

A new interior-point approach for large  
two-stage stochastic problems

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# A new interior-point approach for large two-stage stochastic problems

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Two-stage stochastic models give rise to very large optimization problems. Several approaches have been devised for efficiently solving them, including interior-point methods (IPMs). However, using IPMs, the linking columns associated to first-stage decisions cause excessive fill-in for the solution of the normal equations. This downside is usually alleviated if variable splitting is applied to first-stage variables. This work presents a specialized IPM that applies variable splitting and exploits the structure of the deterministic equivalent of the stochastic problem. The specialized IPM combines Cholesky factorizations and preconditioned conjugate gradients for solving the normal equations. This specialized IPM outperforms other approaches when the number of first-stage variables is large enough. This paper provides computational results for two stochastic problems: (1) a supply chain system and (2) capacity expansion in an electric system. Both linear and convex quadratic formulations were used, obtaining instances of up to 38 million variables and six million constraints. The computational results show that our procedure is more efficient than alternative state-of-the-art IPM implementations (e.g., CPLEX) and other specialized solvers for stochastic optimization.

**Keywords:** interior-point methods; stochastic optimization; structured problems; large-scale optimization

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## 1. Introduction

Stochastic programming is a framework for modelling optimization problems with uncertain data, and it has a wide range of applications in real-life situations. The variability in these models may appear in the objective or the constraints of the decision-making process, and it is due to their relationships to random elements [4]. Such problems can be solved for realistic instances—which are both large and complex—because advances in computing power and algorithm development allow finding solutions in a reasonable amount of time. However, larger instances increasingly demand more efficient solution techniques [1].

A two-stage stochastic model includes two types of variables. The *first-stage variables* occur at the beginning of the period and are the decisions to be made before the outcome of random events; and the *second-stage variables* are decided after the outcomes of uncertain events are known. In the extensive form of the model, all scenarios (i.e., possible realization of the random elements) are incorporated explicitly. This entails a special structure of the constraints matrix with separable blocks.

Two of the most common structures in large-scale optimization correspond to problems with *linking variables* between groups of constraints blocks (also named *dual block-angular*

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structure), and *linking constraints* between blocks (or *primal block-angular* structure) [5]. Two-stage stochastic problems belong to the class of problems with linking variables. The two classical methods of resolution for each type of structure are Benders decomposition for dual block-angular [3], and Dantzig-Wolfe for primal block-angular [11]. Interior-point methods (IPMs) [27] exploiting problem structure is a third technique that has also proven to be useful for these kinds of problems [6, 15–17]. Hence, large-scale refers not only to size but also to problem structure.

Linking variables of two-stage stochastic problems are usually dense columns, thus making them difficult to be solved by standard IPMs. The reformulation technique used in [21], named *splitting formulation*, removes this issue by considering copies of the first-stage variables, equating them through simple and sparse linking constraints. The specialized IPM implemented in the BlockIP package is tailored for problems with primal block-angular structure and linking constraints. It solves the normal equations by combining Cholesky factorizations for the diagonal blocks and a preconditioned conjugate gradient (PCG) for the linking constraints [6]. The preconditioner considered has proven to perform efficiently when the linking constraints are, precisely, simple and sparse [8, 9].

Therefore, the approach we propose in this work aims to efficiently solve two-stage stochastic problems by using the algorithm of the BlockIP solver for a *splitting formulation* of two-stage stochastic problems. Although this approach is not expected to be the most efficient algorithm for any stochastic problem, we will show that, in problems with a large enough number of first-stages variables, it outperforms the alternative state-of-the-art methods. Two standard problems from the literature were used to test this approach: a supply chain model, and a capacity expansion in an electric system. Both linear and quadratic formulations were used to obtain instances of up to 38 million variables and six million constraints.

The remainder of this document is organized as follows. Section 2 reviews the two-stage stochastic problem and its splitting formulation. Section 3 outlines the specialized interior-point method in BlockIP. Section 4 provides preliminary computational results comparing our approach with other specialized methods for two-stage stochastic problems in the solution of standard instances from the literature. Next, Section 5 presents the two stochastic problems and the computational results obtained, for both linear and quadratic instances. Finally, Section 6 summarizes the main outcomes of this paper and suggests directions for further research.

## 2. Two-stage stochastic optimization problems

The general idea behind a two-stage stochastic formulation is to compute some initial (first-stage) decisions that must be made prior to the occurrence of uncertainty, taking into account the expected value of future recourse (second-stage) actions. This work focuses on two-stage stochastic problems with either linear or convex quadratic objective functions. The *first-stage* problem manages the so-called “here-and-now” decisions. The *second-stage problem* refers to “wait-and-see” decisions, which represent the recourse actions. It deals with uncertainty by analyzing possible outcomes [4].

The convex quadratic two-stage stochastic problem can be formulated in standard form as

$$\begin{aligned} \min_x \quad & c^\top x + \frac{1}{2}x^\top Fx + Q(x) \\ \text{s. to} \quad & Mx = b \\ & u_x \geq x \geq 0, \end{aligned} \tag{1}$$

where

$$\mathcal{Q}(x) = E_{\xi}[\mathcal{Q}(x, \xi)] \quad \text{and} \quad \begin{aligned} \mathcal{Q}(x, \xi) = \min_y & q_{\xi}^{\top} y + \frac{1}{2} y^{\top} G_{\xi} y \\ \text{s. to } & W y = h_{\xi} - T_{\xi} x \\ & u_y \geq y \geq 0. \end{aligned} \quad (2)$$

The variables  $x \in \mathbb{R}^{n_x}$  and  $y \in \mathbb{R}^{n_y}$  are, respectively, the first- and second-stage decisions. The number of first- and second-stage constraints are, respectively,  $m_x$  and  $m_y$ . The first-stage vectors and matrices are  $c \in \mathbb{R}^{n_x}$ ,  $F \in \mathbb{R}^{n_x \times n_x}$ ,  $b \in \mathbb{R}^{m_x}$ ,  $M \in \mathbb{R}^{m_x \times n_x}$ , where  $F$  is symmetric and positive definite.  $\mathcal{Q}(x)$ , known as the recourse function, is the future average cost of our second-stage decisions, for all scenarios (i.e., for all realizations of  $\xi$ ). In the second stage,  $q_{\xi} \in \mathbb{R}^{n_y}$ ,  $G_{\xi} \in \mathbb{R}^{n_y \times n_y}$ ,  $W \in \mathbb{R}^{m_y \times n_y}$ ,  $T_{\xi} \in \mathbb{R}^{m_y \times n_x}$ , and  $h_{\xi} \in \mathbb{R}^{m_y}$ ;  $G_{\xi}$  is symmetric and positive definite. The stochastic random vector is comprised of  $q_{\xi}$ ,  $G_{\xi}$ ,  $h_{\xi}$  and  $T_{\xi}$ .

For some particular problems, a closed form solution can be obtained for  $\mathcal{Q}(x, \xi) = q_{\xi}^{\top} y^* + \frac{1}{2} y^{*\top} G_{\xi} y^*$ , where  $y^*$  is the optimum of the second-stage future decisions. In these cases, it is possible to compute  $\mathcal{Q}(x) = E_{\xi}[\mathcal{Q}(x, \xi)]$ , which allows for the solution of (1) only in terms of the first-stage decisions. In general, however, no closed form exists for  $\mathcal{Q}(x, \xi)$ , which forces us to use the extensive form of the stochastic problem. For this purpose, let us consider that  $\xi$  is a discrete random variable of  $k$  values  $\xi_1, \dots, \xi_k$  with probabilities  $p_1, \dots, p_k$  (if  $\xi$  is continuous, it must be previously discretized). Each particular value  $\xi_i$ ,  $i = 1, \dots, k$  is usually known as a scenario. Next, second-stage variables and constraints are replicated for each scenario (i.e.,  $y_i$ ,  $i = 1, \dots, k$  for variables), combining problems (1) and (2) into a single one, as follows:

$$\begin{aligned} \min_{x, y_i} & c^{\top} x + \frac{1}{2} x^{\top} F x + \sum_{i=1}^k p_i \left( q_{\xi_i}^{\top} y_i + \frac{1}{2} y_i^{\top} G_{\xi_i} y_i \right) \\ \text{s. to } & M x = b \\ & u_x \geq x \geq 0 \\ & \left. \begin{aligned} T_{\xi_i} x + W y_i &= h_{\xi_i} \\ u_y &\geq y_i \geq 0 \end{aligned} \right\} i = 1, \dots, k. \end{aligned} \quad (3)$$

For real problems, (3) can be very large and needs to be solved by specialized procedures that exploit the particular problem structure [4, Ch. 5–8], [19, 24].

