Fast Beeping Protocols for Deterministic MIS and $(\Delta + 1)$-Coloring in Sparse Graphs
(Extended Version)

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Abstract—The beeping model is an extremely restrictive broadcast communication model that relies only on carrier sensing. We consider two problems in this model: $(\Delta+1)$-vertex coloring and maximal independent set (MIS), for a network of unknown size $n$ and unknown maximum degree $\Delta$. Solving these problems allows to overcome communication interferences, and to break symmetry, a core component of many distributed protocols. The presented results apply to general graphs, but are efficient in graphs with low edge density (sparse graphs), such as bounded degree graphs, planar graphs and graphs of bounded arboricity. We present $O(\Delta^2 \log n + \Delta^3)$ time deterministic uniform MIS and coloring protocols, which are asymptotically time optimal for bounded degree graphs. Furthermore, we devise $O(a^2 \Delta^2 \log^2 n + a^2 \Delta n \log n)$ time MIS and coloring protocols, as well as $O(a^2 \Delta^2 \log^2 n + a^2 \Delta n \log n)$ time 2-hop MIS and 2-hop coloring protocols, where $a$ is the arboricity of the communication graph. Building upon the 2-hop coloring protocols, we show how the strong CONGEST model can be simulated and by using this simulation we obtain an $O(a)$-coloring protocol. No results about coloring with less than $\Delta + 1$ colors were known up to now in the beeping model.

I. INTRODUCTION

The discrete beeping model was introduced by Cornejo and Kuhn [1] to provide a convenient formal framework for studying radio networks having severe restrictions on communication capabilities, yet where widely applicable protocols can be designed and analytically proven in an efficient manner. Since protocol executions in distributed computing are frequently hard to grasp (even with simulations or experiments), having formal models is crucial for both practical and theoretical reasons. In the discrete beeping model, time is divided into synchronous rounds, and in each round, a node can either listen or transmit a unary signal (beep) to all its neighbors. The possibility to directly transmit a beep to a node is defined by a static communication graph, and nodes have absolutely no knowledge of this graph. A beeping node receives no feedback, while a silent one can only detect that either at least one of its neighbors beeped or that all of them were silent. A listening node does not receive the identifiers of its beeping neighbors, as a beep is merely a detectable burst of energy. Protocols can use the synchronous nature of the rounds to transmit information through beeps, but doing so impacts the time complexity in a quantifiable manner. This work studies how this difficulty can be overcome.

Applications of this model range from radio networks with reduced network stacks [1], such as energy-limited sensor networks, which can provide improved speed, low cost and less transmission errors, to biological networks [2], where the beeping model allows to study the efficiency of natural protocols. Indeed, most biological systems communicate in a primitive manner. Fireflies communicate through flashes of light [3] and cells through the diffusion of specific chemical markers [4].

Different applications will result in different communication graphs. Graphs with low edge density are said to be sparse. The maximum degree and the arboricity of a graph are measures of its edge density, where low values indicate sparse graphs. Contrarily to graphs with low maximum degree, low arboricity graphs can be seen as graphs which are “globally” sparse but may be “locally” dense. Many real-world networks are sparse [5]. In particular, graphs embedded in some surface, for example the plane, have low arboricity.

The distributed vertex coloring and maximal independent set (MIS) problems are fundamental building blocks in protocol design. The coloring problem consists in assigning colors to nodes such that no two neighboring nodes (sharing an edge in the communication graph) have the same color. The MIS problem consists in choosing a set of nodes in the communication graph such that no two nodes in the set are neighbors, and such that any node not in the set has a neighbor in that set. Solving these problems is important for dealing with the interferences inherent to the beeping model. More generally, a coloring can be used to allocate resources that cannot be shared by neighboring nodes. Nodes in an MIS can act as cluster heads in order to coordinate actions, and participate in a network backbone construction.

Serving as important primitives for protocol design in the beeping model, MIS and coloring problems have received a lot of attention (see Sect. I-B). Efficient probabilistic solutions were proposed for general graphs. However, the more difficult deterministic case, useful whenever random behavior is inappropriate or deterministic guarantees are required, has received much less attention. In this work, we are interested in designing deterministic protocols having efficient time complexity.
A. Preliminaries

Let \([k]\) be the set \([1, \ldots, k]\). For any two integers \(a, b (\in \mathbb{Z})\) and any positive integer \(k (\in \mathbb{N}^+)\), let \(a \equiv b \mod k\) denote the congruence relationship between \(a\) and \(b\) such that \(a \mod k = b \mod k\). The operator \(\equiv\) is used for the string concatenation. For any positive integer \(k\), \(l(k)\) is the length of the binary representation of \(k\), i.e., \(l(k) = 1 + \lfloor \log_2 k \rfloor\). For any function \(f : \mathbb{N}^+ \rightarrow \mathbb{N}^+\) and any positive integer \(k\), \(f_i(k)\), where \(i \in [l(k)]\), denotes the \(i\)th most significant bit of \(f(k)\)’s binary representation.

The communication network is represented by a simple connected undirected graph \(G = (V, E)\), where \(V\) is the node set and \(E\) the edge set. The network size \(|V|\) is also denoted by \(n\), the diameter by \(D\) and the maximum degree by \(\Delta\). For a node \(v \in V\), the neighbors of \(v\) are \(N(v) = \{u \in V \text{ s.t. } (u, v) \in E\}\) and its degree is \(\deg(v) = |N(v)|\). Nodes have unique identifiers (ids). This property is essential in order to break symmetry in deterministic protocols. The identifier of a node \(u \in V\), \(id(u)\), is an integer from \([N]\) where \(N = n^c\) with a constant \(c > 1\). \(N\) is an upper bound on the total number of nodes in \(G\). The length of \(id(u)\) is denoted by \(l(u)\). Then, the maximum length over all ids in \(G\) is \(l_{\text{max}} = \max_{a \in V} l(a)\). We have \(l_{\text{max}} = O(\log N) = O(\log n)\).

The distance between two nodes \(u\) and \(v\) in \(G\) is \(d_{\text{ist}}(u, v)\).

The square graph of \(G\) is the graph \(G^2 = (V, E_s)\), where \(E_s = \{(u, v) \mid u, v \in V, d_{\text{ist}}(u, v) \leq 2\}\). \(G[R]\) denotes the subgraph of \(G\) induced by \(R \subseteq V\). Its edges \((E_G[R])\) are the edges of \(G\) connecting two vertices in \(R\). The arboricity of \(G\), denoted by \(a(G)\) or just \(a\), is the minimum number of disjoint forests into which the edge set \(E\) can be partitioned. Equivalently, Nash-Williams [6] proved that arboricity is also a measure of density, i.e., \(a = \max_{R \subseteq V, |R| \geq 2} \frac{|E_G[R]|}{|R| - 1}\).

B. Related Work

In [7], round complexity lower bounds are given for the MIS and \((\Delta + 1)\)-coloring problems. These bounds are \(\Omega(\log n)\) and \(\Omega(\Delta + \log n)\) respectively. They were obtained assuming randomized algorithms, and thus apply to both deterministic and randomized algorithms. In the latter case, the solution or the running time is guaranteed with high probability (w.h.p.). More specifically, these bounds apply to a stronger variant of the beeping model (with collision detection). In this variant, listening nodes can distinguish between a single beep and the superposition of multiple beeps (a collision).

In [1], the authors present the first coloring protocol for the beeping model. It outputs a correct coloring after \(O(\Delta + \log n)\) rounds w.h.p. Following this paper, randomized MIS and coloring protocols were designed for the beeping model with collision detection, in a series of papers ([4], [8], [9]). These protocols achieve optimal round complexity, but only with collision detection. Moreover, the resulting colorings often employ more than \(\Delta + 1\) colors. These protocols can be translated to the weaker beeping model (with no collision detection) with an \(\Omega(\log n)\) multiplicative factor.

Schneider and Wattenhofer [10] solve deterministic MIS in radio networks with collision detection. Although the term “beeping model” does not appear in [10], the presented protocol straightforwardly works in this model. It is time optimal for growth-bounded graphs (GBG). These are graphs where, for any given node \(v\) and integer \(r\), the number of nodes in any independent set (see definition in Sect. 1-D) within distance \(r\) of \(v\) is bounded by \(f(r)\) (polynomial in \(r\)). However, this property does not cover trees, planar graphs, or more generally, sparse graphs.

The round complexities of different MIS and coloring protocols are compared below (see respectively Figure 1 and 2).

![Fig. 1: MIS protocols](image)

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<td>anonymous nodes</td>
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<td>(O(\log n))</td>
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![Fig. 2: Coloring protocols](image)

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<tr>
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<td>(\Delta + \log n) colors</td>
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<td>Here</td>
<td>(O(\Delta^2 \log n + \Delta^3)), deterministic</td>
<td>(\Delta + 1) colors</td>
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C. Protocol-related Definitions

In the beeping model, an execution proceeds in synchronous rounds (there are synchronized local clocks and all nodes start at the same time: \emph{synchronous start}). In each round, nodes synchronously execute the following steps:

1. \textbf{Send:} Each node beeps (instruction \emph{BEEP} in protocols) or listens (\emph{LISTEN} in protocols). Beeps are transmitted to all neighbors of the beeping node.
2. \textbf{Receive:} If a node beeped in the previous step, then it learns no information from its neighbors. Otherwise, it knows whether or not at least one of its neighbors beeped during the previous step of the same round.
3. \textbf{Process:} Each node performs local computations.

