# Critical path estimation in heterogeneous scheduling heuristics

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## 1 Introduction

The critical path of a project is defined as the longest sequence of constituent tasks that must be done in order to complete it [9, 8]. If we express the project in the form of a task graph, then the critical path is simply the longest—i.e., costliest—path through it. This is useful because the time it takes to execute the tasks on the critical path of the graph is therefore a lower bound on the makespan of any possible schedule for the project, no matter how many processors are available. In the context of a listing heuristic for scheduling the task graph on a set of parallel processors, a natural choice then is to prioritize all tasks according to the length of the critical path from that task to the end, the idea being that tasks with the greatest downward path length contribute most heavily to the makespan and should therefore be processed as soon as possible. This approach has a long and successful history in scheduling for homogeneous processors, and is in fact provably optimal for two-processor systems [4].

Now, in the homogeneous case, all tasks have the same cost on all processors, so that there is only one possible cost each task may take. In particular this means that the node weights in the task graph are fixed so, disregarding the edge weights for the moment, the longest path through the DAG is also fixed, no matter what schedule we ultimately follow. Therefore, when we refer to the critical path of the task graph it is clear what is meant. Unfortunately, the concept of the critical path is not so clearly defined for heterogeneous processing environments. The problem is, there are now multiple possible costs that each task may take—depending on the schedule—and likewise many different communication costs that may be incurred. In particular this means that the weights of the task DAG are not fixed and we cannot simply compute a longest path. Consider

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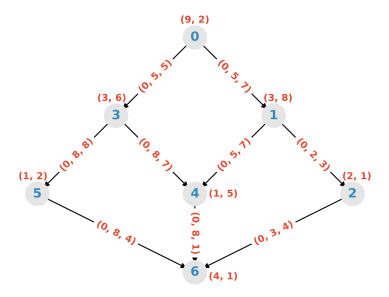


Figure 1: Simple task graph with costs on a two-processor target platform.

for example the simple DAG shown in Figure 1, where the labels represent all the possible weights each task/edge may take on a two-processor heterogeneous target platform; the red labels near the nodes represent the computation costs on processors P1 and P2 in the form  $(W_i^1, W_i^2)$ , while the edge labels represent the possible communication costs in the form  $(W_{ik}^{11} = W_{ik}^{22} = 0, W_{ik}^{12}, W_{ik}^{21})$ . How should the longest path through a graph like that in Figure 1 be defined?

The HEFT approach, as described in previous chapters, is to use average values over all sets of possible costs in order to fix the DAG weights and then compute the critical path in a standard dynamic programming manner. In particular, for each i = 1, ..., n, we compute a number  $u_i$ , called the upward rank, which purportedly represents the critical path length from task  $t_i$  to the end and is taken to be the task's priority. For example, for the graph in Figure 1, HEFT first (implicitly) converts the DAG to the fixed-cost one shown in Figure 2, and then calculates the task ranks like so:

$$\begin{aligned} u_6 &= 2.5, \\ u_5 &= 1.5 + 3 + 2.5 \\ &= 7, \\ u_4 &= 3 + 2.25 + 2.5 \\ &= 7.75, \\ u_2 &= 1.5 + 1.75 + 2.5 \\ &= 5.75, \\ u_3 &= 4.5 + \max\{4 + u_5, \ 3.75 + u_4\} \end{aligned}$$

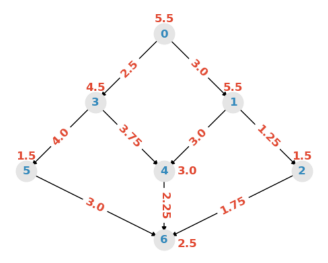


Figure 2: Fixed-cost counterpart of task DAG from Figure 1 which is implicitly used in HEFT.

$$= 4.5 + \max\{11, 11.5\}$$

$$= 16,$$

$$u_1 = 5.5 + \max\{3 + u_4, 1.25 + u_2\}$$

$$= 5.5 + \max\{10.75, 7\}$$

$$= 16.25,$$

$$u_0 = 5.5 + \max\{2.5 + u_3, 3 + u_1\}$$

$$= 5.5 + \max\{18.25, 19.25\}$$

$$= 24.75.$$

These ranks give a scheduling priority list of  $\{0, 1, 3, 4, 5, 2, 6\}$ . The resulting schedule length obtained by HEFT with these priorities is 22, as shown in Figure 3. Interestingly, we see that  $u_0$ , the rank of the single entry task, is greater than the schedule makespan—and therefore obviously not a lower bound on it. The question then is, what quantity do the  $u_i$  values actually represent? Now, because of the averaging of the costs, perhaps the most intuitive interpretation is that the  $u_i$  are in turn estimates of the average critical path lengths, taken over the set of all possible critical paths —but there is no robust mathematical justification for believing that this definition of the critical path is more useful than other possibilities.

Given this ambiguity, alternative ways to define the critical path for the ranking phase in HEFT have been considered before, most notably by Zhao and Sakellariou [14], who empirically compared the performance of HEFT when averages other than the mean (e.g., median, maximum, minimum) are used to compute upward (or downward) ranks. Their conclusions were that using the

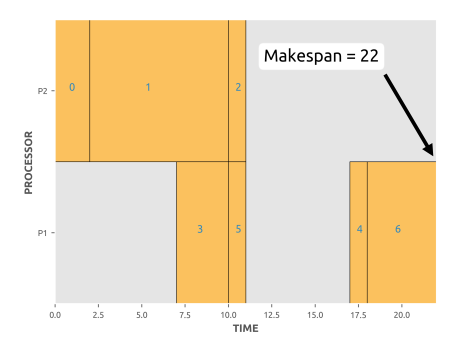


Figure 3: HEFT schedule for DAG in Figure 1.

mean is not clearly superior to other averages, although none of the other options they considered were consistently better. Indeed, perhaps the biggest takeaway from their investigation was that HEFT is very sensitive to how priorities are computed, with significant variation being seen for different graphs and target platforms. In this chapter we undertake a similar investigation with the aim of establishing if there are choices which do consistently outperform the standard upward ranking in HEFT. In addition, we consider how critical path estimates can be used to determine processor selection, as well as task prioritization, following the approach of the *Predict Earliest Finish Time* (PEFT) heuristic [1], and attempt to ascertain which definition leads to the best practical performance.

This will be an empirically-driven study, as is common in this area. To facilitate this investigation we created a software package that simulates heterogeneous scheduling problems, much like that described in the previous chapter, although not restricted to accelerated target platforms. As before, the (Python) source code for this simulator can be found on Github<sup>1</sup> and all of the results presented here can be re-run from scripts contained therein.

