# Similarity-based Decomposition Algorithm for Two-stage Stochastic Scheduling

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#### Abstract

This paper presents a novel decomposition method for two-stage stochastic mixedinteger optimization problems. The algorithm builds upon the idea of similarity between finite sample sets to measure how similar the first-stage decisions are among the uncertainty realization scenarios. Using such a Similarity Index, the non-anticipation constraints are removed from the problem formulation so that the original problem becomes block-separable on a scenario basis. Then, a term for maximizing the Similarity Index is included in all the sub-problems objective functions. Such sub-problems are solved iteratively in parallel, so that their solutions are used to update the weighing parameter for maximizing the Similarity Index. The algorithm gets a feasible solution when full similarity among scenario first stages is reached, that is, when the incumbent solution is non-anticipative. The proposal is tested in four instances of different size of an industrial-like scheduling problem. Comparison results show that the Similarity Index Decomposition provides significant speed-ups compared with the monolithic problem formulation, and provides simpler tuning and improved convergence over the Progressive Hedging Algorithm.

Keywords: Production planning, Mathematical programming, Uncertainty, Progressive Hedging, Mixed-Integer Optimization

# 1. Introduction and Problem Statement

Planning and scheduling deal with the optimal assignment of limited resources to processing tasks over time. Although deterministic scheduling formulations are mature, with broad industrial acceptance, they only translate to actual benefits in practice if the involved model parameters and inputs are precisely known.

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However, uncertainty over a future time horizon is inherent to all industrial processes. Some of the uncertain inputs that have important effects on the operations planning include: equipment uptime and availability, raw material arrival, prices, demands, weather, etc. Failing to correctly predict or estimate these values leads to suboptimal operation and can even result in infeasibility [3], where some operation constraints might need to be ignored to obtain a numerical solution

A well-known approach to incorporate uncertainty into scheduling applications is multi-stage stochastic optimization. In particular, two-stage formulations split the decision variables in two: first (here and now) and second (wait and see) stage variables, and considers a set of scenarios, associated with probabilities, for the realization of the uncertain parameters. Unfortunately, these formulations increase the number of variables of the problem proportionally to the number of scenarios, making it very difficult, or even impossible, to solve in reasonable times for online applications [12, 13, 23, 21]. This issue has received significant attention over the years: scenario bundling and reduction [10], reformulations, decomposition methods, etc.

Let us consider the following two-stage stochastic scheduling linear formulation in discrete time and extensive form:

$$\underset{y, z_e, r_e}{\text{minimize}} \quad \mathbf{c}^T y + \sum_{e \in \mathcal{E}} \mathbf{p}_e \mathbf{q}^T \begin{bmatrix} z_e \\ r_e \end{bmatrix} \tag{1a}$$

subject to

$$Ay \ge b$$
 (1b)

$$T_e y + W_e z_e + R_e r_e \ge h_e$$
 ,  $\forall e \in \mathcal{E}$  (1c)

$$y := \{ y_t \in \{0, 1\}^{n_y} | t \in \mathcal{T}_1 \} \tag{1d}$$

$$z_e := \{ z_{te} \in \{0, 1\}^{n_z} | t \in \mathcal{T}_2, e \in \mathcal{E} \}$$
 (1e)

$$r_e := \{ r_{te} \in \mathbb{R}^{n_s} | t \in \mathcal{T}_2, e \in \mathcal{E} \}$$
 (1f)

The scenarios  $e \in \mathcal{E}$  represent the future uncertainty realizations have and associated probabilities  $\mathbf{p}_e$ , such that  $\sum_e \mathbf{p}_e = 1$ . Time is discretized in periods  $t \in \mathcal{T}$ . The first stage is defined in the time periods  $\mathcal{T}_1 \subset \mathcal{T}$  by the parameters in  $\mathbf{A} \in \mathbb{R}^{m_1 \times n_y}$  and  $\mathbf{b} \in \mathbb{R}^{m_1}$ , weights  $\mathbf{c} \in \mathbb{R}^{n_y}$ , and variables  $y_t$ . The second stage spreads over the remaining time periods  $\mathcal{T}_2 \subset \mathcal{T}$  and involves constraints per scenario defined by parameters  $\mathbf{T}_e \in \mathbb{R}^{m_2 \times n_y}$ ,  $\mathbf{W}_e \in \mathbb{R}^{m_2 \times n_z}$ ,  $\mathbf{R}_e \in \mathbb{R}^{m_2 \times n_s}$ ,  $\mathbf{h}_e \in \mathbb{R}^{m_2}$ , weights  $\mathbf{q} \in \mathbb{R}^{n_z + n_s}$ , as well as variables  $z_{te}$  and  $r_{te}$ .

To exploit the problem structure for decomposition purposes, the first-stage variables are expanded to the scenarios so that they become  $y_{te}$  as in (2). However, for such problem to be equivalent to (1), the so-called non-anticipativity constraints (2d) need to be enforced too.

$$\underset{y_e, z_e, r_e}{\text{minimize}} \quad \sum_{e \in \mathcal{E}} p_e \left( c^T y_e + q^T \begin{bmatrix} z_e \\ r_e \end{bmatrix} \right)$$
subject to

$$Ay_e \ge b$$
 ,  $\forall e \in \mathcal{E}$  (2b)

$$T_e y_e + W_e z_e + R_e r_e \ge h_e$$
 ,  $\forall e \in \mathcal{E}$ , (2c)

$$y_{te_1} = y_{te_2} = \dots = y_{t|\mathcal{E}|}$$
,  $\forall t \in \mathcal{T}_1$  (2d)

$$y_e := \{ y_{te} \in \{0, 1\}^{n_y} | t \in \mathcal{T}_1, \forall e \in \mathcal{E} \}$$
 (2e)

$$z_e := \{ z_{te} \in \{0, 1\}^{n_z} | t \in \mathcal{T}_2, e \in \mathcal{E} \}$$

$$(2f)$$

$$r_e := \{ r_{te} \in \mathbb{R}^{n_s} | t \in \mathcal{T}_2, e \in \mathcal{E} \}$$
 (2g)

Similarly, a wide variety of scheduling models can be written in the state-space form to fit within the framework of model predictive control (MPC) [24]. This allows applying techniques and theoretical results from MPC to guarantee performance and nominal feasibility of closed-loop scheduling. Equation (3) shows a two-stage MILP scheduling problem in state-space form. Time is discretized in periods  $t \in \mathcal{T}$ , with  $\mathcal{T}_1 \subset \mathcal{T}$  defining the first-stage horizon.  $x_e := (x_e(0), x_e(1), \dots, x_E(N))$  and  $u_e := (u_e(0), u_e(1), \dots, u_e(N-1))$  represent the state and input trajectories for each scenario  $e \in \mathcal{E}$ .  $d_e := (d_e(0), d_e(1), \dots, d_e(N-1))$  denotes the measured and unmeasured disturbances.

$$\underset{u_e}{\text{minimize}} \quad \sum_{e \in \mathcal{E}} p_e \left( \mathbf{q}^T x_e + \mathbf{w}^T u_e \right) \tag{3a}$$

subject to

$$x_e(k+1) = Ax_e(k) + Bu_e(k) + Dd_e(k)$$
,  $\forall k \in \mathcal{T} \setminus N, e \in \mathcal{E}$  (3b)

$$\mathbf{E}^{(x)}x_e + \mathbf{E}^{(u)}u_e + \mathbf{E}^{(d)}\mathbf{d}_e + \mathbf{g} \le 0 \qquad , \forall e \in \mathcal{E}$$
(3c)

$$x_e(0) = \mathbf{x}(0)$$
 ,  $\forall e \in \mathcal{E}$  (3d)

$$x_e(N) \in \mathcal{X}_f(N)$$
 ,  $\forall e \in \mathcal{E}$  (3e)

$$u_1(k) = u_2(k) = \dots = u_{n_0}(k) \qquad , \forall k \in \mathcal{T}_1$$
 (3f)

Where (3b) predicts the evolution of the state trajectory depending on the previous state, input and disturbances. (3c) represents the constraint set that bounds the feasibility region of the problem. The terminal constraints of the problem are denoted by  $\mathcal{X}_f(N)$ . Finally, the non-anticipativity constraints are represented by (3f).