One of the most common message passing models is the CONGEST model of edge bandwidth \(B\) [11]. It is stronger than the beeping model, as nodes communicate by sending messages of maximum length \(B\) (commonly \(O(\log n)\)) in a round. Different messages can be sent to different neighbors and nodes receive the full content of all incoming messages.

We adopt the classical definitions. The state of a node is the vector of the values of its variables. A variable \emph{var} of a node \(v\) is explicitly associated to \(v\) using a subscript \emph{var}_v. A configuration is a vector of the states of all nodes. An execution proceeds in rounds and is defined by the sequence
of the configurations at the end of each round, starting from an initial configuration. If the same configuration (resp. state of a node) is repeated indefinitely at the end of each round, we say that this configuration (resp. the state) is terminal. When such a configuration is reached, it is said that the system/protocol has terminated, or that termination has occurred. A problem is given as a first order predicate over configurations. A protocol is said to solve a problem if each execution terminates, and each terminal configuration satisfies the predicate of the problem specification. The round complexity (time complexity) of a protocol is the number of rounds needed to reach a terminal configuration in the worst case. A protocol is said to be uniform in a parameter $p$ if it does not depend on the value of $p$. It is said to be locally termination detecting, or simply locally terminating, if for any given node $v$, $v$ detects if it has reached a terminal state.

In the beeping model, protocols must specify what is done in each round. Due to the nature of the communication model, each action is performed on a sequence of consecutive rounds. For instance, a node may have to wait for a round of silence, or beep only every $k$ rounds. At the code level, this type of action is expressed by a loop. As it will appear later, in some complex protocols, such loops are nested. For the sake of clarity, we will name the sequence of rounds in the innermost loop the $L_1$-phase, the sequence of loops in the loop just above, the $L_2$-phase, and so on.

We extend previous definitions concerning protocols to $L_i$-phases, in particular uniformity and termination. We consider terminal $L_i$-phase states (states that no longer change in this $L_i$-phase), locally terminating $L_i$-phases (any given node $v$ detects when it has reached a terminal $L_i$-phase state) and uniform $L_i$-phases (when the range of the loop index is unknown). Not being able to detect when a uniform $L_i$-phase has ended for all nodes, raises the question of synchronizing the start of the following $L_{i+1}$-phase.

We solve this problem by using $L_i$-synchronization points, represented by $\frac{\tau}{i}$ in the code. Upon reaching an $L_i$-synchronization point (after having reached a terminal $L_i$-phase state), any given node $v$ waits for all of its neighbors to reach a terminal $L_i$-phase state before executing the following $L_{i+1}$-phase, if there is any. $L_i$-synchronization points require locally terminating $L_i$-phases, so that any given node $v$ can detect when all of its neighbors have reached the synchronization point. The method for detecting that was first introduced in Förster et al. [12], with the "Balanced Execution Technique" (BET). However, BET only guarantees $L_i$-synchronization points. In appendix A, we extend BET to guarantee $L_i$-synchronization points for any $i \geq 1$. The extension, referred to as EBET, is crucial in the design of complex uniform protocols in the beeping model.

We call a protocol a competition protocol when nodes are "eliminated" round after round until the "surviving" nodes form an independent set (possibly empty). In this paper, we only consider competition protocols where the elimination process is deterministic and depends on identifier comparison.

D. Problem Specifications

The predicates (over configurations) defining the problems considered in the paper can be naturally obtained from the definitions given below.

A set $I \subseteq V$ of vertices is said to be an independent set if for any $u, v \in I$, $u$ and $v$ are not neighbors in $G$. An independent set $I$ is maximal (MIS) if any vertex in $V \setminus I$ has a neighbor in $I$. A 2-hop MIS of $G$ is an MIS of its square graph $G^2$.

A set $J \subseteq V$ of vertices is said to be a $(t, s)$-ruling set [13], if for any two vertices $u, v \in J$, $\text{dist}(u, v) \geq t$, and for any vertex $v \in V \setminus J$, there exists a vertex $u \in J$ such that $\text{dist}(u, v) \leq s$.

With this definition, an MIS is a $(2, 1)$-ruling set. A forest is said to be a $(t, s)$-ruling forest if the roots are a $(t, s)$-ruling set and the trees are of depth at most $s$.

A $c$-coloring $\text{col}$ is a function from $V$ into a set of colors $[c]$ such that $\forall (u, v) \in E \text{\ col}(u) \neq \text{\ col}(v)$. Notice that in a $c$-coloring, nodes with the same color constitute an independent set. It is thus possible to construct an MIS from them. A 2-hop coloring of $G$ is a coloring of its square graph $G^2$.

Any given function $\text{color}$ is a $d$-defective $c$-coloring [14] if $\forall v \in V$, $\text{color}(v) \in [c]$ and $v$ has at most $d$ neighbors colored with $\text{color}(v)$. We say that $\text{color}$ has a defect of $d$. An edge where both endpoints have different colors is said to be a non defective edge, otherwise it is said to be a defective edge. With this definition, a (proper) coloring is a 0-defective coloring.

E. Contributions

The first contribution of this paper is a tool for analyzing competition protocols: the labeled “deterministic” competition graphs (Sect. II). Using such graphs in protocol analysis is inspired by [15]. Here we adapt this technique to the case of deterministic competition protocols. This general tool may be useful also in future studies of MIS and coloring.

The second and main contribution of the paper is a series of uniform MIS and coloring protocols. First, we present $O(\Delta^2 \log n + \Delta^3)$ deterministic uniform protocols for MIS and $(\Delta + 1)$-coloring, where $\Delta$ is the unknown maximum degree (Sect. III). These protocols are time optimal for bounded degree graphs. They also scale well to graphs with polylogarithmic $\Delta$. Indeed, in these graphs, the time complexity is polylogarithmic in regards to $n$, which is very efficient.

Then, we extend the previous protocols with time complexity dependent on $\Delta$, to protocols with time complexity dependent on the arboricity $a$ (Sect. IV). We get $O(a^2 \Delta^2 \log^2 n + a^3 \Delta^3 \log n)$ time MIS and $(\Delta + 1)$-coloring uniform protocols. This results in efficient polylogarithmic time complexity for the large family of graphs where $a = O(\log^c n)$. Finally, we extend the previous protocols into $O(a^2 \Delta^2 \log^2 n + a^3 \Delta^3 \log n)$ time 2-hop coloring and 2-hop MIS uniform protocols (Sect. V). Given a 2-hop coloring, we prove that the CONGEST model can be simulated with some multiplicative overhead (Sect. VI). Using this simulation and the algorithm proposed in [16] for the CONGEST model, we get an $O((a^2 \Delta^2 + a^3 \Delta^3) \cdot \log^2 n + a^3 \Delta^3 \log n) \cdot a(a)$-coloring protocol in the beeping model, for any given positive constant $\mu < 1$. To the best of our
knowledge, this is the first coloring protocol using less than \( \Delta + 1 \) colors in the beeping model.

II. RULING SET PROTOCOL AND COMPETITION GRAPHS

Ruling sets serve as building blocks to construct complex protocols. They have been used to compute MIS [13] and colorings [17], [18]. In these papers, the ruling sets are used to decompose the network, and nodes in the ruling set (the “local leaders”) take care of solving the problem for the nodes within a certain distance. In the beeping model, doing so is more difficult. We show in the next section, how ruling sets can still be used to design an efficient coloring protocol.

In this section, we introduce a competition protocol (RulingSet - Protocol 1 in Sect. II-B) computing a \( (2,2 \log N) \)-ruling set. This protocol can be considered as a variant of the ruling set algorithm from [13]. That algorithm is heavily recursive, requiring concurrent communications, which are incompatible with the beeping model. Therefore, we adapt it and provide a non-recursive competition protocol with a similar behavior. To prove correctness (Sect. II-C), we use competition graphs, which are directed graphs that serve to model the behavior of competition protocols and help analyzing them. They were first used in [15], but in association with a non-deterministic elimination process. As we are interested in deterministic protocols, we use the nodes’ identifiers to label the edges of a competition graph with \( \alpha \)-encodings (Sect. II-A), and these values determine a deterministic elimination process. The resulting labeled competition graphs allow to compute the surviving nodes in a convenient way.

The encoding starts with as many 1’s as the length of the binary representation, followed by a 0.