<sup>1</sup>https://github.com/mcsweeney90/critical-path-estimation

## 2 A universal lower bound

Functionally, the critical path is used in HEFT as a lower bound on the makespan, so that minimizing the critical path gives us the most scope to minimize the makespan, assuming we make good use of our parallel resources. With this in mind, there are many different ways we can define the critical path so that it gives a lower bound on the makespan of any possible schedule. The most straightforward approach would be to just set all weights to their minimal values but a tighter bound can be computed in the following manner. First, define  $\ell_i^a$  for all tasks  $t_i$  and processors  $p_a$  to be the critical path length from  $t_i$  to the end (inclusive), assuming that it is scheduled on processor  $p_a$ . These values can easily be computed recursively by setting  $\ell_i^a = W_i^a$ ,  $a = 1, \ldots, q$ , for all exit tasks then moving up the DAG and calculating

$$\ell_i^a = W_i^a + \max_{k \in S_i} \left( \min_{b=1,\dots,q} \left( \ell_k^b + W_{ik}^{ab} \right) \right), \quad a = 1,\dots,q,$$
 (1)

for all other tasks. Then, for each i = 1, ..., n,

$$\ell_i = \min_{a=1,\dots,q} \ell_i^a \tag{2}$$

gives a true lower bound on the remaining cost of any schedule once the execution of task  $t_i$  begins. These  $\ell_i$  values could be useful as alternative task priorities in HEFT, especially since the cost of computing all of the  $\ell_i^a$  in this manner is only  $O((m+n)q) \approx O(n^2q)$  so in particular is the same order as the usual HEFT prioritization phase.

For example, for the simple DAG shown in Figure 1, we find that the  $\ell_i$  values are as given in Table 1 (with the  $u_i$  included for comparison). Interestingly, we see that tasks 1 and 3 have the same lower bound (8 units) and the performance of the alternative ranks relative to the standard  $u_i$  sequence in HEFT depends on which is chosen to be scheduled first: if task 1, the priority list does not change so the schedule makespan is 22 units, but if task 3 is selected instead, the final schedule makespan is 20 units—i.e., smaller than the original—as illustrated in Figure 4. Of course, this is only one example: there is no mathematically valid reason to suppose that using the  $\ell_i$  sequence instead of  $u_i$  as the task ranks in HEFT will actually lead to superior performance in general. Still, it seems worthwhile to investigate this empirically, which we do in Section 4.

(Note that the lower bound on the critical path as defined here is very similar to the optimistic cost used in PEFT; this will be discussed further in Section 5.)

# 3 A stochastic interpretation

In this section we propose a family of alternative task ranking phases in HEFT based on the following interpretation of the standard ranking. Given the complex

Table 1: Upward ranks  $u_i$  and lower bounds  $\ell_i$  for the DAG in Figure 1.

| Task: | 0           | 1 | 2 | 3 | 4 | 5 | 6 |
|-------|-------------|---|---|---|---|---|---|
| · ·   | 24.75<br>16 |   |   |   |   |   |   |

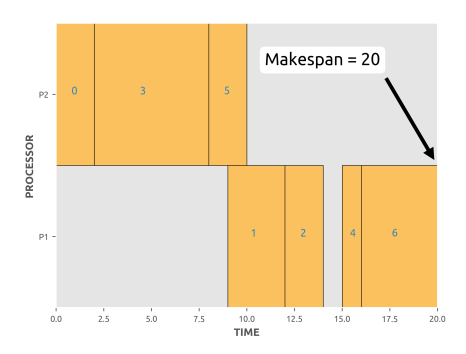


Figure 4: Alternative HEFT schedule for DAG in Figure 1.

interplay between the ranking and processor selection phases, it is impossible to predict exactly which values all DAG weights will take at runtime, at least without modifying the latter. Now, by using average values over all sets of possible costs, HEFT in some sense implicitly assumes that all members of each set are equally likely to be incurred. Conceptually, we can view this as an attempt to model the processor selection phase in order to predict which values the weights will assume. This model takes the form of a  $stochastic\ graph$ —i.e., all weights are assumed to be independent (discrete) random variables with associated probability mass functions (pmfs) given by the assumption of equal likelihood. More precisely, let  $m_i$  be the pmf corresponding to the task weight variable  $w_i$  and  $m_{ik}$  that for the edge weight  $w_{ik}$ , then

$$m_i(W_i^a) := \mathbb{P}[w_i = W_i^a] = \frac{1}{q}, \quad a = 1, \dots, q,$$

and

$$m_{ik}(W_{ik}^{ab}) = m_i(W_i^a) \cdot m_k(W_k^b)$$
$$= \frac{1}{q^2}, \quad \forall a, b.$$

It is important to note here that the model defined by these pmfs is clearly not accurate. In particular, all of the node and edge weight variables are independent of one another. This is induced by the averaging but does not reflect, for example, the fact that edge weights are fully determined once the node weights are realized. Still, as the saying goes, all models are wrong, some are just more useful than others; we attempt to establish exactly how useful this one is later in this chapter.

Note also that the expected values of the node and edge weight variables are given by

$$\mathbb{E}[w_i] = \sum_{a=1}^q W_i^a m_i(W_i^a) = \frac{1}{q} \sum_{a=1}^q W_i^a, \tag{3}$$

$$\mathbb{E}[w_{ik}] = \sum_{a=1}^{q} \sum_{b=1}^{q} W_{ik}^{ab} m_{ik}(W_{ik}^{ab}) = \frac{1}{q^2} \sum_{a,b} W_{ik}^{ab}.$$
 (4)

This means that  $\mathbb{E}[w_i] = \overline{w_i}$  and  $\mathbb{E}[w_{ik}] = \overline{w_{ik}}$ . So the computation of the upward ranks  $u_i$  in HEFT can instead be done by setting  $u_i = \mathbb{E}[w_i]$  for all exit tasks, then moving up the DAG and recursively computing

$$u_i = \mathbb{E}[w_i] + \max_{k \in S_i} \left( u_k + \mathbb{E}[w_{ik}] \right) \tag{5}$$

for all other tasks.

In summary, since all possible node and edge weights of a task graph G are known but their actual values at runtime aren't, one possible interpretation of the standard HEFT task prioritization phase is that critical path lengths in G are estimated through a two-step process:

- 1. An associated stochastic graph  $G_s$  is implicitly constructed with node and edge pmfs  $m_i$  and  $m_{ik}$  as defined above.
- 2. The numbers  $u_i$  are recursively computed for all tasks in  $G_s$  using (5), and taken as the critical path lengths from the corresponding tasks in G.

In the following two sections, we propose modifications of both steps so as to obtain different critical path estimates that may be used as task ranks in HEFT. The performance of these will then be evaluated through extensive numerical simulations in Section 4.

