

# Hierarchical Constraint-Based Scheduling for Multi-Robot Observation Missions

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**Abstract.** In this paper, we consider scheduling problems involving tasks (*area observations*) that can be realized by several resources (*robots*), together with setup operations between tasks (*robot moves between observations*) that are decomposed into low level operations (*moves on edges of a waypoint graph through multiple possible paths*). To deal with such problems, we introduce a framework for representing hierarchical scheduling problems, and for which constraint programming models can be automatically generated. We also define an iterative two-layer constraint-based decision process that alternates between the fast synthesis of high-level schedules based on a coarse-grain model of setup operations, and the production of a detailed schedule based on a fine-grain model of these operations. Experiments realized on representative benchmarks of a multi-robot application show the efficiency of the approach.

**Keywords:** Hierarchical Scheduling · Constraint Programming · Multi-Robot Missions.

## 1 Introduction

In this paper, we deal with so-called *hierarchical scheduling problems*, that have a hierarchical structure in multiple ways. To present our approach, we focus on one of the real case-studies that motivates this work, namely a multi-robot deployment mission in which a fleet of robots must perform a set of observations on specific areas of a field, for situation awareness issues. Our problem is to allocate each candidate observation to a robot, schedule the sequence of observation tasks realized by each robot, and plan navigation tasks between observation locations. The objective is to realize all observations as quickly as possible. It is assumed that robots cannot perform more than one observation at a time. They must also transfer observation data in real time to the mission center, and for this purpose each robot uses a specific emission frequency. To avoid interferences, two robots that use the same frequency cannot transfer observation data simultaneously. Redundancy is also useful in this kind of application, therefore each observation target must be observed by several different robots. Moreover, some precedence constraints can be imposed over observations. Finally, a graph of waypoints is

used to represent the structure of the field, and the movements of robots between observation locations can be broken down into successive movements between pairs of adjacent waypoints. To avoid collisions, each link between two waypoints cannot be occupied by more than one robot at a time.

These specifications lead to a task scheduling problem that includes (1) *tasks which can be realized by several candidate resources*, (2) *setup operations* between the tasks realized by a resource (the moves of robots between observation locations), (3) *several alternatives for realizing each setup operation* (several candidate navigation paths between two distant observations), and (4) *for each setup alternative, a decomposition into low level tasks* (in our case, each path traversal is decomposed into a set of moves on edges of the waypoint graph).

The first contribution of this paper is the introduction of a framework for representing Hierarchical Scheduling Problems (HSPs), to deal with such applications in a generic way. This framework is inspired by *Hierarchical Task Networks* (HTNs [9]) used in the planning community. The main difference is that HSPs define scheduling problems based on tasks and resources, while HTNs are basically suitable for planning problems based on actions and states. We also introduce techniques to automatically generate *flat* constraint programming encodings from HSPs. For non-experts in Constraint Programming (CP), this allows to benefit both from a hierarchical view of the domain in the form of HSPs and from the strength of CP techniques. One of our goals is to push further the applicability of standard powerful CP solvers to hierarchical scheduling problems.

The second contribution is the definition of a hierarchical decision strategy that splits the decision process into several scheduling layers, to deal with HSPs having a large number of candidate task decompositions. Basically, we divide the process into (1) a layer L1 that produces high-level decisions based on a coarse-grain model of all setup operations, and (2) a layer L2 that produces detailed schedules based on a fine-grain model of all low-level tasks realized during setups. We then iteratively solve the scheduling problems of layers L1 and L2, given that at each step the planning data used by L1 (such as the approximate durations of setups) is updated according to the detailed schedules returned by L2.

The paper is organized as follows. Section 2 describes the HSP framework and the proposed CP encoding. Section 3 defines the iterative two-layer decision process. Section 4 provides experimental results obtained on representative benchmarks of the robotic application. Section 5 discusses related works and Section 6 gives some perspectives.

## 2 Hierarchical Scheduling Problems (HSPs)

### 2.1 General Scheduling Model

We consider a set of disjunctive resources  $\mathcal{R}$ , which cannot be used by several tasks simultaneously. This set is partitioned between the set  $\mathcal{R}^d$  of simple disjunctive resources and the set  $\mathcal{R}^s$  of disjunctive resources with setup times, for

which a transition duration is required between successive tasks realized by the resource. Each resource  $r \in \mathcal{R}^s$  has  $S$  possible running states and each task consuming  $r$  requires a particular resource state  $s \in [1..S]$ . For every pair of resource states  $(s, s') \in [1..S]^2$ , a minimum setup duration function  $setup_r$  gives the duration  $setup_r(s, s') \in \mathbb{N}$  required between the end of a task using  $r$  in state  $s$  and the start of a task using  $r$  in state  $s'$ .

Resources in  $\mathcal{R}$  are used to execute *tasks*. A task  $t$  is defined by:

- a release date  $rd_t$  after which it can start and a due date  $dd_t$  before which it must end;
- the set of resources  $R_t \subseteq \mathcal{R}$  it consumes all along its execution (from the start of the task to its end), with for every resource with setup  $r \in R_t \cap \mathcal{R}^s$  the state  $st_{t,r}$  required for  $r$  during the execution of  $t$ .