**Fig. 3: Description of \( \alpha \)-encoding**

### A. \( \alpha \)-encoding

We \( \alpha \)-encode integers to design uniform competition protocols, as in Casteigts et al. [19]. This encoding allows to compare integers (identifiers) bit by bit in a uniform manner. Indeed, when using \( \alpha \)-encodings (of integers from \([N]\)), such protocols do not need to know the binary integers’ lengths (depending on \( \log N \)) to compare them bit by bit. The \( \alpha \)-encoding of integer \( i \) is made up of two parts, as explained in Figure 3. Comparing two \( \alpha \)-encodings \( \alpha(i) \) and \( \alpha(j) \) consists of first comparing the minimum number of bits necessary to encode the integers, and if it is the same, comparing the binary representations of \( i \) and \( j \) (the first bit is unnecessary, because it is 1 and the length is already known).

**Definition 1.** Let \( i \) be a positive integer, \( bin \) its binary representation and \( l(i) \) the length of \( bin \). Let \( bin_{-1} \) be the binary representation without the leading 1 and \( 1^{l(i)} \) a binary string of \( l(i) \) 1’s. The \( \alpha \)-encoding of \( i \), denoted by \( \alpha(i) \), is \( 1^{l(i)} \| 0 \| bin_{-1} \) where \( \| \) denotes string concatenation.

\( \alpha \)-encoding preserves the order between two integers.

**Lemma 1.** For any \( i, j \in \mathbb{N}^>0 \): \( i < j \Leftrightarrow \alpha(i) < \alpha(j) \), where \( <| \) is the lexicographical order on \( \alpha \)-encodings.

### B. Uniform Competition Protocol for Computing a Ruling Set

Nodes use their unique identifiers for comparison and survivors of the elimination process constitute the output set. Each node \( v \) has a unique identifier \( id(v) \). The identifiers are encoded on at most \( l_{max} \) bits, but \( l_{max} \) is unknown to the nodes and thus the binary representations of the identifiers do not necessarily have the same length. Every node \( v \) computes the \( \alpha \)-encoding \( \alpha(id(v)) \) (or \( \alpha(v) \) for short, by notation abuse) and outputs a boolean value \( survived_v \). We prove that the output is a \( (2,2 \log N) \)-ruling set (Theorem 2).

#### Protocol 1 RulingSet

1: \( IN: id, \text{Integer} \) \( \quad OUT: survived, \text{Boolean} \)
2: \( sur\text{vived} := \text{true} \) \( \quad \alpha := \alpha(id) \) \( \triangleright \) Get \( \alpha \)-encoding
3: \( \text{for round } r := 1 \text{ : } r \leq \ell(\alpha) : r++ \) \( \triangleright r \) is incremented after each iteration
4: \( \text{if } v_r \text{ is } 1 \text{ then BEEP} \) \( \triangleright \) Consider the \( r^{th} \) most significant bit
5: \( \text{else} \)
6: \( \text{LISTEN} \)
7: \( \text{if beep heard then} \) \( \triangleright \) If a neighbor has a higher identifier
8: \( survived := \text{false} \)
9: \( \text{EndProtocol} \)
10: \( \text{EndProtocol} \) \( \triangleright \) No beep heard

The following lemma is straightforward.

**Lemma 2.** RulingSet has a round complexity of \( max_{v \in G}(\ell(\alpha(v))) = O(\log N) \).

#### C. Correctness Analysis of Protocol 1

The output set of RulingSet is analyzed through a game, which we refer to as the “elimination game”. This game is enacted on an edge-labeled directed acyclic graph \( G_{dag} \), the labeled competition graph, constructed from the original communication graph \( G \) and the nodes’ unique identifiers. This construction process is adapted here to the RulingSet protocol, but it applies to any competition protocol. \( G_{dag} = (V,E_{dag},\text{label}) \), where \( E_{dag} \) is the set of directed edges and \( \text{label} \) an edge labeling function. \( G_{dag} \) is constructed from the \( \alpha \)-encodings of the identifiers, encoded on a maximum of \( 2l_{max} \) bits.

- Let \( (u,v) \) be an edge of \( G \) with \( \alpha(u) \triangleright \alpha(v) \). Then, \( (u,v) \) is a directed edge in \( G_{dag} \), directed from \( u \) to \( v \).
- Let \( (u,v) \) be an edge of \( G_{dag} \). For the smallest index \( i \in [2l_{max}] \) such that \( \alpha_i(u) = 1 \) and \( \alpha_i(v) = 0 \), set \( \text{label}(u,v) = i \): the edge \( (u,v) \) is labeled (with) \( i \).

For any edge \( e = (u,v) \) of \( G_{dag} \), \( u \) is called the origin and \( v \) the extremity of \( e \). The following lemma is straightforward.

The elimination game is played by the nodes of \( G_{dag} \), round by round, in the following way: on round \( r \), all surviving nodes
We define the same notions for outgoing edges.

Theorem 1. There is a unique label sequence \( \{\text{lab}_1, \ldots, \text{lab}_r\} \) of an elimination game played on \( G \). The game finishes when no more node can be eliminated (thus after at most \( 2l_{\max} \) rounds). A node’s survival is stored as a boolean in the \( \text{survived} \) variable.

Definition 2. Let \( v \) be a vertex in \( G_{\text{dag}} \). Let \( e \) be an incoming edge. We say that \( e \) is acting if the origin of \( e \) is not eliminated before round \( \text{label}(e) \), and non acting otherwise. If \( e = (u, v) \) is an acting incoming edge, then \( u \) eliminates \( v \) at round \( \text{label}(e) \) if and only if \( v \) has not already been eliminated.

Definition 3. Let \( \Pi = (v_1, \ldots, v_l) \) be a directed path in \( G_{\text{dag}} \). There is a unique label sequence \( S_{\text{lab}}(\Pi) = (s_1, \ldots, s_{l-1}) \) s.t. \( \forall r \in [l-1], s_r = (v_r, v_{r+1}) \) and \( s_r = \text{label}(e_r) \).

Results similar to the following lemma and theorem are proven in [20] for a more limited case. Lemmas 3 and 4 are straightforward.

Lemma 3. Let \( \Pi = (v_1, \ldots, v_l) \) be a directed path in \( G_{\text{dag}} \). \( S_{\text{lab}}(\Pi) \) has no consecutive equal labels: \( \forall r \in [l-1] s_r \neq s_{r+1} \).

Theorem 1. Let \( v \) be any node from \( G_{\text{dag}} \) not surviving the elimination game. There exists a surviving node \( u \) such that \( \text{dist}(u, v) \leq 2l_{\max} \), where \( l_{\max} = O(\log N) \).

Proof. First, a path \( \Pi \) from a surviving node \( u \) to node \( v \) is constructed, then we prove that \( \Pi \)’s length is at most \( 2l_{\max} \). \( \Pi \) is constructed by induction. Node \( v \) did not survive, so there exists an acting incoming edge. The acting incoming edge \( (w, v) \) with the smallest label is added to \( \Pi \). If \( w \) does not survive the elimination game, the previous actions are repeated and an acting incoming edge is added to \( \Pi \). This is done until a surviving node is reached. Since at least one node survives the elimination game, \( \Pi \)’s construction is well-defined and \( \Pi = (e_1, \ldots, e_l) \).

Now, let us prove by contradiction that \( l \leq 2l_{\max} \). Suppose \( l > 2l_{\max} \) and focus on \( S_{\text{lab}}(\Pi) \). Because the edge-labels are integers from \( [2l_{\max}] \) and consecutive labels are non equal by Lemma 3, there exists an extremum \( s_i \) indexed by \( r \in \{2, \ldots, 2l_{\max}\} \). Thus there exists \( i \in \{r-1, r\} \) such that \( s_i > s_{i+1} \). However, both \( e_i \) and \( e_{i+1} \) are acting incoming edges, by construction. Thus, the origin of \( e_i \) is eliminated in round \( s_{i+1} \), which contradicts the fact that \( e_i \) is acting. Hence, we have a contradiction.

Lemma 4. Let \( I = \{v \in V \text{ s.t. } \text{survived}_v = \text{true}\} \) at the termination of \( \text{RulingSet} \). Let \( S \) be the set of survivor nodes of an elimination game played on \( G_{\text{dag}} \). We have \( I = S \).
The defect of \( v \) at the start (and end) of an upper bounded by \( \Delta + 1 \) termination detection and make nodes start the next step of the terminating. Therefore we can use BET to perform neighboring terminating (if executed alone). However, they are locally ruling set (for function ColorReduction(color, new color (the smallest available color in \( [\Delta + 1] \)), given by a function \( U \) node's color (function \( U \) stores unavailable colors). Nodes broadcast colors from \( [\Delta + 1] \) rounds, after which, nodes with color = \( c \) change their color to the smallest available color in \( [\Delta + 1] \). The output is the node's new color (color), given by a \( d \)-defective (c - 1)-coloring).

Now, let us present the ColorReduction function invoked in the color reduction \( L_2 \)-phase. Its input parameters are an integer value given by a \( d \)-defective \( c \)-coloring (color), the maximum degree \( \Delta \) and the maximum color \( c \) (in the coloring), where \( c > \Delta + 1 \). Nodes broadcast colors from \( [\Delta + 1] \) in \( \Delta + 1 \) rounds, after which, nodes with color = \( c \) change their color to the smallest available color in \( [\Delta + 1] \). The output is the node's new color (color), given by a \( d \)-defective (c - 1)-coloring).