### 2.1. The constraints structure of the two-stage problem

The constraints matrix in (3) shows the following dual block-angular structure

$$A = \begin{bmatrix} & x & y_1 & y_2 & \cdots & y_k \\ M & & & & & \\ T_1 & W & & & & \\ T_2 & & W & & & \\ \vdots & & & \ddots & & \\ T_k & & & & W & \end{bmatrix}, \quad (4)$$

where  $T_{\xi_i}$  in (3) has been renamed as  $T_i$  to simplify the notation. As shown in the top picture of Figure 1, the linking columns associated with the first-stage variables gives rise to large fill-in when computing the Cholesky factorization of the normal equations  $PA\Theta A^{\top} P^{\top}$  in an IPM—with  $\Theta$  being a positive diagonal matrix and  $P$  a row permutation

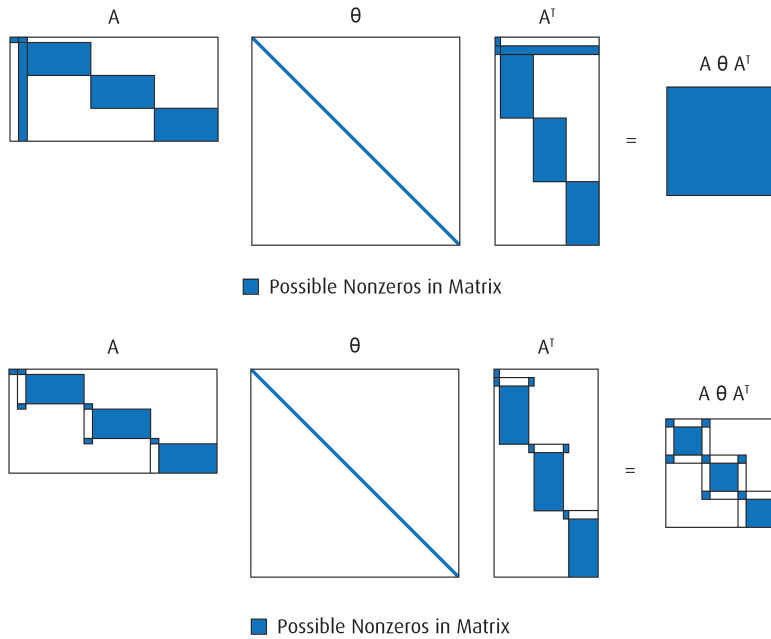


Figure 1.: Multiplication of  $A\Theta A^T$  with and without splitting [21]

matrix. This is true even for good permutation matrices. We note that  $F$  and  $G_\xi$  must be diagonal matrices if  $\Theta$  is assumed to be diagonal; indeed,  $F$  and  $G_\xi$  are considered diagonal in the specialized IPM of Section 3 for reasons of efficiency. Solving the normal equations system constitutes the main computational burden on any IPM. The solution time critically depends upon preserving the sparsity in matrix  $AA^T$  [21].

### 2.1.1. Splitting Formulation

The linking columns ensure that the first-stage decision variables  $x$  are feasible under each future scenario. Since the same first-stage decision variables  $x$  are used throughout, those necessarily satisfy the nonanticipativity requirement. Second-stage decisions  $y_i, i = 1, \dots, k$ , depend on both the first-stage variables and the realization of stochastic events.

In [21], variable splitting was suggested for reducing fill-in. Two variants were attempted: full and partial splitting. The full version split all first-stage variables, while in partial splitting only the dense first-stage variables were replicated. We will make use of full splitting, which is more appropriate than partial splitting for the specialized interior-point algorithm in this work, because it causes more highly sparse (but also much larger) matrices. This is different from general interior-point solvers, where partial splitting was shown to be superior in [21]. The bottom picture of Figure 1 shows the effect of splitting on the normal equations matrix.

Replicating the first-stage variables for each scenario one obtains the following equiva-

lent extensive form formulation:

$$\begin{aligned}
& \min_{x_i, y_i} c^\top x_1 + \frac{1}{2} x_1^\top F x_1 + \sum_{i=1}^k p_i \left( q_{\xi_i}^\top y_i + \frac{1}{2} y_i^\top G_{\xi_i} y_i \right) \\
& \text{s. to } M x_1 = b \\
& \quad \left. \begin{aligned} u_x &\geq x_1 \geq 0 \\ T_{\xi_i} x_i + W y_i &= h_{\xi_i} \\ u_y &\geq y_i \geq 0 \end{aligned} \right\} i = 1, \dots, k \\
& \quad \left. \begin{aligned} x_1 - x_i &= 0 \\ u_x &\geq x_i \geq 0 \end{aligned} \right\} i = 2, \dots, k.
\end{aligned} \tag{5}$$

Reordering columns by the number of scenarios, the constraint matrix in (5) can be written as:

$$A_{(5)} = \begin{bmatrix} & x_1 & y_1 & x_2 & y_2 & \cdots & x_k & y_k \\ M & & & & & & & \\ T_1 & W & & & & & & \\ & & T_2 & W & & & & \\ & & & & \ddots & & & \\ & & & & & T_k & W & \\ I & -I & & & & & & \\ \vdots & & & & \ddots & & & \\ I & & & & & & -I & \end{bmatrix}. \tag{6}$$

Alternative formulations can be obtained by different linking constraints that force the same values for copies of the first-stage variables. For instance, another equivalent problem is obtained by replacing the last group of constraints in (5) with  $x_i = x_{i+1}$ ,  $i = 1, \dots, k-1$ , thus obtaining

$$\begin{aligned}
& \min_{x_i, y_i} c^\top x_1 + \frac{1}{2} x_1^\top F x_1 + \sum_{i=1}^k p_i \left( q_{\xi_i}^\top y_i + \frac{1}{2} y_i^\top G_{\xi_i} y_i \right) \\
& \text{s. to } M x_1 = b \\
& \quad \left. \begin{aligned} u_x &\geq x_1 \geq 0 \\ T_{\xi_i} x_i + W y_i &= h_{\xi_i} \\ u_y &\geq y_i \geq 0 \end{aligned} \right\} i = 1, \dots, k \\
& \quad \left. \begin{aligned} x_i - x_{i+1} &= 0 \\ u_x &\geq x_i \geq 0 \end{aligned} \right\} i = 1, \dots, k-1.
\end{aligned} \tag{7}$$

The constraints matrix of formulation (7) is

$$A_{(7)} = \begin{bmatrix} & x_1 & y_1 & x_2 & y_2 & x_3 & y_3 & \cdots & x_{k-1} & y_{k-1} & x_k & y_k \\ M & & & & & & & & & & & \\ T_1 & W & & & & & & & & & & \\ & & T_2 & W & & & & & & & & \\ & & & & T_3 & W & & & & & & \\ & & & & & & \ddots & & & & & \\ & & & & & & & T_{k-1} & W & & & \\ & & & & & & & & & T_k & W & \\ I & & -I & & & & & & & & & \\ & & & I & & -I & & & & & & \\ & & & & & & \ddots & & & & & \\ & & & & & & & I & & -I & & \end{bmatrix}. \quad (8)$$

Although (5) and (7) are equivalent, the latter is computationally more efficient for the specialized IPM that will be used, as will be shown in Subsection 3.2.

