Principles of
Distributed Computing

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## Contents

1 Vertex Coloring .............................................. 5  
  1.1 Problem & Model ......................................... 5  
  1.2 Coloring Trees ........................................... 8  

2 Leader Election ............................................ 15  
  2.1 Anonymous Leader Election ............................. 15  
  2.2 Asynchronous Ring ....................................... 16  
  2.3 Lower Bounds ............................................ 18  
  2.4 Synchronous Ring ....................................... 20  

3 Tree Algorithms .......................................... 23  
  3.1 Broadcast ................................................ 23  
  3.2 Convergecast ............................................. 24  
  3.3 BFS Tree Construction .................................. 25  
  3.4 MST Construction ....................................... 27  

4 Distributed Sorting ...................................... 31  
  4.1 Array & Mesh ............................................. 31  
  4.2 Sorting Networks ........................................ 34  
  4.3 Counting Networks ...................................... 37  

5 Shared Memory ............................................ 45  
  5.1 Introduction ............................................. 45  
  5.2 Mutual Exclusion ....................................... 46  
  5.3 Store & Collect .......................................... 49  
    5.3.1 Problem Definition .................................. 49  
    5.3.2 Splitters ............................................ 50  
    5.3.3 Binary Splitter Tree ................................ 51  
    5.3.4 Splitter Matrix ..................................... 53  

6 Shared Objects ........................................... 57  
  6.1 Introduction ............................................. 57  
  6.2 Arrow and Friends ..................................... 58  
  6.3 Ivy and Friends ......................................... 63  

7 Maximal Independent Set ................................ 69  
  7.1 MIS ....................................................... 69  
  7.2 Original Fast MIS ....................................... 71  
  7.3 Fast MIS v2 .............................................. 74
CONTENTS

7.4 Applications . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . 77

8 Locality Lower Bounds 83
8.1 Locality . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . 84
8.2 The Neighborhood Graph . . . . . . . . . . . . . . . . . . . . . . 86

9 Social Networks 91
9.1 Small World Networks . . . . . . . . . . . . . . . . . . . . . . . . 92
9.2 Propagation Studies . . . . . . . . . . . . . . . . . . . . . . . . . 98

10 Synchronization 101
10.1 Basics . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . 101
10.2 Synchronizer $\alpha$ . . . . . . . . . . . . . . . . . . . . . . . . . . 102
10.3 Synchronizer $\beta$ . . . . . . . . . . . . . . . . . . . . . . . . . . . 103
10.4 Synchronizer $\gamma$ . . . . . . . . . . . . . . . . . . . . . . . . . . 104
10.5 Network Partition . . . . . . . . . . . . . . . . . . . . . . . . . . . 106
10.6 Clock Synchronization . . . . . . . . . . . . . . . . . . . . . . . . . 108

11 Hard Problems 115
11.1 Diameter & APSP . . . . . . . . . . . . . . . . . . . . . . . . . . . 115
11.2 Lower Bound Graphs . . . . . . . . . . . . . . . . . . . . . . . . . . 117
11.3 Communication Complexity . . . . . . . . . . . . . . . . . . . . . . 120
11.4 Distributed Complexity Theory . . . . . . . . . . . . . . . . . . . . 126

12 Stabilization 129
12.1 Self-Stabilization . . . . . . . . . . . . . . . . . . . . . . . . . . . 129
12.2 Advanced Stabilization . . . . . . . . . . . . . . . . . . . . . . . . 134

13 Wireless Protocols 139
13.1 Basics . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . 139
13.2 Initialization . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . 141
13.2.1 Non-Uniform Initialization . . . . . . . . . . . . . . . . . . 141
13.2.2 Uniform Initialization with CD . . . . . . . . . . . . . . . . . 141
13.2.3 Uniform Initialization without CD . . . . . . . . . . . . . . . 142
13.3 Leader Election . . . . . . . . . . . . . . . . . . . . . . . . . . . . . 143
13.3.1 With High Probability . . . . . . . . . . . . . . . . . . . . . . 143
13.3.2 Uniform Leader Election . . . . . . . . . . . . . . . . . . . . 143
13.3.3 Fast Leader Election with CD . . . . . . . . . . . . . . . . . 144
13.3.4 Even Faster Leader Election with CD . . . . . . . . . . . . . 145
13.3.5 Lower Bound . . . . . . . . . . . . . . . . . . . . . . . . . . . 147
13.3.6 Uniform Asynchronous Wakeup without CD . . . . . . . . . 147
13.4 Useful Formulas . . . . . . . . . . . . . . . . . . . . . . . . . . . . 148

14 Peer-to-Peer Computing 151
14.1 Introduction . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . 151
14.2 Architecture Variants . . . . . . . . . . . . . . . . . . . . . . . . . . 152
14.3 Hypercubic Networks . . . . . . . . . . . . . . . . . . . . . . . . . . 153
14.4 DHT & Churn . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . 159
14.5 Storage and Multicast . . . . . . . . . . . . . . . . . . . . . . . . . 161
Introduction

What is Distributed Computing?

In the last few decades, we have experienced an unprecedented growth in the area of distributed systems and networks. Distributed computing now encompasses many of the activities occurring in today’s computer and communications world. Indeed, distributed computing appears in quite diverse application areas: Typical “old school” examples are parallel computers, or the Internet. More recent application examples of distributed systems include peer-to-peer systems, sensor networks, and multi-core architectures.

These applications have in common that many processors or entities (often called nodes) are active in the system at any moment. The nodes have certain degrees of freedom: they may have their own hardware, their own code, and sometimes their own independent task. Nevertheless, the nodes may share common resources and information, and, in order to solve a problem that concerns several—or maybe even all—nodes, coordination is necessary.

Despite these commonalities, a peer-to-peer system, for example, is quite different from a multi-core architecture. Due to such differences, many different models and parameters are studied in the area of distributed computing. In some systems the nodes operate synchronously, in other systems they operate asynchronously. There are simple homogeneous systems, and heterogeneous systems where different types of nodes, potentially with different capabilities, objectives etc., need to interact. There are different communication techniques: nodes may communicate by exchanging messages, or by means of shared memory. Sometimes the communication infrastructure is tailor-made for an application, sometimes one has to work with any given infrastructure. The nodes in a system sometimes work together to solve a global task, occasionally the nodes are autonomous agents that have their own agenda and compete for common resources. Sometimes the nodes can be assumed to work correctly, at times they may exhibit failures. In contrast to a single-node system, distributed systems may still function correctly despite failures as other nodes can take over the work of the failed nodes. There are different kinds of failures that can be considered: nodes may just crash, or they might exhibit an arbitrary, erroneous behavior, maybe even to a degree where it cannot be distinguished from malicious (also known as Byzantine) behavior. It is also possible that the nodes follow the rules indeed, however they tweak the parameters to get the most out of the system; in other words, the nodes act selfishly.

Apparently, there are many models (and even more combinations of models) that can be studied. We will not discuss them in greater detail now, but simply
define them when we use them. Towards the end of the course a general picture
should emerge. Hopefully!

This course introduces the basic principles of distributed computing, high-
lighting common themes and techniques. In particular, we study some of the
fundamental issues underlying the design of distributed systems:

• Communication: Communication does not come for free; often communi-
cation cost dominates the cost of local processing or storage. Sometimes
we even assume that everything but communication is free.

• Coordination: How can you coordinate a distributed system so that it
performs some task efficiently?

• Fault-tolerance: As mentioned above, one major advantage of a distrib-
uted system is that even in the presence of failures the system as a whole
may survive.

• Locality: Networks keep growing. Luckily, global information is not always
needed to solve a task, often it is sufficient if nodes talk to their neighbors.
In this course, we will address the fundamental question in distributed
computing whether a local solution is possible for a wide range of problems.

• Parallelism: How fast can you solve a task if you increase your computa-
tional power, e.g., by increasing the number of nodes that can share the
workload? How much parallelism is possible for a given problem?

• Symmetry breaking: Sometimes some nodes need to be selected to or-
chestrate the computation (and the communication). This is typically
achieved by a technique called symmetry breaking.

• Synchronization: How can you implement a synchronous algorithm in an
asynchronous system?

• Uncertainty: If we need to agree on a single term that fittingly describes
this course, it is probably “uncertainty”. As the whole system is distrib-
uted, the nodes cannot know what other nodes are doing at this exact
moment, and the nodes are required to solve the tasks at hand despite the
lack of global knowledge.

Finally, there are also a few areas that we will not cover in this course,
mostly because these topics have become so important that they deserve and
have their own courses. Examples for such topics are distributed programming,
software engineering, as well as security and cryptography.

In summary, in this class we explore essential algorithmic ideas and lower
bound techniques, basically the “pearls” of distributed computing and network
algorithms. We will cover a fresh topic every week.

Have fun!

Chapter Notes

Many excellent text books have been written on the subject. The book closest
to this course is by David Peleg [Pel00], as it shares about half of the material.
A main focus of Peleg’s book are network partitions, covers, decompositions, spanners, and labeling schemes, an interesting area we will only touch in this course. There exist a multitude of other text books that overlap with one or two chapters of this course, e.g., [Lei92, Bar96, Lyn96, Tel01, AW04, HKP+05, CLRS09, Suo12].

Some chapters of this course have been developed in collaboration with (former) Ph.D. students, see chapter notes for details. Many students have helped to improve exercises and script. Thanks go to Philipp Brandes, Raphael Eidenbenz, Roland Flury, Klaus-Tycho Förster, Stephan Holzer, Barbara Keller, Fabian Kuhn, Christoph Lenzen, Thomas Locher, Remo Meier, Thomas Moscibroda, Regina O’Dell, Yvonne Anne Pignolet, Jochen Seidel, Stefan Schmid, Johannes Schneider, Jara Uitto, Pascal von Rickenbach (in alphabetical order).

Bibliography


Chapter 1

Vertex Coloring

1.1 Problem & Model

Vertex coloring is an infamous graph theory problem. It is also a useful toy example to see the style of this course already in the first lecture. Vertex coloring does have quite a few practical applications, for example in the area of wireless networks where coloring is the foundation of so-called TDMA MAC protocols. Generally speaking, vertex coloring is used as a means to break symmetries, one of the main themes in distributed computing. In this chapter we will not really talk about vertex coloring applications, but treat the problem abstractly. At the end of the class you probably learned the fastest (but not constant!) algorithm ever! Let us start with some simple definitions and observations.

**Problem 1.1 (Vertex Coloring).** Given an undirected graph $G = (V,E)$, assign a color $c_u$ to each vertex $u \in V$ such that the following holds: $e = (v,w) \in E \Rightarrow c_v \neq c_w$.

**Remarks:**

- Throughout this course, we use the terms vertex and node interchangeably.
- The application often asks us to use few colors! In a TDMA MAC protocol, for example, less colors immediately imply higher throughput. However, in distributed computing we are often happy with a solution which is sub-optimal. There is a tradeoff between the optimality of a solution (efficacy), and the work/time needed to compute the solution (efficiency).

![Figure 1.1: 3-colorable graph with a valid coloring.](image-url)
Assumption 1.2 (Node Identifiers). Each node has a unique identifier, e.g., its IP address. We usually assume that each identifier consists of only \( \log n \) bits if the system has \( n \) nodes.

Remarks:

- Sometimes we might even assume that the nodes exactly have identifiers \( 1, \ldots, n \).
- It is easy to see that node identifiers (as defined in Assumption 1.2) solve the coloring problem 1.1, but not very well (essentially requiring \( n \) colors). How many colors are needed is a well-studied problem.

Definition 1.3 (Chromatic Number). Given an undirected Graph \( G = (V, E) \), the chromatic number \( \chi(G) \) is the minimum number of colors to solve Problem 1.1.

To get a better understanding of the vertex coloring problem, let us first look at a simple non-distributed (“centralized”) vertex coloring algorithm:

Algorithm 1 Greedy Sequential
1. while \( \exists \) uncolored vertex \( v \) do
2. color \( v \) with the minimal color (number) that does not conflict with the already colored neighbors
3. end while

Definition 1.4 (Degree). The number of neighbors of a vertex \( v \), denoted by \( \delta(v) \), is called the degree of \( v \). The maximum degree vertex in a graph \( G \) defines the graph degree \( \Delta(G) = \Delta \).

Theorem 1.5 (Analysis of Algorithm 1). The algorithm is correct and terminates in \( n \) “steps”. The algorithm uses at most \( \Delta + 1 \) colors.

Proof: Correctness and termination are straightforward. Since each node has at most \( \Delta \) neighbors, there is always at least one color free in the range \( \{1, \ldots, \Delta + 1\} \).

Remarks:

- In Definition 1.7 we will see what is meant by “step”.
- For many graphs coloring can be done with much less than \( \Delta + 1 \) colors.
- This algorithm is not distributed at all; only one processor is active at a time. Still, maybe we can use the simple idea of Algorithm 1 to define a distributed coloring subroutine that may come in handy later.

Now we are ready to study distributed algorithms for this problem. The following procedure can be executed by every vertex \( v \) in a distributed coloring algorithm. The goal of this subroutine is to improve a given initial coloring.
### Procedure 2 First Free

**Require:** Node Coloring (e.g., node IDs as defined in Assumption 1.2)

Give \( v \) the smallest admissible color \( \{i.e., \text{the smallest node color not used by any neighbor}\} \)

**Remarks:**
- With this subroutine we have to make sure that two adjacent vertices are not colored at the same time. Otherwise, the neighbors may at the same time conclude that some small color \( c \) is still available in their neighborhood, and then at the same time decide to choose this color \( c \).

### Definition 1.6 (Synchronous Distributed Algorithm)

In a synchronous algorithm, nodes operate in synchronous rounds. In each round, each processor executes the following steps:

1. Do some local computation (of reasonable complexity).
2. Send messages to neighbors in graph (of reasonable size).
3. Receive messages (that were sent by neighbors in step 2 of the same round).

**Remarks:**
- Any other step ordering is fine.
- What does “reasonable” mean in this context? We are somewhat flexible here, and different model variants exist. Generally, we will deal with algorithms that only do very simple computations (a comparison, an addition, etc.). Exponential-time computation is usually considered cheating in this context. Similarly, sending a message with a node ID, or a value is considered okay, whereas sending really long messages is considered cheating. We will do more exact definitions later, when we need them.

### Algorithm 3 Reduce

1: Assume that initially all nodes have ID’s (Assumption 1.2)
2: Each node \( v \) executes the following code
3: node \( v \) sends its ID to all neighbors
4: node \( v \) receives IDs of neighbors
5: while node \( v \) has an uncolored neighbor with higher ID do
   6: node \( v \) sends “undecided” to all neighbors
   7: node \( v \) receives new decisions from neighbors
8: end while
9: node \( v \) chooses a free color using subroutine First Free (Procedure 2)
10: node \( v \) informs all its neighbors about its choice

### Definition 1.7 (Time Complexity)

For synchronous algorithms (as defined in 1.6) the time complexity is the number of rounds until the algorithm terminates.
CHAPTER 1. VERTEX COLORING

 Remarks:

- The algorithm terminates when the last processor has decided to terminate.
- To guarantee correctness the procedure requires a legal input (i.e., pairwise different node IDs).

Theorem 1.8 (Analysis of Algorithm 3). Algorithm 3 is correct and has time complexity $n$. The algorithm uses at most $\Delta + 1$ colors.

 Remarks:

- Quite trivial, but also quite slow.
- However, it seems difficult to come up with a fast algorithm.
- Maybe it’s better to first study a simple special case, a tree, and then go from there.

1.2 Coloring Trees

Lemma 1.9. $\chi(\text{Tree}) \leq 2$

Constructive Proof: If the distance of a node to the root is odd (even), color it 1 (0). An odd node has only even neighbors and vice versa. If we assume that each node knows its parent (root has no parent) and children in a tree, this constructive proof gives a very simple algorithm:

Algorithm 4 Slow Tree Coloring
1: Color the root 0, root sends 0 to its children
2: Each node $v$ concurrently executes the following code:
3: if node $v$ receives a message $x$ (from parent) then
4: node $v$ chooses color $c_v = 1 - x$
5: node $v$ sends $c_v$ to its children (all neighbors except parent)
6: end if
Remarks:

• With the proof of Lemma 1.9, Algorithm 4 is correct.

• How can we determine a root in a tree if it is not already given? We will figure that out later.

• The time complexity of the algorithm is the height of the tree.

• If the root was chosen unfortunately, and the tree has a degenerated topology, the time complexity may be up to \( n \), the number of nodes.

• Also, this algorithm does not need to be synchronous . . . !

Definition 1.10 (Asynchronous Distributed Algorithm). In the asynchronous model, algorithms are event driven (“upon receiving message . . . , do . . . ”). Processors cannot access a global clock. A message sent from one processor to another will arrive in finite but unbounded time.

Remarks:

• The asynchronous model and the synchronous model (Definition 1.6) are the cornerstone models in distributed computing. As they do not necessarily reflect reality there are several models in between synchronous and asynchronous. However, from a theoretical point of view the synchronous and the asynchronous model are the most interesting ones (because every other model is in between these extremes).

• Note that in the asynchronous model, messages that take a longer path may arrive earlier.

Definition 1.11 (Time Complexity). For asynchronous algorithms (as defined in 1.6) the time complexity is the number of time units from the start of the execution to its completion in the worst case (every legal input, every execution scenario), assuming that each message has a delay of at most one time unit.

Remarks:

• You cannot use the maximum delay in the algorithm design. In other words, the algorithm has to be correct even if there is no such delay upper bound.

Definition 1.12 (Message Complexity). The message complexity of a synchronous or asynchronous algorithm is determined by the number of messages exchanged (again every legal input, every execution scenario).

Theorem 1.13 (Analysis of Algorithm 4). Algorithm 4 is correct. If each node knows its parent and its children, the (asynchronous) time complexity is the tree height which is bounded by the diameter of the tree; the message complexity is \( n - 1 \) in a tree with \( n \) nodes.
CHAPTER 1. VERTEX COLORING

Remarks:

• In this case the asynchronous time complexity is the same as the synchronous time complexity.

• Nice trees, e.g. balanced binary trees, have logarithmic height, that is we have a logarithmic time complexity.

• This algorithm is not very exciting. Can we do better than logarithmic?!?

The following algorithm terminates in $\log^* n$ time. Log-Star?! That’s the number of logarithms (to the base 2) you need to take to get down to at least 2, starting with $n$:

**Definition 1.14 (Log-Star).**

$\forall x \leq 2 : \log^* x := 1$

$\forall x > 2 : \log^* x := 1 + \log^*(\log x)$

Remarks:

• Log-star is an amazingly slowly growing function. Log-star of all the atoms in the observable universe (estimated to be $10^{80}$) is 5! There are functions which grow even more slowly, such as the inverse Ackermann function, however, the inverse Ackermann function of all the atoms is 4. So log-star increases indeed very slowly!

Here is the idea of the algorithm: We start with color labels that have $\log n$ bits. In each synchronous round we compute a new label with exponentially smaller size than the previous label, still guaranteeing to have a valid vertex coloring! But how are we going to do that?

**Algorithm 5 “6-Color”**

1: Assume that initially the vertices are legally colored. Using Assumption 1.2 each label only has $\log n$ bits

2: The root assigns itself the label 0.

3: Each other node $v$ executes the following code (synchronously in parallel)

4: send $c_v$ to all children

5: repeat

6: receive $c_p$ from parent

7: interpret $c_v$ and $c_p$ as little-endian bit-strings: $c(k), \ldots, c(1), c(0)$

8: let $i$ be the smallest index where $c_v$ and $c_p$ differ

9: the new label is $i$ (as bitstring) followed by the bit $c_v(i)$ itself

10: send $c_v$ to all children

11: until $c_w \in \{0, \ldots, 5\}$ for all nodes $w$

**Example:**

Algorithm 5 executed on the following part of a tree:

<table>
<thead>
<tr>
<th>Grand-parent</th>
<th>Parent</th>
<th>Child</th>
</tr>
</thead>
<tbody>
<tr>
<td>0010110000</td>
<td>1010010000</td>
<td>0110010000</td>
</tr>
<tr>
<td>$\rightarrow$</td>
<td>$\rightarrow$</td>
<td>$\rightarrow$</td>
</tr>
<tr>
<td>10010</td>
<td>01010</td>
<td>10001</td>
</tr>
<tr>
<td>$\rightarrow$</td>
<td>$\rightarrow$</td>
<td>$\rightarrow$</td>
</tr>
</tbody>
</table>

**Theorem 1.15.** Algorithm 5 terminates in $\log^* n$ time.
Remarks:

• Colors 11* (in binary notation, i.e., 6 or 7 in decimal notation) will not be chosen, because the node will then do another round. This gives a total of 6 colors (i.e., colors 0,..., 5).

• Can one reduce the number of colors in only constant steps? Note that algorithm 3 does not work (since the degree of a node can be much higher than 6)! For fewer colors we need to have siblings monochromatic!

• Before we explore this problem we should probably have a second look at the end game of the algorithm, the UNTIL statement. Is this algorithm truly local?! Let’s discuss!

Algorithm 6 Shift Down
1: Root chooses a new (different) color from \{0, 1, 2\}
2: Each other node \(v\) concurrently executes the following code:
3: Recolor \(v\) with the color of parent

Lemma 1.16 (Analysis of Algorithm 6). Algorithm 6 preserves coloring legality; also siblings are monochromatic.

Now Algorithm 3 (Reduce) can be used to reduce the number of used colors from six to three.

Algorithm 7 Six-2-Three
1: Each node \(v\) concurrently executes the following code:
2: Run Algorithm 5 for \(\log^* n\) rounds.
3: for \(x = 5, 4, 3\) do
4: Perform subroutine Shift down (Algorithm 6)
5: if \(c_v = x\) then
6: choose new color \(c_v \in \{0, 1, 2\}\) using subroutine First Free (Algorithm 2)
7: end if
8: end for

Theorem 1.17 (Analysis of Algorithm 7). Algorithm 7 colors a tree with three colors in time \(O(\log^* n)\).

Remarks:

• The term \(O()\) used in Theorem 1.15 is called “big O” and is often used in distributed computing. Roughly speaking, \(O(f)\) means “in the order of \(f\), ignoring constant factors and smaller additive terms.” More formally, for two functions \(f\) and \(g\), it holds that \(f \in O(g)\) if there are constants \(x_0\) and \(c\) so that \(|f(x)| \leq c|g(x)|\) for all \(x \geq x_0\). For an elaborate discussion on the big O notation we refer to other introductory math or computer science classes.
• As one can easily prove, a fast tree-coloring with only 2 colors is more than exponentially more expensive than coloring with 3 colors. In a tree degenerated to a list, nodes far away need to figure out whether they are an even or odd number of hops away from each other in order to get a 2-coloring. To do that one has to send a message to these nodes. This costs time linear in the number of nodes.

• The idea of this algorithm can be generalized, e.g., to a ring topology. Also a general graph with constant degree $\Delta$ can be colored with $\Delta + 1$ colors in $O(\log^* n)$ time. The idea is as follows: In each step, a node compares its label to each of its neighbors, constructing a logarithmic difference-tag as in 6-color (Algorithm 5). Then the new label is the concatenation of all the difference-tags. For constant degree $\Delta$, this gives a $3\Delta$-label in $O(\log^* n)$ steps. Algorithm 3 then reduces the number of colors to $\Delta + 1$ in $2^{3\Delta}$ (this is still a constant for constant $\Delta$) steps.

• Unfortunately, coloring a general graph is not yet possible with this technique. We will see another technique for that in Chapter 7. With this technique it is possible to color a general graph with $\Delta + 1$ colors in $O(\log n)$ time.
• A lower bound shows that many of these log-star algorithms are asymptotically (up to constant factors) optimal. We will also see that later.

Chapter Notes

The basic technique of the log-star algorithm is by Cole and Vishkin [CV86]. The technique can be generalized and extended, e.g. to a ring topology or to graphs with constant degree [GP87, GPS88, KMW05]. Using it as a subroutine, one can solve many problems in log-star time. For instance, one can color so-called growth bounded graphs (a model which includes many natural graph classes, for instance unit disk graphs) asymptotically optimally in $O(\log^* n)$ time [SW08]. Actually, Schneider et al. show that many classic combinatorial problems beyond coloring can be solved in log-star time in growth bounded and other restricted graphs.

In a later chapter we learn a $\Omega(\log^* n)$ lower bound for coloring and related problems [Lin92]. Linial’s paper also contains a number of other results on coloring, e.g. that any algorithm for coloring $d$-regular trees of radius $r$ that run in time at most $2r/3$ require at least $\Omega(\sqrt{d})$ colors.

For general graphs, later we will learn fast coloring algorithms that use a maximal independent sets as a base. Since coloring exhibits a trade-off between efficacy and efficiency, many different results for general graphs exist, e.g. [PS96, KSOS06, BE09, Kuh09, SW10, BE11b, KP11, BE11a].

Some parts of this chapter are also discussed in Chapter 7 of [Pel00], e.g., the proof of Theorem 1.15.

Bibliography


CHAPTER 1. VERTEX COLORING


Chapter 2

Leader Election

2.1 Anonymous Leader Election

Some algorithms (e.g. the slow tree coloring algorithm) ask for a special node, a so-called “leader”. Computing a leader is a very simple form of symmetry breaking. Algorithms based on leaders do generally not exhibit a high degree of parallelism, and therefore often suffer from poor time complexity. However, sometimes it is still useful to have a leader to make critical decisions in an easy (though non-distributed!) way.

The process of choosing a leader is known as leader election. Although leader election is a simple form of symmetry breaking, there are some remarkable issues that allow us to introduce notable computational models.

In this chapter we concentrate on the ring topology. Many interesting challenges in distributed computing already reveal the root of the problem in the special case of the ring. Paying special attention to the ring also makes sense from a practical point of view as some real world systems are based on a ring topology, e.g., the token ring standard for local area networks.

Problem 2.1 (Leader Election). Each node eventually decides whether it is a leader or not, subject to the constraint that there is exactly one leader.

Remarks:

• More formally, nodes are in one of three states: undecided, leader, not leader. Initially every node is in the undecided state. When leaving the undecided state, a node goes into a final state (leader or not leader).

Definition 2.2 (Anonymous). A system is anonymous if nodes do not have unique identifiers.

Definition 2.3 (Uniform). An algorithm is called uniform if the number of nodes n is not known to the algorithm (to the nodes, if you wish). If n is known, the algorithm is called non-uniform.

Whether a leader can be elected in an anonymous system depends on whether the network is symmetric (ring, complete graph, complete bipartite graph, etc.) or asymmetric (star, single node with highest degree, etc.). Simplifying slightly, in this context a symmetric graph is a graph in which the extended neighborhood
of each node has the same structure. We will now show that non-uniform anonymous leader election for synchronous rings is impossible. The idea is that in a ring, symmetry can always be maintained.

**Lemma 2.4.** After round \( k \) of any deterministic algorithm on an anonymous ring, each node is in the same state \( s_k \).

Proof by induction: All nodes start in the same state. A round in a synchronous algorithm consists of the three steps sending, receiving, local computation (see Definition 1.6). All nodes send the same message(s), receive the same message(s), do the same local computation, and therefore end up in the same state.

**Theorem 2.5** (Anonymous Leader Election). Deterministic leader election in an anonymous ring is impossible.

Proof (with Lemma 2.4): If one node ever decides to become a leader (or a non-leader), then every other node does so as well, contradicting the problem specification 2.1 for \( n > 1 \). This holds for non-uniform algorithms, and therefore also for uniform algorithms. Furthermore, it holds for synchronous algorithms, and therefore also for asynchronous algorithms.

**Remarks:**

- Sense of direction is the ability of nodes to distinguish neighbor nodes in an anonymous setting. In a ring, for example, a node can distinguish the clockwise and the counterclockwise neighbor. Sense of direction does not help in anonymous leader election.

- Theorem 2.5 also holds for other symmetric network topologies (e.g., complete graphs, complete bipartite graphs, ...).

- Note that Theorem 2.5 does not hold for randomized algorithms; if nodes are allowed to toss a coin, some symmetries can be broken.

### 2.2 Asynchronous Ring

We first concentrate on the asynchronous model from Definition 1.10. Throughout this section we assume non-anonymity; each node has a unique identifier as proposed in Assumption 1.2. Having ID’s seems to lead to a trivial leader election algorithm, as we can simply elect the node with, e.g., the highest ID.

**Theorem 2.6** (Analysis of Algorithm 8). Algorithm 8 is correct. The time complexity is \( O(n) \). The message complexity is \( O(n^2) \).

Proof: Let node \( z \) be the node with the maximum identifier. Node \( z \) sends its identifier in clockwise direction, and since no other node can swallow it, eventually a message will arrive at \( z \) containing it. Then \( z \) declares itself to be the leader. Every other node will declare non-leader at the latest when forwarding message \( z \). Since there are \( n \) identifiers in the system, each node will at most forward \( n \) messages, giving a message complexity of at most \( n^2 \).

We start measuring the time when the first node that "wakes up" sends its identifier. For asynchronous time complexity (Definition 1.11) we assume that each message takes at most one time unit to arrive at its destination. After at
Algorithm 8 Clockwise

1: Each node $v$ executes the following code:
2: $v$ sends a message with its identifier (for simplicity also $v$) to its clockwise neighbor. (If node $v$ already received a message $w$ with $w > v$, then node $v$ can skip this step; if node $v$ receives its first message $w$ with $w < v$, then node $v$ will immediately send $v$.)
3: if $v$ receives a message $w$ with $w > v$ then
4: $v$ forwards $w$ to its clockwise neighbor
5: $v$ decides not to be the leader, if it has not done so already.
6: else if $v$ receives its own identifier $v$ then
7: $v$ decides to be the leader
8: end if

most $n - 1$ time units the message therefore arrives at node $z$, waking $z$ up. Routing the message $z$ around the ring takes at most $n$ time units. Therefore node $z$ decides no later than at time $2n - 1$. Every other node decides before node $z$.

Remarks:

- Note that in Algorithm 8 nodes need to distinguish between clockwise and counterclockwise neighbors. In fact they do not: It is okay to simply send your own identifier to any neighbor, and forward a message $m$ to the neighbor you did not receive the message $m$ from. So nodes only need to be able to distinguish their two neighbors.

- Careful analysis shows, that while having worst-case message complexity of $O(n^2)$, Algorithm 8 has an average message complexity of $O(n \log n)$. Can we improve this algorithm?

Theorem 2.7 (Analysis of Algorithm 9). Algorithm 9 is correct. The time complexity is $O(n)$. The message complexity is $O(n \log n)$.

Proof: Correctness is as in Theorem 2.6. The time complexity is $O(n)$ since the node with maximum identifier $z$ sends messages with round-trip times $2, 4, 8, 16, \ldots, 2 \cdot 2^k$ with $k \leq \log(n + 1)$. (Even if we include the additional wake-up overhead, the time complexity stays linear.) Proving the message complexity is slightly harder: if a node $v$ manages to survive round $r$, no other node in distance $2^r$ (or less) survives round $r$. That is, node $v$ is the only node in its $2^r$-neighborhood that remains active in round $r + 1$. Since this is the same for every node, less than $n/2^r$ nodes are active in round $r + 1$. Being active in round $r$ costs $2 \cdot 2 \cdot 2^r$ messages. Therefore, round $r$ costs at most $2 \cdot 2 \cdot 2^r \cdot \frac{n}{2^r} = 8n$ messages. Since there are only logarithmic many possible rounds, the message complexity follows immediately.

Remarks:

- This algorithm is asynchronous and uniform as well.

- The question may arise whether one can design an algorithm with an even lower message complexity. We answer this question in the next section.
Algorithm 9 Radius Growth (For readability we provide pseudo-code only; for a formal version please consult [Attiya/Welch Alg. 3.1])

1. Each node $v$ does the following:
2. Initially all nodes are active. {all nodes may still become leaders}
3. Whenever a node $v$ sees a message $w$ with $w > v$, then $v$ decides to not be a leader and becomes passive.
4. Active nodes search in an exponentially growing neighborhood (clockwise and counterclockwise) for nodes with higher identifiers, by sending out probe messages. A probe message includes the ID of the original sender, a bit whether the sender can still become a leader, and a time-to-live number ($TTL$). The first probe message sent by node $v$ includes a TTL of 1.
5. Nodes (active or passive) receiving a probe message decrement the TTL and forward the message to the next neighbor; if their ID is larger than the one in the message, they set the leader bit to zero, as the probing node does not have the maximum ID. If the TTL is zero, probe messages are returned to the sender using a reply message. The reply message contains the ID of the receiver (the original sender of the probe message) and the leader-bit. Reply messages are forwarded by all nodes until they reach the receiver.
6. Upon receiving the reply message: If there was no node with higher ID in the search area (indicated by the bit in the reply message), the TTL is doubled and two new probe messages are sent (again to the two neighbors). If there was a better candidate in the search area, then the node becomes passive.
7. If a node $v$ receives its own probe message (not a reply) $v$ decides to be the leader.

2.3 Lower Bounds

Lower bounds in distributed computing are often easier than in the standard centralized (random access machine, RAM) model because one can argue about messages that need to be exchanged. In this section we present a first lower bound. We show that Algorithm 9 is asymptotically optimal.

Definition 2.8 (Execution). An execution of a distributed algorithm is a list of events, sorted by time. An event is a record (time, node, type, message), where type is “send” or “receive”.

Remarks:

- We assume throughout this course that no two events happen at exactly the same time (or one can break ties arbitrarily).

- An execution of an asynchronous algorithm is generally not only determined by the algorithm but also by a “god-like” scheduler. If more than one message is in transit, the scheduler can choose which one arrives first.

- If two messages are transmitted over the same directed edge, then it is sometimes required that the message first transmitted will also be received first (“FIFO”).

For our lower bound, we assume the following model:
• We are given an asynchronous ring, where nodes may wake up at arbitrary times (but at the latest when receiving the first message).

• We only accept uniform algorithms where the node with the maximum identifier can be the leader. Additionally, every node that is not the leader must know the identity of the leader. These two requirements can be dropped when using a more complicated proof; however, this is beyond the scope of this course.

• During the proof we will “play god” and specify which message in transmission arrives next in the execution. We respect the FIFO conditions for links.

Definition 2.9 (Open Schedule). A schedule is an execution chosen by the scheduler. An open (undirected) edge is an edge where no message traversing the edge has been received so far. A schedule for a ring is open if there is an open edge in the ring.

The proof of the lower bound is by induction. First we show the base case:

Lemma 2.10. Given a ring $R$ with two nodes, we can construct an open schedule in which at least one message is received. The nodes cannot distinguish this schedule from one on a larger ring with all other nodes being where the open edge is.

Proof: Let the two nodes be $u$ and $v$ with $u < v$. Node $u$ must learn the identity of node $v$, thus receive at least one message. We stop the execution of the algorithm as soon as the first message is received. (If the first message is received by $v$, bad luck for the algorithm!) Then the other edge in the ring (on which the received message was not transmitted) is open. Since the algorithm needs to be uniform, maybe the open edge is not really an edge at all, nobody can tell. We could use this to glue two rings together, by breaking up this imaginary open edge and connect two rings by two edges. An example can be seen in Figure 2.1.

Lemma 2.11. By gluing together two rings of size $n/2$ for which we have open schedules, we can construct an open schedule on a ring of size $n$. If $M(n/2)$ denotes the number of messages already received in each of these schedules, at least $2M(n/2) + n/4$ messages have to be exchanged in order to solve leader election.

Proof by induction: We divide the ring into two sub-rings $R_1$ and $R_2$ of size $n/2$. These subrings cannot be distinguished from rings with $n/2$ nodes if no
messages are received from “outsiders”. We can ensure this by not scheduling such messages until we want to. Note that executing both given open schedules on $R_1$ and $R_2$ “in parallel” is possible because we control not only the scheduling of the messages, but also when nodes wake up. By doing so, we make sure that $2M(n/2)$ messages are sent before the nodes in $R_1$ and $R_2$ learn anything of each other!

Without loss of generality, $R_1$ contains the maximum identifier. Hence, each node in $R_2$ must learn the identity of the maximum identifier, thus at least $n/2$ additional messages must be received. The only problem is that we cannot connect the two sub-rings with both edges since the new ring needs to remain open. Thus, only messages over one of the edges can be received. We look into the future: we check what happens when we close only one of these connecting edges.

Since we know that $n/2$ nodes have to be informed in $R_2$, there must be at least $n/2$ messages that must be received. Closing both edges must inform $n/2$ nodes, thus for one of the two edges there must be a node in distance $n/4$ which will be informed upon creating that edge. This results in $n/4$ additional messages. Thus, we pick this edge and leave the other one open which yields the claim.

**Lemma 2.12.** Any uniform leader election algorithm for asynchronous rings has at least message complexity $M(n) \geq \frac{n}{4} (\log n + 1)$.

Proof by induction: For the sake of simplicity we assume $n$ being a power of 2. The base case $n = 2$ works because of Lemma 2.10 which implies that $M(2) \geq 1 = \frac{2}{4} (\log 2 + 1)$. For the induction step, using Lemma 2.11 and the induction hypothesis we have

$$M(n) = 2 \cdot M\left(\frac{n}{2}\right) + \frac{n}{4}$$

$$\geq 2 \cdot \left(\frac{n}{8} \left(\log \frac{n}{2} + 1\right)\right) + \frac{n}{4}$$

$$= \frac{n}{4} \log n + \frac{n}{4} = \frac{n}{4} (\log n + 1).$$

$\square$

**Remarks:**

- To hide the ugly constants we use the “big Omega” notation, the lower bound equivalent of $O()$. A function $f$ is in $\Omega(g)$ if there are constants $x_0$ and $c > 0$ such that $|f(x)| \geq c|g(x)|$ for all $x \geq x_0$.Again we refer to standard text books for a formal definition. Rewriting Lemma 2.12 we get:

**Theorem 2.13** (Asynchronous Leader Election Lower Bound). Any uniform leader election algorithm for asynchronous rings has $\Omega(n \log n)$ message complexity.