The scheduling problems (2) and (3) are similar in the sense that if the non-anticipativity (2d) and (3f) constraints were removed from the formulations, each scenario would represent an independent optimization problem. This is exactly what enables some of the decomposition methods available in the literature.

The Progressive Hedging Algorithm [19] and Dual Decomposition in stochastic integer optimization [5] are the most widely used decomposition methods for two-stage scheduling problems and much of the current research focus on building advanced algorithms upon them. Benders Decomposition [2] is also often used, but its main drawback is that the solution of the master problem becomes a limiting factor for large-scale problems [14].

The Progressive Hedging Algorithm (PHA) combines the proximal method with the alternating direction method of the multipliers [4]. It is often used for

solving multi-stage stochastic optimization problems as it enables scenario-based decomposition by relaxing the non-anticipation constraints. It is based on penalizing the deviation of the first-stage solution from an aggregated reference one computed from the results got at a previous iteration. Although it was originally devised to handle convex problems involving only continuous variables, it is widely used as a heuristic for problems with integer variables [18]. Consequently, the convergence and optimality properties are not guaranteed for this last kind of problems. Nonetheless, according to [8] it is possible to assess the quality of obtained solutions by computing lower bounds at PHA iterations, and [27] proposed a set of improvements focusing on updating the penalty parameter, accelerating convergence, and enhancements to the termination criteria, albeit the main underlying limitations remain.

Dual Decomposition is a popular method for decomposing problems with continuous variables. [5] combined the Dual Decomposition method with a branch and bound algorithm to achieve convergence in two-stage and multi-stage stochastic problems. In this method, the non-anticipativity constraints are relaxed and a lower bound of the problem is obtained by solving the respective Lagrangian dual problem. This lower bound can be used to guide the computation of feasible solutions (upper bounds). The main difficulty associated with the Dual Decomposition method is that getting feasible solutions from problems without complete recourse is not guaranteed [11]. To overcome this, [11] proposed to add Benders-like cuts in order to tighten the subproblems and exclude infeasible first-stage solutions. The Dual Decomposition method has also been used together with the Benders Decomposition method in a hybrid method to overcome the main drawbacks of the latter [6].

Although the literature is extensive, the success and performance of the available heuristic decomposition methods for mixed-integer programming is often problem dependent, and strongly depends on finding a good parameter tuning, which is not an easy task [26]. Therefore, we aim to provide the operations-research community with a tool that facilitates dealing with two-stage stochastic scheduling problems of industrial size in real time.

The Jaccard, a.k.a. Tanimoto, coefficient [25, 9] is a widely used index in information and computer sciences to measure *similarity* among sets of binarized data [28]. In brief, it is defined as the size of the intersection divided by the size of the union of the sample sets. Alike this concept, this paper presents a decomposition method based on a Similarity Index (SI) that arises as a good alternative to handle two-stage stochastic scheduling problems of the form (2). A single SI index is used to replace the non-anticipativity constraints, enabling scenario-based decomposition in a natural way. The similarity between the scenarios solutions is then maximized in each iteration of the procedure until the first stage fits equal for all.

This paper is the extension of the authors' preliminary idea in [15, 16]. Here the SI decomposition algorithm is formally stated, its convergence is analyzed, and its performance, comparing with other alternatives, is assessed in a realistic case study together with an analysis on the tuning parameters. Results show good performance in terms of convergence and optimality. Furthermore, the SI

algorithm turned out to be easier to tune than the PHA.

The paper is organized as follows. Section 2 explains how to use the Similarity Index to compare different scenario solutions (schedules). Section 3 describes the novel decomposition algorithm based on such index and analyzes its convergence properties. Section 4 presents an industrial-like case study that is used in Section 5 to test different aspects of the proposed decomposition method and to compare it with the PHA. The paper closes on Section 6 with some concluding remarks and proposals for future work.

# 2. Similarity Index

The authors of [17] were interested in obtaining solutions for two-stage stochastic scheduling problems up to a desired robustness level. For such aim, they first proposed a Similarity Index as a way to quantify the risk of the computed stochastic schedule not being optimal (or even feasible) in practice if the realized values of the uncertain inputs/parameters were not explicitly included within the, usually rough, scenario-tree approximation. In this context, the SI is a measurement of the similarity among the second-stage decisions. Hence, a desired minimum robustness level can be enforced via lower bounding the SI in the mathematical MILP problem. In such an instance, specifying SI = 100% implies getting the risk-averse solution, whereas  $SI \geq 0\%$  results in obtaining the less conservative two-stage stochastic solution.

The core idea behind the SI is basically the same of the Jaccard index, applied to compare discrete decisions among scenario schedules (binary decision sets) and weigh similarity if they are made in close, or neighboring, time periods. In scheduling context, a given decision is said to be equal in two scenarios if it is made exactly at the same time period in both. Therefore, the Jaccard index could directly serve for the task. However, what if, in one of those scenarios, such decision is made in the previous or following time period? The schedules would not be the same, but they might be considered *similar*. Nevertheless, the Jaccard index weighs zero in such a case, because the intersection is null. An idea for providing the MIP optimizer with a better conditioned problem and sensitivity, is assigning positive weights  $w \in \mathbb{R}$  (not depending upon scenarios) to each decision taken in the schedules to compare, and then compute the Jaccard index from the resulting non-binary sets [28]. The proposed Similarity Index assigns such weights w by fuzzifying the discrete decisions along the surrounding time periods for each scenario, as in Figure 1. Hence, the contribution of a decision to the overall similarity decreases the further away it is made from the time period under comparison (that is, it is less similar).

Now, consider that there is a decision to be compared at time t among two uncertainty realization scenarios (A and B). However, the decision is made at t-2 and t+2 in scenario A and B respectively. The result of fuzzifying such decisions along two surrounding time periods is shown in Figure 2a. It is clear that the fuzzified areas do not overlap at instant t so, consequently, the schedules are not similar at all. Then, SI=0.

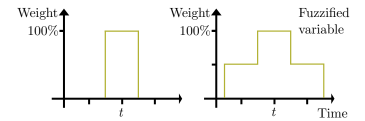
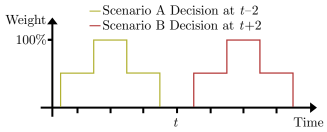
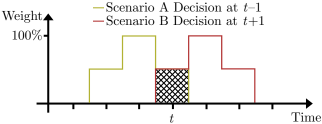


Figure 1: Fuzzifying a decision along the following and previous time periods.



(a) No similarity when the decision is made at  $t\pm 2$  in scenarios A and B.



(b) Similarity when a decision made at  $t\pm 1$ . The fuzzified areas partially overlap, so they are similar up to a certain extent.

Figure 2: Examples of similarity computation between two scenarios.

Figure 2b shows the fuzzified areas generated by a decision made at  $t \pm 1$  in scenarios A and B. The areas overlap partially in t, and one can say that the scheduled decisions are similar to a certain extent. The intersection area is 25% of the total area generated by the fuzzification. Then, SI = 0.25.

When the decisions are made at the same time instant there is complete overlap. Then, the generated intersection area at time t is the total one defined by the fuzzification. Thus, the schedules are 100% similar, that is, SI=1.

The overall Similarity Index is conceptually defined as the total intersection area divided by the maximum intersection possible, i.e., the total area generated by the fuzzified decisions:

$$SI := \frac{\text{Intersection Area}}{\text{Maximum Area Possible}} \tag{4}$$

Hence, given a set of schedules corresponding to different scenarios, the

overall SI is computed by aggregation of the intersected areas for each binary variable and time period in the schedule, divided by the total full-intersection area.