A task is either *primitive* or *compound*:

- a *primitive task*  $o$ , also called an *operation*, has a fixed duration  $du_o$ ;
- a *compound task*  $c$  has a list of possible *decomposition methods*  $M_c = [m_{c,1}, \dots, m_{c,k}]$  usable for realizing  $c$ ; each decomposition method  $m \in M_c$  corresponds to a so-called *task network*  $(T_m, \Psi_m)$ .

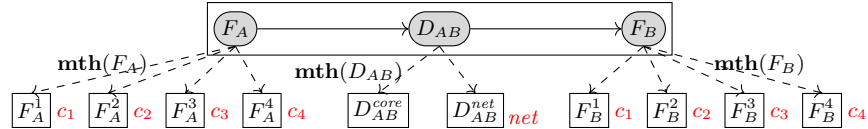
Formally, a task network  $(T, \Psi)$  is composed of a set of tasks  $T$  and a set of constraints  $\Psi$  over these tasks. We consider two types of constraints in  $\Psi$ :

- *temporal constraints* over tasks in  $T$ , denoted  $\Psi_{temp}$ ; in this paper, we only consider a set of acyclic precedence constraints between pairs of tasks, but the approach can deal with any minimum and maximum distance constraint between the start and end time-points of tasks in  $T$ ;
- *decomposition constraints*, denoted  $\Psi_{decomp}$ , that restrict the possible combinations of decompositions of compound tasks in  $T$ ; they are expressed over  $\{\mathbf{mth}(c) \mid c \in T \text{ s.t. } c \text{ is compound}\}$ , where  $\mathbf{mth}(c)$  denotes the index  $i$  of the method  $m_{c,i}$  chosen for realizing task  $c$ .

A *Hierarchical Scheduling Problem* (HSP) is then defined by a *task network*  $(T_0, \Psi_0)$ , called the *root task network*, which represents the set of high-level tasks to be realized. In the following, we denote by  $\mathcal{O}$ ,  $\mathcal{C}$ , and  $\mathcal{M}$  respectively the set of all operations, compound tasks, and methods involved in the full hierarchical decomposition of the root task network. Moreover, for any resource  $r \in \mathcal{R}$ ,  $\mathcal{T}_r$  denotes the set of tasks  $t$  (primitive or compound) that consume resource  $r$ , i.e. such as  $r \in R_t$ .

To illustrate the previous definitions, we consider an HSP associated with the placement of embedded functions over a multi-core platform. Consider two functions  $F_A$  and  $F_B$  that can be allocated to four distinct cores  $c_1 \dots c_4$ , and such that  $F_A$  must transfer a data  $D_{AB}$  to  $F_B$ . If  $F_A$  and  $F_B$  are placed over the same core, this transfer takes 1 time unit, otherwise it uses the embedded network and takes 10 time units. Each core is viewed as a disjunctive resource, as well as the network between cores. Fig. 1 provides a graphical description of the hierarchical problem that can be built for this simple problem (real instances

involve thousands of functions and data transfers). In this problem, the root task network ( $T_0, \Psi_0$ ) contains three compound tasks ( $T_0 = \{F_A, D_{AB}, F_B\}$ ) and two precedence constraints  $F_A \rightarrow D_{AB}$  and  $D_{AB} \rightarrow F_B$ . Task  $F_A$  has four decomposition methods  $M_{F_A} = [m_{F_A,1}, \dots, m_{F_A,4}]$ , where the  $i$ th method corresponds to an allocation of the function on the  $i$ th core. Method  $m_{F_A,i}$  points to a task network  $(T, \Psi) = (\{F_A^i\}, \emptyset)$  which contains a unique primitive task  $F_A^i$  of fixed duration that consumes core  $c_i$ . The modeling is similar for  $F_B$ . Data transfer task  $D_{AB}$  has two possible decomposition methods. The first one points to a task network reduced to a single primitive task  $D_{AB}^{core}$  of duration 1 and corresponds to a data transfer over the same core. The second one also points to a task network reduced to a single primitive task  $D_{AB}^{net}$  of duration 10 which is used for a data transfer over the network, and which consumes the network resource. At the level of the root task network ( $T_0, \Psi_0$ ), one decomposition constraint is added to  $\Psi_0$  to express that a data transfer over the network is used when the two functions are allocated to distinct cores ( $(\text{mth}(D_{AB}) = 2) \leftrightarrow (\text{mth}(F_A) \neq \text{mth}(F_B))$ ).



**Fig. 1.** HSP for the deployment of embedded functions over a multi-core platform

Additionally, we assume that the addressed HSPs are *well formed*. More precisely, it is assumed that each task belongs to a unique task network and that the total number of tasks in an HSP is finite, which implies that the set of candidate decompositions of the root tasks is finite. It is also assumed that given a task  $t$  consuming a resource with setup times  $r$ , its release date  $rd_t$  is greater than or equal to the setup duration required to put resource  $r$  in state  $st_{t,r}$  from the beginning of the schedule.

## 2.2 Encoding in Constraint Programming

Given an HSP, it is possible to generate a *flat* CP encoding for minimizing the makespan. In this encoding, each task is represented by an *interval variable*. For each interval variable  $itv$ , **start**( $itv$ ), **end**( $itv$ ) and **pres**( $itv$ ) respectively denote the start date, the end date and the presence of the interval in the solution. Then, we introduce the following variables:

- for each task  $t \in \mathcal{O} \cup \mathcal{C}$ , an interval variable  $itv_t \in [rd_t..dd_t]$ ;
- for each compound task  $c \in \mathcal{C}$ , an integer variable  $mth_c \in [0..|M_c|]$  that represents the index of the decomposition method chosen for realizing  $c$  (value 0 used when the task is not present).

