

Semi-matching algorithms for scheduling parallel tasks under resource constraints

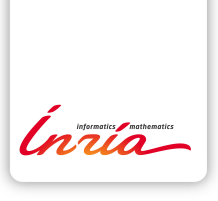
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Abstract: We study the problem of minimum makespan scheduling when tasks are restricted to subsets of the processors (resource constraints), and require either one or multiple distinct processors to be executed (parallel tasks). This problem is related to the minimum makespan scheduling problem on unrelated machines, as well as to the concurrent job shop problem, and it amounts to finding a semi-matching in bipartite graphs or hypergraphs. The problem is known to be NP-complete for bipartite graphs with general vertex (task) weights, and solvable in polynomial time for unweighted graphs (i.e., unit-weight tasks). We prove that the problem is NP-complete for hypergraphs even in the unweighted case. We design several greedy algorithms of low complexity to solve two versions of the problem, and assess their performance through a set of exhaustive simulations. Even though there is no approximation guarantee for these low-complexity algorithms, they return solutions close to the optimal (or a known lower bound) in average.

Key-words: semi-matching, bipartite graphs, hypergraphs, scheduling, parallel tasks, resource constraints.

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Algorithmes de couplage partiel pour l'ordonnancement de tâches parallèles sous contraintes de ressources

Résumé: On étudie le problème d'ordonnancement visant à minimiser le temps d'exécution lorsque les tâches peuvent être exécutées soit sur un seul processeur, soit sur plusieurs processeurs en parallèle, et sont restreintes à certains sous-ensembles de processeurs. Le problème se ramène à devoir trouver un couplage partiel dans un graphe biparti ou dans un hypergraphe. Sur les graphes bipartis, le problème général est NP-complet, mais il est possible de résoudre en temps polynomial les instances de problèmes où toutes les tâches sont de poids unitaire. Nous montrons dans ce rapport que le problème devient NP-complet pour les hypergraphes même dans le cas unitaire. Nous proposons plusieurs algorithmes gloutons pour résoudre deux versions du problème, et nous étudions leur performance à travers des simulations. Bien qu'il n'y ait aucune garantie d'approximation sur ces algorithmes, ils retournent en moyenne des solutions proches de l'optimal (ou d'une borne inférieure connue), avec un temps d'exécution très rapide.

Mots-clés : couplage partiel, graphes bipartis, hypergraphes, ordonnancement, tâches parallèles, contraintes de ressources.

1 Introduction

The Minimum Makespan Scheduling Problem on Unrelated Machines is a classical topic in scheduling [12]. It can be described as follows: given a set of tasks and a set of processors, assign the tasks to the processors such that the load among the processors is balanced, i.e., the maximum load of a processor is minimized. The tasks usually differ in their processing time, i.e., in the load that they create on the processor they are assigned to.

For today's high performance computing environment dominated by server virtualization, cloud computing, application accelerators and emerging architectures, we need to refine the problem formulation. Indeed, in classical scheduling, while the need for different computational resources can be expressed through the difference in processing time, it does not express the fact that a single task may have a choice among combinations of multiple computational resources. We therefore consider the MULTIPROC problem where (i) tasks are parallel, i.e., a task can be split in several identical parts, each part being computed independently on a different processor; (ii) tasks are subject to resource constraints, i.e., several configurations of processors, leading to different execution times, are possible for each task. The goal is to find one configuration for each task, in order to minimize the makespan.

MULTIPROC is related to the concurrent job shop problem [1]. In this problem, a job consists of multiple different components, each of which is to be processed on a specific dedicated processor. Components of the same job can be processed in parallel on their respective processors. A job is completed once all of its components are completed. According to Roemer [24], the problem was introduced by Ahmadi and Bagchi [1]. It has been studied widely [19, 20, 27] and was proved to be strongly NP-complete [24]. The main difference between MULTIPROC and the concurrent job shop problem is the resource constraints, i.e., whether components are restricted to a specific processor or not. In Mul-TIPROC, each task has the choice among multiple different sets of processors (different configurations). Those sets can differ in size (i.e., number of processors), but processing times are equal for all processors in each set. Most of the time, if there are more processors in a set, then the execution time becomes smaller on each processor of the set. Similarly to the concurrent job shop problem, the components of the tasks (i.e., the elements of the processor sets for each task) are independent: they do not require execution at the same time, and no order of execution is specified. In this setting, the overall makespan is the objective function.

In graph theoretical terms, an instance of the MULTIPROC problem can be modeled as a hypergraph, and finding a schedule of minimum makespan amounts to finding a semi-matching in the hypergraph, where the matching hyperedges are to be disjoint when restricted to a subset of vertices. As far as we know, this problem has not been considered before by the graph theory community.

We also consider a simplified variant of the problem, SINGLEPROC. Here, each task is sequential, and it can be scheduled on a single processor among a set of possible ones (with corresponding execution times). In this case, the hypergraph is in fact a bipartite graph, and the problem consists therefore of finding a semi-matching in a bipartite graph. This has been studied intensively in the case of unweighted bipartite graphs, i.e., the problem SINGLEPROC-UNIT with unit-weight tasks: several polynomial-time algorithms were pro-

posed [9, 13, 21, 22]. The weighted version is NP-complete [23]. Our intention in studying SingleProc at the beginning was to gauge the heuristics for MultiProc in a simplified setting by comparing their results with an exact solution for SingleProc-Unit.

Our contribution is twofold, and stands at the crossroad between scheduling and graph theory.

- 1. For the SingleProc-Unit problem, we propose an exact polynomial-time algorithm that is efficient in practice and conceptually very simple. We describe heuristics for this problem that will then be extended for MultiProc. We examine those heuristics theoretically (we build examples in which the heuristics are as far as possible from the optimal), and practically (by comparing them to the optimal solution). Practical experiments are done to validate the interest of the ideas used in these heuristics for the MultiProc problem.
- 2. The core of the paper resides in the study of the MULTIPROC problem. First, we prove that it is NP-complete even in the unweighted case (while SINGLEPROC-UNIT was solvable in polynomial time). Moreover, we show that for all $\epsilon > 0$, there is no (2ϵ) -approximation algorithm unless P=NP. We design a set of heuristics, building upon the SINGLEPROC-UNIT heuristics. We describe a lower bound to assess the performance of the heuristics for MULTIPROC, and present exhaustive simulation results.

The remainder of this paper is organized as follows. We start with a formal description of the optimization problems and a summary of some related work in Section 2. We then prove in Section 3 that the general problem MULTIPROC is NP-complete, even with unit-weight tasks. We design an optimal algorithm for SingleProc-Unit and several greedy algorithms of low complexity for the two variants of the problem in Section 4, and then assess the performance of these heuristics in Section 5. Finally, we conclude in Section 6 with a summary and plans on further investigations of the addressed problems.

2 Framework

We consider the problem of scheduling n independent tasks onto a set of p processors, with the objective of minimizing the makespan, i.e., the maximum load of a processor. Let T_1, \ldots, T_n be the set of tasks, and let P_1, \ldots, P_p be the set of processors.

For $1 \leq i \leq n$, task T_i is subject to resource constraints: it can be executed only on some of the processors, and possibly in parallel on several processors (parallel task). There is therefore a set of possible configurations for each task, e.g., task T_1 can be processed either on processor P_1 , or concurrently on processors P_2 and P_3 . Let S_i be the different configurations for T_i , i.e., the collection of sets of processors on which T_i can be executed. Back to our example, $S_1 = \{\{P_1\}, \{P_2, P_3\}\}$. Task T_i is executed on a set of processors $alloc(i) \in S_i$, and it takes a time $w_i^{alloc(i)}$ on each of the processors $P_u \in alloc(i)$. The processing can be done at different time steps on the processors of alloc(i), since the task is executed in parallel, and we assume that the different parts of the task are independent, similarly to the concurrent job shop problem. The goal is

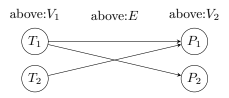


Figure 1: A sample bipartite graph for SingleProc-Unit.

to find a mapping of tasks to processors, i.e., decide the set alloc(i) of processors on which T_i is executed, for $1 \le i \le n$.

We define the load l(u) of processor P_u as its execution time:

$$l(u) = \sum_{i \mid P_u \in alloc(i)} w_i^{alloc(i)}.$$

The goal is to complete all tasks as soon as possible, i.e., minimize the $makespan\ M = \max_{1 \leq u \leq p} l(u)$. We consider several problem variants in the following, and describe the problems in the graph and hypergraph formalisms, so that the scheduling problem amounts to finding a semi-matching.

