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An Integrated Learning and Progressive Hedging Matheuristic for Stochastic Network Design Problem¹

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Abstract. We address the Multicommodity Capacitated Fixed-charge Network Design problem with uncertain demands, which we formulate as a two-stage stochastic program. We rely on the progressive hedging (PH) algorithm of Rockafellar and Wets where the subproblems are defined using scenario groups. To address the problem, we propose an efficient matheuristic approach which we refer to as the Integrated Learning and Progressive Hedging. The proposed method takes advantage of a specialized learningbased matheuristic that is able to quickly produce high-quality solutions to multi-scenario subproblems. Furthermore, we propose a novel reference point definition, at each aggregation step of the PH algorithm, which leverages subproblem information regarding promising design variables. Extensive computational experiments illustrate that the proposed approach should be the method of choice when high-quality solutions to large instances of stochastic network problems need to be found quickly.

Keywords: Network design, uncertain multicommodity demand, two-stage formulation, progressive hedging method, learning-based matheuristic.

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1 Introduction

Multicommodity Capacitated Fixed-charge Network Design ((MCND, Crainic et al., 2021a) models are used to address many important planning problems in a variety of applications, including transportation, logistics, and telecommunications (e.g., Magnanti and Wong, 1984; Crainic, 2000; Crainic et al., 2021c). MCND models formulate the requirement to design a network, i.e., select among a set of potential arcs a subset of arcs, and associated capacity, to satisfy, at minimum total cost, the demand for transportation of a set of commodities between their respective origin-destination node pairs. Selecting an arc incurs a so-called *fixed cost*, while a *unit transportation cost* is associated to each commodity and arc. The goal is thus to find an optimal design that minimizes the sum of the fixed and transportation costs.

Uncertainty often characterizes the problems addressed. Demand, the volume of each commodity in particular, generally involves a certain level of uncertainty in network design problems. Notably, strategic and tactical planning for complex systems such as the ones mentioned above, yield plans defining the structure of the system or the service, respectively, which are then used repeatedly over medium to long periods of time. Some form of forecast then needs to be built to represent the future demand and its variation when the plan is built and the network is designed. Yet, it is well known that solving a deterministic model using single-point estimates of the uncertain parameters, may yield arbitrarily bad solutions (Wallace, 2000; Higle and Wallace, 2003). Models explicitly accounting for uncertainty are then recommended. Stochastic programming and, in particular, two-stage formulations of the MCND with stochastic demand, SMCND in the following, has become the methodology of choice to account for demand uncertainty when building such plans. This *a-priori* optimization approach (Birge and Louveaux, 2011)) mimics the planning decision process, the first stage design decisions being made prior to the realization of demand, while the second stage decides on the best utilization of the first-stage design (possibly, adjusted marginally) to satisfy the revealed demand. The goal is thus to design a network that remains feasible and cost effective when different demand realizations are encountered.

Demand uncertainty is very often represented through a finite set of discrete scenarios, suitably generated to collectively approximate the uncertainty present in the problem setting, and their associated probabilities. For network design, the design arcs are selected in the first stage, while flow-distribution decisions for each scenario belong to the second stage, resulting in a mixed integer formulation with an objective function minimizing the sum of the fixed costs of the selected design arcs and the expectation of the cost of the flow distribution computed over the scenarios considered. This formulation is quite difficult to address using exact methods as, on the one hand, the MCND is a difficult combinatorial problem even when uncertainty is not considered and, on the other hand, using scenarios to model uncertainty yields a large-scale model. Heuristic approaches are thus attractive to produce good-quality solutions within reasonable computing efforts.

The previously proposed methods for stochastic MCND problems have shown great promise in terms of their ability to successfully address the models, but they also have their own limits (e.g., their computational scalability, when they are used to address instances that involve a larger number of scenarios). This is specifically the case for the progressive hedging-based methods, initially proposed by Rockafellar and Wets (1991), which traditionally apply scenario decomposition to separate the stochastic MCND models into a series of single scenario subproblems. The main advantage of these approaches is that each of the obtained subproblems can then be more efficiently addressed (i.e., possibly leveraging algorithms that were proposed for the deterministic MCND variants). However, considering that each subproblem may produce a solution that is tailored to the specific scenario involved, there might be a large number of discrepancies between the solutions of all the scenario subproblems. The stochastic model seeks to find a compromise (i.e., a single consensus solution) that is efficient with respect to the expected cost function. Thus, a progressive and gradual search for such a consensus solution is applied by these methods, i.e., a search process that does not force agreement too early and seeks to identify the solution characteristics that properly hedge against the random changes that may affect the values of the stochastic parameters. When a large number of scenarios are involved and there are large discrepancies in the scenario solutions at the beginning of the solution process, the search for consensus might require a significant amount of effort (thus defining a computational bottleneck for the solution method). Therefore, in the present paper, we propose a series of novel strategies to efficiently apply progressive hedging for stochastic network design models that involve a large scenario set.

We thus propose the Integrated Learning and Progressive Hedging (ILPH) solution method, a new PH-based matheuristic, which is able to handle large scale instances of stochastic network design problems efficiently. ILPH integrates a generalized Learn&Optimize matheuristic (Sarayloo et al., 2021a), which addresses efficiently multi-scenario network design problems, producing rapidly high-quality solutions. The Learn&Optimize procedure provides ILPH with learning capabilities regarding promising design variables gathered during the search paths taken to find high-value solutions to each multi-scenario subproblem. Moreover, we introduce a new reference point in the aggregation step of the proposed ILPH by exploiting the information garnered from subproblems, and using this information to update the PH parameters. This contrasts with the traditional reference-point definition which is based exclusively on the subproblem designs available at each iteration, ignoring all possible information on the evolution of the subproblem designs. Consequently, the ILPH is governed and guided by the local information provided by the learning procedure, resulting in an integrated approach that accelerates the search for network characteristics that better hedge against the random demand changes and is thus capable of handling both larger and more challenging instances to address.

Extensive computational experiments show the efficiency of these methodological contributions, the proposed ILPH matheuristic producing a more efficient overall consensus search, both in terms of overall solution quality and total computational effort, compared to the literature. They thus indicate that ILPH should be the method of choice when high-quality solutions to large instances of stochastic network problems need to be found quickly.

The rest of the paper is organized as follows. We recall the two-stage formulation for the stochastic network design problem in Section 2. We briefly review some relevant literature in Section 3. We then introduce the main ideas and a detailed description of our solution methodology in Section 4. Finally, we present and analyze the experimental results in Section 5 and provide concluding remarks in Section 6.

2 SMCND Formulation

We briefly recall the two-stage stochastic formulation of the MCND with uncertain demand, where the first-stage decisions concern the network configuration selecting the design arcs, while the commodity flows are determined at the second stage, given the first-stage design and the realized random demands. Uncertainty is modeled through a set of scenarios. The formulation, detailed in Crainic et al. (2011), minimizes the sum of the total cost of designing the network and the expected cost of servicing the possible demand expressed through the considered scenarios. The following notation is used:

- \mathcal{N} : Set of nodes $i \in |\mathcal{N}|$.
- $\mathcal{N}^+(i)$: Outward neighbors of node i (i.e., $\{j : (i, j) \in \mathcal{A}\}$).
- $\mathcal{N}^{-}(i)$: Inward neighbors of node i (i.e., $\{j : (j,i) \in \mathcal{A}\}$).
- \mathcal{A} : Set of potential arcs $(i, j) \in \mathcal{A}$.
- f_{ij} : Fixed cost of arc $(i, j) \in \mathcal{A}$ if included in the final design.
- u_{ij} : Capacity on arc $(i, j) \in \mathcal{A}$.
- \mathcal{K} : Set of commodities $k \in \mathcal{K}$ with origin and destination nodes o(k), s(k).
- c_{ij}^k : Unit routing cost for commodity $k \in \mathcal{K}$ on arc $(i, j) \in \mathcal{A}$.
- \mathcal{S} : Set of scenarios $s \in \mathcal{S}$ with probabilities $p^1, \ldots, p^{|\mathcal{S}|}$.
- d_i^{ks} : Volume of commodity $k \in \mathcal{K}$ at node $i \in \mathcal{A}$ in scenario $s \in \mathcal{S}$.
- y_{ij} : Binary design variable, equal to 1 if arc $(i, j) \in \mathcal{A}$ is included in the network at the first stage.

• x_{ij}^{ks} : Continuous flow variable representing the amount of commodity k on arc $(i, j) \in \mathcal{A}$ in scenario $s \in \mathcal{S}$.

