with a specific emphasis on stochastic linear programs. Karabuk’s primary focus is on supporting relaxation-based decompositions in general, and the L-shaped method in particular, although his design would create elements that could be used to construct other algorithms as well. The vision expressed in [Karabuk, 2008] is one where the boundary between model and algorithm must be crossed so that the algorithm can be expressed in terms of model elements. This approach is also possible using Pyomo and PySP, but it is not the underlying philosophy of PySP. Rather, we are interested in enabling the separation of model, data, and algorithm except when the users wish to create model specific algorithm enhancements.

SPInE SPInE [Mitra et al., 2005] provides an integrated modeling and solver environment for stochastic programming. Models are specified in an extension to AMPL called SAMPL (other base modeling languages are provided), which can in turn be solved via a number of built-in solvers. In contrast to PySP, the solvers are not specifically designed to be customizable, and are generally limited to specific problem classes. For example, multi-stage stochastic linear programs are solved via nested Benders decomposition, while Lagrangian relaxation is the only option for two-stage mixed-integer stochastic programs. SPInE is primarily focused on providing an out-of-the-box solution for stochastic linear programs, which is consistent with the lack of emphasis on customizable solution strategies.

SLP-IOR Similar to SPInE, SLP-IOR [Kall and Mayer, 2005b] is an integrated modeling and solver environment for stochastic programming, with a strong emphasis on the linear case. In contrast to SPInE, SLP-IOR is based on the GAMS AML, and provides a broader range of solvers. However, as with SPInE, the focus is not on easily customizable solvers (most of the solver codes are written in FORTRAN). Further, the solvers for the integer case is largely ignored.

6 Progressive Hedging: A Generic Decomposition Strategy

We now transition from modeling stochastic programs and solving them via the extensive form to decomposition-based strategies, which are in practice typically required to efficiently solve large-scale instances with large numbers of scenarios, discrete variables, or decision stages. There are two broad classes of decomposition-based strategies: horizontal and vertical. *Vertical* strategies decompose a stochastic program by time stages; Van Slyke and Wets’ L-shaped method is the primary method in this class [Slyke and Wets, 1969]. In contrast, *horizontal* strategies decompose a stochastic program by scenario; Rockafellar and Wets’ Progressive Hedging algorithm [Rockafellar and Wets, 1991] and Caroe and Schultz’s Dual Decomposition (DD) algorithm [Caroe and Schultz, 1999] are the two notable methods in this class.

Currently, there is not a large body of literature to provide an understanding of practical, computational aspects of stochastic programming solvers, particularly in the mixed integer case. For any given problem class, there are few heuristics to guide selection of the algorithm likely to be most effective. Similarly, while stochastic programming solvers are typically parameterized and/or configurable, there is little guidance available regarding how to select particular parameter values or configurations for a specific problem. Lacking such knowledge, the interface to solver libraries must provide facilities to allow for easily selecting parameters and configurations.

Beyond the need for highly configurable solvers, solvers should also be generic, i.e., independent of any particular AML description. Decomposition strategies are non-trivial to implement, requiring significant development time – especially when more advanced features are considered. The lack of generic decomposition solvers is a known impediment to the broader adoption of stochastic programming. Thénier et al. [2007] concisely summarize the challenge as follows: “Devising efficient solution methods is still an open field. It is thus important to give the user the opportunity to experiment with solution methods of his choice.” By introducing both customizable and generic solvers, our goal is to promote the broader use of and experimentation with stochastic programming by significantly reducing the barrier to entry.

In this section, we discuss the interface to and implementation of a generic implementation of Progressive Hedging. Our selection of this particular decomposition algorithm is based largely on our successful experience with PH in solving difficult, multi-stage mixed-integer stochastic programs. In Section 6.1 we introduce the Progressive Hedging algorithm, and discuss its use in both linear and mixed-integer stochastic programming contexts. The interface to the PySP script for executing PH given an arbitrary PySP model is described in Section 6.2. Finally, we present an overview of the generic implementation in Section 6.3.

6.1 The Progressive Hedging Algorithm

Progressive Hedging (PH) was initially introduced as a decomposition strategy for solving large-scale stochastic linear programs [Rockafellar and Wets, 1991]. PH is a horizontal or scenario-based decomposition technique, and possesses theoretical convergence properties when all decision variables are continuous. In particular, the algorithm converges in linear time given a convex reference scenario optimization model.

Despite its introduction in the context of stochastic linear programs, PH has proved to be a very effective heuristic for solving stochastic mixed-integer programs. PH is particularly effective in this context when there exist computationally efficient techniques for solving the deterministic single-scenario optimization problems. A key advantage of PH in the mixed-integer case is the absence of requirements concerning the number of stages or the type of variables allowed in each stage – as is common for many proposed stochastic mixed-integer algorithms. A disadvantage is the current lack of provable convergence and optimality results. However, PH has been used as an effective heuristic for a broad range of stochastic mixed-integer programs [Fan and Liu, 2010, Listes and Dekker, 2005, Løkketangen and Woodruff, 1996, Cranic et al., 2009, Hvattum and Løkketangen, 2009]. For large, real-world stochastic mixed-integer programs, the determination of optimal solutions is generally not computationally tractable.

The basic idea of PH for the linear case is as follows:

1. For each scenario s , solutions are obtained for the problem of minimizing, subject to the problem constraints, the deterministic f_s (Formulation P_s).
2. The variable values for an implementable – but likely not admissible – solution are obtained by averaging over all scenarios at a scenario tree node.
3. For each scenario s , solutions are obtained for the problem of minimizing, subject to the problem constraints, the deterministic f_s (Formulation P_s) plus terms that penalize the lack of implementability using a sub-gradient estimator for the non-anticipativity constraints and a squared proximal term.
4. If the solutions have not converged sufficiently and the allocated compute time is not exceeded, goto Step 2.
5. Post-process, if needed, to produce a fully admissible and implementable solution.