2.1 With a single processor

In some cases, tasks cannot be executed in parallel, and S_i is just a set of processors on which T_i can be executed, instead of a *set of sets*. This problem is called SINGLEPROC. This amounts to finding a semi-matching in a bipartite graph. We recall some graph definitions below to ease the description.

In a bipartite graph $G = (V_1 \cup V_2, E)$, the vertex sets V_1 and V_2 are disjoint and for all edges in E, one of the endpoints belongs to V_1 and the other belongs to V_2 . In our problem, V_1 is the set of tasks, V_2 is the set of processors, and an edge $e = (T_i, P_u) \in E$ between a task $T_i \in V_1$ and a processor $P_u \in V_2$ means that P_u is in the set S_i (see Fig. 1, where $S_1 = \{P_1, P_2\}$ and $S_2 = \{P_1\}$). We use d_v to refer to the number of neighbors of a vertex $v \in V_1 \cup V_2$. Moreover, we can add weights to the edges, that correspond to execution times: $w(e) = w(T_i, P_u) = w_i^{P_u}$.

Given a bipartite graph $G = (V_1 \cup V_2, E)$, a semi-matching \mathcal{M} in G is a set of edges $\mathcal{M} \subseteq E$ such that each vertex $v \in V_1$ is incident to exactly one edge in \mathcal{M} , i.e., it corresponds to the allocation function alloc(i). Given a semi-matching \mathcal{M} , the load l(u) of $u \in V_2$ is the sum of the weights of the edges in \mathcal{M} incident on u. The objective is to find a semi-matching \mathcal{M} such that $\max_{u \in V_2} l(u)$ is minimized.

This SINGLEPROC problem was shown to be NP-complete [23] (see also a related note [16]) by reduction from the *Minimum Makespan Scheduling Problem on Identical Machines*, which differs from SINGLEPROC in the fact that the tasks can be run on any machine (i.e., no resource constraints). It was also noted there that SINGLEPROC can be reduced to the *Minimum Makespan Scheduling Problem on Unrelated Machines*, a more general formulation where the tasks can vary in execution time on different processors. For this problem, a 2-approximation algorithm was given by Graham et al. [12], which was subsequently improved to $2 - \frac{1}{p}$ by Shchepin and Vakhania [26].

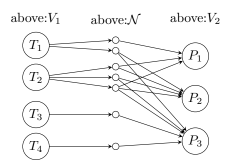


Figure 2: A sample hypergraph for MULTIPROC.

We consider also the unweighted version, that amounts to having $w_i^{P_u} = 1$ for $1 \le i \le n$ and $P_u \in \mathcal{S}_i$. This corresponds to unit tasks, and the problem is called SingleProc-Unit. This simpler instance of the problem can be solved via bipartite graph matching algorithms in polynomial time, as shown in [13]. In Section 4, we design several heuristics of low complexity for this SingleProc-Unit problem, since this allows us to compare the heuristics with the optimal solution in a reasonable time. Heuristics are then extended to solve the most general problem, that we detail below.

2.2 With multiple processors

The general problem is called MULTIPROC, and S_i is now a set of *sets of processors*. Back to the graph theory, this problem can then be seen as a matching problem in hypergraphs. A hypergraph $\mathcal{H} = (V, \mathcal{N})$ consists of a set of vertices V and a set of hyperedges \mathcal{N} . Each hyperedge is a subset of vertices.

A MULTIPROC problem instance can be modeled as a bipartite hypergraph $\mathcal{H} = (V, \mathcal{N})$ of the following form: the vertex set is bipartite $(V = V_1 \cup V_2, V_1 \cap V_2 = \emptyset)$, and each hyperedge $h \in \mathcal{N}$ satisfies $|V_1 \cap h| = 1$, i.e., one single task $T_i \in V_1$ is associated to a set of processors in V_2 through an hyperedge. In the example of Fig. 2, tasks T_3 and T_4 have only one configuration (they are in a single hyperedge), and must therefore be executed on P_3 . Tasks T_1 and T_2 can be executed in parallel and have the choice between several configurations. For example, T_1 can be executed by P_1 sequentially or by P_2 and P_3 collectively.

The problem now amounts to finding a semi-matching in a hypergraph, i.e., a set of hyperedges $\mathcal{M} \subseteq \mathcal{N}$ where the hyperedges in \mathcal{M} are disjoint on the vertices in V_1 . Thus, for all $T_i \in V_1$, there must be exactly one hyperedge h_i in \mathcal{M} such that $h_i \cap V_1 = T_i$. We then define $alloc(i) = h_i \cap V_2$. The processor load l(u) of $u \in V_2$ is equal to the sum of the weights of hyperedges incident on u in \mathcal{M} . For the ease of notations, we let $w_h = w_{h \cap V_1}^{h \cap V_2}$ be the weight associated to hyperedge $h \in \mathcal{N}$. We use d_v to denote the number of hyperedges containing the vertex $v \in V_1$.

The unweighted version of this problem, where all weights are 1, is called MULTIPROC-UNIT. While it is possible to solve SINGLEPROC-UNIT in polynomial time, MULTIPROC-UNIT turns out to be NP-complete (see Section 3).

2.3 Some related work

The problem SINGLEPROC is well known and widely studied (see a recent survey [18], the references therein, and a more recent study [17] dealing with online and offline algorithms for different processor set restrictions).

The problem MULTIPROC is a variant of scheduling with processing set restrictions [18] which is referred to as $P|set_j|C_{\max}$ by Drozdowski [7]. In heterogeneous computer systems, restrictions apply to task to processing unit assignments, as not all processors could be capable of meeting the task requirements. Furthermore, each processing unit could be a set of (sometimes hierarchical [18, p.260]) machines that collectively execute the job.

A more immediate problem for MULTIPROC is also known as scheduling independent tasks with multiple modes. In this setting, each task has a possible set of modes, where each mode is a set of resources simultaneously required by the task for a period of time [4]. The aim is to find a non-preemptive schedule of task executions in the decided modes so that the resource requirements at a time period do not exceed the available ones, and the makespan is minimized. A set of reasonable heuristics for this problem proceeds in two steps (see the discussions in a recent paper [5]), where the first step's goal can be formulated as MULTIPROC problem. In other words, the MULTIPROC problem arises as a subproblem in multimode scheduling problems.

The two problems are also related to scheduling problems with setup times/costs, where the setup cost occurs before a job is processed at a machine [2]. Our formulation in MULTIPROC asks for the minimization of the maximum setup cost across machines, where the tasks are splittable (Aubry et al. [3] discuss such a setting and propose a mixed integer linear program formulation for minimizing the total setup cost under load balance constraints with preemption) and the setup costs are sequence-independent [2].

3 NP-completeness of Multiproc-Unit

It was shown that SINGLEPROC is NP-complete [23], while SINGLEPROC-UNIT is solvable in polynomial time [13]. We show that MULTIPROC is NP-complete even for the unweighted version MULTIPROC-UNIT. Moreover, the reduction implies approximation hardness.

Theorem 1. The problem MULTIPROC-UNIT is NP-complete, and for all $\epsilon > 0$, there is no $(2 - \epsilon)$ -approximation algorithm unless P = NP.

Proof. We consider the associated decision problem: given an instance of MULTIPROC-UNIT and a bound on the makespan D, is there a solution of makespan not larger than D? This problem is obviously in NP, since the makespan can be computed in polynomial time, given an assignment of tasks to processors.

To establish the completeness, we use a reduction from Exact Cover by 3-Sets (X3C) [10, p. 53]. We consider an instance \mathcal{I}_1 of X3C: given a finite set X of elements where |X| = 3q and a collection C of 3-element subsets of X, does C contain an exact cover $C' \subseteq C$ such that every element of X occurs in one member of C'.

We build an instance \mathcal{I}_2 of MULTIPROC-UNIT: the set of elements of \mathcal{I}_1 are the processors, i.e., the vertex set V_2 in the hypergraph formulation. There are

q tasks to be mapped on these 3q processors, i.e., $|V_1| = q$. Each of these tasks can be mapped onto the sets of processors corresponding to the collection C, i.e., $S_i = C$ for $1 \le i \le n$. Moreover, we set the deadline D = 1.

Clearly, the size of \mathcal{I}_2 is polynomial in the size of \mathcal{I}_1 . We show that \mathcal{I}_1 has a solution if and only if \mathcal{I}_2 does.