### 2.4 Synchronous Ring

The lower bound relied on delaying messages for a very long time. Since this is impossible in the synchronous model, we might get a better message complexity.
in this case. The basic idea is very simple: In the synchronous model, not receiving a message is information as well! First we make some additional assumptions:

- We assume that the algorithm is non-uniform (i.e., the ring size $n$ is known).
- We assume that every node starts at the same time.
- The node with the minimum identifier becomes the leader; identifiers are integers.

Algorithm 10 Synchronous Leader Election

1: Each node $v$ concurrently executes the following code:
2: The algorithm operates in synchronous phases. Each phase consists of $n$ time steps. Node $v$ counts phases, starting with 0.
3: if phase = $v$ and $v$ did not yet receive a message then
4: $v$ decides to be the leader
5: $v$ sends the message “$v$ is leader” around the ring
6: end if

Remarks:

- Message complexity is indeed $n$.
- But the time complexity is huge! If $m$ is the minimum identifier it is $m \cdot n$.
- The synchronous start and the non-uniformity assumptions can be dropped by using a wake-up technique (upon receiving a wake-up message, wake up your clockwise neighbors) and by letting messages travel slowly.
- There are several lower bounds for the synchronous model: comparison-based algorithms or algorithms where the time complexity cannot be a function of the identifiers have message complexity $\Omega(n \log n)$ as well.
- In general graphs efficient leader election may be tricky. While time-optimal leader election can be done by parallel flooding-echo (see next chapter), bounding the message complexity is generally more difficult.

Chapter Notes

[Ang80] was the first to mention the now well-known impossibility result for anonymous rings and other networks, even when using randomization. The first algorithm for asynchronous rings was presented in [Lan77], which was improved to the presented clockwise algorithm in [CR79]. Later, [HS80] found the radius growth algorithm, which decreased the worst case message complexity. Algorithms for the unidirectional case with runtime $O(n \log n)$ can be found in [DKR82, Pet82]. The $\Omega(n \log n)$ message complexity lower bound for comparison based algorithms was first published in [FL87]. In [Sch89] an algorithm with constant error probability for anonymous networks is presented. General results about limitations of computer power in synchronous rings are in [ASW88, AS88].
CHAPTER 2. LEADER ELECTION

Bibliography


Chapter 3

Tree Algorithms

In this chapter we learn a few basic algorithms on trees, and how to construct trees in the first place so that we can run these (and other) algorithms. The good news is that these algorithms have many applications, the bad news is that this chapter is a bit on the simple side. But maybe that's not really bad news?!

3.1 Broadcast

Definition 3.1 (Broadcast). A broadcast operation is initiated by a single processor, the source. The source wants to send a message to all other nodes in the system.

Definition 3.2 (Distance, Radius, Diameter). The distance between two nodes \( u \) and \( v \) in an undirected graph \( G \) is the number of hops of a minimum path between \( u \) and \( v \). The radius of a node \( u \) is the maximum distance between \( u \) and any other node in the graph. The radius of a graph is the minimum radius of any node in the graph. The diameter of a graph is the maximum distance between two arbitrary nodes.

Remarks:

- Clearly there is a close relation between the radius \( R \) and the diameter \( D \) of a graph, such as \( R \leq D \leq 2R \).

- The world is often fascinated by graphs with a small radius. For example, movie fanatics study the who-acted-with-whom-in-the-same-movie graph. For this graph it has long been believed that the actor Kevin Bacon has a particularly small radius. The number of hops from Bacon even got a name, the Bacon Number. In the meantime, however, it has been shown that there are “better” centers in the Hollywood universe, such as Sean Connery, Christopher Lee, Rod Steiger, Gene Hackman, or Michael Caine. The center of other social networks has also been explored, Paul Erdős for instance is well known in the math community.

Theorem 3.3 (Broadcast Lower Bound). The message complexity of broadcast is at least \( n - 1 \). The source’s radius is a lower bound for the time complexity.
Proof: Every node must receive the message.

Remarks:

- You can use a pre-computed spanning tree to do broadcast with tight message complexity. If the spanning tree is a breadth-first search spanning tree (for a given source), then the time complexity is tight as well.

**Definition 3.4 (Clean).** A graph (network) is clean if the nodes do not know the topology of the graph.

**Theorem 3.5 (Clean Broadcast Lower Bound).** For a clean network, the number of edges is a lower bound for the broadcast message complexity.

Proof: If you do not try every edge, you might miss a whole part of the graph behind it.

Remarks:

- This lower bound proof directly brings us to the well known flooding algorithm.

**Algorithm 11 Flooding**

1: The source (root) sends the message to all neighbors.
2: Each other node $v$ upon receiving the message the first time forwards the message to all (other) neighbors.
3: Upon later receiving the message again (over other edges), a node can discard the message.

Remarks:

- If node $v$ receives the message first from node $u$, then node $v$ calls node $u$ parent. This parent relation defines a spanning tree $T$. If the flooding algorithm is executed in a synchronous system, then $T$ is a breadth-first search spanning tree (with respect to the root).

- More interestingly, also in asynchronous systems the flooding algorithm terminates after $R$ time units, $R$ being the radius of the source. However, the constructed spanning tree may not be a breadth-first search spanning tree.

### 3.2 Convergecast

Convergecast is the same as broadcast, just reversed: Instead of a root sending a message to all other nodes, all other nodes send information to a root. The simplest convergecast algorithm is the echo algorithm:
Algorithm 12 Echo

Require: This algorithm is initiated at the leaves.
1: A leave sends a message to its parent.
2: If an inner node has received a message from each child, it sends a message to the parent.

Remarks:

• Usually the echo algorithm is paired with the flooding algorithm, which is used to let the leaves know that they should start the echo process; this is known as flooding/echo.

• One can use convergecast for termination detection, for example. If a root wants to know whether all nodes in the system have finished some task, it initiates a flooding/echo; the message in the echo algorithm then means “This subtree has finished the task.”

• Message complexity of the echo algorithm is $n - 1$, but together with flooding it is $O(m)$, where $m = |E|$ is the number of edges in the graph.

• The time complexity of the echo algorithm is determined by the depth of the spanning tree (i.e., the radius of the root within the tree) generated by the flooding algorithm.

• The flooding/echo algorithm can do much more than collecting acknowledgements from subtrees. One can for instance use it to compute the number of nodes in the system, or the maximum ID (for leader election), or the sum of all values stored in the system, or a route-disjoint matching.

• Moreover, by combining results one can compute even fancier aggregations, e.g., with the number of nodes and the sum one can compute the average. With the average one can compute the standard deviation. And so on . . .

3.3 BFS Tree Construction

In synchronous systems the flooding algorithm is a simple yet efficient method to construct a breadth-first search (BFS) spanning tree. However, in asynchronous systems the spanning tree constructed by the flooding algorithm may be far from BFS. In this section, we implement two classic BFS constructions—Dijkstra and Bellman-Ford—as asynchronous algorithms.

We start with the Dijkstra algorithm. The basic idea is to always add the “closest” node to the existing part of the BFS tree. We need to parallelize this idea by developing the BFS tree layer by layer:

Theorem 3.6 (Analysis of Algorithm 13). The time complexity of Algorithm 13 is $O(D^2)$, the message complexity is $O(m + nD)$, where $D$ is the diameter of the graph, $n$ the number of nodes, and $m$ the number of edges.

Proof: A broadcast/echo algorithm in $T_p$ needs at most time $2D$. Finding new neighbors at the leaves costs 2 time units. Since the BFS tree height is bounded
CHAPTER 3. TREE ALGORITHMS

Algorithm 13 Dijkstra BFS

1. The algorithm proceeds in phases. In phase $p$ the nodes with distance $p$ to the root are detected. Let $T_p$ be the tree in phase $p$. We start with $T_1$ which is the root plus all direct neighbors of the root. We start with phase $p = 1$:
2. repeat
3. The root starts phase $p$ by broadcasting “start $p$” within $T_p$.
4. When receiving “start $p$” a leaf node $u$ of $T_p$ (that is, a node that was newly discovered in the last phase) sends a “join $p + 1$” message to all quiet neighbors. (A neighbor $v$ is quiet if $u$ has not yet “talked” to $v$.)
5. A node $v$ receiving the first “join $p + 1$” message replies with “ACK” and becomes a leaf of the tree $T_{p+1}$.
6. A node $v$ receiving any further “join” message replies with “NACK”.
7. The leaves of $T_p$ collect all the answers of their neighbors; then the leaves start an echo algorithm back to the root.
8. When the echo process terminates at the root, the root increments the phase
9. until there was no new node detected

by the diameter, we have $D$ phases, giving a total time complexity of $O(D^2)$. Each node participating in broadcast/echo only receives (broadcasts) at most 1 message and sends (echoes) at most once. Since there are $D$ phases, the cost is bounded by $O(nD)$. On each edge there are at most 2 “join” messages. Replies to a “join” request are answered by 1 “ACK” or “NACK”, which means that we have at most 4 additional messages per edge. Therefore the message complexity is $O(m + nD)$.

Remarks:
- The time complexity is not very exciting, so let’s try Bellman-Ford!

The basic idea of Bellman-Ford is even simpler, and heavily used in the Internet, as it is a basic version of the omnipresent border gateway protocol (BGP). The idea is to simply keep the distance to the root accurate. If a neighbor has found a better route to the root, a node might also need to update its distance.

Algorithm 14 Bellman-Ford BFS

1. Each node $u$ stores an integer $d_u$ which corresponds to the distance from $u$ to the root. Initially $d_{\text{root}} = 0$, and $d_u = \infty$ for every other node $u$.
2. The root starts the algorithm by sending “1” to all neighbors.
3. if a node $u$ receives a message “$y$” with $y < d_u$ from a neighbor $v$ then
4. node $u$ sets $d_u := y$
5. node $u$ sends “$y + 1$” to all neighbors (except $v$)
6. end if

Theorem 3.7 (Analysis of Algorithm 14). The time complexity of Algorithm 14 is $O(D)$, the message complexity is $O(nm)$, where $D, n, m$ are defined as in Theorem 3.6.

Proof: We can prove the time complexity by induction. We claim that a node at distance $d$ from the root has received a message “$d$” by time $d$. The root
knows by time 0 that it is the root. A node \( v \) at distance \( d \) has a neighbor \( u \) at distance \( d - 1 \). Node \( u \) by induction sends a message “\( d \)” to \( v \) at time \( d - 1 \) or before, which is then received by \( v \) at time \( d \) or before. Message complexity is easier: A node can reduce its distance at most \( n - 1 \) times; each of these times it sends a message to all its neighbors. If all nodes do this we have \( O(nm) \) messages.

Remarks:
- Algorithm 13 has the better message complexity and Algorithm 14 has the better time complexity. The currently best algorithm (optimizing both) needs \( O(m + n \log^3 n) \) messages and \( O(D \log^3 n) \) time. This “trade-off” algorithm is beyond the scope of this chapter, but we will later learn the general technique.

### 3.4 MST Construction

There are several types of spanning trees, each serving a different purpose. A particularly interesting spanning tree is the minimum spanning tree (MST). The MST only makes sense on weighted graphs, hence in this section we assume that each edge \( e \) is assigned a weight \( \omega_e \).

**Definition 3.8 (MST).** Given a weighted graph \( G = (V, E, \omega) \), the MST of \( G \) is a spanning tree \( T \) minimizing \( \omega(T) \), where \( \omega(G') = \sum_{e \in G'} \omega_e \) for any subgraph \( G' \subseteq G \).

**Remarks:**
- In the following we assume that no two edges of the graph have the same weight. This simplifies the problem as it makes the MST unique; however, this simplification is not essential as one can always break ties by adding the IDs of adjacent vertices to the weight.
- Obviously we are interested in computing the MST in a distributed way. For this we use a well-known lemma:

**Definition 3.9 (Blue Edges).** Let \( T \) be a spanning tree of the weighted graph \( G \) and \( T' \subseteq T \) a subgraph of \( T \) (also called a fragment). Edge \( e = (u, v) \) is an outgoing edge of \( T' \) if \( u \in T' \) and \( v \notin T' \) (or vice versa). The minimum weight outgoing edge \( b(T') \) is the so-called blue edge of \( T' \).

**Lemma 3.10.** For a given weighted graph \( G \) (such that no two weights are the same), let \( T \) denote the MST, and \( T' \) be a fragment of \( T \). Then the blue edge of \( T' \) is also part of \( T \), i.e., \( T' \cup b(T') \subseteq T \).

Proof: For the sake of contradiction, suppose that in the MST \( T \) there is edge \( e \neq b(T') \) connecting \( T' \) with the remainder of \( T \). Adding the blue edge \( b(T') \) to the MST \( T \) we get a cycle including both \( e \) and \( b(T') \). If we remove \( e \) from this cycle we still have a spanning tree, and since by the definition of the blue edge \( \omega_e > \omega_{b(T')}, \) the weight of that new spanning tree is less than than the weight of \( T \). We have a contradiction.
Remarks:

• In other words, the blue edges seem to be the key to a distributed algorithm for the MST problem. Since every node itself is a fragment of the MST, every node directly has a blue edge! All we need to do is to grow these fragments! Essentially this is a distributed version of Kruskal’s sequential algorithm.

• At any given time the nodes of the graph are partitioned into fragments (rooted subtrees of the MST). Each fragment has a root, the ID of the fragment is the ID of its root. Each node knows its parent and its children in the fragment. The algorithm operates in phases. At the beginning of a phase, nodes know the IDs of the fragments of their neighbor nodes.

Algorithm 15 GHS (Gallager–Humblet–Spira)

1: Initially each node is the root of its own fragment. We proceed in phases:
2: repeat
3: All nodes learn the fragment IDs of their neighbors.
4: The root of each fragment uses flooding/echo in its fragment to determine the blue edge \( b = (u, v) \) of the fragment.
5: The root sends a message to node \( u \) while forwarding the message on the path from the root to node \( u \) all parent-child relations are inverted {such that \( u \) is the new temporary root of the fragment}
6: node \( u \) sends a merge request over the blue edge \( b = (u, v) \).
7: if node \( v \) also sent a merge request over the same blue edge \( b = (v, u) \)
   then
   either \( u \) or \( v \) (whichever has the smaller ID) is the new fragment root
   the blue edge \( b \) is directed accordingly
   else
   node \( v \) is the new parent of node \( u \)
   end if
11: the newly elected root node informs all nodes in its fragment (again using flooding/echo) about its identity
13: until all nodes are in the same fragment (i.e., there is no outgoing edge)

Remarks:

• Algorithm 15 was stated in pseudo-code, with a few details not really explained. For instance, it may be that some fragments are much larger than others, and because of that some nodes may need to wait for others, e.g., if node \( u \) needs to find out whether neighbor \( v \) also wants to merge over the blue edge \( b = (u, v) \). The good news is that all these details can be solved. We can for instance bound the asynchronicity by guaranteeing that nodes only start the new phase after the last phase is done, similarly to the phase-technique of Algorithm 13.

Theorem 3.11 (Analysis of Algorithm 15). The time complexity of Algorithm 15 is \( O(n \log n) \), the message complexity is \( O(m \log n) \).

Proof: Each phase mainly consists of two flooding/echo processes. In general, the cost of flooding/echo on a tree is \( O(D) \) time and \( O(n) \) messages. However,
the diameter $D$ of the fragments may turn out to be not related to the diameter of the graph because the MST may meander, hence it really is $O(n)$ time. In addition, in the first step of each phase, nodes need to learn the fragment ID of their neighbors; this can be done in 2 steps but costs $O(m)$ messages. There are a few more steps, but they are cheap. Altogether a phase costs $O(n)$ time and $O(m)$ messages. So we only have to figure out the number of phases: Initially all fragments are single nodes and hence have size 1. In a later phase, each fragment merges with at least one other fragment, that is, the size of the smallest fragment at least doubles. In other words, we have at most $\log n$ phases. The theorem follows directly.

Remarks:

- The GHS algorithm can be applied in different ways. GHS for instance directly solves leader election in general graphs: The leader is simply the last surviving root!

Chapter Notes

Trees are one of the oldest graph structures, already appearing in the first book about graph theory [Koe36]. Broadcasting in distributed computing is younger, but not that much [DM78]. Overviews about broadcasting can be found for example in Chapter 3 of [Pel00] and Chapter 7 of [HKP+05]. For a introduction to centralized tree-construction, see e.g. [Eve79] or [CLRS09]. Overviews for the distributed case can be found in Chapter 5 of [Pel00] or Chapter 4 of [Lyn96]. The classic papers on routing are [For56, Bel58, Dij59]. In a later chapter, we will later learn a general technique to derive algorithms with an almost optimal time and message complexity.

Algorithm 15 is called “GHS” after Gallager, Humblet, and Spira, three pioneers in distributed computing [GHS83]. Their algorithm won the prestigious Edsger W. Dijkstra Prize in Distributed Computing in 2004, among other reasons because it was one of the first non-trivial asynchronous distributed algorithms. As such it can be seen as one of the seeds of this research area. We presented a simplified version of GHS. The original paper featured an improved message complexity of $O(m + n \log n)$. Later, Awerbuch managed to further improve the GHS algorithm to get $O(n)$ time and $O(m + n \log n)$ message complexity, both asymptotically optimal [Awe87].

Bibliography


Chapter 4

Distributed Sorting

“Indeed, I believe that virtually every important aspect of programming arises somewhere in the context of sorting [and searching]!”

– Donald E. Knuth, The Art of Computer Programming

In this chapter we study a classic problem in computer science—sorting—from a distributed computing perspective. In contrast to an orthodox single-processor sorting algorithm, no node has access to all data, instead the to-be-sorted values are distributed. Distributed sorting then boils down to:

**Definition 4.1 (Sorting).** We choose a graph with $n$ nodes $v_1, \ldots, v_n$. Initially each node stores a value. After applying a sorting algorithm, node $v_k$ stores the $k^{th}$ smallest value.

Remarks:

• What if we route all values to the same central node $v$, let $v$ sort the values locally, and then route them to the correct destinations?! According to the message passing model studied in the first few chapters this is perfectly legal. With a star topology sorting finishes in $O(1)$ time!

**Definition 4.2 (Node Contention).** In each step of a synchronous algorithm, each node can only send and receive $O(1)$ messages containing $O(1)$ values, no matter how many neighbors the node has.

Remarks:

• Using Definition 4.2 sorting on a star graph takes linear time.

4.1 Array & Mesh

To get a better intuitive understanding of distributed sorting, we start with two simple topologies, the array and the mesh. Let us begin with the array:

31
Algorithm 16 Odd/Even Sort

1: Given an array of \( n \) nodes \( \{v_1, \ldots, v_n\} \), each storing a value (not sorted).
2: repeat
3:   Compare and exchange the values at nodes \( i \) and \( i + 1 \), \( i \) odd
4:   Compare and exchange the values at nodes \( i \) and \( i + 1 \), \( i \) even
5: until done

Remarks:

- The compare and exchange primitive in Algorithm 16 is defined as follows: Let the value stored at node \( i \) be \( v_i \). After the compare and exchange node \( i \) stores value \( \min(v_i, v_{i+1}) \) and node \( i + 1 \) stores value \( \max(v_i, v_{i+1}) \).
- How fast is the algorithm, and how can we prove correctness/efficiency?
- The most interesting proof uses the so-called 0-1 Sorting Lemma. It allows us to restrict our attention to an input of 0’s and 1’s only, and works for any “oblivious comparison-exchange” algorithm. (Oblivious means: Whether you exchange two values must only depend on the relative order of the two values, and not on anything else.)

Lemma 4.3 (0-1 Sorting Lemma). If an oblivious comparison-exchange algorithm sorts all inputs of 0’s and 1’s, then it sorts arbitrary inputs.

Proof. We prove the opposite direction (does not sort arbitrary inputs \( \Rightarrow \) does not sort 0’s and 1’s). Assume that there is an input \( x = x_1, \ldots, x_n \) that is not sorted correctly. Then there is a smallest value \( k \) such that the value at node \( v_k \) after running the sorting algorithm is strictly larger than the \( k \)th smallest value \( x(k) \). Define an input \( x^* \) such that \( x_i = x^* \) if \( x_i \leq x(k) \) and \( x_i = 1 \) else. Whenever the algorithm compares a pair of 1’s or 0’s, it is not important whether it exchanges the values or not, so we may simply assume that it does the same as on the input \( x \). On the other hand, whenever the algorithm exchanges some values \( x^*_i = 0 \) and \( x^*_j = 1 \), this means that \( x_i \leq x(k) < x_j \). Therefore, in this case the respective compare-exchange operation will do the same on both inputs. We conclude that the algorithm will order \( x^* \) the same way as \( x \), i.e., the output with only 0’s and 1’s will also not be correct.

Theorem 4.4. Algorithm 16 sorts correctly in \( n \) steps.

Proof. Thanks to lemma 4.3 we only need to consider an array with 0’s and 1’s. Let \( j_1 \) be the node with the rightmost (highest index) 1. If \( j_1 \) is odd (even) it will move in the first (second) step. In any case it will move right in every following step until it reaches the rightmost node \( v_n \). Let \( j_k \) be the node with the \( k \)th rightmost 1. We show by induction that \( j_k \) is not “blocked” anymore (constantly moves until it reaches destination!) after step \( k \). We have already anchored the induction at \( k = 1 \). Since \( j_{k-1} \) moves after step \( k - 1 \), \( j_k \) gets a right 0-neighbor for each step after step \( k \). (For matters of presentation we omitted a couple of simple details.)
Algorithm 17 Shearsort

1: We are given a mesh with $m$ rows and $m$ columns, $m$ even, $n = m^2$.
2: The sorting algorithm operates in phases, and uses the odd/even sort algorithm on rows or columns.
3: repeat
4: In the odd phases 1, 3, \ldots we sort all the rows, in the even phases 2, 4, \ldots we sort all the columns, such that:
5: Columns are sorted such that the small values move up.
6: Odd rows (1, 3, \ldots, $m - 1$) are sorted such that small values move left.
7: Even rows (2, 4, \ldots, $m$) are sorted such that small values move right.
8: until done

Remarks:

- Linear time is not very exciting, maybe we can do better by using a different topology? Let’s try a mesh (a.k.a. grid) topology first.

Theorem 4.5. Algorithm 17 sorts $n$ values in $\sqrt{n}(\log n + 1)$ time in snake-like order.

Proof. Since the algorithm is oblivious, we can use lemma 4.3. We show that after a row and a column phase, half of the previously unsorted rows will be sorted. More formally, let us call a row with only 0’s (or only 1’s) clean, a row with 0’s and 1’s is dirty. At any stage, the rows of the mesh can be divided into three regions. In the north we have a region of all-0 rows, in the south all-1 rows, in the middle a region of dirty rows. Initially all rows can be dirty. Since neither row nor column sort will touch already clean rows, we can concentrate on the dirty rows.

First we run an odd phase. Then, in the even phase, we run a peculiar column sorter: We group two consecutive dirty rows into pairs. Since odd and even rows are sorted in opposite directions, two consecutive dirty rows look as follows:

\[
\begin{align*}
&00000 \ldots 11111 \\
&11111 \ldots 00000
\end{align*}
\]

Such a pair can be in one of three states. Either we have more 0’s than 1’s, or more 1’s than 0’s, or an equal number of 0’s and 1’s. Column-sorting each pair will give us at least one clean row (and two clean rows if “$|0| = |1|$”). Then move the cleaned rows north/south and we will be left with half the dirty rows.

At first glance it appears that we need such a peculiar column sorter. However, any column sorter sorts the columns in exactly the same way (we are very grateful to have lemma 4.3!).

All in all we need $2 \log m = \log n$ phases to remain only with 1 dirty row in the middle which will be sorted (not cleaned) with the last row-sort. \qed
CHAPTER 4. DISTRIBUTED SORTING

Remarks:

• There are algorithms that sort in $3m + o(m)$ time on an $m$ by $m$ mesh (by diving the mesh into smaller blocks). This is asymptotically optimal, since a value might need to move $2m$ times.

• Such a $\sqrt{n}$-sorter is cute, but we are more ambitious. There are non-distributed sorting algorithms such as quicksort, heapsort, or mergesort that sort $n$ values in (expected) $O(n \log n)$ time. Using our $n$-fold parallelism effectively we might therefore hope for a distributed sorting algorithm that sorts in time $O(\log n)!$

4.2 Sorting Networks

In this section we construct a graph topology which is carefully manufactured for sorting. This is a deviation from previous chapters where we always had to work with the topology that was given to us. In many application areas (e.g. peer-to-peer networks, communication switches, systolic hardware) it is indeed possible (in fact, crucial!) that an engineer can build the topology best suited for her application.

Definition 4.6 (Sorting Networks). A comparator is a device with two inputs $x, y$ and two outputs $x', y'$ such that $x' = \min(x, y)$ and $y' = \max(x, y)$. We construct so-called comparison networks that consist of wires that connect comparators (the output port of a comparator is sent to an input port of another comparator). Some wires are not connected to comparator outputs, and some are not connected to comparator inputs. The first are called input wires of the comparison network, the second output wires. Given $n$ values on the input wires, a sorting network ensures that the values are sorted on the output wires. We will also use the term width to indicate the number of wires in the sorting network.

Remarks:

• The odd/even sorter explained in Algorithm 16 can be described as a sorting network.

• Often we will draw all the wires on $n$ horizontal lines ($n$ being the “width” of the network). Comparators are then vertically connecting two of these lines.

• Note that a sorting network is an oblivious comparison-exchange network. Consequently we can apply lemma 4.3 throughout this section. An example sorting network is depicted in figure 4.1.

Definition 4.7 (Depth). The depth of an input wire is 0. The depth of a comparator is the maximum depth of its input wires plus one. The depth of an output wire of a comparator is the depth of the comparator. The depth of a comparison network is the maximum depth (of an output wire).

Definition 4.8 (Bitonic Sequence). A bitonic sequence is a sequence of numbers that first monotonically increases, and then monotonically decreases, or vice versa.
4.2. SORTING NETWORKS

Figure 4.1: A sorting network.

Remarks:
• $< 1, 4, 6, 8, 3, 2 >$ or $< 5, 3, 2, 1, 4, 8 >$ are bitonic sequences.
• $< 9, 6, 2, 3, 5, 4 >$ or $< 7, 4, 2, 5, 9, 8 >$ are not bitonic.
• Since we restrict ourselves to 0's and 1's (lemma 4.3), bitonic sequences have the form $0^i1^j0^k$ or $1^i0^j1^k$ for $i, j, k \geq 0$.

Algorithm 18 Half Cleaner
1: A half cleaner is a comparison network of depth 1, where we compare wire $i$ with wire $i + n/2$ for $i = 1, \ldots, n/2$ (we assume $n$ to be even).

Lemma 4.9. Feeding a bitonic sequence into a half cleaner (Algorithm 18), the half cleaner cleans (makes all-0 or all-1) either the upper or the lower half of the $n$ wires. The other half is bitonic.

Proof. Assume that the input is of the form $0^i1^j0^k$ for $i, j, k \geq 0$. If the midpoint falls into the 0’s, the input is already clean/bitonic and will stay so. If the midpoint falls into the 1’s the half cleaner acts as Shearsort with two adjacent rows, exactly as in the proof of theorem 4.5. The case $1^i0^j1^k$ is symmetric. □

Algorithm 19 Bitonic Sequence Sorter
1: A bitonic sequence sorter of width $n$ ($n$ being a power of 2) consists of a half cleaner of width $n$, and then two bitonic sequence sorters of width $n/2$ each.
2: A bitonic sequence sorter of width 1 is empty.

Lemma 4.10. A bitonic sequence sorter (Algorithm 19) of width $n$ sorts bitonic sequences. It has depth $\log n$. 
Proof. The proof follows directly from the Algorithm 19 and lemma 4.9.

Remarks:

- Clearly we want to sort arbitrary and not only bitonic sequences! To do this we need one more concept, merging networks.

Algorithm 20 Merging Network

1: A merging network of width \( n \) is a merger of width \( n \) followed by two bitonic sequence sorters of width \( n/2 \). A merger is a depth-one network where we compare wire \( i \) with wire \( n - i + 1 \), for \( i = 1, \ldots, n/2 \).

Remarks:

- Note that a merging network is a bitonic sequence sorter where we replace the (first) half-cleaner by a merger.

Lemma 4.11. A merging network of width \( n \) (Algorithm 20) merges two sorted input sequences of length \( n/2 \) each into one sorted sequence of length \( n \).

Proof. We have two sorted input sequences. Essentially, a merger does to two sorted sequences what a half cleaner does to a bitonic sequence, since the lower part of the input is reversed. In other words, we can use same argument as in theorem 4.5 and lemma 4.9: Again, after the merger step either the upper or the lower half is clean, the other is bitonic. The bitonic sequence sorters complete sorting.

Remarks:

- How do you sort \( n \) values when you are able to merge two sorted sequences of size \( n/2 \)? Piece of cake, just apply the merger recursively.

Algorithm 21 Batcher’s “Bitonic” Sorting Network

1: A batcher sorting network of width \( n \) consists of two batcher sorting networks of width \( n/2 \) followed by a merging network of width \( n \). (See figure 4.2.)

2: A batcher sorting network of width 1 is empty.

Theorem 4.12. A sorting network (Algorithm 21) sorts an arbitrary sequence of \( n \) values. It has depth \( O(\log^2 n) \).

Proof. Correctness is immediate: at recursive stage \( k \) (\( k = 1, 2, 3, \ldots, \log n \)) we merge \( 2^k \) sorted sequences into \( 2^{k-1} \) sorted sequences. The depth \( d(n) \) of the sorting network of level \( n \) is the depth of a sorting network of level \( n/2 \) plus the depth \( m(n) \) of a merging network with width \( n \). The depth of a sort of level 1 is 0 since the sorter is empty. Since a merging network of width \( n \) has the same depth as a bitonic sequence sorter of width \( n \), we know by lemma 4.10 that \( m(n) = \log n \). This gives a recursive formula for \( d(n) \) which solves to \( d(n) = \frac{1}{2} \log^2 n + \frac{1}{2} \log n \).
4.3 COUNTING NETWORKS

Remarks:

- Simulating Batcher’s sorting network on an ordinary sequential computer takes time $O(n \log^2 n)$. As said, there are sequential sorting algorithms that sort in asymptotically optimal time $O(n \log n)$. So a natural question is whether there is a sorting network with depth $O(\log n)$. Such a network would have some remarkable advantages over sequential asymptotically optimal sorting algorithms such as heapsort. Apart from being highly parallel, it would be completely oblivious, and as such perfectly suited for a fast hardware solution. In 1983, Ajtai, Komlos, and Szemeredi presented a celebrated $O(\log n)$ depth sorting network. (Unlike Batcher’s sorting network the constant hidden in the big-O of the “AKS” sorting network is too large to be practical, however.)

- It can be shown that Batcher’s sorting network and similarly others can be simulated by a Butterfly network and other hypercubic networks, see next chapter.

- What if a sorting network is asynchronous?? Clearly, using a synchronizer we can still sort, but it is also possible to use it for something else. Check out the next section!

4.3 Counting Networks

In this section we address distributed counting, a distributed service which can for instance be used for load balancing.

**Definition 4.13 (Distributed Counting).** A distributed counter is a variable that is common to all processors in a system and that supports an atomic test-and-increment operation. The operation delivers the system’s counter value to the requesting processor and increments it.
Remarks:

- A naive distributed counter stores the system’s counter value with a distinguished central node. When other nodes initiate the test-and-increment operation, they send a request message to the central node and in turn receive a reply message with the current counter value. However, with a large number of nodes operating on the distributed counter, the central processor will become a bottleneck. There will be a congestion of request messages at the central processor, in other words, the system will not scale.

- Is a scalable implementation (without any kind of bottleneck) of such a distributed counter possible, or is distributed counting a problem which is inherently centralized?!?

- Distributed counting could for instance be used to implement a load balancing infrastructure, i.e. by sending the job with counter value \(i\) (modulo \(n\)) to server \(i\) (out of \(n\) possible servers).

**Definition 4.14 (Balancer).** A balancer is an asynchronous flip-flop which forwards messages that arrive on the left side to the wires on the right, the first to the upper, the second to the lower, the third to the upper, and so on.

**Algorithm 22** Bitonic Counting Network.

1. Take Batcher’s bitonic sorting network of width \(w\) and replace all the comparators with balancers.
2. When a node wants to count, it sends a message to an arbitrary input wire.
3. The message is then routed through the network, following the rules of the asynchronous balancers.
4. Each output wire is completed with a “mini-counter.”
5. The mini-counter of wire \(k\) replies the value “\(k + i \cdot w\)” to the initiator of the \(i\)th message it receives.

**Definition 4.15 (Step Property).** A sequence \(y_0, y_1, \ldots, y_{w-1}\) is said to have the step property, if \(0 \leq y_i - y_j \leq 1\), for any \(i < j\).

Remarks:

- If the output wires have the step property, then with \(r\) requests, exactly the values \(1, \ldots, r\) will be assigned by the mini-counters. All we need to show is that the counting network has the step property. For that we need some additional facts...

**Facts 4.16.** For a balancer, we denote the number of consumed messages on the \(i\)th input wire with \(x_i, i = 0, 1\). Similarly, we denote the number of sent messages on the \(i\)th output wire with \(y_i, i = 0, 1\). A balancer has these properties:

1. A balancer does not generate output-messages; that is, \(x_0 + x_1 \geq y_0 + y_1\) in any state.
2. Every incoming message is eventually forwarded. In other words, if we are in a quiescent state (no message in transit), then \(x_0 + x_1 = y_0 + y_1\).
(3) The number of messages sent to the upper output wire is at most one higher than the number of messages sent to the lower output wire: in any state $y_0 = [(y_0 + y_1)/2]$ (thus $y_1 = [(y_0 + y_1)/2]$).

**Facts 4.17.** If a sequence $y_0, y_1, \ldots, y_{w-1}$ has the step property,

(1) then all its subsequences have the step property,

(2) then its even and odd subsequences satisfy

$$\sum_{i=0}^{\lfloor (w-1)/2 \rfloor} y_{2i} = \left\lfloor \frac{1}{2} \sum_{i=0}^{w-1} y_i \right\rfloor$$

and

$$\sum_{i=0}^{\lfloor (w-1)/2 \rfloor} y_{2i+1} = \left\lceil \frac{1}{2} \sum_{i=0}^{w-1} y_i \right\rceil.$$  

**Facts 4.18.** If two sequences $x_0, x_1, \ldots, x_{w-1}$ and $y_0, y_1, \ldots, y_{w-1}$ have the step property,

(1) and $\sum_{i=0}^{w-1} x_i = \sum_{i=0}^{w-1} y_i$, then $x_i = y_i$ for $i = 0, \ldots, w-1$.

(2) and $\sum_{i=0}^{w-1} x_i = \sum_{i=0}^{w-1} y_i + 1$, then there exists a unique $j$ ($j = 0, 1, \ldots, w-1$) such that $x_j = y_j + 1$, and $x_i = y_i$ for $i = 0, \ldots, w-1$, $i \neq j$.

**Remarks:**

- That’s enough to prove that a Merger preserves the step property.

**Lemma 4.19.** Let $M[w]$ be a Merger of width $w$. In a quiescent state (no message in transit), if the inputs $x_0, x_1, \ldots, x_{w/2-1}$ resp. $x_{w/2}, x_{w/2+1}, \ldots, x_{w-1}$ have the step property, then the output $y_0, y_1, \ldots, y_{w-1}$ has the step property.

**Proof.** By induction on the width $w$.

For $w = 2$: $M[2]$ is a balancer and a balancer’s output has the step property (fact 4.16.3).

For $w > 2$: Let $x_0, \ldots, x_{w/2-1}$ resp. $z_0', \ldots, z'_{w/2-1}$ be the output of the upper respectively lower $M[w/2]$ subnetwork. Since $x_0, x_1, \ldots, x_{w/2-1}$ and $x_{w/2}, x_{w/2+1}, \ldots, x_{w-1}$ both have the step property by assumption, their even and odd subsequences also have the step property (fact 4.17.1). By induction hypothesis, the output of both $M[w/2]$ subnetworks have the step property. Let $Z := \sum_{i=0}^{w/2-1} x_i$ and $Z' := \sum_{i=0}^{w/2-1} z'_i$. From fact 4.17.2 we conclude that $Z = \lfloor \frac{1}{2} \sum_{i=0}^{w/2-1} x_i \rfloor + \lfloor \frac{1}{2} \sum_{i=0}^{w/2-1} x_i \rfloor$ and $Z' = \lfloor \frac{1}{2} \sum_{i=0}^{w/2-1} x_i \rfloor + \lfloor \frac{1}{2} \sum_{i=0}^{w/2-1} x_i \rfloor$. Since $[a] + [b]$ and $[a] + [b]$ differ by at most 1 we know that $Z$ and $Z'$ differ by at most 1.

If $Z = Z'$, fact 4.18.1 implies that $z_i = z'_i$ for $i = 0, \ldots, w/2 - 1$. Therefore, the output of $M[w]$ is $y_i = z_i[2]$ for $i = 0, \ldots, w - 1$. Since $z_0, \ldots, z_{w/2-1}$ has the step property, so does the output of $M[w]$ and the lemma follows.

