On the Distributed Computation of Fractional Connected Dominating Set Packings

Fabian Fuchs, Matthias Wolf
Karlsruhe Institute for Technology
Karlsruhe, Germany
fabian.fuchs@kit.edu, matthias.wolf3@student.kit.edu

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Abstract

One of the most fundamental problems in wireless networks is to achieve high throughput. Fractional Connected Dominating Set (FCDS) Packings can achieve a throughput of \( \Theta(\frac{k}{\log n}) \) messages for networks with node connectivity \( k \), which is optimal regarding routing-based message transmission. FCDS were proposed by Censor-Hillel et al. [SODA’14,PODC’14] and are a natural generalization to Connected Dominating Sets (CDS), allowing each node to participate with a fraction of its weight in multiple FCDS. Thus, \( \Omega(\frac{k}{\Delta}) \) co-existing transmission backbones are established, taking full advantage of the networks connectivity. We propose a modified distributed algorithm that improves upon previous algorithms for \( k \Delta \in o(\min\{\frac{n \log n}{\log n}, D, \sqrt{n \log n \log^* n} \log n\}) \), where \( \Delta \) is the maximum node degree, \( D \) the diameter and \( n \) the number of nodes in the network. We achieve this by explicitly computing connections between tentative dominating sets.

1 Introduction

Wireless ad hoc and sensor networks are used to monitor the environment, industry processes and even large parts of infrastructure. In order to cope with the growing networks size and its demand for efficient communication throughout the network, algorithms and protocols that utilize the capacity available in the network optimally are required. One of the standard methods to manage high throughput in the network is to compute a backbone structure. Recently, Censor-Hillel et al. [1,2] proposed an algorithm that allows to build a network topology based on Fractional Connected Dominating Set (FCDS, see Section 2 for a definition), which can be seen as a generalized Connected Dominating Set (CDS). Such fractional connected dominating sets can be used to achieve a broadcast throughput of \( \Theta(\frac{k}{\log n}) \) messages in networks with \( n \) nodes and vertex-connectivity \( k \), which is optimal regarding
routing-based approaches [2]. This improves on the standard method of using one backbone by, intuitively, replacing it with as many fractional backbones as the network can fit due to its connectivity. To give further intuition, we show an example network that admits multiple FCDS (a so-called FCDS packing) in Fig. 1.

In this work we propose an improved version of the distributed algorithm originally proposed in [1]. Our algorithm is especially beneficial for future large-scale wireless networks, as such networks are expected to consist of a huge number of small wireless nodes deployed on a relatively large area. We present a distributed algorithm that computes a FCDS packing by explicitly computing the connector paths between not-yet connected components of the respective FCDS. Our algorithm runs in the message-passing model $V$-$CONGEST$ and has a round complexity of $O(\log^2 n \min\{\frac{n \log n}{k}, D + \sqrt{n \log n \log^* n} + k\Delta\})$. This improves the runtime of previously $O(\log^3 n \cdot \min\{\frac{n \log n}{k}, D + \sqrt{n \log n \log^* n}\})$ for large and relatively sparse networks with moderate connectivity $k$. More precisely our variant leads to an improvement regarding the runtime if $k\Delta \in o(\min\{\frac{n \log n}{k}, D, \sqrt{n \log n \log^* n}\} \log n)$, which is for example true for logarithmic $k$ and $\Delta$ combined with a diameter such that $k\Delta \in O(D)$ (e.g., in the order of $\sqrt{n}$). An example of a network with network parameters beneficial for our algorithm is described in Appendix A.1. Intuitively the achieved complexity is beneficial for networks with large diameter and moderate density, and generally for large but sparse networks.

Our algorithm is based on the virtual graph structure Censor-Hillel et al. [1, 2] use to compute a FCDS packing. In their distributed implementation they do not explicitly compute the connector paths between the components but rely on the fact that a sufficient number of paths exist, which requires additional coordination within tentatively established dominating sets (so-called components). In [1], the approach of explicitly computing the paths is rendered as probably too expensive. However, our algorithm improves the runtime while explicitly computing possible connector paths between tentative components. Let us now introduce the layered approach used in both their algorithm and ours. For each node in the network we introduce a set of $O(\log n)$ virtual nodes, each virtual node shall be assigned to one FCDS, resulting in a FCDS packing in which each FCDS has at least weight $1/O(\log n)$. The virtual nodes are assembled to a virtual graph $\mathcal{G}$, and partitioned in layers such that there are 1 to 3 copies of a node in each layer. A nodes copy is connected to all copies of the nodes neighbors in $\mathcal{G}$ (cf. Section 2 for details). Now, layer for layer, the virtual nodes are assigned different classes, each of which shall result in a FCDS once the algorithm finishes. Using the first half of the layers, dominating sets are formed, which are then connected by selecting so-called connector paths between components using nodes from the remaining layers.