If \mathcal{I}_1 has a solution, i.e., there is an exact cover, then we assign each task to the set of processors corresponding to one member of C'. Each processor is therefore processing exactly one task, and the makespan is $1 \leq D$, therefore \mathcal{I}_2 has a solution.

Suppose now that \mathcal{I}_2 has a solution. Since the makespan is at most 1, each processor can process at most one task, and since each task is executed on three distinct processors by construction of \mathcal{S}_i , the allocation of \mathcal{I}_2 forms a cover for \mathcal{I}_1 , and hence the result.

There remains to prove the inapproximability result. Let us assume that there is a $(2 - \epsilon)$ -approximation algorithm of MULTIPROC-UNIT, with $\epsilon > 0$. Then, we use this algorithm to solve instance \mathcal{I}_2 , hence obtaining a makespan $M \leq (2 - \epsilon)M_{opt}$, where M_{opt} is the optimal makespan. Since $M_{opt} = 1$, we obtain M < 2, and hence M = 1 since all weights are unit. The algorithm has therefore found an optimal solution, that corresponds to a cover, in polynomial time. This cannot hold unless P=NP.

Since the MULTIPROC problem is NP-complete even with unit weights (MULTIPROC-UNIT), we propose efficient heuristics to tackle it. We design similar heuristics also for the simpler version of the problem SINGLEPROC-UNIT, since we are then able to compare heuristics to the optimal solution.

4 Algorithms

The organization of this section is from the simpler problem SINGLEPROC-UNIT to our main problem MULTIPROC. We propose an exact polynomial-time algorithm for SINGLEPROC-UNIT first. We then discuss heuristics for SINGLEPROC-UNIT as they are simplified versions of the proposed heuristics for MULTIPROC. We also discuss a lower bound for MULTIPROC. The exact algorithm for SINGLEPROC-UNIT and the lower bound for MULTIPROC are developed to obtain baseline values for evaluating the proposed heuristics.

4.1 Exact algorithm for SingleProc-Unit

As shown by Harvey et al. [13], SingleProc-Unit can be solved using some modified versions of the standard matching algorithms in $O(|V_1||E|)$ time. We propose a conceptually simpler algorithm by making use of the standard matching algorithms as a black box (see a relatively recent survey [8] on augmenting-path based ones, and two other studies [11, 14] on push-relabel based ones). Assume a deadline D=1 and run the push-relabel algorithm on G. If a perfect matching is found, we have found a schedule of makespan 1, hence an optimal schedule. Otherwise, increase D by 1 and run the push-relabel algorithm on G_D , where G_D is identical to G except that it contains a total of D copies of each vertex in $u \in V_2$, each having the same neighborhood as the original vertex u. Repeat this process until a matching covering all the task vertices is found, at which time D is equivalent to the optimal makespan.

```
Input: A bipartite graph G = (V_1 \cup V_2, E)
Output: A matching \mathcal{M} in G
1: for all v \in V_1 do
2: find an edge e = \{v, u\} \in E for which l(u) is minimum
3: \mathcal{M} \leftarrow \mathcal{M} + e
4: l(u) \leftarrow l(u) + 1
5: return \mathcal{M}
```

Algorithm 1: basic-greedy

At step D, the algorithm has a complexity of $O(\sqrt{|V_1|}|E|D)$, and there are M_{opt} steps, where M_{opt} is the optimal makespan. Hence the algorithm has a running time complexity of $O(\sqrt{|V_1|}|E||M_{opt}^2)$. Note that $M_{opt} \leq |V_1|$, since the worst case is when all tasks are mapped on the same machine. Note that a bisection search for optimal makespan would yield improved worst-case running time complexity (in practice the algorithm is very fast).

4.2 Greedy algorithms for SingleProc-Unit

4.2.1 Basic-greedy

The basic greedy algorithm is straightforward (see Algorithm 1). It loops through the tasks in V_1 and assigns each task $v \in V_1$ to a processor in the neighborhood of v that has the smallest current load. The running time is O(|E|). Even though this algorithm often performs reasonably well, there are some instances in which it performs poorly. A toy example with two tasks (on the left) and two processors is shown in Fig. 1. If T_1 is mapped to P_1 , with T_2 having a single choice, the basic-greedy algorithm can assign the two tasks to processor P_1 and reach a makespan of 2 (versus 1 for the optimal solution). We show below that the basic greedy algorithm does not have any approximation guarantee.

4.2.2 Sorted-greedy

We improve basic-greedy by sorting tasks by non-decreasing out-degrees. The idea is to schedule the tasks that have less freedom first, e.g., task T_2 in the example of Fig. 1. The only modification to Algorithm 1 consists in visiting the tasks according to a non-decreasing order of degrees (d_v) at line 1. Unfortunately, this sorted-greedy algorithm may also take wrong decisions. We show here an example where it is at a factor k from the optimal solution, for any k (this is an example also showing that basic-greedy can be arbitrarily far from the optimal).

Consider that there are 2^k-1 tasks to be mapped onto 2^k processors. For the ease of reading, tasks are named $T_i^{(\ell)}$, with $0 \le \ell \le k-1$, and $1 \le i \le 2^{k-1-\ell}$. Task $T_i^{(\ell)}$ can be placed either on processor P_i , or on processor $P_{i+2^{k-1-\ell}}$ (see Fig. 3, for k=3).

The optimal solution places $T_i^{(\ell)}$ on $P_{i+2^{k-1-\ell}}$, for $0 \leq \ell \leq k-1$, and $1 \leq i \leq 2^{k-1-\ell}$. There is only one task per processor, hence an optimal schedule has a makespan of 1. However, the *sorted-greedy* algorithm starts by placing

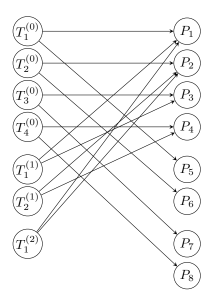


Figure 3: Example where basic-greedy and sorted-greedy obtain a makespan of k = 3, while the optimal makespan is 1.

tasks $T_i^{(0)}$ on processors P_1 through $P_{2^{k-1}}$, and then all processors that can be used for tasks $T_i^{(1)}$ have already a makespan of 1, it places them on processors P_1 through $P_{2^{k-2}}$, and so on. Finally, task $T_1^{(k-1)}$ is also mapped on P_1 , and processor P_1 achieves a makespan of k.

4.2.3 Double-sorted

From the example above, it seems better to also sort processors by increasing in-degrees, when there is a tie (i.e., edges leading to identical loads). This double-sorted algorithm is detailed as Algorithm 2.

```
Input: A bipartite graph G = (V_1 \cup V_2, E)
Output: A matching \mathcal{M} in G
 1: for all v \in V_1, sorted by non-decreasing out-degree do
        \min \leftarrow n
 2:
  3:
        \min \leftarrow n
        for all e = \{v, u\} \in E do
           if l(u) < \min or (l(u) = \min and d_u \le \min) then
              \min l \leftarrow l(u)
 6:
              mind \leftarrow d_u
 7:
              \min \leftarrow e
 8:
        \mathcal{M} \leftarrow \mathcal{M} + \text{mine}
 9:
        l(u) \leftarrow l(u) + 1
10:
11: return \mathcal{M}
```

Algorithm 2: double-sorted

This algorithm may also take wrong decisions. We can for instance generalize the example of Fig. 3 by adding extra tasks with higher degree and processors, so that all processors from the previous example have the same degree and the newcomers have a smaller degree. We illustrate this counter-example only for the case k=3, see Fig. 4. Tasks T_9 to T_{12} have a larger out-degree, so they are considered last. Then, since processors P_1 to P_8 have an identical in-degree of 3, double-sorted may take the same wrong decisions as sorted-greedy does, and obtain a makespan of 3 (while the optimal makespan is 1).

```
Input: A bipartite graph G = (V_1 \cup V_2, E)
Output: A matching \mathcal{M} in G
 1: for all u \in V_2 do
       o(u) \leftarrow 0
    for all v \in V_1 do
        for all \{v, u\} \in E do
          o(u) \leftarrow o(u) + 1/d_v
 6: for all v \in V_1, sorted by non-decreasing out-degree do
        find an edge e = \{v, u\} \in E for which o(u) is minimum
        \mathcal{M} \leftarrow \mathcal{M} + e
       o(u) \leftarrow o(u) + 1 - 1/d_v
 9:
        for all \{v, u\} \in E \setminus \{e\} do
10:
          o(u) \leftarrow o(u) - 1/d_v
11:
12: return \mathcal{M}
```

Algorithm 3: expected-greedy

4.2.4 Expected-greedy

As we have seen above, a weakness of the greedy algorithms is their inability to predict load that will arrive at a given vertex later during the execution of the algorithm. In this last greedy algorithm, we add a simple load prediction technique to *sorted-greedy* and adapt the strategy for the assignment of matching vertices. The resulting algorithm shown in Algorithm 3 is referred to as *expected-greedy*.