Both constraint matrices (6) and (8) have a primal block-angular structure with  $(k - 1)n_x$  very sparse linking constraints. The linear programming problems (5) and (7) match the following general block-angular formulation:

$$\begin{aligned} \min_{x^1, \dots, x^k} \quad & \sum_{i=1}^k \left( c^i \top x^i + \frac{1}{2} x^i \top Q^i x^i \right) \\ \text{s. to} \quad & \begin{bmatrix} N_1 & & & & \\ & N_2 & & & \\ & & \ddots & & \\ & & & N_k & \\ R_1 & R_2 & \dots & R_k & I \end{bmatrix} \begin{bmatrix} x^1 \\ x^2 \\ \vdots \\ x^k \\ x^0 \end{bmatrix} = \begin{bmatrix} b^1 \\ b^2 \\ \vdots \\ b^k \\ b^0 \end{bmatrix} \\ & 0 \leq x^i \leq u^i \quad i = 1, \dots, k \\ & 0 \leq x^0 \leq b^0. \end{aligned} \quad (9)$$

For both formulations (5) and (7)  $x^i, u^i \in \mathbb{R}^{n_x+n_y}$  are, respectively,  $(x_i^\top \ y_i^\top)^\top$  and  $(u_x^\top \ u_y^\top)^\top$   $i = 1, \dots, k$ ;  $x^0$  are the slacks of the linking constraints (they are zero since linking constraints are equalities in the splitting formulation of two-stage stochastic problems). The linear cost vectors  $c^i \in \mathbb{R}^{n_x+n_y}$  are defined as  $c^1 = (c^\top \ p_1 q_{\xi_1}^\top)^\top$ , whereas  $c^i = (0 \ p_i q_{\xi_i}^\top)^\top$  for  $i = 2, \dots, k$ . The quadratic cost matrices  $Q^i \in \mathbb{R}^{(n_x+n_y) \times (n_x+n_y)}$  are  $Q^1 = \begin{bmatrix} F \\ G_{\xi_1} \end{bmatrix}$ , and  $Q^i = \begin{bmatrix} \mathbf{0} \\ G_{\xi_i} \end{bmatrix}$  for  $i = 2, \dots, k$ ; we assume  $Q^i$  are diagonal matrices. The right-hand-side terms are  $\mathbb{R}^{m_x+m_y} \ni b^1 = (b^\top \ h_{\xi_1}^\top)^\top$ , while  $\mathbb{R}^{m_y} \ni b^i = h_{\xi_i}$   $i = 2, \dots, k$ , and  $\mathbb{R}^{(k-1)n_x} \ni b^0 = 0$ . As for the diagonal blocks,  $N_1 \in \mathbb{R}^{(m_x+m_y) \times (n_x+n_y)}$  is  $\begin{bmatrix} M \\ T_1 \ W \end{bmatrix}$ , and  $N_i \in \mathbb{R}^{m_y \times (n_x+n_y)}$ , are  $[T_i \ W]$ ,  $i = 2, \dots, k$ .



Linking constraints for (5) are defined by

$$R_1 = \begin{bmatrix} I_1 & 0_1^y \\ \vdots & \vdots \\ I_{k-1} & 0_{k-1}^y \end{bmatrix} \quad R_i = \begin{bmatrix} 0_1^x & 0_1^y \\ \vdots & \vdots \\ 0_{i-2}^x & 0_{i-2}^y \\ -I_{i-1} & 0_{i-1}^y \\ 0_i^x & 0_i^y \\ \vdots & \vdots \\ 0_{k-1}^x & 0_{k-1}^y \end{bmatrix} \quad i = 2, \dots, k, \quad (10)$$

where  $0_i^x, I_i \in \mathbb{R}^{n_x \times n_x}$ ,  $0_i^y \in \mathbb{R}^{n_x \times n_y}$ , and the subindex denotes the block row position. For (7) the linking constraints are defined by

$$R_1 = \begin{bmatrix} I_1 & 0_1^y \\ 0_2^x & 0_2^y \\ \vdots & \vdots \\ 0_{k-1}^x & 0_{k-1}^y \end{bmatrix} \quad R_i = \begin{bmatrix} 0_1^x & 0_1^y \\ \vdots & \vdots \\ 0_{i-2}^x & 0_{i-2}^y \\ -I_{i-1} & 0_{i-1}^y \\ I_i & 0_i^y \\ 0_{i+1}^x & 0_{i+1}^y \\ \vdots & \vdots \\ 0_{k-1}^x & 0_{k-1}^y \end{bmatrix} \quad R_k = \begin{bmatrix} 0_1^x & 0_1^y \\ \vdots & \vdots \\ 0_{k-2}^x & 0_{k-2}^y \\ -I_{k-1} & 0_{k-1}^y \end{bmatrix}, \quad (11)$$

$$i = 2, \dots, k-1,$$

where  $I_i, 0_i^x$  and  $0_i^y$  have the same dimensions as above. We will denote as  $m_i$  and  $n_i$  the number of rows and columns of each diagonal block  $N_i$ , and by  $l$  the number of linking constraints. The next section outlines the specialized interior-point approach for the efficient solution of (9).

### 3. The specialized interior-point approach

The specialized IPM is based on a primal-dual path-following algorithm [26], that solves the normal equations by means of a scheme that sensibly combines direct sparse Cholesky factorizations [23] and iterative preconditioned conjugate gradient (PCG for short) solvers. This procedure was initially suggested for multicommodity flow problems [9] and later extended to primal block-angular problems [8]. Thus, it can be applied to problem (9). This specialized IPM was recently implemented in the BlockIP solver [6].

Denoting by  $A$  the constraint matrix in (9), the normal equations for the dual variables direction  $\Delta\lambda$  are

$$(A\Theta A^\top)\Delta\lambda = g, \quad (12)$$

where  $\Theta$  is a diagonal matrix of positive entries which are related to variables  $x^i$  and diagonal matrices  $Q^i$ ; and  $g$  is an appropriate right-hand-side. Exploiting the structure

of  $A$  and appropriately partitioning  $\Theta$ , the matrix of system (12) can be recast as

$$\begin{aligned}
A\Theta A^\top &= \left[ \begin{array}{c|c} N_1\Theta_1N_1^\top & N_1\Theta_1R_1^\top \\ \vdots & \vdots \\ N_k\Theta_kN_k^\top & N_k\Theta_kR_k^\top \\ \hline R_1\Theta_1N_1^\top \dots R_k\Theta_kN_k^\top & \Theta_0 + \sum_{i=1}^k R_i\Theta_iR_i^\top \end{array} \right] \\
&= \begin{bmatrix} B & C \\ C^\top & E \end{bmatrix},
\end{aligned} \tag{13}$$

$B \in \mathbb{R}^{\tilde{m} \times \tilde{m}}$  ( $\tilde{m} = \sum_{i=1}^k m_i$ ),  $C \in \mathbb{R}^{\tilde{m} \times l}$  and  $E \in \mathbb{R}^{l \times l}$  being the blocks of  $A\Theta A^\top$  and  $\Theta_i$ ,  $i = 0, \dots, k$ , the submatrices of  $\Theta$  associated with the  $k + 1$  groups of variables in (9). Appropriately partitioning  $g$  and  $\Delta\lambda$ , the normal equations can be written as

$$\begin{bmatrix} B & C \\ C^\top & E \end{bmatrix} \begin{bmatrix} \Delta\lambda_1 \\ \Delta\lambda_2 \end{bmatrix} = \begin{bmatrix} g_1 \\ g_2 \end{bmatrix}. \tag{14}$$

Eliminating  $\Delta\lambda_1$  from the first group of equations of (14), we get

$$(E - C^\top B^{-1}C)\Delta\lambda_2 = (g_2 - C^\top B^{-1}g_1) \tag{15}$$

$$B\Delta\lambda_1 = (g_1 - C\Delta\lambda_2). \tag{16}$$

System (16) is solved by performing one Cholesky factorization for each diagonal block  $N_i\Theta_iN_i^\top$ ,  $i = 1, \dots, k$  of  $B$ . System (15) is solved by a PCG. The dimension of this system is  $l$ , i.e., the number of linking constraints. A good preconditioner is imperative for the efficient solution of (15).

The preconditioner obtained in [9] for multicommodity flows can be applied to any primal block-angular problem [8]. The Neumann series preconditioner is based on the following result (see [9] for a proof):

$$(E - C^\top B^{-1}C)^{-1} = \left( \sum_{i=0}^{\infty} (E^{-1}(C^\top B^{-1}C))^i \right) E^{-1}. \tag{17}$$

The preconditioner, an approximation of  $(E - C^\top B^{-1}C)^{-1}$ , is thus obtained by truncating the infinite power series (17) at some term  $\phi$ . From (17), the quality of the preconditioner depends on the spectral radius  $\rho$  (i.e., the largest eigenvalue) of matrix  $(E^{-1}(C^\top B^{-1}C))$ , which is known to be in  $[0, 1)$  [9]. When  $\rho$  is not too close to 1, the contribution of higher-order terms of the Neumann series can be neglected, and just a few terms (i.e., a small  $\phi$ ) are enough for a good preconditioner. Since the preconditioner is used at each iteration of PCG for the solution of system  $(E - C^\top B^{-1}C)z = r$  (for some vectors  $z$  and  $r$ ), increasing  $\phi$  by one means solving an additional system with matrices  $B$  and  $E$  at each PCG iteration. Therefore, even though it is problem-dependent, we can consider as a rule of thumb  $\phi = 0$  or  $\phi = 1$  are reasonable choices.  $\phi = 0$  has been used for all the computational results in this paper. However, note that even for  $\phi = 0$ , one system with matrix  $E$  needs to be solved at each PCG iteration. Therefore, the efficient solution to systems with  $E$  is indispensable for the performance of the method. In next two sections we analyze the structure and efficient factorization of  $E$  for both formulations (5) and (7).

