Replicated Data Placement for Uncertain Scheduling

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Abstract—Scheduling theory is a common tool to analyze the performance of parallel and distributed computing systems, such as their load balance. How to distribute the input data to be able to execute a set of tasks in a minimum amount of time can be modeled as a scheduling problem. Often these models assume that the computation time required for each task is known accurately. However in many practical case, only approximate values are available at the time of scheduling.

In this paper, we investigate how replicating the data required by the tasks can help coping with the inaccuracies of the processing times. In particular, we investigate the problem of scheduling independent tasks to optimize the makespan on a parallel system where the processing times of the tasks are only known up to a multiplicative factor. The problem is decomposed in two phases: a first offline phase where the data of the tasks are placed and a second online phase where the tasks are actually scheduled.

For this problem we investigate three different strategies, each allowing a different degree of replication of tasks: a) No Replication b) Replication everywhere and c) Replication in groups. We propose approximation algorithms and theoretical lower bound on achievable approximation ratios. This allows us to study the tradeoff between the number of replication and the guarantee on the makespan.

Keywords-Scheduling; Uncertainty; Robustness; Replication; Approximation Algorithms; Parallel System

I. INTRODUCTION

Parallel and distributed computing systems are often modeled using tasks that are processed simultaneously on different machines. Studying the balance of the load of the various component of the system is often key in understanding the performance one obtains in practice. A system typically schedules the set of tasks with the goal of optimizing the load balance (or makespan) of the system or some other metric. A key information these systems use to plan the execution is the time tasks will take to be processed. However, this information is typically not precisely known in practice: because the user can only make a wild guess on the runtime of her task [16], because prediction is hard in the general case [24], or because underlying models of a particular algorithm can only predict runtime within a

given range [7]. Whichever the reason is, not knowing accurately the processing time can significantly impact the performance obtained from the machine.

Though, the uncertainty of processing times is only a real problem if the tasks are pinned to particular computing units. If a task could be moved as the system sees fit without incurring an extra cost, the problem would be vastly alleviated. The problem is that, in practice, a task has to run on a particular machine because the input data for this task are only available on this machine. This is the case in most parallel and distributed systems and it is particularly true in out-of-core computing systems. For instance in out-of-core sparse linear algebra, executing a task where the data are not locally available would have a prohibitive overhead [28], [27].

One approach for dealing with the uncertainty of processing times is to build a robust schedule [2], [9], [6], that is, building a schedule that can naturally cope with variations in the processing times. These techniques often use sensitivity analysis to determine the robustness of the schedule. However, a better approach would be to be able to dynamically change the schedule.

The idea we are pursuing in this paper is to replicate the input data of the tasks onto multiple machines. This way, when the actual processing times of the tasks are too different from their estimations, the system will have some room to adapt at runtime. This is certainly feasible in practice as many system have more memory than the computation use. For instance, most Hadoop system replicates the data for the purpose of tolerating hardware faults [22]. And it has been shown that launching the same task multiple times can help cope with hardware differences [21] but increases resource usage. The cost of replicating the data might be amortized in many applications where the application will iterate over the data multiple times (e.g., in an iterative solver [27], [28]). In this paper, we answer the question "can data replication help cope with the uncertainty of processing time?" And the answer is that it can. (The technical contribution of this paper are summarized in Table 3.)

The remaining of the paper is organized as follows: we describe system model and notations in Section II. Related works are presented in Section III. Section IV investigates what can be done if the tasks can only be deployed on a single machine, we provide a guaranteed algorithm and provide a lower bound on the best guarantee that one can achieve in this case. Section V takes the reverse case and investigates what can be achieved if the data are replicated everywhere, leaving the maximum flexibility at runtime. We investigate one algorithm in this case and analyze its performance guarantee. Section VI investigates grouping machines together and replicating data in these groups as an intermediate between the previous two strategies and provide a guaranteed algorithm in that case. Section VII summarizes the various results we derived and study the tradeoff between performance guarantee and data replication.