### 3.1 The critical path of $G_s$

Now, since all of its weights are RVs, the critical path of the stochastic graph  $G_s$  is clearly itself a random variable. But a natural question arises from the interpretation outlined in the previous section: what is the relationship between the sequence of numbers  $u_i$  as defined by (5) and the critical path of  $G_s$ ? In fact, it has long been known in the context of Program Evaluation and Review Technique (PERT) network analysis that the numbers  $u_i$  are lower bounds on the expected value of the critical path lengths of the stochastic DAG. This result dates back at least as far as Fulkerson [6], who referred to it as already being widely-known and gave a simple proof. This prompts another question: does using the actual expected values lead to superior performance in HEFT?

Unfortunately, computing the moments of the critical path length of a graph whose weights are discrete RVs was shown to be a #P-complete problem by Hagstrom [7]. This means that it is generally impractical to compute the true expected values. However, efficient methods which yield better approximations than the  $u_i$  numbers are known; we discuss examples in the following two sections.

#### 3.1.1 Monte Carlo sampling

Monte Carlo (MC) methods have a long history as a means of approximating the critical path distribution for PERT networks, dating back to at least the early 1960s [13]. The idea is to simulate the realization of all RVs (according to their pmfs) and then evaluate the critical path of the resulting deterministic graph. This is done repeatedly, giving a set of critical path instances whose empirical distribution function is guaranteed to converge to the true distribution by the Glivenko-Cantelli Theorem [2]. Furthermore, analytical results allow us to quantify the approximation error for any given the number of realizations—and therefore the number of realizations needed to reach a desired accuracy.

Table 2 illustrates how our estimates of the expected critical path lengths evolve as the number of realizations increases for the stochastic graph corresponding to that in Figure 1. The relevant  $u_i$  numbers are also included to show that they do indeed appear to be a lower bound on the values the Monte Carlo

Table 2: Monte Carlo estimates of critical path lengths for example graph.

| Task | $u_i$ | MC1 | MC10 | MC100 | MC1000 |
|------|-------|-----|------|-------|--------|
| 0    | 24.75 | 29  | 26.8 | 29.9  | 28.9   |
| 1    | 16.25 | 22  | 17.0 | 17.4  | 17.0   |
| 2    | 5.75  | 3   | 5.4  | 6.0   | 5.8    |
| 3    | 16    | 20  | 18.9 | 19.0  | 18.6   |
| 4    | 7.75  | 14  | 7.2  | 7.9   | 7.8    |
| 5    | 7     | 6   | 5.3  | 7.1   | 7.0    |
| 6    | 2.5   | 1   | 1.9  | 2.5   | 2.5    |

method appears to be converging toward. Note that the critical path length estimate for task 3 eventually exceeds that of task 1 so it has a higher priority if these estimates are taken as task ranks in HEFT; as noted in the previous section, interchanging these two tasks—and keeping the same ordering for all others—leads to a smaller schedule makespan (20 units) compared to the standard ranking (22 units). Of course, this is only one example, but it does serve to illustrate that in some cases a tighter bound on the critical path of  $G_s$  can be useful.

The downside of Monte Carlo sampling is its cost. While modern architectures are well-suited to this approach because of their parallelism, it still may be impractical in the context of a scheduling heuristic, especially when the DAG is large; we often found this to be the case for the examples discussed in Section 4. Hence in this report we typically only use the Monte Carlo method as a means of obtaining a reference solution; see, for example, the following section.

#### 3.1.2 Fulkerson's bound

Before introducing the alternative bounds on the critical path lengths proposed by Fulkerson, we first describe how the stochastic graph  $G_s$  can be expressed in an equivalent formulation with only edge weights. This step is not mathematically necessary but simply makes the elucidation much cleaner; it should be emphasized that all of the following still holds, with only minor adjustments, if this is not done. The most straightforward approach is to simply redefine the edge weights so that they also include the computation cost of the parent task and, if the child task is an exit, the computation cost of the child as well. More precisely, we define a new set of edge weight variables  $\tilde{w}_{ik}$  which take values

$$\tilde{W}_{ik}^{ab} := W_i^a + W_{ik}^{ab} + \delta_k W_k^b, \quad \forall a, b, i, k,$$

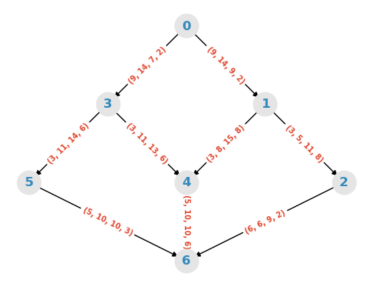


Figure 5: Edge weight-only equivalent of task DAG from Figure 1.

where  $\delta_k = 1$  if  $t_k$  is an exit task and zero otherwise. Figure 5 illustrates how the graph in Figure 1 would be transformed in this manner, where the edge labels are in the form  $(\tilde{W}_{ik}^{11}, \tilde{W}_{ik}^{12}, \tilde{W}_{ik}^{22}, \tilde{W}_{ik}^{21})$ .

are in the form  $(\tilde{W}_{ik}^{11}, \tilde{W}_{ik}^{12}, \tilde{W}_{ik}^{22}, \tilde{W}_{ik}^{21})$ . Note that  $\mathbb{P}[w_{ik} = W_{ik}^{ab}] = \mathbb{P}[\tilde{w}_{ik} = \tilde{W}_{ik}^{ab}]$  for all a, b, i and k, so that  $m_{ik}(\tilde{W}_{ik}^{ab}) \equiv m_{ik}(W_{ik}^{ab})$ . However, we do have to make a minor adjustment to how the  $u_i$  numbers are computed since the expected value of the node weights no longer has any meaning. In particular, we set  $u_i = 0$  for all exit tasks then recursively compute

$$u_i = \max_{k \in S_i} \left( u_k + \mathbb{E}[\tilde{w}_{ik}] \right) \tag{6}$$

for all others. It can readily be verified that this sequence is identical to that defined by (5) with the exception of the exit tasks, for which the corresponding numbers are now zero. (Technically we should perhaps rename this number sequence but given the fact that it is essentially identical we felt this was unnecessary.)

