Fast Distributed Construction of Small $k$-Dominating Sets and Applications

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Abstract

This paper\(^1\) presents a fast distributed algorithm to compute a small $k$-dominating set $D$ (for any fixed $k$) and its induced graph partition (breaking the graph into radius $k$ clusters centered around the vertices of $D$). The time complexity of the algorithm is $O(k \log^4 n)$.

Small $k$-dominating sets have applications in a number of areas, including routing with sparse routing tables, the design of distributed data structures, and center selection in a distributed network. The main application described in this paper concerns a fast distributed algorithm for constructing a minimum-weight spanning tree (MST). On an $n$-vertex network of diameter $d$, the new algorithm constructs an MST in time $O(\sqrt{n} \log^4 n + d)$, improving on previous results.

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1 Introduction

1.1 Motivation and Results

This paper concerns the fast construction of small $k$ dominating sets, and its applications. A $k$-dominating set is a set $D$ satisfying that for every node $v$ in the graph, there is a node $w \in D$ at distance at most $k$ from it.

We introduce a new fast distributed algorithm for the problem of computing a small $k$-dominating set in a network (where “small” here means a set of size at most $n/(k+1)$ for $n \geq k+1$; it is easy to see that this is a lower bound). The algorithm also partitions the network into clusters, such that each cluster has a member of the $k$-dominating set as its center. The time complexity of the algorithm is $O(k \log^8 n)$.

This new algorithm is useful for speeding up a number of distributed tasks. For example, this type of clusters was used in [PU] in the context of routing with sparse routing tables, although no distributed protocol was given there for computing them. Hence the new construction can serve to speed up the preprocessing stage of that routing scheme.

Efficient construction for $k$-dominating sets is also applicable in the context of distributed data structures [P2], where it is proposed that a set of $k$-dominating centers can be selected for locating copies of a distributed directory. Likewise, such sets are useful for efficient selection of network centers for server placement, where it is desired to ensure that each node in the network is sufficiently close to some server (cf. [BKP]).

One application that we describe in detail in the paper concerns speeding up an algorithm for constructing a Minimum Spanning Tree (MST). Note that, informally speaking, MST can be thought of as a more “global” problem, as opposed to the $k$-Dominating Set problem which is more “local” in nature. We now present a formal notion of what we mean by a “fast” algorithm, which applies to both.

Linial [Li] proved lower bounds on the time complexity of distributed algorithms, even when assuming a very strong computational model (not assumed here) where messages can be of arbitrary length. Yet sending a message from a node to a distance $d$ still takes time $d$ in this model. Thus the lower bounds of [Li] actually correspond only to the radius (around each node) from which information must be fetched in order to solve a given problem $P$. Let this radius be $R(P)$ for a given problem $P$. We term an algorithm for $P$ neighborhood optimal if its complexity is $O(R(P))$. We use the word “fast” with respect to this notion, namely, with regard to algorithms aimed at achieving neighborhood optimality. We note that it can be shown that, assuming messages are of bounded size, there exist problems for
which no neighborhood optimal algorithm exists, due to inherent congestion. This is obvious in cases where the problem at hand involves moving around large amounts of data. However, it holds also for certain Boolean decision problems, in which the input at each vertex is a single bit, and the output is a single bit as well. In particular, it is well known that most $n$-variable Boolean functions have communication complexity $\Omega(n)$. It follows that there exist (constant diameter) $n$-vertex graphs $G$ and Boolean $n$-variable problems $P$ such that solving $P$ on $G$ requires as much as $\Omega(n^{\frac{n}{\log n}})$ time (assuming that transmitting an $O(\log n)$ bit message requires one time unit), even though $R(P)$ is clearly constant.

Note that the inherent information radius of our $k$ Dominating Set problem is $k$, hence our algorithm is neighborhood optimal up to a factor of $\log^* n$, which we term “fast”.

A question raised in [GKP] is whether the distributed MST problem can be solved in $O(Diam)$ time, where $Diam$ is the network Diameter. Clearly $R(MST) = Diam$ in certain graphs. Thus in terms of the definition above one can rephrase the question of [GKP]: “does there exist a neighborhood optimal algorithm for MST”. The algorithm presented in [GKP] has time complexity $O(n^{614} + Diam)$, hence it is neighborhood optimal for the case that $Diam > n^{614}$. Previous distributed MST algorithms had running time $O(n \log n)$ [GHS], $O(n \log^* n)[CT, G]$, and $O(n)$ [A2]. Using the fast $k$ Dominating Set algorithm we manage to present an improved MST construction algorithm whose time complexity is $O(\sqrt{n} \log^* n + Diam(G))$. Thus the MST algorithm is now neighborhood optimal for all graphs with $Diam > n^{1/2} \log^* n$. Let us hint about how the improved MST algorithm was obtained. As noted in [GKP], the main obstacle for improving the time complexity is the congestion caused by the need to send the description of many edges. To (partially) overcome the congestion, the solution presented in [GKP] used several techniques to reduce the amount of information sent by combining it. One such technique was clustering, since the construction of a cluster can be relatively local (thus consuming little time). The idea was that the cluster can represent many edges and nodes, thus reducing the amount of information to be sent. To improve the results of [GKP] we essentially replace the clustering technique used there by the clustering constructed by the $k$ Dominating Set algorithm.

1.2 Model and Definitions

We focus on the problem of devising a time efficient algorithm to construct a small $k$ Dominating Set. The network is modeled as an undirected graph $G = (V, E)$, where $V$ is the set of nodes, and $E$ is the set of communication links between them. The nodes can communicate only by sending and receiving messages over the communication links.
A \emph{k-dominating set} for a graph $G$ is a subset $D$ of vertices with the property that for every $v \in V$ there is some $u \in D$ at distance at most $k$ from $v$, measuring distance in the unweighted sense, i.e., in number of hops. (This definition is identical to that of [CN, PU] for example, but differs from other previous usages of this notion, e.g., [CGS]土.) For every such $k$-dominating set we shall define a partition $P$ of the nodes, by associating with each node $v \in V$ a dominator $D(v) \in D$, which is the node closest to $v$ in the dominating set (breaking ties arbitrarily). A small $k$ Dominating Set is of size less than or equal to $max\{1, \left\lceil \frac{k}{k+1} \right\rceil \}$. Clearly one cannot do better than that in the general case.

We use the common assumption that nodes have unique identifiers, and that each edge $e \in E$ is associated with a distinct weight $\omega(e)$, known to the adjacent nodes. One can do without (either) one of these two assumptions. Having distinct edge weights helps as it guarantees the uniqueness of the MST. Yet as pointed out in [GHS], this is not essential, as such weights can always be generated artificially using the distinct node identifiers. However, if the graph has neither distinct edge weights nor distinct node identifiers, then no distributed algorithm can compute an MST with a bounded number of messages [An80].

For every subgraph $F$ of the network, let $Diam(F)$ denote $F$’s diameter, i.e., the maximum distance on $F$ between any two vertices of $F$. Similarly, let $Rad(F)$ denote the radius of $F$, i.e., the maximum distance on $F$ from any vertex of $F$ to some center vertex of $F$ (where the center of $F$ is chosen so as to minimize this distance). For a collection $P$ of subgraphs, let $Diam(P)$ (resp., $Rad(P)$) denote the maximum diameter (resp., radius) of a subgraph in $P$.