To begin the PH implementation for solving formulation (P), we first organize the scenarios and decision times into a tree. The leaves correspond to scenario realizations, such that each leaf is connected to exactly one node at time $t \in \mathcal{T}$ and each of these nodes represents a unique realization up to time t . The leaf nodes are connected to nodes at time $t-1$, such that each scenario associated with a node at time $t-1$ has the same realization up to time $t-1$. This process is iterated back to time 1 (i.e., “now”). Two scenarios whose leaves are both connected to the same node at time t have the

same realization up to time t . Consequently, in order for a solution to be implementable it must be true that if two scenarios are connected to the same node at some time t , then the values of $x_i(t')$ must be the same under both scenarios for all i and for $t' \leq t$.

Progressive Hedging is a technique to iteratively and gradually enforce implementability, while maintaining admissibility at each step in the process. For each scenario s , approximate solutions are obtained for the problem of minimizing, subject to the constraints, the deterministic f_s plus terms that penalize the lack of implementability. These terms strongly resemble those found when the method of augmented Lagrangians is used [Bertsekas, 1996]. The method makes use of a system of row vectors, w , that have the same dimension as the column vector system X , so we use the same shorthand notation. For example, $w(s)$ denotes $(w(s, 1), \dots, w(s, |\mathcal{T}|))$ in the multiplier system.

To provide an formal algorithm statement of PH, we first formalize some of the scenario tree concepts. We use $\Pr(\mathcal{A})$ to denote the sum of $\Pr(s)$ over all s for scenarios emanating from node \mathcal{A} (i.e., those s that are the leaves of the sub-tree having \mathcal{A} as a root, also referred to as $s \in \mathcal{A}$). We use $t(\mathcal{A})$ to indicate the time index for node \mathcal{A} (i.e., node \mathcal{A} corresponds to time t). We use $X(t; \mathcal{A})$ on the left hand side of a statement to indicate assignment to the vector $(x_1(s, t), \dots, x_{N(t)}(s, |\mathcal{T}|))$ for each $s \in \mathcal{A}$. We refer to vectors at each iteration of PH using a superscript; e.g., $w^{(0)}(s)$ is the multiplier vector for scenario s at PH iteration zero. The PH iteration counter is k .

If we briefly defer the discussion of termination criteria, a formal version of the algorithm (with step numbering that matches in the informal statement just given) can be stated as follows, taking $\rho > 0$ as a parameter.

1. $k \leftarrow 0$
2. For all scenario indices, $s \in \mathcal{S}$:

$$X^{(0)}(s) \leftarrow \operatorname{argmin} f_s(X(s)) : X(s) \in \Omega_s \quad (1)$$

and

$$w^{(0)}(s) \leftarrow 0$$

3. $k \leftarrow k + 1$
4. For each node, \mathcal{A} , in the scenario tree, and for all $t = t(\mathcal{A})$:

$$\bar{X}^{(k-1)}(t; \mathcal{A}) \leftarrow \sum_{s \in \mathcal{A}} \Pr(s) X(t; s)^{(k-1)} / \Pr(\mathcal{A})$$

5. For all scenario indices, $s \in \mathcal{S}$:

$$w^{(k)}(s) \leftarrow w^{(k-1)}(s) + (\rho) \left(X^{(k-1)}(s) - \bar{X}^{(k-1)} \right)$$

and

$$X^k(s) \leftarrow \operatorname{argmin} f_s(X(s)) + w^{(k)}(s)X(s) + \rho/2 \left\| X(s) - \bar{X}^{k-1} \right\|^2 : X(s) \in \Omega_s. \quad (2)$$

6. If the termination criteria are not met (e.g., solution discrepancies quantified via a metric $g^{(k)}$), then goto Step 3.

The termination criteria are based mainly on convergence, but we must also allow for the use of time-based termination because non-convergence is a possibility. Iterations are continued until k reaches some pre-determined limit or the algorithm has *converged* – which we take to indicate that the set of scenario solutions s is sufficiently homogeneous. One possible definition is to require the inter-solution distance (e.g., Euclidean) to be less than some parameter.

The value of the perturbation vector ρ strongly influences the actual convergence rate of PH: if ρ is small, the penalty coefficients will vary little between consecutive iterations. To achieve tractable PH run-times, significant tuning and problem-dependent strategies for computing ρ are often required; mechanisms to support such tuning are described in Section 6.2.

6.2 The **runph** Script

Analogous to the **runef** script for generating and solving the extensive form, PySP provides a script – **runph** – to solve and post-process stochastic programs via PH. We now briefly describe the general usage of this script, followed by a discussion of some generally effective options to customize the execution of PH. As is the case with the **runef** script, all options begin with a double dash prefix. A number of key options are shared with the **runef** script: `--verbose`, `--model-directory`, `--instance-directory`, and `--solver`. In particular, the `--model-directory` and `--instance-directory` options are used to specify the PySP problem instance, while the `--solver` option is used to specify the solver applied to individual scenario sub-problems. The most general PH-specific options are:

`--max-iterations=MAX_ITERATIONS`

The maximal number of PH iterations. Defaults to 100.

`--default-rho=DEFAULT_RHO`

The default (global) ρ scalar parameter value for all variables with the exception of those appearing in the final stage. Defaults to 1.

`--termdiff-threshold=TERMDIFF_THRESHOLD`

The convergence threshold used to terminate PH (Step 6 of the pseudocode). Convergence is by default quantified via the following formula:

$$g^k = \sum_{s \in \mathcal{S}} \Pr(s) \|X^{(k)}(t; s) - \bar{X}^{(k)}(\mathcal{A})\|.$$

Defaults to 0.01. This quantity is known as the *termdiff*.

In general, the default values for the maximum allowable iteration count, ρ , and convergence threshold are likely to yield slow convergence of PH; for any real application, experimentation and analysis should be applied to obtain a more computationally effective configuration.