In DegreeColoring, we only require \( L_1 \)-synchronization points (for RulingSet and ColorByBFS), introduced in Sect. I-C. Both functions are uniform in \( N \), and thus are not explicitly terminating (if executed alone). However, they are locally terminating. Therefore we can use BET to perform neighboring termination detection and make nodes start the next step of the protocol synchronously. On the other hand, as the time lengths of all \( L_2 \), \( L_3 \) and \( L_4 \)-phases (and ColorReduction calls) are upper bounded by \( \Delta + 1 \), their termination is completely synchronized at all nodes and we do not need \( L_2 \), \( L_3 \) and \( L_4 \)-synchronization points.

Lemma 5. At the start (and end) of an \( L_3 \)-phase, color \( \in [\Delta + 1] \).

Lemma 6. The defect of color is reduced by one per \( L_3 \)-phase.

Proof. Let color be \( d \)-defective at the start of \( L_3 \)-phase \( \rho_3 \). For any given node \( v \), \( v \) has at most \( d \) defective edges. It is easy to see that non-defective edges remain non-defective. In a non-defective edge \( (v, u) \), let \( v \) be the node with the higher color w.l.o.g. During \( L_3 \)-phase \( p_3 \), \( v \) stores a set of unavailable newColor values, including newColor. As such, when \( v \) executes ColorByBFS, newColor \( v \neq \) newColor. All endpoints of the defective edges of \( v \), and \( v \) itself, execute RulingSet and ColorByBFS in the same \( L_2 \)-phase. If DistC(\( v \)) denotes \( v \)'s distance to the nearest BFS tree root (RulingSet survivor), there is at least one endpoint \( u \) with \( |\text{DistC}(u) - \text{DistC}(v)| = 1 \). Because of the difference in the parity of these distances, \( u \) and \( v \) choose different values in \( [\Delta + 2] \), and at least one edge becomes non-defective. \( \square \)

Theorem 3. DegreeColoring solves \((\Delta + 1)\)-coloring in \( O(\Delta^2 \log n + \Delta^3) \) rounds.

Given a \( (\Delta + 1) \)-coloring, it is simple to compute an MIS in \( \Delta + 1 \) rounds. Nodes with the same color form an independent set. Adding iteratively (at each round) nodes from each such set to a common independent set results in an MIS. Thus, MIS can also be solved in \( O(\Delta^2 \log n + \Delta^3) \) rounds.

B. Uniform \((\Delta + 1)\)-coloring

Now, we wish to transform DegreeColoring into UnifDegreeColoring, which is uniform in both \( \Delta \) and \( n \). The first step is to replace the functions used in DegreeColoring by uniform functions, and to synchronize them using synchronization points. Then, every non-uniform stopping condition of a loop appearing in DegreeColoring should be eliminated and replaced by a so called local termination component. This component is an \( L_{i-2} \)-phase executed at the end of each iteration (\( L_{i-1} \)-phase) of the loop (\( L_i \)-phase). It serves to detect if the executing node has finished the ongoing loop. More formally, this component serves to detect whether the executing node has reached a terminal \( L_i \)-phase state, and makes the \( L_i \)-phase locally-terminating.

First, let us present UnifBroadcastColors, a uniform version of BroadcastColors (since BroadcastColors requires \( \Delta \)). UnifBroadcastColors is an \( L_3 \)-phase, made of consecutive \( L_1 \)-phases, each composed of 2 rounds. In the
first round, the executing node \(v\) beeps if it has not yet communicated newColor\(_v\). Otherwise, it listens so it can detect if all of its neighbors have already communicated their newColor value, and if so, \(v\) terminates. In the second round, we have the round behavior of BroadcastColors. In such a way, we obtain a uniform function having the same behavior as BroadcastColors. Moreover, in this particular case, since all \(L_1\)-phases contain exactly 2 rounds, it is also locally synchronized, even without using EBET, and therefore there is no need to indicate synchronization points explicitly.

```
function UniBroadcastColors(newColor, changingColor, U): U
1: for \(L_1\)-phase \(p1 := 1; p1++\) do \(\triangleright \) \(L_1\)-phase consists of two rounds
2: \(\triangleright \) First round
3: if newColor >\(= p1\) and changingColor then BEEP \(\triangleright \) Not finished yet
4: else
5: \(\triangleright \) Listen
6: if no beep heard then Return U \(\triangleright \) If all neighbors beeped their colors
7: \(\triangleright \) Second round
8: if newColor = \(p1\) and changingColor then
9: \(\triangleright \) BEEP \(\triangleright \) Communicate your color
10: else
11: \(\triangleright \) Listen
12: if beep heard then U := \(U \cup \{p1\}\) \(\triangleright \) Keep neighbors’ newColor values
```

Next, we design a uniform version of ColorReduction. It is used in ReduceColors, a uniform version of the color reduction \(L_2\)-phase from DegreeColoring. UnifColorReduction has two input parameters: the node’s color (color), given by a \(d\)-defective \(c\)-coloring, and a set of unavailable colors (U). It also has two output parameters: the node’s new color color, given by a \(d\)-defective \(c’\)-coloring (with \(c’ = min(c - 1, D + 1)\)), and a boolean sameColor indicating whether color changed. Every node \(v\) transmits its color value to its neighbors by beeping in the first round of the \(L_1\)-phase indexed by color. Nodes with the highest color in their neighborhood choose the smallest available color (colors previously transmitted by neighbors are forbidden). If that color is the node’s current color, then sameColor is assigned to true. Other nodes do not change their color (and end with sameColor equal to false). Here again, there is no need to indicate synchronization points explicitly, since all \(L_1\)-phases contain exactly 2 rounds.

ReduceColors is an \(L_4\)-phase. It has two input parameters: the node’s color (color), given by a \(d\)-defective \(c\)-coloring, and a set of unavailable colors (U). It has a single output parameter: the node’s new color (color), given by a \(d\)-defective (\(D + 1\))-coloring. The main idea is to have the nodes with the highest color in their neighborhood change their color to the smallest available color (in [\(D + 1\)]). At some point, they can no longer improve their color (finished is true). These nodes terminate, allowing the other nodes in their neighborhood to change their color value. Here, it is crucial to put \(L_2\)-synchronization points after the UnifColorReduction and UniBroadcastColors calls, because these functions are uniform. Thus, different nodes can finish executing these functions at different times, i.e., not synchronously. As these functions are locally terminating, EBET can be used to ensure the synchronization points explicitly.

Following this, let us describe the functions used for UnifDegreeColoring’s local termination component. These functions are used to detect when the executing node’s color is proper, i.e., no neighbor has the same color. Then, the executing node can exit the outermost loop and thus locally terminate the protocol (see lines 24 to 29).

```
function UnifColorReduction(color, U); color, sameColor
1: sameColor := false
2: for \(L_1\)-phase \(p1 := 1; p1++\) do \(\triangleright \) \(L_1\)-phase consists of two rounds
3: \(\triangleright \) First round
4: if color = \(p1\) then BEEP
5: else
6: LISTEN
7: if beep heard then U := \(U \cup \{p1\}\)
8: \(\triangleright \) Second round
9: if color > \(p1\) then BEEP
10: else
11: \(\triangleright \) Only a node with the highest color in its neighborhood hears no beep
12: LISTEN
13: if beep heard then Return (color, sameColor)
14: else
15: color := min([\(p1\)] \(\backslash\) U)
16: if color = \(p1\) then sameColor := true
17: Return (color, sameColor)
```

```
function ReduceColors(color, U); color
1: finished := false
2: while not finished do \(\triangleright \) At most \(c\) \(L_3\)-phases
3: \(\triangleright \) (color, finished) := UnifColorReduction(color, U)
4: \(\triangleright \)
5: U := UnifBroadcastColors(color, finished, U)
6: \(\triangleright \)
7: Return color
```

```
function UniCollisionBeep(pid: collision)
1: collision := false
2: for \(L_1\)-phase \(p1 := 1; p1++\) do \(\triangleright \) \(L_1\)-phase consists of two rounds
3: \(\triangleright \)
4: if \(p1 > \ell(a(id))\) then Return collision
5: \(\triangleright \)
6: BEEP ; LISTEN
7: if beep heard in the second round then collision := true
8: \(\triangleright \)
9: if beep heard in the first round then collision := true
```
Finally we describe UnifDegreeColoring. The main idea is the same as in DegreeColoring: we refine the initial \( \Delta \)-defective coloring until the coloring is proper. The main differences are the local termination components. The \( \mathcal{L}_1 \)-phase’s (\( L_3 \) loop) local termination component is similar to the local termination component in UnifBroadcastColors.

A node has finished an \( \mathcal{L}_4 \)-phase if all of its neighbors have chosen a new color. The protocol’s local termination component is described previously. The additional \( U \) variable is used to store unavailable colors that have already been chosen by neighboring nodes which have terminated the protocol.