The improvement in our algorithm is achieved by improving the process of how connector paths are matched to existing components. In [1], paths are matched to components by building a so-called bridging graph. In the bridging graph, whole components (which may span large parts of the network) are simulated by a virtual
node and participate in a matching. Thus, the matching algorithm must communicate through the component, which may require time in the order of $\Omega(D)$ in each step of the matching. In our algorithm we construct a matching graph that can directly be executed by transmitting one message to each neighbor in each round. Overall finding the connector paths requires $O(k\Delta \log n)$ rounds for each one of the $O(\log n)$ layers. Once the connector paths are found, they can be used to connect the components, in order to create multiple CDS is the virtual graph. By translating each CDS to a FCDS in the network, $\Theta(k)$ communication backbones are established in the network, each with weight $1/\log n$.

Related Work: Research on FCDS was started by Censor-Hillel et al. in [2]. They propose a centralized method to compute CDS partitions of size $\Omega(k/\log^5 n)$, as well as FCDS packings of size $\Omega(k/\log n)$, where $k$ is the vertex-connectivity of the network. FCDS packings are the natural generalization of CDS partitions, which allows each node to participate in multiple CDS with a fractional weight between 0 and 1 per CDS such that the sum of the weights is at most 1. Their approach is based on a layered virtual graph, consisting of $\Theta(\log n)$ virtual copies of each node. Each virtual node selects one of $\Theta(k)$ classes, which form the FCDSs later. Using the first $\log n$ layers they achieve domination by assigning random classes to the nodes, the remaining layer are used to connect the existing dominating sets to one connected dominating set per class. Additional to computing FCDS packings, they show that the broadcast throughput using a FCDS packing of size $\Omega(k/\log n)$ is $\Omega(k/\log n)$ messages per round, which is optimal if restricted to routing-based approaches. In contrast to, for example, network coding [4], such approaches consider messages as atomic tokens and use simple store-and-forward methods to route the message. The throughput achievable though routing-based approaches is a logarithmic factor less than the $\Theta(k)$ messages that can be achieved using network coding [2], however, without introducing further challenges (see [4] for example). Ene, Korula and Vakilian [3] consider FCDS packings under the constraint that each node has a capacity. Also using centralized algorithms they compute FCDS packings of size $\Omega(k)$ for planar and minor-closed families of graphs, and $\Omega(k/\log n)$ for the general case. The first distributed implementation is again due to Censor-Hillel, Ghaffari and Kuhn [1]. In this work they consider both vertex- and edge-connectivity. For vertex-connectivity they compute a FCDS packing (or a fractionally disjoint weighted dominating tree, which is a similar concept) of size $\Omega(k/\log n)$, building on the initial approach in [2] as we do in this work.

2 Preliminaries

Our algorithms operate on the communication graph $G = (V, E)$ of a wireless sensor network, where $V$ is the set of nodes or actors in the network, and $E$ the set of edges. An edge $e = (u, v) \in E$ is in the communication graph if $v, u \in V$ can communicate in the network. We assume the communication to be bidirectional, and hence the
communication graph to be undirected. We assume a standard message-passing model, known as \((V-)CONGEST\). Communication is based on synchronous rounds, in which each node can receive the messages of its neighbors as well as transmit one identical message to all neighbors itself. In contrast to the \(E\)-CONGEST model, in which the nodes may transmit different messages to their neighbors, the congestion is on the nodes instead of the edges of the corresponding communication graph. Note that this fits the broadcast nature of wireless networks.

A dominating set \(S \subseteq V\) is a set of nodes in the network such for each node \(v \in V\) it holds that either \(v \in S\) or a neighbor of \(v\) is in \(S\). If such a set is connected we denote it by connected dominating set (CDS). Fractional CDS (FCDS) packings are the natural generalization of CDS. In an FCDS packing, each node can participate in multiple FCDSs with a weight \(x_i \in [0, 1]\) for each FCDS, such that \(\sum_i x_i \leq 1\) for each node. The virtual graph \(G = (V, E)\) used in the construction of the FCDS was first introduced in [2]. For each node \(v \in V\) we introduce \(O(\log n)\) copies in the set of virtual nodes \(\mathcal{V}\). Each copy of \(v\) is connected to all other copies of \(v\) in \(\mathcal{V}\) and to each copy of a neighbor \(w \in V\) of \(v\) in \(G\). We denote the neighbors of \(v\) in \(G\) by \(N_v\), and in \(G\) by \(\mathcal{N}_v\). In contrast to the original description, which was also used for the first distributed implementation in [1], we use \(3L\) copies of each node instead of \(4L\) for \(L \in O(\log n)\), however, this is a minor technical detail. In total the virtual graph has \(3L\) copies of the original graph, plus some additional edges. We subdivide the virtual graph in layers and call the first \(L\) copies of \(V\) in \(\mathcal{V}\) the lower layers. Each so-called upper layer consists of two copies of \(V\). We call the nodes of the first copy type-1 nodes, and the nodes of the second copy type-2 nodes. For each layer \(l\) we denote the nodes of layers 1 to \(l\) by \(\mathcal{V}_l\) and the subgraph induced by these nodes by \(G_l\).