In this algorithm, o(u) represents the expected load of a vertex in V_2 (or processor). The values o(u) can be interpreted as the expected load a vertex u would have if the remaining matchings were performed uniformly at random. Actually, matching v to u can be seen as the collapse of the probability function. Consequently u, i.e., the possibility that was realized is assigned a probability of 1 and all other possibilities (i.e., neighbors of v) are assigned a probability of 0. The values of o are updated accordingly. When the algorithm terminates, the values o(u) are equivalent to actual loads l(u), and their maximum is equal to the makespan. One immediately verifies that the running time remains O(|E|).

On the example of Fig. 4, the values of o(u) differ since tasks T_9 to T_{12} are of degree 3, while the others are of degree 2. Therefore, expected-greedy places $T_1^{(0)}$ on P_5 , as in the optimal solution, and reaches the optimal makespan of 1. However, it is possible to modify the example so that expected-greedy also takes the wrong decisions, by having 16 tasks and 16 processors, and all tasks of out-degree 2, as in Fig. 5. Tasks T_9 to T_{16} can be assigned to their own

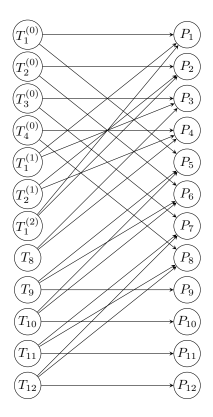


Figure 4: Example where double-sorted obtains a makespan of k=3, while the optimal solution is 1.

processor, and processors P_1 to P_8 have an in-degree 3. Therefore, the same wrong decisions will be taken by *expected greedy* and by *double-sorted* (i.e., tasks $T_i^{(0)}$ will be mapped on P_i , for $1 \le i \le 4$, and so on).

Note that those worst cases are however extremely unlikely in practical scenarios. Therefore, we study the quality offered by the heuristics and contrast it with the optimum values obtained from the exact algorithm. Results are discussed in Section 5.

4.3 Lower bound for MultiProc

Since we cannot compute the optimal solution for MULTIPROC as in the SINGLEPROC-UNIT case, we derive a lower bound so that we will be able to assess the performance of the heuristics, later in Section 5. For each task $T_i \in V_1$, we find a hyperedge $h_i \in \mathcal{N}$ such that $T_i \in h_i$, and $w_{h_i} \times |h_i \cap V_2|$ is minimum. We then define

$$time_i = \min_{h_i \in \mathcal{N}: T_i \in h_i} w_{h_i} \times |h_i \cap V_2|$$
.

Since all tasks must be executed on the p processors, the ideal case is when all processors achieve an identical load, equal to the makespan. Therefore, an obvious lower bound is one in which each task is in the best configuration in terms of the global load, leading to a total execution time of $time_i$, and where the load is equally shared between processors:

$$LB = \frac{1}{p} \sum_{1 \le i \le n} time_i . \tag{1}$$

4.4 Greedy algorithms for MultiProc

In this section, we aim at adapting the previous greedy algorithms for the MULTIPROC problem. On one hand, we need to account for the fact that a task may be executed on several processors. On the other hand, we need to account for task incurring different weights in different configurations, since we considered only the unweighted case previously.

We consider four heuristics for MULTIPROC, mainly based on the *sorted-greedy* algorithm for SingleProc-Unit. We exploit the hypergraph structure by introducing a new way of deciding which set of processors to choose (i.e., which hyperedge) for a given task (see the *vector* heuristics).

4.4.1 Sorted-greedy-hyp

Adapting sorted-greedy for MULTIPROC requires only minimal effort. Instead of choosing a neighbor of v having minimum current load, we chose a hyperedge h that minimizes $\max_{u \in h} l(u)$, among all hyperedges incident to v (see Algorithm 4). We also consider the weights in the new version of the algorithm, when computing the load l(u).

The running time now depends on the number of V_2 vertices in the hyperedges being inspected. In the worst case, the running time becomes $O(\sum_{h \in \mathcal{N}} |h|)$. Since bipartite graph semi-matching is a special case of hypergraph semi-matching.

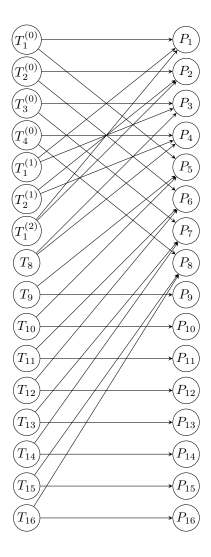


Figure 5: Example where *expected-greedy* obtains a makespan of k=3, while the optimal solution is 1. Note that all tasks are of degree 2, processors P_1 to P_8 are of degree 3, and processors P_8 to P_{16} are of degree 1.

```
Input: A hypergraph \mathcal{H} = (V_1, V_2, \mathcal{N})
Output: A matching \mathcal{M}
1: for all v \in V_1, sorted by non-decreasing out-degree do
2: find a hyperedge h: v \in h for which \max_{u \in h} l(u) is minimum.
3: \mathcal{M} \leftarrow \mathcal{M} + h
4: return \mathcal{M}
```

Algorithm 4: sorted-greedy-hyp

this algorithm also does not have an approximation guarantee. However, approximation of hypergraph semi-matching faces the additional difficulty that a single task can increase the load on multiple processors.

4.4.2 Expected-greedy-hyp

The expected-greedy algorithm can also be naturally extended to hypergraphs. In this case, when computing the values o(u), a hyperedge h containing v assigns its value of w_h/d_v to all vertices of V_2 contained in h, where w_h is the weight associated to the hyperedge, i.e., the execution time of task corresponding to v on each processor of the hyperedge. Other computations of $o(\cdot)$ are performed accordingly, as shown in Algorithm 5.

```
Input: A hypergraph \mathcal{H} = (V_1, V_2, \mathcal{N})
Output: A matching \mathcal{M}
 1: for all u \in V_2 do
        o(u) \leftarrow 0
 3: for all v \in V_1 do
        for all h \in \mathcal{N} : v \in h do
           for all u \in V_2 : u \in h do
 5:
              o(u) \leftarrow o(u) + w_h/d_v
 6:
 7: for all v \in V_1, sorted by non-decreasing out-degree do
        find a hyperedge h: v \in h for which \max_{u \in h} o(u) is minimum.
 8:
        \mathcal{M} \leftarrow \mathcal{M} + h
 9:
        for all u \in V_2 : u \in h do
10:
           o(u) \leftarrow o(u) + w_h - w_h/d_v
11:
        for all h' \in \mathcal{N} \setminus \{h\} : v \in h' do
12:
           for all u \in V_2 : u \in h' do
13:
14:
              o(u) \leftarrow o(u) - w_{h'}/d_v
15: return \mathcal{M}
```

Algorithm 5: expected-greedy-hyp

Similarly to sorted-greedy-hyp, this algorithm has a running time of $O\left(\sum_{h\in\mathcal{N}}|h|\right)$. Due to the updates of o, its running time will generally be in $O\left(\sum_{h\in\mathcal{N}}|h|\right)$ since hyperedges cannot be skipped during the updates. As discussed above, in the hypergraph case the additional information provided by the o values is far more important, since the potential mistakes the basic greedy algorithm can make are far greater.

4.4.3 Vector-greedy-hyp

Consider the sorted-greedy-hyp algorithm. At line 2, instead of looking at the loads increased by the hyperedge, we can look at the current bottleneck value, e.g., $\max_{u \in V_2} l(u)$. Clearly there will be many ties. In these case, we can favor the hyperedge that has the smaller second largest load. This tie breaking mechanism can be extended to check the load vectors sorted in descending order lexicographically. That is, among the hyperedges, choose the ones that yield the smallest largest $l(\cdot)$ value; among the alternatives choose the ones that yield the smallest second largest $l(\cdot)$ value and so on.