**Theorem 4.** MIS and \((\Delta + 1)\)-coloring can be solved with \(O(a^2 \log^2 n + a^3 \log n)\) round complexity in the beeping model, where \( a \) is the arboricity of the communication graph.

To support this theorem, we design two coloring protocols with the above round complexity: one is uniform in \( N \) but not in \( a \), and the other is uniform in \( a \) but not in \( N \). It is important to have a protocol uniform in \( a \), since \( a \) may be harder to obtain than an upper bound on \( N \). The following results from [24] are used to obtain these protocols.

**Lemma 7.** [24] If \( G \) is of arboricity \( a \), at least \( \frac{|V|}{2a} \) nodes have a degree less than \((2 + \epsilon)a\).

**Theorem 6.** [24] If \( G \) is of arboricity \( a \), it can be decomposed into \( l = O(\log n) \) sets of nodes \( H_1, \ldots, H_l \) such that each set \( H_i \) has maximum degree \( O(a) \) in the induced subgraph \( G[A_i] \).

The LimitedDegreeColoring function is the main component of both protocols. It colors all participating low-degree nodes, if it is given an upper bound on the arboricity \( a \). A node \( v \) is considered to be a low-degree node if it has deg \((v) \leq \Delta_a \), where \( \Delta_a = (2 + \epsilon) \cdot a \) for a parameter \( \epsilon > 0 \). Contrary to DegreeColoring, it may happen that some nodes have no available colors in \([\Delta_a + 1]\), due to their high degree, and end the function uncolored, represented by the color 0. We use LimitedColorReduction, a slightly modified version of ColorReduction. It is not presented here, but the only change is that \( color \) is set to 0 if \([\Delta_a + 1] \} \setminus U \) is an empty set.

**Theorem 5.** MIS and \((\Delta + 1)\)-coloring can be solved with \(O(a^2 \log^2 n + a^3 \log n)\) round complexity in the beeping model, where \( a \) is the arboricity of the communication graph.

IV. IMPROVEMENTS FOR GRAPHS WITH SMALL ARBORICITY

DegreeColoring is efficient for graphs with polylogarithmic maximum degree \( \Delta \). However, not all graphs have a low maximum degree, and in these graphs, Protocol 3 is less efficient. Using ideas from [23] and [24], it is possible to design a \((\Delta + 1)\)-coloring protocol which is efficient on graphs with low arboricity \( a \) (more specifically, with polylogarithmic \( a \)). Notice that some important topologies like trees and planar graphs have an arboricity of 1 and 3 respectively, while their maximum degree can be arbitrarily large.

**Lemma 8.** Let \( \Delta_a = (2 + \epsilon) \cdot a \), with \( \epsilon > 0 \). Given the input \( c = \Delta_a \), LimitedDegreeColoring outputs a \((\Delta_a + 1)\)-coloring on a subgraph of nodes, which includes all nodes with degree less than or equal to \( \Delta_a \). All other nodes have output 0. The round complexity is \(O(a^2 \cdot \log n + a^3)\).

**Proof.** The round complexity is straightforward. LimitedDegreeColoring outputs a \((\Delta_a + 1)\)-coloring on the subgraph of nodes with non-zero colors because all colors are chosen from \([\Delta_a + 2]\), if available. And are then reduced to \([\Delta_a + 1]\). ColorCollision ensures that the coloring is valid.
Now, let us prove by contradiction that for any given node \( u \) with \( \deg(u) \leq \Delta_a \), the output is a non-zero color. \( u \) outputs 0 due LimitedColorReduction, ColorByBFS or ColorCollision. The first two cases are impossible because \( |U(u)| \leq \Delta_a \). In the last case, ColorCollision is executed after \( \Delta_a \) \( \mathcal{L}_2 \)-phases. In each \( \mathcal{L}_4 \)-phase, incident non-defective edges remain non-defective, and at least one incident defective edge becomes non-defective. Since after \( \Delta_a \) \( \mathcal{L}_3 \)-phases \( u \) has no defective edges, \( u \) has no neighbor \( v \) with \( \text{color}_u = \text{color}_v \). \( \square \)

### A. \((\Delta+1)\)-coloring Uniform in \( N \)

First, let us focus on the first protocol, uniform in \( N \). LimitedDegreeColoring is executed iteratively by uncolored nodes until all nodes are properly colored. Since \( a \) is known and by Lemma 7, each invocation of the function colors a constant fraction of the nodes of the communication graph. Colored nodes no longer participate in subsequent LimitedDegreeColoring calls. By Theorem 6, executing LimitedDegreeColoring \( l = O(\log N) \) times (or more) colors all nodes with \( O(a \cdot \log N) \) colors. As \( N \) is unknown, invocations of LimitedDegreeColoring continue until the executing node is colored properly (local termination component). When this happens for all nodes, the \( O(a \cdot \log N) \)-coloring is transformed into a \((\Delta+1)\)-coloring by ReduceColors, as in Protocol 3. This takes an additional \( O(a^2 \cdot \log^2 N) \) rounds.

#### Protocol 4 UnifNArbColoring

1. **IN:** id: Identifier, \( N \): Polynomial upper bound on \( n \), \( e \): Parameter
2. **OUT:** color: Integer value
3. \( \Delta_a := (2 + e) \cdot a \)
4. for \( \mathcal{L}_4 \)-phase \( p4 := 1 : p4++ \) do
   5. \( \text{color} := \text{LimitedDegreeColoring}(\text{id}, \Delta_a) \)
   6. \( \mathcal{L}_4 \) loop
   7. if \( \text{color} \neq 0 \) then
      8. \( \text{color} := \text{color} + (p4 - 1) \cdot (\Delta_a + 1) \)
5. \( \mathcal{L}_4 \) loop
10. \( \text{color} := \text{ReduceColors}(\text{color}, 0) \)
11. \( \text{color} \) is an \( O(a \cdot \log N) \)-coloring
12. EndProtocol

#### Theorem 7. Protocol 4 solves MIS and \((\Delta+1)\)-coloring with \( O(a^2 \log^2 N + a^2 \log n) \) round complexity. This protocol is uniform in \( N \) but non-uniform in \( a \).

**Proof.** Let us prove that after all \( \mathcal{L}_4 \)-phases, arbColor is an \( O(a \cdot \log n) \)-coloring. In each \( \mathcal{L}_4 \)-phase of Protocol 4, only uncolored nodes \( (V_{rem}) \) participate in LimitedDegreeColoring. Since the subgraph induced by \( V_{rem} \) also has arboricity at most \( a \), by Lemmas 7 and 8, \( \frac{2}{\log n} |V_{rem}| \) nodes have a degree less than \( \Delta_a \) and thus are part of the subgraph with a \((\Delta_a + 1)\)-coloring. They exit the \( \mathcal{L}_4 \) loop, thus by Theorem 6, there are at most \( \frac{2}{\log n} = O(\log n) \) \( \mathcal{L}_4 \)-phases. Since we use non-overlapping \( \Delta_a + 1 \) colors for each \( \mathcal{L}_4 \)-phase, arbColor is an \( O(a \cdot \log n) \)-coloring. The round complexity follows from the number of \( \mathcal{L}_4 \)-phases and Lemma 8. \( \square \)

### B. \((\Delta+1)\)-coloring Uniform in \( a \)

In the second protocol (uniform in \( a \)), we compute an upper bound on \( a \). This is done by estimating \( a \) iteratively. At each iteration (\( \mathcal{L}_5 \)-phase) \( i \), \( a \) is estimated to be \( 2^i \) and LimitedDegreeColoring is executed \( l = O(\log N) \) times, given this estimation. After \( O(\log a) \) iterations, the estimation is at least as large as the actual arboricity. When this happens, LimitedDegreeColoring executed \( O(\log N) \) times provides a proper coloring (followed by the color range reduction) as in the first protocol.

#### Theorem 8. Protocol 5 solves MIS and \((\Delta+1)\)-coloring with \( O(a^2 \log^2 N + a^3 \log n) \) round complexity. This protocol is uniform in arboricity \( a \) but non-uniform in \( N \).

**Proof.** Let us first prove that Protocol 5 solves \((\Delta+1)\)-coloring. At the end of \( \mathcal{L}_5 \)-phase \( p5 \), by Lemma 8, all nodes with degree less than \( \Delta_p5 = (2 + e) \cdot 2^p5 \) are colored. By Lemmas 7 and 8, and Theorem 6, at \( \mathcal{L}_5 \)-phase \( p5 = 1 + \lceil \log a \rceil \), \( \Delta_p5 \geq (2 + e) \cdot a \) and all nodes are colored after \( \frac{2}{\log N} \cdot \log N \) \( \mathcal{L}_1 \)-phase. Since the color ranges from different \( \mathcal{L}_1 \)-phases or different \( \mathcal{L}_5 \)-phases do not overlap, and the non-zero colors returned by LimitedDegreeColoring form a coloring, arbColor is an \( O(a \cdot \log N) \)-coloring. And after the ReduceColors call, the coloring is reduced to a \((\Delta+1)\)-coloring.