To illustrate the execution **runph** on a stochastic linear program, we again consider Birge and Louveaux’s farmer problem. To solve the farmer problem with PySP, a user simply executes the following:

```
runph --model-directory=models --instance-directory=scenariodata
```

which will result in eventual convergence to an optimal, admissible, and implementable solution – subject to the numerical tolerance issues. For the sake of brevity, we do not illustrate the output here; the final solution is reported in a format identical to that

illustrated in Section 4. The quantity of information generated by PH can be significant, e.g., including the penalty weights and solutions for each scenario problem $s \in \mathcal{S}$ at each iteration. However, this information is not generated by default. Rather, simple summary information, including the value of $g^{(k)}$ at each PH iteration k , is output. As is theoretically guaranteed in the case of stochastic linear programs, **runph** does converge given a linear PySP input model. The exact number of iterations depends in part on the precise solver used; on our test platform, for example, convergence is achieved in 48 iterations using CPLEX 11.2.1. It should be noted that for many stochastic linear – and even small, mixed-integer – programs (including the farmer example), any implementation of PH may solve significantly *slower* than the extensive form. This behavior is primarily due to the overhead associated with communicating with solvers for each scenario, for each PH iteration. However, this overhead is not significant with larger and/or more difficult scenario problems.

Having described the basic **runph** functionality, we now transition to a discussion of some issues with PH that can arise in practice, and their resolution via the **runph** script. More comprehensive configuration methods are discussed in Section 7, to address more complex PH issues.

Setting Variable-Specific ρ : In many applications, no single value of ρ for all variables yields a computationally efficient PH configuration. Consider the situation in which the objective is to minimize expected investment costs in a spare parts supply chain, e.g., for maintaining an aircraft fleet. The acquisition cost for spare parts is highly variable, ranging from very expensive (engines) to very cheap (gaskets). If ρ values are too small, e.g., on the order of the price of a tire, PH will require large iteration counts to achieve changes – let alone convergence – in the decision variables associated with engine procurement counts. If ρ values are too high, e.g., on the order of the price of an engine, then the PH weights w associated with gasket procurement counts will converge too quickly, yielding sub-optimal variable values. Alternatively, PH sub-problem solves may “over-shoot” the optimal variable value, resulting in oscillation. Various strategies for computing variable-specific ρ are discussed in [Watson and Woodruff, 2010].

To support the implementation of variable-specific ρ strategies in PySP, we define the following command-line option to **runph**:

`--rho-cfgfile=RHO.CFGFILE`

The name of a configuration script to compute PH rho values. Default is None.

The rho configuration file is a piece of executable Python code that computes the desired ρ . This allows for the expression of arbitrarily complex formulas and procedures. An example of such a configuration file, used in conjunction with the PySP SIZES example [Jorjani et al., 1999], is as follows:

```
model_instance = self._model_instance # syntatic sugar

for i in model_instance.ProductSizes:
    self.setRhoAllScenarios(model_instance.ProduceSizeFirstStage[i], \
                           model_instance.SetupCosts[i] * 0.001)
    self.setRhoAllScenarios(model_instance.NumProducedFirstStage[i], \
                           model_instance.UnitProductionCosts[i] * 0.001)
for j in model_instance.ProductSizes:
```

```

if j <= i:
    self.setRhoAllScenarios(model_instance.NumUnitsCutFirstStage[i,j], \
                           model_instance.UnitReductionCost * 0.001)

```

The *self* object in the script refers to the PH object itself, which in turn possesses an attribute *_model_instance*. The *_model_instance* attribute represents the deterministic reference model instance, from which the full set of problem variables can be accessed. The example script implements a simple cost-proportional ρ strategy, in which ρ is specified as a function of a variable's objective function cost coefficient. Once the appropriate ρ value is computed, the script invokes the *setRhoAllScenarios* method of the PH object, which distributes the computed ρ value to the corresponding parameter of each of the scenario problem instances. It is also possible to set the ρ values on a per-variable, per-scenario basis; however, there are currently no reported strategies that effectively use this mechanism.

The customization strategy underlying the PySP variable-specific ρ mechanism is a limited form of callback, in which the core PH code temporarily hands control back to a user script to set specific model parameters. While the code is necessarily executable Python, the constrained scope is such that very limited knowledge of the Python language is required to write such an extension.

Linearization of the Proximal Penalty Terms: At each iteration $k \geq 1$ of PH, scenario sub-problem solves involve an augmented form of the original optimization objective, with both linear and quadratic penalty terms. The presence of the quadratic terms can cause significant practical difficulties. At present, no open-source linear or mixed-integer solvers currently support quadratic objective terms in an integrated, robust manner. While most commercial solvers can handle problems with quadratic linear and mixed-integer objectives, solver efficiency is often dramatically worse relative to the linear case: we have consistently observed scenario sub-problem solve times an order of magnitude or larger on quadratic mixed-integer stochastic programs relative to their linearized counterparts.

To address this issue, the **runph** script provides for automatic linearization of quadratic penalty terms in PH. We first observe that a linear expression results from the expansion of any quadratic penalty term involving binary variables. Consequently, the default behavior is to linearize these terms for binary variables. To linearize penalty terms involving continuous and general integer variables, the **runph** script allows specification of the following options:

```
--linearize-nonbinary-penalty-terms=BPTS
```

Approximate the PH quadratic term for non-binary variables with a piece-wise linear function. The argument BPTS gives the number of breakpoints in the linear approximation. Defaults to 0, indicating linearization is disabled.

```
--breakpoint-strategy=BREAKPOINT_STRATEGY
```

Specify the strategy to distribute breakpoints on the $[lb, ub]$ interval of each variable when linearizing. Defaults to 1.

To linearize a proximal term, **runph** requires that both lower and upper bounds (respectively denoted *lb* and *ub*) be specified for each variable in each scenario instance. This is most straightforwardly accomplished by specifying bounds or rules for computing bounds in each of the variable declarations appearing in the base deterministic scenario model. In reality, lower and upper bounds can be specified for all variables,

even if trivially. If for some reason bounds are not easily specified in the deterministic scenario model, the option `--bounds-cfgfile` option is available, which functions in a fashion similar to the mechanism for setting variable-specific ρ described above. Note that if a breakpoint would be very close to a variable bound, then the breakpoint is omitted. In other words, the BPTS parameter serves as an upper bound on the number of actual breakpoints.

Three breakpoint strategies are provided. A value of 1 indicates a uniform distribution of the BPTS points between lb and ub . A value of 2 indicates a uniform distribution of the BPTS points between the current minimum and maximum values observed for the variable at the corresponding node in the scenario tree; segments between the node min/max values and lb/ub are also automatically generated. Finally, a value of 3 places the half of the BPTS breakpoints on either side of the observed variable average at the corresponding node in the scenario tree, with exponentially increasing distance from the mean. Other, more experimental strategies are also provided.