In order to be able to concentrate on the central issue of time complexity, we shall follow the common trend of stripping away unessential complications. In particular, we ignore the communication cost of our algorithm (i.e., the number of messages it uses). We also assume that the computation performed by the network is \emph{synchronous}. (This assumption is not essential, since our decision to ignore communication costs allows us to freely use a \emph{synchronizer} of our choice; for example, we can use the simple synchronizer $\alpha$ of [A1] whose cost in an asynchronous network is one message over each edge in each direction per round of the synchronous algorithm.) We note that the synchronous nature of the network enables all the nodes to start a subroutine simultaneously. This is very useful when dealing with subroutines whose running time is less than the diameter, where it does not make sense to start them by a “wake up” broadcast (that, by itself, may take diameter time).

Still, we shall not adopt the extreme model employed in previous studies of locality issues [Li], in which messages of arbitrary size are allowed to be transmitted in a single time unit, since in this model the refined distinctions we focus on here disappear. Clearly, if unbounded-
size messages are allowed, then the MST problem, for instance, can be trivially solved in time $O(Diam(G))$ by collecting the entire graph's topology into a central node, computing an MST locally and broadcasting the result throughout the network.

Consequently, we will assume the more realistic and more common model in which messages have size $O(\log n)$, and a node may send at most one message over each edge at each time unit. We will also make the assumption that edge weights are polynomial in $n$, so an edge weight can be sent in a single message. (This assumption is required for the time analysis of the previous algorithms that use edge weights or nodes identity, e.g. [GHS, A2, GKP].)

Let us now define the Minimum Spanning Tree task solved later on as an application of the $k$ Dominating Set algorithm. The goal is to have the nodes (processors) cooperate to construct a minimum weight spanning tree (MST) for $G$, namely, a tree covering the nodes in $V$ whose total edge weight is no greater than any other spanning tree for $G$.

1.3 Paper Structure

In the next section we describe some tools. We then use these tools in Section 3 for developing an algorithm for constructing a small $k$ Dominating Set on a tree. Next, in Section 4 we use the previous algorithm to develop an algorithm for $k$-dominating set construction on a general network. The new MST algorithm is described in Section 5.

2 Basic Algorithms for Small $k$-Dominating Sets

As discussed next in Subsection 2.1, it is known that a “small” ($\text{size } n/(k+1)$) $k$-dominating set always exists in an $n$-node graph. The fast distributed algorithm for computing such a set is developed in stages. In this section we only present the existence proof and then (in Subsection 2.2) give a “slow” distributed procedure, computing a $k$-dominating set on a graph $G$ in time $O(Diam(G))$. In the next section we give a fast algorithm for computing a $k$-dominating set on an $n$-node tree $T$ in time $O(k \log^4 n)$. In the following section we extend the result to a general graph.

2.1 Existence of Small $k$-Dominating Sets

The following is well-known (cf. [PU]).
For every connected graph $G$ of $n$ vertices and for every $k \geq 1$ there exists a $k$-dominating set $D$ such that $|D| \leq \max\{1, \left\lfloor \frac{n}{k+1} \right\rfloor \}$.

To motivate the distributed algorithm developed for the $k$-dominating set problem in Subsection 2.2, let us overview the (standard) proof outlined for the above lemma in [PU].

Let $T$ be an arbitrary rooted spanning tree for $G$ and denote its depth (the distance from the root) by $h$. If $k \geq h$ then $D$ may consist of the root alone. Otherwise, divide the vertices of $V$ into levels $T_0, \ldots, T_h$ according to their height in the tree, assigning all the vertices of height $i$ to $T_i$. Now merge these sets into $k+1$ sets $D_0, \ldots, D_k$ by taking

$$D_i = \bigcup_{j \geq 0} T_{i+j(k+1)}$$

(i.e., $T_i$ and every $(k+1)$st level thereafter). See Figure 1. Clearly every $D_i$ is a $k$-dominating set, and since these sets form a complete disjoint partition of $V$, at least one of the sets is of size at most $\left\lfloor \frac{n}{k+1} \right\rfloor$.

### 2.2 Distributed Computation of the Set in Diameter time

The proof of Lemma 2.1 suggests a sequential algorithm to find the $k$-dominating set claimed in the lemma. However, we need to show a distributed algorithm that can implement this computation in time $O(Diam(G))$ given a graph $G$ and a root node $r$.

To initialize this process, it is necessary to construct the BFS tree, mark the levels and let each node know its depth, the total tree depth, and the set $D_i$ it belongs to. This is
1. Perform a distributed breadth-first search (BFS) on the graph, starting from the root $r$ (terminating at $r$). Label every node $v$ by its distance from $r$, $\text{Depth}(v)$.

2. A node $v$ joins the unique set $D_l$ for $0 \leq l \leq k$ such that $\text{Depth}(v) = l \pmod{(k + 1)}$.

3. Perform a “broadcast and echo” process from $r$ to learn the largest distance label $M = \max_v \{\text{Depth}(v)\}$ (namely, the depth of the tree). Specifically, $r$ broadcasts a request throughout the tree. Each leaf receiving the request echos back a message containing its depth. An internal node that received echos from all its children sends an echo to its parent, containing the maximum depth label received from its children.

4. Broadcast the tree depth $M$ to all nodes.

Figure 2: Procedure Initialize.

done by Procedure Initialize, whose high-level description is given in Fig. 2. We assume standard distributed implementations for basic tasks such as broadcast on a tree, or BFS tree construction. For instance, the latter task can be performed (in a synchronous setting) by elementary flooding techniques, in time $O(\text{Diam})$.

For clarity of presentation, the algorithm of Fig. 2, as well as other algorithms later in the paper, are given by a high-level description, omitting the distributed implementation of well-understood stages of the algorithm. For example, we omit the distributed implementation of BFS in its first stage, as well as the termination detection performed by each node for stage 1, in order to perform its stage 2.

Next, we give (in Fig. 3) a procedure $\text{Census}(l)$ whose task is to count the number of vertices in the set $D_l$. The procedure operates by a simple convergecast process on the tree from the leaves upwards. Each node $v$ holds a counter $\text{counter}(v, l)$, used by the procedure for counting the number of $D_l$ nodes in the subtree rooted at $v$.

**Lemma 2.2** At the end of Procedure $\text{Census}(l)$, $\text{counter}(v, l)$ holds the number of $D_l$ nodes in the subtree rooted at $v$, for each node $v$. In particular, for the root $r$, $\text{counter}(r, l) = |D_l|$.

We combine the above procedures into an algorithm named $\text{Diam\_DOM}$ (in Fig. 4) for selecting a small $k$-dominating set. The key to an efficient combination lies in staggering the executions of the different $\text{Census}$ operations, so that they are pipelined, and do not disrupt one another.

**Lemma 2.3** Algorithm $\text{Diam\_DOM}$ computes the $k$-dominating set claimed in Lemma 2.1, and its time complexity is $6 \cdot \text{Diam}(G) + k$. 

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**Code** for a node $v$ in $\text{Depth}(v) = i$, with children $u_1, \ldots, u_p$.

**Assumption:** The procedure starts operating at time $t_0$.

```
/* M is the tree depth */
```

1. If $v$ is not a leaf: At time $t_0 + (M - i) - 1$, receive from children messages containing the values of $\text{counter}(u_j, l)$ for $1 \leq j \leq p$.

2. At time $t_0 + (M - i)$ do:
   (a) Compute $\text{Below} = \sum_j \text{counter}(u_j, l)$.
   (b) Set
       
       $$
       \text{counter}(v, l) = \begin{cases} 
       1 + \text{Below}, & \text{if } v \in D_i; \\
       \text{Below}, & \text{otherwise}. 
       \end{cases}
       $$
   (c) If not the root, then send $\text{counter}(v, l)$ to parent in the tree.

**Figure 3:** Procedure $\text{Census}(l)$.

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1. Apply Procedure $\text{Initialize}$.