The worst case running time complexity of this heuristic is $O\left(\sum_{v\in V_1} d_v |V_2| \log |V_2| + \sum_{h\in \mathcal{N}} |h|\right)$, as checking the sorted load vectors lexicographically requires a sort operation (on $|V_2|$ items) for each hyperedge. Two improvements of this running time are immediate. First, if the hyperedges are unit weighted, then the sort operation can be done in linear time using bucket sort (or counting sort). Furthermore, one can keep the current load vector sorted as a list and then obtain the sorted load vector of an hyperedge by modifying the positions of modified loads. Here one can take advantage of the list already being sorted to reduce the sort operation as the merge of two lists (one of them is the processors in the hyperedge, the other is the remaining ones). This variant has the worst case time complexity of $O\left(\sum_{v\in V_1} d_v |V_2| + \sum_{h\in \mathcal{N}} |h|\right)$.

4.4.4 Expected-vector-greedy-hyp

This last heuristic is a combination of the expected and vector greedy heuristics on hypergraphs. There is one difficulty though. The current expected load vector of the processors contains contributions from each hyperedge associated with the task to be assigned. In order to differentiate between the hyperedges, one of them should be tentatively realized, and the others should be tentatively discarded so that the effect of each hyperedge can be measured. For a vertex with d_v hyperedges, this requires $O(d_v \sum_{v \in h} |h|)$ operations. The overall complexity of the algorithm with the list representation would be $O\left(\sum_{v \in V_1} d_v |V_2| + \sum_v d_v \sum_{v \in h} |h|\right)$. The first term would likely be the dominant one in reasonable settings but the overhead due to the second term (with respect to the vector-greedy-hyp) can be significant.

5 Experiments

We have implemented the proposed heuristics in Matlab and run the codes on a MacBook Pro equipped with a 2.7 GHz Intel Core i7 processor and 8GBytes of 1333 MHz DDR3 ram. We used an implementation of the push-relabel algorithm [14] provided in MatchMaker suite [8]. The reported running times of algorithms are in seconds and obtained by tic-toc routines of Matlab.

Below, we first explain how the data has been generated. Then we detail experimental results for SingleProc-Unit and MultiProc.

5.1 Data set

We have simulated the algorithms in settings where the number of tasks n is in $\{1280, 5120, 20480\}$, and the number of processors p is in 256, 1024, 4096; we did not test the cases where $n < 5 \times p$. The size of the problem instances is comparable to the numbers in recent simulation studies [25]. We implemented, in Matlab, two random bipartite graph generators [6] to create the structure of bipartite graphs and the hypergraphs used in the experiments. These generators are widely used in testing matching [8, 15] and semi-matching algorithms [13]. The generators take a number of parameters and create an instance of the problems at hand. In order to remove statistical bias, we create 10 instances

with a given parameter set and report the median of measurements in those 10 random instances for the given parameter set.

5.1.1 Bipartite graphs and SINGLEPROC-UNIT instances

The HiLo generator has been used in the cited resources to create bipartite graphs with $|V_1| = |V_2|$ where the resulting bipartite graph has a unique maximum matching with cardinality $|V_1|$. The associated task-processor bipartite graphs admit therefore a trivial makespan of one. We use this generator to create task-processor graphs with many more tasks than processors, hence possibly having many maximum matchings (with cardinality $|V_2|$). A little precision is necessary to describe the resulting random bipartite graphs resulting from this generator for the case $|V_1| \neq |V_2|$. There are four parameters to the HiLo bipartite graph generator: n, the number of vertices in V_1 ; p, the number of vertices in V_2 ; g the number of groups in which the vertices of V_1 and V_2 are divided; and d, a parameter used in defining the neighbors of a vertex in V_1 . Let x_i^j be the ith vertex in the jth vertex group of V_1 and v_2^j be the ith vertex in the ith vertex group of ith vertex ith vertex group of ith vertex ith vertex group of ith vertex group of ith vertex ith vertex ith vertex group of ith vertex group of ith vertex ith vertex group of ith vertex group of ith vertex ith vertex ith vertex group of ith vertex group of ith vertex ith vertex ith vertex group of ith

The FewgManyg generator [6] also has four parameters: n, the number of vertices in V_1 ; p, the number of vertices in V_2 ; g the number of groups in which the vertices of V_1 and V_2 are divided; and d, the average degree of a vertex in V_1 . First, the number of neighbors d_i of each vertex $x_i^j \in V_1$ is determined by sampling from a binomial distribution with mean d. Then for a vertex $x_i^j \in V_1$, d_i vertices are randomly chosen (without replacement) among the V_2 vertices in the j-1st to j+1st groups with wrap-around. In cases where d_i is bigger than 3p/g, vertices are chosen with replacement. In the original description of the FewgManyg generator, g=32 was used to refer to bipartite graph instances with few groups, and g=256 was used to refer to bipartite graph instances with few groups. We use FewgManyg(n, p, g, d) to denote a generic instance from this family created according to the four parameters.

In our study, we use all combinations of $d \in \{2, 5, 10\}$ and $g \in \{32, 128\}$ for the two generators to create instances of the problem SingleProc-Unit. We present detailed results for only d = 10, as this choice of d is more common for these generators [6, 13], and give a short summary of the results for other combinations in the appendix.

5.1.2 Hypergraphs and MULTIPROC instances

The hypergraph corresponding to the MULTIPROC instances can conveniently be represented by two bipartite graphs. The first one represents the connections between V_1 and \mathcal{N} , the second one represents the connections between \mathcal{N} and V_2 . By exploiting this fact, we create the instances for the problem MULTIPROC in two steps using five parameters: n, the number of tasks; p, the number of processors; d_v , the average degree of a task; d_h , a parameter used in defining the processor vertices in an hyperedge; and g the number of groups in which the processors and the hyperedges are divided.

In the first step, we choose the degrees of vertices in V_1 by random sampling a binomial distribution with mean d_v . Since the set of hyperedges of each vertex in V_1 are disjoint from the others, the degrees of vertices is enough to form the set of hyperedges. That is, we create $|\mathcal{N}| \approx |V_1|d_v$ hyperedges, each containing a unique vertex from V_1 . Then, in the second step, given the total number of hyperedges from the first step, we call $HiLo(|\mathcal{N}|, p, g, d_h)$ or $FewgManyg(|\mathcal{N}|, p, g, d_h)$ to add the processor vertices to each hyperedge.

In our study, we use all combinations of $d_v, d_h \in \{2, 5, 10\}$ and $g \in \{32, 128\}$ for the two generators to create instances of the problem MultiProc. In one set of data created with these parameters, we used unit hyperedge weights, essentially creating instances of MULTIPROC-UNIT. In the second set of experiments, we deterministically assigned the weight w_h to an hyperedge h as follows. Let $s_h = |h \cap V_2|$; then $w_h = \left\lceil \frac{\min_{j \in \mathcal{N}} \{s_j\} \times \max_{j \in \mathcal{N}} \{s_j\}}{s_h} \right\rceil$. These weights are related: if a task is assigned to more processors, its computation time gets smaller (as would be the case in most realistic settings). Note that the problem with related weights is also NP-complete, since all hyperedges have the same degree in the proof of Theorem 1. In the third data sets, weights are random so that we can double check the results; those experiments are reported in the appendix. Note that this results in almost linear speed-up for all configurations represented by hyperedges (with some overhead increasing with the degree of parallelism). In all combinations of d_v, d_h , the ranking of the heuristics according to the mean average quality were the same. We present detailed results for only $d_v = 5$ and $d_h = 10$, as this choice of d_h is more common with the use of the generators and $d_v = 5$ seems reasonable with respect to d_h . We give a short summary of the results for other combinations of d_v and d_h in the appendix.

5.2 Experimental results for SingleProc

We now present our experimental results. First, an overview over the test instances is shown in Table 1. Instances from the FewgManyg generator are named for number of groups with FG for few groups (g=32) and MG for many groups (g=128). Instances from the HiLo generator are named HLF for g=32 and HLM for g=128. The following two numbers denote the number of tasks and the number of processors, both in multiples of 256. Finally, the appended -SP denotes a SingleProc problem. In this table, |E| is the median value of the number of edges in 10 random instances for a given parameter set.

Results are shown in Table 2. The tables report the optimal makespan M computed with the exact algorithm of Section 4.1, and the ratio of each heuristic solution compared to the optimal makespan. The algorithms are those of Section 4.2: basic-greedy (BG), sorted-greedy (SG) double-sorted greedy (DG), and expected-greedy (EG).