By introducing automatic linearization of the proximal penalty term, PySP enables both a much broader base of solvers to be used in conjunction with PH and more efficient utilization of those solvers. In particular, it facilitates the use of open-source solvers – which can be critical in parallel environments in which it may be infeasible to procure large numbers of commercial solver licenses for concurrent use (see Section 8).

Other Command-Line Options: While not discussed here, the **runph** script also provides options to control the type and extent of output at each iteration (weights and/or solutions), specify solver options, report exhaustive timing information, and tracking intermediary solver files. In general, these are provided for more advanced users; more information can be obtained by supplying the `--help` option to **runph**.

6.3 Implementation Details

We now discuss high-level aspects of the implementation of the **runph** script, emphasizing the mechanisms linking the PH implementation with a generic Pyomo specification of the stochastic program. In doing so, our objective is to illustrate the power of embedding an algebraic modeling language within a high-level programming language, and specifically one that enables object introspection.

The PySP PH initialization process is similar to that for the EF writer/solver: the scenario tree, reference Pyomo instance, and scenario Pyomo instances are all created and initialized from user-supplied data. Without loss of generality, we assume two-stage problems in the following discussion. Following this general initialization, PH must for each first-stage variable create: ρ , node average vectors \bar{x} , and PH weight vectors w . This is accomplished by accessing the information in the *StageVariables* set, e.g., which in the farmer example contains the singleton (string) “DevotedAcreage[*]”. The “*” in this example indicates that non-anticipativity must be enforced at the root node for all indices of the variable *DevotedAcreage*, i.e., for all crop types (specifically, for variables *DevotedAcreage*[CORN], *DevotedAcreage*[SUGAR_BEETS], and *DevotedAcreage*[WHEAT]). Using Python introspection (via the `getattr` built-in function to query object attributes by name), PySP accesses the corresponding variable object in the reference model instance. From the variable object, the index set (also a first-class Python object) is extracted and cloned, eliminating all indices (none, in the case of a template equal to “*”) not matching the specified template.

PySP uses the newly constructed index set to create new parameter objects representing the ρ , weight w , and node average \bar{x} corresponding to the identified variable; the index set is the first argument to the parameter class constructor. Using the Python *setattr* method, the ρ and w parameters are attached to the appropriate scenario instance (the process is repeated for each scenario), while the node average \bar{x} is attached to the root node object in the scenario tree. The ability to create object attributes on-the-fly is directly supported in dynamic languages such as Python or Java, as opposed to C++ or other static and compiled strongly typed languages.

Following initialization, PH solves the original scenario sub-problems and loads the resulting solutions into the corresponding Pyomo instances. Using the same dynamic object query mechanism, PySP computes the first-stage variable averages and stores the result in the newly created parameters in the scenario tree. An analogous process is then used to compute and store the current w parameter values for each scenario. Before executing PH iterations $k \geq 1$, PySP must augment the original objective expressions with the linear and quadratic penalty terms discussed in Section 6.1. Because the Pyomo scenario instances and their attributes (e.g., parameters, variables, constraints, and objectives) are first-class Python objects, their contents can be programmatically modified at run-time. Consequently, it is straightforward to – for each first-stage variable – identify the corresponding variable, weight parameter, and average parameter objects, create objects representing the penalty terms, and augment the original optimization objective.

In summary, the processes described above rely on three capabilities explicitly facilitated through the use of Python. First, user-specified strings (e.g., first stage variables names) can be manipulated to dynamically identify attributes of objects (e.g., variables of scenario instances). Second, all elements of the Pyomo algebraic modeling language (e.g., parameters, variables, constraints, and objectives) are first-class Python objects, and as a consequence can be programmatically queried, cloned, and – most importantly – modified. Third, Python allows for the dynamic addition of attributes to objects (e.g., weight and ρ parameters to scenario instances). None of these enabling features of Python are particularly advanced, and are in general easy to use. Rather, these are key properties of a dynamic high-level programming language, that can be effectively leveraged to construct generic solvers for stochastic programming.

7 Progressive Hedging Extensions: Advanced Configuration

The basic PySP PH implementation is by design customizable to a rather limited degree: mechanisms are provided to allow for specification of ρ values and linearization of the PH objective. In either case, core PH functionality is not perturbed. We now describe more extensive and intrusive customization of the PySP PH behavior. In Section 7.1, we describe the interface to a PH extension providing functionality that is often critical to achieving good performance on stochastic mixed-integer programs. We then discuss in Sections 7.2 and 7.3 command-line options that enable functionality commonly often used in PH practice. Finally, we discuss in Section 7.4 the programmatic facilities that PySP provides to users (typically programmers) that want to develop their own extensions.

7.1 Convergence Accelerators and Mixed-Integer Heuristics

The basic PH algorithm can converge slowly, even if appropriate values of ρ have been computed. Further, in the mixed-integer case, PH can exhibit cyclic behavior, preventing convergence. Consequently, PH implementations in practice are augmented with methods to both accelerate convergence and prevent cycling. Many of these extensions are either described or introduced by Watson and Woodruff [2010].

The PySP implementation of PH provides these extensions in the form of a *plugin*, i.e., a piece of code that extends the core functionality of the underlying algorithm, at well-defined points during execution. This “Watson-Woodruff” (WW) plugin generalizes the accelerator and cycle-avoidance mechanisms described in Watson and Woodruff [2010]. The Python module implementing this plugin is named `wwextension.py`; general users do not need to understand the contents of this module.

The `runph` script provides three command-line options to control the execution of the Watson-Woodruff extensions plugin:

```
--enable-ww-extensions
Enable the Watson-Woodruff PH extensions plugin. Defaults to False.
--ww-extension-cfgfile=WW_EXTENSION_CFGFILE
The name of a configuration file for the Watson-Woodruff PH extensions plugin.
Defaults to “wwph.cfg”.
--ww-extension-suffixfile=WW_EXTENSION_SUFFIXFILE
The name of a variable suffix file for the Watson-Woodruff PH extensions plugin.
Defaults to “wwph.suffixes”.
```

As discussed in Section 7.4, user-defined extensions can co-exist with the Watson-Woodruff extension.