- for each compound task  $c \in \mathcal{C}$  and each decomposition method  $m \in M_c$ , an interval variable  $itv_m \in [rd_c..dd_c]$ .

In the CP encoding, we define Constraints (1) to (11) given below, which use several constraints available in the CpOptimizer tool.<sup>1</sup> They express that all root tasks must be executed (1) and that every compound task is realized using exactly one of its decomposition methods (2). The latter uses the *alternative*( $itv, I$ ) constraint available in CpOptimizer, which specifies that an interval  $itv$  is equal to exactly one of the intervals in  $I$ . Also, each decomposition interval must cover all its subtasks (3). The latter uses the *span* constraint specifying that an interval must cover all present intervals belonging to a given set. The presence of subtasks of a method must also coincide with the presence of the method (4). Constraints (1)-(4) actually correspond to already existing CP encodings of work breakdown structures.<sup>2</sup>

The other constraints introduced are used to represent the scheduling and decomposition constraints of the problem. First, every operation has a fixed duration (5). Also, if a compound task is not present in a solution, then the method choice index for this task is set to its default value (6), otherwise this index is consistent with the presence of the possible decomposition intervals (7). Constraints (8)-(9) enforce that tasks realized by disjunctive resources must not overlap, taking into account setup durations for resources with setup operations. For every task network  $(T, \Psi)$  used in the model, all precedence constraints in  $\Psi_{temp}$  must be satisfied (10). The latter uses the *endBeforeStart* constraint of CpOptimizer, which imposes that the end of a first interval must precede the start of a second one. Last, decomposition constraints  $\psi \in \Psi_{decomp}$  must be satisfied (11). For this, we directly write constraint  $\psi$  conditioned by the presence of the decomposition method from which the task network comes (or unconditioned for the root task network). For the experiments, decomposition constraints are manually added to the CP model generated, but there is no technical difficulty to integrate them in the generation procedure.

$$\forall t \in T_0, \mathbf{pres}(itv_t) = 1 \quad (1)$$

$$\forall c \in \mathcal{C}, \text{alternative}(itv_c, \{itv_m \mid m \in M_c\}) \quad (2)$$

$$\forall m \in \mathcal{M}, \text{span}(itv_m, \{itv_t \mid t \in T_m\}) \quad (3)$$

$$\forall m \in \mathcal{M}, \forall t \in T_m, \mathbf{pres}(itv_m) = \mathbf{pres}(itv_t) \quad (4)$$

$$\forall o \in \mathcal{O}, \text{duration}(itv_o, du_o) \quad (5)$$

$$\forall c \in \mathcal{C}, (\mathbf{pres}(itv_c) = 0) \leftrightarrow (mth_c = 0) \quad (6)$$

$$\forall c \in \mathcal{C}, \forall k \in [1..|M_c|], \mathbf{pres}(itv_{m_{c,k}}) = (mth_c = k) \quad (7)$$

$$\forall r \in \mathcal{R}^d, \text{noOverlap}(\{itv_t \mid t \in \mathcal{T}_r\}) \quad (8)$$

$$\forall r \in \mathcal{R}^s, \text{noOverlap}(\{(itv_\tau, st_{t,r}) \mid t \in \mathcal{T}_r\}, setup_r) \quad (9)$$

for every task network  $(T, \Psi)$ ,  $\forall \psi = (t, t') \in \Psi_{temp}$ ,

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<sup>1</sup> <https://www.ibm.com/analytics/cplex-cp-optimizer>

<sup>2</sup> <http://gdrro.lip6.fr/sites/default/files/JourneeROIA-IBM.pdf>

$$\text{endBeforeStart}(itv_t, itv_{t'}) \quad (10)$$

for every task network  $(T, \Psi)$ , constraints in  $\Psi_{decomp}$  (11)

This encoding can be simplified in the following specific cases. If there is a unique decomposition method  $m$  for a compound task  $c$ , intervals  $itv_m$  and  $itv_c$  are equal. In this case, only one interval instance is defined and Constraint (2) is not generated. Similarly, if a decomposition method  $m$  has a unique subtask  $t$  (*i.e.*  $T_m = \{t\}$ ), intervals  $itv_m$  and  $itv_t$  are equal. In this case, only one interval instance is defined and Constraints (3)-(4) are not generated. Also, the integer variable  $mh_c$  is introduced only to clearly illustrate the layers model, since in practice, constraints on the boolean variables  $\text{pres}(itv_m)$ , are better exploited by CP tools.

When the goal is to minimize the makespan, the objective function generated consists in minimizing the maximum end time of a task in the root network, that is to minimize

$$\max_{t \in T_0} \mathbf{end}(itv_t). \quad (12)$$

Note that the model and the encoding can easily be generalized to cumulative resources having a finite capacity or to non renewable resources. This allows to deal with *Resource Constrained Project Scheduling Problems (RCPSPs [5]) with modes*, and more generally with kinds of hierarchical RCPSPs. In the end, the representation framework obtained is both simple and easy to use by non-CP experts, and quite expressive.