The SMCND formulation becomes

Minimize
$$\sum_{(i,j)\in\mathcal{A}} f_{ij}y_{ij} + \sum_{s\in\mathcal{S}} p^s \sum_{k\in\mathcal{K}} \sum_{(i,j)\in\mathcal{A}} c_{ij}x_{ij}^{ks}$$
(1)

Subject to
$$\sum_{i \in \mathcal{N}^+(i)} x_{ij}^{ks} - \sum_{i \in \mathcal{N}^-(i)} x_{ji}^{ks} = d_i^{ks}, \qquad \forall i \in \mathcal{N}, \ \forall k \in \mathcal{K}, \ \forall s \in \mathcal{S},$$
(2)

$$\sum_{k \in \mathcal{K}} x_{ij}^{ks} \le u_{ij} y_{ij}, \qquad \forall (i,j) \in \mathcal{A}, \ \forall s \in \mathcal{S}, \quad (3)$$

$$y_{ij} \in \{0, 1\}, \qquad \forall (i, j) \in \mathcal{A}, \quad (4)$$

$$x_{ij}^{ks} \ge 0,$$
 $\forall (i,j) \in \mathcal{A}, \ \forall k \in \mathcal{K}, \ \forall s \in \mathcal{S}.$ (5)

The objective function (1) minimizes the total system cost, i.e., the sum of the fixed cost of the selected design arcs and the expectation of the routing costs taken over all the demand scenarios. Constraints (2) represent the flow conservation equations in each scenario, requiring that the volume d^{ks} of demand k in scenario s be moved from its origin $(d_i^{ks} = d^{ks}, \text{ for } i = o(k))$ to its destination $(d_i^{ks} = -d^{ks}, \text{ for } i = s(k))$ and $d_i^{ks} = 0$ for all other nodes, $\forall i \in \mathcal{N}$ such that $i \notin o(k)$ and $i \notin s(k)$. Constraints (3) enforce the linking relation between design and utilization, restricting flows to the capacity of selected arcs only. Notice that the definition of a single design decision variable y together with constraints (3) guarantee the respect of the non-anticipativity conditions requiring a single design network for all considered demand realization (design decisions must not be tailored to the observed demands of particular scenarios). Constraints (4) - (5) impose integrality and non-negativity restrictions on decision variables.

The SMCND model (1)-(5) is a large-scale mixed integer program with a blockdiagonal structure, each block being defined by constraints (2) - (3). We present the solution method we propose following a brief literature review.

3 Literature review

When surveying the literature, one finds a limited number of methodologies to address stochastic network design problems. As mentioned earlier, when a finite set of scenarios is used to estimate the stochastic parameters (see Dupačová et al. (2000) and King and Wallace (2012) for an overview on scenario generation methods), a stochastic program can be formulated as an equivalent (multi-scenario) deterministic model. Considering the large scale of the resulting model, taking advantage of the structure through decomposition-based approaches is especially beneficial and is the focus of much of the algorithmic work in this area. The goal of decomposition-based approaches is to divide the complex problem into subproblems to be able to address them more efficiently. In the case of stochastic programs, such decomposition strategies can be categorized into two types. The first type decomposes the problem via decisional stages while the second type decomposes by scenarios. The former category (referred to as the L-shape method introduced in Van Slyke and Wets (1969)) is a cutting-plane method that is based on the application of Benders decomposition to the equivalent (multi-scenario) deterministic model. Detailed reviews on this type of decomposition approach for stochastic MCND may be found in Rahmaniani et al. (2017, 2018); Hewitt et al. (2021); Crainic et al. (2021b).

In the second category of decomposition strategies, referred to as scenario decomposition, the original problem is decomposed by scenarios by applying an augmented Lagrange relaxation strategy to the non-anticipativity constraints (i.e., the constraints ensuring that a single design is used under all considered scenarios). Once the problem is decomposed, each scenario becomes a deterministic problem (i.e., a single-scenario subproblem (SSP) defined for each scenario). The resulting scenario subproblems can then be used in various ways when solving stochastic network design problems. First, they enable a general lower bound to be obtained for the stochastic model by solving the Lagrange dual, as proposed in Schütz et al. (2009). In turn, such lower bounds can be applied to design exact solution methods for stochastic network design models, e.g., Escudero et al. (2012) and Alonso-Ayuso et al. (2003). Furthermore, considering that each subproblem takes the form of scenario-specific deterministic model, this decomposition approach also enables heuristics to be defined that leverage existing solution methods for deterministic network design models, e.g., the progressive hedging-based metaheuristics developed by Crainic et al. (2011). However, in the follow up study of Crainic et al. (2014), it was shown that defining the subproblems using multiple scenarios (i.e., multi-scenario subproblems (MSSP)) clearly improved the computational performance of this type of progressive hedging-based solution method. When applying progressive hedging-based methods, one thus may leverage efficient metaheuristics that are available for deterministic network design models (in the case of single scenario-specific subproblems), or, for SMCND models to address the subproblems (in the case of MSSP).

Although the literature on efficient metaheuristic methods proposed for deterministic MCND problems is very rich (e.g., (e.g., Crainic et al., 2000; Ghamlouche et al., 2003, 2004; Hewitt et al., 2010; Crainic and Gendreau, 2021), there are only limited contributions on efficient heuristic methods for solving the SMCND problem. For example, Sarayloo et al. (2021a) proposed a learning based matheuristic approach where the main novelty is a learning heuristic able to effectively identify structures of good-quality solutions when the scenarios and their influences on design decisions are gradually considered.

The proposed matheuristic produces information on promising design variables related to the considered scenarios. As previously stated, we believe that this can be efficiently used in the PH algorithm to gather more refined and useful information regarding the solutions to the subproblems that will improve the search for a good overall solution to the SMCND.

Exploiting common solution structures that exist between deterministic and stochastic solutions is another feature that may be employed in the solution methods based on scenario decomposition. Due to the high complexity and difficulty of stochastic network design problems, a number of attempts in the literature on stochastic network design have been devoted to investigate how the solution to the deterministic model relates to the stochastic counterpart. It has been shown that, despite the fact that often times solutions to deterministic models behave badly in stochastic settings (Wallace, 2000; Higle and Wallace, 2003), there are situations where the deterministic solutions share some properties with the corresponding stochastic solutions (Lium et al., 2009; Thapalia et al., 2011, 2012; Crainic et al., 2018). In these studies, the authors show that the deterministic solutions carry useful information (i.e., some structural patterns) which can be extracted to simplify the stochastic case. Following this insight, Sarayloo et al. (2021b) proposed a number of strategies to extract reduced cost information from good quality solutions to be used as a guide to fix the variables in the SMCND problems.

Revisiting the PH algorithm, as previously stated, the algorithm enables available solution methods for the deterministic MCND to be directly applied to address the stochastic version of the problem. There are a number of challenges to applying the PH idea to the SMCND. First, the size of the problem and the number of iterations needed to reach consensus grow with the number of scenarios, while addressing each subproblem means tackling a difficult MCND problem. As shown by Crainic et al. (2014), the number of iterations can be reduced by grouping scenarios to build multi-scenario subproblems. This has the benefit of forcing a level of local consensus, but makes the subproblems harder to address. Second, the traditional reference-point definition is based exclusively on the subproblem designs available at each iteration, ignoring all possible information on the evolution of the subproblem designs. Therefore, in the next section, we propose an integrated progressive hedging solution method which is able to 1) quickly produce high quality solutions for multi-scenario subproblems and 2) extract more refined information from the subproblems and produce a novel reference point leading to a more efficient overall consensus search.

4 Solution methodology

We propose a matheuristic that combines and enhances the Lagrangian-based multiscenario decomposition strategy of progressive hedging and the information extraction and learning capabilities of Learn&Optimize. We cast the SMCND model (1)-(5) in a form suitable for multi-scenario decomposition and provide the outline of the proposed ILPH method in Section 4.1. Sections 4.2 and 4.3 describe the methodological developments and strategies proposed for each step of ILPH. The former describes the Learn&Optimize matheuristic generalized to address multi-scenario subproblems and gather relevant information while doing so. We then explain, in Section 4.3, how we exploit that information to compute a new reference point and update the Lagrange multipliers and penalties modifying the fixed costs used in the subproblems and, thus, guiding the search toward consensus. We provide the detailed pseudocode of ILPH in Section 4.4.

4.1 Outline of the proposed ILPH

Let $\{C_1, \ldots, C_{|G|}\}$ be a partition of the scenario set S, where G is the set of group indices and each group $C_g \subset S$, $\forall g \in G$. Let $p^g = \sum_{s \in C_g} p^s$. Then, in order to apply the PH idea, we make explicit the non-anticipativity restrictions. Define the first-stage variables y_{ij}^g for each scenario group $C_g, \forall g \in G$. Let \bar{y}_{ij} be the *reference point* standing for the design decision that is feasible in all scenarios and is the goal of the formulation:

Minimize
$$\sum_{g \in G} p^g \Big(\sum_{(i,j) \in \mathcal{A}} f_{ij} y_{ij}^g + \sum_{s \in C_g} \frac{p^s}{p^g} \sum_{k \in \mathcal{K}} \sum_{(i,j) \in \mathcal{A}} c_{ij}^k x_{ij}^{ks} \Big)$$
(6)