If $Z$ and $Z'$ differ by 1, fact 4.18.2 implies that $z_i = z'_i$ for $i = 0, \ldots, w/2 - 1$, except a unique $j$ such that $z_j$ and $z'_j$ differ by only 1, for $j = 0, \ldots, w/2 - 1$. Let $l := \min(z_j, z'_j)$. Then, the output $y_i$ (with $i < 2j$) is $l + 1$. The output $y_{2i}$ (with $i > 2j + 1$) is $l$. The output $y_{2j}$ and $y_{2j+1}$ are balanced by the final balancer resulting in $y_{2j} = l + 1$ and $y_{2j+1} = l$. Therefore $M[w]$ preserves the step property. \qed
A bitonic counting network is constructed to fulfill lemma 4.19, i.e., the final output comes from a Merger whose upper and lower inputs are recursively merged. Therefore, the following theorem follows immediately.

**Theorem 4.20 (Correctness).** In a quiescent state, the $w$ output wires of a bitonic counting network of width $w$ have the step property.

**Remarks:**
- Is every sorting network also a counting network? No. But surprisingly, the other direction is true!

**Theorem 4.21 (Counting vs. Sorting).** If a network is a counting network then it is also a sorting network, but not vice versa.

**Proof.** There are sorting networks that are not counting networks (e.g. odd/even sort, or insertion sort). For the other direction, let $C$ be a counting network and $I(C)$ be the isomorphic network, where every balancer is replaced by a comparator. Let $I(C)$ have an arbitrary input of 0’s and 1’s; that is, some of the input wires have a 0, all others have a 1. There is a message at $C$’s $i$th input wire if and only if $I(C)$’s $i$ input wire is 0. Since $C$ is a counting network, all messages are routed to the upper output wires. $I(C)$ is isomorphic to $C$, therefore a comparator in $I(C)$ will receive a 0 on its upper (lower) wire if and only if the corresponding balancer receives a message on its upper (lower) wire. Using an inductive argument, the 0’s and 1’s will be routed through $I(C)$ such that all 0’s exit the network on the upper wires whereas all 1’s exit the network on the lower wires. Applying lemma 4.3 shows that $I(C)$ is a sorting network.

**Remarks:**
- We claimed that the counting network is correct. However, it is only correct in a quiescent state.

**Definition 4.22 (Linearizable).** A system is linearizable if the order of the values assigned reflects the real-time order in which they were requested. More formally, if there is a pair of operations $o_1, o_2$, where operation $o_1$ terminates before operation $o_2$ starts, and the logical order is “$o_2$ before $o_1$”, then a distributed system is not linearizable.

**Lemma 4.23 (Linearizability).** The bitonic counting network is not linearizable.

**Proof.** Consider the bitonic counting network with width 4 in figure 4.3: Assume that two inc operations were initiated and the corresponding messages entered the network on wire 0 and 2 (both in light gray color). After having passed the second resp. the first balancer, these traversing messages “fall asleep”; In other words, both messages take unusually long time before they are received by the next balancer. Since we are in an asynchronous setting, this may be the case.

In the meantime, another inc operation (medium gray) is initiated and enters the network on the bottom wire. The message leaves the network on wire 2, and the inc operation is completed.
Figure 4.3: Linearizability Counter Example.

Strictly afterwards, another inc operation (dark gray) is initiated and enters the network on wire 1. After having passed all balancers, the message will leave the network wire 0. Finally (and not depicted in figure 4.3), the two light gray messages reach the next balancer and will eventually leave the network on wires 1 resp. 3. Because the dark gray and the medium gray operation do conflict with Definition 4.22, the bitonic counting network is not linearizable.

Remarks:

- Note that the example in figure 4.3 behaves correctly in the quiescent state: Finally, exactly the values 0, 1, 2, 3 are allotted.
- It was shown that linearizability comes at a high price (the depth grows linearly with the width).

Chapter Notes

The technique used for the famous lower bound of comparison-based sequential sorting first appeared in [FJ59]. Comprehensive introductions to the vast field of sorting can certainly be found in [Knu73]. Knuth also presents the 0/1 principle in the context of sorting networks, supposedly as a special case of a theorem for decision trees of W. G. Bouricius, and includes a historic overview of sorting network research.

Using a rather complicated proof not based on the 0/1 principle, [Hab72] first presented and analyzed Odd/Even sort on arrays. Shearsort for grids first appeared in [SS86] as a sorting algorithm both easy to implement and to prove correct. Later it was generalized to meshes with higher dimension in [SS89]. A bubble sort based algorithm is presented in [SI86]: it takes time $O(\sqrt{n} \log n)$, but is fast in practice. Nevertheless, already [TK77] presented an asymptotically optimal algorithms for grid network which runs in $3n + O(n^{2/3} \log n)$ rounds for an $n \times n$ grid. A simpler algorithm was later found by [SS86] using $3n + O(n^{3/4})$ rounds.

Batcher presents his famous $O(\log^2 n)$ depth sorting network in [Bat68]. It took until [AKS83] to find a sorting network with asymptotically optimal depth $O(\log n)$. Unfortunately, the constants hidden in the big-O-notation render it rather impractical.
The notion of counting networks was introduced in [AHS91], and shortly afterward the notion of linearizability was studied by [HSW91]. Follow-up work in [AHS94] presents bitonic counting networks and studies contention in the counting network. An overview of research on counting networks can be found in [BH98].

Bibliography


Chapter 5

Shared Memory

5.1 Introduction

In distributed computing, various different models exist. So far, the focus of the course was on loosely-coupled distributed systems such as the Internet, where nodes asynchronously communicate by exchanging messages. The “opposite” model is a tightly-coupled parallel computer where nodes access a common memory totally synchronously—in distributed computing such a system is called a Parallel Random Access Machine (PRAM).

A third major model is somehow between these two extremes, the shared memory model. In a shared memory system, asynchronous processes (or processors) communicate via a common memory area of shared variables or registers:

**Definition 5.1 (Shared Memory).** A shared memory system is a system that consists of asynchronous processes that access a common (shared) memory. A process can atomically access a register in the shared memory through a set of predefined operations. An atomic modification appears to the rest of the system instantaneously. Apart from this shared memory, processes can also have some local (private) memory.

Remarks:

- Various shared memory systems exist. A main difference is how they allow processes to access the shared memory. All systems can atomically read or write a shared register $R$. Most systems do allow for advanced *atomic* read-modify-write (RMW) operations, for example:

  - test-and-set($R$): $t := R$; $R := 1$; return $t$
  - fetch-and-add($R, x$): $t := R$; $R := R + x$; return $t$
  - compare-and-swap($R, x, y$): if $R = x$ then $R := y$; return true; else return false; endif;
  - load-link($R$)/store-conditional($R, x$): Load-link returns the current value of the specified register $R$. A subsequent store-conditional to the same register will store a new value $x$ (and return true) only if no updates have occurred to that register since the load-link. If any updates have occurred, the store-conditional is guaranteed to fail.
The power of RMW operations can be measured with the so-called consensus-number: The consensus-number $k$ of a RMW operation defines whether one can solve consensus for $k$ processes. Test-and-set for instance has consensus-number 2 (one can solve consensus with 2 processes, but not 3), whereas the consensus-number of compare-and-swap is infinite. It can be shown that the power of a shared memory system is determined by the consensus-number (“universality of consensus”). This insight has a remarkable theoretical and practical impact. In practice for instance, after this was known, hardware designers stopped developing shared memory systems supporting weak RMW operations.

Many of the results derived in the message passing model have an equivalent in the shared memory model. Consensus for instance is traditionally studied in the shared memory model.

Whereas programming a message passing system is rather tricky (in particular if fault-tolerance has to be integrated), programming a shared memory system is generally considered easier, as programmers are given access to global variables directly and do not need to worry about exchanging messages correctly. Because of this, even distributed systems which physically communicate by exchanging messages can often be programmed through a shared memory middleware, making the programmer’s life easier.

We will most likely find the general spirit of shared memory systems in upcoming multi-core architectures. As for programming style, the multi-core community seems to favor an accelerated version of shared memory, transactional memory.

From a message passing perspective, the shared memory model is like a bipartite graph: On one side you have the processes (the nodes) which pretty much behave like nodes in the message passing model (asynchronous, maybe failures). On the other side you have the shared registers, which just work perfectly (no failures, no delay).

5.2 Mutual Exclusion

A classic problem in shared memory systems is mutual exclusion. We are given a number of processes which occasionally need to access the same resource. The resource may be a shared variable, or a more general object such as a data structure or a shared printer. The catch is that only one process at the time is allowed to access the resource. More formally:

**Definition 5.2** (Mutual Exclusion). We are given a number of processes, each executing the following code sections:

\[
\text{<Entry>} \rightarrow \text{<Critical Section>} \rightarrow \text{<Exit>} \rightarrow \text{<Remaining Code>}
\]

A mutual exclusion algorithm consists of code for entry and exit sections, such that the following holds...
5.2. MUTUAL EXCLUSION

- **Mutual Exclusion**: At all times at most one process is in the critical section.

- **No deadlock**: If some process manages to get to the entry section, later some (possibly different) process will get to the critical section.

Sometimes we in addition ask for

- **No lockout**: If some process manages to get to the entry section, later the same process will get to the critical section.

- **Unobstructed exit**: No process can get stuck in the exit section.

Using RMW primitives one can build mutual exclusion algorithms quite easily. Algorithm 23 shows an example with the test-and-set primitive.

**Algorithm 23 Mutual Exclusion: Test-and-Set**

<table>
<thead>
<tr>
<th>Input: Shared register $R := 0$</th>
</tr>
</thead>
<tbody>
<tr>
<td>&lt;Entry&gt;</td>
</tr>
<tr>
<td>1: repeat</td>
</tr>
<tr>
<td>2: $r := \text{test-and-set}(R)$</td>
</tr>
<tr>
<td>3: until $r = 0$</td>
</tr>
<tr>
<td>&lt;Critical Section&gt;</td>
</tr>
<tr>
<td>4: ...</td>
</tr>
<tr>
<td>&lt;Exit&gt;</td>
</tr>
<tr>
<td>5: $R := 0$</td>
</tr>
<tr>
<td>&lt;Remainder Code&gt;</td>
</tr>
<tr>
<td>6: ...</td>
</tr>
</tbody>
</table>

**Theorem 5.3.** Algorithm 23 solves the mutual exclusion problem as in Definition 5.2.

**Proof.** Mutual exclusion follows directly from the test-and-set definition: Initially $R$ is 0. Let $p_i$ be the $i^{th}$ process to successfully execute the test-and-set, where successfully means that the result of the test-and-set is 0. This happens at time $t_i$. At time $t'_i$ process $p_i$ resets the shared register $R$ to 0. Between $t_i$ and $t'_i$ no other process can successfully test-and-set, hence no other process can enter the critical section concurrently.

Proving no deadlock works similar: One of the processes loitering in the entry section will successfully test-and-set as soon as the process in the critical section exited.

Since the exit section only consists of a single instruction (no potential infinite loops) we have unobstructed exit.

**Remarks:**

- No lockout, on the other hand, is not given by this algorithm. Even with only two processes there are asynchronous executions where always the same process wins the test-and-set.

- Algorithm 23 can be adapted to guarantee fairness (no lockout), essentially by ordering the processes in the entry section in a queue.
A natural question is whether one can achieve mutual exclusion with only reads and writes, that is without advanced RMW operations. The answer is yes!

Our read/write mutual exclusion algorithm is for two processes $p_0$ and $p_1$ only. In the remarks we discuss how it can be extended. The general idea is that process $p_i$ has to mark its desire to enter the critical section in a “want” register $W_i$ by setting $W_i := 1$. Only if the other process is not interested ($W_1 - i = 0$) access is granted. This however is too simple since we may run into a deadlock. This deadlock (and at the same time also lockout) is resolved by adding a priority variable $\Pi$. See Algorithm 24.

**Algorithm 24** Mutual Exclusion: Peterson’s Algorithm

**Initialization:** Shared registers $W_0, W_1, \Pi$, all initially 0.

**Code for process** $p_i$, $i = \{0, 1\}$

<**Entry**>

1: $W_i := 1$
2: $\Pi := 1 - i$
3: repeat until $\Pi = i$ or $W_1 - i = 0$

<**Critical Section**>

4: ...

<**Exit**>

5: $W_i := 0$

<**Remainder Code**>

6: ...

Remarks:

- Note that line 3 in Algorithm 24 represents a “spinlock” or “busy-wait”, similarly to the lines 1-3 in Algorithm 23.

**Theorem 5.4.** Algorithm 24 solves the mutual exclusion problem as in Definition 5.2.

**Proof.** The shared variable $\Pi$ elegantly grants priority to the process that passes line 2 first. If both processes are competing, only process $p_1$ can access the critical section because of $\Pi$. The other process $p_{1 - \Pi}$ cannot access the critical section because $W_{\Pi} = 1$ (and $\Pi \neq 1 - \Pi$). The only other reason to access the critical section is because the other process is in the remainder code (that is, not interested). This proves mutual exclusion!

No deadlock comes directly with $\Pi$: Process $p_1$ gets direct access to the critical section, no matter what the other process does.

Since the exit section only consists of a single instruction (no potential infinite loops) we have unobstructed exit.

Thanks to the shared variable $\Pi$ also no lockout (fairness) is achieved: If a process $p_i$ loses against its competitor $p_{1 - i}$ in line 2, it will have to wait until the competitor resets $W_{1 - i} := 0$ in the exit section. If process $p_i$ is unlucky it will not check $W_{1 - i} = 0$ early enough before process $p_{1 - i}$ sets $W_{1 - i} := 1$ again in line 1. However, as soon as $p_{1 - i}$ hits line 2, process $p_i$ gets the priority due to $\Pi$, and can enter the critical section. □
Remarks:
- Extending Peterson’s Algorithm to more than 2 processes can be done by a tournament tree, like in tennis. With \( n \) processes every process needs to win \( \log n \) matches before it can enter the critical section. More precisely, each process starts at the bottom level of a binary tree, and proceeds to the parent level if winning. Once winning the root of the tree it can enter the critical section. Thanks to the priority variables \( \Pi \) at each node of the binary tree, we inherit all the properties of Definition 5.2.

5.3 Store & Collect

5.3.1 Problem Definition

In this section, we will look at a second shared memory problem that has an elegant solution. Informally, the problem can be stated as follows. There are \( n \) processes \( p_1, \ldots, p_n \). Every process \( p_i \) has a read/write register \( R_i \) in the shared memory where it can store some information that is destined for the other processes. Further, there is an operation by which a process can collect (i.e., read) the values of all the processes that stored some value in their register.

We say that an operation \( op_1 \) precedes an operation \( op_2 \) iff \( op_1 \) terminates before \( op_2 \) starts. An operation \( op_2 \) follows an operation \( op_1 \) iff \( op_1 \) precedes \( op_2 \).

**Definition 5.5** (Collect). There are two operations: A store(val) by process \( p_i \) sets val to be the latest value of its register \( R_i \). A collect operation returns a view, a partial function \( V \) from the set of processes to a set of values, where \( V(p_i) \) is the latest value stored by \( p_i \) for each process \( p_i \). For a collect operation \( cop \), the following validity properties must hold for every process \( p_i \):

- If \( V(p_i) = \bot \), then no store operation by \( p_i \) precedes \( cop \).
- If \( V(p_i) = v \neq \bot \), then \( v \) is the value of a store operation \( sop \) of \( p_i \) that does not follow \( cop \), and there is no store operation by \( p_i \) that follows \( sop \) and precedes \( cop \).

Hence, a collect operation \( cop \) should not read from the future or miss a preceding store operation \( sop \).

We assume that the read/write register \( R_i \) of every process \( p_i \) is initialized to \( \bot \). We define the step complexity of an operation \( op \) to be the number of accesses to registers in the shared memory. There is a trivial solution to the collect problem as shown by Algorithm 25.

---

Algorithm 25 Collect: Simple (Non-Adaptive) Solution

**Operation** store(val) (by process \( p_i \)):

1: \( R_i := val \)

**Operation** collect:

2: for \( i := 1 \) to \( n \) do
3: \( V(p_i) := R_i \) // read register \( R_i \)
4: end for
Remarks:

- Algorithm 25 clearly works. The step complexity of every store operation is 1, the step complexity of a collect operation is $n$.
- At first sight, the step complexities of Algorithm 25 seem optimal. Because there are $n$ processes, there clearly are cases in which a collect operation needs to read all $n$ registers. However, there are also scenarios in which the step complexity of the collect operation seems very costly. Assume that there are only two processes $p_i$ and $p_j$ that have stored a value in their registers $R_i$ and $R_j$. In this case, a collect in principle only needs to read the registers $R_i$ and $R_j$ and can ignore all the other registers.
- Assume that up to a certain time $t$, $k \leq n$ processes have finished or started at least one operation. We call an operation op at time $t$ adaptive to contention if the step complexity of op only depends on $k$ and is independent of $n$.
- In the following, we will see how to implement adaptive versions of store and collect.

5.3.2 Splitters

Algorithm 26 Splitter Code

**Shared Registers:** $X : \{\bot\} \cup \{1, \ldots, n\}; \ Y : boolean$

**Initialization:** $X := \bot; \ Y := false$

**Splitter access by process $p_i$:**

1. $X := i$
2. if $Y$ then
3. return right
4. else
5. $Y := true$
6. if $X = i$ then
7. return stop
8. else
9. return left
10. end if
11. end if

To obtain adaptive collect algorithms, we need a synchronization primitive, called a splitter.

**Definition 5.6 (Splitter).** A splitter is a synchronization primitive with the following characteristic. A process entering a splitter exits with either stop, left, or right. If $k$ processes enter a splitter, at most one process exits with stop and at most $k - 1$ processes exit with left and right, respectively.

Hence, it is guaranteed that if a single process enters the splitter, then it obtains stop, and if two or more processes enter the splitter, then there is at most one process obtaining stop and there are two processes that obtain
5.3. STORE & COLLECT

Figure 5.1: A Splitter

different values (i.e., either there is exactly one stop or there is at least one left and at least one right). For an illustration, see Figure 5.1. The code implementing a splitter is given by Algorithm 26.

Lemma 5.7. Algorithm 26 correctly implements a splitter.

Proof. Assume that $k$ processes enter the splitter. Because the first process that checks whether $Y = \text{true}$ in line 2 will find that $Y = \text{false}$, not all processes return right. Next, assume that $i$ is the last process that sets $X := i$. If $i$ does not return right, it will find $X = i$ in line 6 and therefore return stop. Hence, there is always a process that does not return left. It remains to show that at most 1 process returns stop. For the sake of contradiction, assume $p_i$ and $p_j$ are two processes that return stop and assume that $p_i$ sets $X := i$ before $p_j$ sets $X := j$. Both processes need to check whether $Y$ is true before one of them sets $Y := \text{true}$. Hence, they both complete the assignment in line 1 before the first one of them checks the value of $X$ in line 6. Hence, by the time $p_i$ arrives at line 6, $X \neq i$ ($p_j$ and maybe some other processes have overwritten $X$ by then). Therefore, $p_i$ does not return stop and we get a contradiction to the assumption that both $p_i$ and $p_j$ return stop.

5.3.3 Binary Splitter Tree

Assume that we are given $2^n - 1$ splitters and that for every splitter $S$, there is an additional shared variable $Z_S : \{\perp\} \cup \{1, \ldots, n\}$ that is initialized to $\perp$ and an additional shared variable $M_S : \text{boolean}$ that is initialized to false. We call a splitter $S$ marked if $M_S = \text{true}$. The $2^n - 1$ splitters are arranged in a complete binary tree of height $n - 1$. Let $S(v)$ be the splitter associated with a node $v$ of the binary tree. The store and collect operations are given by Algorithm 27.

Theorem 5.8. Algorithm 27 correctly implements store and collect. Let $k$ be the number of participating processes. The step complexity of the first store of a process $p_i$ is $O(k)$, the step complexity of every additional store of $p_i$ is $O(1)$, and the step complexity of collect is $O(k)$.

Proof. Because at most one process can stop at a splitter, it is sufficient to show that every process stops at some splitter at depth at most $k - 1 \leq n - 1$ when invoking the first store operation to prove correctness. We prove that at most $k - i$ processes enter a subtree at depth $i$ (i.e., a subtree where the root has distance $i$ to the root of the whole tree). By definition of $k$, the number of
Algorithm 27 Adaptive Collect: Binary Tree Algorithm

Operation store(val) (by process \( p_i \)):
1: \( R_i := \text{val} \)
2: if first store operation by \( p_i \) then
3: \( v := \text{root node of binary tree} \)
4: \( \alpha := \text{result of entering splitter } S(v) \)
5: \( M_{S(v)} := \text{true} \)
6: while \( \alpha \neq \text{stop} \) do
7: if \( \alpha = \text{left} \) then
8: \( v := \text{left child of } v \)
9: else
10: \( v := \text{right child of } v \)
11: end if
12: \( \alpha := \text{result of entering splitter } S(v) \)
13: \( M_{S(v)} := \text{true} \)
14: end while
15: \( Z_{S(v)} := i \)
16: end if

Operation collect:
Traverse marked part of binary tree:
17: for all marked splitters \( S \) do
18: if \( Z_S \neq \perp \) then
19: \( i := Z_S; V(p_i) := R_i \)  // read value of process \( p_i \)
20: end if
21: end for  // \( V(p_i) = \perp \) for all other processes

Processes entering the splitter at depth 0 (i.e., at the root of the binary tree) is \( k \). For \( i > 1 \), the claim follows by induction because of the at most \( k-i \) processes entering the splitter at the root of a depth \( i \) subtree, at most \( k-i-1 \) obtain \text{left} and \text{right}, respectively. Hence, at the latest when reaching depth \( k-1 \), a process is the only process entering a splitter and thus obtains \text{stop}. It thus also follows that the step complexity of the first invocation of store is \( O(k) \).

To show that the step complexity of collect is \( O(k) \), we first observe that the marked nodes of the binary tree are connected, and therefore can be traversed by only reading the variables \( M_S \) associated to them and their neighbors. Hence, showing that at most \( 2k-1 \) nodes of the binary tree are marked is sufficient. Let \( x_k \) be the maximum number of marked nodes in a tree, where \( k \) processes access the root. We claim that \( x_k \leq 2k-1 \), which is true for \( k = 1 \) because a single process entering a splitter will always compute \text{stop}. Now assume the inequality holds for \( 1, \ldots, k-1 \). Not all \( k \) processes may exit the splitter with \text{left} (or \text{right}), i.e., \( k_l \leq k-1 \) processes will turn left and \( k_r \leq \min\{k-k_l, k-1\} \) turn right. The left and right children of the root are the roots of their subtrees, hence the induction hypothesis yields

\[
x_k = x_{k_l} + x_{k_r} + 1 \leq (2k_l - 1) + (2k_r - 1) + 1 \leq 2k - 1,
\]

concluding induction and proof.
5.3. STORE & COLLECT

Remarks:

- The step complexities of Algorithm 27 are very good. Clearly, the step complexity of the collect operation is asymptotically optimal. In order for the algorithm to work, we however need to allocate the memory for the complete binary tree of depth \( n - 1 \). The space complexity of Algorithm 27 therefore is exponential in \( n \). We will next see how to obtain a polynomial space complexity at the cost of a worse collect step complexity.

5.3.4 Splitter Matrix

Instead of arranging splitters in a binary tree, we arrange \( n^2 \) splitters in an \( n \times n \) matrix as shown in Figure 5.2. The algorithm is analogous to Algorithm 27. The matrix is entered at the top left. If a process receives left, it next visits the splitter in the next row of the same column. If a process receives right, it next visits the splitter in the next column of the same row. Clearly, the space complexity of this algorithm is \( O(n^2) \). The following theorem gives bounds on the step complexities of store and collect.

**Theorem 5.9.** Let \( k \) be the number of participating processes. The step complexity of the first store of a process \( p_i \) is \( O(k) \), the step complexity of every additional store of \( p_i \) is \( O(1) \), and the step complexity of collect is \( O(k^2) \).

**Proof.** Let the top row be row 0 and the left-most column be column 0. Let \( x_i \) be the number of processes entering a splitter in row \( i \). By induction on \( i \), we show that \( x_i \leq k - i \). Clearly, \( x_0 \leq k \). Let us therefore consider the case \( i > 0 \). Let \( j \) be the largest column such that at least one process visits the splitter in row \( i - 1 \) and column \( j \). By the properties of splitters, not all processes entering the splitter in row \( i - 1 \) and column \( j \) obtain left. Therefore, not all processes entering a splitter in row \( i - 1 \) move on to row \( i \). Because at least one processes
stays in every row, we get that \( x_i \leq k - i \). Similarly, the number of processes entering column \( j \) is at most \( k - j \). Hence, every process stops at the latest in row \( k - 1 \) and column \( k - 1 \) and the number of marked splitters is at most \( k^2 \). Thus, the step complexity of \textsc{collect} is at most \( O(k^2) \). Because the longest path in the splitter matrix is \( 2k \), the step complexity of \textsc{store} is \( O(k) \).

\[ \square \]

Remarks:

- With a slightly more complicated argument, it is possible to show that the number of processes entering the splitter in row \( i \) and column \( j \) is at most \( k - i - j \). Hence, it suffices to only allocate the upper left half (including the diagonal) of the \( n \times n \) matrix of splitters.

- The binary tree algorithm can be made space efficient by using a randomized version of a splitter. Whenever returning left or right, a randomized splitter returns left or right with probability \( 1/2 \). With high probability, it then suffices to allocate a binary tree of depth \( O(\log n) \).

- Recently, it has been shown that with a considerably more complicated deterministic algorithm, it is possible to achieve \( O(k) \) step complexity and \( O(n^2) \) space complexity.

Chapter Notes

Already in 1965 Edsger Dijkstra gave a deadlock-free solution for mutual exclusion [Dij65]. Later, Maurice Herlihy suggested consensus-numbers [Her91], where he proved the “universality of consensus”, i.e., the power of a shared memory system is determined by the consensus-number. For this work, Maurice Herlihy was awarded the Dijkstra Prize in Distributed Computing in 2003. Peterson’s Algorithm is due to [PF77, Pet81], and adaptive collect was studied in the sequence of papers [MA95, AFG02, AL05, AKP+06].

Bibliography


Chapter 6

Shared Objects

6.1 Introduction

Assume that there is a common resource (e.g. a common variable or data structure), which different nodes in a network need to access from time to time. If the nodes are allowed to change the common object when accessing it, we need to guarantee that no two nodes have access to the object at the same time. In order to achieve this mutual exclusion, we need protocols that allow the nodes of a network to store and manage access to such a shared object. A simple and obvious solution is to store the shared object at a central location (see Algorithm 28).

Algorithm 28 Shared Object: Centralized Solution

**Initialization:** Shared object stored at root node \( r \) of a spanning tree of the network graph (i.e., each node knows its parent in the spanning tree).

**Accessing Object:** (by node \( v \))

1: \( v \) sends request up the tree
2: request processed by root \( r \) (atomically)
3: result sent down the tree to node \( v \)

**Remarks:**

- Instead of a spanning tree, one can use routing.

- Algorithm 28 works, but it is not very efficient. Assume that the object is accessed by a single node \( v \) repeatedly. Then we get a high message/time complexity. Instead \( v \) could store the object, or at least cache it. But then, in case another node \( w \) accesses the object, we might run into consistency problems.

- Alternative idea: The accessing node should become the new master of the object. The shared object then becomes mobile. There exist several variants of this idea. The simplest version is a home-based solution like in Mobile IP (see Algorithm 29).
Algorithm 29 Shared Object: Home-Based Solution

Initialization: An object has a home base (a node) that is known to every node. All requests (accesses to the shared object) are routed through the home base.

Accessing Object: (by node $v$)
1: $v$ acquires a lock at the home base, receives object.

Remarks:
- Home-based solutions suffer from the triangular routing problem. If two close-by nodes access the object on a rotating basis, all the traffic is routed through the potentially far away home-base.

6.2 Arrow and Friends

We will now look at a protocol (called the Arrow algorithm) that always moves the shared object to the node currently accessing it without creating the triangular routing problem of home-based solutions. The protocol runs on a precomputed spanning tree. Assume that the spanning tree is rooted at the current position of the shared object. When a node $u$ wants to access the shared object, it sends out a find request towards the current position of the object. While searching for the object, the edges of the spanning tree are redirected such that in the end, the spanning tree is rooted at $u$ (i.e., the new holder of the object). The details of the algorithm are given by Algorithm 30. For simplicity, we assume that a node $u$ only starts a find request if $u$ is not currently the holder of the shared object and if $u$ has finished all previous find requests (i.e., it is not currently waiting to receive the object).

Remarks:
- The parent pointers in Algorithm 30 are only needed for the find operation. Sending the variable to $u$ in line 13 or to $v$.successor in line 23 is done using routing (on the spanning tree or on the underlying network).
- When we draw the parent pointers as arrows, in a quiescent moment (where no “find” is in motion), the arrows all point towards the node currently holding the variable (i.e., the tree is rooted at the node holding the variable)
- What is really great about the Arrow algorithm is that it works in a completely asynchronous and concurrent setting (i.e., there can be many find requests at the same time).

Theorem 6.1. (Arrow, Analysis) In an asynchronous and concurrent setting, a “find” operation terminates with message and time complexity $D$, where $D$ is the diameter of the spanning tree.
Algorithm 30 Shared Object: Arrow Algorithm

Initialization: As for Algorithm 28, we are given a rooted spanning tree. Each node has a pointer to its parent, the root $r$ is its own parent. The variable is initially stored at $r$. For all nodes $v$, $v$.successor := null, $v$.wait := false.

Start Find Request at Node $u$:

1: do atomically
2: $u$ sends “find by $u$” message to parent node
3: $u$.parent := $u$
4: $u$.wait := true
5: end do

Upon $w$ Receiving “Find by $u$” Message from Node $v$:

6: do atomically
7: if $w$.parent $\neq w$ then
8: $w$ sends “find by $u$” message to parent
9: $w$.parent := $v$
10: else
11: $w$.parent := $v$
12: if not $w$.wait then
13: send variable to $u$ // $w$ holds var. but does not need it any more
14: else
15: $w$.successor := $u$ // $w$ will send variable to $u$ a.s.a.p.
16: end if
17: end if
18: end do

Upon $w$ Receiving Shared Object:

19: perform operation on shared object
20: do atomically
21: $w$.wait := false
22: if $w$.successor $\neq$ null then
23: send variable to $w$.successor
24: $w$.successor := null
25: end if
26: end do
Before proving Theorem 6.1, we prove the following lemma.

**Lemma 6.2.** An edge \( \{u, v\} \) of the spanning tree is in one of four states:

1. Pointer from \( u \) to \( v \) (no message on the edge, no pointer from \( v \) to \( u \))
2. Message on the move from \( u \) to \( v \) (no pointer along the edge)
3. Pointer from \( v \) to \( u \) (no message on the edge, no pointer from \( u \) to \( v \))
4. Message on the move from \( v \) to \( u \) (no pointer along the edge)

**Proof.** W.l.o.g., assume that initially the edge \( \{u, v\} \) is in state 1. With a message arrival at \( u \) (or if \( u \) starts a “find by \( u \)” request, the edge goes to state 2. When the message is received at \( v \), \( v \) directs its pointer to \( u \) and we are therefore in state 3. A new message at \( v \) (or a new request initiated by \( v \)) then brings the edge back to state 1.

**Proof of Theorem 6.1.** Since the “find” message will only travel on a static tree, it suffices to show that it will not traverse an edge twice. Suppose for the sake of contradiction that there is a first “find” message \( f \) that traverses an edge \( e = \{u, v\} \) for the second time and assume that \( e \) is the first edge that is traversed twice by \( f \). The first time, \( f \) traverses \( e \). Assume that \( e \) is first traversed from \( u \) to \( v \). Since we are on a tree, the second time, \( e \) must be traversed from \( v \) to \( u \). Because \( e \) is the first edge to be traversed twice, \( f \) must re-visit \( e \) before visiting any other edges. Right before \( f \) reaches \( v \), the edge \( e \) is in state 2 (\( f \) is on the move) and in state 3 (it will immediately return with the pointer from \( v \) to \( u \)). This is a contradiction to Lemma 6.2.

**Remarks:**

- Finding a good tree is an interesting problem. We would like to have a tree with low stretch, low diameter, low degree, etc.
- It seems that the Arrow algorithm works especially well when lots of “find” operations are initiated concurrently. Most of them will find a “close-by” node, thus having low message/time complexity. For the sake of simplicity we analyze a synchronous system.

**Theorem 6.3.** (Arrow, Concurrent Analysis) Let the system be synchronous. Initially, the system is in a quiescent state. At time 0, a set \( S \) of nodes initiates a “find” operation. The message complexity of all “find” operations is \( O(\log |S| \cdot m^*) \) where \( m^* \) is the message complexity of an optimal (with global knowledge) algorithm on the tree.

**Proof Sketch.** Let \( d \) be the minimum distance of any node in \( S \) to the root. There will be a node \( u_1 \) at distance \( d \) from the root that reaches the root in \( d \) time steps, turning all the arrows on the path to the root towards \( u_1 \). A node \( u_2 \) that finds (is queued behind) \( u_1 \) cannot distinguish the system from a system where there was no request \( u_1 \), and instead the root was initially located at \( u_1 \). The message cost of \( u_2 \) is consequentially the distance between \( u_1 \) and \( u_2 \) on the spanning tree. By induction the total message complexity is exactly as if a collector starts at the root and then “greedily” collects tokens located at the nodes in \( S \) (greedily in the sense that the collector always goes towards the closest token). Greedy collecting the tokens is not a good strategy in general because it will traverse the same edge more than twice in the worst
An asymptotically optimal algorithm can also be translated into a depth-first-search collecting paradigm, traversing each edge at most twice. In another area of computer science, we would call the Arrow algorithm a nearest-neighbor TSP heuristic (without returning to the start/root though), and the optimal algorithm TSP-optimal. It was shown that nearest-neighbor has a logarithmic overhead, which concludes the proof.

Remarks:

- An average request set $S$ on a not-too-bad tree gives usually a much better bound. However, there is an almost tight $\log |S|/\log \log |S|$ worst-case example.

- It was recently shown that Arrow can do as good in a dynamic setting (where nodes are allowed to initiate requests at any time). In particular the message complexity of the dynamic analysis can be shown to have a $\log D$ overhead only, where $D$ is the diameter of the spanning tree (note that for logarithmic trees, the overhead becomes $\log \log n$).

- What if the spanning tree is a star? Then with Theorem 6.1, each find will terminate in 2 steps! Since also an optimal algorithm has message cost 1, the algorithm is 2-competitive...? Yes, but because of its high degree the star center experiences contention...It can be shown that the contention overhead is at most proportional to the largest degree $\Delta$ of the spanning tree.

- Thought experiment: Assume a balanced binary spanning tree—by Theorem 6.1, the message complexity per operation is $\log n$. Because a binary tree has maximum degree 3, the time per operation therefore is at most $3 \log n$.

- There are better and worse choices for the spanning tree. The stretch of an edge $\{u,v\}$ is defined as distance between $u$ and $v$ in a spanning tree. The maximum stretch of a spanning tree is the maximum stretch over all edges. A few years ago, it was shown how to construct spanning trees that are $O(\log n)$-stretch-competitive.

What if most nodes just want to read the shared object? Then it does not make sense to acquire a lock every time. Instead we can use caching (see Algorithm 31).

**Theorem 6.4.** Algorithm 31 is correct. More surprisingly, the message complexity is 3-competitive (at most a factor 3 worse than the optimum).

**Proof.** Since the accesses do not overlap by definition, it suffices to show that between two writes, we are 3-competitive. The sequence of accessing nodes is $w_0, r_1, r_2, \ldots, r_k, w_1$. After $w_0$, the object is stored at $w_0$ and not cached anywhere else. All reads cost twice the smallest subtree $T$ spanning the write $w_0$ and all the reads since each read only goes to the first copy. The write $w_1$ costs $T$ plus the path $P$ from $w_1$ to $T$. Since any data management scheme must use an edge in $T$ and $P$ at least once, and our algorithm uses edges in $T$ at most 3 times (and in $P$ at most once), the theorem follows. \qed
Algorithm 31 Shared Object: Read/Write Caching

- Nodes can either read or write the shared object. For simplicity we first assume that reads or writes do not overlap in time (access to the object is sequential).
- Nodes store three items: a parent pointer pointing to one of the neighbors (as with Arrow), and a cache bit for each edge, plus (potentially) a copy of the object.
- Initially the object is stored at a single node $u$; all the parent pointers point towards $u$, all the cache bits are false.
- When initiating a read, a message follows the arrows (this time: without inverting them!) until it reaches a cached version of the object. Then a copy of the object is cached along the path back to the initiating node, and the cache bits on the visited edges are set to true.
- A write at $u$ writes the new value locally (at node $u$), then searches (follow the parent pointers and reverse them towards $u$) a first node with a copy. Delete the copy and follow (in parallel, by flooding) all edge that have the cache flag set. Point the parent pointer towards $u$, and remove the cache flags.

Remarks:

- Concurrent reads are not a problem, also multiple concurrent reads and one write work just fine.

- What about concurrent writes? To achieve consistency writes need to invalidate the caches before writing their value. It is claimed that the strategy then becomes 4-competitive.