### 3 Iterative Hierarchical Decision Strategy

We now come back to the robotic application presented in the introduction. To tackle this problem, a first option is to define a unique HSP containing, for each robot, all potential setup operations that might be used between any two observation tasks, all possible decompositions of these potential setup operations, and all primitive tasks involved in these decompositions. For our application, this leads to a number of tasks in  $\Theta(RO^2PL)$  with  $R$  the number of robots,  $O$  the number of observations,  $P$  the maximum number of candidate paths between two observations, and  $L$  the maximum number of network links on a single path.

To avoid handling such a huge number of tasks, another approach is to explicitly break down the problem to be solved into several sub-problems and to define a dedicated solution technique for each of these sub-problems. In this direction, we can use for our applications a two-layer decision strategy that first synthesizes a high-level schedule based on a coarse-grain model of setup operations (decision layer L1, which approximates these operations by simple setup durations), and then details this schedule based on a fine-grain model (decision layer L2, which only considers setup operations that are actually used in the coarse-grain solution). Such a top-down approach is quite commonly used in practice for hierarchical decision making. But as high-level decisions are computed from a coarse-grain model, it can fail to reach the highest quality solutions.

This is why we propose to iteratively use the two scheduling layers. More precisely, each time a new detailed schedule is produced by layer L2, input data of the imperfect coarse-grain model of layer L1 is updated, and a new high-level solution is looked for. Doing so, layer L1 iteratively learns a better approximation of the content of layer L2, the goal being to converge towards better full solutions. Before detailing these mechanisms, we first use the hierarchical scheduling framework introduced in the previous section for defining the models of decision layers L1 and L2 for a multi-robot application.

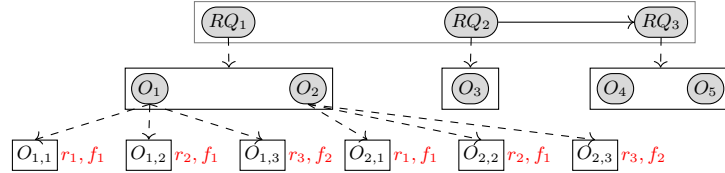
### 3.1 High-level Scheduling Model: Layer L1

*Inputs* The inputs of the first layer of the multi-robot observation mission are:

- a set of *frequencies*  $\mathcal{F} = [1..|\mathcal{F}|]$  for transferring observation data;
- a set of *robots*  $\mathcal{Rob} = [1..|\mathcal{Rob}|]$ ;  $\forall r \in \mathcal{Rob}$ ,  $freq_r \in \mathcal{F}$  is the frequency used by  $r$  to emit data during observations;
- a set of *observation requests*  $\mathcal{Req} = [1..|\mathcal{Req}|]$ , corresponding to areas of the field that must be observed. For each request  $j \in \mathcal{Req}$ ,  $duReq_j \in \mathbb{N}$  is the duration required to observe the area;
- a set of *(atomic) observations*  $\mathcal{Obs} = [1..|\mathcal{Obs}|]$  to be performed. For each request  $j \in \mathcal{Req}$ , it is required that different robots observe the corresponding area (redundancy). For each observation  $i \in \mathcal{Obs}$ ,  $req_i \in \mathcal{Req}$  denotes the request associated with  $i$ ;
- a set of acyclic precedence constraints  $\mathcal{P} \subseteq \mathcal{Req}^2$  between requests of  $\mathcal{Req}$ ;
- for each robot  $r \in \mathcal{Rob}$ ,  $duM_r : \mathcal{Obs} \times \mathcal{Obs} \rightarrow \mathbb{N}$  denotes the function such that  $duM_r(i, i')$  gives the shortest duration required by  $r$  to move from the location of observation  $i$  to the location of observation  $i'$  over all possible paths of the waypoint network; we extend  $duM_r$  so that  $duM_r(0, i')$  gives the shortest duration required to move from the initial location of  $r$  to observation  $i'$ ;
- a temporal horizon  $H \in \mathbb{N}$  for the whole mission.

*Hierarchical scheduling problem* The HSP built for layer L1 is illustrated in Fig. 2. It contains two kinds of resources: the frequency resources (disjunctive), used to emit observation data, and the robot position resources (disjunctive with setup times). For the latter, the states are the set of observations  $[1..|\mathcal{Obs}|]$ , and for each robot  $r$  the setup duration function is  $duM_r$ . The root task network  $(T_0, \Psi_0)$  contains one compound task  $RQ_j$  per request  $j \in \mathcal{Req}$ , and the set of precedence constraints  $\mathcal{P}$  that hold over the requests. Each compound task  $RQ_j$  has a unique decomposition method. The latter defines a task network which contains one compound task  $O_i$  per observation  $i \in \mathcal{Obs}$  such that  $req_i = j$ . Each compound task  $O_i$  has as many decomposition methods as the number of robots, and each of these decomposition methods defines a task network which contains a unique operation  $O_{i,r}$  representing the realization of observation  $i$  by robot  $r$ . Task  $O_{i,r}$  consumes one frequency resource (the frequency used by  $r$ ), and it uses the position resource of robot  $r$  in state  $i$ . Its duration is given by

the duration of request  $j$ . In this particular case, resources are consumed only by operations, and not by compound tasks. Release dates are determined based on the duration required by each robot to go from its initial location to every observation area. The due date of all tasks is set to temporal horizon  $H$ . Last, at the level of the task network associated with the decomposition of a request  $j$ , a decomposition constraint is added to ensure that observations associated with  $j$  are realized by distinct robots (e.g.,  $\text{mth}(O_1) \neq \text{mth}(O_2)$  for the example given in Fig. 2, and more generally we impose an *alldiff* constraint between the indices of the chosen decomposition methods), to meet the requirement to use different robots to observe a certain area (redundancy). As explained in the previous section, a CP model can be directly generated from this hierarchical description.