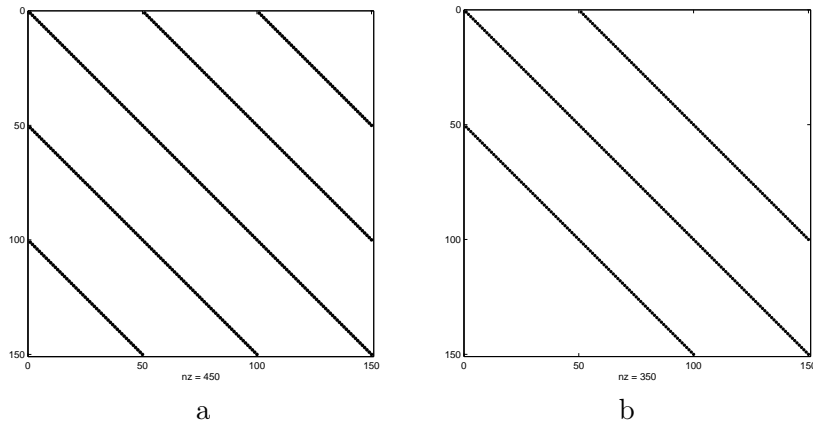


Figure 2.: Problem with  $n_x = 50$  first-stage variables and  $k = 4$  scenarios: a) Structure of (18) for formulation (5); b) Structure of (19) for formulation (7)

One of the important parameters for the efficient solution of (15) is the tolerance requested to the PCG solution. This tolerance is dynamically updated at each interior-point iteration  $i$  as  $\epsilon_i = \max\{\beta\epsilon_{i-1}, \min_\epsilon\}$ , where  $\epsilon_0$  is the initial tolerance (by default  $10^{-2}$  for linear problems, and  $10^{-3}$  for quadratic ones),  $\min_\epsilon$  is the minimum allowed tolerance (by default is  $10^{-8}$ ), and  $\beta$  is a tolerance reduction factor at each interior-point iteration (by default 0.95). Unless otherwise stated, the above default values were used in the computational results of Sections 4 and 5.

### 3.1. Structure of $E$ for formulation (5)

$\Theta_i$  is made of two diagonal submatrices,  $\Theta_i^x$  and  $\Theta_i^y$ , which are, respectively, related to first- and second-stage variables  $x_i$  and  $y_i$ . Since  $E = \Theta_0 + \sum_{i=1}^k R_i \Theta_i R_i^\top$ , and using (10), the structure of  $E$  is given by

$$E = \Theta_0 + \begin{bmatrix} \Theta_1^x + \Theta_2^x & \Theta_1^x & \cdots & \Theta_1^x & \Theta_1^x \\ \Theta_1^x & \Theta_1^x + \Theta_3^x & \cdots & \Theta_1^x & \Theta_1^x \\ \vdots & \vdots & \ddots & \vdots & \vdots \\ \Theta_1^x & \Theta_1^x & \cdots & \Theta_1^x + \Theta_{k-1}^x & \Theta_1^x \\ \Theta_1^x & \Theta_1^x & \cdots & \Theta_1^x & \Theta_1^x + \Theta_k^x \end{bmatrix}. \quad (18)$$

$E$  in (18) is a symmetric positive definite matrix and it is also partially banded, with only  $2(k-2) + 1$  non-zero diagonals:  $E_{ij} > 0$  if  $|i - j|$  is a multiple of  $n_x$  and  $E_{ij} = 0$  elsewhere. This is a particular case of the general band matrix (see, e.g., [12, Ch. 4]). Figure 2.a shows the structure of (18) for a problem with  $n_x = 50$  first-stage variables and  $k = 4$  scenarios, as well as  $2(k-2) + 1 = 5$  non-zero diagonals and 450 non-zero elements.

It is not difficult to prove that either the  $LDL^\top$  or Cholesky factorizations preserve the sparsity (zero fill-in) of (18); this guarantees an efficient factorization. However, we will omit such details. Instead, our focus in the next subsection will be on the  $E$  matrix of formulation (7), which is a more efficient alternative.

### 3.2. Structure of $E$ for formulation (7)

From the definition of  $E$  in (13) and those of  $R_i$  in (11), and considering as above that  $\Theta_i$  is made up of  $\Theta_i^x$  and  $\Theta_i^y$ , by block multiplication we get

$$E = \Theta_0 + \begin{bmatrix} \Theta_1^x + \Theta_2^x & -\Theta_2^x & & & & \\ -\Theta_2^x & \Theta_2^x + \Theta_3^x & -\Theta_3^x & & & \\ & & \ddots & \ddots & & \\ & & & -\Theta_{k-2}^x & \Theta_{k-2}^x + \Theta_{k-1}^x & -\Theta_{k-1}^x \\ & & & & -\Theta_{k-1}^x & \Theta_{k-1}^x + \Theta_k^x \end{bmatrix}. \quad (19)$$

Figure 2.b shows the structure of (19) for the same problem in Subsection 3.1, reformulated as in (7). The fundamental difference between (18) and (19) is that the latter has only three non-zero diagonals, independently of the number of scenarios. In practice, this leads to the efficient solution of systems  $Ez = r$  (for some  $z$  and  $r$ ).

Matrix (19) is a symmetric positive definite “ $n_x$ -shifted tridiagonal matrix”, a generalization of a tridiagonal matrix where the superdiagonal (nonzero diagonal above the main diagonal) and subdiagonal (nonzero diagonal below the main nonzero diagonal) are shifted  $n_x$  positions from the main diagonal, i.e., elements  $(i, j)$  are non-zero only if  $|i - j|$  is either 0 or  $n_x$ . Matrices with such a structure can be efficiently factorized with zero fill-in by extending a standard factorization for tridiagonal matrices. This particular factorization has been added to the BlockIP solver.

## 4. Preliminary computational results

The problem formulation (7) was implemented and solved using the BlockIP package [6, 8], which is based on the the specialized IPM of Section 3. The instances used for preliminarily testing the performance of the approach was a subset of the instances proposed in [1] and [13]. They are publicly available in SMPS format [14]. These two-stage stochastic instances have been used to measure performance in several solvers, such as in [15] and [28].

For testing the performance of the proposed specialized interior-point method, an initial comparison was made using the state-of-the-art IBM ILOG CPLEX (v.12.7) barrier optimizer. Unlike BlockIP, CPLEX computes directions by Cholesky factorizations, instead of a combination of Cholesky and PCG. CPLEX executions were made without crossover for a fair comparison with BlockIP. In this preliminary experiments, for BlockIP the problems were modelled using the splitting formulation (7), whereas for CPLEX the dual of the extensive form (3) was used, in an attempt to avoid dense columns due to the first-stage variables. All the experiments in this work were carried out on a Fujitsu Primergy RX2540 M1 4X server with two 2.6 GHz Intel Xeon E5-2690v3 CPUs (48 cores) and 192 Gigabytes of RAM, under a GNU/Linux operating system (openSuse 13.2), without exploitation of multithreading capabilities.

