

# An Extension of Edge Zeroing Heuristic for Scheduling Precedence Constrained Task Graphs on Parallel Systems Using Cluster Dependent Priority Scheme

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**Abstract.** Sarkar's edge zeroing heuristic for scheduling precedence constrained task graphs on parallel systems can be viewed as a priority based algorithm in which the priority is assigned to edges. In this algorithm, the priority is taken as the edge weight. This can also be viewed as a task dependent priority function that is defined for pairs of tasks. We have extended this idea in which the priority is a cluster dependent function of pairs of clusters (of tasks). Using this idea we propose an algorithm of complexity O(|V||E|(|V|+|E|)) and compare it with some well known algorithms.

Keywords: clustering, homogeneous systems, parallel processing, scheduling, task allocation.

## 1. Introduction

A parallel system is designed so that it can execute the applications faster than a sequential system. For this we need to parallelize the program. There are three steps involved in the parallelization of a program (Sinnen [27]). The first step is called *task decomposition* in which the application is divided into tasks. The *degree of concurrency* is the number of tasks that can be executed simultaneously (Grama et al. [10]).

The tasks generated may have interdependencies between them that will decide the partial execution order of tasks. The determination of precedence constraints between the tasks is the second step of parallelization and is called *dependence analysis* (Banerjee et al. [2], Wolfe [29]).

A dependence relation among the tasks is represented as a directed acyclic graph, also known as the *task graph*. Nodes in the task graph represent the tasks and have a weight associated with them that represents the execution time of the task. Edges in the task graph represent the dependence relation between the tasks and have a weight associated with them that represents the communication time between the tasks.

The final step of parallelization is the *scheduling* of tasks to the processors. By scheduling we mean both the spatial assignment (task allocation), and the temporal assignment (assigning start time) of tasks to the processors.

The problem of finding a scheduling for a given task graph on a given set of processors that takes minimum time is *NP-Complete* (Sarkar [26], Papadimitriou and Yannakakis [24]). Therefore several heuristics are applied for solving this problem in polynomial time (Yang and Gerasoulis [32], Gerasoulis and Yang [8], Dikaiakos et al. [7], Kim and Browne [16], Kwok and Ahmed [17], [18], Lo [19], Malloy et al. [22], Wu and Gajski [30], Kadamuddi and Tsai [14], Sarkar [26], Yang and Gerasoulis [33], Wu et al. [31], Sih and Lee [28]). The solutions generated by using these algorithms are generally suboptimal.

Our heuristic is an extension of Sarkar's edge zeroing heuristic [26] for scheduling precedence constrained task graphs on parallel systems. Sarkar's algorithm can be viewed as a task dependent priority based algorithm in which the priority (in this case edge weight) is a function of pairs of *tasks*. We extend this concept and define the priority as a cluster dependent function of pairs of *clusters*. Using this concept we propose an algorithm of complexity O(|V|/E|(|V|+|E|)) and compare it with some well known algorithms.

The remainder of the paper is organized in the following manner. Section 2 presents an overview of the

related literature. Section 3 defines a cluster dependent priority scheme that is dependent on cluster-pairs and also presents the proposed algorithm. Section 4 presents a detailed description of the algorithms used. Section 5 gives a sample run of the algorithm. Section 6 presents some experimental results. And finally in section 7 we conclude our work.

### 2. Literature Overview

Most scheduling algorithms for parallel systems in the literature are based on an idealized model of the target parallel system also referred to as the *classic model* (Sinnen [27]). It is a set of identical processors with fully connected dedicated communication subsystem. Local communications are cost-free and we also have concurrent inter-processor communications.

A fundamental scheduling heuristic is called the *list scheduling* heuristic. In list scheduling, first we assign a priority scheme to the tasks. Then we sort the tasks according to the priority scheme, while respecting the precedence constraints of the tasks. Finally each task is successively scheduled on a processor chosen for it. Some examples of list scheduling algorithms are: Adam et al. [1], Coffman and Graham [3], Graham [9], Hu [13], Kasahara and Nartia [15], Lee et al. [20], Liu et al. [21], Wu and Gajski [30], Yang and Gerasoulis [34].

Another fundamental scheduling heuristic is called *clustering*. Basically it is a scheduling technique for an unlimited number of processors. It is often proposed as an initial step in scheduling for a limited number of processors. A cluster is a set of tasks that are scheduled on the same processor. Clustering based scheduling algorithms generally consist of three steps. The first step finds a clustering of the task graph. The second step finds an allocation of clusters to the processors. The last step finds a scheduling of the tasks. Some examples of clustering based scheduling algorithms are: Mishra et al. [23], Yang and Gerasoulis [32], Kim and Browne [16], Kadamuddi and Tsai [14], Sarkar [26], Hanen and Munier [11].