As we compute multiple FCDS simultaneously in the virtual graph, we distinguish each FCDS by a class \(i \leq t \in \Theta(k)\). We denote the subset of nodes of class \(i\) up to layer \(l\) by \(\mathcal{V}_l^i\), and the induced subgraph by \(G_l^i\). We use \(\Psi(v_l) = v\) to project from nodes (or a set of nodes) of the virtual graph to the corresponding real node(s). Throughout the rest of the paper we shall use the term node to refer to virtual nodes in \(G\), and real node to refer to a node of network. During the execution of the algorithm, we aim at connecting not-yet-connected components of the dominating set of a class \(i\) to other components of the same class. Given a connected compo-
nent $C$, we use so-called connector paths to identify vertices that could connect $C$ to another component $C'$ with $\Psi(C) \cap \Psi(C') = \emptyset$, both of the same class. In compliance with [2] we call a path $P$ a connector path for component $C$ if it satisfies the following conditions:

a) $P$ has one endpoint in $C$ and the other endpoint in $C'$
b) $P$ has at most two internal vertices and
c) $P$ cannot be shortened, i.e., for $P = (s, v, w, u)$ with $s \in C$ and $u \in C'$, $u$ does not have a neighbor in $C'$ and $v$ does not have a neighbor in $C$.
d) if $P = (s, v, w, u)$ with $s \in C$ and $u \in C'$ has two internal vertices, $v$ is of type-2 and $w$ of type-1.
e) if $P = (s, v, u)$ with $s \in C$ and $u \in C'$ has one internal vertices, $v$ is of type-1.

Connector paths can have length two or three as the components of each class are already dominating. We call connector paths of length two short and those of length three long. For a path $(v_1, v_2, \ldots, v_{i-1}, v_i)$, we call the set of nodes $\{v_2, \ldots, v_{i-1}\}$ the internal nodes. We call a set of paths $\{P_1, \ldots, P_j\}$ internally vertex-disjoint if the internal nodes of $\{P_1, \ldots, P_j\}$ are mutually disjoint.

The following lemma states that we always find at least $k$ connector paths for each component in a $k$-connected graph.

**Lemma 1** (Lemma 4.3 in [1]). For each component $C$ of an arbitrary class $i$ at an arbitrary level $l$ it holds that $C$ has at least $k$ internally vertex-disjoint connector paths.

Note that the algorithms proof of correctness requires the connector paths of one component to be internally vertex-disjoint. We assume our connector paths to have this property in the following section. It is easy to verify that enough short connector paths are available. For long connector paths, we shall explain how a sufficient number of internally vertex-disjoint long connector paths can be found in Section 4. In the virtual graph $\mathcal{G}$, our algorithm computes a CDS partition, as each node may select exactly one class $i$. Let us now briefly consider how this translates to a FCDS packing in $G$. Let $v \in G$ be a node of the network and $v_1, \ldots, v_{3L}$ the corresponding virtual nodes in $\mathcal{G}$. Given $\Theta(k)$ CDSs in $\mathcal{G}$, we can construct a FCDS in $G$ by weighting the class of each virtual node $v_i$ by $1/3L$ at the real node $v$. As there are $3L$ virtual copies of $v$, the weight constraint is satisfied, and the CDS partition translates to a FCDS packing.

### 3 Distributed FCDS Computation

Our algorithm consists of two main components to construct $t = \Theta(k)$ CDSs in $\mathcal{G}$. Recall that we assign each virtual node to one of $t$ classes, which shall form
the CDSs after the execution of the algorithm. The first $O(\log n)$ layers of virtual nodes establishes that each class dominates the whole graph with high probability (cf. Lemma 2). This is surprisingly simple and can be achieved by having each virtual node select one of the classes at random. For the second $O(\log n)$ layers we aim at connecting a constant fraction of the connected components in each layer (with constant probability). This leads to connectivity of each class with high probability after the last layer, yielding the desired CDSs. The existing distributed algorithm to compute FCDS packings uses the same approach for the lower layers, and (essentially) matches existing components in each of the upper layers without computing the connector paths. Our approach on the other hand explicitly computes the (long) connector paths by constructing a helper graph in which a matching algorithm finds $\Omega(k)$ such paths. Thus we do not require communication through existing components, which is beneficial for many networks, especially if they are large with respect to the diameter. Our algorithm consists of the following steps. Note that the overall design of the algorithm is similar to that of [1], however, we use a different method to connect the components of each class, which is one of the key parts of the algorithm.

A) Each virtual node in the lower layers randomly selects one of the $t$ classes. This leads to domination of each class whp, cf. Lemma 2.