Subject to $\sum_{j \in \mathcal{N}^+}$

$$\sum_{(i)} x_{ij}^{ks} - \sum_{j \in \mathcal{N}^{-}(i)} x_{ji}^{ks} = d_i^{ks}, \ \forall i \in \mathcal{N}, \ \forall k \in \mathcal{K}, \forall s \in C_g, \forall g \in G,$$
(7)

$$\sum_{k \in \mathcal{K}} x_{ij}^{ks} \le u_{ij} y_{ij}^g, \ \forall (i,j) \in \mathcal{A}, \forall s \in C_g, \forall g \in G,$$
(8)

$$y_{ij}^g = \bar{y}_{ij}, \ \forall (i,j) \in \mathcal{A}, \ \forall g \in G,$$

$$(9)$$

$$y_{ij}^g \in \{0,1\}, \ \forall (i,j) \in \mathcal{A}, \ \forall g \in G,$$

$$(10)$$

$$x_{ij}^{ks} \ge 0, \ \forall (i,j) \in \mathcal{A}, \ \forall k \in \mathcal{K}, \forall s \in C_g, \forall g \in G,$$

$$(11)$$

where the non-anticipativity constraints (9) guarantee that design decisions are not tailored for each particular scenario and aim to yield a single implementable design. The augmented Lagrangian-relaxation of these constraints then yields the objective function

$$\operatorname{Minimize} \sum_{g \in G} p^{g} \left(\sum_{(i,j) \in \mathcal{A}} f_{ij} y_{ij}^{g} + \sum_{s \in C_{g}} \frac{p^{s}}{p^{g}} \sum_{k \in \mathcal{K}} \sum_{(i,j) \in \mathcal{A}} c_{ij}^{k} x_{ij}^{ks} + \sum_{(i,j) \in \mathcal{A}} \lambda_{ij}^{g} (y_{ij}^{g} - \bar{y}_{ij}) + \frac{\rho}{2} (y_{ij}^{g} - \bar{y}_{ij})^{2} \right), \quad (12)$$

where the Lagrange multipliers λ_{ij}^g , $\forall (i, j) \in \mathcal{A}, \forall g \in G$, are associated with the relaxed constraints (9) and ρ is a penalty ratio. Given the binary requirements for the design

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variables, after rearranging the terms, the function may be reformulated as

$$\text{Minimize} \sum_{g \in G} p^g \left(\sum_{(i,j) \in \mathcal{A}} (f_{ij} + \lambda_{ij}^g - \rho \bar{y}_{ij} + \frac{\rho}{2}) y_{ij}^g + \sum_{s \in C_g} \frac{p^s}{p^g} \sum_{k \in \mathcal{K}} \sum_{(i,j) \in \mathcal{A}} c_{ij}^k x_{ij}^{ks} \right) \\
- \sum_{(i,j) \in \mathcal{A}} \lambda_{ij}^g \bar{y}_{ij} + \sum_{(i,j) \in \mathcal{A}} \frac{\rho}{2} \bar{y}_{ij} \qquad (13)$$

Model (13), ((7)-(11)) decomposes along the scenario groups when the overall design is fixed to the reference point. One thus obtains a series of subproblems, each taking the form of a SMCND problem formulated using the scenarios of the corresponding group $g \in G$ with fixed costs defined as $f_{ij} + \lambda_{ij}^g - \rho \bar{y}_{ij} + \frac{\rho}{2}$, $\forall (i, j) \in \mathcal{A}$. The subproblem SP_g associated to group $g \in G$ can then be expressed as:

$$SP_g: \text{Minimize} \qquad \sum_{(i,j)\in\mathcal{A}} \left(f_{ij} + \lambda_{ij}^g - \rho \bar{y}_{ij} + \frac{\rho}{2} \right) y_{ij}^g + \sum_{s\in C_g} \frac{p^s}{p^g} \sum_{k\in\mathcal{K}} \sum_{(i,j)\in\mathcal{A}} c_{ij}^k x_{ij}^{ks} \tag{14}$$

Subject to

$$\sum_{j \in \mathcal{N}^+(i)} x_{ij}^{ks} - \sum_{j \in \mathcal{N}^-(i)} x_{ji}^{ks} = d_i^{ks}, \ \forall i \in \mathcal{N}, \ \forall k \in \mathcal{K}, \ \forall s \in C_g,$$
(15)

$$\sum_{k \in \mathcal{K}} x_{ij}^{ks} \le u_{ij} y_{ij}, \ \forall (i,j) \in \mathcal{A}, \ \forall s \in C_g,$$
(16)

$$y_{ij}^g \in \{0,1\}, \ \forall (i,j) \in \mathcal{A},\tag{17}$$

$$x_{ij}^{ks} \ge 0, \ \forall (i,j) \in \mathcal{A}, \ \forall k \in \mathcal{K}, \ \forall s \in C_g.$$
 (18)

In the following we first describe the general steps of the PH algorithm applied on the SMCND with single scenario subproblems and then discuss the outline of our proposed method (with multi-scenario subproblems). As originally introduced in Rockafellar and Wets (1991), the PH solution method first decomposes the stochastic problem by scenarios by applying an augmented Lagrange relaxation strategy. When applied to stochastic network design, this enables the problem to be separated into a series of subproblems, each defined as a deterministic network design model using a specific scenario (Crainic et al. (2011)). A solution to the original stochastic model can then be obtained through an iterative process that applies the following three algorithmic steps: 1) Each scenario subproblem is solved separately, thus producing a specific (and possibly different) network. 2) A reference point, indicating the level of consensus among the subproblem solutions is obtained by calculating the weighted average overall scenario networks. 3) The subproblem formulations are then adjusted to promote solution consensus with respect to the current reference point. This is specifically done by adjusting, for each scenario subproblem, the values of the fixed cost associated with the arcs to penalize the differences that are present between the current scenario networks and the reference point.

There are a number of challenges to apply the PH approach to solve the SMCND. Among these challenges there is the process to reach consensus that may require a large number of iterations, where a series of hard deterministic combinatorial optimization problems need to be solved each time. As shown by Crainic et al. (2014), the number of iterations can be reduced by grouping the scenarios in the subproblems, thus forcing a level of local consensus (i.e., obtained within each subproblem). However, this is done at the expense of producing multi-scenario subproblems, which are even more challenging to address. To meet this general challenge, our proposed ILPH method applies an efficient solution method that is able to quickly produce high-quality solutions for multi-scenario network design models at the subproblem level.

Furthermore, in the original PH method, the reference point is obtained by calculating the weighted average overall subproblem networks. By proceeding in this way, one effectively ignores all information that is gathered in the process of solving the subproblems. We are of the opinion that this is an important shortcoming of the PH method when it is applied to stochastic combinatorial optimization problems. Specifically, when multi-scenario subproblems are considered, there may be important information obtained during the search paths that are taken to find high-value solutions for the subproblems that may be helpful in searching for a good consensus. We therefore propose a novel reference point definition which leverages subproblem information regarding promising design variables by exploiting efficient learning mechanisms.

Algorithm 1 displays the outline of our proposed ILPH method. We generalize the Learn&Optimize matheuristic (Sarayloo et al., 2021a) and use it as an efficient method to address the multi-scenario subproblems at each iteration of the PH search (Line 4). We exploit the knowledge learned through the Learn&Optimize procedure in computing the new value of the reference point (Line 5) and in updating the penalties used to modify the fixed costs in the subproblems (Line 6). We thus integrate the new information learned while working on the subproblem formulations to the progressive-hedging search strategy, for increased efficiency. This integrated methodology is our main contribution in this paper. Given the fact that the PH algorithm applied to integer problems does generally not converge to a single solution, the proposed matheuristic proceeds to the second phase when a consensus solution is not obtained once the stopping criterion is met. Phase II (Line 9) follows the general canvas of the literature (Crainic et al., 2011): 1) identify a set of design variables for which the level of consensus is sufficiently high; 2) Fix the values of these variables in the original stochastic model; 3) address the resulting restricted formulation using a commercial MIP solver.

Algorithm 1 The outline of the proposed ILPH	
1: Initialization	
2: Phase I - Multi-scenario PH Search	
3: while Stopping criteria not met do	
4: Solve heuristically the multi-scenario subproblems	\triangleright Section 4.2
5: Aggregation - Compute new reference point	\triangleright Section 4.3
6: Penalty update	\triangleright Section 4.3
7: end while	
8: Phase II - Fix and Solve	

9: Fix the design variables for which consensus is obtained and solve the restricted problem

4.2 Addressing subproblems with the *Learn&Optimize* matheuristic

One of the most challenging parts of the PH algorithm is addressing the multi-scenario stochastic network design problem SP_g^{ν} : (14)-(18) defining the subproblems at iteration ν . As defined, the subproblems appear as large-scaled integer programs, which are hard to solve to optimality. This being said, as evidenced by Kall and Wallace (1994), we may not need to obtain optimal solutions to the subproblems. Heuristics may find satisfactory subproblem solutions that produce an efficient search process for the PH algorithm (Løkketangen and Woodruff, 1996).

We generalize the Learn&Optimize procedure (Sarayloo et al., 2021a) to address the subproblems of the PH algorithm. The Learn&Optimize provides a novel systematic learning mechanism capable of identifying what are the efficient solution structures for the overall stochastic problem by analyzing the solutions obtained for subproblems. In the context of network design, this information takes the form of design decisions that are common to high-quality (i.e., optimal or near-optimal) solutions obtained by gradually considering scenarios and their interactions. Two important motivations made us apply the Learn&Optimize in this PH-base matheuristic. First, it provides an efficient heuristic to address the subproblems. Second, we aim to exploit the knowledge learned through the Learn&Optimize procedure in the aggregation step of the PH algorithm to further improve its performance. Towards this end, we inject the local information regarding what network characteristics (i.e., arcs to either include or exclude) work "best" for each multi-scenario subproblem that is gathered by Learn&Optimize method into the overall PH search procedure.