Now, for all i = 1, ..., n, let  $c_i$  be the critical path length from task  $t_i$  to the end and let  $e_i = \mathbb{E}[c_i]$  be its expected value. Define  $Z_i$  to be the set of all weight RVs corresponding to edges downward of  $t_i$  (i.e., the remainder of the graph). Let  $R_i$  be the set of all possible realizations of the RVs in  $Z_i$ . Given a realization  $z_i \in R_i$ , let  $\ell(z_i)$  be the critical path length from task  $t_i$  to the end (which is a scalar because all weights have been realized). Then by the definition of the expected value we have

$$e_i = \sum_{z_i \in R_i} \mathbb{P}[Z_i = z_i] \ell(z_i). \tag{7}$$

Suppose we define a sequence of numbers by  $f_i = 0$ , if  $t_i$  is an exit task, and

$$f_i = \sum_{z_i \in R_i} \mathbb{P}[Z_i = z_i] \max_{k \in S_i} \{ f_k + z_{ik} \}$$
 (8)

for all other tasks, where  $z_{ik}$  is the realization of the edge weight RV  $w_{ik}$  under the realization  $z_k$ . Then Fulkerson showed that  $u_i \leq f_i \leq e_i$  holds for all *i*—i.e, the  $f_i$  give a tighter bound on the expected values of the critical path lengths.

#### Proof of bound

Computing the Fulkerson numbers To compute each of the  $f_i$  using (8) we need to do an awful lot of work: suppose  $t_i$  has K children, then for any one of them  $t_k$  we need to do  $O(|\tilde{L}_{ik}|)$  operations, i.e.,  $O(|\tilde{L}_{ik}|^K)$  in total. In general,  $|\tilde{L}_{ik}|$  can be  $O(p^2)$  and K can be  $O(n^2)$  so computing the  $f_i$  in this manner is not always practical. Fortunately, a more efficient method was given by Clingen [3] in the context of extending Fulkerson's method to the case where edge weights are modeled as continuous random variables, although here we follow the slightly more compact notation of Elmaghraby [5].

It is well-known that the cumulative probability mass function of the maximum of a finite set of discrete RVs is equal to the product of the individual cumulative pmfs of the RVs. Let  $M_{ik}$  be the cumulative pmf along edge  $(t_i, t_k)$ , so that  $M_{ik}(x) = \mathbb{P}[\tilde{w}_{ik} \leq x]$ . Define the related function  $M_{ik}^*(x) = \mathbb{P}[\tilde{w}_{ik} < x]$ . Let  $Z_i$  be the set of all possible values of  $f_k + \tilde{w}_{ik}$ , for  $k \in S_i$ , and let z run over all elements of  $Z_i$ . For  $i = 1, \ldots, n$ , define

$$\alpha_i = \max_{k \in S_i} (f_k + \min(\tilde{L}_{ik})). \tag{9}$$

Then, with the cost independence assumptions we have already made, we can rewrite equation (8) as

$$f_i = \sum_{z \ge \alpha_i} z \left( \prod_{k \in S_i} M_{ik}(z - f_k) - \prod_{k \in S_i} M_{ik}^*(z - f_k) \right).$$
 (10)

A complete description of a practical procedure for computing the Fulkerson numbers  $f_i$  is given in Algorithm 1. At first blush this may not appear to be any simpler than before but, crucially, the number of operations required to compute each of the  $f_i$  is now  $O(|\tilde{L}_{ik}| \cdot K)$ , where K is the number of child tasks, rather than the first term being exponential in the second as before. Of course, it should also be noted that this procedure is still more expensive than computing the  $u_i$  sequence.

Once all of the  $f_i$  have been computed, they can be taken as alternative task ranks in HEFT (or indeed any other listing heuristic). In Table 3 we compare

Table 3: Fulkerson numbers for example graph.

| Task | $u_i$ | $f_i$ | MC1000 |
|------|-------|-------|--------|
| 0    | 24.75 | 28.2  |        |
| 1    | 16.25 | 16.9  |        |
| 2    | 5.75  | 5.75  |        |
| 3    | 16    | 18.1  |        |
| 4    | 7.75  | 7.75  |        |
| 5    | 7     | 7     |        |
| 6    | 2.5   | 0     |        |

the  $f_i$  values to the standard  $u_i$  for the small DAG from Figure 1; also included is MC1000, the critical path estimates based on 1000 Monte Carlo realizations (see Section 3.1.1), as a reference for the true critical path of the stochastic graph. We see that the  $f_i$  values do indeed give tighter bounds than the  $u_i$  and, more pertinently, if we use the former as task ranks then task 3 is scheduled before task 1 and we obtain the smaller schedule makespan of 20 rather than the standard HEFT makespan of 22.

Again, this is a single example, deliberately chosen for this behavior: although the  $f_i$  give tighter bounds on the critical path lengths of the stochastic DAG  $G_s$  there is absolutely no guarantee that this will lead to superior performance in general. After all,  $G_s$  itself is only a model of how we expect the processor selection phase to proceed—one that we know is inaccurate since, for example, it implicitly assumes that all task and edge weights are independent. Indeed, without this independence assumption it is well-known that the relation  $u_i \leq f_i$  does not necessarily hold even for  $G_s$ ; Fulkerson himself presented examples [6]. Still, we think this is a reasonable enough basis for an alternative ranking method in HEFT, so we investigate its performance compared to the usual  $u_i$  ranks via numerical simulation in Section 4.

Two refinements of Fulkerson's method were proposed by Elmaghraby [5]. The first involves computing each of the  $f_i$  numbers in the aforementioned manner and then reversing the direction of the remaining subgraph in order to calculate an intermediate result which can be used to improve the quality of the bound. The second is a more general approach based on using two or more point estimates of  $e_i$ , rather than just  $f_i$ , a method that was later generalized by Robillard and Trahan [10]. In both cases Elmagharaby proved that the new number sequences achieve tighter bounds on  $e_i$  than the Fulkerson numbers  $f_i$ . However, small-scale experimentation suggested that the improvement of Elmaghraby's new bounds over Fulkerson's were typically minor compared to the improvement of the latter over the standard HEFT  $u_i$  sequence so we chose to only evaluate here whether

tightening the bounds at all is useful.

**Algorithm 1:** Computing the Fulkerson numbers using Clingen's method.

```
1 for i=n,\ldots,1 do
           f_i = 0, \ \alpha_i = 0, \ Z_i = \{\}
 \mathbf{2}
           for k \in S_i do
 3
                \ell_m = \infty
  4
                 for \ell \in L_{ik} do
  \mathbf{5}
                      \ell_m \leftarrow \min(\ell_m, \ell)
  6
                      if f_k + \ell \notin Z_i then
  7
                       Z_i \leftarrow Z_i \cup \{f_k + \ell\}
  8
  9
                 end
10
                 \alpha_i \leftarrow \max(\alpha_i, f_k + \ell_m)
11
           end
12
           for z \in Z_i do
13
                if z \geq \alpha_i then
14
                      g = 1, d = 1
15
                       for k \in S_i do
16
                       g \leftarrow g \times M_{ik}(z - f_k) 
 d \leftarrow d \times M_{ik}^*(z - f_k) 
end
17
18
19
20
                 end
\mathbf{21}
           end
22
23 end
```