**Fig. 2.** HSP for layer L1

### 3.2 Low-level Scheduling Model: Layer L2

*Inputs* Layer L2 is responsible for detailing the routing of robots in the waypoint network and for managing navigation conflicts. To do this, it first receives from L1 a high-level schedule specifying one decomposition method for each observation  $O_i$  and an associated start date. From this, it is possible to derive, for each robot  $r \in \mathcal{Rob}$ , the list of observations that must be successively realized by  $r$ , written as  $\text{obsSeq}_r = [\text{obsSeq}_{r,1}, \dots, \text{obsSeq}_{r,|\text{obsSeq}_r|}]$  where  $\text{obsSeq}_{r,i} \in \mathcal{Obs}$ .

Layer L2 also takes as an input a representation of the navigation paths available in the waypoint graph, given by:

- a set of *links*  $\mathcal{L} = [1..|\mathcal{L}|]$ , where a link is a direct connection between two adjacent waypoints;
- for each robot  $r$  and each pair of successive observations  $(i, i')$  realized by  $r$ , a list of  $n$  candidate paths

$$P_{r,i,i'} = [P_{r,i,i',1}, \dots, P_{r,i,i',n}]$$

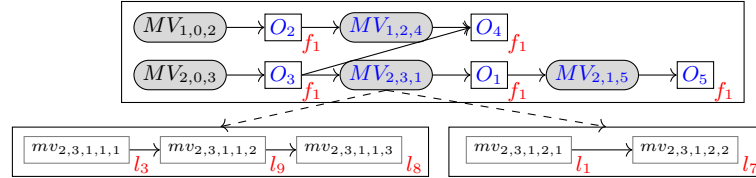
which can be used by  $r$  to go from  $i$  to  $i'$  (typically the  $n$  shortest paths); each path  $P_{r,i,i',p}$  is specified by a sequence of  $m$  links

$$\text{LinkSeq}_{r,i,i',p} = [\text{LinkSeq}_{r,i,i',p,1}, \dots, \text{LinkSeq}_{r,i,i',p,m}]$$

successively traversed on that path, where  $\text{LinkSeq}_{r,i,i',p,k} \in \mathcal{L}$ .

*Hierarchical scheduling problem* The HSP built for layer L2 is illustrated in Fig. 3, where robot  $r_1$  realizes observations  $O_2$  and  $O_4$ , and robot  $r_2$  observations  $O_3$ ,  $O_1$  and  $O_5$ . It contains two kinds of disjunctive resources: the frequency resources, used to emit observation data as in L1, and the link resources, consumed by the robot moves (two robots cannot simultaneously be on the same link). The root task network ( $T_0, \Psi_0$ ) contains the sequence of high-level moves and observations realized by each robot (two robots in the example provided). In Fig. 3, the sequences of observations of robots are  $obsSeq_1 = [2, 4]$  and  $obsSeq_2 = [3, 1, 5]$ . Robot 1 must therefore realize a compound move  $MV_{1,0,2}$  from its initial location to observation number 2, then a primitive observation task  $O_2$ , then a move  $MV_{1,2,4}$  from observation 2 to observation 4, and last a primitive observation task  $O_4$ . Each observation task consumes the frequency used by the robot. In the root task network, precedence constraints are imposed over these tasks to guarantee that each robot realizes an adequate move activity between two successive observations. The root task network also contains precedence constraints between some observations (coming from the precedence required between observation requests).

Each compound move  $MV_{r,i,i'}$  between two observations  $i, i'$  has as many decomposition methods as the number of possible paths  $P_{r,i,i',p}$  usable between the two observations (two possible paths in the case of compound move  $MV_{2,3,1}$ ). A decomposition using path  $P_{r,i,i',p}$  points to a task network which specifies a sequence of atomic moves  $mv_{r,i,i',p,l}$  required on links of the waypoint graph. Each atomic move consumes the link resource number  $LinkSeq_{r,i,i',p,l}$ . For instance, the first path for  $MV_{2,3,1}$  is the sequence of links  $[l_3, l_9, l_8]$  and each subtask  $mv_{2,3,1,1,k}$  consumes the  $k$ th link of the sequence. As for layer L1, a CP encoding can then be directly obtained from all these specifications, which are expressed in a framework that remains readable for non CP experts.



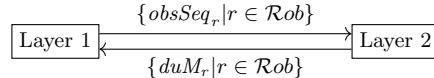
**Fig. 3.** HSP for layer L2

The problem solved in layer L1 can be associated with the Sequence Dependent Setup Time Job Shop Scheduling Problem presented in [15], but the complete problem we address, more than merely temporal constraint propagation and verification, implies decisions about which waypoints to visit between successive operations in layer L2.