- Is the algorithm also time competitive? Well, not really: The optimal algorithm that we compare to is usually offline. This means it knows the whole access sequence in advance. It can then cache the object before the request even appears!

- Algorithms on trees are often simpler, but have the disadvantage that they introduce the extra stretch factor. In a ring, for example, any tree has stretch $n - 1$; so there is always a bad request pattern.
6.3 IVY AND FRIENDS

Algorithm 32 Shared Object: Pointer Forwarding

Initialization: Object is stored at root \( r \) of a precomputed spanning tree \( T \) (as in the Arrow algorithm, each node has a parent pointer pointing towards the object).

Accessing Object: (by node \( u \))
1. follow parent pointers to current root \( r \) of \( T \)
2. send object from \( r \) to \( u \)
3. \( r.\text{parent} := u; \ u.\text{parent} := u; \) \quad \// \ u \text{ is the new root}

Algorithm 33 Shared Object: Ivy

Initialization: Object is stored at root \( r \) of a precomputed spanning tree \( T \) (as before, each node has a parent pointer pointing towards the object). For simplicity, we assume that accesses to the object are sequential.

Start Find Request at Node \( u \):
1. \( u \) sends “find by \( u \)” message to parent node
2. \( u.\text{parent} := u \)

Upon \( v \) receiving “Find by \( u \)” Message:
3. if \( v.\text{parent} = v \) then
4. send object to \( u \)
5. else
6. send “find by \( u \)” message to \( v.\text{parent} \)
7. end if
8. \( v.\text{parent} := u \) \quad \// \ u \text{ will become the new root}

6.3 Ivy and Friends

In the following we study algorithms that do not restrict communication to a tree. Of particular interest is the special case of a complete graph (clique). A simple solution for this case is given by Algorithm 32.

Remarks:

- If the graph is not complete, routing can be used to find the root.

- Assume that the nodes line up in a linked list. If we always choose the first node of the linked list to acquire the object, we have message/time complexity \( n \). The new topology is again a linear linked list. Pointer forwarding is therefore bad in a worst-case.

- If edges are not FIFO, it can even happen that the number of steps is unbounded for a node having bad luck. An algorithm with such a property is named “not fair,” or “not wait-free.” (Example: Initially we have the list \( 4 \rightarrow 3 \rightarrow 2 \rightarrow 1 \); 4 starts a find; when the message of 4 passes 3, 3 itself starts a find. The message of 3 may arrive at 2 and then 1 earlier, thus the new end of the list is \( 2 \rightarrow 1 \rightarrow 3 \); once the message of 4 passes 2, the game re-starts.)

There seems to be a natural improvement of the pointer forwarding idea. Instead of simply redirecting the parent pointer from the old root to the new root, we can redirect all the parent pointers of the nodes on the path visited...
during a find message to the new root. The details are given by Algorithm 33. Figure 6.1 shows how the pointer redirecting affects a given tree (the right tree results from a find request started at node $x_0$ on the left tree).

Remarks:

- Also with Algorithm 33, we might have a bad linked list situation. However, if the start of the list acquires the object, the linked list turns into a star. As the following theorem shows, the search paths are not long on average. Since paths sometimes can be bad, we will need amortized analysis.

**Theorem 6.5.** If the initial tree is a star, a find request of Algorithm 33 needs at most $\log n$ steps on average, where $n$ is the number of processors.

**Proof.** All logarithms in the following proof are to base 2. We assume that accesses to the shared object are sequential. We use a potential function argument. Let $s(u)$ be the size of the subtree rooted at node $u$ (the number of nodes in the subtree including $u$ itself). We define the potential $\Phi$ of the whole tree $T$ as ($V$ is the set of all nodes)

$$\Phi(T) = \sum_{u \in V} \log \frac{s(u)}{2}.$$ 

Assume that the path traversed by the $i^{th}$ operation has length $k_i$, i.e., the $i^{th}$ operation redirects $k_i$ pointers to the new root. Clearly, the number of steps of the $i^{th}$ operation is proportional to $k_i$. We are interested in the cost of $m$ consecutive operations, $\sum_{i=1}^{m} k_i$.

Let $T_0$ be the initial tree and let $T_i$ be the tree after the $i^{th}$ operation. Further, let $a_i = k_i - \Phi(T_{i-1}) + \Phi(T_i)$ be the *amortized cost* of the $i^{th}$ operation. We have

$$\sum_{i=1}^{m} a_i = \sum_{i=1}^{m} (k_i - \Phi(T_{i-1}) + \Phi(T_i)) = \sum_{i=1}^{m} k_i - \Phi(T_0) + \Phi(T_m).$$

For any tree $T$, we have $\Phi(T) \geq \log(n)/2$. Because we assume that $T_0$ is a star, we also have $\Phi(T_0) = \log(n)/2$. We therefore get that

$$\sum_{i=1}^{m} a_i \geq m k_1.$$
6.3. IVY AND FRIENDS

Hence, it suffices to upper bound the amortized cost of every operation. We thus analyze the amortized cost \( a_i \) of the \( i \)th operation. Let \( x_0, x_1, x_2, \ldots, x_{k_i} \) be the path that is reversed by the operation. Further for \( 0 \leq j \leq k_i \), let \( s_j \) be the size of the subtree rooted at \( x_j \) before the reversal. The size of the subtree rooted at \( x_0 \) after the reversal is \( s_{k_i} \) and the size of the one rooted at \( x_j \) after the reversal, for \( 1 \leq j \leq k_i \), is \( s_j - s_{j-1} \) (see Figure 6.1). For all other nodes, the sizes of their subtrees are the same, therefore the corresponding terms cancel out in the amortized cost \( a_i \). We can thus write \( a_i \) as

\[
a_i = k_i - \left( \sum_{j=0}^{k_i} \frac{1}{2} \log s_j \right) + \left( \frac{1}{2} \log s_{k_i} + \sum_{j=1}^{k_i} \frac{1}{2} \log (s_j - s_{j-1}) \right)
\]

\[
= k_i + \frac{1}{2} \sum_{j=0}^{k_i-1} \left( \log (s_{j+1} - s_j) - \log s_j \right)
\]

\[
= k_i + \frac{1}{2} \sum_{j=0}^{k_i-1} \log \left( \frac{s_{j+1} - s_j}{s_j} \right).
\]

For \( 0 \leq j \leq k_i - 1 \), let \( \alpha_j = s_{j+1}/s_j \). Note that \( s_{j+1} > s_j \) and thus that \( \alpha_j > 1 \). Further note, that \( (s_{j+1} - s_j)/s_j = \alpha_j - 1 \). We therefore have that

\[
a_i = k_i + \frac{1}{2} \cdot \sum_{j=0}^{k_i-1} \log (\alpha_j - 1)
\]

\[
= \sum_{j=0}^{k_i-1} \left( 1 + \frac{1}{2} \log (\alpha_j - 1) \right).
\]

For \( \alpha > 1 \), it can be shown that \( 1 + \log(\alpha - 1)/2 \leq \log \alpha \) (see Lemma 6.6). From this inequality, we obtain

\[
a_i \leq \sum_{j=0}^{k_i-1} \log \alpha_j = \sum_{j=0}^{k_i-1} \log \left( \frac{s_{j+1}}{s_j} \right) = \sum_{j=0}^{k_i-1} (\log s_{j+1} - \log s_j)
\]

\[
= \log s_{k_i} - \log s_0 \leq \log n,
\]

because \( s_{k_i} = n \) and \( s_0 \geq 1 \). This concludes the proof. \( \square \)

**Lemma 6.6.** For \( \alpha > 1 \), \( 1 + \log(\alpha - 1)/2 \leq \log \alpha \).

**Proof.** The claim can be verified by the following chain of reasoning:

\[
0 \leq (\alpha - 2)^2
\]

\[
0 \leq \alpha^2 - 4\alpha + 4
\]

\[
4(\alpha - 1) \leq \alpha^2
\]

\[
\log_2 (4(\alpha - 1)) \leq \log_2 (\alpha^2)
\]

\[
2 + \log_2(\alpha - 1) \leq 2 \log_2 \alpha
\]

\[
1 + \frac{1}{2} \log_2(\alpha - 1) \leq \log_2 \alpha.
\]

\( \square \)
Remarks:

- Systems guys (the algorithm is called Ivy because it was used in a system with the same name) have some fancy heuristics to improve performance even more: For example, the root every now and then broadcasts its name such that paths will be shortened.

- What about concurrent requests? It works with the same argument as in Arrow. Also for Ivy an argument including congestion is missing (and more pressing, since the dynamic topology of a tree cannot be chosen to have low degree and thus low congestion as in Arrow).

- Sometimes the type of accesses allows that several accesses can be combined into one to reduce congestion higher up the tree. Let the tree in Algorithm 28 be a balanced binary tree. If the access to a shared variable for example is “add value $x$ to the shared variable”, two or more accesses that accidentally meet at a node can be combined into one. Clearly accidental meeting is rare in an asynchronous model. We might be able to use synchronizers (or maybe some other timing tricks) to help meeting a little bit.

Chapter Notes

The Arrow protocol was designed by Raymond [Ray89]. There are real life implementations of the Arrow protocol, such as the Aleph Toolkit [Her99]. The performance of the protocol under high loads was tested in [HW99] and other implementations and variations of the protocol were given in, e.g., [PR99, HTW00].

On the theory side, it was shown that the find operations of the protocol do not backtrack, i.e., the time and message complexities are in $O(D)$ [DH98], and that the Arrow protocol is fault tolerant [HT01]. Given a set of concurrent request, Herhily et al. [HTW01] showed that the time and message complexities are within factor $\log R$ from the optimal, where $R$ is the number of requests. Later, this analysis was extended to long-lived and asynchronous systems. In particular, Herhily et al. [HKTW06] showed that the competitive ratio in this asynchronous concurrent setting is $O(\log D)$. Thanks to the lower bound of the greedy TSP heuristic, this is almost tight.

The Ivy system was introduced in [Li88, LH89]. On the theory side, it was shown by Ginat et al. [GST89] that the amortized cost of a single request of the Ivy protocol is $\Theta(\log n)$. Closely related work to the Ivy protocol on the practical side is research on virtual memory and parallel computing on loosely coupled multiprocessors. For example [BB81, LSL82, FR86] contain studies on variations of the network models, limitations on data sharing between processes and different approaches.

Later, the research focus shifted towards systems where most data operations were read operations, i.e., efficient caching became one of the main objects of study, e.g., [MMVW97].
Bibliography


Chapter 7

Maximal Independent Set

In this chapter we present a highlight of this course, a fast maximal independent set (MIS) algorithm. The algorithm is the first randomized algorithm that we study in this class. In distributed computing, randomization is a powerful and therefore omnipresent concept, as it allows for relatively simple yet efficient algorithms. As such the studied algorithm is archetypal.

A MIS is a basic building block in distributed computing, some other problems pretty much follow directly from the MIS problem. At the end of this chapter, we will give two examples: matching and vertex coloring (see Chapter 1).

7.1 MIS

Definition 7.1 (Independent Set). Given an undirected Graph $G = (V,E)$ an independent set is a subset of nodes $U \subseteq V$, such that no two nodes in $U$ are adjacent. An independent set is maximal if no node can be added without violating independence. An independent set of maximum cardinality is called maximum.

![Figure 7.1: Example graph with 1) a maximal independent set (MIS) and 2) a maximum independent set (MaxIS).](image)
Remarks:

- Computing a maximum independent set (MaxIS) is a notoriously difficult problem. It is equivalent to maximum clique on the complementary graph. Both problems are NP-hard, in fact not approximable within $n^{\frac{1}{2}-\epsilon}$.

- In this course we concentrate on the maximal independent set (MIS) problem. Please note that MIS and MaxIS can be quite different, indeed e.g. on a star graph the MIS is $\Theta(n)$ smaller than the MaxIS (cf. Figure 7.1).

- Computing a MIS sequentially is trivial: Scan the nodes in arbitrary order. If a node $u$ does not violate independence, add $u$ to the MIS. If $u$ violates independence, discard $u$. So the only question is how to compute a MIS in a distributed way.

Algorithm 34 Slow MIS

Require: Node IDs

Every node $v$ executes the following code:

1: if all neighbors of $v$ with larger identifiers have decided not to join the MIS then
2: $v$ decides to join the MIS
3: end if

Remarks:

- Not surprisingly the slow algorithm is not better than the sequential algorithm in the worst case, because there might be one single point of activity at any time. Formally:

**Theorem 7.2** (Analysis of Algorithm 34). Algorithm 34 features a time complexity of $O(n)$ and a message complexity of $O(m)$.

Remarks:

- This is not very exciting.

- There is a relation between independent sets and node coloring (Chapter 1), since each color class is an independent set, however, not necessarily a MIS. Still, starting with a coloring, one can easily derive a MIS algorithm: We first choose all nodes of the first color. Then, for each additional color we add “in parallel” (without conflict) as many nodes as possible. Thus the following corollary holds:

**Corollary 7.3.** Given a coloring algorithm that needs $C$ colors and runs in time $T$, we can construct a MIS in time $C + T$.

Remarks:

- Using Theorem 1.17 and Corollary 7.3 we get a distributed deterministic MIS algorithm for trees (and for bounded degree graphs) with time complexity $O(\log^* n)$. 
With a lower bound argument one can show that this deterministic MIS algorithm for rings is asymptotically optimal.

There have been attempts to extend Algorithm 5 to more general graphs, however, so far without much success. Below we present a radically different approach that uses randomization. Please note that the algorithm and the analysis below is not identical with the algorithm in Peleg’s book.

7.2 Original Fast MIS

Algorithm 35 Fast MIS

The algorithm operates in synchronous rounds, grouped into phases. A single phase is as follows:

1) Each node $v$ marks itself with probability $\frac{1}{2d(v)}$, where $d(v)$ is the current degree of $v$.

2) If no higher degree neighbor of $v$ is also marked, node $v$ joins the MIS. If a higher degree neighbor of $v$ is marked, node $v$ unmarks itself again. (If the neighbors have the same degree, ties are broken arbitrarily, e.g., by identifier).

3) Delete all nodes that joined the MIS and their neighbors, as they cannot join the MIS anymore.

Remarks:

- Correctness in the sense that the algorithm produces an independent set is relatively simple: Steps 1 and 2 make sure that if a node $v$ joins the MIS, then $v$’s neighbors do not join the MIS at the same time. Step 3 makes sure that $v$’s neighbors will never join the MIS.

- Likewise the algorithm eventually produces a MIS, because the node with the highest degree will mark itself at some point in Step 1.

- So the only remaining question is how fast the algorithm terminates. To understand this, we need to dig a bit deeper.

Lemma 7.4 (Joining MIS). A node $v$ joins the MIS in step 2 with probability $p \geq \frac{1}{4d(v)}$.

Proof: Let $M$ be the set of marked nodes in step 1. Let $H(v)$ be the set of neighbors of $v$ with higher degree, or same degree and higher identifier. Using independence of the random choices of $v$ and nodes in $H(v)$ in Step 1 we get

$$P[v \notin \text{MIS}|v \in M] = P[\exists w \in H(v), w \in M|v \in M] = P[\exists w \in H(v), w \in M]$$

$$\leq \sum_{w \in H(v)} P[w \in M] = \sum_{w \in H(v)} \frac{1}{2d(w)}$$

$$\leq \sum_{w \in H(v)} \frac{1}{2d(v)} \leq \frac{d(v)}{2d(v)} = \frac{1}{2}.$$
Then
\[
P [\{\pi \in V \mid i \in \text{MIS}\}] = P [\pi \in \text{MIS}|\{\pi \in M\}] \cdot P [\pi \in M] \geq \frac{1}{2} \cdot \frac{1}{2d(i)}.
\]

\[\mathbf{Lemma 7.5} \text{ (Good Nodes). A node } v \text{ is called good if }
\[
\sum_{w \in N(v)} \frac{1}{2d(w)} \geq \frac{1}{6}.
\]
Otherwise we call } v \text{ a bad node. A good node will be removed in Step 3 with probability } p \geq \frac{1}{36}.

Proof: Let node } v \text{ be good. Intuitively, good nodes have lots of low-degree neighbors, thus chances are high that one of them goes into the independent set, in which case } v \text{ will be removed in step 3 of the algorithm.

If there is a neighbor } w \in N(v) \text{ with degree at most 2 we are done: With Lemma 7.4 the probability that node } w \text{ joins the MIS is at least } \frac{1}{8}, \text{ and our good node will be removed in Step 3.}

So all we need to worry about is that all neighbors have at least degree 3: For any neighbor } w \text{ of } v \text{ we have } \frac{1}{2d(w)} \leq \frac{1}{6}. \text{ Since } \sum_{w \in N(v)} \frac{1}{2d(w)} \geq \frac{1}{6} \text{ there is a subset of neighbors } S \subseteq N(v) \text{ such that } \frac{1}{6} \leq \sum_{w \in S} \frac{1}{2d(w)} \leq \frac{1}{3}.

We can now bound the probability that node } v \text{ will be removed. Let therefore } R \text{ be the event of } v \text{ being removed. Again, if a neighbor of } v \text{ joins the MIS in Step 2, node } v \text{ will be removed in Step 3. We have }

\[
P [R] \geq P [\exists u \in S, u \in \text{MIS}]
\geq \sum_{u \in S} P [u \in \text{MIS}] - \sum_{u, w \in S; u \neq w} P [u \in \text{MIS and } w \in \text{MIS}].
\]

For the last inequality we used the inclusion-exclusion principle truncated after the second order terms. Let } M \text{ again be the set of marked nodes after Step 1. Using } P [u \in M] \geq P [u \in \text{MIS}] \text{ we get }

\[
P [R] \geq \sum_{u \in S} P [u \in \text{MIS}] - \sum_{u, w \in S; u \neq w} P [u \in M \text{ and } w \in M]
\geq \sum_{u \in S} P [u \in \text{MIS}] - \sum_{u \in S} \sum_{w \in S} P [u \in M] \cdot P [w \in M]
\geq \sum_{u \in S} \frac{1}{4d(u)} - \sum_{u \in S} \sum_{w \in S} \frac{1}{2d(u)} \cdot \frac{1}{2d(w)}
\geq \sum_{u \in S} \frac{1}{2d(u)} \left( \frac{1}{2} - \sum_{w \in S} \frac{1}{2d(w)} \right) \geq \frac{1}{6} \left( \frac{1}{2} - \frac{1}{3} \right) = \frac{1}{36}.
\]

\[\Box\]
Remarks:

• We would be almost finished if we could prove that many nodes are good in each phase. Unfortunately this is not the case: In a star-graph, for instance, only a single node is good! We need to find a work-around.

Lemma 7.6 (Good Edges). An edge \( e = (u, v) \) is called bad if both \( u \) and \( v \) are bad; else the edge is called good. The following holds: At any time at least half of the edges are good.

Proof: For the proof we construct a directed auxiliary graph: Direct each edge towards the higher degree node (if both nodes have the same degree direct it towards the higher identifier). Now we need a little helper lemma before we can continue with the proof.

Lemma 7.7. A bad node has outdegree (number of edges pointing away from bad node) at least twice its indegree (number of edges pointing towards bad node).

Proof: For the sake of contradiction, assume that a bad node \( v \) does not have outdegree at least twice its indegree. In other words, at least one third of the neighbor nodes (let’s call them \( S \)) have degree at most \( d(v) \). But then

\[
\sum_{w \in N(v)} \frac{1}{2d(w)} \geq \sum_{w \in S} \frac{1}{2d(w)} \geq \sum_{w \in S} \frac{1}{2d(v)} \geq \frac{d(v)}{3} \cdot \frac{1}{2d(v)} = \frac{1}{6}
\]

which means \( v \) is good, a contradiction. \( \square \)

Continuing the proof of Lemma 7.6: According to Lemma 7.7 the number of edges directed into bad nodes is at most half the number of edges directed out of bad nodes. Thus, the number of edges directed into bad nodes is at most half the number of edges. Thus, at least half of the edges are directed into good nodes. Since these edges are not bad, they must be good.

Theorem 7.8 (Analysis of Algorithm 35). Algorithm 35 terminates in expected time \( O(\log n) \).

Proof: With Lemma 7.5 a good node (and therefore a good edge!) will be deleted with constant probability. Since at least half of the edges are good (Lemma 7.6) a constant fraction of edges will be deleted in each phase.

More formally: With Lemmas 7.5 and 7.6 we know that at least half of the edges will be removed with probability at least \( 1/36 \). Let \( R \) be the number of edges to be removed. Using linearity of expectation we know that \( \mathbb{E}[R] \geq m/72 \), \( m \) being the total number of edges at the start of a phase. Now let \( p := P[R \leq \mathbb{E}[R]/2] \). Bounding the expectation yields

\[
\mathbb{E}[R] = \sum_{r} P[R = r] \cdot r \leq p \cdot \mathbb{E}[R]/2 + (1 - p) \cdot m.
\]

Solving for \( p \) we get

\[
p \leq \frac{m - \mathbb{E}[R]}{m - \mathbb{E}[R]/2} \leq \frac{m - \mathbb{E}[R]/2}{m} \leq 1 - 1/144.
\]

In other words, with probability at least \( 1/144 \) at least \( m/144 \) edges are removed in a phase. After expected \( O(\log m) \) phases all edges are deleted. Since \( m \leq n^2 \) and thus \( O(\log m) = O(\log n) \) the Theorem follows. \( \square \)
Remarks:

• With a bit of more math one can even show that Algorithm 35 terminates in time $O(\log n)$ "with high probability".

7.3 Fast MIS v2

Algorithm 36 Fast MIS 2

The algorithm operates in synchronous rounds, grouped into phases.

A single phase is as follows:
1) Each node $v$ chooses a random value $r(v) \in [0, 1]$ and sends it to its neighbors.
2) If $r(v) < r(w)$ for all neighbors $w \in N(v)$, node $v$ enters the MIS and informs its neighbors.
3) If $v$ or a neighbor of $v$ entered the MIS, $v$ terminates ($v$ and all edges adjacent to $v$ are removed from the graph), otherwise $v$ enters the next phase.

Remarks:

• Correctness in the sense that the algorithm produces an independent set is simple: Steps 1 and 2 make sure that if a node $v$ joins the MIS, then $v$’s neighbors do not join the MIS at the same time. Step 3 makes sure that $v$’s neighbors will never join the MIS.

• Likewise the algorithm eventually produces a MIS, because the node with the globally smallest value will always join the MIS, hence there is progress.

• So the only remaining question is how fast the algorithm terminates. To understand this, we need to dig a bit deeper.

• Our proof will rest on a simple, yet powerful observation about expected values of random variables that may not be independent:

Theorem 7.9 (Linearity of Expectation). Let $X_i, i = 1, \ldots, k$ denote random variables, then

$$\mathbb{E} \left[ \sum_i X_i \right] = \sum_i \mathbb{E} [X_i].$$

Proof. It is sufficient to prove $\mathbb{E} [X + Y] = \mathbb{E} [X] + \mathbb{E} [Y]$ for two random variables $X$ and $Y$, because then the statement follows by induction. Since

$$P [(X, Y) = (x, y)] = P [X = x] \cdot P [Y = y | X = x] = P [Y = y] \cdot P [X = x | Y = y]$$
we get that

\[ E[X + Y] = \sum_{(X,Y) = (x,y)} P[(X,Y) = (x,y)] \cdot (x + y) \]

\[ = \sum_{X=x} \sum_{Y=y} P[X = x] \cdot P[Y = y | X = x] \cdot x \]

\[ + \sum_{Y=y} \sum_{X=x} P[Y = y] \cdot P[X = x | Y = y] \cdot y \]

\[ = \sum_{X=x} P[X = x] \cdot x + \sum_{Y=y} P[Y = y] \cdot y \]

\[ = E[X] + E[Y]. \]

\[ \square \]

Remarks:

- How can we prove that the algorithm only needs \( O(\log n) \) phases in expectation? It would be great if this algorithm managed to remove a constant fraction of nodes in each phase. Unfortunately, it does not.

- Instead we will prove that the number of edges decreases quickly. Again, it would be great if any single edge was removed with constant probability in Step 3. But again, unfortunately, this is not the case.

- Maybe we can argue about the expected number of edges to be removed in one single phase? Let’s see: A node \( v \) enters the MIS with probability \( 1/(d(v) + 1) \), where \( d(v) \) is the degree of node \( v \). By doing so, not only are \( v \)'s edges removed, but indeed all the edges of \( v \)'s neighbors as well – generally these are much more than \( d(v) \) edges. So there is hope, but we need to be careful: If we do this the most naïve way, we will count the same edge many times.

- How can we fix this? The nice observation is that it is enough to count just some of the removed edges. Given a new MIS node \( v \) and a neighbor \( w \in N(v) \), we count the edges only if \( r(v) < r(w) \) for all neighbors \( x \) of a neighbor \( w \) of \( v \). The probability of event \( (v \to w) \) is at least \( 1/(d(v) + d(w)) \), since \( d(v) + d(w) \) is the maximum number of nodes adjacent to \( v \) or \( w \) (or both). As \( v \) joins the MIS, all (directed) edges \((w,x)\) with \( x \in N(w) \) will be removed; there are \( d(w) \) of these edges.

We now count the removed edges. Whether we remove the edges adjacent to \( w \) because of event \( (v \to w) \) is a random variable \( X_{(v \to w)} \). If event \( (v \to w) \)
occurs, $X_{(v \rightarrow w)}$ has the value $d(w)$, if not it has the value 0. For each undirected edge $\{v, w\}$ we have two such variables, $X_{(v \rightarrow w)}$ and $X_{(w \rightarrow v)}$. Due to Theorem 7.9, the expected value of the sum $X$ of all these random variables is at least

$$E[X] = \sum_{\{v, w\} \in E} E[X_{(v \rightarrow w)}] + E[X_{(w \rightarrow v)}]$$

$$= \sum_{\{v, w\} \in E} P[\text{Event } (v \rightarrow w)] \cdot d(w) + P[\text{Event } (w \rightarrow v)] \cdot d(v)$$

$$\geq \sum_{\{v, w\} \in E} \frac{d(w)}{d(v) + d(w)} + \frac{d(v)}{d(w) + d(v)}$$

$$= \sum_{\{v, w\} \in E} 1 = |E|.$$

In other words, in expectation $|E|$ directed edges are removed in a single phase! Note that we did not double count any edge removals, as a directed edge $(v, w)$ can only be removed by an event $(u \rightarrow v)$. The event $(u \rightarrow v)$ inhibits a concurrent event $(u' \rightarrow v)$ since $r(u) < r(u')$ for all $u' \in N(v)$. We may have counted an undirected edge at most twice (once in each direction). So, in expectation at least half of the undirected edges are removed. □

Remarks:
- This enables us to follow a bound on the expected running time of Algorithm 36 quite easily.

**Theorem 7.11 (Expected running time of Algorithm 36).** Algorithm 36 terminates after at most $3 \log_{4/3} m + 1 \in O(\log n)$ phases in expectation.

Proof: The probability that in a single phase at least a quarter of all edges are removed is at least $1/3$. For the sake of contradiction, assume not. Then with probability less than $1/3$ we may be lucky and many (potentially all) edges are removed. With probability more than $2/3$ less than $1/4$ of the edges are removed. Hence the expected fraction of removed edges is strictly less than $1/3 \cdot 1 + 2/3 \cdot 1/4 = 1/2$. This contradicts Lemma 7.10.

Hence, at least every third phase is “good” and removes at least a quarter of the edges. To get rid of all but two edges we need $\log_{4/3} m$ good phases in expectation. The last two edges will certainly be removed in the next phase. Hence a total of $3 \log_{4/3} m + 1$ phases are enough in expectation.

Remarks:
- Sometimes one expects a bit more of an algorithm: Not only should the expected time to terminate be good, but the algorithm should always terminate quickly. As this is impossible in randomized algorithms (after all, the random choices may be “unlucky” all the time!), researchers often settle for a compromise, and just demand that the probability that the algorithm does not terminate in the specified time can be made absurdly small. For our algorithm, this can be deduced from Lemma 7.10 and another standard tool, namely Chernoff’s Bound.

**Definition 7.12 (W.h.p.).** We say that an algorithm terminates w.h.p. (with high probability) within $O(t)$ time if it does so with probability at least $1 - 1/n^c$.
for any choice of $c \geq 1$. Here $c$ may affect the constants in the Big-O notation because it is considered a “tunable constant” and usually kept small.

**Definition 7.13 (Chernoff’s Bound).** Let $X = \sum_{i=1}^{k} X_i$ be the sum of $k$ independent $0 – 1$ random variables. Then Chernoff’s bound states that w.h.p.

$$|X - \mathbb{E}[X]| \leq O\left(\log n + \sqrt{\mathbb{E}[X] \log n}\right).$$

**Corollary 7.14 (Running Time of Algorithm 36).** Algorithm 36 terminates w.h.p. in $O(\log n)$ time.

Proof: In Theorem 7.11 we used that independently of everything that happened before, in each phase we have a constant probability $p$ that a quarter of the edges are removed. Call such a phase good. For some constants $C_1$ and $C_2$, let us check after $C_1 \log n + C_2 \in O(\log n)$ phases, in how many phases at least a quarter of the edges have been removed. In expectation, these are at least $p(C_1 \log n + C_2)$ many. Now we look at the random variable $X = \sum_{i=1}^{C_1 \log n + C_2} X_i$, where the $X_i$ are independent $0 – 1$ variables being one with exactly probability $p$. Certainly, if $X$ is at least $x$ with some probability, then the probability that we have $x$ good phases can only be larger (if no edges are left, certainly “all” of the remaining edges are removed). To $X$ we can apply Chernoff’s bound. If $C_1$ and $C_2$ are chosen large enough, they will overcome the constants in the Big-O from Chernoff’s bound, i.e., w.h.p. it holds that $|X - \mathbb{E}[X]| \leq \mathbb{E}[X]/2$, implying $X \geq \mathbb{E}[X]/2$. Choosing $C_1$ large enough, we will have w.h.p. sufficiently many good phases, i.e., the algorithm terminates w.h.p. in $O(\log n)$ phases.

**Remarks:**

- The algorithm can be improved a bit more even. Drawing random real numbers in each phase for instance is not necessary. One can achieve the same by sending only a total of $O(\log n)$ random (and as many non-random) bits over each edge.

- One of the main open problems in distributed computing is whether one can beat this logarithmic time, or at least achieve it with a deterministic algorithm.

- Let’s turn our attention to applications of MIS next.

### 7.4 Applications

**Definition 7.15 (Matching).** Given a graph $G = (V, E)$ a matching is a subset of edges $M \subseteq E$, such that no two edges in $M$ are adjacent (i.e., where no node is adjacent to two edges in the matching). A matching is maximal if no edge can be added without violating the above constraint. A matching of maximum cardinality is called maximum. A matching is called perfect if each node is adjacent to an edge in the matching.
Remarks:

- In contrast to MaxIS, a maximum matching can be found in polynomial time (Blossom algorithm by Jack Edmonds), and is also easy to approximate (in fact, already any maximal matching is a 2-approximation).

- An independent set algorithm is also a matching algorithm: Let $G = (V,E)$ be the graph for which we want to construct the matching. The auxiliary graph $G'$ is defined as follows: for every edge in $G$ there is a node in $G'$; two nodes in $G'$ are connected by an edge if their respective edges in $G$ are adjacent. A (maximal) independent set in $G'$ is a (maximal) matching in $G$, and vice versa. Using Algorithm 36 directly produces an $O(\log n)$ bound for maximal matching.

- More importantly, our MIS algorithm can also be used for vertex coloring (Problem 1.1):

**Definition 7.16.** An approximation algorithm $A$ for a maximization problem $\Pi$ has an approximation factor of $r$ if the following condition holds for all instances $I \in \Pi$:

$$\frac{OPT(I)}{A(I)} \leq r.$$ 

**Algorithm 37** General Graph Coloring

1. Given a graph $G = (V,E)$ we virtually build a graph $G' = (V',E')$ as follows:
2. Every node $v \in V$ clones itself $d(v) + 1$ times ($v_0, \ldots, v_{d(v)} \in V'$), $d(v)$ being the degree of $v$ in $G$.
3. The edge set $E'$ of $G'$ is as follows:
4. First all clones are in a clique: $(v_i, v_j) \in E'$, for all $v \in V$ and all $0 \leq i < j \leq d(v)$
5. Second all $i^{th}$ clones of neighbors in the original graph $G$ are connected: $(u_i, v_j) \in E'$, for all $(u,v) \in E$ and all $0 \leq i \leq \min(d(u), d(v))$.
6. Now we simply run (simulate) the fast MIS Algorithm 36 on $G'$.
7. If node $v_i$ is in the MIS in $G'$, then node $v$ gets color $i$.

**Theorem 7.17** (Analysis of Algorithm 37). Algorithm 37 $(\Delta + 1)$-colors an arbitrary graph in $O(\log n)$ time, with high probability, $\Delta$ being the largest degree in the graph.

Proof: Thanks to the clique among the clones at most one clone is in the MIS. And because of the $d(v) + 1$ clones of node $v$ every node will get a free color! The running time remains logarithmic since $G'$ has $O(n^2)$ nodes and the exponent becomes a constant factor when applying the logarithm.

Remarks:

- This solves our open problem from Chapter 1.1!

- Together with Corollary 7.3 we get quite close ties between $(\Delta+1)$-coloring and the MIS problem.
7.4. APPLICATIONS

- Computing a MIS also solves another graph problem on graphs of bounded independence.

**Definition 7.18** (Bounded Independence). A graph \( G = (V, E) \) is of bounded independence, if each neighborhood contains at most a constant number of independent (i.e., mutually non-adjacent) nodes.

**Definition 7.19** ((Minimum) Dominating Sets). A dominating set is a subset of the nodes such that each node is in the set or adjacent to a node in the set. A minimum dominating set is a dominating set containing the least possible number of nodes.

**Remarks:**

- In general, finding a dominating set less than factor \( \log n \) larger than an minimum dominating set is NP-hard.
- Any MIS is a dominating set: if a node was not covered, it could join the independent set.
- In general a MIS and a minimum dominating sets have not much in common (think of a star). For graphs of bounded independence, this is different.

**Corollary 7.20.** On graphs of bounded independence, a constant-factor approximation to a minimum dominating set can be found in time \( O(\log n) \) w.h.p.

Proof: Denote by \( M \) a minimum dominating set and by \( I \) a MIS. Since \( M \) is a dominating set, each node from \( I \) is in \( M \) or adjacent to a node in \( M \). Since the graph is of bounded independence, no node in \( M \) is adjacent to more than constantly many nodes from \( I \). Thus, \( |I| \in O(|M|) \). Therefore, we can compute a MIS with Algorithm 36 and output it as the dominating set, which takes \( O(\log n) \) rounds w.h.p.

**Chapter Notes**

The fast MIS algorithm is a simplified version of an algorithm by Luby [Lub86]. Around the same time there have been a number of other papers dealing with the same or related problems, for instance by Alon, Babai, and Itai [ABI86], or by Israeli and Itai [II86]. The analysis presented in Section 7.2 takes elements of all these papers, and from other papers on distributed weighted matching [WW04]. The analysis in the book by David Peleg [Pel00] is different, and only achieves \( O(\log^2 n) \) time. The new MIS variant (with the simpler analysis) of Section 7.3 is by Métivier, Robson, Saheb-Djahromi and Zemmari [MRSDZ11]. With some adaptions, the algorithms [Lub86, MRSDZ11] only need to exchange a total of \( O(\log n) \) bits per node, which is asymptotically optimum, even on unoriented trees [KSOS06]. However, the distributed time complexity for MIS is still somewhat open, as the strongest lower bounds are \( \Omega(\sqrt{\log n}) \) or \( \Omega(\log \Delta) \) [KMW04]. Recent research regarding the MIS problem focussed on improving the \( O(\log n) \) time complexity for special graph classes, for instances growth-bounded graphs [SW08] or trees [LW11]. There are also results that depend on the degree of the graph [BE09, Kuh09]. Deterministic MIS algorithms are
still far from the lower bounds, as the best deterministic MIS algorithm takes $2^{O(\sqrt{\log n})}$ time [PS96].

Bibliography


CHAPTER 7. MAXIMAL INDEPENDENT SET
Chapter 8

Locality Lower Bounds

In Chapter 1, we looked at distributed algorithms for coloring. In particular, we saw that rings and rooted trees can be colored with 3 colors in $\log^* n + O(1)$ rounds. In this chapter, we will reconsider the distributed coloring problem. We will look at a classic lower bound that shows that the result of Chapter 1 is tight: Coloring rings (and rooted trees) indeed requires $\Omega(\log^* n)$ rounds. In particular, we will prove a lower bound for coloring in the following setting:

- We consider deterministic, synchronous algorithms.
- Message size and local computations are unbounded.
- We assume that the network is a directed ring with $n$ nodes.
- Nodes have unique labels (identifiers) from 1 to $n$.