# 3.2 Adjusting the pmfs

In some sense, the purpose of the node and edge pmfs  $m_i$  and  $m_{ik}$  is to simulate the dynamics of the processor selection phase of HEFT—i.e.,  $m_i(W_i^a)$  should represent the probability that task  $t_i$  is scheduled on processor  $p_a$ , and so on. In HEFT, tasks are assigned to the processor that is estimated to complete their execution at the earliest time and attempting to model this accurately beforehand can quickly get messy and expensive—especially given the interaction between the two phases of the algorithm. However, a sensible idea may be to simply weight the processor selection probabilities according to their respective task computation costs: if, say, a task is 10 times faster on one processor than another then it seems more likely it will be scheduled on the former than the latter, even once the effect

of contention is taken into account. More precisely, for all tasks  $t_i$  let

$$s_i = \sum_a \frac{1}{W_i^a}$$

and define a new set of pmfs by

$$\hat{m}_i(W_i^a) = \frac{1}{W_i^a s_i} \ \forall i, a$$

and

$$\hat{m}_{ik}(W_{ik}^{ab}) = \hat{m}_i(W_i^a) \cdot \hat{m}_k(W_k^b)$$
$$= \frac{1}{W_i^a W_k^b s_i s_k} \quad \forall i, k, a, b.$$

Note that we take the reciprocal of the costs in order to reflect the idea that processors with smaller costs are more likely to be chosen than larger ones.

These modified pmfs can be used in conjunction with either upward ranking, as defined by equation (5), Fulkerson's bound, or Monte Carlo methods. For example, in the first instance, the expectations simply become

$$\mathbb{E}[w_i] = \sum_{\ell \in L_i} \ell \hat{m}_i(\ell) = \frac{q}{s_i},\tag{11}$$

$$\mathbb{E}[w_{ik}] = \sum_{\ell \in L_{ik}} \ell \hat{m}_{ik}(\ell) = \frac{1}{s_i s_k} \sum_{a,b} \frac{W_{ik}^{ab}}{W_i^a W_k^b},\tag{12}$$

and these can be used with equation (5) to compute an alternative sequence of task ranks  $\hat{u}_i$ . Of course, this is slightly more computationally expensive than computing the standard  $u_i$  ranks but only by a constant factor. Similarly, by using the modified pmfs in conjunction with equation (10) we can define alternative Fulkerson numbers  $\hat{f}_i$ , and by sampling realizations according to  $\hat{m}$  rather than m we can incorporate this idea into the Monte Carlo approach for finding longest paths. All three of these possibilities are evaluated as alternative task ranks for HEFT in the following section. Note that since equations (11) and (12) are simply weighted averages we do not believe that computing  $\hat{u}_i$  instead of  $u_i$  is a new idea: although we could not find any explicit references in the literature, we suspect this has been done before in practice.

# 4 Experimental comparison of rankings

In order to evaluate the alternative task prioritization phases in HEFT that we have outlined so far, we make use of our software simulator. However, first we need to define a suitably large and diverse set of graphs to compare them on.

### 4.1 Testing environment

We decided to evaluate the new rankings using the same two types of graphs as in the previous chapter: a set of Cholesky graphs, with real costs based on timings from an accelerated machine that we have access to, and several sets of randomly-generated graphs based on topologies provided by the STG [11]. The former are identical to those described in Section X so note that in particular they target only the two accelerated platforms detailed there: one comprising 1 GPU and 7 CPU cores and the other 4 GPUs and 28 CPU cores.

Since we are now interested in more general heterogeneous platforms we consider multiple sets of graphs based on randomly-generated graph topologies from the STG which differ from those in the previous chapter. In particular, a set is defined by the following parameters:

- $n \in \{100, 1000\}$ , the number of non-entry/exit tasks in each of the graphs. Each graph also has a single entry and exit task so that altogether it has e.g., 102 tasks rather than 100. Once this parameter has been specified, we use the topologies of the corresponding set of that size from the STG.
- $q \in \{2, 4, 8\}$ , the number of processors in the target platform.
- $\beta \in \{0.1, 1, 10\}$ , the computation-to-communication ratio (CCR). Defined as before, although the manner used to generate costs that give the target CCR differs (see below).
- $h \in \{1.0, 2.0\}$ , the *heterogeneity factor* of the processors. This basically determines how similar costs on different processors are to one another (again, see below for more detail).
- $m \in \{R, UR\}$ , the method used to generate the costs.

Once all of the other parameters are chosen, if m = UR (for unrelated), then we use the same method as in the original HEFT paper [12] or [1] to determine all of the task computation costs on each of the processors. To wit, first an average computation cost for the entire graph  $\overline{w_G}$  is chosen randomly (in our case an integer in the interval [1,100]). Then, for all  $i=1,\ldots,n$ , the average computation cost  $\overline{w_i}$  of task  $t_i$ , is chosen uniformly at random from the interval  $[0,\overline{w_G}]$ . Finally, for all  $a=1,\ldots,q$ ,  $W_i^a$  is also chosen uniformly at random but from the interval

$$\left[\overline{w_i} \times (1 - h/2), \ \overline{w_i} \times (1 + h/2)\right].$$

This method is perhaps somewhat unrealistic since task costs are generated independently of whichever processor they represent; typically costs are determined at least in part by the relative processor *powers*. With this in mind, if m = R (for related), then the method proceeds by first selecting an average power  $\bar{p}$  across the

set of processors uniformly at random from the interval [1, 100]. Then for each processor  $p_a$ , a = 1, ..., q, its power  $r_a$  is in turn chosen uniformly at random from the interval

$$[\overline{p} \times (1 - h/2), \ \overline{p} \times (1 + h/2)].$$

Now an average task cost  $\bar{t}$  is also chosen uniformly at random from the interval [1,100]. For each task  $t_i$ ,  $i=1,\ldots,n$ , choose  $x_i \in [0,2\bar{t}]$  uniformly at random and for all  $a=1,\ldots,q$ , realize  $g_a \sim \Gamma(1,r_a)$  (i.e., choose  $g_a$  from a Gamma distribution with mean and variance  $r_a$ ). This is done to ensure that the computation costs are not entirely determined by the power; a Gamma distribution was chosen because it is always positive and heavy-tailed, so has roughly the shape we're after. Finally, let  $W_i^a = x_i g_a$  be the computation cost of task  $t_i$  on processor  $p_a$ .