B) For each upper layer $l$ from $L$ to $2L$ we try to connect existing connected components of each class in the nodes of layers $1$ to $l - 1$ using nodes of layer $l$. We call the nodes of the previous layers $1$ to $l - 1$ old nodes and the nodes of layer $l$ new nodes. For each layer we execute steps B.1 to B.4.

B.1) Identify connected components of old nodes. We use the protocol described in [1]. To be self-contained, we describe the protocol in Appendix A.6.

B.2) Let nodes of type-1 select a random class

B.3) For each class $i$: If the nodes component is not yet connected by short connector paths, find $\Omega(k)$ internally vertex-disjoint long connector paths. We construct a helper graph $H_i$ and run a simple matching algorithm to find the long connector paths. For details on this step we refer to Section 4.

B.4) If the type-2 node is on long connector paths, the node discards the paths for which the type-1 node selected a wrong class, and selects the class of one of the remaining paths at random. If no path remains a random class is selected.

After executing this algorithm each virtual node in $G$ is assigned itself to one of the $t$ classes. Each class dominates the whole graph (Step A) and is connected (Step B). Thus, the nodes computed $t = O(k)$ CDSs in the virtual graph $G$. The CDSs can be converted to one FCDS of size $\Omega(k/\log n)$ by assigning each CDS a weight of $1/3L$ (cf. Section 2). Note that the matching algorithm in Step B.2 matches type-1 with type-2 nodes, thus it does not require communication and coordination within large components. Let us now briefly reference the result that achieves dominance in the lower layers.

**Lemma 2** (Lemma 4.1 [1]). For each class $i$, $V_i^t$ is a dominating set in $G$ w.h.p.
The proof idea is based on the fact that, for class $i$ and a node $v \in V$, the probability that $v$'s virtual copy on layer $l$ selects $i$ is at least $1/t = 1/O(k)$. As each node has at least $k$ neighbors on $l$, this yields constant probability per layer, and w.h.p. over all $\log n$ layers.

4 Finding Connector Paths

In this section we show how our algorithm computes internally vertex-disjoint connector paths for each component in order to connect a constant fraction of the components in each upper layer. We begin this section by giving a high-level proof showing that we can indeed connect a constant fraction of the components with each new layer. In the next sections we introduce the necessary tools and prove the remaining results. In Section 4.1 we introduce the graph $H_i$, which helps to reduce the problem of finding long paths for each component to a matching problem. The matching problem is discussed in more detail in Section 4.3.

As introduced in Section 2, connector paths can have one or two internal nodes, we call them short and long connector paths, respectively. To prove correctness for the algorithm, it must hold for each one of the upper layers that at least a constant fraction of the components (formed by old nodes) of each class are connected to another component of the same class using nodes from the current layer with at least constant probability. We shall now state the overall result of this section, which was first obtained and proven in [1]. Note that there is a minor flaw in the original proof regarding the number of missing connections $M_l$ in layer $l$, see [9, p. 21] for details and a corrected proof. Due to space constraints we sketch the proof of the following lemma in Appendix A.2

**Lemma 3** (Lemma 4.4 in [1]). Let $l \in [L, 3L]$. Then $M_{l+1} \leq (1 - \delta)M_l$ with probability at least $\rho$.

4.1 Helper Graph $H_i$

Finding internally vertex-disjoint long connector paths is only relevant if a component has less than $k/2$ internally vertex-disjoint short connector paths. As each component has at least $k$ internally vertex disjoint connector paths, the component must have at least $k/2$ long connector paths in this case. We introduce a helper graph in this section, which is defined such that a maximum matching in this graph corresponds to finding a maximum number of internally vertex-disjoint long connector paths.

For each class $i$ on an upper layer $l$ we define the helper graph $H_i^l$ as the union of the helper graphs $H_i^l[C]$ constructed for each component $C$ of class $i$ on layer $l$. Note that although the helper graphs are constructed for each layer, we omit $l$ in the following as the helper graphs are used only in the layer in which they are constructed. Thus, we always refer to the helper graph of the current level.
We define the helper graph $H_i[C]$ for class $i$ and component $C$. For each type-2 node $v$ of layer $l$ we add a node $v_C$ to $H_i[C]$ iff the following conditions are met:

1) $\Psi(v) \notin \Psi(C)$
2) $v$ has a neighbor in $C$
3) $v$ does not have a neighbor belonging to another component of class $i$

For each node $v_C$ we added to $H_i[C]$, we add for each type-1 neighbor $w$ of $v$ a node $w_C$ to $H_i[C]$, if $w$ has a neighbor in another component $C'$ of class $i$ but no neighbor in component $C$. Intuitively, this procedure ensures that we added the potential long connector path of component $C$ to $H_i[C]$ using a type-2 node as the node closer to $C$ and a type-1 node as the node closer to the neighboring component of the same class. An edge between $v_C$ and $w_C$ is added to $H_i[C]$ as there is an edge between $v$ and $w$ in $G$.