Remarks:

- A generalization of the lower bound to randomized algorithms is possible.
- Except for restricting to deterministic algorithms, all the conditions above make a lower bound stronger: Any lower bound for synchronous algorithms certainly also holds for asynchronous ones. A lower bound that is true if message size and local computations are not restricted is clearly also valid if we require a bound on the maximal message size or the amount of local computations. Similarly also assuming that the ring is directed and that node labels are from 1 to $n$ (instead of choosing IDs from a more general domain) strengthen the lower bound.
- Instead of directly proving that 3-coloring a ring needs $\Omega(\log^* n)$ rounds, we will prove a slightly more general statement. We will consider deterministic algorithms with time complexity $r$ (for arbitrary $r$) and derive a lower bound on the number of colors that are needed if we want to properly color an $n$-node ring with an $r$-round algorithm. A 3-coloring lower bound can then be derived by taking the smallest $r$ for which an $r$-round algorithm needs 3 or fewer colors.
8.1 Locality

Let us for a moment look at distributed algorithms more generally (i.e., not only at coloring and not only at rings). Assume that initially, all nodes only know their own label (identifier) and potentially some additional input. As information needs at least \( r \) rounds to travel \( r \) hops, after \( r \) rounds, a node \( v \) can only learn about other nodes at distance at most \( r \). If message size and local computations are not restricted, it is in fact not hard to see, that in \( r \) rounds, a node \( v \) can exactly learn all the node labels and inputs up to distance \( r \).

As shown by the following lemma, this allows to transform every deterministic \( r \)-round synchronous algorithm into a simple canonical form.

**Lemma 8.1.** If message size and local computations are not bounded, every deterministic, synchronous \( r \)-round algorithm can be transformed into an algorithm of the form given by Algorithm 38 (i.e., it is possible to first communicate for \( r \) rounds and then do all the computations in the end).

**Proof.** Consider some \( r \)-round algorithm \( A \). We want to show that \( A \) can be brought to the canonical form given by Algorithm 38. First, we let the nodes communicate for \( r \) rounds. Assume that in every round, every node sends its complete state to all of its neighbors (remember that there is no restriction on the maximal message size). By induction, after \( i \) rounds, every node knows the initial state of all other nodes at distance at most \( i \). Hence, after \( r \) rounds, a node \( v \) has the combined initial knowledge of all the nodes in its \( r \)-neighborhood.

We want to show that this suffices to locally (at node \( v \)) simulate enough of Algorithm \( A \) to compute all the messages that \( v \) receives in the \( r \) communication rounds of a regular execution of Algorithm \( A \).

Concretely, we prove the following statement by induction on \( i \). For all nodes at distance at most \( r - i + 1 \) from \( v \), node \( v \) can compute all messages of the first \( i \) rounds of a regular execution of \( A \). Note that this implies that \( v \) can compute all the messages it receives from its neighbors during all \( r \) rounds. Because \( v \) knows the initial state of all nodes in the \( r \)-neighborhood, \( v \) can clearly compute all messages of the first round (i.e., the statement is true for \( i = 1 \)). Let us now consider the induction step from \( i \) to \( i + 1 \). By the induction hypothesis, \( v \) can compute the messages of the first \( i \) rounds of all nodes in its \((r - i + 1)\)-neighborhood. It can therefore compute all messages that are received by nodes in the \((r - i)\)-neighborhood in the first \( i \) rounds. This is of course exactly what is needed to compute the messages of round \( i + 1 \) of nodes in the \((r - i)\)-neighborhood.
Remarks:

- It is straightforward to generalize the canonical form to randomized algorithms: Every node first computes all the random bits it needs throughout the algorithm. The random bits are then part of the initial state of a node.

**Definition 8.2** (r-hop view). We call the collection of the initial states of all nodes in the r-neighborhood of a node \( v \), the r-hop view of \( v \).

Remarks:

- Assume that initially, every node knows its degree, its label (identifier) and potentially some additional input. The r-hop view of a node \( v \) then includes the complete topology of the r-neighborhood (excluding edges between nodes at distance \( r \)) and the labels and additional inputs of all nodes in the r-neighborhood.

Based on the definition of an r-hop view, we can state the following corollary of Lemma 8.1.

**Corollary 8.3.** A deterministic r-round algorithm \( A \) is a function that maps every possible r-hop view to the set of possible outputs.

**Proof.** By Lemma 8.1, we know that we can transform Algorithm \( A \) to the canonical form given by Algorithm 38. After \( r \) communication rounds, every node \( v \) knows exactly its r-hop view. This information suffices to compute the output of node \( v \).

**Remarks:**

- Note that the above corollary implies that two nodes with equal r-hop views have to compute the same output in every r-round algorithm.

- For coloring algorithms, the only input of a node \( v \) is its label. The r-hop view of a node therefore is its labeled r-neighborhood.

- If we only consider rings, r-hop neighborhoods are particularly simple. The labeled r-neighborhood of a node \( v \) (and hence its r-hop view) in an oriented ring is simply a \((2r + 1)\)-tuple \((\ell_{-r}, \ell_{-r+1}, \ldots, \ell_0, \ldots, \ell_r)\) of distinct node labels where \( \ell_0 \) is the label of \( v \). Assume that for \( i > 0 \), \( \ell_i \) is the label of the \( i \)th clockwise neighbor of \( v \) and \( \ell_{-i} \) is the label of the \( i \)th counterclockwise neighbor of \( v \). A deterministic coloring algorithm for oriented rings therefore is a function that maps \((2r + 1)\)-tuples of node labels to colors.

- Consider two r-hop views \( V_r = (\ell_{-r}, \ldots, \ell_r) \) and \( V'_r = (\ell'_{-r}, \ldots, \ell'_r) \). If \( \ell'_{i} = \ell_{i+1} \) for \(-r \leq i \leq r - 1\) and if \( \ell'_{r} \neq \ell_{i} \) for \(-r \leq i \leq r\), the r-hop view \( V'_r \) can be the r-hop view of a clockwise neighbor of a node with r-hop view \( V_r \). Therefore, every algorithm \( A \) that computes a valid coloring needs to assign different colors to \( V_r \) and \( V'_r \). Otherwise, there is a ring labeling for which \( A \) assigns the same color to two adjacent nodes.
8.2 The Neighborhood Graph

We will now make the above observations concerning colorings of rings a bit more formal. Instead of thinking of an $r$-round coloring algorithm as a function from all possible $r$-hop views to colors, we will use a slightly different perspective. Interestingly, the problem of understanding distributed coloring algorithms can itself be seen as a classical graph coloring problem.

**Definition 8.4** (Neighborhood Graph). For a given family of network graphs $\mathcal{G}$, the $r$-neighborhood graph $N_r(\mathcal{G})$ is defined as follows. The node set of $N_r(\mathcal{G})$ is the set of all possible labeled $r$-neighborhoods (i.e., all possible $r$-hop views). There is an edge between two labeled $r$-neighborhoods $V_r$ and $V'_r$ if $V_r$ and $V'_r$ can be the $r$-hop views of two adjacent nodes.

**Lemma 8.5.** For a given family of network graphs $\mathcal{G}$, there is an $r$-round algorithm that colors graphs of $\mathcal{G}$ with $c$ colors iff the chromatic number of the neighborhood graph is $\chi(N_r(\mathcal{G})) \leq c$.

**Proof.** We have seen that a coloring algorithm is a function that maps every possible $r$-hop view to a color. Hence, a coloring algorithm assigns a color to every node of the neighborhood graph $N_r(\mathcal{G})$. If two $r$-hop views $V_r$ and $V'_r$ can be the $r$-hop views of two adjacent nodes $u$ and $v$ (for some labeled graph in $\mathcal{G}$), every correct coloring algorithm must assign different colors to $V_r$ and $V'_r$. Thus, specifying an $r$-round coloring algorithm for a family of network graphs $\mathcal{G}$ is equivalent to coloring the respective neighborhood graph $N_r(\mathcal{G})$. □

Instead of directly defining the neighborhood graph for directed rings, we define directed graphs $B_{k,n}$ that are closely related to the neighborhood graph.

Let $k$ and $n$ be two positive integers and assume that $n \geq k$. The node set of $B_{k,n}$ contains all $k$-tuples of increasing node labels ($\mathbb{N} = \{1, \ldots, n\}$):

$$V[B_{k,n}] = \{(\alpha_1, \ldots, \alpha_k) : \alpha_i \in \mathbb{N}, i < j \rightarrow \alpha_i < \alpha_j\} \quad (8.1)$$

For $\alpha = (\alpha_1, \ldots, \alpha_k)$ and $\beta = (\beta_1, \ldots, \beta_k)$ there is a directed edge from $\alpha$ to $\beta$ iff

$$\forall i \in \{1, \ldots, k-1\} : \beta_i = \alpha_{i+1}. \quad (8.2)$$

**Lemma 8.6.** Viewed as an undirected graph, the graph $B_{2r+1,n}$ is a subgraph of the $r$-neighborhood graph of directed $n$-node rings with node labels from $\mathbb{N}$.

**Proof.** The claim follows directly from the observations regarding $r$-hop views of nodes in a directed ring from Section 8.1. The set of $k$-tuples of increasing node labels is a subset of the set of $k$-tuples of distinct node labels. Two nodes of $B_{2r+1,n}$ are connected by a directed edge iff the two corresponding $r$-hop views are connected by a directed edge in the neighborhood graph. Note that if there is an edge between $\alpha$ and $\beta$ in $B_{k,n}$, $\alpha_1 \neq \beta_k$ because the node labels in $\alpha$ and $\beta$ are increasing. □

To determine a lower bound on the number of colors an $r$-round algorithm needs for directed $n$-node rings, it therefore suffices to determine a lower bound on the chromatic number of $B_{2r+1,n}$. To obtain such a lower bound, we need the following definition.
8.2. THE NEIGHBORHOOD GRAPH

**Definition 8.7 (Diline Graph).** The directed line graph (diline graph) $\mathcal{DL}(G)$ of a directed graph $G = (V, E)$ is defined as follows. The node set of $\mathcal{DL}(G)$ is $V[\mathcal{DL}(G)] = E$. There is a directed edge $((w, x), (y, z))$ between $(w, x) \in E$ and $(y, z) \in E$ iff $x = y$, i.e., if the first edge ends where the second one starts.

**Lemma 8.8.** If $n > k$, the graph $B_{k+1,n}$ can be defined recursively as follows:

$$B_{k+1,n} = \mathcal{DL}(B_{k,n}).$$

**Proof.** The edges of $B_{k,n}$ are pairs of $k$-tuples $\alpha = (\alpha_1, \ldots, \alpha_k)$ and $\beta = (\beta_1, \ldots, \beta_k)$ that satisfy Conditions (8.1) and (8.2). Because the last $k-1$ labels in $\alpha$ are equal to the first $k-1$ labels in $\beta$, the pair $(\alpha, \beta)$ can be represented by a $(k+1)$-tuple $\gamma = (\gamma_1, \ldots, \gamma_{k+1})$ with $\gamma_i = \alpha_i$, $\gamma_i = \beta_{i-1} = \alpha_i$ for $2 \leq i \leq k$, and $\gamma_{k+1} = \beta_k$. Because the labels in $\alpha$ and the labels in $\beta$ are increasing, the labels in $\gamma$ are increasing as well. The two graphs $B_{k+1,n}$ and $\mathcal{DL}(B_{k,n})$ therefore have the same node sets. There is an edge between two nodes $(\alpha_1, \beta_1)$ and $(\alpha_2, \beta_2)$ of $\mathcal{DL}(B_{k,n})$ if $\beta_1 = \alpha_2$. This is equivalent to requiring that the two corresponding $(k+1)$-tuples $\gamma_1$ and $\gamma_2$ are neighbors in $B_{k+1,n}$, i.e., that the last $k$ labels of $\gamma_1$ are equal to the first $k$ labels of $\gamma_2$. □

The following lemma establishes a useful connection between the chromatic numbers of a directed graph $G$ and its diline graph $\mathcal{DL}(G)$.

**Lemma 8.9.** For the chromatic numbers $\chi(G)$ and $\chi(\mathcal{DL}(G))$ of a directed graph $G$ and its diline graph, it holds that

$$\chi(\mathcal{DL}(G)) \geq \log_2 (\chi(G)).$$

**Proof.** Given a $c$-coloring of $\mathcal{DL}(G)$, we show how to construct a $2^c$ coloring of $G$. The claim of the lemma then follows because this implies that $\chi(G) \leq 2^{\chi(\mathcal{DL}(G))}$.

Assume that we are given a $c$-coloring of $\mathcal{DL}(G)$. A $c$-coloring of the diline graph $\mathcal{DL}(G)$ can be seen as a coloring of the edges of $G$ such that no two adjacent edges have the same color. For a node $v$ of $G$, let $S_v$ be the set of colors of its outgoing edges. Let $u$ and $v$ be two nodes such that $G$ contains a directed edge $(u, v)$ from $u$ to $v$ and let $x$ be the color of $(u, v)$. Clearly, $x \in S_u$ because $(u, v)$ is an outgoing edge of $u$. Because adjacent edges have different colors, no outgoing edge $(u, w)$ of $v$ can have color $x$. Therefore $x \notin S_v$. This implies that $S_u \neq S_v$. We can therefore use these color sets to obtain a vertex coloring of $G$, i.e., the color of $u$ is $S_u$ and the color of $v$ is $S_v$. Because the number of possible subsets of $[c]$ is $2^c$, this yields a $2^c$-coloring of $G$. □

Let $\log^{(i)} x$ be the $i$-fold application of the base-2 logarithm to $x$:

$$\log^{(i)} x = \log_2 x, \quad \log^{(i+1)} x = \log_2(\log^{(i)} x).$$

Remember from Chapter 1 that

$$\log^* x = 1 \text{ if } x \leq 2, \quad \log^* x = 1 + \min\{i : \log^{(i)} x \leq 2\}.$$

For the chromatic number of $B_{k,n}$, we obtain

**Lemma 8.10.** For all $n \geq 1$, $\chi(B_{1,n}) = n$. Further, for $n \geq k \geq 2$, $\chi(B_{k,n}) \geq \log^{(k-1)} n$. 
Proof. For $k = 1$, $B_{k,n}$ is the complete graph on $n$ nodes with a directed edge from node $i$ to node $j$ iff $i < j$. Therefore, $\chi(B_{1,n}) = n$. For $k > 2$, the claim follows by induction and Lemmas 8.8 and 8.9.

This finally allows us to state a lower bound on the number of rounds needed to color a directed ring with 3 colors.

**Theorem 8.11.** Every deterministic, distributed algorithm to color a directed ring with 3 or less colors needs at least $(\log^* n)/2 - 1$ rounds.

**Proof.** Using the connection between $B_{k,n}$ and the neighborhood graph for directed rings, it suffices to show that $\chi(B_{2r+1,n}) > 3$ for all $r < (\log^* n)/2 - 1$. From Lemma 8.10, we know that $\chi(B_{2r+1,n}) \geq \log(2^r) n$. To obtain $\log(2^r) n \leq 2$, we need $r \geq (\log^* n)/2 - 1$. Because $\log_2 3 < 2$, we therefore have $\log(2^r) n > 3$ if $r < \log^* n/2 - 1$.

**Corollary 8.12.** Every deterministic, distributed algorithm to compute an MIS of a directed ring needs at least $\log^* n/2 - O(1)$ rounds.

**Remarks:**

- It is straightforward to see that also for a constant $c > 3$, the number of rounds needed to color a ring with $c$ or less colors is $\log^* n/2 - O(1)$.

- There basically (up to additive constants) is a gap of a factor of 2 between the $\log^* n + O(1)$ upper bound of Chapter 1 and the $\log^* n/2 - O(1)$ lower bound of this chapter. It is possible to show that the lower bound is tight, even for undirected rings (for directed rings, this will be part of the exercises).

- Alternatively, the lower bound can also be presented as an application of Ramsey’s theory. Ramsey’s theory is best introduced with an example: Assume you host a party, and you want to invite people such that there are no three people who mutually know each other, and no three people which are mutual strangers. How many people can you invite? This is an example of Ramsey’s theorem, which says that for any given integer $c$, and any given integers $n_1, \ldots, n_c$, there is a Ramsey number $R(n_1, \ldots, n_c)$, such that if the edges of a complete graph with $R(n_1, \ldots, n_c)$ nodes are colored with $c$ different colors, then for some color $i$ the graph contains some complete subgraph of color $i$ of size $n_i$. The special case in the party example is looking for $R(3,3)$.

- Ramsey theory is more general, as it deals with hyperedges. A normal edge is essentially a subset of two nodes; a hyperedge is a subset of $k$ nodes. The party example can be explained in this context: We have (hyper)edges of the form $\{i, j\}$, with $1 \leq i, j \leq n$. Choosing $n$ sufficiently large, coloring the edges with two colors must exhibit a set $S$ of 3 edges $\{i, j\} \subset \{v_1, v_2, v_3\}$, such that all edges in $S$ have the same color. To prove our coloring lower bound using Ramsey theory, we form all hyperedges of size $k = 2r+1$, and color them with 3 colors. Choosing $n$ sufficiently large, there must be a set $S = \{v_1, \ldots, v_{k+1}\}$ of $k + 1$ identifiers, such that all $k + 1$ hyperedges consisting of $k$ nodes from $S$ have the same color. Note
that both \{v_1, \ldots, v_k\} and \{v_2, \ldots, v_{k+1}\} are in the set \(S\), hence there will be two neighboring views with the same color. Ramsey theory shows that in this case \(n\) will grow as a power tower (tetration) in \(k\). Thus, if \(n\) is so large that \(k\) is smaller than some function growing like \(\log^* n\), the coloring algorithm cannot be correct.

- The neighborhood graph concept can be used more generally to study distributed graph coloring. It can for instance be used to show that with a single round (every node sends its identifier to all neighbors) it is possible to color a graph with \((1+o(1))\Delta^2 \ln n\) colors, and that every one-round algorithm needs at least \(\Omega(\Delta^2/\log^2 \Delta + \log \log n)\) colors.

- One may also extend the proof to other problems, for instance one may show that a constant approximation of the minimum dominating set problem on unit disk graphs costs at least \(\Omega(\Delta^2/\log^2 \Delta + \log \log n)\) colors.

- Using \(r\)-hop views and the fact that nodes with equal \(r\)-hop views have to make the same decisions is the basic principle behind almost all locality lower bounds (in fact, we are not aware of a locality lower bound that does not use this principle). Using this basic technique (but a completely different proof otherwise), it is for instance possible to show that computing an MIS (and many other problems) in a general graph requires at least \(\Omega(\sqrt{\log n})\) and \(\Omega(\log \Delta)\) rounds.

Chapter Notes

The lower bound proof in this chapter is by Linial [Lin92], proving asymptotic optimality of the technique of Chapter 1. This proof can also be found in Chapter 7.5 of [Pel00]. The lower bound is also true for randomized algorithms [Nao91]. Recently, this lower bound technique was adapted to other problems [CHW08, LW08]. In some sense, Linial’s seminal work raised the question of what can be computed in \(O(1)\) time [NS93], essentially starting distributed complexity theory.

More recently, using a different argument, Kuhn et al. [KMW04] managed to show more substantial lower bounds for a number of combinatorial problems including minimum vertex cover (MVC), minimum dominating set (MDS), maximal matching, or maximal independent set (MIS). More concretely, Kuhn et al. showed that all these problems need polylogarithmic time (for a polylogarithmic approximation, in case of approximation problems such as MVC and MDS). For recent surveys regarding locality lower bounds we refer to e.g. [KMW10, Suo12].

Ramsey theory was started by Frank P. Ramsey with his 1930 article called “On a problem of formal logic” [Ram30]. For an introduction to Ramsey theory we refer to e.g. [NR90, LR03].

Bibliography

CHAPTER 8. LOCALITY LOWER BOUNDS


Chapter 9

Social Networks

Distributed computing is applicable in various contexts. This lecture exemplarily studies one of these contexts, social networks, an area of study whose origins date back a century. To give you a first impression, consider Figure 9.1.

Figure 9.1: This graph shows the social relations between the members of a karate club, studied by anthropologist Wayne Zachary in the 1970s. Two people (nodes) stand out, the instructor and the administrator of the club, both happen to have many friends among club members. At some point, a dispute caused the club to split into two. Can you predict how the club partitioned? (If not, just search the Internet for Zachary and Karate.)
9.1 Small World Networks

Back in 1929, Frigyes Karinthy published a volume of short stories that postulated that the world was “shrinking” because human beings were connected more and more. Some claim that he was inspired by radio network pioneer Guglielmo Marconi’s 1909 Nobel Prize speech. Despite physical distance, the growing density of human “networks” renders the actual social distance smaller and smaller. As a result, it is believed that any two individuals can be connected through at most five (or so) acquaintances, i.e., within six hops.

The topic was hot in the 1960s. For instance, in 1964, Marshall McLuhan coined the metaphor “Global Village”. He wrote: “As electrically contracted, the globe is no more than a village”. He argues that due to the almost instantaneous reaction times of new (“electric”) technologies, each individual inevitably feels the consequences of his actions and thus automatically deeply participates in the global society. McLuhan understood what we now can directly observe – real and virtual world are moving together. He realized that the transmission medium, rather than the transmitted information is at the core of change, as expressed by his famous phrase “the medium is the message”.

This idea has been followed ardently in the 1960s by several sociologists, first by Michael Gurevich, later by Stanley Milgram. Milgram wanted to know the average path length between two “random” humans, by using various experiments, generally using randomly chosen individuals from the US Midwest as starting points, and a stockbroker living in a suburb of Boston as target. The starting points were given name, address, occupation, plus some personal information about the target. They were asked to send a letter to the target. However, they were not allowed to directly send the letter, rather, they had to pass it to somebody they knew on first-name basis and that they thought to have a higher probability to know the target person. This process was repeated, until somebody knew the target person, and could deliver the letter. Shortly after starting the experiment, letters have been received. Most letters were lost during the process, but if they arrived, the average path length was about 5.5. The observation that the entire population is connected by short acquaintance chains got later popularized by the terms “six degrees of separation” and “small world”.

Statisticians tried to explain Milgram’s experiments, by essentially giving network models that allowed for short diameters, i.e., each node is connected to each other node by only a few hops. Until today there is a thriving research community in statistical physics that tries to understand network properties that allow for “small world” effects.

One of the keywords in this area are power-law graphs, networks were node degrees are distributed according to a power-law distribution, i.e., the number of nodes with degree $\delta$ is proportional to $\delta^{-\alpha}$, for some $\alpha > 1$. Such power-law graphs have been witnessed in many application areas, apart from social networks also in the web, or in Biology or Physics.

Obviously, two power-law graphs might look and behave completely differently, even if $\alpha$ and the number of edges is exactly the same.

One well-known model towards this end is the Watts-Strogatz model. Watts and Strogatz argued that social networks should be modeled by a combination of two networks: As the basis we take a network that has a large cluster coefficient
9.1 SMALL WORLD NETWORKS

**Definition 9.1.** The cluster coefficient of a network is defined by the probability that two friends of a node are likely to be friends as well, averaged over all the nodes.

\[ \text{Definition 9.1.} \]

\[ \text{... then we augment such a graph with random links, every node for instance points to a constant number of other nodes, chosen uniformly at random. This augmentation represents acquaintances that connect nodes to parts of the network that would otherwise be far away.} \]

**Remarks:**

- Without further information, knowing the cluster coefficient is of questionable value: Assume we arrange the nodes in a grid. Technically, if we connect each node to its four closest neighbors, the graph has cluster coefficient 0, since there are no triangles; if we instead connect each node with its eight closest neighbors, the cluster coefficient is 3/7. The cluster coefficient is quite different, even though both networks have similar characteristics.

This is interesting, but not enough to really understand what is going on. For Milgram’s experiments to work, it is not sufficient to connect the nodes in a certain way. In addition, the nodes themselves need to know how to forward a message to one of their neighbors, even though they cannot know whether that neighbor is really closer to the target. In other words, nodes are not just following physical laws, but they make decisions themselves.

Let us consider an artificial network with nodes on a grid topology, plus some additional random links per node. In a quantitative study it was shown that the random links need a specific distance distribution to allow for efficient greedy routing. This distribution marks the sweet spot for any navigable network.

**Definition 9.2 (Augmented Grid).** We take \( n = m^2 \) nodes \((i, j) \in V = \{1, \ldots, m\}^2 \) that are identified with the lattice points on an \( m \times m \) grid. We define the distance between two nodes \((i, j)\) and \((k, \ell)\) as \( d((i, j), (k, \ell)) = |k - i| + |\ell - j| \) as the distance between them on the \( m \times m \) lattice. The network is modeled using a parameter \( \alpha \geq 0 \). Each node \( u \) has a directed edge to every lattice neighbor. These are the local contacts of a node. In addition, each node also has an additional random link (the long-range contact). For all \( u \) and \( v \), the long-range contact of \( u \) points to node \( v \) with probability proportional to \( d(u, v)^{-\alpha} \), i.e., with probability \( d(u, v)^{-\alpha} / \sum_{w \in V \setminus \{u\}} d(u, w)^{-\alpha} \). Figure 9.2 illustrates the model.

**Remarks:**

- The network model has the following geographic interpretation: nodes (individuals) live on a grid and know their neighbors on the grid. Further, each node has some additional acquaintances throughout the network.

- The parameter \( \alpha \) controls how the additional neighbors are distributed across the grid. If \( \alpha = 0 \), long-range contacts are chosen uniformly at random (as in the Watts-Strogatz model). As \( \alpha \) increases, long-range contacts become shorter on average. In the extreme case, if \( \alpha \to \infty \), all long-range contacts are to immediate neighbors on the grid.
It can be shown that as long as $\alpha \leq 2$, the diameter of the resulting graph is polylogarithmic in $n$ (polynomial in $\log n$) with high probability. In particular, if the long-range contacts are chosen uniformly at random ($\alpha = 0$), the diameter is $O(\log n)$.

Since the augmented grid contains random links, we do not know anything for sure about how the random links are distributed. In theory, all links could point to the same node! However, this is almost certainly not the case. Formally this is captured by the term with high probability.

**Definition 9.3 (With High Probability).** Some probabilistic event is said to occur with high probability (w.h.p.), if it happens with a probability $p \geq 1 - 1/n^c$, where $c$ is a constant. The constant $c$ may be chosen arbitrarily, but it is considered constant with respect to $\mathcal{O}$ notation.

**Remarks:**

- For instance, a running time bound of $c \log n$ or $e^{c} \log n + 5000c$ with probability at least $1 - 1/n^c$ would be $O(\log n)$ w.h.p., but a running time of $n^c$ would not be $\mathcal{O}(n)$ w.h.p. since $c$ might also be 50.

- This definition is very powerful, as any polynomial (in $n$) number of statements that hold w.h.p. also holds w.h.p. at the same time, regardless of any dependencies between random variables!
9.1. SMALL WORLD NETWORKS

**Theorem 9.4.** The diameter of the augmented grid with $\alpha = 0$ is $O(\log n)$ with high probability.

**Proof Sketch.** For simplicity, we will only show that we can reach a target node $t$ starting from some source node $s$. However, it can be shown that (essentially) each of the intermediate claims holds with high probability, which then by means of the union bound yields that all of the claims hold simultaneously with high probability for all pairs of nodes (see exercises).

Let $N_s$ be the $\lceil \log n \rceil$-hop neighborhood of source $s$ on the grid, containing $\Omega(\log^2 n)$ nodes. Each of the nodes in $N_s$ has a random link, probably leading to distant parts of the graph. As long as we have reached only $o(n)$ nodes, any new random link will with probability $1-o(1)$ lead to a node for which none of its grid neighbors has been visited yet. Thus, in expectation we find almost $|N_s|$ new nodes whose neighbors are “fresh”. Using their grid links, we will reach $(4-o(1))|N_s|$ more nodes within one more hop. If bad luck strikes, it could still happen that many of these links lead to a few nodes, already visited nodes, or nodes that are very close to each other. But that is very unlikely, as we have lots of random choices! Indeed, it can be shown that not only in expectation, but with high probability $(5-o(1))|N_s|$ many nodes are reached this way (see exercises).

Because all the new nodes have (so far unused) random links, we can repeat this reasoning inductively, implying that the number of nodes grows by (at least) a constant factor for every two hops. Thus, after $O(\log n)$ hops, we will have reached $n/\log n$ nodes (which is still small compared to $n$). Finally, consider the expected number of links from these nodes that enter the $(\log n)$-neighborhood of some target node $t$ with respect to the grid. Since this neighborhood consists of $\Omega(\log^2 n)$ nodes, in expectation $\Omega(\log n)$ links come close enough to target $t$. This is large enough to almost guarantee that this happens (see exercises). Summing everything up, we still used merely $O(\log n)$ hops in total to get from $s$ to $t$.

This shows that for $\alpha = 0$ (and in fact for all $\alpha \leq 2$), the resulting network has a small diameter. Recall however that we also wanted the network to be navigable. For this, we consider a simple greedy routing strategy (Algorithm 39).

**Algorithm 39 Greedy Routing**

1: while not at destination do
2: go to a neighbor which is closest to destination (considering grid distance only)
3: end while

**Lemma 9.5.** In the augmented grid, Algorithm 39 finds a routing path of length at most $2(m-1) \in O(\sqrt{n})$.

**Proof.** Because of the grid, there is always a neighbor which is closer to the destination. Since with each hop we reduce the distance to the target at least by one in one of the two grid dimensions, we will reach the destination within $2(m-1)$ steps.

\qed
This is not really what Milgram’s experiment promises. We want to know how much the additional random links speed up the process. To this end, we first need to understand how likely it is that the random link of node $u$ points to node $v$, in terms of their grid distance $d(u, v)$, the number of nodes $n$, and the constant parameter $\alpha$.

**Lemma 9.6.** Node $u$’s random link points to a node $v$ with probability

- $\Theta(1/(d(u, v)^{\alpha}m^{2-\alpha}))$ if $\alpha < 2$.
- $\Theta(1/(d(u, v)^{2} \log n))$ if $\alpha = 2$.
- $\Theta(1/d(u, v)^{\alpha})$ if $\alpha > 2$.

Moreover, if $\alpha > 2$, the probability to see a link of length at least $d$ is in $\Theta(1/d^{\alpha-2})$.

**Proof.** For a constant $\alpha \neq 2$, we have that

$$\sum_{u \in V \setminus \{u\}} \frac{1}{d(u, w)^{\alpha}} \in \sum_{r=1}^{m} \frac{\Theta(r)}{r^{\alpha}} = \Theta \left( \int_{r=1}^{m} \frac{1}{r^{\alpha-1}} \, dr \right) = \Theta \left( \left[ \frac{r^{2-\alpha}}{2-\alpha} \right]_{1}^{m} \right).$$

If $\alpha < 2$, this gives $\Theta(m^{2-\alpha})$, if $\alpha > 2$, it is in $\Theta(1)$. If $\alpha = 2$, we get

$$\sum_{u \in V \setminus \{u\}} \frac{1}{d(u, w)^{\alpha}} \in \sum_{r=1}^{m} \frac{\Theta(r)}{r^{2}} = \Theta(1) \cdot \sum_{r=1}^{m} \frac{1}{r} = \Theta(\log m) = \Theta(\log n).$$

Multiplying with $d(u, v)^{\alpha}$ yields the first three bounds. For the last statement, compute

$$\sum_{\substack{v \in V \backslash \{u\} \atop d(u, v) \geq d}} \Theta(1/d(u, v)^{\alpha}) = \Theta \left( \int_{r=d}^{m} \frac{r}{r^{\alpha}} \, dr \right) = \Theta \left( \left[ \frac{r^{2-\alpha}}{2-\alpha} \right]_{d}^{m} \right) = \Theta(1/d^{\alpha-2}).$$

\[\square\]

**Remarks:**

- If $\alpha > 2$, according to the lemma, the probability to see a random link of length at least $d = m^{1/(\alpha-1)}$ is $\Theta(1/d^{\alpha-2}) = \Theta(1/m^{(\alpha-2)/(\alpha-1)})$. In expectation we have to take $\Theta(m^{(\alpha-2)/(\alpha-1)})$ hops until we see a random link of length at least $d$. When just following links of length less than $d$, it takes more than $m/d = m/m^{1/(\alpha-1)} = m^{(\alpha-2)/(\alpha-1)}$ hops. In other words, in expectation, either way we need at least $m^{(\alpha-2)/(\alpha-1)} = m^{\Omega(1)}$ hops to the destination.

- If $\alpha < 2$, there is a (slightly more complicated) argument. First we draw a border around the nodes in distance $m^{(2-\alpha)/3}$ to the target. Within this border there are about $m^{2(2-\alpha)/3}$ many nodes in the target area. Assume that the source is outside the target area. Starting at the source, the probability to find a random link that leads directly inside the target area is according to the lemma at most $m^{2(2-\alpha)/3} \cdot \Theta(1/m^{2-\alpha}) = \Theta(1/m^{(2-\alpha)/3})$. In other words, until we find a random link that leads into the target area,
9.1. SMALL WORLD NETWORKS

in expectation, we have to do $\Theta(m^{(2-\alpha)/3})$ hops. This is too slow, and our greedy strategy is probably faster, as thanks to having $\alpha < 2$ there are many long-range links. However, it means that we will probably enter the border of the target area on a regular grid link. Once inside the target area, again the probability of short-cutting our trip by a random long-range link is $\Theta(1/m^{(2-\alpha)/3})$, so we probably just follow grid links, $m^{(2-\alpha)/3} = m^\Omega(1)$ many of them.

- In summary, if $\alpha \neq 2$, our greedy routing algorithm takes $m^\Omega(1) = n^\Omega(1)$ expected hops to reach the destination. This is polynomial in the number of nodes $n$, and the social network can hardly be called a “small world”.

- Maybe we can get a polylogarithmic bound on $n$ if we set $\alpha = 2$?

**Definition 9.7 (Phase).** Consider routing from source $s$ to target $t$ and assume that we are at some intermediate node $w$. We say that we are in phase $j$ at node $w$ if the lattice distance $d(w, t)$ to the target node $t$ is between $2^j < d(w, t) \leq 2^{j+1}$.

**Remarks:**

- Enumerating the phases in decreasing order is useful, as notation becomes less cumbersome.

- There are $\lceil \log m \rceil \in O(\log n)$ phases.

**Lemma 9.8.** Assume that we are in phase $j$ at node $w$ when routing from $s$ to $t$. The probability for getting (at least) to phase $j - 1$ in one step is at least $\Omega(1/\log n)$.

**Proof.** Let $B_j$ be the set of nodes $x$ with $d(x, t) \leq 2^j$. We get from phase $j$ to (at least) phase $j - 1$ if the long-range contact of node $w$ points to some node in $B_j$. Note that we always make progress while following the greedy routing path. Therefore, we have not seen node $w$ before and the long-range contact of $w$ points to a random node that is independent of anything seen on the path from $s$ to $w$.

For all nodes $x \in B_j$, we have $d(w, x) \leq d(w, t) + d(x, t) \leq 2^{j+1} + 2^j < 2^{j+2}$. Hence, for each node $x \in B_j$, the probability that the long-range contact of $w$ points to $x$ is $\Omega(1/2^{j+2} \log n)$. Further, the number of nodes in $B_j$ is at least $(2^j)^2/2 = 2^{2j-1}$. Hence, the probability that some node in $B_j$ is the long range contact of $w$ is at least

$$\Omega \left( \frac{|B_j| \cdot \frac{1}{2^{2j+4} \log n}}{2^{2j+4} \log n} \right) = \Omega \left( \frac{2^{2j-1}}{2^{2j+4} \log n} \right) = \Omega \left( \frac{1}{\log n} \right).$$

**Theorem 9.9.** Consider the greedy routing path from a node $s$ to a node $t$ on an augmented grid with parameter $\alpha = 2$. The expected length of the path is $O(\log^2 n)$.

**Proof.** We already observed that the total number of phases is $O(\log n)$ (the distance to the target is halved when we go from phase $j$ to phase $j - 1$). At each point during the routing process, the probability of proceeding to the next phase is at least $\Omega(1/\log n)$. Let $X_j$ be the number of steps in phase $j$. Because...
the probability for ending the phase is \( \Omega(1/\log n) \) in each step, in expectation we need \( O(\log n) \) steps to proceed to the next phase, i.e., \( E[X_j] \in O(\log n) \). Let 
\[ X = \sum_j X_j \]
be the total number of steps of the routing process. By linearity of expectation, we have
\[
E[X] = \sum_j E[X_j] \in O(\log^2 n).
\]

Remarks:

• One can show that the \( O(\log^2 n) \) result also holds w.h.p.
• In real world social networks, the parameter \( \alpha \) was evaluated experimentally. The assumption is that you are connected to the geographically closest nodes, and then have some random long-range contacts. For Facebook grandpa LiveJournal it was shown that \( \alpha \) is not really 2, but rather around 1.25.

9.2 Propagation Studies

In networks, nodes may influence each other’s behavior and decisions. There are many applications where nodes influence their neighbors, e.g., they may impact their opinions, or they may bias what products they buy, or they may pass on a disease.

On a beach (modeled as a line segment), it is best to place an ice cream stand right in the middle of the segment, because you will be able to “control” the beach most easily. What about the second stand, where should it settle? The answer generally depends on the model, but assuming that people will buy ice cream from the stand that is closer, it should go right next to the first stand.

Rumors can spread surprisingly fast through social networks. Traditionally this happens by word of mouth, but with the emergence of the Internet and its possibilities new ways of rumor propagation are available. People write email, use instant messengers or publish their thoughts in a blog. Many factors influence the dissemination of rumors. It is especially important where in a network a rumor is initiated and how convincing it is. Furthermore the underlying network structure decides how fast the information can spread and how many people are reached. More generally, we can speak of diffusion of information in networks. The analysis of these diffusion processes can be useful for viral marketing, e.g., to target a few influential people to initiate marketing campaigns. A company may wish to distribute the rumor of a new product via the most influential individuals in popular social networks such as Facebook. A second company might want to introduce a competing product and has hence to select where to seed the information to be disseminated. Rumor spreading is quite similar to our ice cream stand problem.