Communication costs are generated in the same manner for both choices of m. First, an average edge cost  $\overline{e}$  is computed such that the specified CCR is (approximately) achieved. Then for all edges  $(t_i, t_k)$ , we choose  $\overline{w_{ik}}$  uniformly at random from the interval  $[0, 2\overline{e}]$ . Then specific communication costs between the tasks on all possible pairs of distinct processors are chosen uniformly at random from the interval

$$\left[\overline{w_{ik}} \times (1 - h/2), \ \overline{w_{ik}} \times (1 + h/2)\right].$$

(Recall that costs are assumed to be zero when both tasks are scheduled on the same processor.) Note that the randomness here means that sometimes the target CCR is not precisely achieved, although it is usually acceptably close.

## 4.2 Benchmarking

First we want to establish how effective the HEFT task prioritization phase actually is for the sets of graphs described above. More so than the speedup and schedule length ratio, which were described in Section X, the metric that we perhaps really want to use here is how well the task list computed by HEFT compares to other topological sorts—orderings which respects precedence constraints—of the tasks. Now, the Networkx function topological\_sort returns a topologically sorted list of nodes in an input graph. Furthermore, the algorithm which this implements does not consider any objective other than meeting the precedence constraints, so we can in some sense regard this as a random sample from the set of all possible topological orderings for the graph. Hence to gauge the effectiveness of the HEFT task prioritization phase, we compare the makespan obtained with the standard ranking with that which would be obtained when using the task priority list returned by topological\_sort instead. Figure 6 shows the average makespan reduction, as a percentage, for all of the subsets of DAGs from the STG with 100 tasks.

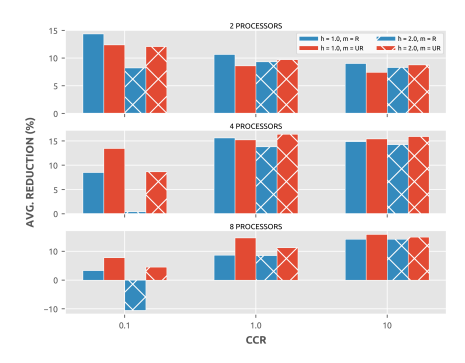


Figure 6: Average makespan reduction (%) for standard HEFT ranking phase vs random topological sort.

Clearly the most interesting takeaway from the figure is the negative reduction that we see for one of the DAG sets in the bottom-left corner—i.e., the random sort did better on average for those DAGs with CCR 0.1 and h=2.0 for which costs were generated using the *related* method. Interestingly, the effect appears to become more pronounced as the ratio of tasks to processors decreases: the reduction is always positive for the larger DAG sets (n=1000) with the same parameters, although the raw number of instances for which the random sort outperformed standard HEFT is high for those as well.

It is not obvious why this should be so but we suspect it is related to the similar phenomenon remarked upon in the previous chapter, where HEFT failed altogether due to difficulty managing communication costs because of its greedy processor selection phase. In fact, both standard HEFT and the random ranking alternative failed altogether in the vast majority of instances for which the former was worse than the latter; the average reduction is much smaller for certain subsets because they are precisely the ones for which HEFT is most likely to struggle in general. Disregarding those instances in which both failed, the percentage of graphs for which the random sort outperformed the standard ranking was roughly 0–2% for all subsets (of both sizes). Given this, we conclude that HEFT's task prioritization phase is clearly useful except for those circumstances when the algorithm itself struggles because of its greedy selection phase.

#### 4.3 Results

In this section we compare the performance of HEFT with the following task prioritization phases:

- the standard upward ranks  $u_i$ ,
- LB, the optimistic ranks  $\ell_i$  as defined by (2),
- F, the Fulkerson ranks  $f_i$  as defined by (10),
- W, the weighted means  $\hat{u}_i$  as defined in Section 3.2,
- WF, the weighted Fulkerson numbers  $\hat{f}_i$  as defined in Section 3.2.

First, we consider the sets of DAGs based on the STG with 100 tasks as an exploratory example. Ultimately, when deciding whether to use an alternative ranking we want to know whether it's more likely to help rather than harm performance, so Figure 7 shows the difference between the percentage of DAGs for which each of the alternative rankings obtained a schedule makespan better than the standard  $u_i$  ranks and the percentage for which they were worse. Note that we combine all four of the subsets (h = 1.0 or 2.0, m = R or UR) for each CCR and processor number combination, so that each of the bars represents a percentage over  $4 \times 180 = 720$  DAGs.

Of course, the figure does not tell the whole story: what about the relative magnitudes of the makespan reductions and increases? In general, both tended to be fairly small, with makespan changes greater than 5% either way being rare. Overall, across the entire set, the F and WF rankings obtained an average reduction of around 1%, while the LB and F rankings on average did no better than the standard ranking. Of course, a 1% reduction might not seem significant, but in certain situations—such as when the same application has to be executed repeatedly—this may be worthwhile. Bear in mind also that this is an average value: more significant reductions can be seen, for example, as the number of processors increases. Note that while large negative average reductions can be seen for some of the rankings on certain graph sets with  $\beta = 0.1$  (i.e., high communication), this is in fact down to only a handful of graphs. The issue is related to the previously-discussed problem of HEFT failing altogether in some cases. Typically, all of the rankings tend to fail for the same instances, but sometimes one or more of the rankings fails spectacularly whereas the HEFT ranking doesn't fail at all; this leads to a very large percentage difference in makespan which somewhat distorts the average. These cases are clearly important so should still be taken into account but they were very infrequent; overall, the number of failures was similar for all of the rankings. All of the data from the experiments described here can be found in the Github repository<sup>2</sup> associated with this chapter.

<sup>&</sup>lt;sup>2</sup>critical-path-estimation/scripts

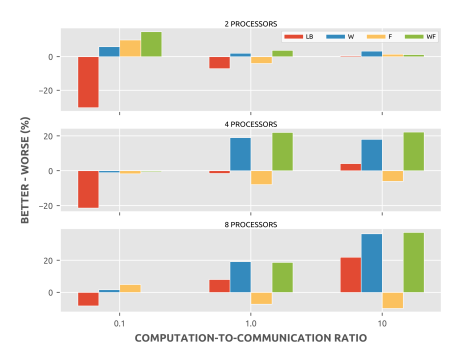


Figure 7: Percentage of instances for which the alternative rankings are better than the default, minus the percentage for which they were worse.

Results so far seem to suggest that the Fulkerson ranking in particular does not seem to be an improvement on the standard ranking. Given how much more expensive it can be, this is a big problem: only large makespan reductions can really justify the extra cost. Indeed, although the weighted Fulkerson ranking WF did much better, both compared to F and the standard ranking, it was not a significant improvement over the W ranking. This suggests that there is little advantage if any to obtaining a tighter bound on the associated stochastic graph (see Section 3.1). To confirm this, we repeated the experiments described above when the Monte Carlo approach (with a sufficient number of samples) is used to compute even tighter bounds on the critical path lengths (which are then used as ranks in HEFT). We used both the original pmfs m and the weighted pmfs  $\hat{m}_i$  for this. Our results can be found in full at the Github repository but overall they supported our conclusions from the example above: tightening the bound alone led to no consistent performance gains, whereas weighting the pmfs was more promising, with a roughly 2\% average makespan reduction across all of the graphs we considered.