Note that the Similarity Index was first proposed to compare second-stage solutions. However, the same concept can be applied to the first-stage variables to replace, or approximate, the non-anticipativity constraints. These constraints restrict the first-stage solution to be equal among the scenarios, which would imply that the Similarity Index is equal to 100%. Our decomposition method is based precisely on this idea. The non-anticipativity constraints are replaced by the Similarity Index, which is then iteratively pushed up through the objective function until it reaches 100%.

**Definition.** The overall Similarity Index for the first-stage discrete variables of a decomposed stochastic scheduling (2) is computed by:

$$SI := \frac{\sum_{t=1}^{t_R} \left\| \min_{e \in \mathcal{E}} \left\{ y_{te} + \sum_{\tau=1}^{\Delta} \frac{\Delta - \tau}{\Delta} (y_{(t-\tau)e}|_{t>\tau} + y_{(t+\tau)e}|_{t+\tau \le t_R}) \right\} \right\|_{1}}{t_R \Delta - 2 \sum_{\tau=1}^{\Delta} \tau \frac{\Delta - \tau}{\Delta}}$$
(5)

Where  $y_{te} \in \{0,1\}$  are the binary decisions at time t in scenario e.  $\Delta \in \mathbb{N}$  is the length of the fuzzification horizon in time periods. It includes the current time period plus the ones to the right (or the left) which are considered for weighing similarity.  $t_R \in \mathbb{N}$  is the number of time periods in the first stage, and  $t_1$  is the first time period of this stage.  $\tau \in \mathbb{N}$  is an auxiliary time index used to move along the periods in the fuzzification horizon and weigh the made decision accordingly.

In this way, setting  $\Delta = 2$  results in weighting only the immediate neighboring periods to the current one for similarity; if the two previous and following periods, besides the current one, are considered, then  $\Delta = 3$ , and so on. For instance, when  $\Delta = 2$ , (5) reduces to:

$$SI = \frac{1}{2t_R - 1} \sum_{t=1}^{t_R} \left\| \min_{e \in \mathcal{E}} \left\{ y_{te} + 0.5 y_{(t-1)e}|_{t > t_1} + 0.5 y_{(t+1)e}|_{t < t_R} \right\} \right\|_1$$
 (6)

Of course, the area of the extreme tails that would lie outside the robust horizon is not included in the total intersection area. Hence,  $\Delta \leq \lceil t_R/2 \rceil$ , as there is no sense in fuzzifying each decision outside the robust horizon.

Remark. Setting  $\Delta=1$  in (5) reduces to the standard Jaccard/Tanimoto index comparing binary data sets. Moreover, the denominator (i.e. the union between given sets in the Jaccard index) here is known beforehand and does not depend on the particular sets to compare, as any schedule is constrained to provide one and just one decision each time period. In this case, the union always coincides with the maximum intersection area. Note that if a particular problem needs to allow a "no decision" possibility, one would only need to add an extra decision variable to the problem  $(y_D)$  in this example which would mean not making any of the other decisions.

Note that the intersection operation, represented by the element-wise operator  $\min\{\cdot\}$  in (5), is not linear. As will be stated later, the SI needs to be computed as part of the optimization problems, so this nonlinearity would become an issue for usual linear scheduling formulations.

**Lemma 1.** The SI (5) can be lower bounded using the set of linear inequalities (7) and (8) together with slack variables  $s \ge 0$ .

$$SI \ge \sum_{t \in \mathcal{T}_1} \frac{\|s_t\|_1}{t_R \Delta - 2\sum_{\tau=1}^{\Delta} \tau \frac{\Delta - \tau}{\Delta}}$$
 (7)

$$s_t \le y_{te} + \sum_{\tau=1}^{\Delta} \frac{\Delta - \tau}{\Delta} \left( y_{(t-\tau)e}|_{t>\tau} + y_{(t+\tau)e}|_{t+\tau \le t_R} \right) \forall e \in \mathcal{E}, \forall t \in \mathcal{T}_1$$
 (8)

*Proof.* On the one hand, by inequalities (8),  $s_t$  are always less than:

$$\min_{e \in \mathcal{E}} \left\{ y_{te} + \sum_{\tau=1}^{\Delta} \frac{\Delta - \tau}{\Delta} (y_{(t-\tau)e}|_{t>\tau} + y_{(t+\tau)e}|_{t+\tau \le t_R}) \right\}$$

On the other hand, as the decisions  $y_{te}$  are binary and  $s_t$  are nonnegative, the 1-norm in both (5) and (7) just performs the sum of its elements. Furthermore,  $s_t$  are always bounded by the maximum possible intersection area:

$$s_t \le 1 + 2\sum_{\tau=1}^{\Delta} \frac{\Delta - \tau}{\Delta}$$

Then, summing up for all  $t \in \mathcal{T}_1$ , excluding the tails that lie outside from the extreme periods, one gets (7).

For the above defined bound to be tight, the slack variables need to be added to the problem objective function for maximization.

# 2.1. SI computation example

Consider a two-stage scheduling problem with three discrete decision variables  $y_A$ ,  $y_B$ , and  $y_C$  to be computed over five time periods (in the first stage). The uncertain parameters are discretized in three uncertainty realization scenarios. Table 1 shows a feasible schedule in which the first-stage decisions are equal among the scenarios (non-anticipativity is met).

Table 2 shows the contribution to the Similarity Index at each time period weighing only the following and previous time periods ( $\Delta = 2$ ), that is, the sum of the fuzzified intersection areas of the decisions taken per scenario at each period. For instance, decision  $y_C$  is made at  $t_4$  and  $t_5$ . There is a contribution of 0.5 at  $t_3$  coming from the fuzzified decision made at the following period. At  $t_4$ , there is a contribution of 1 from the decision made at that period and 0.5 from the  $t_5$  decision for a total contribution of 1.5. The same applies for  $t_5$ . Note that the total area is equal to 2 in the inner time periods, while in the extreme

		$t_1$			$t_2$			$t_3$			$t_4$			$t_5$	
	$e_1$	$e_2$	$e_3$												
$y_A$	1	1	1				1	1	1						
$y_B$				1	1	1									
$y_C$										1	1	1	1	1	1

Table 1: Feasible first-stage schedule. The decisions are equal among the scenarios so that the non-anticipation constraints are met.

	$t_1$	$t_2$	$t_3$	$t_4$	$t_5$
$y_A$	1	0.5 + 0.5	1	0.5	0
$y_B$	0.5	1	0.5	0	0
$y_C$	0	0	0.5	1 + 0.5	0.5 + 1
	1.5	2	2	2	1.5

Table 2: Contribution to the SI of a feasible first-stage schedule (intersection area). The decisions are equal among the scenarios so that the non-anticipation constraints are met.

ones  $(t_1, t_5)$  it is equal to 1.5, as the tails of fuzzified areas would lie outside the first-stage horizon. In this case the intersection area among scenarios is always the maximum, as decisions are equal for all  $e \in \mathcal{E}$ .

Equation (9) shows the computation of the SI for the schedule solution from Table 1 using (6).

$$SI = \frac{1.5 + 2 + 2 + 2 + 1.5}{2 \cdot 5 - 1} = \frac{9}{9} = 1 \tag{9}$$

As expected, the Similarity Index is equal to 100% for the schedule in Table 1. However, Table 3 shows an unfeasible schedule solution where some decisions are not the same among scenarios in the first stage.

		$t_1$			$t_2$			$t_3$			$t_4$			$t_5$	
	$e_1$	$e_2$	$e_3$												
$y_A$	1	1	1				1		1						
$y_B$				1	1	1		1							
$y_C$										1	1	1	1	1	1

Table 3: Unfeasible first-stage schedule. The decisions at  $t_3$  are not equal and the non-anticipation constraints are not met.