![Figure 2](image.png)

Figure 2: A graph $G$ with three components of a class $i$, along with the helper graph $H_i$ and $H_i[C_1]$ restricted to component $C_1$. Note that $w$ is not in the helper graph as it is on a long connector path.

connection between long connector paths of a component and edges in $H_i$ is shown in the following lemma.

**Lemma 4.** There is an edge $(v_C, w_C)$ in $H_i[C]$ iff there is a long connector path from $C$ to another component of class $i$ through $v$ and $w$ on the current layer.

**Proof.** Let us first assume an edge $(v_C, w_C)$ is added to $H_i[C]$. Then $v$ has at least one neighbor in $C$, which we denote by $s$. Also, $v$ has a neighbor $w$ (of type-1) which does not have a neighbor in component $C$ but has at least one in a component $C' \neq C$ of class $i$. Let us denote this neighbor by $u$. We claim that $P = s, v, w, u$ is a long connector path (cf. Section 2 for the definition), which holds as a) $P$ has one endpoint in $C$, the other in $C' \neq C$ of class $i$, b) $P$ has two internal vertices, c) $P$ cannot be shortened as $v$ does not have a neighbor in a component $C' \neq C$ of class $i$ and $w$ does not have a neighbor in $C$, and d) $v$ is of type-2 while $w$ is of type-1.

Let us now assume we have a long connector path $P = s, v, w, u$. It holds that

1) $w$ does not have a neighbor in $C$, 2) $s$ is in $C$, and 3) $w$ is not in $C$ and $v$ of type-2.

Thus, $v_C$ is added to $H_i[C]$. Also, $w$ is of type-1 and has a neighbor $u$ that is in another component $C' \neq C$ of class $i$ but no neighbor of component $C$, which implies that $w_C$ and the corresponding edge $(v_C, w_C)$ are added as well. 

\[\square\]
The matching algorithm is executed on $H_i$, which is the union of helper graphs for each component, however, observe that we know for each edge in $H_i$ from which component it is induced.

**Observation 5.** Given an arbitrary layer $l \geq L$, a class $i$, and the corresponding helper graph $H_i$. Then each edge in the helper graph can be attributed to exactly one component $C$.

We have shown that the construction ensures that there is a vertex disjoint long connector path through $v$ and $w$ for component $C$ iff there is an edge between $v_C$ and $w_C$ in $H_i[C]$. Thus a matching induces long connector paths. We shall argue in Section 4.3 that we can compute a matching of size $\Omega(k)$ in $H_i$ for each component of class $i$ with $\Omega(k)$ long connector paths. However, let us first describe the distributed algorithm to construct $H_i$.

### 4.2 Distributed Construction $H_i$

Due to Step B.1 of the algorithm, which is executed for each layer before constructing the graph $H_i$, each type-2 node $v$ knows the classes and components of its neighbors. Thus $v$ can decide whether a node $v_C$ should be added for neighboring components $C$. Note that due to Lemma 15 a type-2 node lies only on one long connector path for each class, however, up to $t$ components may have a long connector path through $v$, see Observation 16. If $v$ adds $v_C$ for a component $C$ to $H_i[C]$, it transmits this information along with the class $i$ and the id of $C$ (which is also used in Step B.1) to its type-1 neighbors. These type-1 neighbors can now easily check whether they have a neighbor in $C$ and resign, or verify if at least one neighbor is in another component of class $i$ due to information obtained during Step B.1. If so, $w$ adds $w_C$ and the edge between $w_C$ and $v_C$ to $H_i[C]$.

**Lemma 6.** For each class $i$ on layer $l$ it requires $O(\Delta)$ rounds to construct $H_i$.

**Proof.** First note that each real node $v$ simulates exactly the two virtual copies of $v$ on layer $l$. Due to Conditions 2) and 3) in the definition of $H_i$, the type-2 copy of a node participates in $H_i$ only if its neighbors of class $i$ belong to the same component $C$. In this case, $v$ sends the id of $C$, which requires one message. After receiving these messages, each type-1 node $w$ transmits one message for each message they received from a type-2 node. Note that as $w$ has received at most one message from each neighbor, $w$ responds to at most $\Delta$ messages. Hence, this results in $O(\Delta)$ messages.

The following observation follows from the fact that a type-2 node is only added to $H_i$ if all its neighbors of the class $i$ are in one component, while one type-1 copy is added for each message received by another type-2 node. It helps bounding the runtime of our matching algorithm operating on $H_i$. 


Observation 7. For each node \( v \in V \), there is at most one type-2 copy in \( H_i \), but up to \( \Delta \) type-1 copies in \( H_i \).