Table 1 shows a sample of the results with the instances referred to above. The first column indicates the name of each instance; the second column (named *1st*) gives the number of variables in the first stage; columns  $k$ ,  $m$ , and  $n$  refer to, respectively the number of scenarios, total constraints, and total variables. In the case of BlockIP, we show the number of interior-point iterations (column “Iter”), CPU time (column “CPU”) and total number of PCG iterations (column “PCG”). For CPLEX we have the number of interior-point iterations and CPU time. We considered an initial tolerance of  $\epsilon_0 = 10^{-2}$  for the PCG solutions (which is the default in BlockIP for linear problems). The optimality tolerances for BlockIP and CPLEX were set to  $10^{-3}$  (smaller values were difficult to

Instance	1st	k	m	n	BlockIP			CPLEX	
					Iter	CPU	PCG	Iter	CPU
LandS	4	3	23	62	9	0.00	41	6	0.00
AIRL_first	4	25	152	402	12	0.00	121	7	0.00
pltexpA2	188	6	686	2760	60	0.05	492	10	0.01
4node128	52	128	9486	32008	35	0.97	1167	25	0.30
4node4096	52	4096	303118	1024008	49	76.06	2938	45	17.30
4node16384	52	16384	1212430	4096008	59	522.10	5131	43	57.89
env_15	49	15	768	2046	21	0.03	78	14	0.01
env_1875	49	1875	90048	251286	64	9.09	1096	26	1.17
envDiss8232	49	8232	395184	1103124	135	154.36	4066	62	15.02
phone	8	32768	753665	3309569	21	38.02	255	23	26.00

Table 1.: Results of linear instances with BlockIP and CPLEX barrier

Instance	1st	k	m	n	BlockIP			CPLEX	
					Iter	CPU	PCG	Iter	CPU
LandS	4	3	23	62	9	0.00	48	6	0.00
AIRL_first	4	25	152	402	11	0.00	173	10	0.00
pltexpA2	188	6	686	2760	54	0.03	241	10	0.2
4node128	52	128	9486	32008	29	0.61	534	7	0.12
4node4096	52	4096	303118	1024008	31	36.60	984	7	6.81
4node16384	52	16384	1212430	4096008	31	137.87	1219	8	14.29
env_15	49	15	768	2046	14	0.01	46	13	0.02
env_1875	49	1875	90048	251286	15	2.24	261	17	1.11
envDiss8232	49	8232	395184	1103124	17	19.44	393	18	4.60
phone	8	32768	753665	3309569	13	21.95	169	18	12.88

Table 2.: Results of quadratic instances with BlockIP and CPLEX barrier

achieve by BlockIP in those instances due to the approximate solution of the Newton direction by PCG). The results of Table 1 indicate that BlockIP is slower in these kinds of instances where, although with a large number of scenarios, the number of first-stage variables is small.

From the above linear instances we created a set of quadratic stochastic problems by adding convex separable quadratic costs to the first and second stage variables; these quadratic terms were synthetic and of the same order as the linear costs. Table 2 reports the characteristics of these quadratic instances and the results obtained with BlockIP and CPLEX. In these executions the initial PCG tolerance was set to  $\epsilon_0 = 10^{-3}$  (the default value in BlockIP for quadratic problems). The optimality tolerances used for BlockIP and CPLEX were  $5 \cdot 10^{-2}$ . The specialized IPM in BlockIP is known to be more efficient for quadratic than for linear instances [7]. Then, as expected, the performance of BlockIP improved in these quadratic stochastic instances. However, it was still outperformed by CPLEX.

#### 4.1. Comparison with other specialized solvers for stochastic optimization

In addition to the CPLEX general purpose solver, we explored other specialized methods for two-stage stochastic problems, namely: Benders decomposition, and Benders decomposition with regularization by the level set method, as implemented in the FortSP stochastic solver [28]; the primal-dual column generation approach of [15]; and the dual decomposition approach (with an interior-point cutting-plane generator) implemented in the DSP (Decomposition for Structured Programming) stochastic solver [18]. Comparing BlockIP with these other approaches is not straightforward since: the solvers of [28]

Instance	1st	k	m	n	BlockIP			DSP	
					Iter	CPU	PCG	Algorithm	CPU
LandS	4	3	23	62	9	0.00	41	Benders	0
								Dual	0.3
AIRL_first	4	25	152	402	12	0.00	121	Benders	0.8
								Dual	6.6
pltxpA2	188	6	686	2760	60	0.05	492	Benders	0.1
								Dual	0.1
4node128	52	128	9486	32008	35	0.97	1167	Benders	3.7
								Dual	16529.1
4node4096	52	4096	303118	1024008	49	76.06	2938	Benders	1535.1
								Dual	—
phone	8	32768	753665	3309569	21	38.02	255	Benders	—
								Dual	—

— Instance exceeded the maximum memory allotted by Neos Server to a job

Table 3.: Results of linear instances with BlockIP and DSP

and [15] are not freely available; the DSP solver had to be used remotely from the *Neos Server* [10] (we did not succeed in installing it locally due to its many dependencies with third-party software); and the four hardwares (i.e., those of [28], [15], the Neos Server, and our work) were different, so only an indirect comparison can be performed.

Table 3 shows the results obtained with BlockIP and two of the algorithms in the DSP package (namely, a Benders decomposition and a dual decomposition) for a subset of the instances of Table 1. CPU times for DSP were provided by the Neos Server [22], so the comparison between our approach and DSP should be done with caution. When the instance exceeded the maximum memory allotted by the Neos Server to a job, the solution was not found; this is marked with “—” in Table 3. From the times reported, DSP seems not to be competitive with our approach, specially for the largest instances. A possible explanation might be that DSP was mainly designed for mixed integer stochastic problems.

Concerning the other two specialized methods, and according to the information provided by their authors, the processors used in [15] and [28] were, in single-thread mode, respectively a 27% and a 11% faster (in terms of Mflops, millions of floating point operations per second) than the one used in our work. The approach of [15] outperformed in general all the methods tested in [28], from the results in those papers. Therefore we will focus on the primal-dual column generation method of [15]. This approach required (in the hardware used in [15]) for the 10 instances of Table 1, respectively, 0.03, 0.03, 0.11, 0.56, 10.83, 37.97, 0.05, 0.68, 5.86 and 0.76 CPU seconds. Using the 27% correction factor between processors, if we had ran the approach of [15] in our hardware, the CPU times would have been, approximately, 0.04, 0.04, 0.15, 0.76, 14.77, 51.78, 0.07, 0.93, 7.99 and 1.03. We observe from Table 1 that the performance of CPLEX was very similar in those instances (excluding the last two, where the approach of [15] was significantly faster—specially for the last instance). Therefore, although being a general purpose solver, CPLEX can be considered a good candidate for benchmarking (this is also consistent with the results of [28], where the CPLEX barrier ranked among the best three algorithms for stochastic optimization). In addition, CPLEX can solve quadratic instances, while the approaches of [15] and [28] dealt only with linear problems. CPLEX will thus be the solver used to test our approach in next section, for the solution of more difficult instances provided by two particular applications.

## 5. Computational results for two particular applications

The instances of Tables 1 and 2 have a small number of first-stage linking variables, so they are loosely coupled among the different scenarios. For this reason, they can be considered “not too difficult”, and specially tailored for decomposition algorithms. In order to get more difficult instances, we modified two problems from the literature: (1) a stochastic supply chain problem based on [2], which will be described in Section 5.1; (2) and instance *LandS* of Section 4, which corresponds to a stochastic electricity generation problem and it is explained below in Section 5.2. The implementations developed for these two problems allowed us to manage the number of scenarios and the number of first- (mainly) and second-stage variables. Both problems were solved with CPLEX and BlockIP. For the solution with BlockIP they were modelled with the splitting formulation (7), thus obtaining the constraints structure (8). For CPLEX we considered two formulations: the splitting formulation (7) (as for BlockIP); and the dual of the original extensive form (3) (whose constraints matrix is the transpose of (4) for linear problems), such that linking variables are converted into linking constraints which, in principle, avoid the presence of dense columns, thus potentially increasing the performance of the CPLEX barrier algorithm.

### 5.1. The stochastic supply chain problem

This problem was created from the models in [2] and [25]. The original model considered a supply chain network  $\mathcal{G} = (\mathcal{N}, \mathcal{A})$ , where  $\mathcal{N}$  is the set of nodes and  $\mathcal{A}$  is the set of arcs. The set  $\mathcal{N}$  consists of suppliers  $\mathcal{S}$ , potential processing facilities  $\mathcal{P}$  and customer centers  $\mathcal{C}$ , i.e.,  $\mathcal{N} = \mathcal{S} \cup \mathcal{P} \cup \mathcal{C}$ . The decisions for first-stage are associated with a binary variable  $x_i$ , such that  $x_i = 1$  if processing facility  $i$  is built, and 0 otherwise. The second-stage refers to tactical decisions on routing the flow of some product from suppliers to customers. Variable  $y_{ij}$  denotes the flow of the product from a node  $i$  to a node  $j$  in the network, where  $(ij) \in \mathcal{A}$ , and  $z_j$  denotes a shortfall of the product at customer  $j$ , meaning that it is impossible to meet the demand. The random vector  $\xi$  includes the demands, supplies and costs (of processing, transportation and shortage).