As the decisions at  $t_3$  are different, the Similarity Index must be less than 100%. Table 4 shows the intersection area at each time period, which is used in (10) to compute the SI. None of the decisions made at  $t_3$  can contribute to the SI as they are non equal among the scenarios. The only contributions at  $t_3$  come from the decisions made at  $t_2$  and  $t_4$ . Note that the contribution to the SI is computed for each variable at each time period.

$$SI = \frac{1.5 + 1.5 + 1 + 1.5 + 1.5}{2 \cdot 5 - 1} = \frac{7}{9} = 0.778 \tag{10}$$

	$t_1$	$t_2$	$t_3$	$t_4$	$t_5$
$y_A$	1	0.5	0	0	0
$y_B$	0.5	1	0.5	0	0
$y_C$	0	0	0.5	1 + 0.5	0.5 + 1
	1.5	1.5	1	1.5	1.5

Table 4: Contribution to the SI of an unfeasible first-stage schedule (intersection area). Some decisions do not meet the non-anticipativity constraints.

The unfeasible schedules from scenarios in Table 3 are then only 77.8% percent similar.

# 3. Proposed Decomposition Algorithm

As mentioned in Section 1, removing the non-anticipativity constraints from two-stage stochastic scheduling problems allows decomposing the problem into smaller and easier-to-solve subproblems. However, the subproblem solutions would not coincide in the first-stage stage as nothing is enforcing non-anticipativity anymore. The core idea of our decomposition method is to solve the subproblems independently and use the Similarity Index to compare their solutions. An iterative procedure is set up to progressively increase the Similarity Index until complete similarity is reached, hence, meeting the non-anticipation criteria.

However, the SI formula shown in (5) is nonlinear and consequently cannot be used in the usual MILP formulations found in scheduling problems. Then, a local bound for the Similarity Index is computed for each subproblem using inequalities (7) and (8). Anyway, each subproblem requires information from the others to estimate the Similarity Index, and the decomposition would no longer be possible. The PHA and the Dual Decomposition solve this issue by using the solution obtained from each subproblem in the previous iteration. In our case, this is not possible as the inequalities (8) would bound the slack variables  $s_t$  to the worst contribution of all scenarios to the Similarity Index in each time period  $s_t$ . Eventually, the optimizer would be unable to increase the Similarity beyond a certain value. For instance, at time  $t_2$  a scenario j could bound  $s_{t_2}$  to 0 as its solution is too different. Then, at time  $t_3$ , a scenario k could also bound  $s_{t_3}$  to zero. In the end, the optimizer would not be able to adjust the corresponding scenario solution to locally increase the similarity.

To solve this issue, our proposal is that subproblems only compare their decision variables (and thus measuring similarity) to a single reference set of decisions, denoted as  $\bar{y}$ . This reference is intended to be the solution of the subproblem in a previous iteration that is less similar to the rest. With this in mind, the  $|\mathcal{E}|$  subproblems derived from the monolithic problem (2), are formulated as in (11). Note that this is equally applicable to problems in the state-space form (3).

$$\underset{y_e, z_e, r_e, s}{\text{minimize}} \quad p_e \left( \mathbf{c}^T y_e + \mathbf{q}^T \begin{bmatrix} z_e \\ r_e \end{bmatrix} \right) - \lambda \sum_{t \in \mathcal{T}_t} \frac{\|s_t\|_1}{t_R \Delta - 2 \sum_{\tau=1}^{\Delta} \tau \frac{\Delta - \tau}{\Delta}} \tag{11a}$$

subject to

$$Ay_e \ge b \tag{11b}$$

$$T_e y_e + W_e z_e + R_e r_e \ge h_e \tag{11c}$$

$$s_t \le \bar{y}_t + \sum_{\tau=1}^{\Delta} \frac{\Delta - \tau}{\Delta} \left( \bar{y}_{(t-\tau)}|_{t>\tau} + \bar{y}_{(t+\tau)}|_{t+\tau \le t_R} \right) \qquad , \forall t \in \mathcal{T}_1$$
 (11d)

$$s_t \le y_{te} + \sum_{\tau=1}^{\Delta} \frac{\Delta - \tau}{\Delta} \left( y_{(t-\tau)e}|_{t>\tau} + y_{(t+\tau)e}|_{t+\tau \le t_R} \right) \quad , \forall t \in \mathcal{T}_1$$
 (11e)

$$y_e := \{ y_{te} \in \{0, 1\}^{n_y} | t \in \mathcal{T}_1 \} \tag{11f}$$

$$z_e := \{ z_{te} \in \{0, 1\}^{n_z} | t \in \mathcal{T}_2 \}$$
 (11g)

$$r_e := \{ r_{te} \in \mathbb{R}^{n_s} | t \in \mathcal{T}_2 \} \tag{11h}$$

$$s := \{ s_t \in \mathbb{R}^{n_y} | t \in \mathcal{T}_1 \} \tag{11i}$$

The reference values  $\bar{y}_t \in \{0,1\}^{n_y}$  that appear in (11d) refer to the solution in the previous iteration of the scenario sub-problem that yielded the worst *local* Similarity Index. The parameter  $\lambda$  weights the importance of maximizing such index: a value close to zero would mean that the original objective function is prioritized over maximizing the similarity; however, if the Similarity Index is not maximized, it is highly probable that the obtained solutions do not meet the non-anticipation criteria in the end. Contrarily, a large value of  $\lambda$  would give more importance to maximizing the similarity, at the cost of getting to a worse solution in the terms of optimality. Therefore, inspired by the sub-gradient method [22], we propose to update the weighing parameter  $\lambda$  as in (12).

$$\lambda_{k+1} = \lambda_k - \alpha \left( SI_k - 1 \right) \tag{12}$$

Where k represents the current iteration,  $\alpha$  is a tuning parameter that represents the step size, and  $SI_k$  is the exact global Similarity Index computed a posteriori using (5) in each iteration from the subproblems solutions.

Algorithm 1 provides the pseudo-code of the decomposition method. In the algorithm,  $J_e$  stands for the cost of scenario e, that also represents the contribution of each scenario solution to the overall cost according to the probability distribution, i.e.:

$$J_e := p_e \left( c^T y_e + q^T \begin{bmatrix} z_e \\ r_e \end{bmatrix} \right)$$
 (13)

The algorithm begins by initializing the weighing parameter  $\lambda$  and the iteration counter k to zero, and an empty set  $\bar{e}$ . Then, the reference values  $\bar{y}$  are initially set to the solution of the deterministic problem consisting in formulation (1) restricted to the first stage, i.e., just considering variables and constraints

# Algorithm 1 Similarity Index Decomposition

```
Require: \alpha, k_{max}, \Delta
  1: k \leftarrow 0, \lambda_0 \leftarrow 0, \bar{e} \leftarrow \emptyset
                                                                                                         ▶ Initialization
  2: \bar{y} \leftarrow \arg\min c^T y, s.t.: t \in \mathcal{T}_1
                                                               \triangleright Solve first-stage deterministic problem
  3: repeat
            parallel for e in \mathcal{E}/\bar{e} do
  4:
                 y_e^*, z_e^*, r_e^* \leftarrow \arg\min_{y_e, z_e, r_e, s} J_e - \lambda_k SI_e
                                                                                                              ⊳ Solve (11)
  5:
            end parallel for
  6:
           \ell \leftarrow \arg\min_{e \in \mathcal{E}/\ell} SI_e(y_e^*, \bar{y})
                                                                            ▷ Scenario(s) with worst local SI
  7:
           \bar{e} \leftarrow \arg\max_{e \in \ell} J_e(y_e^*, z_e^*, r_e^*)
                                                              ▷ Scenario with highest-cost contribution
  8:
           \bar{y} \leftarrow y_{\bar{e}}^*
  9:
           SI \leftarrow SI(y_{e \in \mathcal{E}}^*)
10:
                                                                                         ▶ Global SI computation
            \lambda_{k+1} \leftarrow \lambda_k - \alpha \cdot (\mathrm{SI} - 1)
11:
            k \leftarrow k + 1
12:
13: until SI = 1 \lor k = k_{max}
                                                                                                ▷ Convergence check
14: if SI=1 then return y_{te}^*, z_{te}^*, r_{te}^*
15: else No feasible solution found
16: end if
```