The Learn&Optimize procedure iteratively executes a learning step, to learn and build statistics on solution characteristics. The learning mechanism embedded in this method gradually builds an image of design variables that potentially belong to good solutions for the original stochastic formulation. The results of this learning mechanism are then

used to guide the overall search toward higher-quality solutions. Sections 4.2.1 and 4.2.2 provide the full description of the generalized *Learn&Optimize* procedure.

4.2.1 Learning step

The authors introduce two main concepts in the Learn&Optimize procedure, the Artificial Demand Scenario (ADS), which is defined as a particular combination of two scenarios, and the Artificial Recourse Problem (ARP), which is a network flow problem (Sarayloo et al., 2021a). The main idea behind the ARP is to send the commodity flows associated with the demand vector of ADS, on a given design, and identify the required corrective actions, taking the form of extra arcs, that are needed to satisfy ADS demands. One then learns by iteratively defining ADSs, solving the associated ARPs (which constitutes the core of our learning mechanism), and gradually building an image of design variables that potentially belong to good solutions for the stochastic formulation. The results of this learning mechanism are then used to guide the overall search towards higher quality solutions.

Considering the fact that our goal is to apply the heuristic procedure repeatedly to the subproblems of the PH algorithm which contain multiple scenarios, we generalize the idea of ADS and propose a generalized *Learn&Optimize*. To do so, we introduce *Group-based Artificial Demand Scenario*, *Gb-ADS*, with the associated ARP.

A Group-based Artificial Demand Scenario Gb-ADS δ^{g} , $g \in G$, given the scenarios in group g, $\{s_1^g, \ldots, s_{|g|}^g\}$, is defined as

$$\boldsymbol{\delta}^{g} = \begin{pmatrix} \delta_{1}^{g}(s_{1}^{g}, \dots, s_{|g|}^{g}) \\ \delta_{2}^{g}(s_{1}^{g}, \dots, s_{|g|}^{g}) \\ \delta_{3}^{g}(s_{1}^{g}, \dots, s_{|g|}^{g}) \\ \delta_{3}^{g}(s_{1}^{g}, \dots, s_{|g|}^{g}) \end{pmatrix}, \text{ such that} \\ \vdots \\ \delta_{|\mathcal{K}|}^{g}(s_{1}^{g}, \dots, s_{|g|}^{g}) \end{pmatrix}$$

Let Δ^g be a set of Gb-ADSs, δ^g , generated for group g containing the scenarios $C_g = \{s_1^g, \ldots, s_{|g|}^g\}$. We use algorithm 2 to construct the set Δ^g of cardinality N_{Δ^g} , through the random selection of the demand values of the scenarios in C_g . In other words, we randomly select $s_i \in C_g$, copy its demand value associated to commodity k, $d_k(s_i)$, and let $\delta_k^g \leftarrow d_k(s_i)$ (Lines 2-4). The procedure stops when it builds N_{Δ^g} Gb-ADSs (Line 6).

Following Sarayloo et al. (2021a), we aim to explore the solution characteristics as-

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Algorithm 2 Construct Δ^g

1: repeat 2: for all $k \in \mathcal{K}$ do 3: Randomly choose $s_i \in C_g$ and let $\delta_k^g \leftarrow d_k(s_i)$ 4: end for 5: Let $\Delta^g \leftarrow \delta^g \bigcup \Delta^g$ 6: until $|\Delta^g| = N_{\Delta^g}$ 7: Return Δ^g

sociated to each Gb-ADS, $\delta^g \in \Delta^g$, to extract information regarding promising design variables. The exploration is performed by solving an *Artificial Recourse Problem*, $ARP(\delta^g, \hat{y})$, for each artificial demand scenario $\delta^g \in \Delta^g$ considering a given design \hat{y} .

To define $ARP_g(\delta^g, \hat{y})$, we separate the set of arcs \mathcal{A} according to \hat{y} . Then, $\mathcal{A} = \mathcal{A}^0 \cup \mathcal{A}^1$, where $\mathcal{A}^0 = \{(i, j) | (i, j) \in \mathcal{A}, \hat{y}_{ij} = 0\}$ and $\mathcal{A}^1 = \{(i, j) | (i, j) \in \mathcal{A}, \hat{y}_{ij} = 1\}$ are the sets of closed and open arcs in \hat{y} , respectively. By considering that the fixed cost f_{ij}^{ν} is updated at each iteration of the PH algorithm, we then define a modified arc variable cost \bar{c}_{ij} by linearizing the fixed cost of the closed arcs

$$\bar{c}_{ij} = \begin{cases} c_{ij} + \frac{f_{ij}^{\nu}}{u_{ij}}, & \forall (i,j) \in \mathcal{A}^0, \\ c_{ij}, & \forall (i,j) \in \mathcal{A}^1, \end{cases}$$
(19)

and solve the $ARP_q(\boldsymbol{\delta}^g, \hat{y})$, which is defined as a multi-commodity network flow problem

$$ARP_g(\boldsymbol{\delta}^g, \hat{y})$$
: Minimize $\sum_{k \in \mathcal{K}} \sum_{(i,j) \in \mathcal{A}} \bar{c}_{ij} x_{ij}^k$ (20)

Subject to

$$\sum_{\mathcal{N}^+(i)} x_{ij}^k - \sum_{i \in \mathcal{N}^-(i)} x_{ji}^k = \delta_k^{ig}, \quad \forall i \in \mathcal{N}, \ \forall k \in \mathcal{K},$$
(21)

$$\sum_{k \in \mathcal{K}} x_{ij}^k \leq u_{ij}, \qquad \forall (i,j) \in \mathcal{A}, \qquad (22)$$

$$x_{ij}^k \ge 0,$$
 $\forall (i,j) \in \mathcal{A}, \ \forall k \in \mathcal{K}.$ (23)

Solving $ARP_g(\delta^g, \hat{y})$ yields $x_{ij}(\delta^g) = \sum_{k \in \mathcal{K}} x_{ij}^k, \forall (i, j) \in A$, with $\mathcal{A}_{\delta^g} = \{(i, j) | x_{ij}(\delta^g) > 0\}$. We then define a corresponding design solution as $y_{ij}^{\delta^g} = 1$, when $x_{ij}(\delta^g) > 0$, and 0, otherwise. It is noteworthy that some of the arcs in \mathcal{A}^0 , closed in \hat{y} , may be open in $y_{ij}^{\delta^g}$ to satisfy the demand vector δ^g . These modifications capture the interactions occurring in the integration of multiple scenarios within δ^g , yielding partial information regarding the design arcs required to address the uncertainty captured by the scenarios involved in group g. Repeating this procedure for different Gb-ADSs builds the knowledge we seek.

4.2.2*Learn&Optimize* procedure

The generalized *Learn&Optimize* procedure iteratively executes the *learning step* to identify a promising set of design variables. Then, it performs the *partial-optimization step* where the identified promising variables are fixed and the reduced-size formulation is solved exactly.

The generalized *Learn&Optimize* procedure is described in Algorithm 3. The sets of Gb-ADSs, $\Delta^{g\nu}$, are reconstructed at each iteration ν of PH algorithm as described in Algorithm 2. Such an approach, i.e., reconstructing a new set of $\Delta^{g\nu}$ at each iteration of ILPH, may allow us to obtain more diverse information that would not be available from using a single set of $\Delta^{g\nu}$ in all iterations. We define the *frequency memory*, $F_{ii}^{g\nu}$, representing how often arc (i, j) has been used in the solutions of the different $ARP_{g}(\boldsymbol{\delta}^{g}, \hat{y})$ and the normalized frequencies values, $\mathbf{f}_{ij}^{g\nu}$, which is computed as $\mathbf{f}_{ij}^{g\nu} := F_{ij}^{g\nu} / max\{F_{ij}^{g\nu} | (i, j) \in \mathcal{A}\}$ (Line 10). We keep the frequency memories built in the previous iterations, i.e., $F_{ij}^{g\nu} \leftarrow F_{ij}^{g\nu-1}$ in order to keep track of promising arcs from the beginning. More specifically, we are implementing a long-term memory strategy on the solution information that is learned at the subproblem level (i.e., the arc frequencies) as a means to both warm start the solution method and to further drive solution consensus at the subproblem level. We also define $\mathcal{A}_{\Delta^{g\nu}}^{\nu}$, the set of design arcs used in at least one $ARP_q(\delta^g, \hat{y})$ at iteration ν , and $\mathcal{A}^{g\nu}$, the set of promising design variables to be identified by the procedure.