## 3. The Cluster Dependent Priority Scheduling Algorithm

#### 3.1. Notation

Let *N* denote the set of natural numbers:  $\{1, 2, 3, ...\}$ . Let *R* denote the set of real numbers, and let  $R^+$  denote the set of non-negative real numbers. For  $(1 \le i \le n)$ , let there be *n* tasks  $M_i$ . Let

$$\mathbf{M} = \{\mathbf{M}_i \mid 1 \le i \le n\} \tag{1}$$

be the set of tasks. Then for  $(1 \le i \le n)$ , the clusters  $C_i \in M$  are such that for  $i \ne j$  and  $(1 \le i \le n, 1 \le j \le n)$ :

$$C_i \cap_{i \neq j} C_j = \phi, \tag{2}$$

and

$$\mathcal{O}^{n}_{i=1}C_{i}=\boldsymbol{M}.$$
(3)

Let

$$\boldsymbol{C} = \{ \boldsymbol{C}_i \mid \boldsymbol{C}_i \in \boldsymbol{M}, \ 1 \leq i \leq n \}$$

$$\tag{4}$$

be a decomposition of M into clusters. Note that some of the  $C_i$ 's may be empty. Let

$$V = \{i \mid 1 \le i \le n\} \tag{5}$$

denote the set of vertices of the task graph. Let the directed edge from *i* to *j* be denoted as  $(i \rightarrow j)$ . Let

$$E = \{(i, j) \mid i \in V, j \in V, \exists (i \to j)\}$$
(6)

denote the set of edges of the task graph. Let  $m_i \in \mathbf{R}^+$  be the execution time of the task  $M_i$ . If  $(i, j) \in E$ , then let  $w_{ij} \in \mathbf{R}^+$  be the communication time from  $M_i$  to  $M_j$ . Let T be the adjacency list representation of the task graph.

Let *cluster* :  $N \rightarrow N$  be a function such that:

$$cluster(i) = j \leftrightarrow M_i \in C_i.$$
<sup>(7)</sup>

For  $C_i \in C$ , let *comp* :  $C \to R^+$  be a function that gives the total computation time of a cluster:

$$comp(C_j) = \sum_{Mi \in Cj} w_i.$$
 (8)

For  $C_i \in C$ , and  $C_j \in C$ , let *comm* :  $C \times C \to R^+$  be a function that gives the total communication time from the first cluster to the second cluster:

$$comm(C_i, C_j) = \sum_{Mp \in C_i, Mq \in C_j, (p, q) \in E} w_{pq}.$$
(9)

We follow the convention that for a given function f, f(...) is a function notation (useful in definition) and f[...] (or f[...][...]) is the corresponding array notation (useful in algorithms for implementing functions as arrays).

#### **3.2.** Sarkar's Edge Zeroing Heuristic

When the two tasks that are connected through a large weight edge, are allocated to different processors, then this will make a large communication delay. To avoid large communication delays, we generally put such tasks together on the same machine, thus avoiding the communication delay between them. This concept is called edge zeroing.

Sarkar's alorithm [26] uses the concept of edge zeroing for clustering of tasks. Edges are sorted in decreasing order of edge weights. Initially each task is in a separate cluster. Edges are examined one-by-one in decreasing order of edge weight. The two clusters connected by the edge are merged together if on doing so, the parallel execution time does not increase. Sarkar's algorithm uses the level information to determine the parallel execution time and the levels are computed for each step. This process is repeated until all the edges are examined. The complexity of Sarkar's algorithm is O(|E|(|V|+|E|)).

We can define a task dependent priority scheme for Sarkar's algorithm. Let  $P_m : M \times M \to R$  be a function such that

$$P_m(M_i, M_j) = w_{ij} \leftrightarrow (i,j) \in E, \tag{10}$$

and

$$P_m(M_i, M_j) = -\infty \leftrightarrow (i, j) \notin E.$$
(11)

Now if we make a descending priority queue of pairs of tasks  $(M_i, M_j)$  based on the value of  $P_m$  such that

$$P_m(M_i, M_j) \ge 0, \tag{12}$$

then this is equivalent to the first step of Sarkar's algorithm in which the edges are sorted in decreasing order of edge weight.

#### **3.3.** An Example of Cluster Dependent Priority Scheme

In order to get a good scheduling algorithm, we should define a good cluster dependent priority scheme  $P_c$ . We use two observations. The first observation is that by merging two clusters that are heavily communicating with each other, we can expect to reduce the parallel execution time by using the edge zeroing principle. The second observation is that we can exploit parallelism in a better way by keeping two heavily computing clusters separated. Therefore a good priority scheme should reflect these two observations.

One such cluster dependent priority scheme is defined by:

$$P_c(C_i, C_i) = comm(C_i, C_i) + comm(C_i, C_i) - comp(C_i) - comp(C_i).$$

$$\tag{13}$$

#### **3.4.** The Cluster-Pair-Priority-Scheduling Algorithm

Cluster-Pair-Priority-Scheduling(T)

```
01 flag \leftarrow true
02 for k \leftarrow l to |V|
03
        do cluster[k] \leftarrow k
04 P_c[...][...] \leftarrow Evaluate-Priority(T, cluster[...])
05 (p, array[...]) \leftarrow Sort(T, P_c[...][...], cluster[...])
06 min \leftarrow Evaluate-Time(T, cluster[...])
07 reach[...][...] \leftarrow Reachability (T)
08 (topology[...], trank[...]) \leftarrow Topological-Sort(T)
09 while flag = true
10
       do flag \leftarrow false
           for m \leftarrow l to p
11
                do if flag = true
12
```

13 *then break* 

])

14	$(i, j) \leftarrow array[m]$
15	Merge(T, i, j, cluster[], topology[], trank[], reach[][
16	$time \leftarrow Evaluate-Time(T, cluster[])$
17	<i>if</i> time < min
18	then $flag \leftarrow true$
19	$min \leftarrow time$
20	$P_c[][] \leftarrow Evaluate-Priority(T, cluster[])$
21	$(p, array[]) \leftarrow Sort(T, P_c[][], cluster[])$
22	else undo step 15
• •	

23 return (min, cluster[...])