2. Let $t_1$ be the time unit just after the completion of this procedure.
   For $l = 0$ to $k$ do:
   At time $t_1 + l$, start Procedure $\text{Census}(l)$.

3. After completion of all $\text{Census}(l)$ executions, the root $r$ computes $l$ such that $\text{counter}(r, l)$ is the minimum, and broadcasts it on the tree. $D_i$ is the output $k$-dominating set.

**Figure 4:** Algorithm $\text{Diam_DOM}$.
Proof: It is immediate from Lemma 2.2 that the algorithm is an implementation of the construction in the proof of Lemma 2.1, and thus it computes a dominating set as claimed therein. The time requirements of Procedure Initialize, constructing the synchronous BSF and performing the “broadcast and echo” process, is $4 \cdot Diam(G)$. At this point every node can calculate the time $t_1$. The lemma now follows from the crucial observation that the separate executions of Procedure Census($l$) for $0 \leq l \leq k$ never disrupt one another, since the participation of a given vertex $v$ of depth $i$ in a particular execution Census($l$) is limited to (possibly) receiving messages at time $(t_1 + l) + (M - i) - 1$ and sending a message at time $(t_1 + l) + (M - i)$, hence no collisions occur. Therefore the last Census operation terminates after $Diam(G) + k$ time, and the final broadcast requires $Diam(G)$ additional time units.

Remark: By employing a slightly more involved pipelining of the Census operations, it is possible to reduce the total completion time to $6 \cdot Diam(G)$. The key observation is that the Census operations can be staggered on the basis of their start level rather than their start time. The point is that not all Census operations need to start at the nodes of depth $M$; in particular, a Census($l$) operation corresponding to a set $D_l$ whose lowest level is at depth $M - j$ can start from that level. Hence the $k + 1$ Census operations can all start at the same time $t_1$, only at different levels of the tree. This improvement does not lead to any asymptotic improvement in the complexity of our algorithm, hence we preferred the simpler version of the algorithm.

3 Distributed Computation of a Small $k$-Dominating Set on a Tree

In this section we concentrate on a fast algorithm for computing a small $k$-dominating set on a tree $T$. (We use the slow algorithm of Subsect. 2.2 as a subroutine.) Let us start with an overview of the algorithm.

Algorithm Diam_DOM of the previous section requires time $O(Diam(G))$. Hence our approach to building a small $k$-dominating set is based on first partitioning the tree into a collection of subtrees (or clusters), such that the depth of each subtree is at most $O(k)$. Then Algorithm Diam_DOM can be applied on each subtree separately, and the resulting dominating sets can be merged to yield the dominating set for the entire graph.

A source of difficulty is that the subtrees we construct must not be too small: for a subtree of $l < k$ vertices, any dominating set must still contain at least one vertex, which
does not meet the desired ratio of $1/(k+1)$ between dominators and dominated vertices. Hence our goal is to partition the tree into subtrees of size $k+1$ or more and depth $O(k)$.

Formally, let us introduce the following definition.

**Definition 3.1** A $(\sigma, \rho)$ spanning forest of a graph $G(V,E)$ is a collection of disjoint trees $\{T_1, \ldots, T_m\}$ with the following properties:

1. the trees consist of edges of $E$ and span all the nodes of $V$,
2. each tree contains at least $\sigma$ nodes, and
3. the radius of each tree is at most $\rho$.

In Subsection 3.1 we show a fast distributed algorithm for computing a 1-dominating set with certain useful properties. This is next used in Subsection 3.2 to construct a fast distributed algorithm for partitioning a tree into $O(k)$-depth, $\Omega(k)$-size clusters. Finally, in Subsection 3.3 we present a fast distributed algorithm that computes the desired “small” $k$-dominating set by first applying the fast partitioning algorithm of Subsection 3.2.3, and then running Procedure $\text{Diam\_DOM}$ of Subsection 2.2 inside each (low-diameter) cluster.

### 3.1 A Distributed Balanced Dominating Set Algorithm

**Definition 3.2** A balanced dominating set of a graph $G$ with $n$ nodes is a set $D$ of nodes in $G$, and an associated partition $P$ of $G$’s nodes with the following properties:

(a) $|D| \leq \lceil \frac{n}{2} \rceil$.

(b) $D$ is a dominating set.

(c) In the partition $P$, each cluster has at least two nodes (including the node in $D$).

Algorithm $\text{Balanced\_DOM}$ is a variant of the dominating set algorithm $\text{Small\_Dom\_Set}$ in [GKP]. That algorithm computed an ordinary (not necessarily balanced) dominating set on a given tree, namely, a set satisfying properties (a) and (b) above, but not property (c). Algorithm $\text{Small\_Dom\_Set}$ made use of a procedure for computing a maximal independent set (MIS) as a black box. (A vertex set $M$ is an MIS if it is independent and cannot be increased, i.e., every node $v \in V \setminus M$ has a neighbor in $M$.) Specifically, it used the deterministic distributed MIS algorithm of [PS]. Since Algorithm $\text{Small\_Dom\_Set}$ uses the MIS procedure only for computing MIS on a tree, it is possible to replace the procedure of
Figure 5: An execution of Algorithm BalancedDOM. Step (1) generates the dominating set $D$ depicted by the black vertices in the first row, and the partition $P$ whose clusters are drawn by dashed lines. Steps (2) and (3) transform $D$ and $P$ into the configuration depicted in the second row, and Step (4) yields the final structure, depicted in the third row of the figure.

[PS] with the faster procedure of [GPS] (which computes an MIS on an $n$-vertex tree in time $O(\log^3 n)^2$. Thus the following can be said about Small-Dom-Set, preparing to use it in Algorithm BalancedDOM:

**Lemma 3.3 [GKP]** There exists a distributed procedure Small-Dom-Set which, applied to a given $n$-vertex tree $T$ for $n \geq 2$, computes (in a synchronous manner) a dominating set $D$ of size at most $\lceil n/2 \rceil$ using $O(\log n)$-bit messages, and its time complexity is $O(\log^3 n)$. Furthermore, the output of the procedure has the property that each node in $D$ has some neighbor outside $D$.

The property described in the last statement of the lemma is not argued in [GKP], but can easily be shown to hold.

Note that even with these changes, Algorithm Small-Dom-Set does not guarantee property (c). The modified variant, Procedure BalancedDOM described in Fig. 6, will ensure also the additional property that every cluster has at least two nodes, namely, the partition $P$ will not contain any singleton set. The process is illustrated in Figure 5.

**Lemma 3.4** On a tree of $n \geq 2$ vertices, Algorithm BalancedDOM constructs a balanced
1. Perform Algorithm Small-Dom-Set of [GKP] on the tree $T$. Let $D$ and $P$ be the output dominating set and partition.

2. For every singleton $\{v\}$ in $P$, $v$ quits the set $D$, and selects an arbitrary neighbor $u \notin D$ as its dominator, $D(u) = u$.

3. Each node $u \notin D$ that was selected as a dominator in step (2) adds itself to $D$, quits its old cluster in $P$, and forms a new cluster, consisting of itself and all the nodes that chose it as their dominator.

4. Consider a node $v \in D$ whose cluster in the modified partition $P$ is now a singleton. Then $v$ chooses arbitrarily one node $u$ that was in $v$'s cluster in the original partition and left it in step (2), and joins $u$'s cluster. Also, $v$ quits the dominating set.

$\text{Figure 6: Algorithm Balanced\_DOM.}$

dominating set and requires time $O(\log^4 n)$.

Proof: We first observe that step (2) can always be performed, by the last property in Lemma 3.3.