- Algorithm 3 The generalized Learn&Optimize procedure to address SP_g^{ν} 1: Initialization: $F_{ij}^{g\nu} \leftarrow F_{ij}^{g\nu-1}, \forall (i,j) \in A, \ \mathcal{A}^{g\nu} \leftarrow \emptyset$, construct $\Delta^{g\nu}$, let \hat{y}^{ν} be the current design;
- 2: Learning and memorizing:

3: repeat

- Randomly choose a Gb-ADS $\delta^g \in \Delta^{g\nu}$: 4:
- 5:
- 6:
- Solve $ARP_g(\delta^g, \hat{y}^{\nu})$ yielding $x_{ij}^{\nu}(\delta^g), \forall (i, j) \in \mathcal{A};$ $Identify \ \mathcal{A}_{\delta^g}^{\nu} \text{ and } compute \ y_{ij}^{\delta^g}, \ \forall (i, j) \in \mathcal{A}_{\delta^g}^{\nu};$ $Update \ \mathcal{A}_{\Delta^{g\nu}}^{\nu} := \mathcal{A}_{\Delta^{g\nu}}^{\nu} \bigcup \mathcal{A}_{\delta^g}^{\nu} \text{ and } F_{ij}^{g\nu} := F_{ij}^{g\nu} + 1, \text{ for all } (i, j) \in \mathcal{A}_{\delta^g}^{\nu};$ 7:
- Remove $\boldsymbol{\delta}^{g}$ from $\Delta^{g\nu}$; 8:
- 9: until $\Delta^{g\nu} = \emptyset;$
- 10: Normalize frequencies $\mathbf{f}_{ij}^{g\nu} := F_{ij}^{g\nu}/max\{F_{ij}^{g\nu}|(i,j) \in \mathcal{A}\}, \ \forall (i,j) \in \mathcal{A};$
- 11: for all $(i, j) \in \mathcal{A}$ do 12: if $\mathbf{f}_{ij}^{g\nu} \geq \tau$ then $\mathcal{A}^{g\nu} \leftarrow \mathcal{A}^{g\nu} \cup \{(i, j)\};$
- 13:end if
- 14: end for
- 15: Partial optimization:
- 16: Solve SP_q^{ν} , by fixing variables belonging to $\mathcal{A}^{g\nu}$ to open, yielding solution $y_{ij}^{g\nu}, \forall (ij) \in$
- 17: Return the normalized frequencies $\mathbf{f}_{ij}^{g\nu}, \forall (ij) \in \mathcal{A}$ and solution $y_{ij}^{g\nu}, \forall (ij) \in \mathcal{A}$.

The main loop (Lines 3 to 9) iterates over the Gb-ADSs in $\Delta^{g\nu}$, each being discarded once examined. The loop stops when the set of artificial demand scenarios becomes empty. The $ARP_g(\delta^g, \hat{y})$ is solved for each $\delta^g \in \Delta^{g\nu}$, to distribute the demand of δ^g (Line 5). The corresponding design vector is created (Line 7), while the set of used design arcs and the frequency memories are updated at Line 6. Once the artificial demand scenarios $\delta^g \in \Delta^{g\nu}$ are treated, a reduced problem, obtained by fixing $\mathcal{A}^{g\nu}$ as the most frequently used arcs (given a threshold τ), is solved using a MIP solver. This yields the design solution $y_{ij}^{g\nu}, \forall (ij) \in \mathcal{A}$. Finally, the procedure returns the normalized frequency values $\mathbf{f}_{ij}^{g\nu}, \forall (i, j) \in \mathcal{A}$, as well as the design solution $y_{ij}^{g\nu}, \forall (ij) \in \mathcal{A}$.

4.3 Aggregation and penalty updates

We exploit and inject the local information obtained by the *Learn&Optimize* procedure into the PH algorithm through the definition of a new reference point, Section 4.3.1, and a new heuristic design. The latter, presented in Section 4.3.2, is created by exploiting the dual information associated with the solutions obtained for the subproblems and serves as the initial solution provided for the search processes of the subproblems at the following iteration.

4.3.1 A new reference point

The reference point is a major component of PH matheuristics as it measures the degree of consensus among subproblems and indicates the current trend for opening and closing arcs amongst the subproblem designs. As mentioned earlier, the reference point in previous PH algorithms for SMCND (e.g., Crainic et al., 2011, 2014) is constructed as $\bar{y}_{ij}^{\nu} \leftarrow \sum_{g \in G} p^g y_{ij}^{g\nu}$, $\forall (i, j) \in \mathcal{A}$. The dual prices $\lambda_{ij}^{g\nu}$ are updated, using the reference solution \bar{y}_{ij}^{ν} and the parameter ρ , as $\lambda_{ij}^{g\nu} \leftarrow \lambda_{ij}^{g\nu-1} + \rho(y_{ij}^{g\nu} - \bar{y}_{ij}^{\nu})$. This strategy considers a single design vector only, obtained by each subproblem, to produce the reference point. Considering single solutions only may result in missing alternative cost-efficient subproblem solutions for which consensus could be more easily reached. Our proposed reference point directly targets this issue by explicitly considering the information learned through the use of the *Learn& Optimize* method regarding what generally constitutes good subproblem solutions, i.e., arcs that often appear in high-quality solutions to the subproblems.

Recall that the *Learn&Optimize* procedure creates a history of promising design variables in each subproblem through frequency memories $F_{ij}^{g\nu}$, and $\mathbf{f}_{ij}^{g\nu} \in [0,1], \forall (i,j) \in \mathcal{A}$. The information provided by $\mathbf{f}_{ij}^{g\nu}$ allows the opportunity to better explore the trend amongst the design variables observed during the iterations of the *Learn&Optimize* matheuristic. We propose $\tilde{y}_{ij}^{\nu} \leftarrow \sum_{g \in G} p^g \mathbf{f}_{ij}^{g\nu}, \forall (i,j) \in \mathcal{A}$, where $p^g = \sum_{s \in C_g} p^s$, as a ref-

erence point in the aggregation step of the ILPH. The Lagrangian multipliers $\lambda_{ij}^{g\nu} \forall (i, j) \in \mathcal{A}$ are updated using the new reference point $\tilde{y}_{ij}^{\nu} \forall (i, j) \in \mathcal{A}$ as $\lambda_{ij}^{g\nu} \leftarrow \lambda_{ij}^{g\nu-1} + \rho(\mathbf{f}_{ij}^{g\nu} - \tilde{y}_{ij}^{\nu})$.

4.3.2 A new design solution to iteratively promote consensus

The second idea is to exploit the dual information provided by the subproblem solutions to create an integer design solution at each iteration of the ILPH. This integer design solution $\hat{y}_{ij}^{\nu}, \forall (i, j) \in \mathcal{A}$, is constructed in the aggregation step of the ILPH algorithm and serves as an initial design solution for the generalized *Learn&Optimize* procedure in the subproblems at the following iteration. Exploiting the reduced cost information associated with multiple solutions is motivated by the observation made in Sarayloo et al. (2021b) suggesting that this strategy allows the identification of good-quality solutions in the context of stochastic network design problems.

Note that the proposed design solution does not necessarily define a feasible solution to the original stochastic problem. However, using the same integer initial solution for all subproblems in the Learn & Optimize procedure, defines a soft strategy to promote consensus at the subproblem level.

To create \hat{y}_{ij}^{ν} , we first compute $\tilde{y}_{ij}^{\nu} \leftarrow \sum_{g \in G} p^g f_{ij}^{g\nu}$, $\forall (i, j) \in \mathcal{A}$, and partition the set of design variables into two disjoint subsets:

• $\hat{\mathcal{A}}_1 = \{(i, j) | \tilde{y}_{ij}^{\nu} \leq l_0 \text{ or } \tilde{y}_{ij}^{\nu} \geq u_1\}$: the set of design variables for which a consensus has been almost obtained (given thresholds l_0 and u_1) among the groups, or in other words, (almost) all groups agree that these arcs have to be opened or closed. The design variables in $\hat{\mathcal{A}}_1$ are then set to the value 0 or 1 as follows

$$\hat{y}_{ij}^{\nu} = \begin{cases} 0, \text{ if } \tilde{y}_{ij}^{\nu} \le l_0, \\ 1, \text{ if } \tilde{y}_{ij}^{\nu} \ge u_1. \end{cases}$$
(24)

• $\hat{\mathcal{A}}_2 = \{(i,j)|l_0 < \bar{y}_{ij}^{\nu} < u_1\}$: the set of the remaining design variables or those for which a consensus has not been obtained.

For the variables in $\hat{\mathcal{A}}_2$, the decision is based on reduced cost information. Let $r_{ij}^g, \forall (i, j) \in \hat{\mathcal{A}}_2$, be the reduced cost associated with $y_{ij}^g, \forall (i, j) \in \hat{\mathcal{A}}_2$; $G_{ij}^1 = \{g | y_{ij}^g = 1\}$ the set of groups where the associated design variables y_{ij}^g are equal to one, and $\bar{r}_{ij}^{g\nu} = \sum_{g \in G_{ij}^1} p^g r_{ij}^{g\nu}, (i, j) \in \hat{\mathcal{A}}$ the average reduced cost over groups $g \in G_{ij}^1$. It should be noted that, given the fact that we are solving the restricted problem SP_g^{ν} with the integrality requirements, we need to perform one additional step to obtain the reduced

cost values. Once the restricted problem SP_g^{ν} is solved and its optimal (integer) solution, $y_{ij}^{g\nu}, \forall (ij) \in \mathcal{A}$, is obtained, we need to solve the LP relaxation of the problem SP_g^{ν} while the design variables are fixed to the values of the obtained optimal solution $y_{ij}^{g\nu}, \forall (ij) \in \mathcal{A}$. In this way, one can obtain the set of reduced cost values associated with design variables.