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Cluster-Pair-Priority-Scheduling(...) (CPPS) is the proposed algorithm that uses the cluster dependent priority scheme. Let  $P_c: C \times C \to R$  be a cluster dependent priority function. flag is a Boolean variable that is used to determine when a clustering is able to reduce the parallel execution time of the system. In line 01, flag is initialized to true. In the for loop from lines 02 to 03, each task is initially kept in a separate cluster. In line 04, the priority of each cluster-pair is calculated using the function Evaluate-Priority(...) that is connected by at least one edge and stored in the array  $P_c[...][...]$ . In line 05, the cluster-pairs are sorted using the function Sort(...) in non-increasing order of their  $P_c[...][...]$  values. Sort(...) returns p and array[...]. p is the number of cluster-pairs between which  $P_c[...][...]$  is defined. In line 06, the parallel execution time of the clustering is evaluated using the function Evaluate-Time(...), and stored in the variable min. min is used to obtain the clustering that gives the minimum parallel execution time. In line 07, the reachability matrix reach[...][...] is evaluated using the function Reachability(...). We have reach[i][j] = true if and only if there exists a path from  $M_i$  to  $M_j$ . In line 08, the task graph T is topologically sorted using the function Topological-Sort(...) and stored in the variable set using the array topology[...]. The topological ranks of the tasks are stored in the array trank[...].

In lines 09 to 22, in the *while* loop, the cluster-pairs are examined one-by-one in non-increasing order of their  $P_c[...][...]$  values (the *for* loop from lines 11 to 22). In line 15, the cluster-pair is merged using the function *Merge(...)*. In line 16, the parallel execution time of the current clustering is evaluated using the function *Evaluate-Time(...)* and the value is stored in the variable *time*. In lines 18 to 21, if the parallel execution time is reduced, then the cluster-pairs are again sorted in non-increasing order of their  $P_c[...][...]$  values, and the *while* loop from lines 09 to 22 is restarted. In line 22, if the parallel execution time of the clustering is not reduced, then the cluster-pair is kept separated. This process is repeated until no further reduction in parallel execution time is possible. Line 23 returns the parallel execution time and the clustering.

Line 01 has complexity O(1). The *for* loop from lines 02 to 03 has complexity O(/V/). In line 04, the algorithm *Evaluate-Priority*(...) has complexity O(/V/ + /E/) (section 4.1). In line 05, the algorithm *Sort*(...) has complexity  $O(/E/\log (/E/))$  (Horowitz et al. [12]). In line 06, the algorithm Evaluate-Time(...) has complexity O(/V/ + /E/) (section 4.3). In line 07, the algorithm *Reachability*(...) has complexity O(/V/ + /E/) (section 4.3). In line 07, the algorithm *Reachability*(...) has complexity O(/V/ + /E/) (Cormen et al. [4]). The complexity of the CPPS algorithm is dominated by the *while* loop from lines 09 to 22 that can iterate a maximum of /V/ times, since after each merging of clusters, one cluster is reduced. The *for* loop from lines 11 to 22 can iterate a maximum of /E/ times, since  $P_c[...][...]$  is defined between the clusters that are having edges between them. Each iteration of the *for* loop has complexity that is dominated by lines 15 and 16, each of which has complexity O(/V/ + /E/) (Sections 4.2 and 4.3 respectively). Therefore, the *while* loop has complexity O(/V/|E/(/V/ + /E/)) that is also the complexity of CPPS algorithm.

## 4. A Detailed Description of the Algorithms Used

#### 4.1. Evaluate-Priority

Evaluate-Priority(T, cluster[...])

- 01 *for* each  $(i,j) \in E$
- 02 *do*  $k \leftarrow cluster[i]$
- 03  $l \leftarrow cluster[j]$
- 04 if  $k \neq l$

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05 then  $P_c[k][l] \leftarrow 0$ 06 *for* each  $(i,j) \in E$ *do*  $k \leftarrow cluster[i]$ 07 08  $l \leftarrow cluster[j]$ 09 if  $k \neq l$ 10 then  $P_c[k][l] \leftarrow P_c[k][l] + w_{ii}$ 11 for  $i \leftarrow l$  to |V|12 do comp[i]  $\leftarrow 0$ 13 for  $i \leftarrow l$  to |V|14 do  $k \leftarrow cluster[i]$ 15  $comp[k] \leftarrow comp[k] + w_i$ 16 for each  $(i,j) \in E$ 17 do  $k \leftarrow cluster[i]$  $l \leftarrow cluster[j]$ 18 19 if  $k \neq l$ 20 then if (comp[k] + comp[l]) was previously not subtracted from  $P_c[k][l]$ 21 then  $P_c[k][l] \leftarrow P_c[k][l] - (comp[k] + comp[l])$ 22 *return P<sub>c</sub>[...][...]* 