### 3.3 Iterative Hierarchical Scheduling

We now present the interaction between the two layers modeled previously. As explained before, we define an iterative process in which the solution found by the second layer is used to update the inputs used by the first one. An illustration is given in Fig. 4 for the robot observation mission: layer L1 transmits the sequence of observations for each robot to layer L2, and the solution produced by the latter is used to update the coarse-grain duration of the setup operations between locations for each robot. The iterations between layers are performed until a given CPU time is reached.

Note that the purpose of this process is not to obtain an optimal solution for the real problem but to get solutions of good quality within a short time, which is more crucial than finding optimality in many multi-robot exploration missions. In fact, as layer L2 solutions are constrained by the solutions of layer L1 that works with an approximation of the global model, the iterative process has no guarantee to find an optimal solution. Note that if durations used by layer L1 are proven to be lower bounds of real durations, then the solution produced by layer L1 gives a lower bound of the expected makespan. That lower bound can be used to evaluate the distance between the final solution of the iterative process and the optimal solution.



**Fig. 4.** Iterative process using layers.

Algorithm 1 presents a generic pseudo-code of a process between two layers that use each others solutions in order to minimize an objective function. We first detail the main lines of this pseudo-code and then detail our implementation for the multi-robot application.

The first step of the algorithm is to initialize several elements. The last solution found by the different layers  $sol_1$  and  $sol_2$  and the best solution found by layer L2  $sol^*$  are all set to *null*. The best objective value found by L2 is denoted  $obj^*$  and is set up to an initial upper bound  $obj^{UP}$  (line 1). The input data for layer L1 are initialized through function **initL1** (line 2).

The process runs until a maximum CPU time is reached (line 3). In order to escape from local optima, we follow a restart strategy modeled by function **shouldRestart** that takes as parameters the last solutions found by the two layers. If a restart is required, function **perturbL1** re-initializes a given percentage of the input data of layer L1 (line 4). Then, if layer L2 has found a solution at the previous iteration, the latter is used by function **updateL1** to update the input data of the first layer (line 5). The problem to solve by L1 is created (line 6) and the associated solution is obtained through function **solve** to which a maximum CPU time is given. In our case, this CPU time is computed by a

function **maxTimeL1** that uses the number of iterations  $nLoops$  desired for the whole process and the CPU time elapsed (line 7). More complex strategies could be defined to share the CPU time between L1 and L2. If a solution exists for layer L1, then it is used to create the problem  $pb_2$  that is solved by layer L2 (lines 8-10). If this problem has a solution, we first compare its objective value to the best one found so far and update the latter if needed (lines 11 - 12). The best solution is finally returned (line 14).

---

**Algorithm 1: IterativeHS( $obj^{UP}$ ,  $cpuMax$ ,  $nLoops$ ):**


---

```

1  $sol_1, sol_2, sol^* \leftarrow null$ ,  $obj^* \leftarrow obj^{UP}$ ,  $it \leftarrow 0$ 
2 initL1()
3 while CpuTimeElapsed() <  $cpuMax$  do
4   if shouldRestart( $sol_1, sol_2$ ) then perturbL1()
5   else if  $sol_2 \neq null$  then updateL1( $sol_2$ )
6    $pb_1 \leftarrow \text{createPbL1}()$ 
7    $sol_1 \leftarrow \text{solve}(pb_1, \text{maxTimeL1}(it, nLoops))$ 
8   if  $sol_1 \neq null$  then
9      $pb_2 \leftarrow \text{createPbL2}(sol_1)$ 
10     $sol_2 \leftarrow \text{solve}(pb_2, \text{maxTimeL2}(it, nLoops))$ 
11    if  $sol_2 \neq null \wedge \text{objective}(sol_2) < obj^*$  then
12       $obj^* \leftarrow \text{objective}(sol_2)$ ,  $sol^* \leftarrow sol_2$ 
13     $it \leftarrow it + 1$ 
14 return  $sol^*$ 

```

---

We now detail how the functions and parameters defined previously are instantiated for the multi-robot application.

- **createPbL1**, **createPbL2** Creates problems along with variables and constraints as described in Sections 3.1 and 3.2.
- **solve** Solves the parameter problem using a CP solver that returns the best solution found until **maxTime**.
- **objective**,  $obj^{UP}$  The objective function is to minimize the makespan and  $H$  (i.e. the length of the temporal horizon) is used as an initial upper bound.
- **initL1** The data to be initialized for layer L1 are the approximate setup time matrices for each robot  $r$ . Depending on how the initialization is chosen, the algorithm can behave differently. If the matrices are initialized with lower bounds of the real duration (for instance,  $duM_r$ ), then layer L1 tends to provide optimistic solutions to layer L2 and the makespan of solutions of layer L2 can be greater than the ones of layer L1. On the contrary, if the matrices are initialized with upper bounds of the real duration, then layer L1 provides pessimistic solutions to layer L2. In our implementation, we have defined an optimistic layer L1 by using the matrices  $duM_r$  that give the shortest duration to go from one waypoint to another when each robot is alone in the waypoint graph.