Next, we claim that whenever some node $v$ is left as a singleton after step (3) of the algorithm, a node $u$ as defined in step (4) of the algorithm exists. To see this, let us examine the status of $v$ in the original partition produced on step (1). First we exclude the possibility that $v$ has joined $D$ in step (3), since in that case there would be at least one other node $u$ that chose $v$ as its dominator in step (2) and joined its cluster. Hence $v$ must have belonged to the original dominating set $D$ produced in step (1). Next, we note that $v$ cannot have belonged to a singleton in the original partition of step (1), since all of those have quit $D$ by step (2). Hence $v$'s original cluster must have contained some other vertex $u$ that has left it by step (3).

Now properties (b) and (c) are immediate from the algorithm. Property (a) follows from property (c). The time bound follows from Lemma 3.3, since all the additional steps require constant time.  

3.2 Fast Distributed Partitioning of a Tree

In this subsection we show how to partition a given tree of size $n \geq k + 1$ into clusters of sufficient size (at least $k + 1$) and small radius. For the sake of clarity we develop the algorithm in several steps. In Subsection 3.2.1 we start by giving a very simple algorithm.
For \( \lceil \log(k + 1) \rceil \) times do:

1. Perform Algorithm $\text{Balanced\_DOM}$, assigning each node not in the dominating set to a cluster with an arbitrary neighbor in the dominating set.

2. Contract each cluster to one node.

End_For

Figure 7: Algorithm $\text{DOM\_Partition}\_1(k)$.

that constructs a \((k + 1, O(k^2))\) spanning forest, with time complexity which is \(O(k^2 \log^* n)\).

In Subsection 3.2.2 we show how to limit the growth of the trees in the forest, so that the output is a \((k + 1, O(k))\) forest, with running time which is \(O(k \log k \log^* n)\). Finally, in Subsection 3.2.3 we show how to improve the running time by a factor of \(\log k\), using an idea similar to those used before, e.g., in [AG, G, CT, A2, JM].

3.2.1 Constructing a \((k + 1, O(k^2))\) Spanning Forest

The algorithm $\text{DOM\_Partition}\_1(k)$ described in Figure 7 for constructing this partition operates on a tree $T$ via repetitive applications of Procedure $\text{Balanced\_DOM}$. Each application constructs a balanced dominating set and an associated partition on the tree, and then contracts each cluster in the partition into a single node, thus forming a (smaller) tree for the next iteration.

Let us remark that the distributed implementation of the contraction in the algorithm requires us to appoint for each cluster a center node, that will from now on perform the operations for the whole cluster, while the other nodes in the cluster will just serve as links between it and the other cluster centers.

For this simple algorithm, the properties of Algorithm $\text{Balanced\_DOM}$ enable us to prove the following two bounds. First, note that in each iteration of the algorithm, the minimum size of a cluster at least doubles. Next, note that in each iteration the maximum radius of a cluster increases by a factor of at most 4. (More precisely, denoting the maximum cluster radius after the $i$th iteration of the algorithm by $f(i)$, we note that $f(i + 1) \leq 3f(i) + 1$.) Fig. 8 illustrates the extreme cases of these two bounds.

Hence as the number of iterations is \(\lceil \log(k + 1) \rceil\), we get the following lemma. (Since we prove later, in Subsection 3.2.3, a much stronger result for Algorithm $\text{DOM\_Partition}(k)$, we omit the formal proof here.)

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Lemma 3.5 Algorithm DOM\_Partition\_1(\(k\)) requires time \(O(k^2 \log^2 n)\). Every cluster \(C\) in the output partition \(P\) satisfies \(|C| \geq k + 1\) and \(\text{Rad}(C) \leq 4k^2\).  

3.2.2 Constructing a \((k + 1, O(k))\) Spanning Forest

The straightforward variant of the algorithm presented in the previous subsection suffices to guarantee the requirements of cluster size \(k + 1\) or more and cluster radius \(O(k^2)\). However, we would like to modify this algorithm so that it has a better running time and a better bound on the radius of each cluster in the partition, namely, \(O(k)\). This is achieved by employing the following simple idea: while constructing the partition, clusters that have reached depth greater than \(k\) are erased from the tree (i.e., they stop participating in the cluster merging process), and hence they stop growing.

The algorithm, named DOM\_Partition\_2(\(k\)), is complicated by the fact that whenever a (sufficiently deep) cluster stops participating, and is effectively eliminated from the tree, it causes the tree to split into a forest of subtrees. Subsequent invocations of Procedure Balanced\_DOM therefore execute on each of these subtrees separately. Hence we formally describe the algorithm as applied to a forest of trees \(T\) (initially containing the single tree \(T\)).

One problem with this idea is that the “shallow” clusters may not be able to grow solely by joining other shallow clusters. In particular, it may be the case that a shallow cluster has no neighboring shallow cluster; all its neighboring clusters have grown beyond the specified depth bound and were consequently eliminated from the tree. On the other hand, if such a shallow cluster is allowed to join a neighboring deep cluster, how can we control the growth of the deep cluster, so that its diameter will not become much larger than \(k\)?

The solution is to attempt to merge shallow trees together as long as possible, and merge
shallow trees to neighboring deep trees only at the very last step of the algorithm (i.e., only once). This maintains a reasonable bound on the depth of the resulting trees.

Algorithm DOM\textsubscript{Partition\_2}(k) is described in detail in Fig. 9. In every intermediate iteration $i$, the algorithm associates with each node $v$ a cluster $C_i(v)$ consisting of the original nodes (of the input tree $T$) of the subtree that was contracted into $v$. Also, the algorithm maintains a forest $\mathcal{T}_i$ of trees to be handled in the $i$th iteration.

The algorithm is again described on a high-level, ignoring issues related to the distributed implementation, such as how individual nodes know which subtree (or cluster) they belong to at any given moment, how a cluster is (logically) contracted into a single node, or how the sets $S$ and $P_{\text{out}}$ are globally maintained. These issues are handled by standard techniques. Let us briefly remark on these implementation issues. Each cluster is maintained in an implicit way, in the graph itself, by the participating nodes. In particular, the actions related to each cluster $C$ in the algorithm are managed by some center node $v$ in it. Every other cluster member maintains a cluster center variable $CC_i$ for each level $i = 1$ to $\lceil \log(k + 1) \rceil$, storing the id of its $i$th level cluster center, and the sets $C_i$ are implicitly defined by these variables. The participation of a non-center node in the protocol is limited to facilitating the communication between its center and (the centers of) other clusters. When a collection of level $i$ clusters $C_1, \ldots, C_p$ get merged into a larger cluster $\hat{C}$ on level $i + 1$, the center of one of the clusters $C_j$ (say, the dominator around which $\hat{C}$ was formed) becomes the center of $\hat{C}$, and this information is disseminated to all the members of the original clusters $C_j$. The global set $S$ (and similarly $P_{\text{out}}$) is maintained in a similar way, via local variables indicating (at each cluster center) whether the cluster belongs to $S$.

Lemma 3.6 The collection $P_{\text{out}}$ output by Algorithm DOM\textsubscript{Partition\_2} is a partition of $T$.

Proof: Note that whenever we erase a node $v$ from some tree during iteration $i$, we make sure to move the corresponding set $C(v)$ of original nodes to $P_{\text{out}}$ or to $S$. Also note that in the final step (4), each cluster $C(v)$ in $S$ is either moved to $P_{\text{out}}$ or merged into some cluster of $P_{\text{out}}$. (Let us remark, b.t.w., that despite the fact that each cluster in $S$ has failed the test in Step 3b, there may be clusters $C(v) \in S$ passing the test in Step 4a, as the tests are slightly different.)