II. PROBLEM DEFINITION

Let J be a set of n tasks which need to be scheduled onto a set M of m machines. Notice that scheduling theory commonly use interchangeably the terms "machines" and "processors" and the terms "jobs" and "tasks". We are considering the problem where the scheduler does not know the processing time p_i of task jprecisely before the task completes. But the scheduler has access to some estimation of the processing time $\tilde{p_i}$ of task j before making any scheduling decisions. We assume that the actual processing time p_i of a task j is within a multiplicative factor α of the estimated processing time $\tilde{p_i}$. (An inaccuracy of the throughput of the system leads to a multiplicative error and any interval of confidence of a runtime can be transformed into a value and a multiplicative error.) α is a quantity known to the scheduler. In other words the scheduler knows that:

$$\frac{\tilde{p}_j}{\alpha} \le p_j \le \alpha \tilde{p}_j \tag{1}$$

Assuming that the processing time of the tasks is known to be in an interval is reasonable in many application scenarios. One could derive bounds experimentally using machine learning techniques: for instance [26] used Support Vector Machines to predict the time it will take to run graph traversal algorithms. Models of runtime of algorithms can also be derived analytically: in [7] the authors provide bounds for the performance of various sparse linear algebra operations using only the size of the matrices and vectors involved.

Scheduling for this problem is performed in two phases. Phase 1 chooses where data are replicated using the estimated processing time \tilde{p}_j , for each of the task

j. The phase takes $\tilde{p_j}$, m and α as inputs and outputs sets of machines, $M_j \subseteq M$ where each task j can be scheduled. This phase is purely offline and corresponds to the operations performed to prepare the execution of the application.

Phase 2 takes the output of phase 1 as its input and maps each task j to a machine within the set of machines M_j . For each machine i, let $E_i \subseteq J$ be the set of tasks assigned to machine i. This phase chooses the actual schedule following an online semiclairvoyant process. Only the approximate processing time is known when a task is placed, but the scheduler can wait for a machine to become idle, to place the next one. Therefore, it can dynamically schedule the tasks and the actual processing time of the tasks are known once they complete.

The problem is to optimize the makespan, $C_{max} = \max_i \sum_{j \in E_i} p_j$ which is the completion time of the last task of the system. C^*_{max} denotes the optimal makespan of an instance of the problem (knowing the actual processing times). An offline algorithm is said to be a ρ -approximation algorithm (or to have an approximation ratio of ρ) if it guarantees for all the instances that $C_{max} \leq \rho C^*_{max}$. When the problem is online, we are talking about competitive ratios.

III. RELATED WORK

When $\alpha = 1$, the problem is exactly the classical independent tasks scheduling problem on identical machines, which is known to be NP-Hard [8]. We use Graham's List Scheduling (LS) [10] and Largest Processing Time (LPT) algorithms [11] to derive approximation ratios in different scenarios. The LS algorithm takes tasks one at a time and assigns them to the machine having the least load at that time. LS is a 2approximation algorithm and is widely used in online scheduling problems. LPT sorts the tasks in a nonincreasing order of processing time and assigns them one at a time in this order to the machine with the smallest current load. The LPT algorithm has a worst case approximation ratio of $\frac{4}{3} - \frac{1}{3m}$ in the offline setting. One can even obtain an arbitrarily good approximation algorithm for this problem by increasing its complexity with a dual approximation algorithm [12].

Based on various models for describing the uncertain input parameters, various methodologies can be used including reactive, stochastic, fuzzy and robust approach [14]. We are using the bounded uncertainty model which assumes that an input parameter has a value between a lower and upper bound. Wierman and Nuyens [23] introduce SMART, a classification to

understand size-based policies and draw analytic corelation between response time and estimated job size in single server problem. Robust approaches to deal with uncertainty are widely used on MapReduce systems [13], [20], in Hadoop [25], [22], on databases [15] and on web servers [3]. The HSFS and FLEX schedulers provide robustness in scheduling against uncertain job size [25], [17]. Cannon and Jeannot [2] analyzed the correlation between various metrics used to measure robustness and provided scheduling heuristics that optimizes both makespan and robustness for scheduling task graph on heterogeneous system.

Most of the work on robust scheduling use scenarios to structure the variability of uncertain parameters. Daniels and Kouvelis [5] used them to optimize the flow-time using a single machine. Davenport, Gefflot, and Bek analyzed slack based technique (adding extra idle time) to cope with uncertainty [6]. Gatto and Widmayer derive bounds on competitive ratio of Graham's online algorithm in scenarios where the processing times of tasks either increase or decrease arbitrarily due to perturbations [9]. These works considered augmenting or decreasing of job processing times as different problem scenarios that need to be optimized. We approach the problem using worst case analysis where some tasks may increase and some may decrease within the same schedule.