In lines 01 to 05, the  $P_c[...][...]$  values between different cluster-pairs are initialized to 0. In lines 06 to 10, the edge weights between the corresponding cluster-pairs are added to the  $P_c[...][...]$  values between them. The *comp[...]* value of each cluster is initialized to 0 in lines 11 to 12. The *comp[...]* value of clusters are calculated in lines 13 to 15. In lines 16 to 21, the *comp[...]* value of corresponding clusters are subtracted from the sum of weights between the clusters to get the  $P_c[...][...]$  values between the cluster-pairs. Line 22 returns the array  $P_c[...][...]$ .

The *for* loop from lines 01 to 05, and 06 to 10, each have complexity O(|E|). The *for* loop from lines 11 to 12, and 13 to 15, each have complexity O(|V|). The *for* loop from lines 16 to 21 has complexity O(|E|). Line 22 has complexity O(1). Therefore the algorithm *Evaluate-Priority*(...) has complexity O(|V| + |E|).

## 4.2. Merge

Merge(T, i, j, cluster[...], topology[...], trank[...], reach[...][...])  $01 flag_1 \leftarrow flag_2 \leftarrow false$  $02 k \leftarrow l \leftarrow 0$ 03  $ablevel[...] \leftarrow Allocated-Bottom-Level(T, cluster[...])$ 04 *while*  $k \leq |V|$  and  $l \leq |V|$ 05 do if flag<sub>1</sub> = false *then while*  $k \leq |V|$ 06 07 do  $k \leftarrow k + 1$ 08 *if* cluster[topology[k]] = cluster[i] 09 then  $p_1 \leftarrow topology[k]$  $flag_1 \leftarrow true$ 10 11 break  $if flag_2 = false$ 12 13 *then while*  $l \leq |V|$ 14 *do*  $l \leftarrow l + l$ 15 *if* cluster[topology[l]] = cluster[j] 16 *then*  $p_2 \leftarrow topology[l]$ 

17	$flag_2 \leftarrow true$
18	break
19	<b>if</b> $flag_1 = true and flag_2 = true$
20	<i>then if</i> $reach[p_1][p_2] = true$
21	then $flag_1 \leftarrow false$
22	$if reach[p_2][p_1] = true$
23	then $flag_2 \leftarrow false$
24	<i>if</i> $flag_1 = true and flag_2 = true$
25	then if $ablevel[p_1] > ablevel[p_2]$
26	<i>then</i> add a pseudo-edge of cost 0 from $p_1$ to $p_2$
27	$flag_1 \leftarrow false$
28	$if$ $ablevel[p_1] < ablevel[p_2]$
29	<i>then</i> add a pseudo-edge of cost 0 from $p_2$ to $p_1$
30	$flag_2 \leftarrow false$
31	$if$ $ablevel[p_1] = ablevel[p_2]$
32	<i>then if</i> $trank[p_1] < trank[p_2]$
33	<i>then</i> add a pseudo-edge of cost 0 from $p_1$ to $p_2$
34	$flag_1 \leftarrow false$
35	else add a pseudo-edge of cost 0 from $p_2$ to $p_1$
36	$flag_2 \leftarrow false$
37 for $m \leftarrow$	1 to  V

 $3 \mid for \ m \leftarrow 1 \ to \ |V|$ 

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38 *do if* cluster[m] = cluster[j]

39 *then*  $cluster[m] \leftarrow cluster[i]$ 

In line 01,  $flag_1$  and  $flag_2$  are initialized to false.  $flag_1$  is used for searching the next task that is in the cluster  $C_i$  in topological order (line 05 and the inner *while* loop from lines 06 to line 11). The task searched is stored in  $p_1$  (line 09).  $flag_2$  is used for searching the next task that is in the cluster  $C_j$  in topological order (line 12 and the inner *while* loop from lines 13 to line 18). The task searched is stored in  $p_2$  (line 16). In line 02, k and l are initialized to 0. k and l are used as indices for searching the tasks in topological order that belong to the clusters  $C_i$  and  $C_j$  respectively. In line 03, the allocated bottom level (Sinnen [27]) of the tasks of the task graph T for the current allocation cluster[...] (before merging  $C_i$  and  $C_j$ ) is evaluated using the function Allocated-Bottom-Level(...) and stored in the array ablevel[...]. Given an allocation of tasks, the *allocated bottom level* of a task  $M_n$  is the length of the longest path starting with  $M_n$  in which the weight of an edge is taken as 0 if the two tasks corresponding to the edge are allocated on the same processor.