Despite this, there are situations for which the Fulkerson ranking is competitive with the standard ranking in terms of both cost and performance. In particular, for accelerated environments, such as in the previous chapter, the set of possible values each weight may take (i.e.,  $\tilde{L}_{ik}$ ) is much smaller, which significantly improves efficiency. Furthermore, the Fulkerson ranking appears to perform

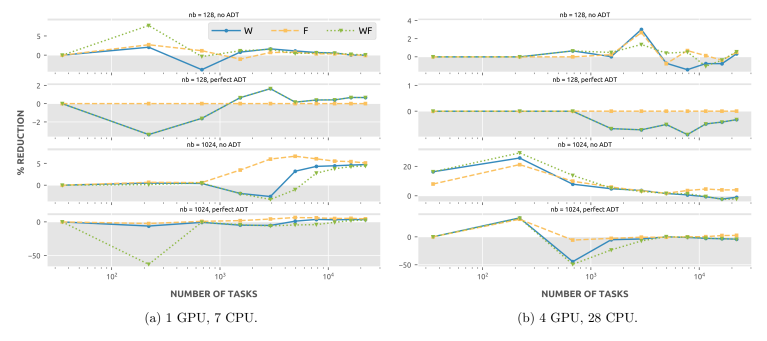


Figure 8: Schedule makespan reduction of HEFT with alternative ranking phases for Cholesky DAGs.

well for certain real application task graphs. For example, Figure 8 shows the makespan reductions of the three rankings W, F and WF compared to the default for the set of Cholesky graphs and (accelerated) target platforms defined in Section X [earlier chapter]. Although there is significant variation, we see that the Fulkerson ranking is superior to the standard ranking phase overall—typically only slightly but occasionally more significantly.

While it may be worthwhile to investigate in future if the performance of the F and WF rankings improves for larger DAGs, given that the results so far suggest they offer no advantage compared to much cheaper alternatives, we decided to omit those two rankings when we repeated our comparison for sets of larger DAGs (n=1000) from the STG. The metric we choose to use now is the average percentage degradation (APD), as defined in the previous chapter. Figure 9 shows the APDs for the standard HEFT ranking and the alternative LB and W rankings. We see roughly the same pattern as for the smaller DAGs: the W ranking consistently outperforms the other two, although the margins are even smaller this time than before.

#### 4.4 Conclusions

To summarize, our main conclusions from all of the experiments described in this section are as follows.

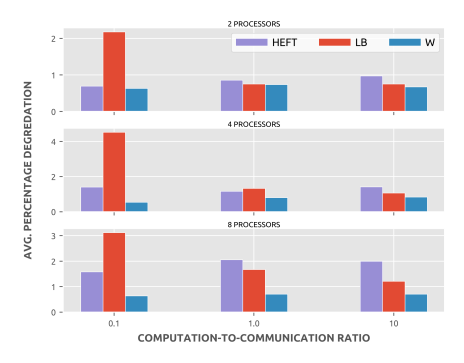


Figure 9: Average percentage degradation for select task prioritization phases in HEFT, size 1000 DAGS from the STG.

- Largely supporting previous investigations along these lines [14], our biggest takeaway is perhaps that how the critical path lengths are estimated seems to make relatively little difference to the schedule makespan overall, with improvements (if any) relative to the default approach typically being fairly minor on average.
- Having said that, the W ranking was consistently more likely to result in a smaller makespan across the majority of the sets considered. The usual caveats about the limitations of our experimental framework hold of course but overall we would softly recommend it be used instead of the default, particularly since it is conceptually simple (just a weighted mean) and not significantly more expensive.
- There is no clear benefit in obtaining tighter bounds on the associated stochastic graph (i.e., Fulkerson's bound). Given the higher time complexity in the general case, this does not therefore seem to be a worthwhile alternative.
- Similarly, there appears to be little advantage in using the LB ranking, although it also wasn't clearly inferior to the standard ranking in terms of performance or cost.

## 5 Processor selection

Critical path estimates are used in HEFT—and many similar listing heuristics—only at the task prioritization phase. This begs the question, can they also be useful for processor selection? Arabnejad and Barbosa's Predict Earliest Finish Time (PEFT) heuristic [1] represents one sensible way this can be done. Recall from Section X that before scheduling begins PEFT computes a table of *optimistic costs*  $C_i^a$  for all task and processor combinations in the following manner. First, set  $C_i^a = 0$ ,  $a = 1, \ldots, q$ , for all exit tasks, then move up the DAG and recursively compute

$$C_i^a = \max_{k \in S_i} \left( \min_{b=1,\dots,q} \left( \delta_{ab} \overline{w_{ik}} + W_k^b + C_k^b \right) \right)$$
 (13)

for all other tasks, where  $\delta_{ab} = 1$  if a = b and 0 otherwise. The  $C_i^a$  values are referred to in PEFT as optimistic costs but can be interpreted as *conditional* critical paths in that they represent some estimate of future schedule costs given a processor selection. When scheduling, say, task  $t_i$ , in PEFT we choose the processor  $p_{opt}$  defined by

$$p_{opt} := \min_{a} \left( F_i^a + C_i^a \right)$$

where, as in previous chapters,  $F_i^a$  is the estimated schedule makespan when  $t_i$  is completed by  $p_a$ . This is a nice extension of the dynamic programming principles underlying HEFT: rather than optimizing the schedule makespan up to the current task (i.e.,  $F_i^a$ ), we extend the horizon and optimize an estimate of the complete schedule makespan. Now, computing the full optimistic cost table is only  $O(n^2q)$ —i.e., the same as HEFT—but since the values within are similar in nature to the upward ranks  $u_i$  it is sensible (and more efficient) to make use of them for prioritizing tasks, rather than going to the effort of computing the upward ranks as well. Hence PEFT defines task priorities  $C_i$  through

$$C_i = \frac{1}{q} \sum_a C_i^a. \tag{14}$$

It is important to note here that the task priorities as computed by (14) do not necessarily respect precedence constraints since the equality  $C_i^a \leq C_k^a$  for  $k \in S_i$  does not hold (because of the internal minimization over all processors). However this is easily remedied by selecting tasks for scheduling from the pool of currently "ready" tasks (i.e., those for which all of their parents have been scheduled) according to their priorities. Note also that although it is arguably more natural that task ranks include the cost of the task itself, which these do not, it is suggested that the savings made through the alternative selection step are more beneficial overall. Certainly, numerical experiments described by

the original authors [1] suggest that PEFT is at least competitive with HEFT, especially when there are a large number of processing resources.