It is straightforward to prove the round complexity. Since arbColor is an \( O(a \cdot \log N) \)-coloring, ReduceColors takes at most \( O(a^2 \log^2 N) \) rounds, while the \( \mathcal{L}_5 \) loop takes at most \( O(a^2 \log^2 N + a^3 \log N) \) rounds. \( \square \)

### V. Uniform Protocols for 2-hop MIS and 2-hop \((\Delta^2+1)\)-coloring

To obtain protocols for 2-hop MIS and 2-hop coloring, we provide and use a general transformer, the SquareSim protocol (Protocol 6), allowing to "simulate \( G^2 \) over \( G \)." The idea is that nodes propagate beeps for an extra round (and therefore contact nodes at distance 2), so that they can simulate a protocol on the square of the communication graph, for a small time multiplicative overhead. SquareSim provides two primitives SquareSim(true) and SquareSim(false) to simulate in \( G \), the BEEP and LISTEN instructions invoked on graph \( G^2 \).

#### Lemma 9. A protocol designed to be executed on \( G^2 \) can be simulated on \( G \) by replacing all BEEP instructions by calls to SquareSim(true) and LISTEN instructions by calls to SquareSim(false).

The maximum degree of the square communication graph is \( \Delta^2 \). By applying Lemma 9 to the previous protocols, we obtain
protocols for solving 2-hop coloring with $(\Delta^2 + 1)$ colors and 2-hop MIS. These protocols are very efficient on bounded degree graphs, and efficient for graphs with polylogarithmic $\Delta$. 2-hop coloring is an important tool in the beeping model, used to break symmetry and to deal with the interferences. In the next section, we show how this can be used to simulate the stronger CONGEST communication model and obtain an $O(a)$-coloring.

**Corollary 1.** 2-hop MIS and 2-hop $(\Delta^2 + 1)$-coloring can be solved in $O(\Delta^4 \log n + \Delta^6)$ rounds.

Instead of the maximum degree of the square of the given graph, consider its arboricity. Using a result from [25], showing that $a(G^2) \leq 2^a \cdot a \cdot \Delta$, we obtain Corollary 2, which provides a more efficient result for graphs with small arboricity.

**Corollary 2.** 2-hop MIS and 2-hop $(\Delta^2 + 1)$-coloring are solved by the two protocols in Sect. IV with an $O(\mu^2 \Delta^3 \log n + a^3 \Delta^3 \log n)$ round complexity. One of them is uniform in $N$ but not in $a$, and the other is uniform in a but not in $N$.

**VI. CONGEST MODEL SIMULATION AND $O(a)$-COLORING**

By using a 2-hop coloring, nodes can simulate the transmission of messages through the edges of the communication graph, like in the CONGEST model with edge bandwidth $B$ (commonly $O(\log N)$). We want to make sure that for any given node $v$, a message can be sent or received along any edge without interference, and that the provenance and destination of the message can be deduced easily.

First, **InitCongest** (Protocol 7) is used at the beginning of the simulation to obtain all possible message provenance and destinations for any given node $v$ (simulated from the 2-hop coloring). After which, the transmission of messages is done through SimCongest.

Our simulation algorithm SimCongest (Protocol 8) is made of two components. The **first component** is used to transmit a $B$ bit message. If we have no interference, a node can transmit $B$ bits during $2B$ rounds (in phases of two rounds, one round for transmitting bit 1 and another one for bit 0).

The **second component**, and the core part of the simulation, deals with the interferences inherent to the beeping model. Here, a 2-hop $c$-coloring (for some constant $c$) is required so that messages can be associated to a pair of colors $p = (\text{colorProvenance}, \text{colorDestination})$, according to their provenance and destination ($c^2$ possibilities). The simulation is composed of phases, each of $c^2$ invocations of the first component. In this way, transmitted bits never collide. The $B$ bit messages are part of the input parameters of SimCongest. They are given through a hash table ($mSend$), with the message destinations (colors) as keys and the messages as values. The messages received are stored in a similar structure ($mRec$), where the message provenances are the keys.

**Protocol 6 Simulating the square communication graph: SquareSim**

1: **IN:** beep: Boolean value  
   **OUT:** detectedBeep: Boolean value  
2: detectedBeep := false  
3: **if** beep then BEEP  
   **then** Transmit beep to neighbor nodes: First round  
4: else  
5:   **LISTEN**  
6:   **if** beep heard then detectedBeep := true  
7:   **if** detectedBeep then BEEP  
   **then** Transmit to distance 2 nodes: Second round  
8: else  
9:   **LISTEN**  
10: **if** beep heard and not beep then detectedBeep := true  
11: **EndProtocol**

The following lemma is straightforward.

**Lemma 10.** Given a 2-hop $c$-coloring, the CONGEST model with edge bandwidth $B$ can be simulated in the beeping model, with an $O(c^2 \cdot B)$ multiplicative factor.

Finally, using the simulation of CONGEST, one can use the result of Barenboim and Elkin [16] (given for CONGEST), to obtain an $O(a)$-coloring in the beeping model. It is done by first computing, in the beeping model, a 2-hop $(\Delta^2 + 1)$-coloring in $O(a^2 \Delta^2 \log^2 n + a^3 \Delta^3 \log n)$ rounds (Corollary 2). Then the $O(a)$-coloring from [16] (with $O(a^6 \log n)$ round complexity) is combined with the CONGEST simulation, using the $(\Delta^2 + 1)$-coloring obtained before. By Lemma 10, the resulting simulation of the $O(a)$-coloring protocol has $O(a^6 \Delta^4 \log^2 n)$ round complexity.

The final result is an $O((a^2 \Delta^2 + a^6 \Delta^4) \cdot \log^2 n + a^3 \Delta^3 \log n)$ time $O(a)$-coloring protocol in the beeping model. Notice that now by using this coloring algorithm, together with the SquareSim protocol, to obtain a 2-hop $O(a \cdot \Delta)$-coloring (see Sect. V), we reduce the time multiplicative factor when simulating CONGEST algorithms. Consequently, one obtains a more efficient simulation.
REFERENCES


APPENDIX A

EBET

We remind that the “Balanced Execution Technique” (BET) from Förster et al. [12] guarantees $L_1$-synchronization points. Now, we present an extension of BET, with which we guarantee $L_i$-synchronization points for all $i \geq 1$. The “Extended Balanced Execution Technique” (EBET) allows the design of complex uniform protocols in the beeping model.

A. Introducing EBET

Synchronization points are not a natural primitive in the beeping model: an $L_1$-synchronization point forces nodes which have reached a terminal $L_1$-phase state (ended the $L_1$-phase) to wait for their neighboring nodes to end the $L_1$-phase, before starting the next one. Some protocols are difficult to design in a uniform manner without the use of synchronization points. Therefore, we want to be able to design a protocol $P$ using synchronization points, and then apply a “technique” on the formal description of $P$, so that the result is a protocol that can be run in the beeping model (not necessarily a formal description). The resulting protocol is called $P_{sim}$. The technique we use for that is EBET.

EBET has two crucial components and a parameter $k \in \mathbb{N}>0$, which controls the small multiplicative overhead of EBET. The first component is a Finite State Machine (FSM), used to stall nodes when they have ended an $L_i$-phase (stall property), for all $i \leq k$, so that other nodes can catch up. The second is a balanced round counter $rC$, which is used so that nodes can reach some agreement on the clock value for the current $L_1$-phase. By balanced counter, we mean that the $rC$ values of two neighbors differ by at most 1 (balancing property). Thus two neighbors participating in the same $L_1$-phase are in the same round, or in consecutive rounds. EBET’s main addition is an extension to the FSM component. As a consequence, EBET provides $L_i$-synchronization points, for all $i \leq k$. For better clarity, we consider EBET with $k = 2$, but it is simple to extend the following techniques for any given positive integer $k$.

We assume that in $P$ and $P_{sim}$, all nodes start synchronously. By using synchronization points, $P$ is simple, easily described, coded and understood. Here we consider $P$ to be a uniform loop of $L_2$-phases (thus a uniform $L_3$-phase). Whereas $P_{sim}$ is a uniform loop of $L_1$-phases, and each of its $L_1$-phase simulates a round of $P$. Since the $L_1$-phases of $P_{sim}$ contain exactly 8 or 11 rounds (referred to as slots to differentiate from the rounds in $P$), $P_{sim}$ can be run in the beeping model. We refer to phases of $P$ as original phases, and to phases of $P_{sim}$ as simulation phases. It is crucial that $P_{sim}$ outputs the same result as $P$, and proving this is the main focus of Sect. A-C.

In the first section, we describe the balanced counter technique (extending that of [12]), which allows EBET to maintain a balanced round counter, and to guarantee the stall property for all $L_1$-phases. When abstracted to a higher level, the stall property is the essence of synchronization points. In the second section, we describe how communication is adapted for EBET. Indeed, nodes do not have perfectly synchronized round counters, so we adapt the manner in which nodes communicate between themselves (having a balanced counter is crucial here).