4.3 Matching internal vertices

Let us now consider how to distributively compute a matching of cardinality \( \Omega(k) \) in the helper graph \( H_i \) for each component. We shall use this in the next section to prove that each component finds a long connector paths with constant probability in each layer. We use a randomized distributed maximal matching algorithm, which was proposed by Censor-Hillel et al. in [1] and is based on Luby’s distributed maximal independent set algorithm [7]. However, in our case each node in the helper graph \( H_i \) is simulated by only one node at not by several nodes that have to coordinate their actions through components. The algorithm makes use of the special structure of \( H_i \).

Lemma 8. The helper graph \( H_i \) is bipartite.

Proof. As described in the previous section, all nodes in the helper graph \( H_i \) are either added by a type-1 or type-2 node of \( G \). Hence, we may say that the nodes of the helper graph also have types, which induce a partitioning of the nodes of \( H_i \). For two nodes \( v_C \) and \( w_C \) to be connected in \( H_i \), it must hold that \( v \) is of type 2 and \( w \) of type 1. Thus, all edges in \( H_i \) connect nodes of different types, which shows that the graph is bipartite.

Using this lemma the matching algorithm operates as follows. A node is active exactly if none of the adjacent edges is matched, and an edge is active if both adjacent nodes are active. In each round, we assign random numbers from a sufficiently large range to all active edges such that no two edges have the same number whp. Since \( H_i \) is bipartite, assigning the numbers is particularly easy as each type-2 node can pick a number for each incident edge. Each active type-2 node then selects the edge with the largest number and sends its choice to its neighbors. In this round only the selected edges may be added to the matching. At this point, there is at most one edge selected at each type-2 node. However, each type-1 node may have received more than one proposal. To satisfy the matching condition, each type-1 node that has received at least one proposal picks the proposed edge with the largest number and adds it to the matching. The two matched nodes and their edges become inactive. It can be shown that after \( O(\log n) \) rounds all edges are deactivated with high probability and a maximal matching is achieved. Let us now show that such a maximal matching is of cardinality \( \Omega(k) \) if the corresponding component has \( \Omega(k) \) long connector paths.

Lemma 9. Given a component \( C \) of class \( i \) with \( \Omega(k) \) long connector paths. A maximal matching in \( H_i[C] \) is of cardinality at least \( \Omega(k) \).
Proof. If follows from Lemma 1 and the one-to-one correspondence of the long connector paths and the edges in $H_i$ of Lemma 4 that there are $\Omega(k)$ independent edges in $H_i[C]$. Thus, the maximum matching is of size at least $\Omega(k)$, as well as the maximal matching as it is a 2-approximation of the maximum matching.

After showing that the matching is of sufficient size, we prove that this allows us to identify the $\Omega(k)$ long connector paths for each component.

Lemma 10. Consider a component $C$ of class $i$ with $\Omega(k)$ long connector paths. Then a maximal matching in $H_i$ identifies $\Omega(k)$ long connector paths for $C$.

Proof. Let us consider a maximal matching in $H_i$, and component $C$ as required. According to Observation 5, we can consider the subgraph $H_i[C]$ of $H_i$ corresponding to component $C$ as disjoint from other parts of $H_i$. Thus, the matching is maximal also in $H_i[C]$. It holds by Lemma 9 that the size of the maximal matching is $\Omega(k)$. It remains to show that two independent edges in $H_i[C]$ correspond to two internally vertex disjoint connector paths. Consider the edges $(v_C, w_C)$ and $(v'_C, w'_C)$ and assume the corresponding long connector paths with internal vertices $v, w$ and $v', w'$ are not internally vertex disjoint. Thus, either $v = v'$ or $w = w'$ which implies either $v_C = v'_C$ or $w_C = w'_C$. This contradicts the assumption as the edges are not independent. As each matched edge is independent, the set of matched edges in $H_i[C]$ corresponds to a set of internally vertex-disjoint long connector paths of cardinality $\Omega(k)$.

The correspondence between a maximal matching in $H_i$ and long connector paths in $G$ is depicted in Fig. 3. Let us now consider the number of time slots required to compute the maximal matching. The matching algorithm is executed once for every class $i$, and operates on the virtual graph $H_i$. The next lemma proves that $O(\Delta)$ time slots are sufficient for each round of the matching algorithm.

Lemma 11. In each round of the matching on $H_i$ we transmit over each real edge at most twice in each direction, resulting in $O(\Delta)$ time slots for each round.

Proof. Let $v$ be an arbitrary real node, and note that there may be up to $\Delta$ copies of $v$ as type-1 node in $H_i$, but only one copy of $v$ as type-2 node in $H_i$, cf. Observation 7.
Consider any real edge from \( v \) to an arbitrary neighbor \( w \). We may assume that there is at least one copy of the edge \((v, w)\) in the helper graph \( H_i \), as otherwise this edge is not used for the matching algorithm at all. Since the type-2 copy of \( v \) sends only one message over one of its incident edges, it uses the edge \((v, w)\) at most once. After the type-1 copies of \( v \) have received the messages from the type-2 nodes, they respond to one of them. Hence, each type-1 copy sends at most one message.