For the FewgManyg generator, basic-greedy is the fastest algorithm but offers the lowest quality. Sorted-greedy significantly improves upon it while taking only marginally longer running time. The same is true for double-sorted greedy, but it offers no benefit in comparison to the standard sorted variant. Finally, it is observed that expected-greedy offers the best approximation but does so at the cost of an immense increase in running time, and its running time is closer to the exact algorithm's running time (which is implemented in C [14]) than to that of sorted-greedy. We note however that this is due to Matlab being an

Table 1: Random bipartite graph instances constructed by the FewgManyg and HiLo generators.

Instance	$ V_1 $	$ V_2 $	E
FG-5-1-SP	1280	256	12446
MG-5-1-SP	1280	256	5552
FG-20-1-SP	5120	256	49775
MG-20-1-SP	5120	256	22165
FG-20-4-SP	20480	256	198779
MG-20-4-SP	20480	256	88705
FG-80-1-SP	5120	1024	51280
MG-80-1-SP	5120	1024	49817
FG-80-4-SP	20480	1024	204874
MG-80-4-SP	20480	1024	198561
FG-80-16-SP	20480	4096	204325
MG-80-16-SP	20480	4096	204853
HLF-5-1-SP	1280	256	18396
HLM-5-1-SP	1280	256	4845
HLF-20-1-SP	5120	256	78876
HLM-20-1-SP	5120	256	20145
HLF-20-4-SP	20480	256	320796
HLM-20-4-SP	20480	256	81345
HLF-80-1-SP	5120	1024	107415
HLM-80-1-SP	5120	1024	74460
HLF-80-4-SP	20480	1024	440055
HLM-80-4-SP	20480	1024	319260
HLF-80-16-SP	20480	4096	440055
HLM-80-16-SP	20480	4096	434775

interpreted language, which cannot do all optimizations to each code. Thus, we can conclude that in this experiment, *sorted-greedy* offers the best combination of quality and speed with the current implementation. *Expected-greedy* should not be too slow with respect to *sorted-greedy* in an implementation using an imperative language. Hence it is also as preferable as the *sorted-greedy*.

For the HiLo generator, the overall picture is similar to that of the Fewg-Manyg instances (see the lower part of Table 2). The difference in approximation quality is more pronounced here. Expected-greedy now offers a significantly better approximation and because values of M are higher, its running time is much faster compared to the exact algorithm. If the running time is of significant importance, sorted-greedy can be considered as the method that offers the best tradeoff between the quality and running time.

We can conclude that *sorted-greedy* is essentially superior to *basic-greedy* and *double-sorted greedy*. Since the problem can be solved exactly in polynomial time, these algorithms are mostly useful in situations where running times is crucial. Therefore, for *expected-greedy* to be useful, it must be implemented in another programming language.

We remind the reader that we have tested with $d \in \{2, 5, 10\}$, and pre-

Table 2: Performance of the greedy algorithms for the FewgManyg and HiLo random bipartite graph instances with respect to the optimal value M. BG: basic-greedy; SG: sorted-greedy DG: double-sorted greedy; EG: expected-greedy.

Instance	M	$_{\mathrm{BG}}$	SG	$\overline{\mathrm{DG}}$	EG
FG-5-1-SP	5	1.20	1.20	1.20	1.20
MG-5-1-SP	5	1.20	1.20	1.20	1.20
FG-20-1-SP	20	1.25	1.05	1.05	1.05
MG-20-1-SP	20	1.25	1.15	1.15	1.05
FG-20-4-SP	80	1.25	1.02	1.04	1.01
MG-20-4-SP	80	1.26	1.14	1.14	1.01
FG-80-1-SP	5	1.40	1.20	1.20	1.20
MG-80-1-SP	5	1.20	1.20	1.20	1.20
FG-80-4-SP	20	1.30	1.05	1.05	1.05
MG-80-4-SP	20	1.25	1.05	1.05	1.05
FG-80-16-SP	5	1.40	1.20	1.20	1.20
MG-80-16-SP	5	1.40	1.20	1.20	1.20
Average quality	1	1.28	1.14	1.14	1.12
Average time (.s)	2.725	0.067	0.073	0.077	1.236
HLF-5-1-SP	5	1.80	1.40	1.40	1.20
HLM-5-1-SP	5	1.80	1.40	1.40	1.20
HLF-20-1-SP	20	1.95	1.50	1.50	1.25
HLM-20-1-SP	20	1.95	1.50	1.50	1.25
HLF-20-4-SP	80	1.99	1.50	1.50	1.25
HLM-20-4-SP	80	1.99	1.50	1.50	1.25
HLF-80-1-SP	12	2.00	1.58	1.58	1.25
HLM-80-1-SP	5	1.80	1.40	1.40	1.20
HLF-80-4-SP	56	2.00	1.50	1.50	1.25
HLM-80-4-SP	20	1.95	1.50	1.50	1.25
HLF-80-16-SP	47	2.00	1.51	1.51	1.26
HLM-80-16-SP	12	2.00	1.58	1.58	1.25
Average quality	1	1.94	1.49	1.49	1.24
Average time (.s)	6.842	0.058	0.067	0.071	1.083

Table 3: Random hypergraph instances constructed by the FewgManyg and HiLo generators.

Instance	$ V_1 $	$ V_2 $	$ \mathcal{N} $	$\sum_{h\in\mathcal{N}} h\cap V_2 $
FG-5-1-MP	1280	256	6368	61643
MG-5-1-MP	1280	256	6400	27705
FG-20-1-MP	5120	256	25504	248683
MG-20-1-MP	5120	256	25600	110817
FG-20-4-MP	5120	1024	25632	256459
MG-20-4-MP	5120	1024	25728	249483
FG-80-1-MP	20480	256	102336	993764
MG-80-1-MP	20480	256	102016	441810
FG-80-4-MP	20480	1024	102112	1021574
MG-80-4-MP	20480	1024	101888	994256
FG-80-16-MP	20480	4096	102176	1022141
MG-80-16-MP	20480	4096	102144	1027001
HLF-5-1-MP	1280	256	6368	99036
HLM-5-1-MP	1280	256	6400	25245
HLF-20-1-MP	5120	256	25472	400428
HLM-20-1-MP	5120	256	25600	101745
HLF-20-4-MP	5120	1024	26016	556479
HLM-20-4-MP	5120	1024	25600	400860
HLF-80-1-MP	20480	256	102752	1612548
HLM-80-1-MP	20480	256	102528	407235
HLF-80-4-MP	20480	1024	102848	2219679
HLM-80-4-MP	20480	1024	102656	1626900
HLF-80-16-MP	20480	4096	102592	2218293
HLM-80-16-MP	20480	4096	101888	2235585

sented only results for d=10. However, the ranking of the heuristics for the SingleProc-Unit problem were always the same as in Table 2, for the two families of the random bipartite graphs.

5.3 Experimental results for MULTIPROC

For MULTIPROC, we study weighted as well as unweighted instances for both types of random hypergraphs. Here, the appended -MP denotes a MULTIPROC problem. The instances otherwise follow the same naming conventions as the SINGLEPROC instances. Weighted instances are denoted by an appended -W, but are otherwise identical to their unweighted counterparts. The instances are listed in Table 3. Instead of indicating the number of edges, we now report the number of hyperedges $|\mathcal{N}|$ and the total number of vertices of V_2 contained in the hyperedges $\sum_{h \in \mathcal{N}} |h \cap V_2|$, where these two last values are the median of the ten random instances generated with the given parameter settings.

Since M is infeasible to compute via exact algorithm in this setting, the lower bound LB (1) described in Section 4.3 is given for comparison. The algorithms are sorted-greedy-hyp (SGH), vector-greedy-hyp (VGH), vector-greedy-hyp (VGH), vector-greedy-hyp (VGH), vector-greedy-hyp (VGH)

Table 4: Performance of the greedy algorithms for the unweighted FewgManyg and HiLo random hypergraphs with respect to the lower bound LB. SGH: sorted-greedy-hyp; VGH: vector-greedy-hyp; EGH: expected-greedy-hyp; EVG: expected-vector-greedy.