In particular, note that in case $C(v)$ is of size $k$ or smaller, step (4b) always succeeds in finding a neighboring cluster $C'$ for $C(v)$. This is because $v$ entered $S$ in step (3c) as a result of remaining a single node in some tree $T''$ in $T''$. Since the input tree was of size $k + 1$ or larger, and $v$ represents a cluster $C(v)$ of fewer than $k + 1$ nodes, $C(v)$ must have some neighboring cluster $C(w)$ that was already removed from the tree earlier (either in this iteration or in an earlier one). This $C(w)$ could not have been moved to $S$, since at that time
1. Set $\mathcal{T}_0 = \{ T \}$ and $S = P_{\text{out}} = \emptyset$.
   /* $P_{\text{out}}$ is the output partition. $S$ collects small isolated trees. */

2. For every vertex $v$ set $C_1(v) = \{ v \}$.

3. For $i = 1$ to $\lceil \log(k + 1) \rceil$ do:
   
   (a) /* Form the new forest $\mathcal{T}_{i+1}$. */
       For every tree $T'$ in the forest $\mathcal{T}_i$ do:
       
       • Perform Algorithm $\text{BalancedDOM}$ on $T'$, yielding a partition $P$.
       • Contract each cluster $C$ of $P$ into a single node $v$, and set
         $C_{i+1}(v) = \bigcup_{w \in C} C_i(w)$.

       End_For
       Let $\mathcal{T}'$ be the resulting forest of contracted trees.

   (b) /* Remove sufficiently large clusters. */
       For every tree $T'$ in the forest $\mathcal{T}'$, and for every node $v$ in $T'$, if the tree spanning
       $C_{i+1}(v)$ has depth $k + 1$ or greater then do:
       
       • Erase $v$ from the tree $T'$ (thus possibly splitting $T'$ into a forest).
       • Make $C_{i+1}(v)$ a cluster in the output partition $P_{\text{out}}$.

       End_For
       Let $\mathcal{T}''$ be the resulting forest.

   (c) /* Remove lone small clusters. */
       For every tree $T''$ in the forest $\mathcal{T}''$, if $T''$ consists of a single node $v$ then do:
       
       • Erase $T''$ from the forest $\mathcal{T}''$.
       • Add the cluster $C_{i+1}(v)$ to the set $S$.

       End_For
       Let $\mathcal{T}_{i+1}$ be the resulting forest.

End_For

4. For each cluster $C(v)$ in the set $S$ do:
   
   (a) If $|C(v)| > k$ then move it to $P_{\text{out}}$ as is.
   (b) Else Merge $C(v)$ into some neighboring cluster $C'$ in $P_{\text{out}}$.

End_For

5. Return $P_{\text{out}}$.

Figure 9: Algorithm DOM\_Partition\_2($k$).
w still had v (or some “ancestor” of v) as a neighbor. Hence the cluster \( C(w) \) was moved to \( P_{out} \), and thus it can be used in step (4b) of the procedure.

It remains to prove only that at the end of the main loop, the collection \( T \) is empty. This is proved as follows. Let \( s_i \) denote the minimum size of a cluster \( C(v) \) corresponding to a node v occurring in some tree \( T \) in \( T_i \), namely,

\[
s_i = \min_{T \in \mathcal{T}_i} \min_{v \in T} |C(v)|.
\]

By Lemma 3.4 and property (c) in Def. 3.2, \( s_i \) at least doubles in every iteration \( i \). Since initially \( s_i = 1 \), at the end of iteration \( i = \lceil \log(k+1) \rceil \) we have \( s_{i+1} \geq k + 1 \), hence \( T_{i+1} \) must be empty.

**Lemma 3.7** If the input tree \( T \) is of size \( n \geq k + 1 \), then every cluster \( C \) in the output collection \( P_{out} \) has the following properties.

(a) \(|C| \geq k + 1\).

(b) \(\text{Rad}(C) \leq 5k + 2\).

**Proof:** Let us first prove Property (a). For the set \( P_{out} \) obtained at the end of the main loop, the claim follows immediately by the rule of step (3b). For the final output, note that the claim must remain true since sets in \( P_{out} \) can only increase in the last step (through the merge of sets from \( S \)), and sets moved from \( S \) intact are sufficiently large too.

Let us now turn to Property (b). Consider first a cluster \( C_{i+1}(v) \) from the set \( P_{out} \) obtained at the end of the main loop. This cluster was constructed in step (3a). The operation that constructed it consisted of merging some clusters \( C_t(w) \) for \( w \in C \). By property (b) in Def. 3.2, the radius of this cluster obeys the relation \( \text{Rad}(C_{i+1}(v)) \leq 3r + 1 \), where \( r = \max_{w \in C} \text{Rad}(C_t(w)) \). (See Figure 8.) Claim (b) now follows for \( C_{i+1}(v) \) by the fact that the clusters \( C_t(w) \) that formed it had radius \( k \) or smaller (since otherwise they would have been removed to \( P_{out} \) in some previous iteration), hence \( \text{Rad}(C_{i+1}(v)) \leq 3k + 1 \).

For the final output, note that the merges performed in the last step are shaped as “stars” composed of an original cluster \( C \in P_{out} \) merged with a collection of neighboring clusters from \( S \). Each such cluster from \( S \) has radius \( k \) or smaller (since, again, had its radius been \( k + 1 \) or more, it would have found its way into \( P_{out} \)). Thus the radius of each set resulting from such a “star merge” is at most \((3k + 1) + 2k + 1\). The claim follows.

We have established that Algorithm DMPartition_2(k) constructs the desired \((k + 1, O(k))\) spanning forest. However, the construction requires time \( O(k \log k \log^* n) \). This is
because each application of Procedure \texttt{Balanced}\_\texttt{DOM} takes time $O(\log^* n)$, except that its
distributed implementation on the $i$th iteration is slowed down by a factor proportional to
the maximum diameter of clusters at that iteration. This diameter is bounded by $k$, hence
the total time bound of $\sum_{i \leq \lceil \log (k+1) \rceil} O(k \log^* n) = O(k \log k \log^* n)$.

### 3.2.3 An $O(k \log^* n)$ Time Construction

In this section we show how to improve the running time of Algorithm \texttt{DOM}\_\texttt{Partition\_2}(k)
by a factor of $\log k$. Let us first give an intuitive explanation for the extra $\log k$ factor in
Algorithm \texttt{DOM}\_\texttt{Partition\_2}. Note that after the $i$th iteration the diameter of a cluster may
be as high as $4^i$ (although no greater than $k$). Thus, after $\frac{1}{2} \log (k + 1)$ iterations there may
be a cluster whose diameter is about $\frac{k}{2}$. Each of the remaining $\Omega(\log k)$ iterations will thus
last $\Omega(k \log^* n)$ time, which brings the total complexity to $O(k \log k \log^* n)$. Note that many
clusters may have grown at a slower pace, and thus $O(\log k)$ iterations may still be necessary.

The idea behind reducing the complexity is to execute each phase $i$ only for $O(2^i)$ time. Clusters that are too large in phase $i$ (namely, whose depth is larger than $2^i$) will thus not participate in the phase (except for the $2^i$ steps required to determine that their depth is indeed too large).

We still face the problem of how to handle a small (hence participating) cluster all of
whose neighboring clusters are large (hence non-participating). As before, the solution we
employ is to allow such a small cluster to merge onto its larger neighbor, but make sure that
this type of merging cannot increase the depth of the formed cluster beyond $k$. Once a depth
of $\Omega(k)$ is reached, the cluster is removed from the tree, as in the previous algorithm, and is
not merged onto again until the very last step.