Given an allocation, a scheduled DAG (Yang and Gerasoulis [32]) is a DAG in which there are no independent tasks in any cluster. If there are independent tasks in a cluster, then we can make them dependent by adding 0-cost pseudo-edges between them. We follow the approach used by Sarkar (Sarkar [26]) in which we add a pseudo-edge of  $\cos \theta$  from a task of higher allocated bottom level to a task of lower allocated bottom level if the two tasks are independent (lines 25 to 30). If the allocated bottom levels of the tasks are same, then we add a pseudo-edge of  $\cos t \theta$  from a task that comes earlier in topological order to the task that comes later in topological order (lines 31 to 36). By following this strategy, it will ensure that there does not exist any cycle in the task graph so that the resulting graph is a scheduled DAG. This is because in a DAG, the successor of a task has lower allocated bottom level and also it comes topologically later. If we always add pseudo-edges of cost 0 from tasks having a higher allocated bottom level to a task having a lower allocated bottom level, then if a cycle exists then some successor of a node will have a higher allocated bottom level than the node itself which is not possible (for the case of cycles having at least one non-pseudoedge). For the case of equal allocated bottom levels (for the case of cycles having all edges as pseudo-edges), since we are always adding a pseudo-edge of  $\cot \theta$  from a task that comes earlier in topological order to the task that comes later in topological order, consider the task in the cycle that comes earliest in topological order. Now the predecessor of that task should come even earlier than the task itself implying a contradiction. Therefore a cycle is not possible.

We are evaluating the reachability matrix reach[...][...] using the function Reachabilty(...) at the initial time only to reduce the complexity of the Merge(...) algorithm. This makes no difference because adding some extra 0-cost pseudo-edges will make no difference to the scheduling. If  $p_2$  is reachable from  $p_1$ , then we again search for the next  $p_1$  in topological order (lines 20 to 21). If  $p_1$  is reachable from  $p_2$ , then we again search for the next  $p_2$  in topological order (lines 22 to 23). If neither  $p_2$  is reachable from  $p_1$ , nor  $p_1$  is reachable from  $p_2$ , then we draw a 0-cost pseudo-edge from the task with higher ablevel[...] to the task with lower ablevel[...] (lines 24 to 30). For the case of equal ablevel[...], we draw a 0-cost pseudo edge from the task with lower trank[...] to the task with higher trank[...]. We continue in this manner until all the nodes of the two clusters are examined (the outer **while** loop from lines 04 to 36). And finally in lines 37 to 39, we merge the two clusters  $C_i$  and  $C_j$ .

Lines 01 to 02 have complexity O(1). In line 03, *Allocated-Bottom-Level*(...) has complexity O(|V| + |E|) (Sinnen [27]). The *while* loop from lines 04 to 36 has complexity O(|V|). The *for* loop from lines 37 to 39 has complexity O(|V|). Therefore the algorithm Merge(...) has complexity O(|V| + |E|).

#### 4.3. Evaluate-Time

```
Evaluate-Time(T, cluster[...])
01 stack \leftarrow empty
02 for k \leftarrow l to |V|
        do at level \lceil k \rceil \leftarrow 0
03
            time[k] \leftarrow 0
04
05
            backlink[k] \leftarrow 0
06 for each (k, m) \in E
07
        do backlink[m] \leftarrow backlink[m] + 1
08 for k \leftarrow l to |V|
09
        do if backlink[k] = 0
10
               then Push(stack, k)
11 while stack \neq empty
12
       do i \leftarrow Pop(stack)
13
           if atlevel[i] \geq time[cluster[i]]
              then time[cluster[i]] \leftarrow atlevel[i] + w_i
14
              else time[cluster[i]] \leftarrow time[cluster[i]] + w_i
15
16
           for each (i, j) \in E
17
                do backlink[j] \leftarrow backlink[j] - 1
           if cluster[i] = cluster[j]
18
19
              then c \leftarrow 0
20
              else c \leftarrow w_{ii}
21
           if time[cluster[i]] + c > atlevel[j]
22
              then at level[i] \leftarrow time[cluster[i]] + c
23
           if backlink[j] = 0
24
              then Push(stack, j)
```

```
25 return Max(time[...])
```

In line 01, *stack* is initially empty. *stack* is used to traverse the nodes of the task graph. In the *for* loop from lines 02 to 05, the arrays *atlevel[...]*, *time[...]*, and *backlink[...]* are initialized to 0. *atlevel[k]* is the allocated top level (Sinnen [27]) of task  $M_k$ . Given an allocation of tasks, the *allocated top level* of a task  $M_k$  is the length of the longest path ending with  $M_k$  (excluding the computation weight of  $M_k$  that is  $w_k$ ) in which the weight of an edge is taken as 0 if the two tasks corresponding to the edge are allocated tasks. *backlink[k]* is the number of edges that are incident on the task  $M_k$ . In the *for* loop from lines 06 to 07, *backlink[...]* of

the tasks is evaluated. In the *for* loop from lines 08 to 10, the initial tasks that are not having any incident edge are pushed onto the stack using the function Push(...). In the *while* loop from lines 11 to 24, we repeatedly pop a task using the function Pop(...) from the stack and compare its allocated top level with the partial execution time of the processor on which it is allocated. We update the *time[...]* value of the processor depending on whether processor has to remain idle for some time or not (lines 12 to 15). In the *for* loop from lines 16 to 17, for each outgoing edge we decrement the corresponding *backlink[...]* value indicating the completion of communication corresponding to the edges. In lines 18 to 20, *c* is the communication time from the task  $M_i$  to the task  $M_j$  depending upon whether the two tasks are allocated on the same processor or not. In lines 21 to 22, the allocated top level of successors is updated according to the definition. In lines 23 to 24, tasks are pushed onto the stack if they have completed all of their communications. And finally in line 25, the maximum of partial execution time of processors (using the function Max(...)) is returned that is the parallel execution time of the given scheduling if the task graph *T* is a scheduled DAG.