for  $t \in \mathcal{T}_1$ . Note that this is a fast-to-obtain initial guess required to compute local similarity indexes  $SI_e$  in the first iteration. Then, the subproblems (11) can be solved in parallel and their solutions are collected (line 4). Afterwards (line 7), the algorithm seeks for the subset of scenarios  $\ell \in \mathcal{E}$  that showed the lowest  $SI_e$  value (computed a posteriori by (5) with just the solution  $y_e^*$  of each subproblem and the reference  $\bar{y}$ ). Subsequently, the scenario from those in  $\ell$  which is contributing most to the overall cost function is selected (line 8) and used to set the new reference  $\bar{y}$  (line 9). The global SI is computed (line 7) and used to update the weighing parameter  $\lambda$  (line 9) by rule (12). The process repeats until full similarity (convergence) or the maximum number of iterations  $k_{max}$  are reached (line 13). If convergence is reached, the algorithm output is the first-stage and recourse decisions.

Remark. Note that, in any iteration k>0, the scenario subproblem  $\bar{e}$  corresponding to the current reference  $\bar{y}$  does not need to be solved in the next iteration. This is because  $\lambda_{k+1}>\lambda_k$  so, evidently, the optimal solution for this subproblem is again  $y_{\bar{e}}^*=\bar{y}$  that makes  $SI_{\bar{e}}=1$ . This is of utmost importance with limited CPU resources, as it frees some threads/cores to be directly assigned to other subproblems. Note also that for k>0, the computations benefit from the native warm-start feature of the MILP solvers and extra time is saved by initializing subproblems at their previous solutions.

# 3.1. Convergence & optimality

The procedure that Algorithm 1 describes seeks to progressively push the scenario different solutions together to reach 100% similarity in the first stage.

In brief, the procedure uses the local  $SI_e$  to select the scenario whose solution is less similar to the reference one, and so on. Informally, this is, the first-stage decisions for the "reference scenario" are frozen and the other scenario solutions are pushed closer to them. Then, at next iteration, such reference is changed to the scenario whose solution in the first stage remained further away from the reference. In case there are multiple scenario solutions at "equally far distance", the one that contributes most to the overall cost function is set as an heuristic for the reference choice. Note that this concept of distance between two discrete-decision schedules (a measure of how "far apart" they are) can be formally stated by its  $symmetric\ difference$ , a.k.a. disjunctive union [7].

**Lemma 2.** Given two vectors  $y_1$  and  $y_2$  of dimension n over the binary numbers and the Similarity Index defined in (5), its symmetric difference

$$\delta(y_1, y_2) := 1 - SI(y_1, y_2) \tag{14}$$

is a proper measure of the distance between them.

*Proof.* The SI is a real-valued function  $\wp : \{0,1\}^n \to \mathbb{R}_+$  upper bounded by 1 by definition, so it is  $\delta$  too. In addition, the following properties hold:

- 1.  $\delta(y,y) = 0 \ \forall y \in \{0,1\}^n$
- 2.  $\delta(y_1, y_2) = \delta(y_2, y_1) \ \forall y_1, y_2 \in \{0, 1\}^n$
- 3.  $\delta(y_1, y_3) \le \delta(y_1, y_2) + \delta(y_2, y_3) \ \forall y_1, y_2, y_3 \in \{0, 1\}^n$

Property 3 (triangle inequality) is proven from the property of the inverses in a Boolean group [20].  $\Box$ 

Accordingly,  $\delta = 0$  means 100% similarity whilst  $\delta = 1$  means no similarity at all.

**Proposition 1.** Assume that a feasible solution for problem (1), equivalently (2), exists. Then:

- The solution of (11) with λ = 0 also exists for each scenario e ∈ ε. Thus, cost (2a) computed from such solutions states a lower bound on the optimal cost (1a).
- 2. Any solution set obtained from independent sub-problems (11) with  $\lambda > 0$ , fulfilling SI = 1 is an upper bound on the optimal cost (1a).

*Proof.* Problem (11) stated for each scenario with  $\lambda = 0$  is equivalent to (2) without non-anticipativity constraints (2d). That is, a relaxation of (1), which proves statement 1. Statement 2 is evident, as any solution set fulfilling all constraints in (2) is a feasible solution for (1).

**Theorem.** If a feasible solution for (2) exists, Algorithm 1 with  $\alpha > 0$  reaches it in a finite number of iterations. Moreover, with  $\alpha$  large enough it reaches a feasible solution in  $|\mathcal{E}| + 1$  iterations at most.

*Proof.* Consider the set of solutions  $y_{e\in\mathcal{E}}^*$  provided by Algorithm 1 at any iteration k. Denote by  $\Theta$  to the sum of all peer-to-peer distances between solutions  $y_e^*$ . Let  $\bar{y}=y_{\bar{e}}^*$  be the solution of the selected reference scenario  $\bar{e}$ . Then,

$$\begin{split} \Theta := \sum_{i=1}^{|\mathcal{E}|} \sum_{j>i}^{|\mathcal{E}|} \delta(y_i^*, y_j^*) &= \sum_{\substack{i,j \in \mathcal{E}/\bar{e} \\ j>i}} \delta(y_i^*, y_j^*) + \sum_{e=1}^{|\mathcal{E}|} \delta(y_e^*, \bar{y}) \\ &\leq \sum_{\substack{i,j \in \mathcal{E}/\bar{e} \\ i>i}} \left( \delta(y_i^*, \bar{y}) + \delta(y_j^*, \bar{y}) \right) + \sum_{e=1}^{|\mathcal{E}|} \delta(y_e^*, \bar{y}) \end{split}$$

by  $\delta(\cdot,\cdot)$  satisfying the triangle inequality (Lemma 2). Algorithm 1 progressively maximizes each  $SI_e(y_e^*,\bar{y})$  through  $\lambda_k$ , thus minimizing  $\delta(y_e^*,\bar{y}) \, \forall e \in \mathcal{E}$  and, in particular, the largest distance by setting  $\bar{y}$  to the scenario with worst local SI. Furthermore, as  $\lambda_k$  increases monotonically with k, Algorithm 1 performs a contractive process. Therefore,  $\Theta_k \leq \Theta_{k-1}$  holds. Consequently, two situations can occur: either  $\Theta$  converges to a positive number or zero. In the first case, if  $\Theta$  converges to a positive number, there is no feasible solution which contradicts the main assumption that a feasible solution exists.

Hence, if a feasible solution for (2) exists, eventually  $\Theta$  reaches zero which, in turn, means the global SI=1 as  $\Theta \leq |\mathcal{E}| (1 - SI(y_{e \in \mathcal{E}}^*))$  by definition (recall the min operator in (5)).

Now, let  $\alpha >> 0$  be large enough such that  $\lambda_k \geq \lambda^* \ \forall k$ , being  $\lambda^*$  the optimal value<sup>1</sup> of the Lagrange multiplier in problem  $(11)^2$  to fulfill:

$$\sum_{t \in \mathcal{T}_1} \frac{\|s_t\|_1}{t_R \Delta - 2\sum_{\tau=1}^{\Delta} \tau^{\frac{\Delta - \tau}{\Delta}}} = 1$$

In this case, problem (11) focuses on fulfilling the equality above. All solutions  $y_{e\in\mathcal{E}}$  are forced to fully meet the reference  $\bar{y}$  (Lemma 1) from k>0. In this setup, if any  $y_{\mathbf{e}}^* \neq \bar{y}$  after solving the sub-problems, it is because  $\bar{y}$  is infeasible for that sub-problem  $\mathbf{e}$ . Consequently, Algorithm 1 sets  $\bar{y}=y_{\mathbf{e}}^*$  for k+1 and forces the remaining  $y_{e\in\mathcal{E}}$  to meet that reference. In the worst case, Algorithm 1 has tested all vertices  $y_{e\in\mathcal{E}}^*$  as reference in  $k=|\mathcal{E}|+1$  iterations, so a feasible solution must have been reached.