More formally, we may study propagation problems in graphs. Given a graph, and two players. Let the first player choose a seed node \( u_1 \); afterwards let the second player choose a seed node \( u_2 \), with \( u_2 \neq u_1 \). The goal of the game is to maximize the number of nodes that are closer to one’s own seed node.

In many graphs it is an advantage to choose first. In a star graph for instance the first player can choose the center node of the star, controlling all but one
node. In some other graphs, the second player can at least score even. But is there a graph where the second player has an advantage?

**Theorem 9.10.** In a two player rumor game where both players select one node to initiate their rumor in the graph, the first player does not always win.

**Proof.** See Figure 9.3 for an example where the second player will always win, regardless of the decision the first player. If the first player chooses the node $x_0$ in the center, the second player can select $x_1$. Choice $x_1$ will be outwitted by $x_2$, and $x_2$ itself can be answered by $z_1$. All other strategies are either symmetric, or even less promising for the first player.

![Figure 9.3: Counter example.](image.png)

**Chapter Notes**

A simple form of a social network is the famous stable marriage problem [DS62] in which a stable matching bipartite graph has to be found. There exists a great many of variations which are based on this initial problem, e.g., [KC82, KMV94, EO06, FKPS10, Hoe11]. Social networks like Facebook, Twitter and others have grown very fast in the last years and hence spurred interest to research them. How users influence other users has been studied both from a theoretical point of view [KKT03] and in practice [CHBG10]. The structure of these networks can be measured and studied [MMG07]. More than half of the users in social networks share more information than they expect to [LGKM11].

The small world phenomenon that we presented in this chapter is analyzed by Kleinberg [Kle00]. A general overview is in [DJ10]. This chapter has been written in collaboration with Michael Kuhn.

**Bibliography**


Chapter 10

Synchronization

So far, we have mainly studied synchronous algorithms. Generally, asynchronous algorithms are more difficult to obtain. Also it is substantially harder to reason about asynchronous algorithms than about synchronous ones. For instance, computing a BFS tree (Chapter 3) efficiently requires much more work in an asynchronous system. However, many real systems are not synchronous, and we therefore have to design asynchronous algorithms. In this chapter, we will look at general simulation techniques, called synchronizers, that allow running synchronous algorithms in asynchronous environments.

10.1 Basics

A synchronizer generates sequences of clock pulses at each node of the network satisfying the condition given by the following definition.

Definition 10.1 (valid clock pulse). We call a clock pulse generated at a node \( v \) valid if it is generated after \( v \) received all the messages of the synchronous algorithm sent to \( v \) by its neighbors in the previous pulses.

Given a mechanism that generates the clock pulses, a synchronous algorithm is turned into an asynchronous algorithm in an obvious way: As soon as the \( i \)th clock pulse is generated at node \( v \), \( v \) performs all the actions (local computations and sending of messages) of round \( i \) of the synchronous algorithm.

Theorem 10.2. If all generated clock pulses are valid according to Definition 10.1, the above method provides an asynchronous algorithm that behaves exactly the same way as the given synchronous algorithm.

Proof. When the \( i \)th pulse is generated at a node \( v \), \( v \) has sent and received exactly the same messages and performed the same local computations as in the first \( i - 1 \) rounds of the synchronous algorithm.

The main problem when generating the clock pulses at a node \( v \) is that \( v \) cannot know what messages its neighbors are sending to it in a given synchronous round. Because there are no bounds on link delays, \( v \) cannot simply wait “long enough” before generating the next pulse. In order satisfy Definition 10.1, nodes have to send additional messages for the purpose of synchronization. The total
complexity of the resulting asynchronous algorithm depends on the overhead introduced by the synchronizer. For a synchronizer $S$, let $T(S)$ and $M(S)$ be the time and message complexities of $S$ for each generated clock pulse. As we will see, some of the synchronizers need an initialization phase. We denote the time and message complexities of the initialization by $T_{\text{init}}(S)$ and $M_{\text{init}}(S)$, respectively. If $T(A)$ and $M(A)$ are the time and message complexities of the given synchronous algorithm $A$, the total time and message complexities $T_{\text{tot}}$ and $M_{\text{tot}}$ of the resulting asynchronous algorithm then become

$$T_{\text{tot}} = T_{\text{init}}(S) + T(A) \cdot (1 + T(S))$$

and

$$M_{\text{tot}} = M_{\text{init}}(S) + M(A) + T(A) \cdot M(S),$$

respectively.

Remarks:

- Because the initialization only needs to be done once for each network, we will mostly be interested in the overheads $T(S)$ and $M(S)$ per round of the synchronous algorithm.

Definition 10.3 (Safe Node). A node $v$ is safe with respect to a certain clock pulse if all messages of the synchronous algorithm sent by $v$ in that pulse have already arrived at their destinations.

Lemma 10.4. If all neighbors of a node $v$ are safe with respect to the current clock pulse of $v$, the next pulse can be generated for $v$.

Proof. If all neighbors of $v$ are safe with respect to a certain pulse, $v$ has received all messages of the given pulse. Node $v$ therefore satisfies the condition of Definition 10.1 for generating a valid next pulse.

Remarks:

- In order to detect safety, we require that all algorithms send acknowledgements for all received messages. As soon as a node $v$ has received an acknowledgement for each message that it has sent in a certain pulse, it knows that it is safe with respect to that pulse. Note that sending acknowledgements does not increase the asymptotic time and message complexities.

10.2 The Local Synchronizer $\alpha$

Algorithm 40 Synchronizer $\alpha$ (at node $v$)

1: wait until $v$ is safe
2: send SAFE to all neighbors
3: wait until $v$ receives SAFE messages from all neighbors
4: start new pulse

Synchronizer $\alpha$ is very simple. It does not need an initialization. Using acknowledgements, each node eventually detects that it is safe. It then reports this fact directly to all its neighbors. Whenever a node learns that all its neighbors are safe, a new pulse is generated. Algorithm 40 formally describes the synchronizer $\alpha$. 
Theorem 10.5. The time and message complexities of synchronizer $\alpha$ per synchronous round are

$$T(\alpha) = O(1) \quad \text{and} \quad M(\alpha) = O(m).$$

Proof. Communication is only between neighbors. As soon as all neighbors of a node $v$ become safe, $v$ knows of this fact after one additional time unit. For every clock pulse, synchronizer $\alpha$ sends at most four additional messages over every edge: Each of the nodes may have to acknowledge a message and reports safety. \hfill $\square$

Remarks:

- Synchronizer $\alpha$ was presented in a framework, mostly set up to have a common standard to discuss different synchronizers. Without the framework, synchronizer $\alpha$ can be explained more easily:
  
  1. Send message to all neighbors, include round information $i$ and actual data of round $i$ (if any).
  2. Wait for message of round $i$ from all neighbors, and go to next round.

- Although synchronizer $\alpha$ allows for simple and fast synchronization, it produces awfully many messages. Can we do better? Yes.

10.3 The Global Synchronizer $\beta$

Algorithm 41 Synchronizer $\beta$ (at node $v$)

1: \textbf{wait} until $v$ is safe
2: \textbf{wait} until $v$ receives SAFE messages from all its children in $T$
3: \textbf{if} $v \neq \ell$ \textbf{then}
4: \textbf{send} SAFE message to parent in $T$
5: \textbf{wait} until PULSE message received from parent in $T$
6: \textbf{end if}
7: \textbf{send} PULSE message to children in $T$
8: start new pulse

Synchronizer $\beta$ needs an initialization that computes a leader node $\ell$ and a spanning tree $T$ rooted at $\ell$. As soon as all nodes are safe, this information is propagated to $\ell$ by a convergecast. The leader then broadcasts this information to all nodes. The details of synchronizer $\beta$ are given in Algorithm 41.

Theorem 10.6. The time and message complexities of synchronizer $\beta$ per synchronous round are

$$T(\beta) = O(\text{diameter}(T)) \leq O(n) \quad \text{and} \quad M(\beta) = O(n).$$

The time and message complexities for the initialization are

$$T_{\text{init}}(\beta) = O(n) \quad \text{and} \quad M_{\text{init}}(\beta) = O(m + n \log n).$$
Proof. Because the diameter of $T$ is at most $n - 1$, the convergecast and the broadcast together take at most $2n - 2$ time units. Per clock pulse, the synchronizer sends at most $2n - 2$ synchronization messages (one in each direction over each edge of $T$).

With an improvement (due to Awerbuch) of the GHS algorithm (Algorithm 15) you saw in Chapter 3, it is possible to construct an MST in time $O(n)$ with $O(m + n \log n)$ messages in an asynchronous environment. Once the tree is computed, the tree can be made rooted in time $O(n)$ with $O(n)$ messages.

Remarks:

- We now got a time-efficient synchronizer ($\alpha$) and a message-efficient synchronizer ($\beta$), it is only natural to ask whether we can have the best of both worlds. And, indeed, we can. How is that synchronizer called? Quite obviously: $\gamma$.

10.4 The Hybrid Synchronizer $\gamma$

Figure 10.1: A cluster partition of a network: The dashed cycles specify the clusters, cluster leaders are black, the solid edges are the edges of the intracluster trees, and the bold solid edges are the intercluster edges.

Synchronizer $\gamma$ can be seen as a combination of synchronizers $\alpha$ and $\beta$. In the initialization phase, the network is partitioned into clusters of small diameter. In each cluster, a leader node is chosen and a BFS tree rooted at this leader node is computed. These trees are called the intracluster trees. Two clusters $C_1$ and $C_2$ are called neighboring if there are nodes $u \in C_1$ and $v \in C_2$ for which $(u, v) \in E$. For every two neighboring clusters, an intercluster edge is chosen, which will serve for communication between these clusters. Figure 10.1 illustrates this partitioning into clusters. We will discuss the details of how to construct such a partition in the next section. We say that a cluster is safe if all its nodes are safe.
Synchronizer $\gamma$ works in two phases. In a first phase, synchronizer $\beta$ is applied separately in each cluster by using the intracluster trees. Whenever the leader of a cluster learns that its cluster is safe, it reports this fact to all the nodes in the clusters as well as to the leaders of the neighboring clusters. Now, the nodes of the cluster enter the second phase where they wait until all the neighboring clusters are known to be safe and then generate the next pulse. Hence, we essentially apply synchronizer $\alpha$ between clusters. A detailed description is given by Algorithm 42.

**Algorithm 42** Synchronizer $\gamma$ (at node $v$)

1: wait until $v$ is safe
2: wait until $v$ receives SAFE messages from all children in intracluster tree
3: if $v$ is not cluster leader then
4: send SAFE message to parent in intracluster tree
5: wait until CLUSTERSAFE message received from parent
6: end if
7: send CLUSTERSAFE message to all children in intracluster tree
8: send NEIGHBORSAFE message over all intercluster edges of $v$
9: wait until $v$ receives NEIGHBORSAFE messages from all adjacent intercluster edges and all children in intracluster tree
10: if $v$ is not cluster leader then
11: send NEIGHBORSAFE message to parent in intracluster tree
12: wait until PULSE message received from parent
13: end if
14: send PULSE message to children in intracluster tree
15: start new pulse

**Theorem 10.7.** Let $m_C$ be the number of intercluster edges and let $k$ be the maximum cluster radius (i.e., the maximum distance of a leaf to its cluster leader). The time and message complexities of synchronizer $\gamma$ are

$$T(\gamma) = O(k) \quad \text{and} \quad M(\gamma) = O(n + m_C).$$

**Proof.** We ignore acknowledgements, as they do not affect the asymptotic complexities. Let us first look at the number of messages. Over every intracluster tree edge, exactly one SAFE message, one CLUSTERSAFE message, one NEIGHBORSAFE message, and one PULSE message is sent. Further, one NEIGHBORSAFE message is sent over every intercluster edge. Hence there are less than $n$ intracluster tree edges and the total message complexity therefore is at most $4n + 2m_C = O(n + m_C)$.

For the time complexity, note that the depth of each intracluster tree is at most $k$. On each intracluster tree, two convergecasts (the SAFE and NEIGHBORSAFE messages) and two broadcasts (the CLUSTERSAFE and PULSE messages) are performed. The time complexity for this is at most $4k$. There is one more time unit needed to send the NEIGHBORSAFE messages over the intercluster edges. The total time complexity therefore is at most $4k + 1 = O(k)$. □
10.5 Network Partition

We will now look at the initialization phase of synchronizer $\gamma$. Algorithm 43 describes how to construct a partition into clusters that can be used for synchronizer $\gamma$. In Algorithm 43, $B(v, r)$ denotes the ball of radius $r$ around $v$, i.e., $B(v, r) = \{ u \in V : d(u, v) \leq r \}$ where $d(u, v)$ is the hop distance between $u$ and $v$. The algorithm has a parameter $\rho > 1$. The clusters are constructed sequentially. Each cluster is started at an arbitrary node that has not been included in a cluster. Then the cluster radius is grown as long as the cluster grows by a factor more than $\rho$.

**Algorithm 43 Cluster construction**

1: while unprocessed nodes do  
2: select an arbitrary unprocessed node $v$;  
3: $r := 0$;  
4: while $|B(v, r + 1)| > \rho |B(v, r)|$ do  
5: $r := r + 1$  
6: end while  
7: makeCluster($B(v, r)$)  
8: end while

Remarks:

- The algorithm allows a trade-off between the cluster diameter $k$ (and thus the time complexity) and the number of intercluster edges $m_C$ (and thus the message complexity). We will quantify the possibilities in the next section.

- Two very simple partitions would be to make a cluster out of every single node or to make one big cluster that contains the whole graph. We then get synchronizers $\alpha$ and $\beta$ as special cases of synchronizer $\gamma$.

**Theorem 10.8.** Algorithm 43 computes a partition of the network graph into clusters of radius at most $\log_\rho n$. The number of intercluster edges is at most $(\rho - 1) \cdot n$.

**Proof.** The radius of a cluster is initially 0 and does only grow as long as it grows by a factor larger than $\rho$. Since there are only $n$ nodes in the graph, this can happen at most $\log_\rho n$ times.

To count the number of intercluster edges, observe that an edge can only become an intercluster edge if it connects a node at the boundary of a cluster with a node outside a cluster. Consider a cluster $C$ of size $|C|$. We know that $C = B(v, r)$ for some $v \in V$ and $r \geq 0$. Further, we know that $|B(v, r + 1)| \leq \rho \cdot |B(v, r)|$. The number of nodes adjacent to cluster $C$ is therefore at most $|B(v, r + 1) \setminus B(v, r)| \leq \rho \cdot |C| - |C|$. Because there is only one intercluster edge connecting two clusters by definition, the number of intercluster edges adjacent to $C$ is at most $(\rho - 1) \cdot |C|$. Summing over all clusters, we get that the total number of intercluster edges is at most $(\rho - 1) \cdot n$.

**Corollary 10.9.** Using $\rho = 2$, Algorithm 43 computes a clustering with cluster radius at most $\log_2 n$ and with at most $n$ intercluster edges.
Corollary 10.10. Using $\rho = n^{1/k}$, Algorithm 43 computes a clustering with cluster radius at most $k$ and at most $O(n^{1+1/k})$ intercluster edges.

Remarks:
- Algorithm 43 describes a centralized construction of the partitioning of the graph. For $\rho \geq 2$, the clustering can be computed by an asynchronous distributed algorithm in time $O(n)$ with $O(m + n \log n)$ (reasonably sized) messages (showing this will be part of the exercises).
- It can be shown that the trade-off between cluster radius and number of intercluster edges of Algorithm 43 is asymptotically optimal. There are graphs for which every clustering into clusters of radius at most $k$ requires $n^{1+c/k}$ intercluster edges for some constant $c$.

The above remarks lead to a complete characterization of the complexity of synchronizer $\gamma$.

Corollary 10.11. The time and message complexities of synchronizer $\gamma$ per synchronous round are

$$T(\gamma) = O(k) \quad \text{and} \quad M(\gamma) = O(n^{1+1/k}).$$

The time and message complexities for the initialization are

$$T_{\text{init}}(\gamma) = O(n) \quad \text{and} \quad M_{\text{init}}(\gamma) = O(m + n \log n).$$

Remarks:
- The synchronizer idea and the synchronizers discussed in this chapter are due to Baruch Awerbuch.
- In Chapter 3, you have seen that by using flooding, there is a very simple synchronous algorithm to compute a BFS tree in time $O(D)$ with message complexity $O(m)$. If we use synchronizer $\gamma$ to make this algorithm asynchronous, we get an algorithm with time complexity $O(n + D\log n)$ and message complexity $O(m + n \log n + D \cdot n)$ (including initialization).
- The synchronizers $\alpha$, $\beta$, and $\gamma$ achieve global synchronization, i.e. every node generates every clock pulse. The disadvantage of this is that nodes that do not participate in a computation also have to participate in the synchronization. In many computations (e.g. in a BFS construction), many nodes only participate for a few synchronous rounds. An improved synchronizer due to Awerbuch and Peleg can exploit such a scenario and achieves time and message complexity $O(\log^3 n)$ per synchronous round (without initialization).
- It can be shown that if all nodes in the network need to generate all pulses, the trade-off of synchronizer $\gamma$ is asymptotically optimal.
- Partitions of networks into clusters of small diameter and covers of networks with clusters of small diameters come in many variations and have various applications in distributed computations. In particular, apart from synchronizers, algorithms for routing, the construction of sparse spanning subgraphs, distributed data structures, and even computations of local structures such as a MIS or a dominating set are based on some kind of network partitions or covers.
10.6 Clock Synchronization

“A man with one clock knows what time it is – a man with two is never sure.”

Synchronizers can directly be used to give nodes in an asynchronous network a common notion of time. In wireless networks, for instance, many basic protocols need an accurate time. Sometimes a common time in the whole network is needed, often it is enough to synchronize neighbors. The purpose of the time division multiple access (TDMA) protocol is to use the common wireless channel as efficiently as possible, i.e., interfering nodes should never transmit at the same time (on the same frequency). If we use synchronizer $\beta$ to give the nodes a common notion of time, every single clock cycle costs $D$ time units!

Often, each (wireless) node is equipped with an internal clock. Using this clock, it should be possible to divide time into slots, and make each node send (or listen, or sleep, respectively) in the appropriate slots according to the media access control (MAC) layer protocol used.

However, as it turns out, synchronizing clocks in a network is not trivial. As nodes’ internal clocks are not perfect, they will run at speeds that are time-dependent. For instance, variations in temperature or supply voltage will affect this clock drift. For standard clocks, the drift is in the order of parts per million, i.e., within a second, it will accumulate to a couple of microseconds. Wireless TDMA protocols account for this by introducing guard times. Whenever a node knows that it is about to receive a message from a neighbor, it powers up its radio a little bit earlier to make sure that it does not miss the message even when clocks are not perfectly synchronized. If nodes are badly synchronized, messages of different slots might collide.

In the clock synchronization problem, we are given a network (graph) with $n$ nodes. The goal for each node is to have a logical clock such that the logical clock values are well synchronized, and close to real time. Each node is equipped with a hardware clock, that ticks more or less in real time, i.e., the time between two pulses is arbitrary between $[1 - \epsilon, 1 + \epsilon]$, for a constant $\epsilon \ll 1$. Similarly as in our asynchronous model, we assume that messages sent over the edges of the graph have a delivery time between $[0, 1]$. In other words, we have a bounded but variable drift on the hardware clocks and an arbitrary jitter in the delivery times. The goal is to design a message-passing algorithm that ensures that the logical clock skew of adjacent nodes is as small as possible at all times.

**Theorem 10.12.** The global clock skew (the logical clock difference between any two nodes in the graph) is $\Omega(D)$, where $D$ is the diameter of the graph.

**Proof.** For a node $u$, let $t_u$ be the logical time of $u$ and let $(u \rightarrow v)$ denote a message sent from $u$ to a node $v$. Let $t(m)$ be the time delay of a message $m$ and let $u$ and $v$ be neighboring nodes. First consider a case where the message delays between $u$ and $v$ are $1/2$. Then all the messages sent by $u$ and $v$ at time $i$ according to the clock of the sender arrive at time $i + 1/2$ according to the clock of the receiver.

Then consider the following cases

- $t_u = t_v + 1/2$, $t(u \rightarrow v) = 1$, $t(v \rightarrow u) = 0$
- $t_u = t_v - 1/2$, $t(u \rightarrow v) = 0$, $t(v \rightarrow u) = 1$,
where the message delivery time is always fast for one node and slow for the other and the logical clocks are off by $1/2$. In both scenarios, the messages sent at time $i$ according to the clock of the sender arrive at time $i + 1/2$ according to the logical clock of the receiver. Therefore, for nodes $u$ and $v$, both cases with clock drift seem the same as the case with perfectly synchronized clocks.

Furthermore, in a linked list of $D$ nodes, the left- and rightmost nodes $l, r$ cannot distinguish $t_l = t_r + D/2$ from $t_l = t_r - D/2$.

Remarks:

- From Theorem 10.12, it directly follows that all the clock synchronization algorithms we studied have a global skew of $\Omega(D)$.

- Many natural algorithms manage to achieve a global clock skew of $O(D)$.

As both the message jitter and hardware clock drift are bounded by constants, it feels like we should be able to get a constant drift between neighboring nodes. As synchronizer $\alpha$ pays most attention to the local synchronization, we take a look at a protocol inspired by the synchronizer $\alpha$. A pseudo-code representation for the clock synchronization protocol $\alpha$ is given in Algorithm 44.

Algorithm 44 Clock synchronization $\alpha$ (at node $v$)

1: repeat
2: send logical time $t_v$ to all neighbors
3: if Receive logical time $t_u$, where $t_u > t_v$, from any neighbor $u$ then
4: \quad $t_v := t_u$
5: end if
6: until done

Lemma 10.13. The clock synchronization protocol $\alpha$ has a local skew of $\Omega(n)$.

Proof. Let the graph be a linked list of $D$ nodes. We denote the nodes by $v_1, v_2, \ldots, v_D$ from left to right and the logical clock of node $v_i$ by $t_i$. Apart from the left-most node $v_1$ all hardware clocks run with speed 1 (real time). Node $v_1$ runs at maximum speed, i.e. the time between two pulses is not 1 but $1 - \epsilon$. Assume that initially all message delays are 1. After some time, node $v_1$ will start to speed up $v_2$, and after some more time $v_2$ will speed up $v_3$, and so on. At some point of time, we will have a clock skew of 1 between any two neighbors. In particular $t_1 = t_D + D - 1$.

Now we start playing around with the message delays. Let $t_1 = T$. First we set the delay between the $v_1$ and $v_2$ to 0. Now node $v_2$ immediately adjusts its logical clock to $T$. After this event (which is instantaneous in our model) we set the delay between $v_2$ and $v_3$ to 0, which results in $v_3$ setting its logical clock to $T$ as well. We perform this successively to all pairs of nodes until $v_{D-2}$ and $v_{D-1}$. Now node $v_{D-1}$ sets its logical clock to $T$, which indicates that the difference between the logical clocks of $v_{D-1}$ and $v_D$ is $T - (T - (D - 1)) = D - 1$. \qed
Remarks:

- The introduced examples may seem cooked-up, but examples like this exist in all networks, and for all algorithms. Indeed, it was shown that any natural clock synchronization algorithm must have a bad local skew. In particular, a protocol that averages between all neighbors is even worse than the introduced $\alpha$ algorithm. This algorithm has a clock skew of $\Omega(D^2)$ in the linked list, at all times.

- Recently, there was a lot of progress in this area, and it was shown that the local clock skew is $\Theta(\log D)$, i.e., there is a protocol that achieves this bound, and there proof that no algorithm can be better than this bound!

- Note that these are worst-case bounds. In practice, clock drift and message delays may not be the worst possible, typically the speed of hardware clocks changes at a comparatively slow pace and the message transmission times follow a benign probability distribution. If we assume this, better protocols do exist.

Chapter Notes

The idea behind synchronizers is quite intuitive and as such, synchronizers $\alpha$ and $\beta$ were implicitly used in various asynchronous algorithms [Gal76, Cha79, CL85] before being proposed as separate entities. The general idea of applying synchronizers to run synchronous algorithms in asynchronous networks was first introduced by Awerbuch [Awe85a]. His work also formally introduced the synchronizers $\alpha$ and $\beta$, whereas other constructions were presented in [AP90, PU87].

Naturally, as synchronizers are motivated by practical difficulties with local clocks, there are plenty of real life applications. Studies regarding applications can be found in, e.g., [SM86, Awe85b, LTC89, AP90, PU87]. Synchronizers in the presence of network failures have been discussed in [AP88, HS94].

It has been known for a long time that the global clock skew is $\Theta(D)$ [LL84, ST87]. The problem of synchronizing the clocks of nearby nodes was introduced by Fan and Lynch in [LF04]; they proved a surprising lower bound of $\Omega(\log D / \log \log D)$ for the local skew. The first algorithm providing a non-trivial local skew of $O(\sqrt{D})$ was given in [LW06]. Later, matching upper and lower bounds of $\Theta(\log D)$ were given in [LLW10]. The problem has also been studied in a dynamic setting [KLO09, KLLO10].

Clock synchronization is a well-studied problem in practice, for instance regarding the global clock skew in sensor networks, e.g. [EGE02, GKS03, MKSL04, PSJ04]. One more recent line of work is focussing on the problem of minimizing the local clock skew [BvRW07, SW09, LSW09, FW10, FZTS11].

Bibliography


Chapter 11

Hard Problems

This chapter is on “hard” problems in distributed computing. In sequential computing, there are NP-hard problems which are conjectured to take exponential time. Is there something similar in distributed computing? Using flooding/echo (Algorithms 11,12) from Chapter 3, everything so far was solvable basically in $O(D)$ time, where $D$ is the diameter of the network.

11.1 Diameter & APSP

But how do we compute the diameter itself?? With flooding/echo, of course!

Algorithm 45 Naive Diameter Construction

1: all nodes compute their radius by synchronous flooding/echo
2: all nodes flood their radius on the constructed BFS tree
3: the maximum radius a node sees is the diameter

Remarks:

- Since all these phases only take $O(D)$ time, nodes know the diameter in $O(D)$ time, which is asymptotically optimal.

- However, there is a problem! Nodes are now involved in $n$ parallel flooding/echo operations, thus a node may have to handle many and big messages in one single time step. Although this is not strictly illegal in the message passing model, it still feels like cheating! A natural question is whether we can do the same by just sending short messages in each round.

- In Definition 1.6 of Chapter 1 we postulated that nodes should send only messages of “reasonable” size. In this chapter we strengthen the definition a bit, and require that each message should have at most $O(\log n)$ bits. This is generally enough to communicate a constant number of ID’s or values to neighbors, but not enough to communicate everything a node knows!

- A simple way to avoid large messages is to split them into small messages that are sent using several rounds. This can cause that messages are
getting delayed in some nodes but not in others. The flooding might not use edges of a BFS tree anymore! These floodings might not compute correct distances anymore! On the other hand we know that the maximal message size in Algorithm 45 is $O(n \log n)$. So we could just simulate each of these “big message” rounds by $n$ “small message” rounds using small messages. This yields a runtime of $O(nD)$ which is not desirable. A third possible approach is “starting each flooding/echo one after each other” and results in $O(nD)$ in the worst case as well.

- So let us fix above algorithm! The key idea is to arrange the flooding-echo processes in a more organized way: Start the flooding processes in a certain order and prove that at any time, each node is only involved in one flooding. This is realized in Algorithm 46.

**Definition 11.1.** $(BFS_v)$ Performing a breath first search at node $v$ produces spanning tree $BFS_v$ (see Chapter 3). This takes time $O(D)$ using small messages.

**Remarks:**

- A spanning tree of a graph $G$ can be traversed in time $O(n)$ by sending a pebble over an edge in each time slot.

- This can be done using e.g. a depth first search (DFS): Start at the root of a tree, recursively visit all nodes in the following way. If the current node still has an unvisited child, then the pebble always visit that child first. Return to the parent only when all children have been visited.

- Algorithm 46 works as follows: Given a graph $G$, first a leader $l$ computes its BFS tree $BFS_l$. Then we send a pebble $P$ to traverse tree $BFS_l$. Each time pebble $P$ enters a node $v$ for the first time, $P$ waits one time slot, and then starts a breath first search (BFS) – using edges in $G$ – from $v$ with the aim of computing the distances from $v$ to all other nodes. Since we start a $BFS_v$ from every node $v$, each node $u$ learns its distance to all these nodes $v$ during the according execution of $BFS_v$. There is no need for a echo-process at the end of $BFS_u$.

**Algorithm 46** Computes APSP on $G$.

1. Assume we have a leader node $l$ (if not, compute one first)
2. **compute** $BFS_l$ of leader $l$
3. **send** a pebble $P$ to traverse $BFS_l$ in a DFS way;
4. **while** $P$ traverses $BFS_l$ **do**
5. **if** $P$ visits a new node $v$ **then**
6. **wait** one time slot; // avoid congestion
7. **start** $BFS_v$ from node $v$; // compute all distances to $v$
8. // the depth of node $u$ in $BFS_v$ is $d(u,v)$
9. **end if**
10. **end while**
11.2. LOWER BOUND GRAPHS

Remarks:

• Having all distances is nice, but how do we get the diameter? Well, as before, each node could just floods its radius (its maximum distance) into the network. However, messages are small now and we need to modify this slightly. In each round a node only sends the maximal distance that it is aware of to its neighbors. After $D$ rounds each node will know the maximum distance among all nodes.

Lemma 11.2. In Algorithm 46, at no time a node $w$ is simultaneously active for both BFS$_u$ and BFS$_v$.

Proof. Assume a BFS$_u$ is started at time $t_u$ at node $u$. Then node $w$ will be involved in BFS$_u$ at time $t_u + d(u, w)$. Now, consider a node $v$ whose BFS$_v$ is started at time $t_v > t_u$. According to the algorithm this implies that the pebble visits $v$ after $u$ and took some time to travel from $u$ to $v$. In particular, the time to get from $u$ to $v$ is at least $d(u, v)$, in addition at least node $v$ is visited for the first time (which involves waiting at least one time slot), and we have $t_v \geq t_u + d(u, v) + 1$. Using this and the triangle inequality, we get that node $w$ is involved in BFS$_v$ strictly after being involved in BFS$_u$ since $t_v + d(v, w) \geq (t_u + d(u, v) + 1) + d(v, w) \geq t_u + d(u, v) + 1 > t_u + d(u, w)$.

Theorem 11.3. Algorithm 46 computes APSP (all pairs shortest path) in time $O(n)$.

Proof. Since the previous lemma holds for any pair of vertices, no two BFS “interfere” with each other, i.e. all messages can be sent on time without congestion. Hence, all BFS stop at most $D$ time slots after they were started. We conclude that the runtime of the algorithm is determined by the time $O(D)$ we need to build tree BFS$_t$, plus the time $O(n)$ that $P$ needs to traverse BFS$_t$, plus the time $O(D)$ needed by the last BFS that $P$ initiated. Since $D \leq n$, this is all in $O(n)$.

Remarks:

• All of a sudden our algorithm needs $O(n)$ time, and possibly $n \gg D$. We should be able to do better, right?!

• Unfortunately not! One can show that computing the diameter of a network needs $\Omega(n/\log n)$ time.

• On the other hand we can check fast whether a graph has diameter 1 or not: each node just checks whether its degree is $n - 1$ and tells the result to its neighbors.

11.2 Lower Bound Graphs

We define a family $G$ of graphs that we use to prove a lower bound on the rounds needed to compute the diameter. To simplify our analysis, we assume that $(n - 2)$ can be divided by 8. We start by defining four sets of nodes, each consisting of $q = q(n) := (n - 2)/4$ nodes. Throughout this chapter we write $[q]$ as a short version of $\{1, \ldots, q\}$ and define:
118 CHAPTER 11. HARD PROBLEMS

\[
L_0 := \{l_i \mid i \in [q]\} \quad \text{// upper left in Figure 11.1}
\]

\[
L_1 := \{l'_i \mid i \in [q]\} \quad \text{// lower left}
\]

\[
R_0 := \{r_i \mid i \in [q]\} \quad \text{// upper right}
\]

\[
R_1 := \{r'_i \mid i \in [q]\} \quad \text{// lower right}
\]

Figure 11.1: The above skeleton \(G'\) contains \(n = 10\) nodes, such that \(q = 2\).

We add node \(c_L\) and connect it to all nodes in \(L_0\) and \(L_1\). Then we add node \(c_R\), connected to all nodes in \(R_0\) and \(R_1\). Furthermore, nodes \(c_L\) and \(c_R\) are connected by an edge. For \(i \in [q]\) we connect \(l_i\) to \(r_i\) and \(l'_i\) to \(r'_i\). Also we add edges such that nodes in \(L_0\) are a clique, nodes in \(L_1\) are a clique, nodes in \(R_0\) are a clique, and nodes in \(R_1\) are a clique. The resulting graph is called \(G'\). Graph \(G'\) is the skeleton of any graph in family \(G\).

More formally skeleton \(G' = (V', E')\) is:

\[
V' := L_0 \cup L_1 \cup R_0 \cup R_1 \cup \{c_L, c_R\}
\]

\[
E' := \bigcup_{v \in L_0 \cup L_1} \{(v, c_L)\} \quad \text{// connections to } c_L
\]

\[
\bigcup_{v \in R_0 \cup R_1} \{(v, c_R)\} \quad \text{// connections to } c_R
\]

\[
\bigcup_{i \in [q]} \{(l_i, r_i), (l'_i, r'_i)\} \cup \{(c_L, c_R)\} \quad \text{// connects left to right}
\]

\[
\bigcup_{S \in \{L_0, L_1, R_0, R_1\}} \bigcup_{u \neq v \in S} \{(u, v)\} \quad \text{// clique edges}
\]

To simplify our arguments, we partition \(G'\) into two parts: Part \(L\) is the subgraph induced by nodes \(L_0 \cup L_1 \cup \{c_L\}\). Part \(R\) is the subgraph induced by nodes \(R_0 \cup R_1 \cup \{c_R\}\).
Family \( \mathcal{G} \) contains any graph \( G \) that is derived from \( G' \) by adding any combination of edges of the form \((l_i, l'_j)\) resp. \((r_i, r'_j)\) with \( l_i \in \mathbf{L}_0, l'_j \in \mathbf{L}_1, r_i \in \mathbf{R}_0, \) and \( r'_j \in \mathbf{R}_1. \)

**Figure 11.2**: The above graph \( G \) has \( n = 10 \) and a member of family \( \mathcal{G} \). What is the diameter of \( G \)?

**Lemma 11.4.** The diameter of a graph \( G = (V,E) \in \mathcal{G} \) is 2 if and only if: For each tuple \((i,j)\) with \( i,j \in [q], \) there is either edge \((l_i, l'_j)\) or edge \((r_i, r'_j)\) (or both edges) in \( E. \)

**Proof.** Note that the distance between most pairs of nodes is at most 2. In particular, the radius of \( c_L \) resp. \( c_R \) is 2. Thanks to \( c_L \) resp. \( c_R \) the distance between any two nodes within **Part L** resp. within **Part R** is at most 2. Because of the cliques \( \mathbf{L}_0, \mathbf{L}_1, \mathbf{R}_0, \mathbf{R}_1, \) distances between \( l_i \) and \( r_j \) resp. \( l'_i \) and \( r'_j \) is at most 2.

The only interesting case is between a node \( l_i \in \mathbf{L}_0 \) and node \( r'_j \in \mathbf{R}_1 \) (or, symmetrically, between \( l'_j \in \mathbf{L}_1 \) and node \( r_i \in \mathbf{R}_0. \)) If either edge \((l_i, l'_j)\) or edge \((r_i, r'_j)\) is present, then this distance is 2, since the path \((l_i, l'_j, r'_j)\) or the path \((l_i, r_i, r'_j)\) exists. If neither of the two edges exist, then the neighborhood of \( l_i \) consists of \( \{c_L, r_i\} \), all nodes in \( \mathbf{L}_0 \), and some nodes in \( \mathbf{L}_1 \setminus \{l'_j\} \), and the neighborhood of \( r'_j \) consists of \( \{c_R, l'_j\} \), all nodes in \( \mathbf{R}_1 \), and some nodes in \( \mathbf{R}_0 \setminus \{r_i\} \) (see for example Figure 11.3 with \( i = 1 \) and \( j = 2 \).) Since the two neighborhoods do not share a common node, the distance between \( l_i \) and \( r'_j \) is (at least) 3.

**Remarks:**

- Each part contains up to \( q^2 \in \Theta(n^2) \) edges.
- There are \( 2q + 1 \in \Theta(n) \) edges connecting the left and the right part.

Since in each round we can transmit \( O(\log n) \) bits over each edge (in each direction), the bandwidth between **Part L** and **Part R** is \( O(n \log n). \)
Figure 11.3: Nodes in the neighborhood of \( l_2 \) are cyan, the neighborhood of \( r_2' \) is white. Since these neighborhoods do not intersect, the distance of these two nodes is \( d(l_2, r_2') > 2 \). If e.g. edge \((l_2, l_2')\) was included, their distance was 2.

- If we transmit the information of the \( \Theta(n^2) \) edges in a naive way with a bandwidth of \( O(n \log n) \), we need \( \Omega(n/\log n) \) time. But maybe we can do better?!? Can an algorithm be smarter and only send the information that is really necessary to tell whether the diameter is 2?

- It turns out that any algorithm needs \( \Omega(n/\log n) \) rounds, since the information that is really necessary to tell that the diameter is larger than 2 contains basically \( \Theta(n^6) \) bits.