It remains to show that no two type-1 copies use the same real edge. Assume that there were two type-1 copies of \( v \) that transmit over the real edge \((v, w)\). This would imply that both type-1 copies have received a message from the type-2 copy of \( w \) over the real edge \((w, v)\). However, we have shown above that each real edge is used at most once by the type-2 nodes, which contradicts our assumption. Thus, in one round each real edge transmits at most one message from a type-1 and one from a type-2 node, resulting in two messages per edge. As we operate in the \( V\text{-CONGEST} \) model, two messages per edge results in \( O(\Delta) \) time per round.

It follows from [7] that \( O(\log n) \) rounds are sufficient to compute a maximal matching with high probability.

**Corollary 12.** Our distributed randomized matching algorithm computes a maximal matching in \( H_i \) in \( O(\Delta \log n) \) time.

This implies that we can find \( \Omega(k) \) long connector paths for all components of one class that have less than \( k/2 \) short connector paths in time \( O(\Delta \log n) \). As we have \( t = \Theta(k) \) classes, this results in \( O(k \Delta \log n) \) for Step B.3 on each layer. Let us now prove that the long connector paths can indeed be used to connect the components with at least constant probability on each layer.

### 4.4 From long connector paths to connected components

As components with at least \( k/2 \) short connector paths are connected using those connector paths, we keep focusing on components with at least \( k/2 \) long connector paths. In the previous section we showed how to find \( \Omega(k) \) vertex-disjoint long connector paths for each such component. As type-1 nodes already selected a random class to connect those components that have a sufficient number of short connector paths, the class of the type-2 nodes on the current layer remains to be selected. Since each type-2 node lies on at most one long connector path per class, there are at most \( t \) long connector paths per type-2 node. On these paths, however, the internal type-1 nodes may have chosen classes that differ from the class of the path. Intuitively, this means that the path cannot be used to connect two components of the same class since one of the internal nodes has already picked the wrong class. Therefore, as described in Step B.4, the type-2 nodes discard these long connector paths and select the class of one of the remaining paths at random. If no long connector paths remains, the node selects a random class.
We show in this section that this is sufficient to guarantee that a constant fraction of the components are connected with constant probability. Let us consider an arbitrary component \( C \) of class \( i \). There are two challenges. The first is to show that each connector path connects to another component of the same class with probability in the order of \( 1/k \). This is non-trivial, as the type-1 node on each long connector path already selected a random class, which upper bounds the probability by \( 1/t \). The second challenge is, that the events that two type-2 nodes on different connector paths of \( C \) selecting class \( i \) are not necessarily independent. This can be circumvented by using a tail bound, once the probability for each event is upper and lower bounded independently of the outcome of other events. Let us now state the result. The proof is based on [1] with some modifications. We sketch the main ideas in Appendix A.5.

**Lemma 13.** Given a component \( C \) of class \( i \) on an upper layer \( l \) with \( \Omega(k) \) long connector paths. The probability that one of the long connector paths is good is at least \( \delta \).

5 Conclusion

The algorithm presented in this work computes a fractional connected dominating set packing in the \( CONGEST \) model of distributed computation. It is based on an algorithm by Censor-Hillel, Ghaffari and Kuhn [1, 2], however, our distributed implementation computes the long connector paths explicitly, instead of matching components under the assumption that sufficient long connector paths exist. The runtime of our algorithm is \( O(\log^2 n(\min(\frac{n \log n}{k}, D + \sqrt{n \log n \log^* n}) + k \Delta)) \), which is beneficial for large networks with moderate density, particularly if \( k \Delta \in o(\min(\frac{n \log n}{k}, D, \sqrt{n \log n \log^* n}) \log n) \). We expect future large-scale wireless sensor networks to satisfy such conditions.

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References

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[2] Censor-Hillel, K., Ghaffari, M., Kuhn, F.: A new perspective on vertex connectivity. In: Proc. 25th. Ann. ACM-SIAM Symp. Discrete Algorithms (SODA’14). pp. 546–561 (2014) [1, 2, 3, 4, 5, 13, 16]
A Appendix

A.1 Example network

Let us consider the following network. The nodes of the network are distributed evenly on $d$ rings such that each node on a ring is connected to its neighbors on the ring. Also, $d$ nodes, one from each of the rings, form a clique. For $d = 4$, an example of such a network is displayed in Fig. 4. We can see that such a network is $d + 1$-connected, and has maximum degree $d + 1$. The diameter of the network, however, is in $\Omega(n/d)$. Let us, for example, consider a network with $d = \sqrt[3]{n}$. Such a configuration results in $k\Delta = D = O(n^{2/3})$. Thus, the runtime of the proposed modification yields an improvement over [1].

A.2 Number of Components decreases

In the following we sketch the proof of Lemma 3. This proves that the number of components decreases in each iteration by a constant factor with at least constant probability.