- **shouldRestart** As we have implemented an optimistic layer L1, we expect the makespan of the solutions it produces to be lower than the one of layer L2. Therefore, we restart whenever layer L1 produces a solution with a makespan bigger than the best one found so far. For layer L2, we allow solutions with lower quality in order for layer L1 to improve its approximation, but we restart if layer L2 has the same makespan during a given number of consecutive iterations.
- **perturbL1** We randomly reinitialize a percentage  $rateReinit$  of the duration matrix of each robot.
- **updateL1** To update the duration matrices of layer L1 based on the solution given by layer L2 at the previous iteration, we have defined a reinforcement learning rate parameter  $\alpha \in [0, 1]$  that represents the influence of the duration of move tasks from a solution of layer L2 (denoted  $duML2_r$ ) on the input values of layer L1 (denoted  $duML1_r$ ). Formally,  $duML1_r$  is updated by  $(1 - \alpha) \cdot duML1_r + \alpha \cdot duML2_r$ .

The functions for creating and solving problems of each layer are implemented along with the description given previously in the paper.

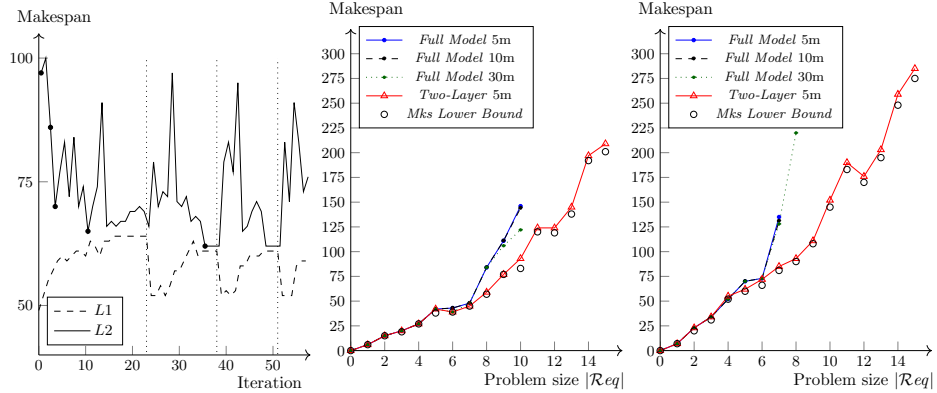
## 4 Experiments

**Instances** Experiments were performed over several multi-robot problem instances generated randomly, where the field structure contains  $3 \times |\mathcal{R}_{eq}|$  waypoints connected to their closest neighbors within a fixed range. The generated instances contain from 1 to 15 *observation requests*, each request requiring observations from 1 to 3 robots. From 1 to 3 *frequencies* are available to transfer observation data, and 3 *robots* are available to carry out the observations, each of them with different specifications (particularly a different speed which determines the duration needed to traverse a link). Function **updateL1** is implemented with a learning rate parameter  $\alpha$  ranging from 0.2 to 1 and the reinitialization rate is  $rateReinit = 0.2$ .

The generated instances were all tested on IBM ILOG CP Optimizer 12.5 on an Intel Xeon CPU E5-1603, 2.80GHz 8GB RAM, setting  $cpuMax = \{5, 10, 30\}$  minutes and an adequate iterations number  $nLoops$ , depending on the number of *observation requests* (size of the problem) and the  $cpuMax$ .

**Interaction between layers L1 and L2** To illustrate the interaction between the two layers, makespan results over iterations of the proposed approach are presented in Fig. 5 for a problem instance containing 5 *observation requests* (each one must be observed by 2 different robots) and 3 available *robots*. In order to accentuate the behavior, we have specifically implemented a very optimistic **initL1** function with matrix values  $duM_r$  equal to zero and we have used a parameter  $\alpha = 1$ . The mean duration of one L1-L2 loop iteration is 1.188 seconds. The best obtained makespan is marked with a filled dot.

Fig. 5 shows that makespans of both layers tend to converge quickly. When solutions cannot be improved, restarts are realized (vertical lines in the figure). More precisely, the first two restarts occurred when layer L1 did not find a better



**Fig. 5.** Interaction between layers

**Fig. 6.** Makespan results with 1 robot per request and precedences

**Fig. 7.** Makespan results with 2 robots per request and no precedences

solution than the best one found so far, and the third one because the makespan of the solution in layer L2 remained the same during several iterations. Since **initL1** is very optimistic, the makespan of solutions of layer L1 tends to increase, given that values of input duration matrices are also increasing.

**Full Model** To compare the hierarchical decision process with a global one-shot resolution strategy, we developed a full CP model of the robot deployment problem. This full model can be used to find an optimal solution. It contains (1) the model of layer L1 and (2) the model of layer L2 duplicated for each possible transition between observations. More precisely, the full CP model involves, for each robot  $r$  and each pair of distinct observations  $i, i'$ , one optional interval  $itv_{r,i,i'}$  representing a global move of robot  $r$  from  $i$  to  $i'$ , plus a huge number of optional intervals  $itv_{r,i,i',p,k}$  modeling the move of robot  $r$  on the  $k$ th link of the  $p$ th path available to transit from  $i$  to  $i'$ . To boost constraint propagation, the model contains, for each robot, both (1) the coarse-grain no overlap constraint of layer L1, which takes into account all observation intervals and minimum setup times between them, and (2) a fine-grain no overlap constraint that takes into account all observations and all move intervals over links associated with each robot. Last, a constraint is added to ensure that the successive fine-grain activities realized by each robot have consistent types, *i.e.* that an observation interval associated with observation  $i$  is preceded by a move interval of the form  $itv_{r,i',i}$ .