### 11.3 Communication Complexity

To prove the last remark formally, we can use arguments from two-party communication complexity. This area essentially deals with a basic version of distributed computation: two parties are given some input each and want to solve a task on this input.

We consider two students (Alice and Bob) at two different universities connected by a communication channel (e.g. via email) and we assume this channel to be reliable. Now Alice and Bob want to check whether they received the same problem set for homework (we assume their professors are lazy and wrote it on the black board instead of putting a nicely prepared document online.) Do Alice and Bob really need to type the whole problem set into their emails? In a more formal way: Alice receives an \( n \)-bit string \( x \) and Bob another \( n \)-bit string \( y \), and the goal is for both of them to compute the equality function.

**Definition 11.5.** (Equality.) We define the equality function \( \text{EQ} \) to be:

\[
\text{EQ}(x, y) := \begin{cases} 
1 & : x = y \\
0 & : x \neq y 
\end{cases}
\]
11.3. COMMUNICATION COMPLEXITY

Remarks:

- In a more general setting, Alice and Bob are interested in computing a certain function $f : \{0,1\}^k \times \{0,1\}^k \rightarrow \{0,1\}$ with the least amount of communication between them. Of course they can always succeed by having Alice send her whole $n$-bit string to Bob, who then computes the function, but the idea here is to find clever ways of calculating $f$ with less than $n$ bits of communication. We measure how clever they can be as follows:

**Definition 11.6.** (Communication complexity $CC$.) The communication complexity of protocol $A$ for function $f$ is $CC(A,f) := \text{minimum number of bits exchanged between Alice and Bob in the worst case when using } A$. The communication complexity of $f$ is $CC(f) := \min\{CC(A,f) \mid A \text{ solves } f\}$. That is the minimal number of bits that the best protocol needs to send in the worst case.

**Definition 11.7.** For a given function $f$, we define a $2^k \times 2^k$ matrix $M_f$ representing $f$. That is $M_f(x,y) := f(x,y)$.

**Example 11.8.** For $EQ$, in case $k = 3$, matrix $M^EQ$ looks like this:

$$
\begin{array}{cccccccc}
\text{EQ} & 000 & 001 & 010 & 011 & 100 & 101 & 110 & 111 \\
000 & 1 & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\
001 & 0 & 1 & 0 & 0 & 0 & 0 & 0 & 0 \\
010 & 0 & 0 & 1 & 0 & 0 & 0 & 0 & 0 \\
011 & 0 & 0 & 0 & 1 & 0 & 0 & 0 & 0 \\
100 & 0 & 0 & 0 & 0 & 1 & 0 & 0 & 0 \\
101 & 0 & 0 & 0 & 0 & 0 & 1 & 0 & 0 \\
110 & 0 & 0 & 0 & 0 & 0 & 0 & 1 & 0 \\
111 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 1 \\
\end{array}
$$

As a next step we define a (combinatorial) monochromatic rectangle. These are “submatrices” of $M^f$ which contain the same entry.

**Definition 11.9.** (Monochromatic rectangle.) A set $R \subseteq \{0,1\}^k \times \{0,1\}^k$ is called a monochromatic rectangle, if

- whenever $(x_1, y_1) \in R$ and $(x_2, y_2) \in R$ then $(x_1, y_2) \in R$.
- there is a fixed $z$ such that $f(x,y) = z$ for all $(x,y) \in R$.

**Example 11.10.** The first three of the following rectangles are monochromatic, the last one is not:

$R_1 = \{011\} \times \{011\}$

$R_2 = \{011, 100, 101, 110\} \times \{000, 001\}$

$R_3 = \{000, 001, 101\} \times \{011, 100, 110, 111\}$

$R_4 = \{000, 001\} \times \{000, 001\}$

Each time Alice and Bob exchange a bit, they can eliminate columns/rows of the matrix $M^f$ and a combinatorial rectangle is left. They can stop communicating when this remaining rectangle is monochromatic. Informally speaking, a fooling set can be used to fool a protocol that wants to be lazy: if the fooling
CHAPTER 11. HARD PROBLEMS

set is large, there will be many maximal monochromatic rectangles (maximal in the sense that they cannot be extended while staying monochromatic). Since by communicating one bit the set of possible monochromatic rectangles does not shrink too much, we can expect that it takes long time until a monochromatic rectangle is found in the worst case.

Definition 11.11. (fooling set.) A set $S \subset \{0,1\}^k \times \{0,1\}^k$ fools $f$ if there is a fixed $z$ such that

- $f(x,y) = z$ for each $(x,y) \in S$
- For any $(x_1,y_1) \neq (x_2,y_2) \in S$, the rectangle $\{x_1,x_2\} \times \{y_1,y_2\}$ is not monochromatic: Either $f(x_1,y_2) \neq z$, $f(x_2,y_1) \neq z$ or both $\neq z$.

Example 11.12. Consider $S = \{(000,000),(001,001)\}$. Take a look at the non-monochromatic rectangle $R_4$ in Example 11.10. Verify that $S$ is indeed a fooling set for $EQ$!

Remarks:

- Can you find a larger fooling set for $EQ$?
- We assume that Alice and Bob take turns in sending a bit. This results in $2$ possible actions (send 0/1) per round and in $2^t$ action patterns during a sequence of $t$ rounds.

Lemma 11.13. If $S$ is a fooling set for $f$, then $\text{CC}(f) = \Omega(\log |S|)$.

Proof. For simplicity we assume that $|S|$ is a power of 2. We prove the statement via contradiction: fix a protocol $A$ and assume that it needs $t := \log_2(|S|/2)$ rounds in the worst case. Then there are $2^t = 2^{\log_2(|S|/9)} = |S|/2$ possible action patterns. On the other hand there are $|S| = 2^{\log_2 |S|}$ elements in $S$ and we conclude that at least two elements (let’s call them $(x_1,y_1),(x_2,y_2)$) in $S$ cause the same action pattern $P$. Naturally, the action pattern on the alternative inputs $(x_1,y_2),(x_2,y_1)$ will be $P$ as well: in the first round Alice and Bob have no information on the other party’s string and send the same bit that was sent in $P$. Based on this, they determine the second bit to be exchanged, which will be the same as the second one in $P$ for a similar reason. This continues for all $t$ rounds. We conclude that after $t$ rounds, Alice does not know whether Bob’s input is $y_1$ or $y_2$ and Bob does not know whether Alice’s input is $x_1$ or $x_2$. By the definition of fooling set, either

- $f(x_1,y_2) \neq f(x_1,y_1)$ in which case Alice (with input $x_1$) does not know the solution yet,

or

- $f(x_2,y_1) \neq f(x_1,y_1)$ in which case Bob (with input $y_1$) does not know the solution yet.

This contradicts the assumption that $A$ leads to a correct decision for all inputs after $t$ rounds. Therefore at least $t + 1$ rounds are necessary, which is

$$t + 1 = \log_2(|S|/2) + 1 = \frac{\log |S|}{\log 2} \in \Omega(\log |S|).$$
11.3. COMMUNICATION COMPLEXITY

**Theorem 11.14.** \( CC(EQ) = \Omega(k) \).

*Proof.* The set \( S := \{ (x, x) \mid x \in \{0, 1\}^k \} \) fools \( EQ \) and has size \( 2^k \). Now apply Lemma 11.13. \( \square \)

**Definition 11.15.** Denote the negation of a string \( z \) by \( \overline{z} \) and by \( x \circ y \) the concatenation of strings \( x \) and \( y \).

**Lemma 11.16.** Let \( x, y \) be \( k \)-bit strings. Then \( x \neq y \) if and only if there is an index \( i \in [2k] \) such that the \( i^{th} \) bit of \( x \circ \overline{x} \) and the \( i^{th} \) bit of \( \overline{y} \circ y \) are both 0.

*Proof.* If \( x \neq y \), there is an \( j \in [k] \) such that \( x \) and \( y \) differ in the \( j^{th} \) bit. Therefore either the \( j^{th} \) bit of both \( x \) and \( \overline{y} \) is 0, or the \( j^{th} \) bit of \( \overline{y} \) and \( y \) is 0. For this reason, there is an \( i \in [2k] \) such that \( x \circ \overline{x} \) and \( \overline{y} \circ y \) are both 0 at position \( i \).

If \( x = y \), then for any \( i \in [2k] \) it is always the case that either the \( i^{th} \) bit of \( x \circ \overline{x} \) is 1 or the \( i^{th} \) bit of \( \overline{y} \circ y \) (which is the negation of \( x \circ \overline{x} \) in this case) is 1. \( \square \)

**Remarks:**

- With these insights we get back to the problem of computing the diameter of a graph and relate this problem to \( EQ \).

**Definition 11.17.** Using the parameter \( q \) defined before, we define a bijective map between all pairs \( x, y \) of \( q^2 \)-bit strings and the graphs in \( \mathcal{G} \): each pair of strings \( x, y \) is mapped to graph \( G_{x,y} \in \mathcal{G} \) that is derived from skeleton \( G' \) by adding

  - edge \( (l_i, l'_j) \) to **Part L** if and only if the \((j + q \cdot i)^{th}\) bit of \( x \) is 1.
  - edge \( (r_i, r'_j) \) to **Part R** if and only if the \((j + q \cdot i)^{th}\) bit of \( y \) is 1.

**Remarks:**

- Clearly, **Part L** of \( G_{x,y} \) depends on \( x \) only and **Part R** depends on \( y \) only.

**Lemma 11.18.** Let \( x \) and \( y \) be \( q^2 \)-bit strings given to Alice and Bob\(^1\). Then graph \( G := G_{x \circ \overline{x}, y \circ \overline{y}} \in \mathcal{G} \) has diameter 2 if and only if \( x = y \).

*Proof.* By Lemma 11.16 and the construction of \( G \), there is neither edge \( (l_i, l'_j) \) nor edge \( (r_i, r'_j) \) in \( E \) if and only if \( x \neq y \). Applying Lemma 11.4 yields: \( G \) has diameter 2 if and only if \( x = y \). \( \square \)

**Theorem 11.19.** Any distributed algorithm \( A \) that decides whether a graph \( G \) has diameter 2 might need \( \Omega \left( \frac{n}{\log n} + D \right) \) time.

\(^1\)That's why we need that \( n - 2 \) can be divided by 8.
Proof. Computing $D$ for sure needs time $\Omega(D)$. It remains to prove $\Omega\left(\frac{n}{\log n}\right)$. Assume there is a distributed algorithm $A$ that decides whether the diameter of a graph is 2 in time $o(n/\log n)$. When Alice and Bob are given $\frac{k}{2}$-bit inputs $x$ and $y$, they can simulate $A$ to decide whether $x = y$ as follows: Alice constructs Part $L$ of $G_{x \circ x, y \circ y}$ and Bob constructs Part $R$. As we remarked, both parts are independent of each other such that Part $L$ can be constructed by Alice without knowing $y$ and Part $R$ can be constructed by Bob without knowing $x$. Furthermore, $G_{x \circ x, y \circ y}$ has diameter 2 if and only if $x = y$ (Lemma 11.18.)

Now Alice and Bob simulate the distributed algorithm $A$ round by round: In the first round, they determine which messages the nodes in their part of $G$ would send. Then they use their communication channel to exchange all $2(2^q + 1) \in \Theta(n)$ messages that would be sent over edges between Part $L$ and Part $R$ in this round while executing $A$ on $G$. Based on this Alice and Bob determine which messages would be sent in round two and so on. For each round simulated by Alice and Bob, they need to communicate $\Theta(\frac{n}{\log n})$ bits: possibly $\Theta(\log n)$ bits for each of $\Theta(n)$ messages. Since $A$ makes a decision after $o(n/\log n)$ rounds, this yields a total communication of $o(n^2)$ bits. On the other hand, Lemma 11.14 states that to decide whether $x$ equals $y$, Alice and Bob need to communicate at least $\Omega\left(\frac{n^2}{2}\right) = \Omega(n^2)$ bits. A contradiction. \hfill $\square$

Remarks:

- Until now we only considered deterministic algorithms. Can one do better using randomness?

Algorithm 47 Randomized evaluation of $EQ$.

1: Alice and Bob use public randomness. That is they both have access to the same random bit string $z \in \{0, 1\}^k$
2: Alice sends bit $a := \sum_{i \in [k]} x_i \cdot z_i \mod 2$ to Bob
3: Bob sends bit $b := \sum_{i \in [k]} y_i \cdot z_i \mod 2$ to Alice
4: if $a \neq b$ then
5: we know $x \neq y$
6: end if

Lemma 11.20. If $x \neq y$, Algorithm 47 discovers $x \neq y$ with probability at least $1/2$.

Proof. Note that if $x = y$ we have $a = b$ for sure.

If $x \neq y$, Algorithm 47 may not reveal inequality. For instance, for $k = 2$, if $x = 01$, $y = 10$ and $z = 11$ we get $a = b = 1$. In general, let $I$ be the set of indices where $x_i \neq y_i$, i.e. $I := \{i \in [k] \mid x_i \neq y_i\}$. Since $x \neq y$, we know that $|I| > 0$. We have

$$|a - b| \equiv \sum_{i \in I} z_i \pmod{2},$$

and since all $z_i$ with $i \in I$ are random, we get that $a \neq b$ with probability at least $1/2$. \hfill $\square$
Remarks:

- By excluding the vector $z = 0^k$ we can even get a discovery probability strictly larger than $1/2$.

- Repeating the Algorithm 47 with different random strings $z$, the error probability can be reduced arbitrarily.

- Does this imply that there is a fast randomized algorithm to determine the diameter? Unfortunately not!

- Sometimes public randomness is not available, but private randomness is. Here Alice has her own random string and Bob has his own random string. A modified version of Algorithm 47 also works with private randomness at the cost of the runtime.

- One can prove an $\Omega(n/\log n)$ lower bound for any randomized distributed algorithm that computes the diameter. To do so one considers the disjointness function $DISJ$ instead of equality. Here, Alice is given a subset $X \subseteq [k]$ and and Bob is given a subset $Y \subseteq [k]$ and they need to determine whether $Y \cap X = \emptyset$. ($X$ and $Y$ can be represented by $k$-bit strings $x, y$.) The reduction is similar as the one presented above but uses graph $G_{x \circ y}$ instead of $G_{x \cup y}$. However, the lower bound for the randomized communication complexity of $DISJ$ is more involved than the lower bound for $CC(EQ)$.

- Since one can compute the diameter given a solution for APSP, an $\Omega(n/\log n)$ lower bound for APSP is implied. As such, our simple Algorithm 46 is almost optimal!

- Many prominent functions allow for a low communication complexity. For instance, $CC(PARITY) = 2$. What is the Hamming distance (number of different entries) of two strings? It is known that $CC(HAM \geq d) = \Omega(d)$. Also, $CC(\text{decide whether } "HAM \geq k/2 + \sqrt{k}" \text{ or } "HAM \leq k/2 - \sqrt{k}") = \Omega(k)$, even when using randomness. This problem is known as the Gap-Hamming-Distance.

- Lower bounds in communication complexity have many applications. Apart from getting lower bounds in distributed computing, one can also get lower bounds regarding circuit depth or query times for static data structures.

- In the distributed setting with limited bandwidth we showed that computing the diameter has about the same complexity as computing all pairs shortest paths. In contrast, in sequential computing, it is a major open problem whether the diameter can be computed faster than all pairs shortest paths. No nontrivial lower bounds are known, only that $\Omega(n^2)$ steps are needed – partly due to the fact that there can be $n^2$ edges/distances in a graph. On the other hand the currently best algorithm uses fast matrix multiplication and terminates after $O(n^{2.3727})$ steps.
CHAPTER 11. HARD PROBLEMS

11.4 Distributed Complexity Theory

We conclude this chapter with a short overview on the main complexity classes of distributed message passing algorithms. Given a network with \( n \) nodes and diameter \( D \), we managed to establish a rich selection of upper and lower bounds regarding how much time it takes to solve or approximate a problem. Currently we know five main distributed complexity classes:

- **Strictly local** problems can be solved in constant \( O(1) \) time, e.g. a constant approximation of a dominating set in a planar graph.

- Just a little bit slower are problems that can be solved in \( \log^* n \) time, e.g. many combinatorial optimization problems in special graph classes such as growth bounded graphs. 3-coloring a ring takes \( O(\log^* n) \).

- A large body of problems is polylogarithmic (or pseudo-local), in the sense that they seem to be strictly local but are not, as they need \( O(\text{polylog } n) \) time, e.g. the maximal independent set problem.

- There are problems which are global and need \( O(D) \) time, e.g. to count the number of nodes in the network.

- Finally there are problems which need polynomial \( O(\text{poly } n) \) time, even if the diameter \( D \) is a constant, e.g. computing the diameter of the network.

Chapter Notes

The linear time algorithm for computing the diameter was discovered independently by [HW12, PRT12]. The presented matching lower bound is by Frischknecht et al. [FHW12], extending techniques by [DHK++11].

Due to its importance in network design, shortest path-problems in general and the APSP problem in particular were among the earliest studied problems in distributed computing. Developed algorithms were immediately used e.g. as early as in 1969 in the ARPANET (see [Lyn96], p.506). Routing messages via shortest paths were extensively discussed to be beneficial in [Ta77, MS79, MRR80, SS80, CM82] and in many other papers. It is not surprising that there is plenty of literature dealing with algorithms for distributed APSP, but most of them focused on secondary targets such as trading time for message complexity. E.g. papers [AR78, Tou80, Che82] obtain a communication complexity of roughly \( O(n \cdot m) \) bits/messages and still require superlinear runtime. Also a lot of effort was spent to obtain fast sequential algorithms for various versions of computing APSP or related problems such as the diameter problem, e.g. [CW90, AGM91, AMGN92, Sei95, SZ99, BVW08]. These algorithms are based on fast matrix multiplication such that currently the best runtime is \( O(n^{2.3727}) \) due to [Wil12].

The problem sets in which one needs to distinguish diameter 2 from 4 are inspired by a combinatorial \( (\times, 3/2) \)-approximation in a sequential setting by Aingworth et. al. [ACIM99]. The main idea behind this approximation is to distinguish diameter 2 from 4. This part was transferred to the distributed setting in [HW12].
Two-party communication complexity was introduced by Andy Yao in [Yao79]. Later, Yao received the Turing Award. A nice introduction to communication complexity covering techniques such as fooling-sets is the book by Nisan and Kushilevitz [KN97].

This chapter was written in collaboration with Stephan Holzer.

Bibliography


Chapter 12

Stabilization

A large branch of research in distributed computing deals with fault-tolerance. Being able to tolerate a considerable fraction of failing or even maliciously behaving ("Byzantine") nodes while trying to reach consensus (on e.g. the output of a function) among the nodes that work properly is crucial for building reliable systems. However, consensus protocols require that a majority of the nodes remains non-faulty all the time.

Can we design a distributed system that survives transient (short-lived) failures, even if all nodes are temporarily failing? In other words, can we build a distributed system that repairs itself?

12.1 Self-Stabilization

Definition 12.1 (Self-Stabilization). A distributed system is self-stabilizing if, starting from an arbitrary state, it is guaranteed to converge to a legitimate state. If the system is in a legitimate state, it is guaranteed to remain there, provided that no further faults happen. A state is legitimate if the state satisfies the specifications of the distributed system.

Remarks:

- What kind of transient failures can we tolerate? An adversary can crash nodes, or make nodes behave Byzantine. Indeed, temporarily an adversary can do harm in even worse ways, e.g. by corrupting the volatile memory of a node (without the node noticing – not unlike the movie Memento), or by corrupting messages on the fly (without anybody noticing). However, as all failures are transient, eventually all nodes must work correctly again, that is, crashed nodes get resurrected, Byzantine nodes stop being malicious, messages are being delivered reliably, and the memory of the nodes is secure.

- Clearly, the read only memory (ROM) must be taboo at all times for the adversary. No system can repair itself if the program code itself or constants are corrupted. The adversary can only corrupt the variables in the volatile random access memory (RAM).
Chapter 12: Stabilization

Definition 12.2 (Time Complexity). The time complexity of a self-stabilizing system is the time that passed after the last (transient) failure until the system has converged to a legitimate state again, staying legitimate.

Remarks:

• Self-stabilization enables a distributed system to recover from a transient fault regardless of its nature. A self-stabilizing system does not have to be initialized as it eventually (after convergence) will behave correctly.

• One of the first self-stabilizing algorithms was Dijkstra’s token ring network. A token ring is an early form of a local area network where nodes are arranged in a ring, communicating by a token. The system is correct if there is exactly one token in the ring. Let’s have a look at a simple solution. Given an oriented ring, we simply call the clockwise neighbor parent \(p\), and the counterclockwise neighbor child \(c\). Also, there is a leader node \(v_0\). Every node \(v\) is in a state \(S(v) \in \{0, 1, \ldots, n\}\), perpetually informing its child about its state. The token is implicitly passed on by nodes switching state. Upon noticing a change of the parent state \(S(p)\), node \(v\) executes the following code:

Algorithm 48 Self-stabilizing Token Ring

```plaintext
1: if \(v = v_0\) then
2:   if \(S(v) = S(p)\) then
3:     \(S(v) := S(v) + 1 \pmod{n}\)
4:   end if
5: else
6:   \(S(v) := S(p)\)
7: end if
```

Theorem 12.3. Algorithm 48 stabilizes correctly.

Proof: As long as some nodes or edges are faulty, anything can happen. In self-stabilization, we only consider the system after it is correct (at time \(t_0\), however starting in an arbitrary state).

Every node apart from leader \(v_0\) will always attain the state of its parent. It may happen that one node after the other will learn the current state of the leader. In this case the system stabilizes after the leader increases its state at most \(n\) time units after time \(t_0\). It may however be that the leader increases its state even if the system is not stable, e.g. because its parent or parent’s parent accidentally had the same state at time \(t_0\).

The leader will increase its state possibly multiple times without reaching stability, however, at some point the leader will reach state \(s\), a state that no other node had at time \(t_0\). (Since there are \(n\) nodes and \(n\) states, this will eventually happen.) At this point the system must stabilize because the leader cannot push for \(s + 1 \pmod{n}\) until every node (including its parent) has \(s\).

After stabilization, there will always be only one node changing its state, i.e., the system remains in a legitimate state.
12.1. SELF-STABILIZATION

Remarks:

- Although one might think the time complexity of the algorithm is quite bad, it is asymptotically optimal.
- It can be a lot of fun designing self-stabilizing algorithms. Let us try to build a system, where the nodes organize themselves as a maximal independent set (MIS, Chapter 7):

Algorithm 49 Self-stabilizing MIS

Require: Node IDs

Every node v executes the following code:

1: do atomically
2: Leave MIS if a neighbor with a larger ID is in the MIS
3: Join MIS if no neighbor with larger ID joins MIS
4: Send (node ID, MIS or not MIS) to all neighbors
5: end do

Remarks:

- Note that the main idea of Algorithm 49 is from Algorithm 34, Chapter 7.
- As long as some nodes are faulty, anything can happen: Faulty nodes may for instance decide to join the MIS, but report to their neighbors that they did not join the MIS. Similarly messages may be corrupted during transport. As soon as the system (nodes, messages) is correct, however, the system will converge to a MIS. (The arguments are the same as in Chapter 7).
- Self-stabilizing algorithms always run in an infinite loop, because transient failures can hit the system at any time. Without the infinite loop, an adversary can always corrupt the solution “after” the algorithm terminated.
- The problem of Algorithm 49 is its time complexity, which may be linear in the number of nodes. This is not very exciting. We need something better! Since Algorithm 49 was just the self-stabilizing variant of the slow MIS Algorithm 34, maybe we can hope to “self-stabilize” some of our fast algorithms from Chapter 7?
- Yes, we can! Indeed there is a general transformation that takes any local algorithm (efficient but not fault-tolerant) and turns it into a self-stabilizing algorithm, keeping the same level of efficiency and efficacy. We present the general transformation below.

Theorem 12.4 (Transformation). We are given a deterministic local algorithm $A$ that computes a solution of a given problem in $k$ synchronous communication rounds. Using our transformation, we get a self-stabilizing system with time complexity $k$. In other words, if the adversary does not corrupt the system for $k$ time units, the solution is stable. In addition, if the adversary does not corrupt any node or message closer than distance $k$ from a node $u$, node $u$ will be stable.
CHAPTER 12. STABILIZATION

Proof: In the proof, we present the transformation. First, however, we need to be more formal about the deterministic local algorithm $\mathcal{A}$. In $\mathcal{A}$, each node of the network computes its decision in $k$ phases. In phase $i$, node $u$ computes its local variables according to its local variables and received messages of the earlier phases. Then node $u$ sends its messages of phase $i$ to its neighbors. Finally node $u$ receives the messages of phase $i$ from its neighbors. The set of local variables of node $u$ in phase $i$ is given by $L_i^u$. (In the very first phase, node $u$ initializes its local variables with $L_1^u$.) The message sent from node $u$ to node $v$ in phase $i$ is denoted by $m_{i,u,v}$. Since the algorithm $\mathcal{A}$ is deterministic, node $u$ can compute its local variables $L_i^u$ and messages $m_{i,u,v}$ of phase $i$ from its state of earlier phases, by simply applying functions $f_L$ and $f_m$. In particular,

$$L_i^u = f_L(u, L_{i-1}^u, m_{i-1}^u), \text{ for } i > 1, \text{ and}$$

$$m_{i,u,v} = f_m(u,v, L_i^u), \text{ for } i \geq 1. \quad (12.1)$$

The self-stabilizing algorithm needs to simulate all the $k$ phases of the local algorithm $\mathcal{A}$ in parallel. Each node $u$ stores its local variables $L_1^u, \ldots, L_k^u$ as well as all messages received $m_1^u, \ldots, m_k^u$ in two tables in RAM. For simplicity, each node $u$ also stores all the sent messages $m_1^u, \ldots, m_k^u$ in a third table. If a message or a local variable for a particular phase is unknown, the entry in the table will be marked with a special value $\perp$ ("unknown"). Initially, all entries in the table are $\perp$.

Clearly, in the self-stabilizing model, an adversary can choose to change table values at all times, and even reset these values to $\perp$. Our self-stabilizing algorithm needs to constantly work against this adversary. In particular, each node $u$ runs these two procedures constantly:

- For all neighbors: Send each neighbor $v$ a message containing the complete row of messages of algorithm $\mathcal{A}$, that is, send the vector $(m_{1,v}^u, \ldots, m_{k,v}^u)$ to neighbor $v$. Similarly, if neighbor $u$ receives such a vector from neighbor $v$, then neighbor $u$ replaces neighbor $v$’s row in the table of incoming messages by the received vector $(m_{1,v}^u, \ldots, m_{k,v}^u)$.

- Because of the adversary, node $u$ must constantly recompute its local variables (including the initialization) and outgoing message vectors using Functions (12.1) and (12.2) respectively.

The proof is by induction. Let $N^i(u)$ be the $i$-neighborhood of node $u$ (that is, all nodes within distance $i$ of node $u$). We assume that the adversary has not corrupted any node in $N^k(u)$ since time $t_0$. At time $t_0$ all nodes in $N^k(u)$ will check and correct their initialization. Following Equation (12.2), at time $t_0$ all nodes in $N^k(u)$ will send the correct message entry for the first round $(m_1^u)$ to all neighbors. Asynchronous messages take at most 1 time unit to be received at a destination. Hence, using the induction with Equations (12.1) and (12.2) it follows that at time $t_0 + i$, all nodes in $N^{k-i}(u)$ have received the correct messages $m_{1}^u, \ldots, m_{k-i}^u$. Consequently, at time $t_0 + k$ node $u$ has received all messages of local algorithm $\mathcal{A}$ correctly, and will compute the same result value as in $\mathcal{A}$. \qed
Remarks:

• Using our transformation (also known as “local checking”), designing self-stabilizing algorithms just turned from art to craft.

• As we have seen, many local algorithms are randomized. This brings two additional problems. Firstly, one may not exactly know how long the algorithm will take. This is not really a problem since we can simply send around all the messages needed, until the algorithm is finished. The transformation of Theorem 12.4 works also if nodes just send all messages that are not ⊥. Secondly, we must be careful about the adversary. In particular we need to restrict the adversary such that a node can produce a reproducible sufficiently long string of random bits. This can be achieved by storing the sufficiently long string along with the program code in the read only memory (ROM). Alternatively, the algorithm might not store the random bit string in its ROM, but only the seed for a random bit generator. We need this in order to keep the adversary from reshuffling random bits until the bits become “bad”, and the expected (or with high probability) efficacy or efficiency guarantees of the original local algorithm $A$ cannot be guaranteed anymore.

• Since most local algorithms have only a few communication rounds, and only exchange small messages, the memory overhead of the transformation is usually bearable. In addition, information can often be compressed in a suitable way so that for many algorithms message size will remain polylogarithmic. For example, the information of the fast MIS algorithm (Algorithm 36) consists of a series of random values (one for each round), plus two boolean values per round. These boolean values represent whether the node joins the MIS, or whether a neighbor of the node joins the MIS. The order of the values tells in which round a decision is made. Indeed, the series of random bits can even be compressed just into the random seed value, and the neighbors can compute the random values of each round themselves.

• There is hope that our transformation as well gives good algorithms for mobile networks, that is for networks where the topology of the network may change. Indeed, for deterministic local approximation algorithms, this is true: If the adversary does not change the topology of a node’s $k$-neighborhood in time $k$, the solution will locally be stable again.

• For randomized local approximation algorithms however, this is not that simple. Assume for example, that we have a randomized local algorithm for the dominating set problem. An adversary can constantly switch the topology of the network, until it finds a topology for which the random bits (which are not really random because these random bits are in ROM) give a solution with a bad approximation ratio. By defining a weaker adversarial model, we can fix this problem. Essentially, the adversary needs to be oblivious, in the sense that it cannot see the solution. Then it will not be possible for the adversary to restart the random computation if the solution is “too good”.
• Self-stabilization is the original approach, and self-organization may be the
general theme, but new buzzwords pop up every now and then, e.g. self-
configuration, self-management, self-regulation, self-repairing, self-healing,
self-optimization, self-adaptivity, or self-protection. Generally all
these are summarized as “self-*”. One computing giant coined the term
“autonomic computing” to reflect the trend of self-managing distributed
systems.

12.2 Advanced Stabilization

We finish the chapter with a non-trivial example beyond self-stabilization, show-
ing the beauty and potential of the area: In a small town, every evening each
citizen calls all his (or her) friends, asking them whether they will vote for the
Democratic or the Republican party at the next election.\footnote{We are in the US, and as we know from The Simpsons, you “throw your vote away” if
you vote for somebody else. As a consequence our example has two parties only.}
In our town citizens
listen to their friends, and everybody re-chooses his or her affiliation according
to the majority of friends.\footnote{Assume for the sake of simplicity that everybody has an odd number of friends.} Is this process going to “stabilize” (in one way or
another)?

Remarks:

• Is eventually everybody voting for the same party? No.
• Will each citizen eventually stay with the same party? No.
• Will citizens that stayed with the same party for some time, stay with
that party forever? No.
• And if their friends also constantly root for the same party? No.
• Will this beast stabilize at all?!? Yes!

Theorem 12.5 (Dems & Reps). Eventually every citizen is rooting for the
same party every other day.

Proof: To prove that the opinions eventually become fixed or cycle every other
day, think of each friendship between citizens as a pair of (directed) edges, one
in each direction. Let us say an edge is currently “bad” if the party of the
advising friend differs from the next-day’s party of the advised friend. In other
words, the edge is bad if the advised friend did not follow the advisor’s opinion
(which means that the advisor was in the minority). An edge that is not bad,
is “good”.

Consider the out-edges of citizen $c$ on day $t$, during which (say) $c$ roots for
the Democrats. Assume that during day $t$, $g$ out-edges of $c$ are good, and $b$
out-edges are bad. Note that $g + b$ is the degree of $c$. Since $g$ out-edges were
good, $g$ friends of $c$ root for the Democrats on day $t + 1$. Likewise, $b$ friends of $c$
root for the Republicans on day $t + 1$. In other words, on the evening of day $t + 1$
citizen $c$ will receive $g$ recommendations for Democrats, and $b$ for Republicans.
We distinguish two cases:
12.2. ADVANCED STABILIZATION

• $g > b$: In this case, citizen $c$ will still (or again) root for the Democrats on day $t + 2$. Note that in this case, on day $t + 1$, exactly $g$ in-edges of $c$ are good, and exactly $b$ in-edges are bad. In other words, the number of bad out-edges on day $t$ is exactly the number of bad in-edges on day $t + 1$.

• $g < b$: In this case, citizen $c$ will root for the Republicans on day $t + 2$. Note that in this case, on day $t + 1$, exactly $b$ in-edges of $c$ are good, and exactly $g$ in-edges are bad. In other words, the number of bad out-edges on day $t$ was exactly the number of good in-edges on day $t + 1$ (and vice versa). Since citizen $c$ is rooting for the Republicans, the number of bad out-edges on day $t$ was strictly larger than the number of bad in-edges on day $t + 1$.

We account for every edge as out-edge on day $t$, and as in-edge on day $t + 1$. Since in both of the above cases the number of bad edges does not increase, the total number of bad edges $B$ cannot increase. In fact, if any node switches its party from day $t$ to $t + 2$, we know that the total number of bad edges strictly decreases. But $B$ cannot decrease forever. Once $B$ hits its minimum, the system stabilizes in the sense that every citizen will either stick with his or her party forever or flip-flop every day – the system “stabilizes”.

Remarks:

• The model can be generalized considerably by, for example, adding weights to vertices (meaning some citizens’ opinions are more important than others), allowing loops (citizens who consider their own current opinions as well), allowing tie-breaking mechanisms, and even allowing different thresholds for party changes.

• How long does it take until the system stabilizes?

• Some of you may be reminded of Conway’s Game of Life: We are given an infinite two-dimensional grid of cells, each of which is in one of two possible states, dead or alive. Every cell interacts with its eight neighbors. In each round, the following transitions occur: Any live cell with fewer than two live neighbors dies, as if caused by loneliness. Any live cell with more than three live neighbors dies, as if by overcrowding. Any live cell with two or three live neighbors lives on to the next generation. Any dead cell with exactly three live neighbors is “born” and becomes a live cell. The initial pattern constitutes the “seed” of the system. The first generation is created by applying the above rules simultaneously to every cell in the seed, births and deaths happen simultaneously, and the discrete moment at which this happens is sometimes called a tick. (In other words, each generation is a pure function of the one before.) The rules continue to be applied repeatedly to create further generations. John Conway figured that these rules were enough to generate interesting situations, including “breeders” with create “guns” which in turn create “gliders”. As such Life in some sense answers an old question by John von Neumann, whether there can be a simple machine that can build copies of itself. In fact Life is Turing complete, that is, as powerful as any computer.
Chapter Notes

Self-stabilization was first introduced in a paper by Edsger W. Dijkstra in 1974 [Dij74], in the context of a token ring network. It was shown that the ring stabilizes in time $\Theta(n)$. For his work Dijkstra received the 2002 ACM PODC Influential Paper Award. Shortly after receiving the award he passed away. With Dijkstra being such an eminent person in distributed computing (e.g. concurrency, semaphores, mutual exclusion, deadlock, finding shortest paths in graphs, fault-tolerance, self-stabilization), the award was renamed Edsger W. Dijkstra Prize in Distributed Computing. In 1991 Awerbuch et al. showed that any algorithm can be modified into a self-stabilizing algorithm that stabilizes in the same time that is needed to compute the solution from scratch [APsV91].

The Republicans vs. Democrats problem was popularized by Peter Winkler, in his column “Puzzled” [Win08]. Goles et al. already proofed in [GO80] that any configuration of any such system will end up in a situation where each citizen votes for the same party every second day. Winkler additionally proofed that the time such a system takes to stabilize is bounded by $O(n^2)$. Closely related to this puzzle is the well known game of life which was described by the mathematician John Conway and made popular by Martin Gardner [Gar70], where cells can be either dead or alive and change their states according to the number of alive neighbors.

Bibliography

[APsV91] Baruch Awerbuch, Boaz Patt-shamir, and George Varghese. Self-Stabilization By Local Checking and Correction. In In Proceedings


Chapter 13

Wireless Protocols

Wireless communication was one of the major success stories of the last decades. Today, different wireless standards such as wireless local area networks (WLAN) are omnipresent. In some sense, from a distributed computing viewpoint wireless networks are quite simple, as they cannot form arbitrary network topologies. Simplistic models of wireless networks include geometric graph models such as the so-called unit disk graph. Modern models are more robust: The network graph is restricted, e.g., the total number of neighbors of a node which are not adjacent is likely to be small. This observation is hard to capture with purely geometric models, and motivates more advanced network connectivity models such as bounded growth or bounded independence.

However, on the other hand, wireless communication is also more difficult than standard message passing, as for instance nodes are not able to transmit a different message to each neighbor at the same time. And if two neighbors are transmitting at the same time, they interfere, and a node may not be able to decipher anything.

In this chapter we deal with the distributed computing principles of wireless communication: We make the simplifying assumption that all \( n \) nodes are in the communication range of each other, i.e., the network graph is a clique. Nodes share a synchronous time, in each time slot a node can decide to either transmit or receive (or sleep). However, two or more nodes transmitting in a time slot will cause interference. Transmitting nodes are never aware if there is interference because they cannot simultaneously transmit and receive.