The structure of PEFT follows a more general heuristic framework which we refer to as *Heterogeneous Smallest Makespan* (HSM) and is defined by the following procedure:

- 1. Compute a table of conditional critical path estimates  $C_i^a$  for all  $i=1,\ldots,n$  and  $a=1,\ldots,q$ .
- 2. Compute all task ranks  $C_i$  as some function of the  $C_i^a$ .
- 3. At the processor selection phase, schedule task  $t_i$  on the processor which minimizes  $F_i^a + C_i^a$ .

PEFT is defined by using equations (13) and (14) for the first and second parts of this framework, respectively. A natural question is, are there any better choices?

### 5.1 Alternative conditional critical paths

All of the methods for estimating critical path lengths at the prioritization phase which were introduced previously can be modified to give conditional critical path estimates (which also disregard the cost of the task itself). No matter which, we first let  $C_i^a = 0$  for  $a = 1, \ldots, q$  and all exit tasks, then move up the DAG and recursively compute the other values, so from now on we focus only on the latter. The most straightforward method to extend is the optimistic lower bound, for which we now compute

$$C_i^a = \max_{k \in S_i} \left( \min_{b=1,\dots,q} \left( C_k^b + W_k^b + W_{ik}^{ab} \right) \right), \quad a = 1,\dots,q,$$
 (15)

for all non-exit tasks. Note that these values are extremely similar to the optimistic costs (13) as used in PEFT, with the exception that the specific communication cost  $W_{ik}^{ab}$  is used in the minimization rather than the average  $\overline{w_{ik}}$ . Indeed, this is arguably the most intuitive way to define the conditional critical path since the value  $C_i^a$  is a true lower bound on the remaining makespan of any schedule which executes task  $t_i$  on processor  $p_a$ ; locally-optimal processor selections are overruled if the best possible final makespan we can hope to achieve given that selection is inferior to other choices.

Alternatively, we could take a similar tack to the standard HEFT upward ranks and use an estimate of what we expect the conditional critical paths to be. More specifically, we move up the DAG and recursively compute

$$C_i^a = \max_{k \in S_i} \left( \frac{1}{q} \sum_{b=1}^q (C_k^b + W_k^b + W_{ik}^{ab}) \right), \quad a = 1, \dots, q.$$
 (16)

In fact, we could just as easily use a weighted average inside the maximization, which is particularly attractive since the corresponding ranking performed well in the previous section. In particular, for all non-exit tasks we recursively compute

$$C_i^a = \max_{k \in S_i} \left( \frac{1}{\hat{s}_k} \sum_{b=1}^q \frac{C_k^b + W_k^b + W_{ik}^{ab}}{W_k^b + C_k^b} \right), \quad a = 1, \dots, q,$$
 (17)

where

$$\hat{s}_k = \sum_{b=1}^q \frac{1}{W_k^b + C_k^b}. (18)$$

This is conceptually similar to the weighted pmf  $\hat{m}$  as defined in Section 3.2, except that we also include the conditional critical path lengths in the weighting. The motivation for the change is that the probability a task  $t_i$  will ultimately be scheduled on a given processor  $p_a$  at the selection phase now also depends on the  $C_i^a$  value.

### 5.2 Computing task priorities

There are many different ways we can use the conditional critical path estimates  $C_i^a$  to derive task priorities  $C_i$ . However, given the superior performance of the weighted average task ranking in Section 4, a similar approach here would seem to be most sensible default choice. In particular, no matter how the conditional critical paths  $C_i^a$  are estimated, we compute the task priorities recursively, starting from the leaves, through

$$C_i = \frac{q}{\hat{s}_i} + \max_{k \in S_i} C_k,\tag{19}$$

where the maximization is taken to be zero if  $S_i$  is empty (i.e., for exit tasks) and  $\hat{s}$  is as defined in (18). All of the results presented in the following section are based on heuristics for which the task priorities are computed in this manner, although it should again be emphasized that others are possible and several were considered in preliminary testing.

# 5.3 Empirical comparison

In order to evaluate the performance of the different choices in the HSM framework that have been described so far, we used the same software simulation environment as in Section 4. As before, all of the scripts used to generate the results presented here are available at the Github repository<sup>3</sup> for this chapter. We consider the following three variants of the HSM framework, defined by how they compute the conditional critical paths since all use the task prioritization given by (19).

<sup>&</sup>lt;sup>3</sup>critical-path-estimation/scripts

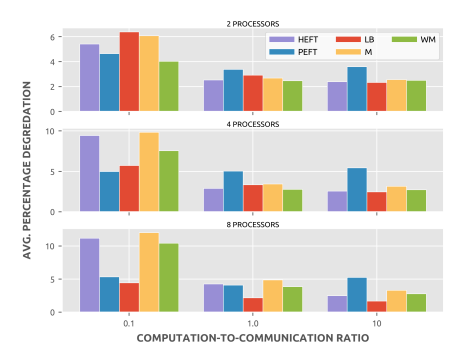


Figure 10: Average percentage degradation for HSM variants, HEFT and PEFT, size 100 DAGS from the STG.

- LB, using (15).
- M, using (16).
- WM, using (17).

Figure 10 shows the average percentage degradation of these three heuristics, as well as HEFT and PEFT, for the set of DAGs based on topologies from the STG with 100 tasks (plus single entry and exit tasks). As before, we combine all four of the subsets (h=1.0 or 2.0, m=R or UR) so that each CCR and processor number combination represents a larger subset comprising 720 DAGs. The most immediate takeaway from the figure is that, with the exception of the subsets with the smallest CCR, PEFT actually does relatively poorly compared to both HEFT and the HSM variants. On the surface, this might seem to contradict results published elsewhere, but it should be noted that those experiments were typically for much larger and more diverse graph sets, so some local variation is to be expected. Note that when communication predominates (i.e.,  $\beta=0.1$ ), PEFT and HSM-LB are superior to the others for both 4 and 8 processors. This is again related to the problem of listing heuristics failing altogether for DAGs with high communication: PEFT and HSM-LB recorded only around half as many failures as the others for these sets.

One obvious question we have not so far addressed is how useful the PEFT-like processor selection phase actually is compared to the simple HEFT-like earliest

finish time alternative. In fact, this was far from clear in our experiments: with a few exceptions, there was usually no statistically significant difference on average between the two. The exceptions were all for the smallest CCR sets, for which the processor selection clearly improved ( $\approx 5\%$  average makespan reduction) on the ranking-only version for the LB variant and was even more significantly worse for the M and WM variants (> 5% average makespan increase).

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