B. Extending the Balanced Counter Technique for EBET

1) Slot Behavior in the Balanced Counter Technique: The balanced counter technique is implemented in the following
manner. Nodes have the following variables: state, rC, p1 and p2. These variables are parametrized by a node v and if unclear, by a simulation \( L_1 \)-phase \( p \), to indicate their value for \( v \) at the start of a simulation \( L_1 \)-phase \( p \). The state variable can be any of the 5 states from Figure 5 \( (CT, R-N, R-W, R-N2 \) and \( R-W2) \). A node \( v \) is said to be participating (in phase \( p \)) if \( state(v, p) = CT \). The \( R-N \) and \( R-W \) states are used to stall nodes starting the next original \( L_1 \)-phase. In the same way, the \( R-N2 \) and \( R-W2 \) states are used to stall nodes starting the next original \( L_2 \)-phase.

Fig. 5: Finite State Machine Component for EBET \((k = 2)\)

Each simulation \( L_1 \)-phase contains exactly 8 slots and is used to transmit a node’s local clock value and its FSM state. Using this information, nodes know if they are ahead or behind of their neighbors, and act accordingly. The first three slots (indexed 0 to 2) are used to transmit the counter value \( (rC) \) modulo 3, and the other slots (indexed 3 to 7) are used to transmit the current FSM state of a node \( (state) \). For any given node \( v \), the information is transmitted in the following manner during each simulation \( L_1 \)-phase:

- If \( state = CT \), then \( v \) beeps in slots \( (rC \mod 3) \) and 3,
- If \( state = R-N \), then \( v \) beeps in slot 4,
- If \( state = R-W \), then \( v \) beeps in slots \( rC \mod 3 \) and 5,
- If \( state = R-N2 \), then \( v \) beeps in slot 6,
- If \( state = R-W2 \), then \( v \) beeps in slots \( rC \mod 3 \) and 7.

Node \( v \) listens in all slots it does not beep in.

Now, we describe the state transitions of the FSM, and their constraints (also shown in Figure 5). These constraints are essential to the balanced counter and \( stall \) properties in EBET. For any given node \( v \), the allowed state transitions are:

1) \( CT \rightarrow R-N \) if no node \( u \in N(v) \) is in \( R-W \),
2) \( R-N \rightarrow R-W \) if no node \( u \in N(v) \) is in \( CT \),
3) \( R-W \rightarrow CT \) if no node \( u \in N(v) \) is in \( R-N \),
4) \( CT \rightarrow R-N2 \) if no node \( u \in N(v) \) is in \( R-W2 \),
5) \( R-N2 \rightarrow R-W2 \) if no node \( u \in N(v) \) is in \( CT2 \),
6) \( R-W2 \rightarrow CT \) if no node \( u \in N(v) \) is in \( R-N2 \).

\( CT2 = \{ CT, R-N, R-W \} \) is a composite state. A node \( v \) with \( state(v) = CT2 \) is simulating an original \( L_2 \)-phase iteration. If its state is \( R-N2 \) or \( R-W2 \), it has “reached” an \( L_2 \)-synchronization point and is waiting for its neighbors.

We define a boolean \( next(v, p) \) for any given simulation \( L_1 \)-phase \( p \) and node \( v \). The boolean is true if and only if all neighboring nodes of \( v \) have equal or greater \( rC \) values. \( v \) learns its \( next \) value after the first three slots of \( p \), since the boolean is true if and only if \( v \) detects no beeps in slot \( rC(v) - 1 \mod 3 \). If \( next(v, p) \) is true, then \( rC(v) \) is incremented at the end of phase \( p \).

2) Functions of the Balanced Counter Technique: Now we describe the functions used by the balanced counter technique. The \( increment \), \( reset \) and \( reset2 \) functions increment \( rC \), \( p1 \) and \( p2 \) such that \( rC \) remains a balanced counter, and \( p1 \) and \( p2 \) are the same for any two participating neighboring nodes. For any given node \( v \), these functions can only be invoked if \( v \) is participating.

- \( increment \) is used to increment \( rC \) without violating the balancing property. \( v \) invokes \( increment \) in the very first phase of \( P_sim \), and calls \( increment \) again whenever the previous call finishes. During these calls, \( v \) simulates \( P \) since \( state(v) = CT \) and the rules described previously are independent of \( increment \). When \( increment \) is invoked by a node \( v \), \( v \) waits for the first simulation \( L_1 \)-phase \( p \) in which no beep is received in slot \( s = rC(v,p) - 1 \mod 3 \). At the end of this phase, \( v \) increments \( rC \).

- \( reset \) is used to start the simulation of the next original \( L_1 \)-phase, without interfering with the other nodes’ simulations of the current original \( L_1 \)-phase. When invoked by \( v \), \( v \) goes through the transitions 1, 2 and 3 as soon as they are available (and communicates according to its state). \( rC(v) \) is reset to 0 after transition 1 succeeds and \( p1(v) \) is incremented after transition 3 succeeds. \( reset2 \) is similar but is used to start the simulation of the next original \( L_2 \)-phase, without interfering with the other nodes’ simulations of the current original \( L_2 \)-phase (which can mean waiting for multiple original \( L_1 \)-phases to be simulated by neighboring nodes). When invoked by \( v \), \( v \) goes through the transitions 4, 5 and 6 as soon as they are available (and communicates according to its state). \( rC(v) \) and \( p1(v) \) are reset to 0 after transition 1 succeeds and \( p2(v) \) is incremented after transition 3 succeeds.

\( reset \) (resp. \( reset2 \)) is invoked by a participating node \( v \) in the round after \( v \) has detected it has ended the current \( L_1 \)-phase (resp. \( L_2 \)-phase). The details are in the next section (Sect. A-C).

3) Properties of the Balanced Counter Technique: First, we prove the \( stall \) property (extending Lemma 21 from [12]), after which we prove that \( rC \) is a balanced counter.

Lemma 11 (Stall property). For any given simulation \( L_1 \)-phase \( p \) and node \( v \), if \( state(v,p) = R-W \) or \( state(v,p) = R-W2 \) then for all \( u \in N(v) \) either \( rC(u,p) = 0 \) or \( rC(u,p) = 1 \).

Proof. For any given node \( w \) and simulation \( L_1 \)-phase \( p \), if \( state(w,p') \neq CT \) then \( rC(w,p') = 0 \). Thus, w.l.o.g., consider \( state(v,p) = CT \). Then \( state(v,x) = R-N \) or \( state(v,x) = R-N2 \) in some simulation \( L_1 \)-phase \( x < p \). Let \( q \) be the largest such \( x \). Without loss of generality, consider \( state(v,q) = R-N \) (the same line of arguments holds for the other case). The state transitions for \( v \) after \( q \) are \( R-N \rightarrow R-W \rightarrow CT \) and when \( v \) arrives in \( CT \) in
some simulation $L_1$-phase $t \geq q$, $rC(v,t) = 0$. This implies that $u$ arrives in $R$-$W$ before simulation $L_1$-phase $t$. Finally, in all simulation $L_1$-phases $y$, where $t \leq y \leq p$, node $v$ receives a beep in slot 0 (sent by $u$) and $rC(v,y) \leq 1$.

**Lemma 12 (Balancing property).** For any given simulation $L_1$-phase $p$ and two neighboring participating nodes $u$ and $v$, 
\[ |rC(v,p) - rC(u,p)| \leq 1. \]

**Proof.** Let us prove that $rC$ satisfies the balancing property by induction on $p$. Node $v$ is participating in simulation $L_1$-phase $p$ if $\text{state}(v,p) = CT$. For $p = 0$, the balancing property is given by the initialization conditions.

For the induction step, consider a simulation $L_1$-phase $p > 0$ and two neighboring nodes $u$ and $v$. If either $u$ or $v$ is not participating, the induction hypothesis holds. Otherwise, there are two cases. In the first case, $u$ and $v$ were participating in simulation $L_1$-phase $p - 1$. Then by the induction hypothesis, 
\[ |rC(u,p - 1) - rC(v,p - 1)| \leq 1. \]
Since counters can only increase by one per simulation $L_1$-phase, and increment stalls nodes which are ahead, the induction hypothesis holds. In the second case, at least one of the nodes was not participating in simulation $L_1$-phase $p - 1$. W.l.o.g., $u$ was not participating. Due to the transition restrictions, $u$ was in $R$-$W$ or $R$-$W$ in $p - 1$. Thus, by Lemma 11, $rC(v,p - 1) \leq 1$. The same line of arguments as above shows that the induction hypothesis holds.