Line 01 has complexity O(1). The *for* loop from lines 02 to 05, and 08 to 10, each have complexity O(|V|). The *for* loop from lines 06 to 07 has complexity O(|E|). The *while* loop from lines 11 to 24 has complexity O(|E|) because of the *for* loop from lines 16 to 17. Line 25 has complexity O(|V|). Therefore the algorithm *Evaluate-Time(...)* has complexity O(|V| + |E|).



# 5. A Sample Run of the CPPS Algorithm

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Fig. 1 (left): The example task graph. Fig. 2 (right): The initial clusters and the priorities between them. Parallel execution time is 39.

As an example for explaining the steps involved in the CPPS algorithm, we will consider the task graph in Fig. 1, that is taken from Kadamuddi and Tsai [14], for clustering. Tasks are shown in circles with their labels and execution times. Communication delays (edge weights) are shown in rectangular boxes along their respective edges.

Initially each task is kept in a separate cluster as shown in Fig. 2, in which clusters are shown in circles listing their constituent tasks, along with the weight of the cluster. In rectangular boxes, along the edges, are shown the priorities between different cluster-pairs. Parallel execution time for this clustering comes out to be *39*.

In Fig. 2, the cluster-pairs  $\{M_2\}$  and  $\{M_5\}$ ; and  $\{M_{14}\}$  and  $\{M_{15}\}$  are examined for merging in nonincreasing order of priorities between them. After merging, their parallel execution time comes out to be 39, and 36 respectively. Therefore the cluster-pair  $\{M_{14}\}$  and  $\{M_{15}\}$  is merged in Fig. 3, since it is the first cluster-pair that is reducing the parallel execution time from the previous value of 39 (Fig. 2) to 36 (Fig. 3).



Fig. 3 (left): Clustering after merging  $\{M_{14}\}$  and  $\{M_{15}\}$ . Parallel execution time is 36. Fig. 4 (right): Clustering after merging  $\{M_1\}$  and  $\{M_2\}$ . Parallel execution time is 33.



Fig. 5 (left): Clustering after merging  $\{M_{13}\}$  and  $\{M_{14}, M_{15}\}$ . Parallel execution time is 30. Fig. 6 (right): Clustering after merging  $\{M_1, M_2\}$  and  $\{M_4\}$ . Parallel execution time is 28.

In Fig. 3, the cluster-pairs  $\{M_2\}$  and  $\{M_5\}$ ;  $\{M_6\}$  and  $\{M_9\}$ ;  $\{M_7\}$  and  $\{M_{11}\}$ ;  $\{M_8\}$  and  $\{M_{12}\}$ ;  $\{M_{11}\}$  and  $\{M_{12}\}$ ; and  $\{M_1\}$  and  $\{M_2\}$  are examined for merging in non-increasing order of priorities between them. After merging, their parallel execution time comes out to be 36, 36, 36, 36, 36, and 33 respectively. Therefore the cluster-pair  $\{M_1\}$  and  $\{M_2\}$  is merged in Fig. 4, since it is the first cluster-pair that is reducing the parallel execution time from the previous value of 36 (Fig. 3) to 33 (Fig. 4).

In Fig. 4, the cluster-pairs  $\{M_6\}$  and  $\{M_9\}$ ;  $\{M_8\}$  and  $\{M_{12}\}$ ;  $\{M_{11}\}$  and  $\{M_{12}\}$ ;  $\{M_1, M_2\}$  and  $\{M_5\}$ ;  $\{M_7\}$  and  $\{M_{11}\}$ ;  $\{M_{10}\}$  and  $\{M_{14}, M_{15}\}$ ; and  $\{M_{13}\}$  and  $\{M_{14}, M_{15}\}$  are examined for merging in non-increasing order of priorities between them. After merging, their parallel execution time comes out to be 33, 33, 33, 33, 33, 33, and 30 respectively. Therefore the cluster-pair  $\{M_{13}\}$  and  $\{M_{14}, M_{15}\}$  is merged in Fig. 5, since it is the first cluster-pair that is reducing the parallel execution time from the previous value of 33 (Fig. 4) to 30 (Fig. 5).

In Fig. 5, the cluster-pairs  $\{M_6\}$  and  $\{M_9\}$ ;  $\{M_8\}$  and  $\{M_{12}\}$ ;  $\{M_{11}\}$  and  $\{M_{12}\}$ ;  $\{M_7\}$  and  $\{M_{11}\}$ ;  $\{M_1, M_2\}$  and  $\{M_5\}$ ;  $\{M_6\}$  and  $\{M_{10}\}$ ;  $\{M_3\}$  and  $\{M_7\}$ ; and  $\{M_1, M_2\}$  and  $\{M_4\}$  are examined for merging in non-increasing order of priorities between them. After merging, their parallel execution time comes out to be 30, 30, 30, 30, 30, 30, 30, 30, and 28 respectively. Therefore the cluster-pair  $\{M_1, M_2\}$  and  $\{M_4\}$  is merged in Fig. 6, since it is the first cluster-pair that is reducing the parallel execution time from the previous value of 30 (Fig. 5) to 28 (Fig. 6).