We represent the set of reduced costs by $R = \{\bar{r}_{ij}^{g\nu} | (i, j) \in \hat{\mathcal{A}}_2\}$. In order to identify good candidate design variables to be fixed to 1 (open), we choose the variables with the smallest reduced cost values. To do so, we sort the reduced cost values $\bar{r}_{ij}^{g\nu} \in R$ in non-decreasing order. Let r_1^{max} and r_1^{min} be the maximum and minimum values in R. We then divide the difference $r_1^{max} - r_1^{min}$ in N_1 equally-sized classes and store the indexes of variables belonging to the classes 1 to p_1 ($1 \leq p_1 \leq N_1$) in R_1 . We then set $\hat{y}_{i,j}^{\nu} \leftarrow 1$, $\forall (i,j) \in R_1$ and $\hat{y}_{ij}^{\nu} \leftarrow 0$, $\forall (i,j) \notin R_1$. Consequently, the solution $\hat{y}_{ij}^{\nu} \forall (i,j) \in \mathcal{A}$, at each iteration ν , is created by exploiting the reduced cost information associated with multiple solutions obtained by all considered groups.

4.4 The algorithm

Algorithm 4 sums up the entire procedure. The algorithm is initialized by constructing the list of scenario groups $\bar{C} = \{C_1, \ldots, C_{|G|}\}$. The scenarios within each group may be chosen using different strategies proposed in Crainic et al. (2014). For each group $g \in G$, we address heuristically subproblem SP_g^0 : Minimize $\sum_{(i,j)\in\mathcal{A}} (f_{ij}y_{ij}^g + \sum_{s\in C_g} \frac{p^s}{p^g} \sum_{k\in\mathcal{K}} \sum_{(i,j)\in\mathcal{A}} c_{ij}^k x_{ij}^{ks})$ st. (15) – (18), using the generalized *Learn&Optimize* procedure described in Section 4.2. Once we address the subproblems $g \in G$, we perform the aggregation step to produce the reference point as well as the integer design solution as explained in Section 4.3.

During Phase I, the multi-scenario subproblems are addressd approximately as explained in Section 4.2. At each aggregation step, \tilde{y} and \hat{y} are constructed and the Lagrangian multipliers are updated as described in Section 4.3. We follow the strategy used in Crainic et al. (2011, 2014) to construct a heuristic feasible network $y^{M\nu}$ at each iteration ν , by setting the design variables $y_{ij}^{M\nu}$, $\forall (i, j) \in \mathcal{A}$ to

$$y_{ij}^{M\nu} = \begin{cases} 1, \text{ if } y_{ij}^{g\nu} = 1, \text{ for any } g \in G, \\ 0, \text{ otherwise }. \end{cases}$$
(25)

The best network found, i.e., y^{Best} , is updated based on the quality (total cost) of the feasible solution $y^{M\nu}$ obtained at each iteration ν . We use similar stopping criteria (Line 9) as those in Crainic et al. (2014), namely, a total of N_{Itr} iterations, N_{Imp} consecutive iterations without improving, the best known solution, t_{max} CPU time, or when there are fewer than γ ($0 \le \gamma \le 1$) percent of the arcs for which a consensus has not been reached. When such a situation occurs, *Phase II* is initiated. We fix the design variables for which

a consensus is obtained and solve the original problem to obtain the final design solution y^{Final} . At Line 27, we update y^{best} , when needed.

Algorithm 4 The proposed integrated learning and progressive hedging matheuristic

1: Initialization 2: Let $\nu \leftarrow 0 \ \lambda_{ij}^{g\nu} \leftarrow 0, \forall (i,j) \in \mathcal{A}, \ \rho^{\nu} \leftarrow \rho^{0}$ 3: Construct the list of scenario groups $\bar{C} = \{C_1, \ldots, C_{|G|}\}$ 4: for each group g do Solve SP_q^0 heuristically by performing Algorithm 3 (Section 4.2.2) 5: 6: end for 7: Construct solutions $y_{ij}^{M\nu}$, \tilde{y}_{ij}^{ν} , and \hat{y}_{ij}^{ν} 8: Phase I: Seek consensus on the arcs (i,j) that should exist in the design 9: while stopping criteria not met do **Iteration update:** 10: $\nu \leftarrow \nu + 1$ 11: Solving subproblems heuristically: 12:13:for each group *q* do Solve SP_q^{ν} heuristically by performing Algorithm 3, considering Lagrangian multi-14:pliers $\lambda_{ij}^{g\nu-1}, \forall (i,j) \in \mathcal{A}$ and solution $\hat{y}_{ij}^{\nu-1}, \forall (i,j) \in \mathcal{A}$, to obtain $\mathbf{f}_{ij}^{g\nu}$ and $y_{ij}^{g\nu} \forall (i,j) \in \mathcal{A}$ end for 15:16:**Aggregation:** Construct solution $y_{ij}^{M\nu}, \forall (i,j) \in \mathcal{A}$ according to (25) 17:Update the best feasible solution $y^{Best} \leftarrow y^{M\nu}$, if appropriate; 18:Let $\tilde{y}_{ij}^{\nu} \leftarrow \sum_{g \in G} p^g \mathbf{f}_{ij}^{g\nu}, \ \forall (i,j) \in \mathcal{A} \text{ where } p^g = \sum_{s \in C_g} p^s$ 19:Update solution $\hat{y}_{ij}^{\nu}, \forall (i,j) \in \mathcal{A}$ (Section 4.3.2) 20: Penalty updates: 21: Adjust penalty values $\lambda_{ij}^{g\nu} \leftarrow \lambda_{ij}^{g\nu-1} + \rho(\mathbf{f}_{ij}^{g\nu} - \tilde{y}_{ij}^{\nu})$ and $\rho^{\nu} \leftarrow \alpha \rho^{\nu-1}$ 22: 23: end while 24: Phase II: Solve a restricted MIP problem 25: Fix the design variables for which consensus is obtained 26: Solve the restricted SMCND model (1)-(5) to obtain a final design y^{final} 27: Update best solution, $y^{Best} \leftarrow y^{final}$ if appropriate.

5 Experimental results

This section presents the results obtained from extensive computational experiments performed to assess the performance of the proposed algorithm. We used two collections of instances which are described in Section 5.1. To evaluate the performance of the proposed ILPH algorithm, we compared its results to those of alternative approaches that were tested on the same instances:

• IBM-ILOG CPLEX 12.6.1 with its default settings (CPLEX in the following) on

the associated MIP,

- The basic progressive hedging with single scenario subproblems, where subproblems are solved using CPLEX (PHS in the following),
- The progressive hedging with multiple scenario subproblems (PHG in the following) proposed in Crainic et al. (2014),
- The reduced cost-based restriction and refinement matheuristic (RCHeur in the following) proposed in Sarayloo et al. (2021b) that currently defines the state-of-the-art heuristic algorithm for the considered problem.

After presenting the two collections of instances and the experimental setting, we start by analyzing the internal performances of the proposed ILPH on the first collection of instances, with small number of scenarios, in Section 5.2. To assess the advantage of the proposed algorithm to efficiently deal with a large number of scenarios, we provide performance comparisons of the proposed ILPH versus PHS, PHG, RCHeur and CPLEX on the second collection of instances in Section 5.3.

5.1 Data and experimental setting

We consider six problem classes (R5-R10) from the set of SMCND R instances introduced in Crainic et al. (2011, 2014). Each class is characterized by a number of nodes $|\mathcal{N}|$, a number of arcs $|\mathcal{A}|$, and a number of commodities $|\mathcal{K}|$, specified in Table 1. For each instance class, we consider five networks, namely, networks 1, 3, 5, 7, and 9 indicating continuously increasing ratios of fixed to variable costs and total demand to total network capacity. In the first collection of instances, for each of these networks, there are instances with 16, 32, and 64 scenarios. Demands were assumed to be linearly correlated, and two different levels of correlations (0.2 and 0.8) were used to create the different instances. A total of 180 instances were thus obtained. In the second collection of instances, for each of the networks, there are 10 instances with 1000 scenarios which were generated according to the procedure in Boland et al. (2016). The data of all instances are available on the website: https://github.com/FSarayloo/JOH-Data.git.

Algorithms were coded in C++ using IBM-ILOG CPLEX 12.6.1 as the MILP solver. All parameters of the PHS and PHG methods (including the stopping criteria parameters) were set to the values originally used in Crainic et al. (2011, 2014). A brief summary of the parameter values is provided in the Appendix. The same parameter values are used in ILPH and PHS with the exception of N_{Imp} . According to the results provided in Sarayloo et al. (2021b), we set the values of N_1 and p_1 to 3 and 2, respectively. The value of $N_{\Delta g}$ is also set to $|\mathcal{K}| * |C_g|$. Hence, almost all parameters of the proposed ILPH (except N_{Imp} and τ) were set to the values established in previous studies including (Sarayloo et al. (2021b,a); Crainic et al. (2011, 2014)) without applying additional parameter tuning procedures.

As for the τ , N_{Imp} parameters, a preliminary analysis was conducted to fine tune them. For this preliminary analysis, the following values were considered:

- $N_{Imp} = 2, 4, 8, 10$
- $\tau = 0.9, \, 0.95, \, 0.98$

for a total of 12 possible combinations of these parameter settings. An exhaustive evaluation of all the combinations was then performed on a subset of 24 instances. This subset of instances was obtained by randomly selecting four instances among each group of instances R5– R10. This analysis enabled us to conclude that the best results were obtained with the combination: $\tau = 0.95$, $N_{Imp} = 4$. Nevertheless, the analysis showed that the difference between the best and the worst optimality gap among the 12 different combinations is less 0.5%, indicating that these combinations could provide about the same average solution quality. We let $\hat{y}^0 \leftarrow y^{exp}$ be the initial integer solution at iteration 0 in the Learn&Optimize procedure, where y^{exp} is the optimal solution to the expected value (EV) problem. The EV problem is obtained by replacing the random demand variables by their expected values and solving the resulting deterministic problem. To reduce the time required to complete Phase I, the optimality tolerance parameter of CPLEX was set to 1% when solving the subproblems. This parameter is set to its default value when solving the restricted problem of the second phase. Unless otherwise specified, all other CPLEX parameters were set to their default values since preliminary experiments indicated that these settings yielded better results. All experiments were performed on a Sun Fire X4100 cluster of 16 computers. Each has two 2.6 GHz Dual-Core AMD Opteron processors and 8192 Megabytes of RAM, operating under Solaris 2.10.