Instance	LB	SGH	VGH	EGH	EVG
FG-5-1-MP	34	1.43	1.33	1.39	1.37
MG-5-1-MP	17	1.43	1.32	1.43	1.38
FG-20-1-MP	135	1.34	1.24	1.32	1.30
MG-20-1-MP	70	1.40	1.27	1.38	1.38
FG-20-4-MP	34	1.41	1.30	1.39	1.37
MG-20-4-MP	34	1.45	1.34	1.39	1.39
FG-80-1-MP	539	1.30	1.22	1.27	1.27
MG-80-1-MP	280	1.39	1.26	1.37	1.36
FG-80-4-MP	136	1.35	1.24	1.32	1.32
MG-80-4-MP	135	1.34	1.25	1.31	1.31
FG-80-16-MP	34	1.42	1.30	1.39	1.39
MG-80-16-MP	34	1.42	1.30	1.39	1.39
Average quality		1.39	1.28	1.36	1.35
Average time (.s)		0.717	5.355	0.732	9.819
HLF-5-1-MP	68	1.18	1.17	1.17	1.18
HLM-5-1-MP	19	1.12	1.12	1.12	1.12
HLF-20-1-MP	291	1.1	1.1	1.1	1.1
HLM-20-1-MP	78	1.04	1.04	1.04	1.04
HLF-20-4-MP	99	2.84	2.84	2.84	2.84
HLM-20-4-MP	72	1.12	1.12	1.12	1.12
HLF-80-1-MP	1182	1.08	1.08	1.08	1.08
HLM-80-1-MP	313	1.03	1.03	1.03	1.03
HLF-80-4-MP	405	3.06	3.06	3.06	3.06
HLM-80-4-MP	307	1.05	1.05	1.05	1.05
HLF-80-16-MP	101	10.54	10.54	10.54	10.54
HLM-80-16-MP	105	2.7	2.69	2.69	2.69
Average quality		2.29	2.29	2.29	2.29
Average time (.s)		0.758	4.944	0.810	9.479

hyp (EGH) for hypergraphs, and expected-vector-greedy-hyp (EVG). The variants using lexicographic ordering of load vectors (VGH and EVG) are not implemented using the asymptotically faster algorithms discussed in the end of Secion 4.4.3. The quality of the four greedy algorithms is given as the ratio of the achieved makespan to the lower bound, where the ratio is taken as the median of the ten random instances. Note that the lower bound is very optimistic and may be far from the optimal solution.

Results for the unweighted instances are reported in Table 4. For the Fewg-Manyg generator, we immediately notice that vector-greedy-hyp provides better quality than the alternatives, but also takes significantly more time while sorted-greedy-hyp and expected-greedy-hyp are rather close. Interestingly, expected vector-greedy-hyp does not attain the good approximation quality of vector-greedy-hyp. For the unweighted HiLo instances, we again observe similar running times. However, all algorithms attain the same approximation quality, which means that neither the expected nor the vector strategy work here.

The weighted results are then reported in Table 5. Interestingly, the weighted FewgManyg results show a very different picture than the unweighted ones. Running times remain similar, but here the expected-greedy-hyp algorithm shows much better quality, while vector-greedy-hyp cannot improve upon sorted-greedy-hyp. Interestingly, expected-vector-greedy-hyp does improve upon the quality of expected-greedy-hyp, although at a steep cost in running time. For the weighted HiLo instances, similarly to the unweighted HiLo case, vector-greedy-hyp is at the same level as sorted-greedy-hyp, while the expected greedy algorithms show better approximation. This is consistent with their behavior in the weighted FewqManyg case.

From the above observation, we can conclude that the expected strategy is helpful in weighted instances. Expected-greedy-hyp showed better quality than sorted-greedy-hyp at the cost of only marginally higher running time. On the other hand, vector-greedy-hyp performed better only for unweighted FewgManyg and was significantly faster. Thus, the vector strategy is useful in weighted instances and is of limited use in unweighted instances. Using it to improve upon the quality of expected-greedy-hyp with expected-vector-greedy-hyp is worthwhile in order to obtain the best performance (see also the experiments with random hyperedge weights in Table 8, where expected-vector-greedy-hyp gives much better results).

Thus, the usefulness of the *vector* strategy is somewhat limited, although considering that these problems are NP-complete, using it to improve upon the quality of *expected-greedy-hyp* with *expected-vector-greedy-hyp* might still be worthwhile in order to obtain the best performance.

We remind the reader that we have tested with all combinations of $d_v, d_h \in \{2, 5, 10\}$. The ranking of the heuristics for the MULTIPROC-UNIT and MULTIPROC problem were always the same as in the Tables 4 and 5, for the two families of the random hypergraphs.

6 Conclusion

We have studied the problem of scheduling parallel tasks under resource constraints to minimize the makespan. We have formulated the problem in terms of bipartite graphs and hypergraphs, and shown that the scheduling problem

Table 5: Performance of the greedy algorithms for the weighted FewgManyg and HiLo random hypergraphs with respect to the lower bound LB. SGH: sorted-greedy-hyp; VGH: vector-greedy-hyp; EGH: expected-greedy-hyp; EVG: expected-vector-greedy.

Instance	LB	SGH	VGH	EGH	EVG
FG-5-1-MP-W	87	1.34	1.3	1.27	1.25
MG-5-1-MP-W	26	1.63	1.59	1.51	1.32
FG-20-1-MP-W	335	1.25	1.24	1.19	1.19
MG-20-1-MP-W	103	1.55	1.55	1.43	1.28
FG-20-4-MP-W	123	1.35	1.35	1.26	1.17
MG-20-4-MP-W	84	1.41	1.36	1.31	1.26
FG-80-1-MP-W	1406	1.19	1.18	1.15	1.15
MG-80-1-MP-W	413	1.54	1.54	1.43	1.27
FG-80-4-MP-W	549	1.24	1.24	1.12	1.11
MG-80-4-MP-W	381	1.22	1.21	1.17	1.15
FG-80-16-MP-W	141	1.36	1.35	1.24	1.17
MG-80-16-MP-W	141	1.35	1.37	1.29	1.17
Average quality		1.37	1.36	1.28	1.21
Average time (.s)		0.717	6.213	0.730	9.816
HLF-5-1-MP-W	80	1.25	1.24	1.12	1.02
HLM-5-1-MP-W	20	1.15	1.15	1.05	1.05
HLF-20-1-MP-W	320	1.17	1.17	1.05	1.02
HLM-20-1-MP-W	80	1.06	1.06	1.03	1.01
HLF-20-4-MP-W	110	2.93	2.93	2.61	2.60
HLM-20-4-MP-W	80	1.18	1.18	1.16	1.02
HLF-80-1-MP-W	1280	1.15	1.15	1.03	1.02
HLM-80-1-MP-W	320	1.04	1.04	1.01	1.01
HLF-80-4-MP-W	440	3.22	3.23	2.87	2.86
HLM-80-4-MP-W	320	1.07	1.06	1.03	1.01
HLF-80-16-MP-W	110	11.07	11.06	9.89	9.85
HLM-80-16-MP-W	110	2.66	2.66	2.57	2.57
Average quality		2.41	2.41	2.20	2.17
Average time (.s)		0.733	4.921	0.780	9.134

amounts to finding semi-matchings in the corresponding graph theoretical formulation. In the case of hypergraphs (i.e., parallel tasks), we have proved that the problem is NP-complete and that for all $\epsilon > 0$, there is no $(2 - \epsilon)$ -approximation algorithm unless P=NP.

For the simplest problem instance corresponding to semi-matchings in unweighted bipartite graphs, we have designed several greedy algorithms of low complexity, and from the simulation results, it turns out that performing a simple sort on the out-degree of the tasks (sorted-greedy algorithm) is very efficient and the execution is much faster than for the optimal algorithm. In addition, expected-greedy, which incorporates expected loads of processors, is shown to be more effective, albeit with an increase in the running time, in the current test platform.

We have extended the heuristics proposed for the bipartite graphs to the general case of weighted hypergraphs. While the adaptation of *sorted-greedy* still performs quite well in this case, the one with the load prediction technique (*expected-greedy-hyp*), obtains better results at the price of only a small increase in the execution time. We have also introduced two new heuristics based on a lexicographic ordering of load vectors instead of just minimizing the maximum load at each step, and the obtained solution is then shown to be even better.

As future work, we plan to further investigate the hypergraph problem, and to design new algorithms with guarantees. Indeed, even if the greedy algorithms perform quite well, their solution may be arbitrarily far from the optimal. Therefore, it seems challenging to obtain approximation algorithms for this problem. We also plan to implement the proposed heuristics in an imperative programming language to perform further tests.