Before discussing the configuration of this extension (which necessarily relies on problem-specific knowledge), we provide more motivation and algorithmic detail underlying the extension:

- **Convergence detection:** A detailed analysis of PH behavior on a variety of problems indicates that individual decision variables frequently converge to specific, fixed values across all scenarios in early PH iterations. Further, despite interactions among the variables, this value frequently does not change in subsequent PH iterations. Such variable “fixing” behaviors lead to a potentially powerful, albeit obvious, heuristic: once a particular variable has converged to an identical value across all scenarios for some number of iterations, fix it to that value. However, the strategy must be used carefully. In particular, for problems where the constraints effectively limit variables x from both sides, these methods may result in PH encountering infeasible scenario sub-problems even though the problem is ultimately feasible.
- **Cycle detection:** In the presence of integer variables, PH occasionally exhibits cycling behavior. Consequently, cycle detection and avoidance mechanisms are required to force eventual convergence of the PH algorithm in the mixed-integer case. To detect cycles, we focus on repeated occurrences of the weight vectors w , heuristically implemented using a simple hashing scheme [Woodruff and Zemel, 1993] to minimize impact on run-time. Once a cycle in the weight vectors associated with any decision variable is detected, the value of that variable is fixed (using problem-specific, user-supplied knowledge) across scenarios in order to break the cycle.

- **Convergence-Based Sub-Problem Optimality Thresholds:** A number of researchers have noted that it is unnecessary to solve scenario sub-problems to optimality in early PH iterations [Helgason and Wallace, 1991]. In these early iterations, the primary objective is to quickly obtain coarse estimates of the PH weight vectors, which (at least empirically) does not require optimal solutions to scenario sub-problems. Once coarse weight estimates are obtained, optimal solutions can then be pursued to tune the weight vectors in the effort to achieve convergence. Given a measure of scenario solution homogeneity (e.g., the convergence threshold $g^{(k)}$), a commonly used strategy is to set the solver *mipgap* – a termination threshold based on the difference in current lower and upper bounds – in proportion to this measure.

Fixing variables aggressively typically results in shorter run-times, but the strategy can also degrade the quality of the obtained solution. Furthermore, for some problems, aggressive fixing can result in infeasible sub-problems even though the extensive form is ultimately feasible. Many of the parameters discussed in the next subsections control fixing of variables.

7.1.1 Mipgap Control and Cycle Detection Parameters

The WW extension defines and exposes a number of key user-controllable parameters, each of which can be optionally specified in the WW PH configuration file (named `wph.cfg` by default). The parameters, and a description of the corresponding feature they control, are given as follows:

- **Iteration0MipGap:** Specifies the mipgap for all PH iteration 0 scenario sub-problem solves. Defaults to 0, indicating that the solver *default* mipgap is used.
- **InitialMipGap:** Specifies the mipgap for all PH iteration 1 scenario sub-problem solves. Defaults to 0.1. A value equal to 0 indicates that the solver default mipgap is used. If a value $z > 0$ is specified, then no PH iteration $k > 1$ scenario sub-problem solves will use a mipgap greater than z . Let $g(1)$ denote the value of the PH convergence metric (independent of the particular metric used) at PH iteration 1. To determine the mipgap for PH iterations $k > 1$, the value of the convergence metric $g(k)$ is used to interpolate between the InitialMipGap and FinalMipGap parameter values; the latter is discussed below. In cases where the convergence metric $g(k)$ increases relative to $g(k - 1)$, the mipgap is thresholded to the value computed during PH iteration $k - 1$.
- **FinalMipGap:** The target final value for all PH iteration k scenario sub-problem solves at PH convergence, i.e., when the value of the convergence metric $g(k)$ is indistinguishable from 0 (subject to tolerances). Defaults to 0.001. The value of this parameter must be less than or equal to the InitialMipGap.
- **hash_hit_len_to_slam:** Ignore possible cycles in the weight vector associated with a variable for which the cycle length is less than this value. Also, ignore cycles if any variables have been fixed in the previous `hash_hit_len_to_slam` PH iterations. Defaults to the number of problem scenarios $|S|$. This default is often not good choice. For many problems with numerous scenarios, fixed constant values (e.g., such as 10 or 20) typically lead to significantly better performance.
- **DisableCycleDetection:** A binary parameter, defaulting to False. If True, cycle detection and the associated slamming logic (described below) are completely

disabled. This parameter cannot be changed during PH execution, as the data structures associated with cycle detection storage and per-iteration computations are bypassed.

Users specify values for these parameters in the WW PH configuration file, which is loaded by specifying the `--ww-extension-cfgfile runph` command-line option. To simplify implementation, the parameters are set directly using Python syntax, e.g., as follows:

```
self.Iteration0MipGap = 0.1
self.InitialMipGap = 0.2
self.FinalMipGap = 0.001
```

The contents of the configuration file are read by the WW extension following initialization. The `self` identifier refers to the WW extension object itself; the file contents are directly executed by the WW extension via the Python `execfile` command. While powerful and simplistic, this approach to initialization is potentially dangerous, as any attribute of the WW extension object is subject to manipulation.

7.1.2 General Variable Fixing and Slamming Parameters

Variable fixing is often an empirically effective heuristic for accelerating PH convergence. Fixing strategies implicitly rely on strong correlation between the converged value of a variable across all scenario sub-problems in an intermediate PH iteration and the value of the variable in the final solution should no fixing be imposed. Variable fixing reduces scenario sub-problem size, accelerating solve times. However, depending on problem structure, the strategy can lead to either sub-optimal solutions (due to premature declarations of convergence) or the failure of PH to converge (due to interactions among the constraints). Consequently, careful and problem-dependent tuning is typically required to achieve an effective fixing strategy. To facilitate such tuning, the WW PH extension allows for specification of the following parameters in the configuration file:

- **Iter0FixIfConvergedAtLB**: A binary-valued parameter indicating whether discrete variables that are at their lower bound in all scenarios after PH iteration 0 solves will be fixed at that bound. Defaults to False.
- **Iter0FixIfConvergedAtUB**: Analogous to the **Iter0FixIfConvergedAtLB** parameter, except applying to discrete variable upper bounds. Defaults to False.
- **Iter0FixIfConvergedAtNB**: Analogous to the **Iter0FixIfConvergedAtLB** parameter, except applying to discrete variable values not equal to either lower or upper bounds. Defaults to False.
- **FixWhenItersConvergedAtLB**: The number of consecutive PH iterations $k \geq 1$ (i.e., the lag) over which discrete variables must be at their lower bound in all scenarios before they will be fixed at that bound. Defaults to 10. A value of 0 indicates discrete variables will never be fixed at this bound.
- **FixWhenItersConvergedAtUB**: Analogous to the **FixWhenItersConvergedAtLB** parameter, except applying to discrete variable upper bounds. Defaults to 10.
- **FixWhenItersConvergedAtNB**: Analogous to the **FixWhenItersConvergedAtLB** parameter, except applying to discrete variable values not equal to either lower or upper bounds. Defaults to 10.

- **FixWhenItersConvergedContinuous:** The number of consecutive PH iterations $k \geq$ that continuous variables must be at the same, consistent value in all scenarios before they will be fixed at that value. Defaults to 0, indicating that continuous variables will not be fixed.

Fixing strategies at iteration 0 are typically distinct from those in subsequent iterations, e.g., iteration 0 agreement of acquisition quantities in a resource allocation problem to a value of 0 may (depending on the problem structure) indicate that no such resources are likely to be required. In general, longer lag times for PH iterations $k \geq 1$ yield better solutions, albeit at the expense of longer run-times; this trade-off is numerically illustrated in Watson and Woodruff [2010]. Differentiation between fixing behaviors at lower bounds, upper bounds, or intermediate values are typically necessary due to variable problem structure (e.g., variables being constrained from lower or upper bounds).

For many mixed-integer problems, PH can spend a disproportionately large number of iterations “fine-tuning” the values of a small number of variables in order to achieve convergence. Consequently, it is often desirable to force early agreement of these variables, even at the expense of sub-optimal final solutions. This mechanism is referred to as *slamming* in Watson and Woodruff [2010]. Slamming is also used to break cycles detected through the mechanisms described above. The WW PH extension supports a number of configuration options to control variable slamming, given as follows:

- **SlamAfterIter:** The PH iteration k after which variables will be slammed to force convergence. After this threshold is passed, one variable every other iteration is slammed to force convergence. Defaults to the number of scenarios $|S|$.
- **CanSlamToLB:** A binary parameter indicating whether any discrete variable can be slammed to its lower bound. Defaults to False.
- **CanSlamToUB:** Analogous to the **CanSlamToLB** parameter, except applies to discrete variable upper bounds. Defaults to False.
- **CanSlamToAnywhere:** Analogous to the **CanSlamToLB** parameter, that the variable can be slammed to its current average value across scenarios. Defaults to False.
- **CanSlamToMin:** A binary parameter indicating whether any discrete variable can be slammed to its current minimum value across scenarios. Defaults to False.
- **CanSlamToMax:** Analogous to the **CanSlamToMin** parameter, except applies to discrete variable maximum values across scenarios. Defaults to False.
- **PH.Iters.Between.Cycle.Slams:** Controls the number of PH iterations to wait between variable slams imposed to break cycles. Defaults to 1, indicating a single variable will be slammed every iteration if a cycle is detected. A value of 0 indicates an unlimited number of variable slams can occur per PH iteration.

Depending on the specified values, it is possible that no variable is allowed to be slammed. Slamming to the minimum and maximum scenario tree node values is often useful in resource allocation problems. For example, it is frequently safe with respect to feasibility to slam a variable value to the scenario maximum in the case of one-side “diet” problems. In the event that multiple slamming options are available, the priority order is given as: lower bound, minimum, upper bound, maximum, and anywhere.

7.1.3 Variable-Specific Fixing and Slamming Parameters

Global controls for variable fixing and slamming are generally useful, but for many problems more fine-grained control is required. For example, in one-sided diet prob-

lems, feasibility can be maintained during slamming by fixing a variable value at the maximal level observed across scenarios (assuming a minimization objective) [Watson and Woodruff, 2010]. Similarly, it is often desirable in a multi-stage stochastic program to fix variables appearing in early stages before those appearing in later stages, or to fix binary variables for siting decisions in facility location prior to discrete allocation variables associated with those sites.

The WW PH extension provides fine-grained, variable-specific control of both fixing and slamming using the concept of *suffixes*, similar to the mechanism employed by AMPL [AMPL, 2010]. Global defaults are established using the mechanisms described in Section 7.1.2, while optional variable-specific over-rides are specified via the suffix mechanism we now introduce.

The specific suffixes recognized by the WW PH extension include the following, with analogous (variable-specific) functionality to that provided by the parameters described in Section 7.1.2: `IterOfFixIfConvergedAtLB`, `IterOfFixIfConvergedAtUB`, `IterOfFixIfConvergedAtNB`, `FixWhenItersConvergedAtLB`, `FixWhenItersConvergedAtUB`, `FixWhenItersConvergedAtNB`, `CanSlamToLB`, `CanSlamToUB`, `CanSlamToAnywhere`, `CanSlamToMin`, and `CanSlamToMax`. In addition, we introduce the suffix `SlammingPriority`, which allows for prioritization of variables slammed during convergence acceleration; larger values indicate higher priority. The latter are particularly useful, for example, in the context of resource allocation problems in which early slamming of lower-cost items tends to yield lower-cost final solutions.

Variable-specific suffixes are supplied to the WW PH extension in a file, the name of which is communicated to the **runph** script through the `--ww-extension-suffixfile` option. An example of a suffix file (notionally implementing the motivational examples described above) is as follows:

```
resource_quantity[* , Stage1] FixWhenItersConvergedAtUB 10
resource_quantity[* , Stage2] FixWhenItersConvergedAtUB 50

resource_quantity[TIRES, Stage1] SlammingPriority 50
resource_quantity[ENGINES, Stage1] SlammingPriority 10

resource_quantity[TIRES, *] CanSlamToMax True
resource_quantity[ENGINES, *] CanSlamToMax True
```

In general, suffixes are specified via (**VARSPEC**, **SUFFIX**, **VALUE**) triples, where **VARSPEC** indicates a variable slice (i.e., a template that matches one or more indices of a variable; if the variable is not indexed, only the variable name is specified), **SUFFIX** indicates the name of a suffix recognized by the WW PH extension, and **VALUE** indicates the quantity associated with the specified suffix (and is expected to be consistent with the type of value expected by the suffix). If no suffix is associated with a given variable, then the global default parameter values are accessed.