Data placement and replication methodologies are highly used in distributed systems including peer-to-peer and Grid systems to achieve effective data management and improve performance [4], [1], [18]. Tse [19] used selective replication of documents to increase the available bandwidth to serve files using web servers and study the problem through bi-criteria optimization techniques to maximize the quality of service and minimize the memory occupation.

IV. STRATEGY 1: NO REPLICATION

This section considers the situation where the data of each task is restricted to be on only one machine, *i.e.*, $\forall j, |M_j| = 1$. We have a set J of n tasks, and a set M of m machines. Let $f: J \mapsto M$ be a function that assigns each job to exactly one machine. The restriction that the data of each task is deployed on a single machine puts all the decisions in phase 1: each task can only be scheduled on one machine in phase 2.

A. Lower Bound

Theorem 1. When $|M_j|=1$, there is no online algorithm having competitive ratio better than $\frac{\alpha^2 m}{\alpha^2 + m - 1}$.

Proof: We use the adversary technique to prove the lower bound of this theorem. An adversary discloses

the input instance piece by piece. It analyzes the choices made by the algorithm to change the part of the instance that has not been disclosed yet. That way it can build an instance that maximizes the competitive ratio of the algorithm.

Let us consider an instance with λm tasks of equal estimated processing time $\forall j, \tilde{p_j}=1$. After phase 1, let i be the most loaded machine which has B tasks. Obviously, $B \geq \lambda$. In phase 2 the adversary increases the processing time of the tasks on machine i by a factor of α and changes the processing time of the other tasks by a factor of $\frac{1}{\alpha}$. So, $C_{max}=\alpha B$. C_{max}^* will be no worse than any feasible solution. In particular, the solution that distributes equally the tasks of size α and the tasks of size $\frac{1}{\alpha}$. Therefore $C_{max}^* \leq \frac{1}{\alpha} \lceil \frac{\lambda m - B}{m} \rceil + \alpha \lceil \frac{B}{m} \rceil$. Figure 1 depicts the online solution and the offline optimal. We have

$$\frac{C_{max}}{C_{max}^*} \geq \frac{\alpha^2 B}{\lceil \frac{\lambda m - B}{m} \rceil + \alpha^2 \lceil \frac{B}{m} \rceil}$$
 Since $\frac{\lambda m - B}{m} + 1 \geq \lceil \frac{\lambda m - B}{m} \rceil$ and $\frac{B}{m} + 1 \geq \lceil \frac{B}{m} \rceil$, we have
$$\frac{C_{max}}{C_{max}^*} \geq \frac{\alpha^2 B}{\frac{\lambda m - B}{m} + 1 + \alpha^2 \frac{B}{m} + \alpha^2}$$

From above expression it is clear that the smaller the value of B, the more the value of the expression decreases. So, any algorithm should minimize B to achieve better performance. For a schedule to be feasible the condition $B \geq \lambda$ must be satisfied. For $B = \lambda$ the value of $\frac{C_{max}}{C_{max}^*}$ is minimum and is equal to $\frac{\alpha^2 m \lambda}{\lambda(\alpha^2 + m - 1) + m(\alpha^2 + 1)}$. The adversary can maximize the ratio of the algorithm by arbitrarily increasing λ . When λ tends to ∞ , we have

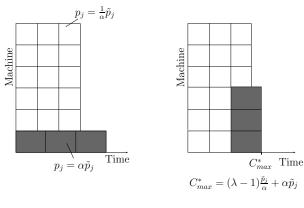
$$\frac{C_{max}}{C_{max}^*} \ge \frac{\alpha^2 m}{\alpha^2 + m - 1}$$

Corollary 1.1. When m goes to ∞ there is no online algorithm having competitive ratio better than α^2 .

B. Algorithm

We present the algorithm **LPT-No Choice**. In phase 1, the algorithm distributes the data of the tasks to the machine using their estimated processing times according to LPT [11]: The tasks are sorted in non-increasing order of their processing time and are greedily scheduled on the machine that minimizes the sum of $\tilde{p_j}$ of the tasks already allocated on that machine. Since there is no replication, there is no decision to take in phase 2.