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Table 6: Performance of the greedy algorithms for the SINGLEPROC-UNIT problem with FewgManyg and HiLo random bipartite graph instances, reported as average quality with respect to the optimal value for the random graph families $FewgManyg(\cdot,\cdot,\cdot,d)$ and $HiLo(\cdot,\cdot,\cdot,d)$ for differing d. BG: basic-greedy; SG: sorted-greedy DG: double-sorted greedy; EG: expected-greedy.

	BG	SG	$\overline{\mathrm{DG}}$	EG
d	Fewg	\overline{Manyg}	instand	ces
2	1.31	1.02	1.02	1.02
5	1.32	1.24	1.24	1.12
_10	1.28	1.14	1.14	1.12
\overline{d}	HiLo	instan	ces	
2	1.98	1.50	1.50	1.25
5	1.98	1.49	1.49	1.23
10	1.94	1.49	1.49	1.24

A Further experimental results

A.1 SINGLEPROC-UNIT

In Section 5.2, we have presented some detailed results for the SINGLEPROCUNIT problem. We have seen results with random bipartite graphs created using generators FewgManyg(n,p,g,d) and HiLo(n,p,g,d), where n, the number of tasks, is in $\{1280,5120,20480\}$; p, the number of processors, is in 256, 1024, 4096 (we did not test the cases where $n < 5 \times p$); d = 10; and the number of groups $g \in \{32,128\}$. In Table 6, we give average quality results for the heuristics with $d \in \{2,5,10\}$. In this table, a row shows the average performance of the heuristic with respect to the optimal value. In a sense each row corresponds to the row "Average quality" given in Table 2, which are repeated here for convenience. As seen from Table 6, whereas the quality of each heuristic changes with respect to the optimal value, the ranking of them remains the same: expected-greedy is the best, basic-greedy is the worst, and the two sorted variants are in the middle, without any difference between them.

A.2 MultiProc

In Section 5.3, we have presented some detailed results for the MULTIPROC-UNIT and MULTIPROC problems. We have seen results with random hypergraphs created using generators $FewgManyg(|\mathcal{N}|, p, g, d_h)$ and $HiLo(|\mathcal{N}|, p, g, d_h)$, where $|\mathcal{N}|$ is the number of hyperedges determined according to a random sampling of the degree of task vertices, n, under a binomial distribution with mean d_v , where the hyperedge weights were of the form $\left\lceil \frac{\min_{j \in \mathcal{N}} \{s_j\} \times \max_{j \in \mathcal{N}} \{s_j\}}{s_h} \right\rceil$. In the presented detailed experiments (see Tables 4 and 5), we had $d_v = 5$ and $d_h = 10$. In Table 7, we give average quality results for the heuristics with $d_v, d_h \in \{2, 5, 10\}$. In this table, a row shows the average performance of the heuristic with respect to the lower bound. In a sense each row corresponds to the row "Average quality" given in Tables 4 and 5, which are repeated here for

Table 7: Performance of the greedy algorithms for the MULTIPROC-UNIT (on the left) and MULTIPROC (on the right) problems with FewgManyg and HiLo random hypergraph instances, reported as average quality with respect to the lower bound value for differing d_v and d_h . SGH: sorted-greedy-hyp; VGH: vector-greedy-hyp; EGH: expected-greedy-hyp; EVG: expected-vector-greedy.

		MultiProc-Unit problem			Mi	ULTIPRO	c probl	em	
		SGH	VGH	EGH	EVG	SGH	VGH	EGH	EVG
d_v	d_h	FewgN	Ianyg in	stances		FewgA	Ianyg in	stances	
2	2	1.39	1.37	1.40	1.40	1.29	1.30	1.25	1.25
2	5	1.28	1.25	1.27	1.27	1.29	1.28	1.22	1.16
2	10	1.31	1.27	1.30	1.30	1.32	1.30	1.29	1.25
5	2	1.47	1.26	1.40	1.40	1.43	1.43	1.13	1.13
5	5	1.49	1.30	1.43	1.43	1.39	1.39	1.19	1.14
5	10	1.39	1.28	1.36	1.35	1.37	1.36	1.28	1.21
10	2	1.51	1.16	1.34	1.33	1.56	1.55	1.12	1.16
10	5	1.74	1.37	1.61	1.61	1.47	1.47	1.19	1.15
10	10	1.55	1.32	1.48	1.48	1.45	1.44	1.31	1.21
$\overline{d_v}$	d_h	HiLo i	nstances	S		HiLo instances			
2	2	6.78	6.78	6.78	6.78	6.92	6.91	6.68	6.66
2	5	3.59	3.59	3.59	3.59	3.65	3.65	3.52	3.51
2	10	2.25	2.25	2.24	2.24	2.30	2.30	2.23	2.19
5	2	7.03	7.02	7.02	7.02	7.36	7.36	6.67	6.65
5	5	3.72	3.72	3.71	3.71	3.87	3.87	3.52	3.50
5	10	2.29	2.29	2.29	2.29	2.41	2.41	2.20	2.17
10	2	7.45	7.45	7.45	7.45	8.05	8.05	6.67	6.65
10	5	3.93	3.93	3.93	3.93	4.22	4.22	3.51	3.50
_10	10	2.45	2.45	2.45	2.45	2.61	2.61	2.19	2.17

convenience. As seen in Table 7, the average quality of the heuristics changes, but their ranking remain the same in different problems and families.

As the hyperedge weights of the form $\left\lceil \frac{\min_{j \in \mathcal{N}}\{s_j\} \times \max_{j \in \mathcal{N}}\{s_j\}}{s_h} \right\rceil$ are special in that they are correlated with the hyperedge sizes, we designed further experiments with random hyperedge weights. In the first set of experiments, the hyperedge weights are uniformly distributed between 0.5 and 2 and in the second set of experiments they are uniformly distributed between 0.1 and 10. In these two cases the variations between the hyperedge weights are low and high. Table 8 shows the results of these experiments. The results again confirm that the ranking of the heuristics matches to the ranking stated before (matches to the order of the columns in Table 8), where the superiority of expected-vector-greedy (EVH) is more pronounced.

Table 8: Performance of the greedy algorithms for the MULTIPROC problem with FewgManyg and HiLo random hypergraph instances, reported as average quality with respect to the lower bound value for differing d_v and d_h with random hyperedge weights. SGH: sorted-greedy-hyp; VGH: vector-greedy-hyp; EGH: expected-greedy-hyp; EVG: expected-greedy-hyp; EVG: expected-greedy-hyp; EVG: expected-greedy-hyp; EVG: expected-greedy-hyp; expected-greedy-

		$w_h \in [0.5, 2]$					$w_h \in [0]$	[0.1, 10]		
		SGH	VGH	EGH	EVG	SGH	VGH	EGH	EVG	
d_v	d_h	FewgN	\overline{Ianyg} in	stances		FewgN	FewgManyg instances			
2	2	1.49	1.49	1.49	1.47	1.74	1.74	1.72	1.69	
2	5	1.41	1.41	1.39	1.35	1.61	1.61	1.57	1.50	
2	10	1.47	1.47	1.46	1.39	1.66	1.66	1.64	1.52	
5	2	1.90	1.90	1.78	1.59	2.81	2.81	2.57	2.03	
5	5	1.82	1.82	1.75	1.54	2.70	2.70	2.55	1.96	
5	10	1.79	1.79	1.74	1.49	2.76	2.76	2.65	1.88	
10	2	2.33	2.33	2.12	1.70	4.95	4.95	4.35	2.66	
10	5	2.29	2.29	2.12	1.70	4.71	4.71	4.27	2.57	
_10	10	2.18	2.18	2.08	1.59	4.91	4.91	4.61	2.47	
$\overline{d_v}$	d_h	HiLo i	nstances	3		HiLo instances				
2	2	4.63	4.63	4.61	4.33	5.26	5.26	5.23	4.61	
2	5	2.56	2.56	2.55	2.40	2.92	2.92	2.90	2.55	
2	10	1.77	1.77	1.76	1.65	2.02	2.02	2.00	1.75	
5	2	6.14	6.14	6.13	4.87	9.63	9.63	9.61	5.82	
5	5	3.40	3.40	3.38	2.68	5.29	5.29	5.27	3.20	
5	10	2.33	2.33	2.33	1.85	3.63	3.63	3.62	2.17	
10	2	7.59	7.59	7.57	4.99	17.92	17.92	17.85	6.74	
10	5	4.17	4.17	4.15	2.74	9.93	9.93	9.86	3.70	
10	10	2.87	2.87	2.86	1.89	6.74	6.74	6.73	2.50	

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