In terms of implementation, suffixes are trivially processed via Python’s dynamic object attribute functionality. For each **VARSPEC** encountered, the index template (if it exists) is expanded and all matching variable value objects are identified. Then, for each variable value, a call to `setattr` is performed to attach the corresponding attribute/value pair to the object. The advantage of this approach is simplicity and generality: any suffix can be applied to any variable. The disadvantage is the lack of

error-checking, in that suffixes unknown to the WW PH extension can inadvertently be specified, e.g., a capitalized **SLAMMINGPRIORITY** suffix.

7.2 Solving a Constrained Extensive Form

A common practice in using PH as a mixed-integer stochastic programming heuristic involves running PH for a limited number of iterations (e.g., via the `--max-iterations` option), fixing the values of discrete variables that appear to have converged, and then solving the significantly smaller extensive form that results [Løkketangen and Woodruff, 1996]. The resulting compressed extensive form is generally far smaller and easier to solve than the original extensive form. This technique directly avoids issues related to the empirically long number of PH iterations required to resolve relatively small remaining discrepancies in scenario sub-problem solutions. Any disadvantage stems from variable fixing itself, i.e., premature fixing of variables can lead to sub-optimal extensive form solutions.

To write and solve the extensive form following PH termination, we provide the following options in the **runph** script:

`--write-ef`

Upon termination, write the extensive form of the model. Disabled by default.

`--solve-ef`

Following write of the extensive form model, solve the extensive form and display the resulting solution. Disabled by default.

`--ef-output-file=EF_OUTPUT_FILE`

The name of the extensive form output file. Defaults to “efout.lp”.

When writing the extensive form, all variables whose value is currently fixed in any scenario sub-problem are automatically (via Pyomo) preprocessed into constant terms in any referencing constraints or the objective. For reasons identical to those discussed in Section 4.1, PySP only supports output of the CPLEX LP file format. Solver selection is controlled with the `--solver` keyword, and is identical to that used for solving scenario sub-problems. The **runph** script additionally provides mechanisms for specifying solver options (including mipgap) specific to the extensive form solve.

7.3 Alternative Convergence Criteria

The PySP PH implementation supports a variety of alternative convergence metrics, enabled via the following **runph** command-line options:

`--enable-termdiff-convergence`

Terminate PH based on the **termdiff convergence metric**, which is defined (in Section 6.2) as the unscaled sum of differences between variable values and the mean. Defaults to True.

`--enable-normalized-termdiff-convergence`

Terminate PH based on the **normalized termdiff convergence metric**. Each term in the termdiff sum is normalized by the average variable value. Defaults to False.

`--enable-free-discrete-count-convergence`

Terminate PH based on the **free discrete variable count convergence metric**. Defaults to False.

```
--free-discrete-count-threshold=FREE_DISCRETE_COUNT_THRESHOLD
```

The convergence threshold associated with the **free discrete variable count convergence metric**. PH will terminate once the number of free discrete variables drops below this threshold.

The termination criterion associated with the free discrete variable count is particularly useful when deployed in conjunction with the capability to solve restricted extensive forms described in Section 7.2.

7.4 User-Defined Extensions

The **Watson-Woodruff PH extensions** described in Section 7.1 are built up a simple, general callback framework in PySP for developing user-defined extensions to the core PH algorithm. While most modelers and typical PySP users would not make use of this feature, programmers and algorithm developers can easily leverage the capability. The interface for user-defined PH extensions is defined in a PySP read-only file called **phextension.py**. The file contents are supplied as follows, to identify the points at which **runph** temporarily transfers control to the user-defined extension:

```
class IPHExtension(Interface):

    def post_ph_initialization(self, ph):
        """ Called after PH initialization."""
        pass

    def post_iteration_0_solves(self, ph):
        """ Called after the iteration 0 solves."""
        pass

    def post_iteration_0(self, ph):
        """ Called after the iteration 0 solves, averages \
            computation, and weight update."""
        pass

    def post_iteration_k_solves(self, ph):
        """ Called after the iteration k solves."""
        pass

    def post_iteration_k(self, ph):
        """ Called after the iteration k solves, averages \
            computation, and weight update."""
        pass

    def post_ph_execution(self, ph):
        """ Called after PH has terminated."""
        pass
```


To create a user-defined extension, one simply needs to define a Python class that implements the PH extension interface shown above, e.g., via the following code fragment:

```
from pyutilib.component.core import *
from coopr.pysp import phextension

class examplephextension(SingletonPlugin):

    implements (phextension.IPHExtension)

    def post_instance_creation(self, ph):
        print "Done creating PH scenario sub-problem instances!"

    # over-ride for each defined callback in phextension.py
    ...
```

The full example PH extension is supplied with PySP, in the form of the Python file `testphextension.py`. All Coopr user plugins are derived from a `SingletonPlugin` base class (indicating that there cannot be multiple instances of each type of user-defined extension), which can for present purposes be viewed simply as a necessary step to integrate the user-defined into the Coopr framework in which PySP is embedded. We defer to Hart and Sirola [2010] for an in-depth discussion of the Coopr plugin framework leveraged by PySP.

Each transfer point (i.e., callback) in the user-defined extension is supplied the PH object, which includes the current state of the scenario tree, reference instance, all scenario instances, PH weights, etc. User code can then be developed to modify the state of PH (e.g., current solver options) or variable attributes (e.g., fixing as in the case of the Watson-Woodruff extension).

To use a customized extension with `runph`, the user invokes the command-line option `--user-defined-extension=EXTENSIONFILE`. Here, `EXTENSIONFILE` is the Python module name, which is assumed to be either in the current directory or in some directory specified via the `PYTHONPATH` environment variable. Finally, we observe that both a user-defined extension and the Watson-Woodruff PH extension can co-exist. However, the Watson-Woodruff extension will be invoked prior to any user-defined extension.