The performance of the algorithm depends mostly on how much the actual processing times of the tasks differ



Online Solution

Offline Optimal

Figure 1: Instance constructed by the adversary in the proof of theorem 1 with $\lambda=3$ and m=6. In the online solution, the adversary increases the processing time of a task of the most loaded machine by a factor of α . If that information was available beforehand, an optimal offline algorithm could have distributed these longer tasks to other machines.

from their estimation. It also depends on the existence of a better arrangement if the actual processing times were known. The following theorem states the theoretical guarantee of the algorithm.

Theorem 2. The **LPT-No Choice** has a competitive ratio of $\frac{2\alpha^2 m}{2\alpha^2 + m - 1}$.

Proof: The algorithm assigns the tasks to machines based on their estimated processing times using LPT in phase 1. So, the planned makespan considering the estimated processing times of tasks, \tilde{C}_{max} have the following relation with the total estimated processing time, $\tilde{p_j}$ and estimated processing time of the task l that reaches \tilde{C}_{max} .

$$\tilde{C}_{max} \le \frac{\sum \tilde{p}_j + (m-1)\tilde{p}_l}{m} \tag{2}$$

The actual makespan of a schedule, C_{max} , obtained using the actual processing times of all the tasks, must be smaller than $C_{max} \leq \alpha \tilde{C}_{max}$ (thanks to Equation 1). We have following inequality:

$$C_{max} \le \alpha \tilde{C}_{max} \le \alpha \left(\frac{\sum \tilde{p}_j + (m-1)\tilde{p}_l}{m} \right)$$
 (3)

The worst case situation is when the tasks of the machine where the sum of estimated processing time is \tilde{C}_{max} sees the actual processing time of its tasks being α times larger than their estimate; meanwhile the processing time of the tasks on the rest of the machines

is $\frac{1}{\alpha}$ times their estimation. The argument behind this statement is that the greater the value of ratio $\frac{C_{max}}{\sum p_j}$, the worse the algorithm approximation ratio will be. So the total actual processing time is given by the following equation.

$$\sum p_j = \frac{\sum \tilde{p_j} - \tilde{C_{max}}}{\alpha} + \alpha \tilde{C}_{max} \tag{4}$$

Also the actual optimal makespan has the following constraint

$$C_{max}^* \ge \frac{\sum p_j}{m}$$

Substituting for $\sum p_j$, we have

$$mC_{max}^* \ge \frac{\sum \tilde{p_j} - \tilde{C_{max}}}{\alpha} + \alpha \tilde{C}_{max}$$

$$mC_{max}^* \ge \frac{\sum \tilde{p_j} - \left(\frac{\sum \tilde{p_j} + (m-1)\tilde{p_l}}{m}\right)}{\alpha} + C_{max}$$

$$mC_{max}^* \ge \frac{m-1}{\alpha m} \left(\sum \tilde{p_j} - \tilde{p_l}\right) + C_{max}$$

By the property of LPT, $\sum \tilde{p}_j - \tilde{p}_l \ge m(\tilde{C}_{max} - \tilde{p}_l)$, we have,

$$mC_{max}^* \ge \frac{m-1}{\alpha} \left(\tilde{C_{max}} - \tilde{p_l} \right) + C_{max}$$

All the instances where there is only one task per machine are always optimal. Therefore, we can restrict our analysis without loss of generality to instances with at least two tasks per machine. (Notice that in the original proof of Graham's LPT [11], an argument is made that all instances with two tasks per machine are optimal. However, the argument does not port in our case where only estimated processing times are known.) For at least two tasks on the machine that reaches \tilde{C}_{max} , the (estimated) processing time of last job is smaller than half the estimated makespan, $\tilde{p}_l \leq \frac{\tilde{C}_{max}}{2}$. Substituting this expression in the above equation, we have

$$mC_{max}^* \ge \frac{m-1}{\alpha} \left(\tilde{C}_{max} - \frac{\tilde{C}_{max}}{2} \right) + C_{max}$$