In Fig. 6, the cluster-pairs  $\{M_6\}$  and  $\{M_9\}$ ;  $\{M_{11}\}$  and  $\{M_{12}\}$ ; and  $\{M_7\}$  and  $\{M_{11}\}$  are examined for merging in non-increasing order of priorities between them. After merging, their parallel execution time comes out to be 28, 28, and 26 respectively. Therefore the cluster-pair  $\{M_7\}$  and  $\{M_{11}\}$  is merged in Fig. 7, since they are the first pair of clusters that is reducing the parallel execution time from the previous value of 28 (Fig. 6) to 26 (Fig. 7).



Fig. 7 (left): Clustering after merging  $\{M_7\}$  and  $\{M_{11}\}$ . Parallel execution time is 26. Fig. 8 (right): Clustering after merging  $\{M_8\}$  and  $\{M_{13}, M_{14}, M_{15}\}$ . Parallel execution time is 25.

In Fig. 7, the cluster-pairs  $\{M_6\}$  and  $\{M_9\}$ ;  $\{M_8\}$  and  $\{M_{12}\}$ ;  $\{M_7\}$  and  $\{M_{11}\}$ ;  $\{M_1, M_2, M_4\}$  and  $\{M_5\}$ ;  $\{M_7, M_{11}\}$  and  $\{M_{12}\}$ ;  $\{M_{10}\}$  and  $\{M_{13}, M_{14}, M_{15}\}$ ; and  $\{M_8\}$  and  $\{M_{13}, M_{14}, M_{15}\}$  are examined for merging in nonincreasing order of priorities between the cluster-pairs. After merging, their parallel execution time comes out to be 26, 26, 26, 26, 26, 26, and 25 respectively. Therefore, the cluster-pair  $\{M_8\}$  and  $\{M_{13}, M_{14}, M_{15}\}$  is merged in Fig. 8, since it is the first cluster-pair that is reducing the parallel execution time from the previous value of 26 (Fig. 7) to 25 (Fig. 8).

In Fig. 8, the cluster-pairs { $M_6$ } and { $M_9$ }; { $M_6$ } and { $M_{10}$ }; { $M_1$ ,  $M_2$ ,  $M_4$ } and { $M_5$ }; { $M_7$ ,  $M_{11}$ } and { $M_{12}$ }; { $M_3$ } and { $M_7$ ,  $M_{11}$ }; { $M_8$ ,  $M_{13}$ ,  $M_{14}$ ,  $M_{15}$ } and { $M_{12}$ }; { $M_1$ ,  $M_2$ ,  $M_4$ } and { $M_3$ }; { $M_7$ ,  $M_{11}$ } and { $M_{10}$ }; { $M_1$ ,  $M_2$ ,  $M_4$ } and { $M_6$ }; { $M_8$ ,  $M_{13}$ ,  $M_{14}$ ,  $M_{15}$ } and { $M_{10}$ }; { $M_7$ ,  $M_{11}$ } and { $M_{10}$ }; { $M_1$ ,  $M_2$ ,  $M_4$ } and { $M_6$ }; { $M_8$ ,  $M_{13}$ ,  $M_{14}$ ,  $M_{15}$ } and { $M_{10}$ }; { $M_7$ ,  $M_{11}$ } and { $M_8$ ,  $M_{13}$ ,  $M_{14}$ ,  $M_{15}$ } are examined for merging in non-increasing order of priorities between the cluster-pairs. After merging, their parallel execution time comes out to be 25, 25, 25, 25, 26, 28, 25, 25, 25, 27, and 23 respectively. Therefore the cluster-pair { $M_1$ ,  $M_2$ ,  $M_4$ } and { $M_8$ ,  $M_{13}$ ,  $M_{14}$ ,  $M_{15}$ } is merged in Fig. 9, since it is the first cluster-pair that is reducing the parallel execution time from the previous value of 25 (Fig. 8) to 23 (Fig. 9).