8 Solving PH Scenario Sub-Problems in Parallel

One immediate benefit to embedding PySP and Pyomo in a high-level language such as Python is the ability to leverage both native and third-party functionality for distributed computation. Specifically, Python's `pickl` module provides facilities for object serialization, which involves encoding complex Python objects – including Pyomo instances – in a form (e.g., a byte stream) suitable for transmission and/or storage. The third-party, open-source Pyro (Python Remote Objects) package [PYRO, 2009] provides capabilities for distributed computing, building on Python's `pickl` serialization functionality.

PySP currently supports distributed solves, accessible from both the **runef** and **runph** scripts. At present, only a simple client-server paradigm is supported in the publically available distribution. The general distributed solver capabilities provided in the Coopr library are discussed in Hart et al. [2010], including the mechanisms and scripts by which name servers (used to locate distributed objects) and solver servers (daemons capable of solving MIPs, for example) are initialized and interact. Here, we simply describe the use of a distributed set of solver servers in the context of PySP.

Both the **runef** and **runph** scripts are implemented such that all requests for the solution of (mixed-integer) linear problems are mediated by a “solver manager”. The default solver manager in both scripts is a serial solver manager, which executes all solves locally. Alternatively, a user can invoke a remote solver manager by specifying the command-line option `--solver-manager=pyro`. The remote (pyro) solver manager identifies available remote solver daemons, serializes the relevant Pyomo model instance for communication, and initiates a solve request with the daemon. After the daemon has solved the instance, the solution is returned to the remote solver manager, which then transfers the solution to the invoking script.

The accessibility of remote solvers within the PySP PH implementation immediately confers the benefit of trivial parallelization of scenario sub-problem solves. In the case of commercial solvers, all available licenses can be leveraged. In the open-source case, cluster solutions can be deployed in a straightforward manner. Parallelism in PySP most strongly benefits stochastic mixed-integer program solves, in which the difficulty of scenario sub-problems masks the overhead associated with object serialization and client-server communication. At the same time, parallel efficiency necessarily *decreases* as the number of scenarios increases, due to high variability in mixed-integer solve times and the presence of barrier synchronization points in PH (after Step 4 in the pseudocode introduced in Section 6.1). However, parallel efficiency is of increasingly diminishing concern relative to the need to support high-throughput computing.

For solving the extensive form, remote solves serve a different purpose: to facilitate access to a central computing resources, with associated solver licenses. For example, our primary workstation for developing PySP is a 16-core workstation with a large amount of RAM (96GB). All commercial solver licenses are localized to this workstation, which in turn exposes parallelism enabled by now-common multi-threaded solver implementations.

Given the growing availability of cluster-based computing resources and the increasing accessibility of solver licenses (as is particularly the case for academics, with access to free licenses for most commercial solvers), the use of distributed computation by PySP users is expected to continue to grow.

9 Conclusions

Despite the potential of stochastic programming to solve real-world decision problems involving uncertainty, the use of this tool in industry is far from widespread. Historical impediments include the lack of stochastic programming support in algebraic modeling tools and the inability to experiment with and customize stochastic programming solvers, which are immature relative to their deterministic counterparts.

We have described PySP, an open-source software tool designed to address both impediments simultaneously. PySP users express stochastic programs using Pyomo, an open-source algebraic modeling language co-developed by the authors. In addition

to exhibiting the benefits of open-source software, Pyomo is based on the Python high-level programming language, allowing generic access and manipulation of model elements. PySP is also embedded within CoopR, which provides a wide range of solver interfaces (both commercial and open-source), problem writers, solution readers, and distributed solvers.

PySP leverages the features of Python, Pyomo, and CoopR to develop model-independent algorithms for generating and solving the extensive form directly, and solving the extensive form via scenario-based decomposition (PH). The resulting PH implementation is highly configurable, broadly extensible, and trivially parallelizable. PySP serves as a novel case study in the design of generic, configurable, and extensible stochastic programming solvers, and illustrates the benefits of integrating the core algebraic modeling language within a high-level programming language.

PySP has been used to develop and solve a number of difficult **stochastic mixed-integer, multi-stage programs**, and is under active use and development by the authors and their collaborators. PySP ships with a number of academic examples that have been used during the development effort, including examples from **Birge and Louveaux's test**, a **network flow problem** [Watson and Woodruff, 2010], and a **production planning problem** [Jorjani et al., 1999]. Real-world applications that have either been completed in PySP or are under active investigation include biofuel network design [Huang, 2010], forest harvesting [Badilla, 2010], wind farm network design, sensor placement, and electrical grid generation expansion.

In-progress and future PySP development efforts include new solvers (e.g., the **L-shaped method**), **scenario bundling techniques**, and support for **large-scale parallelism**. In addition, we have recently integrated capabilities for confidence interval estimation on solution quality.

Both PySP and the Pyomo algebraic modeling language upon which PySP is based are actively developed and maintained by Sandia National Laboratories. Both are packages distributed with the CoopR open-source Python project for optimization, which is now part of the COIN-OR open-source initiative [COIN-OR, 2010].

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Appendix: Getting Started

Both PySP and the underlying Pyomo modeling language are distributed with the CoopR software package. All documentation, information, and source code related to CoopR is maintained on the following web page: <https://software.sandia.gov/trac/>

`coopr`. Installation instructions are found by clicking on the *Download* tab found on the Coop main page, or by navigating directly to <https://software.sandia.gov/trac/coopr/wiki/GettingStarted>. A variety of installation options are available, including directly from the project SVN repositories or via release snapshots found on PyPi. Two Google group e-mail lists are associated with Coop and all contained sub-projects, including PySP. The *coopr-forum* group is the main source for user assistance and release announcements. The *coopr-developers* group is the main source for discussions regarding code issues, including bug fixes and enhancements. Much of the information found at <https://software.sandia.gov/trac/coopr> is mirrored on the COIN-OR web site (<https://projects.coin-or.org/Coopr>); similarly, a mirror of the Sandia SVN repository is maintained by COIN-OR.

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