Using equation 3,

$$mC_{max}^* \ge \frac{m-1}{2\alpha} \frac{C_{max}}{\alpha} + C_{max}$$

$$mC_{max}^* \ge \left(\frac{m-1}{2\alpha^2} + 1\right) C_{max}$$

$$\frac{C_{max}}{C_{max}^*} \le \frac{2\alpha^2 m}{2\alpha^2 + m - 1}$$

V. STRATEGY 2: REPLICATE DATA EVERYWHERE

With this strategy, we put no restriction on phase 2. The tasks are replicated everywhere i.e. $\forall j, |M_j| = |M|$. We introduce the **LPT-No Restriction** which replicates the data of all the tasks on each machine in the first phase. In the second phase we simply use the Longest Processing Time algorithm (LPT) in an online fashion using the estimated processing time of the task. That is to say, the tasks are sorted in non-increasing order of their estimated processing time. Then the task are greedily allocated on the first machine that becomes available. Note that this is done in phase 2, the machine become available when the actual processing time of the task scheduled onto it elapses.

Lemma 3. Let l be the task that reaches C_{max} in the solution constructed by **LPT-No Restriction**. If there are at least two tasks on the machine that executes l in **LPT-No Restriction**, then $C_{max}^* \geq \frac{2}{\alpha^2} p_l$.

Proof: Since there are at least two tasks on the machine that executes l in **LPT-No Restriction**, there are at least m+1 tasks j such that $\tilde{p_j} \geq \tilde{p_l}$. Therefore, in any solution, at least one machine gets two tasks c and d, such that $\tilde{p_c} \geq \tilde{p_l}$ and $\tilde{p_d} \geq \tilde{p_l}$. C^*_{max} must be greater than sum of the processing time of these two tasks.

$$C_{max}^* \geq p_c + p_d$$

As the actual processing time of a task must be greater than $\frac{1}{\alpha}$ times of its estimated value, we have $p_c \geq \frac{1}{\alpha} \tilde{p_c}$ and $p_d \geq \frac{1}{\alpha} \tilde{p_d}$. Using this

$$C_{max}^* \geq \frac{1}{\alpha} \tilde{p}_c + \frac{1}{\alpha} \tilde{p}_d \geq \frac{2}{\alpha} \tilde{p}_l$$

Since, $\tilde{p}_l \geq \frac{1}{\alpha} p_l$, we have

$$C_{max}^* \geq \frac{2}{\alpha^2} p_l$$

Theorem 4. LPT-No Restriction has a competitive ratio of $\frac{C_{max}}{C_{max}^*} \leq 1 + (\frac{m-1}{m})\frac{\alpha^2}{2}$

Proof: The optimal makespan, C^*_{max} must be at least equal to the average load on the m machines. We have

$$C_{max}^* \ge \frac{\sum p_j}{m} \tag{5}$$

By the property of LPT (actually, it is a property of List Scheduling which LPT is a refinement of) the load on each machine i is greater than the load on the machine which reaches C_{max} before the last task l is

scheduled. So for each machine $i, C_{max} \leq \sum_{j \in E_i} p_j + p_l$ holds true. Summing for all the machines we have

$$mC_{max} \le \sum p_j + (m-1)p_l$$

$$C_{max} \le \frac{\sum p_j}{m} + \frac{(m-1)}{m} p_l \tag{6}$$

Using Equations 5 and 6, we have

$$\frac{C_{max}}{C_{max}^*} \le 1 + \frac{m-1}{m} \left(\frac{p_l}{C_{max}^*}\right)$$

Using Lemma 3, we have

$$\frac{C_{max}}{C_{max}^*} \le 1 + \left(\frac{m-1}{m}\right) \frac{\alpha^2}{2}$$

Graham's List Scheduling algorithm always has a competitive ratio of $2-\frac{1}{m}$. For $\alpha^2<2$, the **LPT-No Restriction** algorithm has a better approximation than List Scheduling. For $\alpha^2>2$, List Scheduling has better guarantee than the one expressed in Theorem 4. Since **LPT-No Restriction** is a variant of List Scheduling, the algorithm has a competitive ratio of $\min(1+\frac{m-1}{2m}\alpha^2,2-\frac{1}{m})$.