Tabl	le 1: Chara	cteris	tics of	instances
	Problem	$ \mathcal{N} $	$ \mathcal{A} $	$ \mathcal{K} $
	R05	10	60	25
	R06	10	60	50
	R07	10	82	10
	R08	10	83	25
	R09	10	83	50
	R10	20	120	40

5.2 Internal performance analysis

We first focus on the first collection of instances (with 16, 32, and 64 scenarios) for which CPLEX is able to provide either the optimal solution or at least a feasible solution

within the time limit of 8 hours. In the following, we evaluate different components of the proposed ILPH and analyse their behaviour on these instances.

5.2.1 Effect of using Learn&Optimize

We first evaluated the effect of using the Learn & Optimize on the performance of the proposed ILPH. To do so, we considered a variant of ILPH where the subproblems are solved exactly using CPLEX, which we refer to as ILPH(E). We also investigated the impact of different network characteristics and correlations. The results of these tests are summarized in Table 2. Each row presents the results obtained on 18 instances, which are characterised by different network index values and correlations. We compare the three methods, CPLEX, ILPH and ILPH(E), in terms of optimality gaps (considering the lower bound provided by CPLEX) and computation time (in seconds) denoted by "Gap" and "Time", respectively. The "Time" indicates the time at which the best solution was found (within the time limit of 8 hours).

The results in Table 2 show that using Learn&Optimize instead of an exact method improves the performance of the algorithm. We observe that ILPH dominates ILPH(E) in terms of both solution quality (on average, % gap of 1.41 versus 1.77) and computation time (880 versus 3660 seconds). Furthermore, the results show that the proposed ILPH displays a consistent behaviour in dealing with different network characteristics. In particular, for the instances of index 3 and 5, which are considered as the most difficult instances, ILPH is able to reduce the optimality gap of CPLEX by up to 50%, while also reducing the solution time by a factor of around 30 (on average, 441 vs 14108 seconds). We also observe that the demand correlation has little impact on how difficult the associated problems are to solve. Overall, these results clearly highlight the significant benefit of the proposed algorithm combining PH and Learn&Optimize.

Index	Corr	# of	CF	PLEX	ILF	ILPH		(E)
		Ins	Gap	Time	Gap(%)	Time	Gap(%)	Time
1	0.2	18	0.00	340	0.02	65	0.02	161
1 ¹	0.8	18	0.00	349	0.02	65	0.02	187
2	0.2	18	5.37	14059	2.45	427	2.79	1169
5	0.8	18	5.39	14208	2.45	474	2.79	1279
5	0.2	18	2.67	15686	1.70	914	1.79	5797
	0.8	18	2.69	15890	1.72	963	1.79	6004
7	0.2	18	0.31	7588	0.30	811	0.61	4207
	0.8	18	0.31	7759	0.30	861	0.61	4494
0	0.2	18	2.54	14550	2.05	2045	2.97	6180
9	0.8	18	2.57	14990	2.09	2165	2.97	6842

 Table 2: Effect of using Learn&Optimize

To analyze more thoroughly the behaviour of the proposed algorithm, we aggregate the results according to the number of scenarios (indicated by |S|) in Table 3. The descriptions of the columns 2 to 6 are the same as in Table 2. As expected, increasing the number of scenarios makes the problem more difficult to solve for both ILPH and CPLEX. However, the ILPH outperforms CPLEX on all instances. In particular, for instances with 16 and 32 scenarios, ILPH is able to find the solutions with the same quality as those found by CPLEX in significantly less time (i.e., on average, 463 and 647 seconds with ILPH compared to 6964 and 10134 seconds with CPLEX, for the instances with |S| = 16 and |S| = 32, respectively). Furthermore, over all instances with 64 scenarios, ILPH is able to obtain solutions with on average optimality gap of 1.89% versus 4.31% obtained by CPLEX by again notably reducing the computational time (i.e., on average, 2362 seconds and 13347 seconds for ILPH and CPLEX, respectively). In order to show the capability of the proposed method to address instances with large number of scenarios, we will analyze the performance of our proposed ILPH by solving difficult benchmark instances with 1000 scenarios in Section 5.3.

S	# of	CPL	EX	ILF	Ч
	Ins	Gap(%) Time		Gap(%)	Time
16	60	0.39	6964	0.39	463
32	60	1.52	10134	1.10	647
64	60	4.31	13347	1.89	2362

Table 3: Aggregated results according to the number of scanarios

5.2.2 Effect of using \tilde{y} and \hat{y}

We next study the effect of using the two new important components of the proposed algorithm, i.e., the proposed reference point \tilde{y} and solution \hat{y} , on the performance of ILPH. To evaluate the effect of \tilde{y} , we consider an alternative, which we refer to as ILPH- (\tilde{y}) where the traditional reference solution $\bar{y} \leftarrow \sum_{g \in G} p^g y^g$ is used instead of \tilde{y} . Also, in columns "ILPH- (\hat{y}) ", we evaluate the effect of using \hat{y} by replacing it with the EV solution (y^{exp}) as the initial solution for the *Learn&Optimize* in all iterations of ILPH. The results in Table 4 show that using the proposed \tilde{y} provides better quality solutions with an average gap of 1.41% within almost half the computation time, compared to ILPH- (\tilde{y}) , where the traditional \bar{y} is used. We also observed that using the proposed initial solution \hat{y} in ILPH has a good effect on providing slightly better quality solutions in almost 70% less computation time compared to ILPH- (\hat{y}) .

Pro	# of	CPLEX		ILF	РΗ	ILPH-(\tilde{y})		$ILPH-(\hat{y})$	
	Ins	Gap	Time	Gap(%)	Time	Gap(%)	Time	Gap(%)	Time
R05	30	0.00	1411	0.12	63	0.14	110	0.12	97
R06	30	1.52	11076	0.94	451	1.02	1420	0.98	980
R07	30	0.12	2130	0.31	105	0.35	152	0.32	137
R08	30	0.96	7516	1.08	841	1.08	1764	1.08	1341
R09	30	3.48	16415	1.91	910	1.95	1971	1.91	1710
R10	30	7.01	23557	4.14	4081	4.34	8891	4.18	6170
Avg		2.18	10350	1.41	1075	1.48	2384	1.43	1739

Table 4: Effect of introducing the reference point \tilde{y} and the design \hat{y}

5.2.3 Effect of different grouping strategies

Table 5 reports the results obtained by the proposed ILPH when different grouping strategies, i.e., "Random", "Similar" and "Dissimilar", are considered, which we refer to as ILPH(R), ILPH(S), and ILPH(D), respectively. The description of these grouping strategies is detailed in Crainic et al. (2014). We provide the comparison results in Table 5. We observed that ILPH(R), ILPH(S), and ILPH(D) provide almost similar optimality gaps of 1.41%, 1.50%, and 1.44%, respectively. However, the computation time by ILPH(R) is 45% and 30% faster than ILPH(S) and ILPH(D), respectively. It is also interesting to note that ILPH(R) is almost 10 times (on average) faster than CPLEX in providing better quality solutions (on average, the gap is 1.41% versus 2.18%).

Pro	# of	CPLEX		ILPH	ILPH(R) I		ILPH(S)		ILPH(D)	
	Ins	Gap(%)	Time	Gap(%)	Time	Gap(%)	Time	Gap(%)	Time	
R05	30	0.00	1411	0.12	118	0.14	146	0.12	136	
R06	30	1.52	11076	0.94	951	0.97	1467	0.95	1274	
R07	30	0.12	2130	0.31	105	0.31	149	0.31	128	
R08	30	0.96	7516	1.08	841	1.12	1638	1.10	1168	
R09	30	3.48	16415	1.91	840	2.02	1852	1.92	942	
R10	30	7.01	23557	4.14	4081	4.48	6480	4.24	5218	
Avg		2.18	10350	1.41	1075	1.50	1955	1.44	1470	

Table 5: Performance comparison versus CPLEX on first collection of instances

The results above are encouraging and demonstrate the potential of ILPH, but the instances are inadequate to fully assess the computational advantage of ILPH. Most instances in this set can be solved to optimality by CPLEX in less than 2 hours. This is not the setting for which a decomposition approach is designed. The ILPH is designed to be used in settings where the instances are large, difficult, and cannot be solved in a reasonable amount of time when providing the MILP formulation to a solver. Indeed, solving the root relaxation may already be computationally prohibitive. In the following subsection, we present results on instances that are more appropriate to show the benefits

of ILPH.