Ensuring the depth bound on the clusters is achieved by maintaining an accurate count
of the depth at each node in the cluster. When a small cluster merges onto a large one, it
will update the depth counters at each of its nodes, to reflect the depth w.r.t. the new root
(which is the root of the large fragment). A leaf in a large cluster will permit other clusters
to merge to it only so long as the depth it records is at most $k$.

We use the following notation. For a node $v$ belonging to a cluster $C$, let $\text{Depth}(v)$
denote the depth of $v$ in $C$, namely, its distance from the root. (When the algorithm
starts, $\text{Depth}(v)$ is initialized to zero for every $v$.) We will use a set $\mathcal{W}$ to keep nodes
marked as non-participating (and thus waiting) at a given iteration. Initially, we set $\mathcal{W} = \emptyset$.
Algorithm \texttt{DOM}\_\texttt{Partition}(k) is obtained by adding the following instructions to Algorithm
\texttt{DOM}\_\texttt{Partition\_2}(k). Inside the main loop, just before step (3a), we add the steps listed...
in Fig. 10. (Again, this high-level description is implemented via standard distributed techniques, as briefly discussed earlier.)

(3-I): Return the nodes in the set $W$ to the forest $T_i$. 
/* This may reunite trees that were split in previous iterations */

(3-II): For every cluster $C_i(v)$, check whether the cluster’s radius is larger than $2 \cdot 2^i$. 
If not, then notify all nodes in $C_i(v)$ that the cluster does participate in iteration $i$. 
Term such a cluster a participating cluster. /* Note that within $O(2^i)$ time every node in the cluster 
knows whether the cluster is a participating cluster, or a non-participating one. */

(3-III): Remove from the trees of $T_i$ all the nodes $u$ representing non-participating clusters $C_i(u)$, 
and place them in the set $W$.

(3-IV): For every tree $T'$ in $T_i$, if $T'$ consists of a single node $v$ (representing a lone participating 
cluster $C' = C_i(v)$) then do:

(A) If there exists a node $u \in W$ (representing a non-participating cluster $C'' = C_i(u)$) such that 
$C''$ contains a node $w$ that neighbors $C'$, and is of depth $\text{Depth}(w) \leq k$ in $C''$ then do:

(i) Erase $T'$ (and node $v$) from the forest $T_i$ and move the nodes of $C'$ to the cluster $C''$. 
/* Now $u$ represents a cluster $C_i(u)$ composed of $C' \cup C''$, 
and $w$ becomes the parent of the nodes of $C'$. */

(ii) Use $\text{Depth}(w)$ to assign the correct $\text{Depth}$ values to every node $z$ in $C'$ (i.e., their new depth in the combined cluster $C_i(u)$).

(B) Else /* no such cluster $C''$ exists */

erase $T'$ from the forest $T_i$ and add cluster $C_i(v)$ to the set $S$.

Figure 10: Additional section for Algorithm DOM Partition($k$).

An additional modification to Algorithm DOM Partition.2 is necessary due to the fact 
that the test performed in step (3b) for the radius of the cluster cannot be implemented 
directly, since it might cost $\Omega(k)$ time. Instead, this test is implemented by relying on the 
Depth counters maintained at the nodes. Namely, each node $v \in C$ detecting $\text{Depth}(v) > k$ 
must notify the root of $C$ of that fact. This will still cost $\Omega(k)$ time (or actually, exactly $k$ time units to notify the root), but only once, as this cluster will perform step (3b) immediately 
after that. We do not bring here the pseudo code for this different implementation of the 
Depth test.

Lemma 3.8 The collection $P_{out}$ output by Algorithm DOM Partition($k$) is a partition of 
$T$. Furthermore, if $T$ is of size $n \geq k + 1$, then every cluster $C$ in $P_{out}$ has the following 
properties.

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(a) \(|C| \geq k + 1\).

(b) \(\text{Rad}(C) \leq 5k + 5\).

**Proof:** The proof is similar to that of Lemmas 3.7 and 3.6. Note that a node that does not participate in phase \(i\) has size at least \(2^i\). Thus a cluster which reaches size \(k + 1\) is eventually put in \(P_{out}\) as in Lemma 3.6. For clusters moved to \(S\) in the additional step (3-IV(B)), note that by arguments similar to those in the proof of Lemma 3.6, such a cluster must have a large cluster to merge with in \(P_{out}\), since at the point it was moved to \(S\), all the clusters that are connected to it were either already in \(P_{out}\) or non-participating with depth \(k + 1\) or greater. We conclude that the output is indeed a partition.

Similarly, property (a) follows either as in Lemma 3.7 for a cluster that is put in \(P_{out}\) in step (3b), or else it follows immediately by the rule of step (3-IV)(A)(i), noting that \(C'' = C_i(u)\) is already sufficiently large for every \(u \in W\). Property (b) follows again as before for clusters in \(P_{out}\). For clusters generated in the last iteration by merging some small clusters \(C'\) onto a non-participating cluster \(C'' = C_i(u)\) (for \(u \in W\)) in the additional step (3-IV), the property follows from the rule in step (3-IV)(A)(i), noting that the node \(w\) to which \(C'\) is connected is at depth \(k\) or less, and the radius of \(C'\) itself is at most \(2^{i+1} \leq 2k + 2\), hence the radius of the cluster resulting from this merge step is at most \(5k + 5\). ■

**Lemma 3.9** Algorithm \(\text{DOM-Partition}(k)\) requires time \(O(k \log^* n)\).

**Proof:** This is clear from step (3-I), which ensures that iteration \(i\) takes only \(O(2^i)\) time. ■

### 3.3 Fast Distributed Computation of Small \(k\)-dominating sets

The fast algorithm \(\text{Fast-DOM-T}\) is now obtained simply as the combination of the previous algorithms. Namely:

**Algorithm** \(\text{Fast-DOM-T}\)

1. Perform Algorithm \(\text{DOM-Partition}(k)\), yielding the partition \(P'\).

2. Perform Algorithm \(\text{Diam DOM}\) on each cluster \(C_i\) of the partition \(P'\).

Let \(D_i, P_i\) be the resulting \(k\)-dominating set and partition of the cluster \(C_i\), for each \(i\).

Let \(D = \bigcup_i D_i\), and let \(P\) be the resulting partition of the entire graph (consisting of the clusters of all partitions \(P_i\)).
Corollary 3.10 \((a)\) \(|D| \leq \frac{n}{k+1}\).

\((b)\) \(\text{Rad}(P) \leq k\).

**Proof:** By Lemma 2.1 and part \((a)\) of Lemma 3.8, \(|D_i| \leq \left\lfloor \frac{|C_i|}{k+1} \right\rfloor\). Thus \(|D| = \sum_i |D_i| \leq \frac{1}{k+1} \sum_i |C_i| = \frac{n}{k+1}\). The radius bound on the partition \(P\) follows from Lemma 2.3 for each \(P_i\) separately. \(\blacksquare\)

**Lemma 3.11** The time complexity of Algorithm \texttt{FastDOM.T} is \(O(k \log^4 n)\).

**Proof:** The complexity of Algorithm \texttt{Diam_DOM} on each cluster \(C_i\) is of the order of the diameter of \(C_i\), which is \(O(k)\) by property \((b)\) of Lemma 3.8. Hence the complexity of Algorithm \texttt{DOM_Partition} \((k)\) (which is \(O(k \log^4 n)\) by Lemma 3.9) dominates the total complexity of Algorithm \texttt{Fast_DOM.T}. \(\blacksquare\)

**Theorem 3.12** There exists a distributed algorithm for computing a \(k\)-dominating set of size at most \(n/(k+1)\) on a tree, with time complexity \(O(k \log^4 n)\).