In our model, modifications were made to expand the problem, especially the first-stage variables. First, the number of nodes for each set could be chosen (this is provided as input data), thus allowing to increase the number of first-decision variables as well as the number of scenarios for expanding the problem as much as is necessary. In addition, the capacity decision  $u_{ij}$  for arcs  $(ij) \in \mathcal{A}$  was included, where  $u_{ij}$  cannot be greater than  $M$ . This capacity could not be exceeded during transportation operations. Finally, we added the nodes of suppliers  $\mathcal{S}$  to the decision on whether or not to build the facility. Furthermore, all binary variables  $x_i$  were relaxed, i.e.,  $0 \leq x_i \leq 1$ .

The resulting stochastic supply chain design problem is formulated as follows:

$$\begin{aligned} \min_{x,u} \quad & \sum_{i \in \mathcal{S} \cup \mathcal{P}} c_i x_i + \sum_{(ij) \in \mathcal{A}} f_{ij} u_{ij} + Q(x, u) \\ \text{s. to} \quad & 0 \leq x_i \leq 1 \quad \forall i \in \mathcal{S} \cup \mathcal{P} \\ & 0 \leq u_{ij} \leq M \quad \forall (ij) \in \mathcal{A} \end{aligned} \quad \text{and} \quad Q(x, u) = E_{\xi}[Q(x, u, \xi)], \quad (20)$$

where  $Q(x, u, \xi)$  is the optimal value of the following problem:

$$Q(x, u, \xi) = \min_{y, z} \sum_{(ij) \in \mathcal{A}} q_{ij} y_{ij} + \sum_{j \in \mathcal{C}} h_j z_j \quad (21a)$$

$$\text{s. to } \sum_{i \in \mathcal{N}} y_{ij} - \sum_{l \in \mathcal{N}} y_{jl} = 0 \quad \forall j \in \mathcal{P} \quad (21b)$$

$$\sum_{i \in \mathcal{N}} y_{ij} + z_j \geq d_j \quad \forall j \in \mathcal{C} \quad (21c)$$

$$\sum_{j \in \mathcal{N}} y_{ij} \leq s_i x_i \quad \forall i \in \mathcal{S} \quad (21d)$$

$$\sum_{i \in \mathcal{N}} y_{ij} \leq m_j x_j \quad \forall j \in \mathcal{P} \quad (21e)$$

$$0 \leq y_{ij} \leq u_{ij} \quad \forall (ij) \in \mathcal{A} \quad (21f)$$

$$z_j \geq 0 \quad \forall j \in \mathcal{C}, \quad (21g)$$

whose parameters are:

- $c_i$  is the investment cost for supply and processing facility  $i$ .
- $f_{ij}$  is the cost per-unit of capacity on the arc  $ij$ .
- $q_{ij}$  represents the per-unit cost of transporting the product on arc  $ij$ .
- $h_j$  is the per-unit penalty incurred for failing to meet demand of the product at customer center  $j$ .
- $M$  denotes the maximum capacity per arc.
- $d_j$  is the demand of customer  $j$ .
- $s_i$  is the maximum supply capacity of supplier  $i$ .
- $m_j$  is the maximum processing capacity of processing plant  $j$ .

The first-stage problem (20) consists of choosing the design variables  $x_i$  and  $u_{ij}$ . The objective function minimizes the total investment for nodes  $\mathcal{S}$  and  $\mathcal{P}$ , arc capacities  $u_{ij}$ , and the expected value of the second stage. Binary variables  $x_i$  are relaxed, and the upper bound  $M$  is set for all arc capacities.

The second-stage problem (21) involves the processing and the transportation of the product, where the objective function (21a) minimizes the transportation and shortage costs. The first set of constraints (21b) enforces the flow conservation across each processing node  $j$ . The next group of constraints (21c) requires that the total flow for the customer  $j$  plus its shortfall should be at least the demand  $d_j$ . Constraints (21d) ensure that the total flow from a supplier node  $i$  should not exceed the supply capacity  $s_i$  at that node if it is built. The set of constraints (21e) establishes that the number of units to process should not be greater than the capacity  $m_j$  of facility  $j$  if it is built. If facility  $i \in \mathcal{S} \cup \mathcal{P}$  is not built, the above two groups of constraint will force all flow variables  $y_{ij} = 0$  for all  $i \in \mathcal{S} \cup \mathcal{P}$ . Finally, constraints (21f) and (21g) are the upper and lower bounds.

### 5.1.1. Test instances

The original model in [2] considered a supply chain with four suppliers (A, B, C, and D), four possible processing facilities (E, F, G and H), and three customer centers (L, M and N). This supply network is depicted in Fig. 3. The product considered in [2] was uniform-quality wine in bulk (raw material). Four scenarios were considered (boom, good, fair and poor), with different probabilities associated to each one. It should be stressed that in



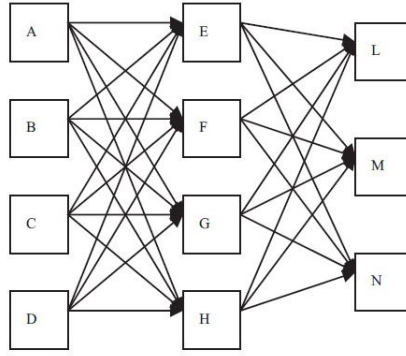


Figure 3.: The network of the wine company used in [2]

[2] the supplies, transportation costs and shortage costs were considered as deterministic parameters for the extensive form (or deterministic equivalent) formulation.

We extended the original model in [2] in several ways. First, all the parameters of the problem were created by means of a random generator considering different ranges of values. For instance, the range of investment costs for building each processing (bottling) plant was between 250,000 and 350,000. The costs per unit of arc capacities were between 1000 and 1,000,000. The unit production costs and market demands under each scenario were in the range of 500–750 and 550–800, respectively. The costs for transporting bulk wine from the suppliers to the processing facilities, and bottled wine from the processing facilities to the clients were, respectively, around 150 and 450. The unit storage costs at each distribution center were in the range of 10–16 thousand units. Furthermore, the maximum amount of bulk wine that can be shipped from the suppliers was between 250 and 350 units.

Next, we increased the number of first-stage variables by considering 10 suppliers, 10 processing facilities and 50 customers. This resulted in a problem with 620 first-stage variables, used as the base case from which the several instances in below Sections 5.3.1 and 5.4.1 are obtained by replicating the number of scenarios.

## 5.2. The stochastic electricity generation problem

This problem (based on [20]) consists of finding the optimal investment over various types of power plants to satisfy an uncertain electricity demand. A two-period model with a set of  $\mathcal{M}$  operating modes and a set of  $\mathcal{T}$  different technologies is considered. The modes are a discrete representation of the true load-duration demand curve, where each mode is a rectangular approximation of a part of this curve. It is necessary to consider large number of modes for a good approximation to the reality. The several power plants use one of the available technologies. See [1, 4] for more details.

Technology  $i \in \mathcal{T}$  has an associated investment cost  $c_i$  (a multiple of €/Mw), and a production cost  $q_i$  (a multiple of €/Mwh). Operating modes are defined by its duration  $t_j$  (hours) and demand  $d_j$  (Mw), for  $j \in \mathcal{M}$ ; the demand  $d_1 = \xi$  of the first mode is the stochastic parameter of this problem. First-stage decisions are the capacities  $x_i$  (Mw) invested in each technology  $i \in \mathcal{T}$ . Second-stage variables are the capacities  $y_{ij}$  effectively operated in each mode  $j \in \mathcal{M}$ , for each technology  $i \in \mathcal{T}$ . We assume technologies are always available, so they can be operated full-time.

The model formulation is the following:

$$\begin{aligned}
& \min_x \quad \sum_{i \in \mathcal{T}} c_i x_i + Q(x) \\
& \text{s. to} \quad \sum_{i \in \mathcal{T}} x_i \geq C \quad \text{and} \quad Q(x) = E_\xi[Q(x, \xi)], \\
& \quad \quad \sum_{i \in \mathcal{T}} c_i x_i \leq B \\
& \quad \quad x_i \geq 0 \quad i \in \mathcal{T}
\end{aligned} \tag{22}$$

where

$$Q(x, \xi) = \min_y \quad \sum_{i \in \mathcal{T}} \sum_{j \in \mathcal{M}} q_i t_j y_{ij} \tag{23a}$$

$$\text{s. to} \quad \sum_{j \in \mathcal{M}} y_{ij} \leq x_i \quad i \in \mathcal{T} \tag{23b}$$

$$\sum_{i \in \mathcal{T}} y_{ij} = \xi \quad j \in \{1\} \tag{23c}$$

$$\sum_{i \in \mathcal{T}} y_{ij} = d_j \quad j \in \mathcal{M} \setminus \{1\} \tag{23d}$$

$$y_{ij} \geq 0 \quad i \in \mathcal{T}, j \in \mathcal{M}, \tag{23e}$$

$B$  and  $C$  being, respectively, a budget limit and a minimum capacity to be provided in the first stage.