Of course, reaching a global feasible solution does not guarantee optimality, as any feasible solution different from the optimal is just an upper bound, see Proposition 1.

**Corollary.** The optimal solution for (2) lies in the convex hull formed by the  $|\mathcal{E}|$  vertices of optimal solutions  $y_{e\in\mathcal{E}}^*$  computed from (11) with  $\lambda = 0$ .

<sup>&</sup>lt;sup>1</sup>Karush-Kuhn-Tucker (KKT) necessary optimality conditions.

<sup>&</sup>lt;sup>2</sup>The duality statement and KKT conditions are extended to problems where some of the variables are integer and the objective and constraints are convex [1].

*Proof.* These scenario optimal solutions  $y_{e\in\mathcal{E}}^*$  define a lower bound on the cost (Proposition 1). Indeed, this bound is the lowest possible to be computed from independent scenario solutions, otherwise would mean that the solution for any scenario is suboptimal. This also means that there cannot exist a different set of solutions  $y_{e\in\mathcal{E}}$  yielding better overall cost with higher peer-to-peer distance sum  $\Theta$ . Consequently, as (2) is a linear programming problem, its optimal solution  $y^*$  must lie in the convex hull defined by  $y_{e\in\mathcal{E}}^*$  obtained with  $\lambda=0$ .

## 3.2. Parameter tuning

Algorithm 1 has mainly two tunable parameters: the step size  $\alpha \in \mathbb{R}$  and the length of the fuzzification horizon  $\Delta \in \{2,3,4,\ldots\}$ . Intuitively, step size  $\alpha$  balances solution optimality with convergence speed: a large value of  $\alpha$  implies that more importance is given to maximizing the Similarity Index at the cost of possibly reaching a suboptimal solution. On the contrary, small  $\alpha$  provides more flexibility for the subproblems to progressively adapt their schedules towards the feasible optimum, but many iterations might be required to converge.

The length of the fuzzification horizon  $\Delta$  affects the granularity of the SI function. Although  $SI \in \mathbb{R}$ , by definition, it can only take a finite number of values. Thus, larger  $\Delta$  increases the number of possible values the SI can take. There is no clear insight on what is the influence of this parameter in the solution quality for any general case. On the one hand, large  $\Delta$  implies that, for each decision that is not coincident in a given period t among the scenarios, more neighboring periods get their SI reduced, thus reducing the overall SI value. This means that finding the right path to improve the SI at first iterations is harder for the optimizer, because the SI sensitivity is lower. On the other hand, the smallest  $\Delta$  may affect SI sensitivity in a similar way if a decision is scheduled very close in time among scenarios but just outside the fuzzification horizon. For instance, if certain constraints in a scenario impede to take as first decision (at  $t_1$ ) the one that is considered optimal for the rest of scenarios (e.g. a maintenance task), and then it is taken with a delay of just  $\Delta$  periods, the schedules could be exactly the same but slightly delayed in time. Clearly some similarity exists in this hypothetical case. However, the overall SI will be zero as only the immediate neighboring period is considered towards similarity.

From this analysis one can infer that the problem definition itself is which could determine the optimal length of the fuzzification horizon: same decision taken with a difference of  $\Delta$  periods may be seen as relatively similar if the scheduling horizon (more precisely,  $|\mathcal{T}_1|$ ) is large, or dissimilar if the opposite. Hence, for problems where the first stage involves many periods,  $\Delta$  shall be increased from the default value 2 (by which only periods t-1 and t+1 are weighted towards similarity) to keep a desired ratio  $\Delta/|\mathcal{T}_1|$ .

#### 4. Case study

The performance of the Similarity Index decomposition was tested on an industrial-like case study that uses a discrete-time formulation. Palacín et al.

[17] studied the evaporation network of a cellulose fibers production plant. Some chemicals (henceforth products) are used in the spinning process to provide the fibers with some desired mechanical properties. Such products have a significant economic value and must be reused. After passing through the main process, the diluted products are sent to an evaporation network where they are concentrated. The network consists of several evaporation plants that consist of a series of heat exchangers, evaporation chambers, condensers, and cooling systems. Palacín et al. [17] used a surrogate model to estimate the operating costs of each evaporation plant that depends on the efficiency of its cooling system, the nominal efficiency of the evaporation plant, the increased costs due to fouling, and the product load.

The goal of the optimization problem is to minimize the operating costs of the evaporation network over a finite horizon denoted by discrete time periods  $t \in \mathcal{T}$ . The evaporation plants  $v \in \mathcal{V}$  can only process a product  $p \in \mathcal{P}$  at a time. Unfortunately, the plants need to be regularly cleaned as fouling decreases their efficiency increasing steam consumption and thus the operating costs. The continuous fouling effect is discretized over a set of consecutive states  $\{s_1, \ldots, s_{40}\} \in \mathcal{S}$  with decreasing plant efficiency. The optimizer needs to decide when to stop the plants to perform either a deep (A) or a light (B) cleaning, which are denoted by states  $s_A, s_B \subseteq \mathcal{S}$ . The plants can also be put on stand-by either before or after a cleaning task. Time is discretized in one-day long periods as that is what usually cleaning the plants takes.

The optimization problem seeks to optimally schedule the evaporation plants and their maintenance (cleaning). Production is modelled by non-negative real variables  $P_{vtp}$ . Product assignment and plants states are decided using binary variables  $A_{vtp}$  and  $E_{vts}$  respectively.

The model accounts for several constraints including:

- Evaporation plants can only be in one and only one state at each time period.
- Evaporation plants can only process one product at a time.
- Due to personnel limitations, only one cleaning task can occur per time period.
- After an evaporation plant is cleaned, it can only be cleaned again after several time periods in operation.
- An evaporation plant in operation can stay in such a state, go to stand by, or get cleaned. A fouled plant on standby can only remain on standby or be cleaned. A clean plant can remain on standby or start operating.
- The evaporation plants cannot change products unless a cleaning task is performed.

Uncertainty is very important for the adequate operation of the network. The product demand  $D_p$  can vary over the prediction horizon depending on

the market and the main process. Additionally, the maximum load of the evaporation plants depends on the ambient air temperature. The model is then formulated as a two-stage stochastic scheduling problem with scenarios  $e \in \mathcal{E}$ . This additional index e is added to the decision variables, together with non-anticipation constraints. For a detailed description of the model and its equations, refer to [17].

The SI formula needs to be updated to the case study nomenclature and additional sets. The SI will only be calculated with respect to the evaporation plant state variables  $E_{vtse}$  (binary). Due to the problem particular structure, fixing  $E_{vtse}$  implies unique values for product allocation  $(A_{vtpe})$  and plant load  $(P_{vtpe})$  decisions<sup>3</sup>. Of course, this would not be the case in general, but the reader may note that the SI-decomposition algorithm can trivially combine with PHA for instance, to handle continuous variables appearing in the first stage. Details omitted for brevity.

Hence, the formula to compute the overall SI in this case, shown in (15), only accounts for  $E_{vtse}$ . The corresponding formula for the local SI  $(SI_e)$  is omitted, as it can be easily derived from (15).

$$SI = \sum_{v \in \mathcal{V}} \sum_{s \in \mathcal{S}} \sum_{t \in \mathcal{T}_1} \frac{\left\| \min_{e \in \mathcal{E}} \left\{ E_{vtse} + 0.5 E_{v(t-1)se} + 0.5 E_{v(t+1)se} \right\} \right\|_1}{n_v \left( 2t_R - 1 \right)}$$
(15)

The main apparent difference of (15) with respect to the formula (6) in the SI definition is that there are two extra summations according to the sets defining the binary variables: the evaporation plants  $\mathcal{V}$  and the operation state  $\mathcal{S}$ . Note that the original denominator is multiplied by the number of evaporators  $n_v$ , but not by the number of states, as the evaporators can only be at one state at a time.