4 Distributed Computation of a Small \(k\)-Dominating Set on a Graph

In this section we describe how to extend the solution to general graphs. The easy solution of constructing a BFS spanning tree \(T\) of the graph \(G\) and then computing a small \(k\)-dominating set on the tree \(T\) has the disadvantage that constructing the spanning tree requires time \(O(\text{Diam}(G))\). We circumvent this problem by constructing in time \(O(k)\) a spanning forest of the graph, composed of large trees, of size \(k+1\) or more, or formally, a \((k+1, n)\) spanning forest. Then the solution of the previous section can be applied to each tree separately.

4.1 Constructing a \((k+1, n)\) Spanning Forest

To construct a \((k+1, n)\) spanning forest for a given graph \(G\), we use a procedure \texttt{Simple_MST}, which we describe below. This procedure is basically a simplified version of the algorithms of [GHS, A2, G]. The minor modifications necessary for the algorithm of [A2] result from the fact that we need to stop the algorithm after \(\log(k+1)\) phases, but to ensure that these phases do not last more than \(O(k)\) time, while in the [A2] algorithm \(O(n)\) time is allowed for \(O(\log n)\) phases. Rather than describing the somewhat involved algorithm of [A2] in its entirety along with our modifications, we present a version that is significantly simplified,
due to our assumptions that the network is synchronized and that message complexity is ignored. (The number of messages in our algorithm can be reduced, at the expense of simplicity.) Algorithms using similar ideas have appeared in the literature also in different contexts (cf. [JM]).

4.2 A High Level Description

The high level description of the Simple MST procedure we use here is similar to the algorithms of [GHS, A2], and the differences concern mainly the termination rule and some implementation details.

Nodes group themselves into fragments of increasing size. Initially, all nodes are in singleton fragments. Nodes in each fragment \( F \) are connected by edges that form a rooted MST, \( T(F) \), for the fragment. (In the sequel we loosely use the word “fragment” to mean both the collection of nodes \( F \) and the corresponding tree \( T(F) \).)

Active fragments grow through merging onto neighboring fragments. A fragment stops participating in this process and becomes terminated once it detects that its depth has reached \( k + 1 \). Such a fragment will stop trying to merge onto other fragments, although it will still allow other (smaller, active) fragments to merge onto it.

A fragment \( F \) is identified as follows. An active fragment (whose depth is still \( k \) or smaller) is identified by the id of its root node. Each node in the fragment knows this identity, as well as the MST edges that are connected to it. In a terminated fragment \( F \) (whose depth is \( k + 1 \) or larger), on the other hand, identity is less clear. Nodes close to the root (i.e., within distance \( k \) from it) still know its identity. However, more remote nodes know only three things: (a) the MST edges connected to them, (b) the fact that the fragment \( F \) is terminated, and (c) an outdated fragment id (identifying some small fragment \( F' \) they belonged to before \( F' \) was consolidated through merging into \( F \)), whose main useful property is that it is different from the id of any other fragment \( F'' \) that is still active in the system.

Fragment merging is performed as follows. The process is carried out in phases. In the beginning of a phase, within each active fragment \( F \), nodes cooperate to find the minimum weight outgoing edge in the entire fragment. (An outgoing edge of a fragment \( F \) is an edge with one endpoint in \( F \) and another at a node outside it.) The strategy for identifying this edge involves a broadcast (initiated by the fragment root) over the fragment’s tree \( T(F) \)

---

\(^3\text{Note that while this procedure superficially resembles also the partitioning algorithm \texttt{DOM\_Partition} of the previous section, there are a number of major differences due to the different tasks performed by these two algorithms; for example, here we do not need an upper bound on the radius of a fragment.}\)
asking each node separately for its own minimum weight outgoing edge. These edges are then sent upwards on the tree \( T(F) \), towards the root. Each intermediate node first collects this information from all its children in the tree, and then upcasts only the lowest-weight edge it has seen (which is therefore the lowest-weight edge in its subtree). The minimum weight outgoing edge is selected by the root to be included in the final MST.

Once an active fragment’s minimum weight outgoing edge is found, a message is sent out over that edge to the fragment on the other side. The two fragments may then combine, possibly along with several other fragments, into a new, larger fragment \( F' \).

The final step in the phase involves deciding whether the resulting fragment \( F' \) will be active or terminated in the next phase. This is decided by the root of \( F' \), which is chosen in a way described later on. The decision is made by performing a process of “broadcast and echo” to depth \( k + 1 \) over the tree, namely, using a hop counter in the broadcast message, which is decremented with each forwarding of the message. This allows us to check whether the entire tree was covered by the broadcast; if the broadcast reaches a non-leaf node when the hop counter is zero, then the root must be notified that the tree is too deep. This information enables the root to decide whether \( F' \) should be active or terminated. At the end of this process, the root broadcasts again to depth \( k \), informing the nodes of the outcome. Nodes not receiving (any of) these messages (by the appropriate time bound, which is \( k + 1 \) time units after the beginning of this step) learn that they belong to a terminated fragment.

The process as outlined here requires \( O(k \log k) \) time. This is because each phase requires \( O(k) \) time for the various broadcast and convergecast processes performed over the fragment trees. In the next subsection we give a more precise description of Procedure \text{Simple\_MST}, resulting in an improved time complexity of \( O(k) \), again using an idea similar to that of \([AG, G, CT, A2, JM]\).

### 4.3 Procedure \text{Simple\_MST}

The (synchronous) algorithm operates in phases \( 1, 2, \ldots, \log k + 1 \), where phase \( i \) lasts exactly \( 5 \cdot 2^i + 2 \) time units.

In phase \( i \) the root first finds out whether the depth of its tree is larger than \( 2^i \). (This is done using a “broadcast and echo” to depth \( 2^i \) over the tree, just as outlined in the previous subsection.) If the depth is larger than \( 2^i \) then the fragment \text{halts} and does nothing more in this phase, but it does not terminate, i.e., it may resume its active status in later phases. (We see below that some of the fragment’s nodes may receive messages from neighbors not in the fragment and answer them.)
If the root finds that the tree depth is not larger than $2^i$ (hence the fragment should remain active during this phase), then it broadcasts its identity over the tree. (This can be combined with the previous operation.) Exactly after $2^i + 1$ time units after the beginning of the identity broadcast, each node knows whether its fragment is active or not in the current phase. At this point, each node belonging to an active fragment transmits the identity of its root over all of its edges. By the next time unit each node $v$ receives over some of its edges identities of the corresponding roots. Those edges over which it received an identity equal to that of its own fragment are recognized as internal edges, namely, they lead to other nodes in the same fragment. The remaining edges are outgoing edges.

Each node (of an active fragment) now selects its minimum (weight) outgoing edge, and convergcasts its weight to the root. During the convergcast, a message containing an edge weight is discarded once a lower weight edge is known. Thus the root learns the minimum weight outgoing edge of the tree $2^i$ time units after the convergcast started. During the convergcast, each node also remembers the child from which it received each message. Thus the route from the root to the minimum weight outgoing edge can be reconstructed.

Exactly $2^i + 1$ time units after the convergcast started, the root starts a process that transfers the “rootship” to the endpoint of the minimum outgoing edge. Exactly $2^i$ time units later that endpoint is already the root. The new root then notifies the other endpoint of the minimum outgoing edge that it wishes to connect.