5.3 Comparative analysis

In this part of experiment, we focus on the difficult instances with 1000 scenarios. We provide the comparative results of the proposed ILPH versus CPLEX in Tables 6 and 7, versus the progressive hedging-based methods (PHS and PHG) and RCHeur in Table 9. Each row of these tables refers to 10 instances with 1000 scenarios with the different characteristics mentioned in Section 5.1. The results displayed in these tables show the advantage of the proposed method in dealing with such difficult instances.

We apply the ILPH considering three different group sizes, of 20, 50, and 100 scenarios, the results being identified as ILPH(20), ILPH(50), and ILPH(100), respectively. We then compare the best obtained solution versus that of CPLEX in Table 6. We report the average optimality gap, OptGap, that CPLEX provides after a time limit of 8 hours. Note that, we consider an optimality gap of 100% for the instances for which CPLEX is unable to provide any feasible solution. In the "Diff" columns, we report the relative difference (%) between the best solutions found by the two algorithms (i.e., the upper bounds), noted z_{ILPH} and z_{CPX} , after 8 hour CPU time limit, computed as $100*(z_{ILPH} - z_{CPX})/z_{ILPH}$. Negative values refer to the cases for which ILPH provides better solutions than CPLEX. The "Time" columns display the respective total computation times.

The results show that ILPH outperforms CPLEX in all the cases, for all group sizes, providing an improvement of at least 24.72% over the best CPLEX solution, within about one third of computation time. Comparing the group sizes, a group size of 100 seems to yield a better performance (especially in the case of difficult instances, e.g., R6 and R9) in terms of both solution quality and computation time. In particular, for R9 instances with an average optimality gap of 66.37%, ILPH(100) provides solutions with an improvement of 63.58% compared to 52.42% when the group size is 20. This is due to the capability of *Learn&Optimize* to handle subproblems with many scenarios, which ultimately leads to good solutions obtained by ILPH. As for the results reported in Tables 5 and 6, they involve the use of "Random" grouping strategy, which is applied to define suproblems with 100 scenarios. This grouping strategy is then used to perform the rest of the analyses.

To highlight the capability of the proposed ILPH to identify good solutions quickly, we report the comparative results of ILPH versus CPLEX in a shorter amount of time. Table 7 displays the results obtained by the two methods within 3 hours. We report in column ILPH/CPLEX(3h) the relative difference (%) between the best solutions found by the two algorithms (i.e., z_{ILPH} and z_{CPX}) by imposing a 3 hour computation time limit. The relative difference is computed as $100^* (z_{ILPH} - z_{CPX})/z_{ILPH}$. Negative values indicate ILPH provides better solutions than CPLEX. In the last columns, we compare the two

				1						
Pro	CPLE	Х		Diff(%)			Time			
	OptGap(%)	Time	ILPH(20)	ILPH(50)	ILPH(100)	ILPH(20)	ILPH(50)	ILPH(100)		
R05	17.94	25805	-12.80	-12.89	-12.95	11292	12829	10842		
R06	47.87	24884	-35.32	-37.12	-37.62	12682	11886	11280		
R07	9.70	24063	-5.94	-5.09	-5.05	7952	7292	7494		
R08	27.72	29013	-17.12	-18.36	-18.40	8682	8604	9258		
R09	66.37	29901	-52.42	-62.42	-63.58	24253	23453	22630		
Avg	33.92	26743	-24.72	-27.40	-27.52	12971	12892	12410		

Table 6: Performance comparison of ILPH versus CPLEX

solution methods based on the percentage of instances for which the considered solution method is able to provide a feasible solution. The average optimality gap provided by CPLEX is 45.59%, indicating the difficulty of these problems. We observe that ILPH is able to find feasible solutions for all considered instances, while CPLEX is unable to do so for 32% of instances. Furthermore, in terms of solution quality, ILPH provides an average improvement of -39.23% versus CPLEX, which is very interesting.

Pro	# of	CPLEX	ILPH/CPLEX(3h)(%)	Sol	L(%)
	Ins	OptGap(%)		ILPH	CPLEX
R05	10	22.56	-12.62	100	100
R06	10	63.90	-62.31	100	40
R07	10	15.97	-4.34	100	100
R08	10	41.36	-35.03	100	80
R09	10	84.18	-81.84	100	20
Avg		45.59	-39.23	100	68

 Table 7: Efficiency of the ILPH versus CPLEX

Table 8 displays the performance results after Phase I of the proposed ILPH compared to PHS and PHG, in terms of solution quality and computation time. In order to investigate how the proposed ILPH performs compared to the other two methods, we consider the objective function value of the best solution provided after the first phase (y^{Best}) , by ILPH, PHS, and PHG. Columns "*ILPH/PHS*" and "*ILPH/PHG*" report the relative improvement (in %), computed as $100 * (z_{ILPH} - z_{PHS})/z_{ILPH}$ and $100 * (z_{ILPH} - z_{PHG})/z_{ILPH}$, respectively. Negative values indicate that ILPH outperforms PHS and PHG. We also report the average computation times in seconds (*Time*) and the percentage of instances with a feasible solution (*Sol.*%) for the three methods.

We observe that the proposed ILPH provides solutions with an average improvement of 15.28% within one third of computation time compared to PHS. Also, ILPH dominates PHG providing 12.52% improvement on solution quality within almost half the computation time. Furthermore, ILPH is able to provide a feasible solution for all considered instances, while, PHS and PHG fail to provide any solution for 8% and 6% of the instances, respectively.

Pro	# of	ILPH/PHS	ILPH/PHG		Time	Sol(%)			
	Ins	(%)		ILPH	PHS	PHG	ILPH	PHS	PHG
R05	10	-8.45	-6.84	582	5325	4140	100	100	100
R06	10	-10.14	-9.27	4121	19277	15842	100	100	100
R07	10	-10.16	-9.20	1210	3576	2386	100	100	100
R08	10	-5.14	-5.14	13755	14822	14196	100	100	100
R09	10	-42.47	-32.17	7611	30613	21640	100	60	70
Avg		-15.28	-12.52	5456	14728	11640	100	92	94

Table 8: Performance comparison of ILPH versus PHS and PHG (after Phase 1)

Finally, Table 9 reports the comparative final results (after Phase II) of ILPH versus that of PHS, PHG, and RCHeur over all 50 instances. The percentages of relative improvements are calculated in the same way as for Table 8. Overall, ILPH outperforms PHS, PHG, and RCHeur with average relative improvements of 18.81%, 14.12%, and 11.37%, respectively. Moreover, ILPH provides these solution much faster than the other methods, with an average computational time of 12470 versus 29645, 2378,1 and 19360, respectively.

Table 9: Performance comparison of ILPH versus PHS, PHG and RCHeur (final solution)

Pro	# of	ILPH/PHS	ILPH/PHG	ILPH/RCHeur		Ti	ime	
	Ins		(%)		ILPH	PHS	PHG	RCHeur
Avg	50	-18.81	-14.12	-11.37	12470	29645	23781	19360

6 Conclusions

This paper explores the development of an efficient optimization approach to address SM-CND problems. We propose a two phase *Integrated Learning and Progressive Hedginig* matheuristic to handle a large number of scenarios in the considered context. We generalize the *Learn&Optimize* procedure to efficiently address multi-scenario subproblems at each iteration of the PH algorithm. We further exploit the knowledge learned through the *Learn&Optimize* to improve the performance of the matheuristic. Specifically, we introduce a new reference point in each aggregation step of ILPH by making full use of the knowledge that is gathered regarding the promising design variables when solving the subproblems. By proceeding in this way, we inject the knowledge learned through the heuristic procedure into the PH algorithm leading to the proposed *ILPH* method, which is the main contribution of this paper.

We show, through computational experiments, that the proposed algorithm performs very well in terms of both solution quality and computation time. We provide a comparative analysis that show the superiority of the proposed approach versus CPLEX, as well as other PH-based algorithms and matheuristics proposed in the literature. The results indicate that the proposed algorithm provides impressive improvements of 27.40% and 18.81% when dealing with large instances with 1000 scenarios, versus CPLEX and the basic PH algorithm, respectively. The analysis also indicates that ILPH should be the method of choice when high-quality solutions to instances of stochastic network problems containing a large number of scenarios need to be found quickly.

We conclude by identifying a few possible directions for future research. These include investigating whether the algorithm would be as successful for different optimization problems or other variants of MCND. Indeed, ILPH is a general-purpose algorithm and can be applied to different stochastic programs. Tailored implementations of the proposed PH algorithm can lead to quite effective matheuristics for very large stochastic programming applications. Other research avenues include considering other strategies for updating the penalties within the PH algorithm and other methods for solving the subproblems. Finally, another important research direction is to develop parallel strategies for the ILPH algorithm, which will further amplify its advantages and benefits.

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Appendix

For implementation of the PHS and PHG in Section 5, we used similar parameters and stopping criteria as presented in Crainic et al. (2011, 2014) as follows: $N_{Itr} = 1000$, $N_{Imp} = 10$, $\gamma = .1$, $\beta = 1.1$, $c^{high} = 0.8$, $c^{high} = 0.2$, $c^{near} = 0.7$, $c^{far} = 0.2$, and $t_{max} = 500$ minutes. Wer effer readers to Crainic et al. (2011, 2014) for more detailed algorithmic implementations and descriptions.

Declaration

Conflict of interest On behalf of all authors, the corresponding author states that there is no conflict of interest.