**Comparison between the Two-Layer process and the Full Model** All generated instances have been tested on the two-layer model and the Full Model with different maximum CPU times. Representative results are presented in Fig. 6 and Fig. 7, where the circle marks correspond to a makespan lower bound obtained by layer L1. For the instances of Fig. 6, only one robot must observe each request, and there are  $0.2 \times |\mathcal{R}_{eq}|^2$  randomly generated precedences

between requests. For Fig. 7, two different robots must observe each request and there are no precedences. Since the same makespan is reached for most of the experiments with several  $\alpha$  values and the impact on the behavior is not significant, we only present results for  $\alpha = 0.7$ . Similarly, giving more CPU time to the two-layers process does not improve significantly the best solution and we only present the results for 5 minutes.

For the Full Model, the generated instances contain from 14 interval variables in the smallest instance (with only one *observation request*) to 139543 ones in the largest instance. For the largest instances, the Full Model does not find any solution, even with a CPU time of 30 minutes. The two-layer approach achieves better makespan results in a significantly shorter time, which are very close to the obtained makespan lower bounds. It provides first solutions of good quality in less than 3 seconds (not represented on the figures), even for the largest instances in which the complete solver is not able to reach any solution after several minutes. For the smallest instances, the two-layer approach manages to find the optimal solution (without proving its optimality).

These results demonstrate that the proposed iterative approach is more effective than the Full Model for solving large hierarchical scheduling problems. It allows to quickly get first solutions with good quality no matter the problem size, which represents the biggest drawback when it comes to solving with the Full Model.

## 5 Related Works

As said before, HSPs are related to HTNs [9, 13], in which high-level tasks are broken down into primitive tasks using a catalog of decomposition methods. One of the main difference is that in HTNs, tasks basically interact through their preconditions and effects on the system state, whereas in HSPs they interact through their consumptions on resources. In standard HTNs, it is still possible to consider state features representing the current amounts of resources available, however this can make the reasoning about resource usages at different levels of the decision hierarchy quite complex. Some recent HTN planners explicitly integrate resources and temporal constraints, such as GSACB-SHOP2 [16]. The latter is however not available for comparing results.

ASPEN [7] and some other planners such as HiPOP [2] and FAPE [8] can reason about global abstract plans and progressively decompose the abstract tasks of these plans. CHIMP [17] is even able to handle online action plans containing subtasks shared between high-level tasks. To our knowledge, these planners do not benefit in their current state from the strength of constraint-based scheduling for reasoning about resources.

On the planning side again, several hierarchical planners such as CHIMP, Meta-CSP [12], ASPEN, EUROPA [1], PLATINUM [18], and other works such as RCPSP with time windows [6], Iterative flattening approaches [15] and the framework extending TFPOP [14], use hybrid domain knowledge mixing symbolic, temporal and resource reasoning. All these ingredients are integrated in an

iterative flaw resolution search technique that tries to repair at each step some flaws in the plan such as resource over-consumptions. The iterations involved in our search scheme are not used to solve flaws but to improve the quality of the coarse-grain model used by the high-level decision layer.

On the algorithmic side, the two-layer scheduling approach we propose can first be related to work on approximation models (or *surrogate models*) for black-box optimization [10]. Indeed, for layer L1, the computation of a solution through the combinatorial model of layer L2 can be seen as the computation of a complex evaluation function. The latter is summarized in layer L1 by a matrix of minimum transition durations between observations. Each time a new detailed schedule is available, this surrogate model is updated based on the real transition times that take into account interferences between atomic moves on links of the waypoint graph. In surrogate models, a key point is the choice of the next parameters for which the complex evaluation function (layer L2) must be computed. In our case, the more promising high-level schedules according to layer L1 are evaluated at each step. This also differs from some iterative incomplete search techniques such as Iterated Greedy Search and Large Neighborhood Search, where at each step, a part of the current solution is modified.

The interaction between layers can also be compared to Bender's decomposition [3], where the solutions produced by a MIP-based first-stage problem are sent to a second-stage problem that might produce new constraints (Benders cuts) which are violated by the current solution of the first-stage problem. In our case, the differences are that decision layer L1 is based on a CP model and not on MIP, and layer L2 does not need to compute cuts. Instead, it just returns a fine-grain plan from which the parameters of the coarse-grain problem of layer L1 are updated.

Last, in CP literature, representation hierarchies have already been used for configuration problems in which the specification of components has to be progressively refined [11]. One of the main differences with our approach is that HSPs cover hierarchies of tasks for scheduling problems, which leads to other solving techniques.

## 6 Conclusion

This paper introduces techniques for scheduling hierarchical tasks. As shown in the experiments, the approach exploits the strengths of already existing CP solvers and gives acceptable computation times, even on problems for which the set of possible task decompositions is large.

On the modeling side, we could add representation features to deal with action preconditions and action effects as in HTNs, and benefit from recent work on the CP encoding of such specifications [4]. On the algorithmic side, we could search for strategies that decompose compound tasks step-by-step, instead of having an arbitrary separation between two decision layers. One common point is that hierarchical description is more readable for the users.

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