VI. STRATEGY 3: REPLICATION IN GROUPS

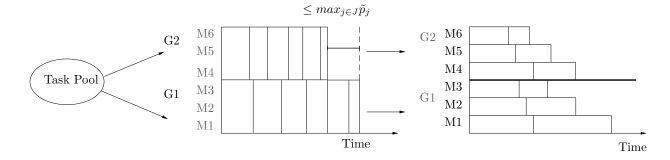
This strategy partitions the machines into k groups G1,G2...Gk. The size of each group is equal and have $\frac{m}{k}$ machines within each group. For the sake of simplicity, we assume that we will only use values of k such that k divides m. In the first phase, the data of each task is replicated on all the machines of one of the k groups, i.e., $\forall j, |M_j| = \frac{m}{k}$. In the second phase the tasks are scheduled within the group they are assigned to in first phase. Figure 2 shows the construction of the two phases.

We propose the **LS-Group** algorithm which is based on Graham's List Scheduling algorithm. In phase 1, we use List Scheduling to distribute the tasks to the k groups of machines. In phase 2 each task is scheduled to a particular machine within the group it was allocated in phase 1 using the online List Scheduling algorithm.

Theorem 5. With k groups, the competitive ratio of **LS-Group** is $\frac{k\alpha^2}{\alpha^2+k-1}(1+\frac{k-1}{m})+\frac{m-k}{m}$

Proof: We assume without loss of generality that C_{max} comes from group G1. C_{max}^* must be greater than the average of the loads on the machines.

$$C_{max}^* \ge \frac{\sum_{j \in J} p_j}{m}$$



Phase 1 Phase 2

Figure 2: An example of replication in groups with m=6, k=2. In phase 1, the data of the tasks are assigned to one of the groups. Phase 2 schedules each task to a machine within the group it was assigned.

 $\sum_{j\in J} p_j$ can be written as sum of load on G1 and load on rest of groups.

$$C_{max}^* \ge \frac{\sum_{j \in G1} p_j + \sum_{l=2}^k \sum_{j \in Gl} p_j}{m}$$
 (7)

As in phase 1 tasks are allocated to different groups using List Scheduling with the estimated processing times of the tasks, the (estimated) load difference between any two groups cannot be greater than the estimated value of largest task $max_{j\in J}\tilde{p}_{j}$. So, for any group $Gl \neq G1$, We have

$$\forall l \in \{2, 3, \dots, k\}, \left| \sum_{j \in G1} \tilde{p}_j - \sum_{j \in Gl} \tilde{p}_j \right| \le \max_{j \in J} \tilde{p}_j$$

Adding for all values of l leads to

$$\left| (k-1) \sum_{j \in G1} \tilde{p}_j - \sum_{l=2}^k \sum_{j \in Gl} \tilde{p}_j \right| \le (k-1) \max_{j \in J} \tilde{p}_j$$

Case 1: If $(k-1) \sum_{j \in G1} \tilde{p}_j > \sum_{l=2}^k \sum_{j \in Gl} \tilde{p}_j$.

$$\sum_{l=2}^{k} \sum_{j \in Gl} \tilde{p}_j \ge (k-1) \left(\sum_{j \in G1} \tilde{p}_j - \max_{j \in J} \tilde{p}_j \right)$$

As the actual processing time of the tasks can vary within a factor α and $\frac{1}{\alpha}$ of their estimated processing times, the following inequality holds

$$\alpha \sum_{l=2}^{k} \sum_{j \in Gl} p_j \ge (k-1) \left(\frac{1}{\alpha} \sum_{j \in G1} p_j - \alpha \max_{j \in J} p_j \right)$$

$$\sum_{l=2}^{k} \sum_{j \in Gl} p_j \ge (k-1) \left(\frac{1}{\alpha^2} \sum_{j \in G1} p_j - \max_{j \in J} p_j \right)$$

Phase 2 applies List Scheduling in the online mode. We assumed that C_{max} comes from G1. Using the guarantees of List Scheduling we can write,

$$C_{max} \le \frac{\sum_{j \in G1} p_j}{m/k} + \frac{m/k - 1}{m/k} p_{max}$$
 (9)

where p_{max} is actual processing time of longest task in G1.