In Fig. 9, the cluster-pairs  $\{M_6\}$  and  $\{M_9\}$ ;  $\{M_6\}$  and  $\{M_{10}\}$ ;  $\{M_3\}$  and  $\{M_7, M_{11}\}$ ;  $\{M_7, M_{11}\}$  and  $\{M_{12}\}$ ;  $\{M_7, M_{11}\}$  and  $\{M_{10}\}$ ;  $\{M_1, M_2, M_4, M_8, M_{13}, M_{14}, M_{15}\}$  and  $\{M_{5}\}$ ;  $\{M_1, M_2, M_4, M_8, M_{13}, M_{14}, M_{15}\}$  and  $\{M_{12}\}$ ;  $\{M_1, M_2, M_4, M_8, M_{13}, M_{14}, M_{15}\}$  and  $\{M_6\}$ ;  $\{M_1, M_2, M_4, M_8, M_{13}, M_{14}, M_{15}\}$  and  $\{M_{10}\}$ ;  $\{M_1, M_2, M_4, M_8, M_{13}, M_{14}, M_{15}\}$  and  $\{M_{13}\}$ ; and  $\{M_1, M_2, M_4, M_8, M_{13}, M_{14}, M_{15}\}$  and  $\{M_3\}$ ; and  $\{M_1, M_2, M_4, M_8, M_{13}, M_{14}, M_{15}\}$  and  $\{M_3\}$ ; and  $\{M_1, M_2, M_4, M_8, M_{13}, M_{14}, M_{15}\}$  and  $\{M_7, M_{11}\}$  are examined for merging in non-increasing order of priorities between the cluster-pairs. After merging, their parallel execution time comes out to be 23, 23, 23, 24, 25, 26, 26, 24, 25, and 28 respectively. Therefore we cannot merge any cluster-pair in Fig. 9, since no cluster-pair is able to reduce the parallel execution time from the current value of 23 (Fig. 9) to a smaller value.



Fig. 9: Clustering after merging  $\{M_1, M_2, M_4\}$  and  $\{M_8, M_{13}, M_{14}, M_{15}\}$ . Parallel execution time is 23.

The CPPS algorithm stops at this point. The clusters generated are:  $\{M_1, M_2, M_4, M_8, M_{13}, M_{14}, M_{15}\}$ ,  $\{M_3\}, \{M_5\}, \{M_6\}, \{M_7, M_{11}\}, \{M_9\}, \{M_{10}\}$ , and  $\{M_{12}\}$ . Parallel execution time comes out to be 23.



# 6. Experimental Results

Fig. 10 (left): Parallel execution time for 50 node task graphs. Average improvement of CPPS over EZ is 15.01%; over LC is 4.57%; and over DSC is 0.59%. Fig. 11 (right): Parallel execution time for 100 node task graphs. Average improvement of CPPS over EZ is 17.43%; over LC is 4.72%; and over DSC is 0.32%.



Fig. 12 (left): Parallel execution time for 200 node task graphs. Average improvement of CPPS over EZ is 17.16%; over LC is 4.68%; and over DSC is 0.45%. Fig. 13 (right): Parallel execution time for 300 node task graphs. Average improvement of CPPS over EZ is 18.88%; over LC is 5.07%; and over DSC is 0.77%.



Fig. 14 (left): Parallel execution time for 400 node task graphs. Average improvement of CPPS over EZ is 17.92%; over LC is 4.86%; and over DSC is 0.32%. Fig. 15 (right): Parallel execution time for 500 node task graphs. Average improvement of CPPS over EZ is 19.13%; over LC is 5.11%; and over DSC is 0.86%.

The CPPS algorithm is compared with three well known algorithms: Sarkar's Edge Zeroing (EZ) algorithm [26], Kim and Browne's Linear Clustering (LC) algorithm [16], and Yang and Gerasoulis' Dominant Sequence Clustering (DSC) algorithm [32]. The algorithms are tested on benchmark task graphs of Tatjana and Gabriel [5, 6]. We have tested for *180* task graphs having the number of nodes as *50*, *100*, *200*, *300*, *400*, and *500* respectively. Each task graph has a label as  $tn_i_j.td$ . Here *n* is the number of nodes. *i* is a

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parameter depending on the edge density. Its possible values are: 20, 40, 50, 60, and 80. For each combination of n and i, there are 6 task graphs that are indexed by j. j ranges from 1 to 6. Therefore, for each n there are 30 task graphs.

We define the *performance improvement ratio* of an algorithm A over an algorithm B(PI(A, B)) for a given instance of task graph as follows:

$$PI(A, B) = 1 - PT(A) / PT(B),$$
 (14)

where PT(X) is the *parallel execution time* of algorithm *X*. For performance evaluation we take the average value of PI(A, B) for a given set of task graphs.

For the values of *n* having 50, 100, 200, 300, 400, and 500 respectively, Fig. 10 to Fig. 15 show the comparison between the four algorithms: EZ, LC, DSC, and CPPS for the parallel execution time. The average improvement of CPPS algorithm over EZ algorithm ranges from 15.01% to 19.13%. The average improvement of CPPS algorithm over LC algorithm ranges from 4.57% to 5.11%. The average improvement of CPPS algorithm ranges from 0.32% to 0.86%.

#### 7. Conclusion

We formulated Sarkar's EZ algorithm [26] for scheduling precedence constrained task graphs on a parallel system as a task dependent priority scheme in which the priority is defined for pairs of tasks. We then extended this idea to cluster dependent priority scheme in which the priority is defined for pairs of clusters. Using this idea we developed the CPPS algorithm. Using a suitable cluster dependent priority scheme of complexity O(|V|/E|(|V| + |E|)) we compared our algorithm with some well known scheduling algorithms. Experimental results demonstrated good performance of the CPPS algorithm.

We give two research directions for future work. The first line of research is to reduce the complexity of the CPPS algorithm without compromising its performance. The second line of research is to study the performance of the CPPS algorithm for more intelligent priority schemes. The example given in this paper is a static priority scheme. We can also think of dynamic priority schemes.

## 8. References

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