If no message from the other endpoint is received by the next time unit then the fragment became a part of the fragment of the other endpoint. Its root thus ceases to be a root. Otherwise, the other endpoint also notified that it wishes to connect. The endpoint with the highest identity becomes the root of the combined fragment.

4.4 Analysis of the Simple_MST Procedure

Since each phase $i$ lasts $O(2^i)$ time, we have

**Lemma 4.1** The Simple_MST procedure terminates after $O(k)$ time.

**Lemma 4.2** When Procedure Simple_MST terminates, the collection of resulting fragments forms a $(k + 1,n)$ spanning forest for the graph. Moreover, each tree of this forest is a fragment of the MST of the graph.

**Proof:** The proof is just a simplified version of the proofs of the previous algorithms, e.g. [GHS, A2]. The lower bound of $k + 1$ on the size of fragments can be proved inductively, arguing that after phase $i$, every fragment is of size at least $2^i$. This is based on the obser-
vations that a fragment at least doubles its size in every iteration in which it is active, and
that if some fragment is halted in phase $i$ then its size was greater than $2^i$ to begin with.
The claim follows since there are $\log(k+1)$ iterations.  

The properties of our simple MST algorithm are summarized in the following lemma.

**Lemma 4.3** Procedure Simple\_MST terminates after $O(k)$ time. Upon termination, the collection of resulting fragments forms a $(k+1,n)$ spanning forest for the graph. Moreover, each tree of this forest is a fragment of the MST of the graph.

### 4.5 Computing a Small $k$-Dominating Set on a Graph

The fast algorithm Fast\_DOM\_G is now obtained as the combination of the partitioning algorithm Simple\_MST of Subsection 4.3 and the $k$-dominating set algorithm Fast\_DOM\_T of Section 3.3. Namely:

**Algorithm Fast\_DOM\_G**

1. Perform Algorithm Simple\_MST, yielding a $(k+1,n)$ spanning forest $F$ for $G$.
2. Perform Algorithm Fast\_DOM\_T on each tree $T_i$ in the spanning forest $F$.

**Theorem 4.4** There exists a distributed algorithm for computing a $k$-dominating set of size at most $n/(k+1)$ on a graph, with time complexity $O(k \log^k n)$.  

### 5 An Application: a Faster MST Algorithm

Our new MST algorithm is composed of two parts. The first part partitions the graph into fragments of the MST, and the second part combines these fragments together. This general structure is similar to that of the MST algorithm of [GKP]. Moreover, the second part of our algorithm is precisely the second part of the MST algorithm of [GKP]. The main change is in offering a new method for accomplishing the partitioning in the first part of the algorithm.

The first part involves simply invoking Algorithm Fast\_DOM\_G of the previous section with parameter $k = \sqrt{n}$. Recall that this algorithm itself is composed of the following three stages: First, execute Procedure Simple\_MST of Subsection 4.3, constructing a $(\sqrt{n}+1,n)$ spanning forest for $G$. Each tree in this forest is a fragment of the MST of the graph.
Next, apply Procedure \texttt{DOM\_Partition}(\sqrt{n}) on each fragment separately, thus yielding a 
\((\sqrt{n} + 1, O(\sqrt{n}))\) spanning forest for \(G\), with each tree in this forest still being a fragment 
of the MST. Finally, apply Procedure \texttt{Diam\_DOM} on each fragment separately\(^4\).

The output of Algorithm \texttt{Fast\_DOM\_G} is a \(\sqrt{n}\)-dominating set \(D\) of size at most \(\sqrt{n}\), and
an accompanying partition \(P\) of the graph into clusters of radius at most \(\sqrt{n}\) around each
of the nodes in \(D\). However, note that each of these clusters is still spanned by a fragment
of the MST, and these fragments were constructed as a byproduct of the algorithm. In
particular, each node belonging to some cluster \(C\) in \(P\) knows which edges belong to the
MST fragment spanning \(C\), and also knows the identity of the fragment’s root.

Thus at the end of the first part, the global state reached by our algorithm is precisely of
the form required for applying the second part of [GKP]. For completeness of the presentation
we now describe the outline of this second part.

Consider the fragment graph \(\tilde{F}\), whose vertex set \(V(\tilde{F})\) is the collection of clusters in \(P\),
containing \(N = O(\sqrt{n})\) fragments of the MST, and whose edge set \(E(\tilde{F})\) is a collection of
(possibly more than \(O(N)\)) inter-fragment edges, which are candidates for joining the MST.

In the second part of the MST algorithm (here, as well as in the algorithm of [GKP]),
a BFS tree is constructed for the graph. Then, each node forwards the description of the
inter-fragment edges it knows of to its parent on the tree. These descriptions are meant to
eventually reach the tree root. However, a node avoids forwarding the description of cycle
heavy edges. That is, a node forwards the description of edges according to their weight, and
does not forward the description of an edge that closes a cycle with edges whose description
has already been forwarded. This seems to suggest that at certain points in time a node
might be forced to wait, due to not having an edge description that is eligible for forwarding.
Thus it seems that the convergecast may not be pipelined, and hence may take a long time.
It is nevertheless proved in [GKP] that this simple convergecast is fully pipelined, and thus
its running time is the one required.

\textbf{Lemma 5.1} [GKP] \textit{When Procedure Pipeline is run on an input composed of a fragment
graph with \(N\) fragments of the MST, its running time is bounded by \(O(N + Diam(G))\), and
its output is an MST for \(G\).}

\section{Algorithm Fast\_MST}

Combining the two parts, we get the following distributed algorithm for MST.

\(^4\)This stage is not essential for the purposes of the current section.
1. Perform Algorithm \texttt{Fast\_DOM\_G} for \( k = \sqrt{n} \).

2. Perform Algorithm Pipeline.

**Theorem 5.2** There exists a distributed Minimum-weight Spanning Tree algorithm with time complexity \( O(\sqrt{n} \log^* n + Diam(G)) \).

**Proof:** By Theorem 4.4 and the choice of \( k = \sqrt{n} \), the execution time of the first stage of Algorithm \texttt{Fast\_MST} is \( O(\sqrt{n} \log^* n) \), and at the end, the graph is decomposed into \( N = \sqrt{n} \) MST fragments. By Lemma 5.1 the time complexity of the second stage of Algorithm \texttt{Fast\_MST} is bounded by \( O(\sqrt{n} + Diam(G)) \). The claim follows.

6 Discussion and Open Problems

There is still a gap of \( \Omega(\log^* n) \) between the upper and lower bounds on the complexity of the \( k \)-Dominating Set construction. It can be interesting to close this gap. In addition, note that the size of the \( k \)-Dominating Set constructed matches the lower bound for a general graph, but nevertheless, a better \( k \)-Dominating Set may exist for any particular given graph. The problem of finding the optimal size set is NP Complete, so it would be interesting to find a distributed algorithm that constructs a good approximation. (As this problem can be formulated as a special case of the set cover problem of [Lo], the greedy sequential algorithm described therein provides an approximation ratio of \( \log n + 1 \). The problem was studied further in the realm of sequential algorithms in [BKP], where its load-balanced case is given an approximate solution.)

As an application of the \( k \)-Dominating Set algorithm we derived a distributed algorithm for constructing an MST, with improved time complexity. This is a classical problem, and it would be interesting to establish a lower bound for its time complexity, and to develop a time-optimal solution (assuming the current solution is not optimal). Neighborhood Optimal (or at least fast) solutions for other problems may speed up other applications, and help fighting congestion. They can also increase our understanding regarding what makes a problem amiable for a neighborhood optimal solution. Finally, it seems very desirable to find characterizations and tools that will make it easier to identify problems for which neighborhood optimal algorithms exist, and to construct them.
References