The PHA was used to assess the performance of the SI decomposition. Our implementation of the PHA is shown in Algorithm 2. Note that no update rule was performed for the parameter  $\rho$ , so it remains constant for every iteration.

#### 5. Benchmark results & discussion

This section presents a comparison between the Similarity Index Decomposition, the Progressive Hedging Algorithm and the Monolithic approach for solving four instances of the case study but with increasing size. Additionally, the effect of varying the main parameters on the SI Decomposition and PHA is analyzed. The four problem instances are defined by:

Instance A: 3 plants, 2 products, 4 scenarios and 25 days horizon. 21869 variables.

<sup>&</sup>lt;sup>3</sup>Plants current state (at  $t_0$ ) is not a decision variable. So, once  $E_{vtse}$  is fixed equal for all  $e \in \mathcal{E}$ , constraints on product changeover and cleaning prevent the existence of different allocations among scenarios

# Algorithm 2 Progressive Hedging Algorithm

```
Require: \rho, k_{max}
 1: k \leftarrow 0, \lambda_{vste}^0 \leftarrow 0
                                                                                                                    ▶ Initialization
  2: for e in \mathcal{E} do
                                                                                                                ▶ Decomposition
  3:
             E_{vste}^* \leftarrow \arg\min_{E_{vste}} J_e
  4: end for
  5: repeat
             k \leftarrow k + 1
  6:
             \widehat{E}_{vst_R} = \frac{1}{|\mathcal{E}|} \sum_{e \in \mathcal{E}} E_{vst_R e}^*
                                                                                                                     ▶ Aggregation
  7:
             \lambda_{vst_{R}e}^{k} \leftarrow \lambda_{vst_{R}e}^{k-1} + \rho \left( E_{vst_{R}e} - \widehat{E}_{vst_{R}} \right)
                                                                                                                   ▷ Price update
  8:
            E_{vste}^* \leftarrow \arg\min_{E_{vste}} J_e + \lambda_{vst_Re}^k \cdot E_{vst_Re} + \frac{1}{2}\rho ||E_{vst_Re} - \hat{E}_{vst_R}||^2 end for
  9:
                                                                                                                ▷ Decomposition
10:
11:
12: until ||E^*_{vst_Re} - \hat{E}_{vst_R}|| = 0 \lor k = \mathbf{k}_{max}
                                                                                                                    ⊳ Stop criteria
13: return E_{vste}^*
```

- Instance B: 9 plants, 3 products, 16 scenarios and 25 days horizon. 187202 variables.
- Instance C: 9 plants, 3 products, 16 scenarios and 30 days horizon. 224642 variables.
- Instance D: 23 plants, 5 products, 16 scenarios and 30 days horizon. 618242 variables.

For the sake of comparison, the robust horizon  $\mathcal{T}_1$  was set to 7 days for all the instances. It shall be noted that approximately 90% of the variables are binary. The modeling environment GAMS 42.1.0 together with Gurobi 10.0 as MIP solver were used for implementation. The monolithic and SI decomposition instances resulted in MILPs, while the PHA ones lead to MIQPs.

# 5.1. Performance assessment

Table 5 presents a comparison of the cost objective value and computation time of the different approaches in all the case study instances. The stop criterion for the optimizer Gurobi was the optimality gap in the decomposition approaches whilst a time limit of 10 days of computation is set for the monolithic approach. The step sizes  $\rho$ ,  $\alpha$ , for the PHA and SI decomposition respectively, were tuned to obtain the minimum cost (objective function) at the lowest possible computation time.  $\Delta=2$  was set for the SI decomposition.

In Instance A, both decomposition methods were considerably slower than their monolithic counterpart. The reported objective values were identical. However, this instance is apparently too small to truly realize the benefits of decomposition. Instance B showed significant time reductions for both decomposition approaches, two orders of magnitude faster than the monolithic approach. Note that this is even stopping the time count in the monolithic resolution at

Instance	Approach	Step size	Cost	Time (s)	Iter.	Gap (%)
	Monolithic	-	10051.6	8	-	0
A	PHA	2.5	10051.6	27	5	0
	SI-D	460	10051.6	24	3	0
	Monolithic	-	91819.6	1305	-	0.49
В	PHA	45	91843.05	37	4	0
	SI-D	700	91819.6	40	3	0
	Monolithic	-	108698.4	48569	-	0.97
$\mathbf{C}$	PHA	45	108698.4	417	4	0
	SI-D	1500	108730.4	565	3	0
	Monolithic	-	527617	864000	-	> 1
D	PHA	45	527298	12737	6	0.01
	SI-D	2500	527037	15470	4	0.01

Table 5: Comparison on the four problem instances solved by the monolithic approach, the PH Algorithm, and the SI Decomposition.

0.49% gap, when the finally proven (after 4412s) optimal value was first found. In this instance, while SI Decomposition reached the optimal cost value, the PHA reported a slightly worse cost. For instance C, both decomposition approaches greatly outperformed the monolithic approach in terms of computation time, again despite of stopping the clock of the monolithic resolution when the optimal solution is first found, almost at 1% gap (to reduce such gap to 0, Gurobi spent 176.5h in total). In this case, PHA slightly outperformed SI Decomposition for the chosen setups. Finally, in the largest instance D, the SI Decomposition only needed 4 iterations to converge to a solution whose cost is better than the one obtained by PHA after comparable computation time, and much better than what the monolithic approach is able to provide after ten days.

For completeness, Figure 3 graphically shows the progress of Algorithm 1 with the iterations. Note that, although some distances  $\delta$  can eventually increase after switching the reference solution,  $\Theta$  always decreases towards zero with the iterations.

All tests and parallel computations were performed on a Dell Precision T5500 workstation with two Intel Xeon X5650 CPUs (24 available logical processors in total), setting three threads in Gurobi to solve each subproblem.

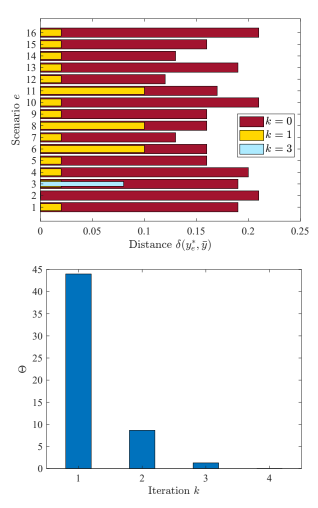


Figure 3: Computed distances through the optimization process of Instance D with SI-D. Solutions  $y_e^* = 2$ , 8 and 3 were chosen as reference in iterations k = 1, 2 and 3, respectively.

# 5.2. Effect of the fuzzification horizon

As discussed in Section 3.2, the length of the fuzzification horizon (in which similarity is considered among variables in different scenarios) is determined by the parameter  $\Delta$ . Hence,  $\Delta=2$  means that similarity is considered only within the previous and following period. For  $\Delta=3$ , two previous and two following days are taken into account, and so on.

To test the influence of  $\Delta$  in the SI Decomposition, all problem instances were solved for  $\Delta \in \{1, 2, 3, 4\}$  so that the number of surrounding days to weight in the SI is between 2 and 7. The objective values and computation times are shown in Table 6.