From Equation 8 and 7, we derive

$$C_{max}^* \geq \frac{\sum_{j \in G1} p_j + (k-1) \left(\frac{1}{\alpha^2} \sum_{j \in G1} p_j - max_{j \in J} p_j\right)}{m}$$

$$\alpha^{2}(mC_{max}^{*} + (k-1)max_{j \in J}p_{j}) \ge (\alpha^{2} + k - 1) \sum_{j \in G_{1}} p_{j}$$

$$\frac{\alpha^2}{\alpha^2 + k - 1} \left(mC_{max}^* + (k - 1)max_{j \in J} p_j \right) \ge \sum_{j \in G1} p_j \tag{10}$$

Using 9 and 10, We have

$$\begin{split} C_{max} & \leq \frac{k\alpha^2}{\alpha^2 + k - 1} \left(C_{max}^* + \frac{k - 1}{m} max_{j \in J} p_j \right) \\ & + \frac{m/k - 1}{m/k} p_{max} \end{split}$$

As $C_{max}^* \geq max_{j \in J} p_j \geq p_{max}$, we have

$$\begin{split} C_{max} & \leq \frac{k\alpha^2}{\alpha^2 + k - 1} \left(C_{max}^* + \frac{k - 1}{m} C_{max}^* \right) \\ & + \frac{m - k}{m} C_{max}^* \end{split}$$

So, in case 1 the algorithm has a competitive ratio of,

$$\frac{C_{max}}{C_{max}^*} \le \frac{k\alpha^2}{\alpha^2 + k - 1} \left(1 + \frac{k - 1}{m} \right) + \frac{m - k}{m}$$

Case 2: If
$$(k-1)\sum_{j\in G1}\tilde{p}_j \leq \sum_{l=2}^k\sum_{j\in Gl}\tilde{p}_j$$
.

Since the processing times of the tasks can vary within a factor α and $\frac{1}{\alpha}$ of their estimated values, the expression for case 2 can be written as

$$\sum_{l=2}^{k} \sum_{j \in Gl} p_j \ge \frac{1}{\alpha^2} (k-1) \sum_{j \in Gl} p_j$$

Putting this value in Equation 7, we have

$$C_{max}^* \ge \frac{\alpha^2 + k - 1}{m\alpha^2} \sum_{j \in G1} p_j \tag{11}$$

Using Equations 9 and 11, and as $C^*_{max} \ge p_{max}$, we have

$$C_{max} \leq \frac{k\alpha^2}{\alpha^2 + k - 1} C_{max}^* + \frac{m - k}{m} C_{max}^*$$

So, in case 2 the algorithm has a competitive ratio of $\frac{k\alpha^2}{\alpha^2+k-1} + \frac{m-k}{m}$.

Clearly, the algorithm has a worst competitive ratio in case 1. So, the algorithm has a competitive approximation ratio of $\frac{C_{max}}{C_{max}^*} \leq \frac{k\alpha^2}{\alpha^2+k-1} \left(1+\frac{k-1}{m}\right) + \frac{m-k}{m}$.

LS-Group uses List Scheduling in both its phases. A LPT-based algorithm may have better guarantee. But without performing any replication, *i.e.* when k=m, the **LS-Group** algorithm has a competitive ratio almost equals to **LPT-No choice**'s when the number of machines m is large and the value of α is within practical range. This indicates an LPT-based algorithm for strategy 3 would likely not have a much more interesting guarantee.

VII. SUMMARY

Table 3 summarizes the results of this paper in terms of approximation ratios. Based on adversary technique, Theorem 1 states that there is no algorithm which can give performance better than $\frac{\alpha^2 m}{\alpha^2 + m - 1}$ for the model where no replication is allowed. **LPT-No Choice** is a $\frac{2\alpha^2 m}{2\alpha^2 + m - 1}$ -approximation that uses that strategy. For the second strategy that replicates the data of all tasks everywhere ($|M_j| = |M|$), **LPT-No Restriction** achieves a competitive ratio of $1 + (\frac{m-1}{m})\frac{\alpha^2}{2}$. The third strategy uses replication within k groups of size m/k (i.e., $|M_j| = m/k$). Using this strategy, the **LS-Group** algorithm has a competitive ratio of $\frac{k\alpha^2}{\alpha^2 + k - 1} \left(1 + \frac{k-1}{m}\right) + \frac{m-k}{m}$.

Of course, there is an inherent tradeoff between replicating data and obtaining good values for the makespan.