The first-stage problem (22) considers the investment cost for each technology, subject to a minimum total capacity and a maximum budget constraints. The objective function (23a) of the second-stage problem consists of minimizing the operational costs for each technology effectively used in every mode. The constraints (23b) impose that available capacities cannot be exceeded, for each technology. The other two sets of constraints (23c)–(23d) impose demand satisfaction for every mode; the first constraint is for the first mode, whose demand is stochastic.

### 5.2.1. Test Instances

The instance presented in [1, 4] considered  $m = 3$  operating modes and  $n = 4$  available technologies, along with demands  $d = (\xi, 2, 3)$  (that is, the demand of the first operating mode is stochastic) and load durations  $t = (10, 6, 1)$ . The three scenarios had the values of  $\xi = (3, 5, 7)$  and  $(0.3, 0.4, 0.3)$  probabilities, respectively. The investment costs were  $c = (10, 7, 16, 6)$  for each technology, with production costs  $q = (4, 4.5, 3.2, 5.5)$ . The budget keeps all investment below  $B = 120$  and minimum capacity is  $C = 12$ .

The previous basic instance was extended by increasing the number of technologies available (which is the same as the number of first-stage constraints), existing operating modes and scenarios for every instance. The rest of parameters were randomly generated from the values of the basic instance. For instance, the budget and minimum capacity varied according to the number of technologies. The random demand took values between 2 and 10, the investment costs  $c$  in the range 2–20,  $q$  between 2 and 6, and  $t$  went up to 10. The number of technologies (that is, first-stage decisions), was increased up to 600 (as it will be seen in below sections of results); in this case, rather than technologies, first-stage decisions could be considered as different power plants.

$k$	$m$	$n$	BlockIP			CPLEX		CPLEX	
			Iter	CPU	PCG	no splitting		with splitting	
						Iter	CPU	Iter	CPU
200	259380	511380	140	28.18	2747	19	149.76	21	109.34
400	519380	1023380	174	59.36	2504	25	424.54	30	262.32
600	779380	1535380	155	77.32	2076	32	616.63	29	250.59
800	1039380	2047380	192	128.81	2565	19	566.50	31	342.79
1000	1299380	2559380	200	312.07	6295	23	843.60	32	428.56

Table 4.: Results for supply chain problem up to 1000 scenarios

$k$	$m$	$n$	BlockIP			CPLEX		CPLEX	
			Iter	CPU	PCG	no splitting		with splitting	
						Iter	CPU	Iter	CPU
2000	2599380	5119380	48	48.34	204	11	647.42	20	627.82
3000	3899380	7679380	43	67.41	197	10	916.60	18	1208.98
4000	5199380	10239380	86	159.94	275	10	1242.50	18	2112.53
5000	6499380	12799380	81	214.08	379	10	1572.92	19	3678.25

Table 5.: Results for supply chain problem up to 5000 scenarios

### 5.3. Results for linear instances

Next two subsections present results for the linear instances of the supply chain and electricity generation problems. Those instances were obtained by increasing the number of first-stage variables and/or scenarios in the base instances described in previous sections. In these runs, the optimality gap was set to  $10^{-6}$  for both BlockIP and CPLEX, unless otherwise stated. For BlockIP, the initial tolerance and the tolerance reduction factor of the PCG were, respectively,  $\epsilon_0 = 10^{-2}$  and  $\beta = 0.95$  (which are BlockIP default values), unless otherwise stated.

#### 5.3.1. Linear instances of the stochastic supply chain problem

From the base instance of 10 suppliers, 10 processing facilities and 50 customers—with a total of 620 first-stage variables—described in above Section 5.1.1, we generated a first set of stochastic instances with 200, 400, 600, 8000 and 1000 scenarios. Table 4 shows the results obtained with BlockIP (splitting formulation), and CPLEX (for both the with and without splitting formulations). The meaning of the columns is the same as in previous tables. These instances resulted to be difficult for BlockIP, as the spectral radius  $\rho$  was close to one in the early iterations. Therefore, the optimality gap was set to  $10^{-5}$  for both BlockIP and CPLEX. The PCG tolerance reduction factor in BlockIP was also set to  $\beta = 1$  (that is, the PCG tolerance was not reduced at each interior-point iteration). From 4 it is clearly observed that BlockIP was more efficient than CPLEX either with or without splitting.

Table 5 reports results for a second set of larger instances with a number of scenarios between 2000 and 5000; the largest instance has 6.4M constraints and 12.8M variables. The optimality tolerance was increased to  $10^{-3}$  for both BlockIP and CPLEX, which can be considered a reasonable choice for stochastic models with a large number of scenarios, and at the same time it allowed us to avoid the last interior-point iterations where the spectral radius  $\rho$  usually tends to one. From 5 we see that BlockIP clearly outperformed both CPLEX variants. It is also worth to note that, unlike in Table 4, the CPLEX runs

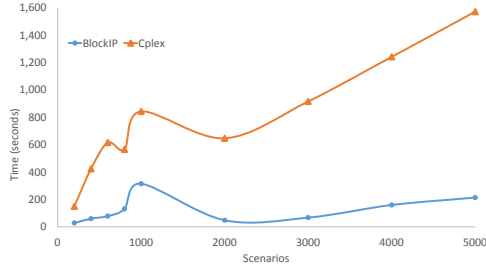


Figure 4.: BlockIP vs CPLEX for linear supply chain problem (CPLEX without splitting)

1st	m	n	BlockIP			CPLEX	
			Iter	CPU	PCG	Iter	CPU
400	80602	519602	83	7.14	559	23	87.50
450	90552	584552	85	8.20	533	20	86.42
500	100502	649502	81	9.02	470	23	110.02
550	110452	714452	90	12.45	714	28	139.01
600	120402	779402	85	12.00	491	19	108.76

Table 6.: Results for electricity generation problem for different number of first-stage variables

for the model with splitting were much slower than those without splitting. This fact will be even more evident in below Table 9 for the quadratic instances of the supply chain problem. This is due to the aggressive preprocessing applied by CPLEX to the splitting model, which removes a significant number of constraints and variables at the expense of modifying the matrix structure and significantly increasing the fill-in of the factorizations (some numbers will be provided below for the results of Table 9 for quadratic instances).

As summarized in Figure 4 (which plots the solution time against the number of scenarios for the instances of Tables 4 and 5), BlockIP always outperformed CPLEX, and one could infer that the difference between them will increase if the number of scenarios grows higher.

### 5.3.2. Linear instances of the stochastic electricity generation problem

We generated a preliminary set of instances where the first-stage variables (number of available technologies) were increased up to 600, the number of modes of electricity demand were set at 10, and the number of scenarios at 100. The default tolerances were used (i.e., optimality tolerance was  $10^{-6}$ ,  $\epsilon_0 = 10^{-2}$  and  $\beta = 0.95$ ). The performance of BlockIP was very promising, as it is shown in Table 6.