The parameter  $\Delta$  did not significantly impact the cost value and computation times in the small problem instances. However, some differences can be observed

Instance	$\Delta$	Objective	Iter.	Time (s)
	1	10051.6	3	28
Α	2	10051.6	3	24
$\mathbf{A}$	3	10051.6	3	27
	4	10051.6	3	28
	1	91819.6	3	38
В	2	91819.6	3	40
Б	3	91819.6	3	45
	4	91819.6	3	43
	1	108698.4	4	630
$\mathbf{C}$	2	108730.4	3	565
C	3	108730.4	3	726
	4	108730.4	3	620
	1	527375.6	3	13347
D	2	527037.1	4	15470
D	3	527375.3	5	21332
	4	527375.3	5	21660

Table 6: Effect of  $\Delta$  on the SI Decomposition algorithm.

both in optimality and computation time for the larger instances, especially instance D. On instance C,  $\Delta=2$  provided the shortest computation time by approximately 10%, but yielded higher cost than with  $\Delta=1$ . Effects look more appreciable on instance D, where computation times monotonically increase with  $\Delta$ , but note that the best cost value is obtained with  $\Delta=2$ .

These numerical tests seem to confirm that the effect of  $\Delta$  on the decomposition method is minor and might depend on the problem structure and size. As a tuning guideline, the modeler may set  $\Delta$  to low values and only increase it to explore if there is an extra improvement to unlock in problems where  $|\mathcal{T}_1|$  is large, i.e., fine discretized robust horizons.

#### 5.3. Effect of the step size in the SI decomposition

Instances A and D were solved for different values of  $\alpha$ . Figure 4 shows the computation time and the iterations required to converge in such instances for the tested values. For this evaluation, the problem objective functions were not scaled in neither case.

The algorithm converged regardless of the value of  $\alpha$ , as expected from theory. Small values of  $\alpha$  lead to many iterations and, consequently, to larger computation times. For large values of  $\alpha$ , although the computation time was shorter in both instances, the resulting solutions deviated slightly from optimality: the gap became 0.2% for  $\alpha >= 500$  in Instance A, whilst an increase of 0.4% for  $\alpha >= 5000$  was reported in Instance D. Values in the ranges of  $100 < \alpha < 500$  in Instance A and of  $1000 < \alpha < 5000$  in Instance D are the ones which reported the

best performance if optimality is prioritized. For larger  $\alpha$  values the computation times continue decreasing, but tend to converge to the minimum number of iterations possible. Consequently, the impact of varying  $\alpha$  within relatively large values is not significant.

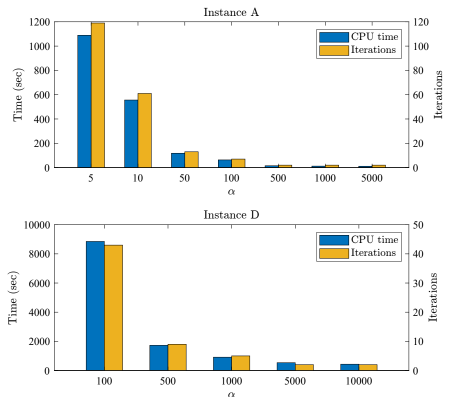


Figure 4: Performance evaluation with  $\alpha$  in SI decomposition.

As a suggestion, the algorithm shall be tuned starting with a large value of  $\alpha$  and then decrease it until the computation time increases significantly (i.e. exceeds what is considered acceptable for a particular application). Then, the value for which the best objective is obtained at the lowest CPU time would be the choice. The opposite path is not recommended, as the algorithm would need many iterations to converge in the first trials.

#### 5.4. Effect of the step size in the PHA

Although the step sizes of the PHA and our algorithm are not directly comparable because of their different meaning in the formulations, the performance of the PHA was evaluated as well using different values of  $\rho$ . Figure 5 shows the computation time and the required number of iterations in Instance A for different values of  $\rho$ .

The PHA converged to the optimal solution for  $\rho \in [0.1, 3)$ . As seen in figure, the computation time tends to decrease as  $\rho$  increases, until reaching a limit

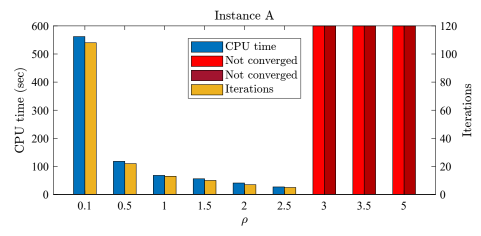


Figure 5: Performance evaluation with  $\rho$  in PHA.

value where the algorithm did not converge to any feasible solution. Outside the provided interval, the error criterion  $||E^*_{vst_Re} - \widehat{E}_{vst_R}||$  gets stuck in a positive value after some iterations. This issue was amplified in instances B, C and D where, despite the different problem sizes and initial conditions, convergence was only achieved in a small range around  $\rho = 45$ .

An explanation for this unwanted behavior is in the discontinuity of the feasible region due to binary decision variables [19]. What probably happens in this case is that, when a few scenario optimal solutions diverge from the rest and cannot approach the reference (mean) values  $\hat{E}_{vst_R}$  for some reason (either a specific constraint or because multiplier  $\lambda$  already became too large), all scenario solutions keep invariant. And this can happen for too large or too small values of  $\rho$ , as scheduling problems often involve numerous lower cost operational decisions. Besides, oscillation phenomena can also occur if  $\lambda$  grows too fast [27]. Consequently, in a complex setting, tuning  $\rho$  is a non-directed trial-and-error process for finding a value that just gets convergence, becoming tougher if best performance-optimality tradeoff is sought.

# 6. Conclusions

The Similarity Index replaces the non-anticipativity constraints of two-stage stochastic scheduling problems. It allows decomposing the monolithic problem into smaller subproblems on a scenario basis. The proposed method significantly reduced the required computation time to obtain a solution in realistic-size instances of the presented case study, without sacrificing the actual optimization objective. The results gathered indicate that such speed up would enable considering uncertainty in near real-time plant scheduling. Note that the largest instance is similar in size to the industrial facility the case study is inspired on.

Comparing to the existing alternatives in the literature, such as the Progressive Hedging Algorithm, this new method is advantageous in the sense that the

formulation is more compact (non-anticipativity in all first-stage variables is merged into a single indicator) and simpler to tune: the modeler can balance between optimality and convergence speed by just varying the step size  $\alpha$ . In contrast, for the PHA to reach optimal (even feasible) solutions at lower times, a tedious trial-and-error process is required to tune its step size  $\rho$ .

Although the proposed SI decomposition algorithm could arrive at least at the same objective value as the monolithic approach in all tested instances, in other cases, large values of the weighing parameter  $\alpha$  can result in an optimality gap. Evidently, the presented analysis about the effect of the tuning parameters on the algorithm performance is preliminary. It is necessary to test the method on more two-stage scheduling problems of different nature in order to draw general conclusions.

Nonetheless, there are also some drawbacks. It is well known that arriving to a proven optimal solution is an extremely time consuming and computationally intensive task in large-scale mixed-integer problems. Therefore, high quality solutions that can be obtained in reasonable amounts of time are usually preferred in the scheduling practice. The monolithic formulation naturally provides this feature by setting a time limit or an optimality gap (gap between the incumbent solution and the theoretically best possible) as termination criteria in the MILP solver. However, the SI Decomposition cannot cope with neither of such criteria: due to its nature, there is no guarantee of having a feasible solution until the algorithm has converged to SI=1, and the number of needed iterations for this is not known beforehand. Nevertheless, and opposed to other decomposition algorithms in the literature, convergence can be forced in a finite number of iterations if  $\alpha$  is large enough and a feasible solution exists, of course.

The reader may note that decomposition algorithms can be speed up by not solving the subproblems to proven optimality (zero gap). However, this does not poses any bound on the actual optimality gap once a solution for the *global* problem is reached. Furthermore, relaxing the local optimality gap criterion too much causes oscillations in decomposition algorithms and hampers convergence.

The proposed decomposition algorithm always converged to a feasible and near-optimal solution in the tested case study, no matter of the values set for the tuning parameters. Future work will focus on extending the method to also handle continuous decision variables in a formal way, as well as on tackling continuous-time scheduling formulations.

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