Combining Lemmas 11 and 12, it is easy to see that nodes in state $R$-$W$ or $R$-$W$ stall the counters of neighboring nodes in $CT$. Moreover, nodes in $CT$ cannot transition to $R$-$N$ if they have neighbors in $R$-$W$, nor can they transition to $R$-$N$ if they have neighbors in $R$-$W$. Finally, a node $v$ cannot transition from $R$-$N$ to $R$-$W$ if there are neighboring nodes in $CT$. As such, we get the following lemma (extended from Lemma 22 in [12]):

**Lemma 13.** For any given simulation $L_1$-phase $p$ and two neighboring nodes $u$ and $v$, if $\text{state}(u,p) = \text{state}(v,p) = CT$ then $p_1(v,p) = p_1(u,p)$ and $p_2(v,p) = p_2(u,p)$. In the same terms as [12], $u$ and $v$ have invoked $\text{Reset}2$ the same number of times, and have invoked $\text{Reset}1$ the same number of times since they last invoked $\text{Reset}2$.

In Lemma 13, we prove that $p_1$ and $p_2$ act as proper indexes for original $L_1$ and $L_2$-phases. Two neighboring participating nodes are simulating the same $L_1$-phase (in the same $L_2$-phase). Moreover, since $rC$ is a balanced counter, we can adapt communications (as in BET) so that EBET simulates the rounds of an original $L_1$-phase.

**C. Balanced Executions in EBET**

We extend the simulation $L_1$-phases with 3 additional slots. Thus, a simulation $L_1$-phase contains 11 slots. The 3 extra slots are dedicated to the simulation of a round $r$ in $\mathcal{P}$. That simulated round is either $rC$ or $rC - 1$, depending on the $rC$ values of the neighboring nodes.

We define a **correct action**, for any given participating node $v$ and simulation $L_1$-phase $p$ of $\mathcal{P}_{\text{sim}}$, $v$’s action when simulating round $r$ in simulation $L_1$-phase $p$ is said to be **correct** if it is the same as $v$’s action in round $r$ of $\mathcal{P}$. We prove that all actions (simulating rounds of $\mathcal{P}$) done by nodes in $\mathcal{P}_{\text{sim}}$ are correct. Thus, $\mathcal{P}_{\text{sim}}$ and $\mathcal{P}$ have the same result.

1) **Rules to ensure Balanced Execution:** We give the following additional rules. They ensure, that for any given participating node $v$ and simulation $L_1$-phase $p$ of $\mathcal{P}_{\text{sim}}$, $v$’s actions in $L_1$-phase $p$ is correct.

- If $\text{next}(v,p) = \text{false}$, $v$ simulates round $rC(v,p) - 1$.
- Otherwise, $v$ simulates round $rC(v,p)$.

A round $r$ is simulated by $v$ in the following way. If $v$’s action for $r$ is $\text{BEEP}$, then $v$ beeps in slot $r \mod 3 + 8$ of simulation $L_1$-phase $p$, and otherwise it listens in that slot.

With the rules above, the following definitions are natural. For any given node $v$ and for any simulated round $r$ of $\mathcal{P}$, we define $p_n(v,r)$ as the first simulation $L_1$-phase $p$ in which $v$ simulates the next round $(r + 1)$. We also define $p_{f}(v,r)$ as the first simulation $L_1$-phase $p$ in which $v$ simulates round $r$.

Now, consider **end of phase rounds** of $\mathcal{P}$ (rounds in which a node ends an $L_n$-phase). A participating node $v$ detects (and chooses) whether it terminates after round $r$ of $\mathcal{P}$ in simulation $L_1$-phase $p_n(v,r)$, since in that phase, $v$ is already done with beeping or listening to beeps for round $r$ (since even the slowest neighbors simulated $r$ in the previous phase). Consequently, consider $rF$ as the round after which $v$ ends an $L_n$-phase in $\mathcal{P}$. $v$ invokes $\text{reset}$ or $\text{reset}2$ in simulation $L_1$-phase $p_n(v,rF)$, which ensures the simulation of $\mathcal{P}$ is correct.

2) **Simulation Proofs:** First, we give the following simple lemma. It states that when a node $v$ is simulating round $rC(v,p) - 1$ in a simulation $L_1$-phase $p$, it has already simulated the round once, in a previous simulation phase. The round is simulated again while $v$ is waiting for the slower nodes (with smaller $rC$ values), until next$(v)$ is true, in which case all neighboring nodes have caught up.

**Lemma 14.** For any given phase $p > 0$ and participating node $v$, $v$ has already simulated round $rC(v) - 1$ at least once.

Now, we prove a crucial lemma. Basically, it states that for any simulation $L_1$-phase $p$, all nodes have correctly simulated $\mathcal{P}$ for all rounds $r < rC(v,p)$. Moreover, in the round in which a participating node $v$ increments $rC(v)$, $rC(v)$ is simulated correctly. This is due to the fact that beeps are transmitted properly despite the different $rC$ values, over multiple simulation $L_1$-phases.

**Lemma 15.** For any given simulation $L_1$-phase $p$ and participating node $v$, all previous actions from $v$ were correct.

1) Moreover, if next$(v,p) = \text{true}$:

   a) If $v$ listens for round $rC(v) - 1$: $\exists u \in N(v)$, $u$ participating, s.t. $u$ beeps for $rC(v) - 1 \leftrightarrow v$ detects a (correct) beep for $rC(v) - 1$ in a phase $p' < p$.

   b) If $v$ beeps for round $rC(v) - 1$: $\exists u \in N(v)$, $u$ participating, s.t. $u$ listens for $rC(v) - 1 \leftrightarrow u$ detects a (correct) beep for $rC(v) - 1$ in a phase $p' < p$.

   c) $v$’s action for round $rC(v,p)$ is correct.
2) Otherwise, v’s action for round $rC(v, p) - 1$ is correct.

Proof. Let us prove this lemma by induction on the simulation $\mathcal{L}_1$-phase $p$. For $p = 0$, the induction hypothesis (IH) holds obviously.

For the induction step, consider a phase $p > 0$ and any given participating node $v$. First, from the IH in phase $p - 1$, we get that all actions done by $v$ previous to phase $p - 1$ were correct, as well as the action $v$ executed in $p - 1$.

Next, let us prove part 1a and 1b of the IH. Consider any given phase $p'$ in which $v$ or any of its neighbors simulates $rC(v, p) - 1$ for the first time. In part 1 of the IH, $\text{next}(v, p) = \text{true}$ thus $p' < p$ (Lemma 14).

Let us prove ($\Rightarrow$) of parts 1a and 1b. Consider $u \in N(v)$ s.t. $u$ beeps (resp. listens) for $rC(v, p) - 1$. We prove $v$ detects $u$’s beep (resp. $v$ detects $v$’s beep). The faster node of the pair ($u$ and $v$) is stalled by the slower node. When the slower node first simulates $rC(v, p) - 1$ in a phase $p' < p$, the faster node $w$ is still simulating $rC(v, p) - 1$ because $\text{next}(w, p')$ is false. Thus, in $p'$, $v$ detects $u$’s beep (resp. $u$ detects $v$’s beep). By the IH, any beep heard is correct.

$(\Leftarrow)$ follows from the fact that beeps are transmitted to neighboring nodes only and because of the manner in which the last three slots are used (and non participating nodes do not use them).

Since all previous actions done by $v$ were correct and part 1a of the IH holds (for phase $p$), part 1c of the IH holds.

Finally, let us prove part 2 of the IH. Suppose $\text{next}(v, p) = \text{false}$. We know $v$’s action for $rC(v, p) - 1$ in phase $p - 1$ is correct, by part 1a of the IH or part 2 of the IH (depending on $\text{next}(v, p - 1)$). Since the action chosen by $v$ for round $rC(v, p) - 1$ does not change, part 2 of the IH holds.

The previous lemma gives us the following theorem:

**Theorem 9.** The outputs of $\mathcal{P}$ and $\mathcal{P}_{sim}$ are identical.

With the previous theorem, there is only one obstacle remaining before we can justify our use of EBET. We need to prove that the round complexity of $\mathcal{P}_{sim}$ is close to that of $\mathcal{P}$. We prove the cost of using EBET is a small multiplicative factor impacting the round complexity.

**Theorem 10.** Let $R_{sim}$ be the round complexity of $\mathcal{P}_{sim}$ and $R$ be that of $\mathcal{P}$. Then $R_{sim} = O(R)$.

Proof. First, there is a constant factor (here 11) between the number of rounds and the number of simulation $\mathcal{L}_1$-phases in $\mathcal{P}_{sim}$. Thus, we compare the number of simulation $\mathcal{L}_1$-phases in $\mathcal{P}_{sim}$ and the number of rounds in $\mathcal{P}$.

Let $L$ be any given original $\mathcal{L}_1$-phase of $\mathcal{P}$. Let $w$ be the node which takes the most rounds to end $L$, that quantity being $r_w$. In $\mathcal{P}_{sim}$, for any given node, at most $r_w$ simulation $\mathcal{L}_1$-phases are used to simulate $L$. Since this holds for all original $\mathcal{L}_1$-phases, and starting the next $\mathcal{L}_1$-phase takes a constant number of simulation $\mathcal{L}_1$-phases, we have $R_{sim} = O(R)$. 

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