Encouraged by the previous results, we worked on the 600 first-stage variables instance while increasing the number of scenarios to 1000. Default tolerances were also used. The results are given in Table 7, which shows that the performance of BlockIP is much better than the one of CPLEX either with or without splitting. It is seen that CPLEX with either model required large executions in those instances. For this reason, the second set of even larger instances obtained by considering a number of scenarios between 2000 and 5000 were only ran with BlockIP. The results are provided in Table 8. BlockIP was very

$k$	$m$	$n$	BlockIP			CPLEX no splitting		CPLEX with splitting	
			Iter	CPU	PCG	Iter	CPU	Iter	CPU
			200	241402	1559402	89	30.07	703	22
400	483402	3119402	107	66.73	790	126	17633.54	40	10547.41
600	725402	4679402	79	81.93	691	87	54178.15	41	33770.16
800	967402	6239402	106	145.09	1036	20	45467.37	40	73447.25
1000	1209402	7799402	84	145.06	761	117	309381.56	44	167873.29

Table 7.: Results for electricity generation problem up to 1000 scenarios

$k$	$m$	$n$	BlockIP		
			Iter	CPU	PCG
2000	2419402	15599402	101	359.41	1074
3000	3629402	23399402	116	660.73	1204
4000	4839402	31199402	108	825.09	1193
5000	6049402	38999402	107	1027.31	1199

Table 8.: Results for electricity generation problem up to 5000 scenarios

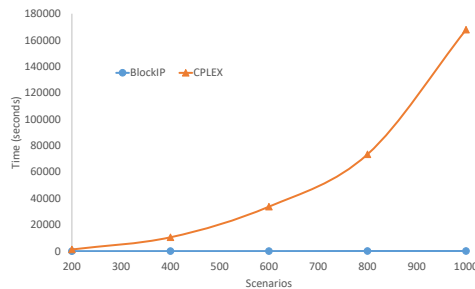


Figure 5.: BlockIP vs CPLEX for linear electricity generation problems (CPLEX with splitting)

efficient, being able to solve the largest case of 6M constraints and 39M variables in less than 18 minutes. It is also worth remarking that in those largest instances CPLEX ran out of memory (it exhausted the available 192 Gigabytes of RAM). The plot in Figure 5 (which reports the solution time against the number of scenarios) helps better appreciate that BlockIP is by far faster than CPLEX in those instances.

#### 5.4. Results for quadratic instances

From the previous linear instances we created a set of quadratic cases by adding convex separable quadratic costs to the first and second stage variables. The quadratic terms were synthetic and of the same order as the linear costs. The interest of testing quadratic instances relies on the fact that BlockIP is known to be faster for quadratic problems than for linear ones, since quadratic terms tend to reduce the spectral radius  $\rho$  (see [7] for details). In all the runs, the optimality tolerance for BlockIP and CPLEX was  $10^{-6}$ . For BlockIP the initial PCG tolerance and reduction factor were  $\epsilon_0 = 10^{-3}$  and  $\beta = 0.95$ ,

$k$	$m$	$n$	BlockIP			CPLEX no splitting		CPLEX with splitting	
			Iter	CPU	PCG	Iter	CPU	Iter	CPU
200	259380	511380	89	26.08	2774	17	132.12	15	1299.24
400	519380	1023380	146	78.34	3952	18	338.61	16	10960.21
600	779380	1535380	112	108.95	3703	20	403.64	15	36211.61
800	1039380	2047380	126	210.26	5661	19	548.12	16	102556.88
1000	1299380	2559380	129	289.83	6048	21	788.77	14	173633.19

Table 9.: Results for supply chain problem up to 1000 scenarios

$k$	$m$	$n$	BlockIP			CPLEX no splitting	
			Iter	CPU	PCG	Iter	CPU
2000	2599380	5119380	170	889.21	9538	22	1199.15
3000	3899380	7679380	193	1052.82	6834	21	1701.36
4000	5199380	10239380	147	1054.70	5206	20	2307.52
5000	6499380	12799380	149	1701.24	5282	19	2680.00

Table 10.: Results for supply chain problem up to 5000 scenarios

which are its default values for quadratic problems. As for the linear instances, the results with BlockIP were computed with the splitting formulation (7), while for CPLEX both the splitting formulation (7) and the (no splitting) dual of the extensive form (3) were considered (note that in the latter case the constraints matrix contains both the transpose of (4) and the terms associated to quadratic costs).

#### 5.4.1. Quadratic instances of the stochastic supply chain problem

As for the linear instances, we considered the base case of 620 first-stage variables and increased the number of scenarios up to 1000. The results can be seen in Table 9. Clearly, BlockIP outperformed both CPLEX runs (with and without splitting). It is also observed that CPLEX is much less efficient with splitting in those instances. After analyzing the CPLEX outputs, we concluded this is due to the large fill-in created to the reduced problem after the aggressive preprocessing applied by CPLEX. For example, for the 1000 scenarios case of Table 9, CPLEX without splitting has after preprocessing a reduced problem with 650620 rows, 1331240 columns, and 3621240 nonzeros, but the number of nonzeros in the lower triangle of  $AA^T$  is 22100000, and the number of nonzeros in the factor (after reordering) is 346453439. For the splitting model of the same instance these numbers are 680000 rows, 650620 columns, and 2970000 nonzeros (therefore, same order as for the no-splitting case), but the number of nonzeros in lower triangle of  $AA^T$  is 312500000 (14 times higher than for the model without splitting), and the number of nonzeros in the factor (after reordering) is 10855849207 (31 times higher than without splitting). This explains the poor performance of the splitting model with CPLEX in some cases.

Next, we raised the number of scenarios up to 5000. The results are shown in Table 10. CPLEX was not executed with the splitting model for these instances to avoid large CPU times. BlockIP is also faster than CPLEX. In short, the differences in this quadratic problem can also be seen in Figure 6, where BlockIP remains always below the time of CPLEX, even in small instances.

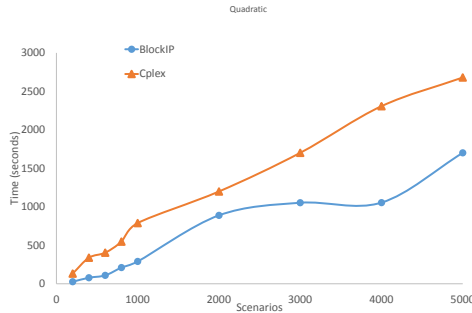


Figure 6.: BlockIP vs CPLEX for quadratic supply chain problems (CPLEX without splitting)

$k$	$m$	$n$	BlockIP			CPLEX no splitting		CPLEX with splitting	
			Iter	CPU	PCG	Iter	CPU	Iter	CPU
200	41902	259902	43	5.45	1760	71	336.45	65	10.02
400	83902	519902	42	11.19	1911	66	652.94	61	18.13
600	125902	779902	43	14.77	1471	67	996.91	69	37.18
800	167902	1039902	46	22.42	1575	73	1447.08	62	48.01
1000	209902	1299902	56	191.98	13604	72	1784.58	62	63.80
2000	419902	2599902	58	258.42	9037	73	3610.31	64	138.01
3000	629902	3899902	60	153.12	3205	74	5544.63	67	211.03
4000	839902	5199902	59	219.14	3493	78	7714.95	67	282.93
5000	1049902	6499902	61	227.20	2738	77	9599.25	67	363.00
10000	2099902	12999902	59	457.10	2509	75	19681.28	70	764.49

Table 11.: Results for electricity generation problem up to 10000 scenarios

#### 5.4.2. Quadratic instances of the stochastic electricity generation problem

We considered as the base case the instance with 100 first-stage variables, such that the number of scenarios can be largely increased without obtaining an out-of-memory error by CPLEX (like it happened for linear instances and the 600 first-stage variables base case). Table 11 presents the results for up to 10000 scenarios. In these executions the model with splitting was the most efficient variant for CPLEX; anyway, it was still outperformed by BlockIP except for the instances with 1000 and 2000 scenarios. In those two instances BlockIP required many PCG iterations and this explains its poor behaviour. The good results obtained in most cases with BlockIP are evident in Figure 7.

## 6. Conclusions

In this work, we explored the performance of a specialized interior-point algorithm for block-angular problems (implemented in the BlockIP package) for two-stage stochastic programming while using a splitting technique. The computational experiments used small- and large-scale instances to test both linear and quadratic objective functions. The performance of BlockIP was compared against alternative state-of-the-art interior-point (CPLEX barrier) and other specialized solvers for stochastic optimization. The results proved that BlockIP is competitive, mainly when the instances have a large number of

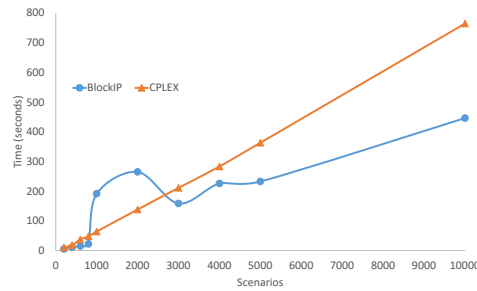


Figure 7.: BlockIP vs CPLEX for quadratic electricity generation problems (CPLEX with splitting)

first-stage variables. However, for the list of (small to medium) problems in [1], BlockIP was outperformed by the other approaches.

This new approach for stochastic optimization has potential in areas where decision-makers need to analyze many first-stage decisions, with a large number of future scenarios, and in a very short time.

Future studies should focus on the applicability of the specialized interior-point algorithm to block-angular reformulations of multistage stochastic problems.

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