Sketch of proof, based on [1, 9]. To proof the theorem we consider each component and show that the component is connected to another component by layer $l+1$ with constant probability. This implies that a constant fraction of the existing components are connected by layer $l + 1$ with constant probability. Given a component $C$ of class $i$ on layer $l$ and assume class $i$ has at least two components. It holds
that $C$ has at least $k$ connector paths, connecting $C$ to another component of class $i$, according to Lemma 1. There are two cases: Either at least $k/2$ of the paths are short connector paths, or at least $k/2$ of the paths are long connector paths.

Let us first consider the case of at least $k/2$ short connector paths. This is intuitively the easier case, as only one node separates $C$ from another component on $\Omega(k)$ paths. Recall that each layer has two copies of each real node: a type-1 and a type-2 node. Let us consider only the type-1 node for now. According to Lemma 14 it is sufficient that all type-1 nodes select a random class to connect $C$ to a neighboring component with constant probability in this case. Intuitively, this holds as $\Omega(k)$ nodes can connect $C$ with another component, and each of these nodes selects one of $t$ classes.

For the case of less than $k/2$ short connector paths, it holds that there are $\Omega(k)$ long connector paths as in total $k$ connector paths exist according to Lemma 1. We use the remainder of this section to prove that a component $C$ selects a long connector path with constant probability. The theorem follows with the result stated in Lemma 13: At least one of the connector paths selects the required class with at least constant probability.

\[ \Box \]

A.3 Enough short connector paths are sufficient

We state the lemma without a formal proof, which is given as part of the proof of Lemma 4.4 in [1].

Lemma 14 (part of Lemma 4.4 in [1]). Given a class $i$ and a component $C$ of layer $l > L \log n$ with at least $k/2$ short connector paths, $C$ has at least one short connector path of class $I$ with probability at least $\delta$.  

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A.4 Number of connector paths for type-2 node

**Lemma 15** (Proposition 4.2 in [2]). For an arbitrary class $i$ and a type-2 node $v$, $v$ lies on at most one long connector path of one component of $C$.

Which implies the following Observation.

**Observation 16.** Each type-2 node lies on at most $t \in \Theta(k)$ long connector paths.

A.5 Selected long connector paths are good

Let us sketch the proof of Lemma 13 in the following. The analysis is structured in three parts. The first part considers how likely it is that a type-2 node $v$ selects another class, given that the corresponding type-1 node is of the correct class. In order to bound this probability, a random discard step for long connector paths is introduced. This allows to show that all other possible long connector paths through $v$ are discarded with constant probability. This results in an probability in the interval $[1/4t, 1/t]$ for the event that both internal vertices of a long connector path of $C$ select class $i$—independent of the class selected by nodes on other long connector paths of $C$. In the second part it is shown that the proven bounds hold even if the random discard step is not used (this is required, as it is not used in the algorithms). The third and final step uses the independent bounds on the probability of a long connector path to select class $i$ with a tail bound to show that for at least one of the $\Omega(k)$ long connector paths both internal vertices selected the same class $i$.

For more technical details we refer to [1]. All required adaptions are outlined Section 4.2 of [9].

A.6 Identifying Connected Components

To identifying and communicate through connected component, we use the protocol described in [1, Theorem B.2]. There are two protocols that can be used, depending on the maximum diameter $D'$ of the components in the virtual graph maximum diameter $D'$ of the components, which is in $O(n \log n)$ whp. If it is relatively small, i.e. $\frac{n \log n}{k} = o(D + \sqrt{n \log n \log^* n})$, a simple protocol can be used, while a variation of a protocol to identify connector components by Thurimella [8] is used otherwise.

Let us now consider the simpler variant. Each node transmits its class, and the smallest node id it received so far (including its own). Nodes discard received ids if they are transmitted by nodes with different classes. After $D' = O\left(\frac{n \log n}{k}\right)$ rounds, each node in each component received the smallest id of the component, which is selected as the component id and the components root node. The union of paths from the root to nodes of the components can be used as communication tree in the component.
The more complex protocol, which is a variation of the algorithm to identify connected components by Thurimella [8] is originally based on an minimum spanning tree (MST) algorithm by Garay, Kutten and Peleg [5], which was improved to the current runtime bound by a new MST algorithm in [6]. The protocol allows each node in a network to learn the smallest id in its component in \( O(D + \sqrt{n} \log^* n) \) rounds. The id of each virtual node \( v_l \) (of layer \( l \)) is set to \((id_v, l, \text{type})\), where \( id_v \) is the id of the corresponding real node, \( l \) the virtual nodes layer, and type its type (either 1 or 2). The algorithm by Thurimella is executed on \( G \), which has a diameter in \( O(D) \), and \( O(n \log n) \) nodes, resulting in \( O(D + \sqrt{n} \log n \log^* n) \) rounds for identifying the connected components.