Replication	Approximation ratio
$ M_j =1$	$\frac{C_{max}}{C_{max}^*} \le \frac{2\alpha^2 m}{2\alpha^2 + m - 1} $ (Th. 2)
	No approximation better than $\frac{\alpha^2 m}{\alpha^2 + m - 1}$ (Th. 1)
$ M_j = m$	$\frac{C_{max}}{C_{max}^*} \le 1 + (\frac{m-1}{m}) \frac{\alpha^2}{2}$ (Th. 4) $\frac{C_{max}}{C^*} \le 2 - \frac{1}{m}$ [10]
$ M_j = \frac{m}{k}$	$\frac{C_{max}}{C_{max}^*} \le \frac{k\alpha^2}{\alpha^2 + k - 1} \left(1 + \frac{k - 1}{m} \right) + \frac{m - k}{m} $ (Th. 5)

Table 3: Summary of the contribution of this paper. Three proposed algorithms have guaranteed performance. One lower bound on approximability has been established.

To better understand the tradeoff we show in Figure 4 how the expressions of the guarantees (or impossibility) translate to actual values in an approximation ratio / replication space. We picked 3 values of α while keeping the number of machines fixed to m=210.

When $\alpha=1.1$, even with multiple groups **LS-Group** provides little improvement over **LPT-No Choice** in terms of makespan guarantee. However there is a significant gap between the guarantee of **LPT-No Choice** and the lower bound on possible approximation. When α is small, there is a significant improvement in using **LPT-No Restriction** over using **LS-Group** with only 1 group.

When α increases to 1.5, there is no more differences in the guarantees of **LS-Group** with 1 group and **LPT-No Restriction**. Also **LS-Group** provides many interesting intermediate solutions between deploying the data on a single machine and deploying them everywhere.

When $\alpha=2$, the range of the approximation ratios widens and the value of the lower bound increases. Now **LS-Group** is able to get a better approximation using less than 50 replications than can be guaranteed by deploying data on a single machine. Also, the approximation ratio quickly improves from more than 7.5 with the data being replicated on 1 machine to a ratio of less than 6 with only replicating the data on 3 machines.

Overall, when α is large, only few replication improve the performance significantly.

VIII. CONCLUSION AND FUTURE WORK

We studied the effect of uncertainty in the processing time of tasks when scheduling for parallel and distributed machines. In particular, we investigated how allowing tasks to execute on different machines can help dealing with not knowing the processing time of tasks accurately. We investigated three different replication strategies, provided an approximation algorithm in each

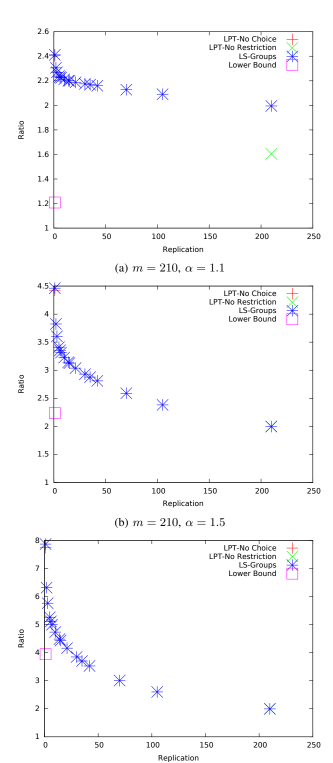


Figure 4: Ratio-Replication graph with m=210 and $\alpha \in \{1.1, 1.5, 2\}$.

(c) $m = 210, \alpha = 2$

case and a lower bound on the best achievable approximation in one of the case. The various strategies allow to trade the number of replication for a better guarantee. In particular, we obtained a better guarantee with fewer replication than can be achieved by putting the data of a task on only one machine. We observed that even a small amount of replications can improve the guarantee significantly. Therefore, we concluded that deploying the data on multiple machines can be an effective way of dealing with processing time uncertainties.

There are some open problems which can be explored further. Better lower bounds might help understanding the problem better: clearly when α is low, the problem is no different than the offline problem, and when it is large, the problem converges to the non-clairvoyant online problem. Having a clearer idea of where the boundary is will certainly prove useful in understanding how much can be gained using data replication. Also, while replicating data using groups of machines proved effective, more general replication policies can certainly lead to better guarantees.

Though, we assumed that every tasks were replicated the same amount of time. A more realistic model would introduce a cost of replicating a task (either global or per machine). This would allow to replicate only some critical tasks and limit memory usage.

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