13.1 Basics

The basic communication protocol in wireless networks is the medium access control (MAC) protocol. Unfortunately it is difficult to claim that one MAC protocol is better than another, because it all depends on the parameters, such as the network topology, the channel characteristics, or the traffic pattern. When it comes to the principles of wireless protocols, we usually want to achieve much simpler goals. One basic and important question is the following: How long does it take until one node can transmit successfully, without interference? This question is often called the wireless leader election problem (Chapter 2), with the node transmitting alone being the leader.
Clearly, we can use node IDs to solve leader election, e.g., a node with ID \(i\) transmits in time slot \(i\). However, this may be incredibly slow. There are better deterministic solutions, but by and large the best and simplest algorithms are randomized.

Throughout this chapter, we use a random variable \(X\) to denote the number of nodes transmitting in a given slot.

**Algorithm 50** Slotted Aloha

1. **Every node** \(v\) executes the following code:
2. **repeat**
3. transmit with probability \(1/n\)
4. **until** one node has transmitted alone

**Theorem 13.1.** Using Algorithm 50 allows one node to transmit alone (become a leader) after expected time \(e\).

**Proof.** The probability for success, i.e., only one node transmitting is

\[
Pr[X = 1] = n \cdot \frac{1}{n} \cdot (1 - \frac{1}{n})^{n-1} \approx \frac{1}{e},
\]

where the last approximation is a result from Theorem 13.23 for sufficiently large \(n\). Hence, if we repeat this process \(e\) times, we can expect one success.

**Remarks:**

- The origin of the name is the ALOHAnet which was developed at the University of Hawaii.

- How does the leader know that it is the leader? One simple solution is a “distributed acknowledgment”. The nodes just continue Algorithm 50, including the ID of the leader in their transmission. So the leader learns that is the leader.

- One more problem?! Indeed, node \(v\) which managed to transmit the acknowledgment (alone) is the only remaining node which does not know that the leader knows that it is the leader. We can fix this by having the leader acknowledge \(v\)’s successful acknowledgment.

- One can also imagine an unslotted time model. In this model two messages which overlap partially will interfere and no message is received. As everything in this chapter, Algorithm 50 also works in an unslotted time model, with a factor 2 penalty, i.e., the probability for a successful transmission will drop from \(1/2\) to \(1/4\). Essentially, each slot is divided into \(t\) small time slots with \(t \to \infty\) and the nodes start a new \(t\)-slot long transmission with probability \(1/2nt\).
13.2 Initialization

Sometimes we want the $n$ nodes to have the IDs $\{1, 2, \ldots, n\}$. This process is called initialization. Initialization can for instance be used to allow the nodes to transmit one by one without any interference.

13.2.1 Non-Uniform Initialization

**Theorem 13.2.** If the nodes know $n$, we can initialize them in $O(n)$ time slots.

**Proof.** We repeatedly elect a leader using e.g., Algorithm 50. The leader gets the next free number and afterwards leaves the process. We know that this works with probability $1/e$. The expected time to finish is hence $e \cdot n$.

Remarks:

- But this algorithm requires that the nodes know $n$ in order to give them IDs from $1, \ldots, n!$ For a more realistic scenario we need a uniform algorithm, i.e, the nodes do not know $n$.

13.2.2 Uniform Initialization with CD

**Definition 13.3** (Collision Detection, CD). Two or more nodes transmitting concurrently is called interference. In a system with collision detection, a receiver can distinguish interference from nobody transmitting. In a system without collision detection, a receiver cannot distinguish the two cases.

Let us first present a high-level idea. The set of nodes is recursively partitioned into two non-empty sets, similarly to a binary tree. This is repeated recursively until a set contains only one node which gets the next free ID. Afterwards, the algorithm continues with the next set.

**Algorithm 51 RandomizedSplit($b$)**

1: Every node $v$ executes the following code:
2: repeat
3: if $b_v = b$ then
4: choose $r$ uniformly at random from $\{0, 1\}$
5: in the next two time slots:
6: transmit in slot $r$, and listen in other slot
7: end if
8: until there was at least 1 transmission in both slots
9: if $b_v = b$ then
10: $b_v := b_v + r$ {append bit $r$ to bitstring $b_v$}
11: end if
12: if some node $u$ transmitted alone in slot $r \in \{0, 1\}$ then
13: node $u$ gets ID $m$ {and becomes passive}
14: $m := m + 1$
15: else
16: RandomizedSplit($b + r$)
17: end if
CHAPTER 13. WIRELESS PROTOCOLS

Remarks:

• In line 8 the transmitting nodes need to know if they were the only one transmitting. Since we have enough time, we can do a leader election first and use a similar trick as before to ensure this.

• In line 12 we check separately for $r = 0$ and $r = 1$

Algorithm 52 Initialization with Collision Detection
1: Every node $v$ executes the following code:
2: global variable $m := 0$ \{number of already identified nodes\}
3: local variable $b_v := ‘’$ \{current bitstring of node $v$, initially empty\}
4: RandomizedSplit("")

Theorem 13.4. Algorithm 52 correctly initializes the set of nodes in $O(n)$.

Proof. A successful split is defined as a split in which both subsets are non-empty. We know that there are exactly $n - 1$ successful splits because we have a binary tree with $n$ leaves and $n - 1$ inner nodes. Let us now calculate the probability for creating two non-empty sets from a set of size $k \geq 2$ as

$$Pr[1 \leq X \leq k - 1] = 1 - Pr[X = 0] - Pr[X = k] = 1 - \frac{1}{2^k} - \frac{1}{2^k} \geq \frac{1}{2}.$$

Thus, in expectation we need $O(n)$ splits.

Remarks:

• What if we do not have collision detection?

13.2.3 Uniform Initialization without CD

Let us assume that we have a special node $\ell$ (leader) and let $S$ denote the set of nodes which want to transmit. We now split every time slot from before into two time slots and use the leader to help us distinguish between silence and noise. In the first slot every node from the set $S$ transmits, in the second slot the nodes in $S \cup \{\ell\}$ transmit. This gives the nodes sufficient information to distinguish the different cases (see Table 13.1).

| $|S| = 0$ | nodes in $S$ transmit | $\times$ | nodes in $S \cup \{\ell\}$ transmit | $\checkmark$ |
|----------|------------------------|--------|-------------------------------------|--------|
| $|S| = 1, S = \{\ell\}$ | $\checkmark$ | $\checkmark$ |
| $|S| = 1, S \neq \{\ell\}$ | $\checkmark$ | $\times$ |
| $|S| \geq 2$ | $\times$ | $\times$ |

Table 13.1: Using a leader to distinguish between noise and silence: $\times$ represents noise/silence, $\checkmark$ represents a successful transmission.
13.3. **LEADER ELECTION**

Remarks:

- As such, Algorithm 52 works also without CD, with only a factor 2 overhead.
- More generally, a leader immediately brings CD to any protocol.
- This protocol has an important real life application, for instance when checking out a shopping cart with items which have RFID tags.
- But how do we determine such a leader? And how long does it take until we are “sure” that we have one? Let us repeat the notion of with high probability.

### 13.3 Leader Election

#### 13.3.1 With High Probability

**Definition 13.5** (With High Probability). Some probabilistic event is said to occur with high probability (w.h.p.), if it happens with a probability \( p \geq 1 - \frac{1}{n^c} \), where \( c \) is a constant. The constant \( c \) may be chosen arbitrarily, but it is considered constant with respect to Big-O notation.

**Theorem 13.6.** Algorithm 50 elects a leader w.h.p. in \( O(\log n) \) time slots.

**Proof.** The probability for not electing a leader after \( c \cdot \log n \) time slots, i.e., \( c \log n \) slots without a successful transmission is

\[
\left(1 - \frac{1}{e}\right)^{c \ln n} = \left(1 - \frac{1}{e}\right)^{c' \ln n} \leq \frac{1}{e^{c' \ln n}} = \frac{1}{n^{c'}}.
\]

\[\square\]

Remarks:

- What about uniform algorithms, i.e. the number of nodes \( n \) is not known?

#### 13.3.2 Uniform Leader Election

**Algorithm 53** Uniform leader election

1: **Every node** \( v \) executes the following code:
2: **for** \( k = 1, 2, 3, \ldots \) **do**
3: **for** \( i = 1 \) **to** \( ck \) **do**
4: transmit with probability \( p := 1/2^k \)
5: if node \( v \) was the only node which transmitted **then**
6: \( v \) becomes the leader
7: **break**
8: **end if**
9: **end for**
10: **end for**
Theorem 13.7. By using Algorithm 53 it is possible to elect a leader w.h.p. in $O(\log^2 n)$ time slots if $n$ is not known.

Proof. Let us briefly describe the algorithm. The nodes transmit with probability $p = 2^{-k}$ for $ck$ time slots for $k = 1, 2, \ldots$. At first $p$ will be too high and hence there will be a lot of interference. But after $\log n$ phases, we have $k \approx \log n$ and thus the nodes transmit with probability $\approx \frac{1}{n}$. For simplicity’s sake, let us assume that $n$ is a power of 2. Using the approach outlined above, we know that after $\log n$ iterations, we have $p = \frac{1}{n}$. Theorem 13.6 yields that we can elect a leader w.h.p. in $O(\log n)$ slots. Since we have to try $\log n$ estimates until $k \approx n$, the total runtime is $O(\log^2 n)$.

Remarks:
- Note that our proposed algorithm has not used collision detection. Can we solve leader election faster in a uniform setting with collision detection?

13.3.3 Fast Leader Election with CD

Algorithm 54 Uniform leader election with CD

1: Every node $v$ executes the following code:
2: repeat
3: transmit with probability $\frac{1}{2}$
4: if at least one node transmitted then
5: all nodes that did not transmit quit the protocol
6: end if
7: until one node transmits alone

Theorem 13.8. With collision detection we can elect a leader using Algorithm 54 w.h.p. in $O(\log n)$ time slots.

Proof. The number of active nodes $k$ is monotonically decreasing and always greater than 1 which yields the correctness. A slot is called successful if at most half the active nodes transmit. We can assume that $k \geq 2$ since otherwise we would have already elected a leader. We can calculate the probability that a time slot is successful as

$$
Pr[1 \leq X \leq \left\lfloor \frac{k}{2} \right\rfloor] \geq \frac{1}{2} - Pr[X = 0] = \frac{1}{2} - \frac{1}{2^k} \geq \frac{1}{4}.
$$

Since the number of active nodes at least halves in every successful time slot, $\log n$ successful time slots are sufficient to elect a leader. Now let $Y$ be a random variable which counts the number of successful time slots after $8 \cdot c \cdot \log n$ time slots. The expected value is $E[Y] \geq 8 \cdot c \cdot \log n \cdot \frac{1}{4} \geq 2 \cdot \log n$. Since all those time slots are independent from each other, we can apply a Chernoff bound (see Theorem 13.22) with $\delta = \frac{1}{4}$ which states

$$
Pr[Y < (1 - \delta)E[Y]] \leq e^{-\frac{\delta^2}{2} E[Y]} = e^{-\frac{1}{2} \cdot 2c \log n} \leq n^{-\alpha}
$$

for any constant $\alpha$. 

13.3. LEADER ELECTION

Remarks:

• Can we be even faster?

13.3.4 Even Faster Leader Election with CD

Let us first briefly describe an algorithm for this. In the first phase the nodes transmit with probability $1/2^0, 1/2^1, 1/2^2, \ldots$ until no node transmits. This yields a first approximation on the number of nodes. Afterwards, a binary search is performed to determine an even better approximation of $n$. Finally, the third phase finds a constant approximation of $n$ using a biased random walk. The algorithm stops in any case as soon as only one node is transmitting which will become the leader.

Algorithm 55 Fast uniform leader election

1: $i := 1$
2: repeat
3: $i := 2 \cdot i$
4: transmit with probability $1/2^i$
5: until no node transmitted
{End of Phase 1}
6: $l := 2^{i-2}$
7: $u := 2^i$
8: while $l + 1 < u$ do
9: $j := \lceil \frac{l + u}{2} \rceil$
10: transmit with probability $1/2^j$
11: if no node transmitted then
12: $u := j$
13: else
14: $l := j$
15: end if
16: end while
{End of Phase 2}
17: $k := u$
18: repeat
19: transmit with probability $1/2^k$
20: if no node transmitted then
21: $k := k - 1$
22: else
23: $k := k + 1$
24: end if
25: until exactly one node transmitted

Lemma 13.9. If $j > \log n + \log \log n$, then $Pr[X > 1] \leq \frac{1}{\log n}$.

Proof. The nodes transmit with probability $1/2^j < 1/2^{\log n + \log \log n} = \frac{1}{n \log n}$. The expected number of nodes transmitting is $E[X] = \frac{n}{n \log n}$. Using Markov’s inequality (see Theorem 13.21) yields $Pr[X > 1] \leq Pr[X > E[X] \cdot \log n] \leq \frac{1}{\log n}$.
Lemma 13.10. If $j < \log n - \log \log n$, then $P[X = 0] \leq \frac{1}{n}$.

Proof. The nodes transmit with probability $1/2^j < 1/2^{\log n - \log \log n} = \frac{\log n}{n}$. Hence, the probability for a silent time slot is $(1 - \frac{\log n}{n})^n = e^{-\log n} = \frac{1}{n}$.

Corollary 13.11. If $i > 2 \log n$, then $Pr[X > 1] \leq \frac{1}{\log n}$.

Proof. This follows from Lemma 13.9 since the deviation in this corollary is even larger.

Corollary 13.12. If $i < \frac{1}{2} \log n$, then $P[X = 0] \leq \frac{1}{n}$.

Proof. This follows from Lemma 13.10 since the deviation in this corollary is even larger.

Lemma 13.13. Let $v$ be such that $2^{v-1} < n \leq 2^v$, i.e., $v \approx \log n$. If $k > v + 2$, then $Pr[X > 1] \leq \frac{1}{4}$.

Proof. Markov’s inequality yields

$$Pr[X > 1] = Pr \left[ X > \frac{2k}{n} E[X] \right] < Pr[X > \frac{2k}{2v} E[X]] < Pr[X > 4E[X]] < \frac{1}{4}.$$ 

Lemma 13.14. If $k < v - 2$, then $P[X = 0] \leq \frac{1}{4}$.

Proof. A similar analysis is possible to upper bound the probability that a transmission fails if our estimate is too small. We know that $k \leq v - 2$ and thus

$$Pr[X = 0] = \left(1 - \frac{1}{2^k}\right)^n < e^{-\frac{n}{2^k}} < e^{-\frac{2^{v-1}}{2^k}} < e^{-2} < \frac{1}{4}. $$

Lemma 13.15. If $v - 2 \leq k \leq v + 2$, then the probability that exactly one node transmits is constant.

Proof. The transmission probability is $p = \frac{1}{2^{v-2} + 1} = \Theta(1/n)$, and the lemma follows with a slightly adapted version of Theorem 13.1.

Lemma 13.16. With probability $1 - \frac{1}{\log n}$ we find a leader in phase 3 in $O(\log \log n)$ time.

Proof. For any $k$, because of Lemmas 13.13 and 13.14, the random walk of the third phase is biased towards the good area. One can show that in $O(\log \log n)$ steps one gets $\Omega(\log \log n)$ good transmissions. Let $Y$ denote the number of times exactly one node transmitted. With Lemma 13.15 we obtain $E[Y] = \Omega(\log \log n)$. Now a direct application of a Chernoff bound (see Theorem 13.22) yields that these transmissions elect a leader with probability $1 - \frac{1}{\log n}$.

Theorem 13.17. The Algorithm 55 elects a leader with probability of at least $1 - \frac{\log \log n}{\log n}$ in time $O(\log \log n)$. 


13.3. LEADER ELECTION

Proof. From Corollary 13.11 we know that after $O(\log \log n)$ time slots, the first phase terminates. Since we perform a binary search on an interval of size $O(\log n)$, the second phase also takes at most $O(\log \log n)$ time slots. For the third phase we know that $O(\log \log n)$ slots are sufficient to elect a leader with probability $1 - \frac{1}{\log n}$ by Lemma 13.16. Thus, the total runtime is $O(\log \log n)$.

Now we can combine the results. We know that the error probability for every time slot in the first two phases is at most $\frac{1}{\log n}$. Using a union bound (see Theorem 13.20), we can upper bound the probability that no error occurred by $\frac{\log \log n}{\log n}$. Thus, we know that after phase 2 our estimate is at most $\log \log n$ away from $\log n$ with probability of at least $1 - \frac{\log \log n}{\log n}$. Hence, we can apply Lemma 13.16 and thus successfully elect a leader with probability of at least $1 - \frac{\log \log n}{\log n}$ (again using a union bound) in time $O(\log \log n)$.

Remarks:

- Tightening this analysis a bit more, one can elect a leader with probability $1 - \frac{1}{\log n}$ in time $\log \log n + o(\log \log n)$.

- Can we be even faster?

13.3.5 Lower Bound

Theorem 13.18. Any uniform protocol that elects a leader with probability of at least $1 - \frac{1}{\log n}$ must run for at least $\log \log n$ time slots.

Proof. The probability that exactly one node transmits is

$$Pr[X = 1] = n \cdot p \cdot (1 - p)^{n-1}.$$ 

Consider now a system with only 2 nodes. The probability that exactly one transmits is at most

$$Pr[X = 1] = p \cdot (1 - p) \leq \frac{1}{2}.$$ 

Thus, after $\log \log n$ time slots the probability that a leader was elected is at most $1 - \frac{1}{2} \log \log n = 1 - \frac{1}{\log n}$.

13.3.6 Uniform Asynchronous Wakeup without CD

Until now we have assumed that all nodes start the algorithm in the same time slot. But what happens if this is not the case? How long does it take to elect a leader if we want an uniform and anonymous (nodes do not have an identifier and thus cannot base their decision on it) algorithm?

Theorem 13.19. If nodes wake up in an arbitrary (worst-case) way, any algorithm may take $\Omega(n/\log n)$ time slots until a single node can successfully transmit.
Proof. Nodes must transmit at some point, or they will surely never successfully transmit. With a uniform protocol, every node executes the same code. We focus on the first slot where nodes may transmit. No matter what the protocol is, this happens with probability $p$. Since the protocol is uniform, $p$ must be a constant, independent of $n$.

The adversary wakes up $w = \frac{c}{n} \ln n$ nodes in each time slot with some constant $c$. All nodes woken up in the first time slot will transmit with probability $p$. We study the event $E_1$ that exactly one of them transmits in that first time slot. Using the inequality $(1 + t/n)^n \leq e^t$ from Lemma 13.23 we get

$$\Pr[E_1] = w \cdot p \cdot (1 - p)^{w-1} = c \ln n (1 - p)^{\frac{1}{2}(c \ln n - p)} \leq c \ln n \cdot e^{-c \ln + p} = c \ln n \cdot n^{-c} e^p = n^{-c} \cdot \Theta(\log n) \leq \frac{1}{n^{c-1}} = \frac{1}{n^{c}}.$$ 

In other words, w.h.p. that time slot will not be successful. Since the nodes cannot distinguish noise from silence, the same argument applies to every set of nodes which wakes up. Let $E_\alpha$ be the event that all $n/w$ time slots will not be successful. Using the inequality $1 - p \leq (1 - p/k)^k$ from Lemma 13.24 we get

$$\Pr[E_\alpha] = (1 - \Pr(E_1))^{n/w} > \left(1 - \frac{1}{n^{c}}\right)^{\Theta(n/\log n)} > 1 - \frac{1}{n^{c'}}.$$ 

In other words, w.h.p. it takes more than $n/w$ time slots until some node can transmit alone.

In this chapter we have used several inequalities in our proofs. For simplicity’s sake we list all of them in this section.

Theorem 13.20. Boole’s inequality or union bound: For a countable set of events $E_1, E_2, E_3, \ldots$, we have

$$\Pr[\bigcup_i E_i] \leq \sum_i \Pr[E_i].$$

Theorem 13.21. Markov’s inequality: If $X$ is any random variable and $a > 0$, then

$$\Pr[|X| \geq a] \leq \frac{E[X]}{a}.$$ 

Theorem 13.22. Chernoff bound: Let $Y_1, \ldots, Y_n$ be a independent Bernoulli random variables let $Y := \sum_i Y_i$. For any $0 \leq \delta \leq 1$ it holds

$$\Pr[Y < (1 - \delta)E[Y]] \leq e^{-\frac{\delta^2}{2} E[Y]}$$
and for \( \delta > 0 \)

\[
\Pr[Y \geq (1 + \delta) \cdot E[Y]] \leq e^{-\min\left( \delta, \frac{\delta^2}{2} \right) E[Y]}
\]

**Theorem 13.23.** We have

\[
e^t \left( 1 - \frac{t^2}{n} \right) \leq \left( 1 + \frac{1}{n} \right)^n \leq e^t
\]

for all \( n \in \mathbb{N} \), \(|t| \leq n\). Note that

\[
\lim_{n \to \infty} \left( 1 + \frac{1}{n} \right)^n = e^t.
\]

**Theorem 13.24.** For all \( p, k \) such that \( 0 < p < 1 \) and \( k \geq 1 \) we have

\[1 - p \leq (1 - p/k)^k.\]

**Chapter Notes**

The Aloha protocol is presented and analyzed in [Abr70, BAK+75, Abr85]; the basic technique that unslotted protocols are twice as bad as slotted protocols is from [Rob75]. The idea to broadcast in a packet radio network by building a tree was first presented in [TM78, Cap79]. This idea is also used in [HNO99] to initialize the nodes. Willard [Wil86] was the first that managed to elect a leader in \( O(\log \log n) \) time in expectation. Looking more carefully at the success rate, it was shown that one can elect a leader with probability \( 1 - \frac{1}{\log n} \) in time \( \log \log n + o(\log \log n) \) [NO98]. Finally, approximating the number of nodes in the network is analyzed in [JKZ02, CGK05]. The lower bound for probabilistic wake-up is published in [JS02]. In addition to single-hop networks, multi-hop networks have been analyzed, e.g. broadcast [BYGI92, KM98, CR06], or deployment [MvRW06].

This chapter was written in collaboration with Philipp Brandes.

**Bibliography**


Chapter 14

Peer-to-Peer Computing

“Indeed, I believe that virtually every important aspect of programming arises somewhere in the context of [sorting and] searching!”

– Donald E. Knuth, The Art of Computer Programming

14.1 Introduction

Unfortunately, the term peer-to-peer (P2P) is ambiguous, used in a variety of different contexts, such as:

- In popular media coverage, P2P is often synonymous to software or protocols that allow users to “share” files, often of dubious origin. In the early days, P2P users mostly shared music, pictures, and software; nowadays books, movies or tv shows have caught on. P2P file sharing is immensely popular, currently at least half of the total Internet traffic is due to P2P!

- In academia, the term P2P is used mostly in two ways. A narrow view essentially defines P2P as the “theory behind file sharing protocols”. In other words, how do Internet hosts need to be organized in order to deliver a search engine to find (file sharing) content efficiently? A popular term is “distributed hash table” (DHT), a distributed data structure that implements such a content search engine. A DHT should support at least a search (for a key) and an insert (key, object) operation. A DHT has many applications beyond file sharing, e.g., the Internet domain name system (DNS).

- A broader view generalizes P2P beyond file sharing: Indeed, there is a growing number of applications operating outside the juridical gray area, e.g., P2P Internet telephony à la Skype, P2P mass player games on video consoles connected to the Internet, P2P live video streaming as in Zattoo or StreamForge, or P2P social storage such as Wuala. So, again, what is P2P?! Still not an easy question... Trying to account for the new applications beyond file sharing, one might define P2P as a large-scale distributed system that operates without a central server bottleneck. However, with
this definition almost everything we learn in this course is P2P! Moreover, according to this definition early-day file sharing applications such as Napster (1999) that essentially made the term P2P popular would not be P2P! On the other hand, the plain old telephone system or the world wide web do fit the P2P definition...

- From a different viewpoint, the term P2P may also be synonymous for privacy protection, as various P2P systems such as Freenet allow publishers of information to remain anonymous and uncensored. (Studies show that these freedom-of-speech P2P networks do not feature a lot of content against oppressive governments; indeed the majority of text documents seem to be about illicit drugs, not to speak about the type of content in audio or video files.)

In other words, we cannot hope for a single well-fitting definition of P2P, as some of them even contradict. In the following we mostly employ the academic viewpoints (second and third definition above). In this context, it is generally believed that P2P will have an influence on the future of the Internet. The P2P paradigm promises to give better scalability, availability, reliability, fairness, incentives, privacy, and security, just about everything researchers expect from a future Internet architecture. As such it is not surprising that new “clean slate” Internet architecture proposals often revolve around P2P concepts.

One might naively assume that for instance scalability is not an issue in today’s Internet, as even most popular web pages are generally highly available. However, this is not really because of our well-designed Internet architecture, but rather due to the help of so-called overlay networks: The Google website for instance manages to respond so reliably and quickly because Google maintains a large distributed infrastructure, essentially a P2P system. Similarly companies like Akamai sell “P2P functionality” to their customers to make today’s user experience possible in the first place. Quite possibly today’s P2P applications are just testbeds for tomorrow’s Internet architecture.

### 14.2 Architecture Variants

Several P2P architectures are known:

- **Client/Server goes P2P**: Even though Napster is known to the be first P2P system (1999), by today’s standards its architecture would not deserve the label P2P anymore. Napster clients accessed a central server that managed all the information of the shared files, i.e., which file was to be found on which client. Only the downloading process itself was between clients (“peers”) directly, hence peer-to-peer. In the early days of Napster the load of the server was relatively small, so the simple Napster architecture made a lot of sense. Later on, it became clear that the server would eventually be a bottleneck, and more so an attractive target for an attack. Indeed, eventually a judge ruled the server to be shut down, in other words, he conducted a juridical denial of service attack.

- **Unstructured P2P**: The Gnutella protocol is the anti-thesis of Napster, as it is a fully decentralized system, with no single entity having a global picture. Instead each peer would connect to a random sample of other
peers, constantly changing the neighbors of this virtual overlay network by exchanging neighbors with neighbors of neighbors. (In such a system it is part of the challenge to find a decentralized way to even discover a first neighbor; this is known as the bootstrap problem. To solve it, usually some random peers of a list of well-known peers are contacted first.) When searching for a file, the request was being flooded in the network (Algorithm 11 in Chapter 3). Indeed, since users often turn off their client once they downloaded their content there usually is a lot of churn (peers joining and leaving at high rates) in a P2P system, so selecting the right “random” neighbors is an interesting research problem by itself. However, unstructured P2P architectures such as Gnutella have a major disadvantage, namely that each search will cost $m$ messages, $m$ being the number of virtual edges in the architecture. In other words, such an unstructured P2P architecture will not scale.

- Hybrid P2P: The synthesis of client/server architectures such as Napster and unstructured architectures such as Gnutella are hybrid architectures. Some powerful peers are promoted to so-called superpeers (or, similarly, trackers). The set of superpeers may change over time, and taking down a fraction of superpeers will not harm the system. Search requests are handled on the superpeer level, resulting in much less messages than in flat/homogeneous unstructured systems. Essentially the superpeers together provide a more fault-tolerant version of the Napster server, all regular peers connect to a superpeer. As of today, almost all popular P2P systems have such a hybrid architecture, carefully trading off reliability and efficiency, but essentially not using any fancy algorithms and techniques.

- Structured P2P: Inspired by the early success of Napster, the academic world started to look into the question of efficient file sharing. The proposal of hypercubic architectures lead to many so-called structured P2P architecture proposals, such as Chord, CAN, Pastry, Tapestry, Viceroy, Kademlia, Koorde, SkipGraph, SkipNet, etc. In practice structured P2P architectures are not yet popular, apart from the Kad (from Kademlia) architecture which comes for free with the eMule client.

14.3 Hypercubic Networks

In this section we will introduce some popular families of network topologies. These topologies are used in countless application domains, e.g., in classic parallel computers or telecommunication networks, or more recently (as said above) in P2P computing. Similarly to Chapter 4 we employ an All-to-All communication model, i.e., each node can set up direct communication links to arbitrary other nodes. Such a virtual network is called an overlay network, or in this context, P2P architecture. In this section we present a few overlay topologies of general interest.

The most basic network topologies used in practice are trees, rings, grids or tori. Many other suggested networks are simply combinations or derivatives of these. The advantage of trees is that the routing is very easy: for every source-destination pair there is only one possible simple path. However, since the root
of a tree is usually a severe bottleneck, so-called fat trees have been used. These trees have the property that every edge connecting a node $v$ to its parent $u$ has a capacity that is equal to all leaves of the subtree routed at $v$. See Figure 14.1 for an example.

![Figure 14.1: The structure of a fat tree.](image)

**Remarks:**

- Fat trees belong to a family of networks that require edges of non-uniform capacity to be efficient. Easier to build are networks with edges of uniform capacity. This is usually the case for grids and tori. Unless explicitly mentioned, we will treat all edges in the following to be of capacity 1. In the following, $[x]$ means the set $\{0, \ldots, x - 1\}$.

**Definition 14.1 (Torus, Mesh).** Let $m, d \in \mathbb{N}$. The $(m, d)$-mesh $M(m, d)$ is a graph with node set $V = [m]^d$ and edge set

$$E = \left\{ ((a_1, \ldots, a_d), (b_1, \ldots, b_d)) \mid a_i, b_i \in [m], \sum_{i=1}^{d} |a_i - b_i| = 1 \right\}.$$

The $(m, d)$-torus $T(m, d)$ is a graph that consists of an $(m, d)$-mesh and additionally wrap-around edges from nodes $(a_1, \ldots, a_{i-1}, m, a_{i+1}, \ldots, a_d)$ to nodes $(a_1, \ldots, a_{i-1}, 1, a_{i+1}, \ldots, a_d)$ for all $i \in \{1, \ldots, d\}$ and all $a_j \in [m]$ with $j \neq i$. In other words, we take the expression $a_i - b_i$ in the sum modulo $m$ prior to computing the absolute value. $M(m, 1)$ is also called a line, $T(m, 1)$ a cycle, and $M(2, d) = T(2, d)$ a $d$-dimensional hypercube. Figure 14.2 presents a linear array, a torus, and a hypercube.

**Remarks:**

- Routing on mesh, torus, and hypercube is trivial. On a $d$-dimensional hypercube, to get from a source bitstring $s$ to a target bitstring $d$ one only needs to fix each “wrong” bit, one at a time; in other words, if the source and the target differ by $k$ bits, there are $k!$ routes with $k$ hops.
14.3. HYPERCUBIC NETWORKS

![Hypercube Diagram]

Figure 14.2: The structure of $M(m, 1)$, $T(4, 2)$, and $M(2, 3)$.

• The hypercube can directly be used for a structured P2P architecture. It is trivial to construct a distributed hash table (DHT): We have $n$ nodes, $n$ for simplicity being a power of 2, i.e., $n = 2^d$. As in the hypercube, each node gets a unique $d$-bit ID, and each node connects to $d$ other nodes, i.e., the nodes that have IDs differing in exactly one bit. Now we use a globally known hash function $f$, mapping file names to long bit strings; SHA-1 is popular in practice, providing 160 bits. Let $f_d$ denote the first $d$ bits (prefix) of the bitstring produced by $f$. If a node is searching for file name $X$, it routes a request message $f(X)$ to node $f_d(X)$. Clearly, node $f_d(X)$ can only answer this request if all files with hash prefix $f_d(X)$ have been previously registered at node $f_d(X)$.

• There are a few issues which need to be addressed before our DHT works, in particular churn (nodes joining and leaving without notice). To deal with churn the system needs some level of replication, i.e., a number of nodes which are responsible for each prefix such that failure of some nodes will not compromise the system. We give some more details in Section 14.4. In addition there are other issues (e.g., security, efficiency) which can be addressed to improve the system. These issues are beyond the scope of this lecture.

• The hypercube has many derivatives, the so-called hypercubic networks. Among these are the butterfly, cube-connected-cycles, shuffle-exchange, and de Bruijn graph. We start with the butterfly, which is basically a “rolled out” hypercube (hence directly providing replication!).

**Definition 14.2 (Butterfly).** Let $d \in \mathbb{N}$. The $d$-dimensional butterfly $BF(d)$ is a graph with node set $V = [d + 1] \times [2]^d$ and an edge set $E = E_1 \cup E_2$ with

$$E_1 = \{((i, \alpha), (i + 1, \alpha)) \mid i \in [d], \alpha \in [2]^d\}$$

and

$$E_2 = \{((i, \alpha), (i + 1, \beta)) \mid i \in [d], \alpha, \beta \in [2]^d, \alpha \text{ and } \beta \text{ differ only at the } i^{th} \text{ position}\}.$$

A node set $\{(i, \alpha) \mid \alpha \in [2]^d\}$ is said to form level $i$ of the butterfly. The $d$-dimensional wrap-around butterfly $W-BF(d)$ is defined by taking the $BF(d)$ and identifying level $d$ with level 0.
Remarks:

- Figure 14.3 shows the 3-dimensional butterfly $BF(3)$. The $BF(d)$ has $(d+1)2^d$ nodes, $2d \cdot 2^d$ edges and degree 4. It is not difficult to check that combining the node sets $\{(i, \alpha) \mid i \in [d]\}$ into a single node results in the hypercube.

- Butterflies have the advantage of a constant node degree over hypercubes, whereas hypercubes feature more fault-tolerant routing.

- The structure of a butterfly might remind you of sorting networks from Chapter 4. Although butterflies are used in the P2P context (e.g. Viceroy), they have been used decades earlier for communication switches. The well-known Benes network is nothing but two back-to-back butterflies. And indeed, butterflies (and other hypercubic networks) are even older than that; students familiar with fast fourier transform (FFT) will recognize the structure without doubt. Every year there is a new application for which a hypercubic network is the perfect solution!

- Indeed, hypercubic networks are related. Since all structured P2P architectures are based on hypercubic networks, they in turn are all related.

- Next we define the cube-connected-cycles network. It only has a degree of 3 and it results from the hypercube by replacing the corners by cycles.

\begin{figure}[h]
\centering
\includegraphics[width=0.5\textwidth]{bf3.png}
\caption{The structure of BF(3).}
\end{figure}

**Definition 14.3** (Cube-Connected-Cycles). Let $d \in \mathbb{N}$. The cube-connected-cycles network $CCC(d)$ is a graph with node set $V = \{(a, p) \mid a \in [2]^d, p \in [d]\}$ and edge set

\[
E = \left\{ \{(a, p), (a, (p + 1) \mod d)\} \mid a \in [2]^d, p \in [d]\right\}
\cup \left\{ \{(a, p), (b, p)\} \mid a, b \in [2]^d, p \in [d], a = b \text{ except for } a_p \right\}.
\]
14.3. HYPERCUBIC NETWORKS

Figure 14.4: The structure of CCC(3).

Remarks:

- Two possible representations of a CCC can be found in Figure 14.4.
- The shuffle-exchange is yet another way of transforming the hypercubic interconnection structure into a constant degree network.

Definition 14.4 (Shuffle-Exchange). Let $d \in N$. The $d$-dimensional shuffle-exchange $SE(d)$ is defined as an undirected graph with node set $V = \{2\}^d$ and an edge set $E = E_1 \cup E_2$ with

$$E_1 = \{(a_1, \ldots, a_d), (a_1, \ldots, \bar{a}_d)\} \mid (a_1, \ldots, a_d) \in \{2\}^d, \bar{a}_d = 1 - a_d\}$$

and

$$E_2 = \{(a_1, \ldots, a_d), (a_d, a_1, \ldots, a_{d-1})\} \mid (a_1, \ldots, a_d) \in \{2\}^d\} \right.$$.

Figure 14.5 shows the 3- and 4-dimensional shuffle-exchange graph.

Figure 14.5: The structure of SE(3) and SE(4).

Definition 14.5 (DeBruijn). The $b$-ary DeBruijn graph of dimension $d$ $DB(b, d)$ is an undirected graph $G = (V, E)$ with node set $V = \{v \in [b]^d\}$ and edge set $E$ that contains all edges $\{v, w\}$ with the property that $w \in \{(x, v_1, \ldots, v_{d-1}) : x \in [b]\}$, where $v = (v_1, \ldots, v_d)$. 
CHAPTER 14. PEER-TO-PEER COMPUTING

Remarks:

- Two examples of a DeBruijn graph can be found in Figure 14.6. The DeBruijn graph is the basis of the Koorde P2P architecture.

- There are some data structures which also qualify as hypercubic networks. An obvious example is the Chord P2P architecture, which uses a slightly different hypercubic topology. A less obvious (and therefore good) example is the skip list, the balanced binary search tree for the lazy programmer:

Definition 14.6 (Skip List). The skip list is an ordinary ordered linked list of objects, augmented with additional forward links. The ordinary linked list is the level 0 of the skip list. In addition, every object is promoted to level 1 with probability 1/2. As for level 0, all level 1 objects are connected by a linked list. In general, every object on level \( i \) is promoted to the next level with probability \( 1/2 \). A special start-object points to the smallest/first object on each level.

Remarks:

- Search, insert, and delete can be implemented in \( O(\log n) \) expected time in a skip list, simply by jumping from higher levels to lower ones when overshoting the searched position. Also, the amortized memory cost of each object is constant, as on average an object only has two forward pointers.

- The randomization can easily be discarded, by deterministically promoting a constant fraction of objects of level \( i \) to level \( i + 1 \), for all \( i \). When inserting or deleting, object \( o \) simply checks whether its left and right level \( i \) neighbors are being promoted to level \( i + 1 \). If none of them is, promote object \( o \) itself. Essentially we establish a MIS on each level, hence at least every third and at most every second object is promoted.

- There are obvious variants of the skip list, e.g., the skip graph. Instead of promoting only half of the nodes to the next level, we always promote all the nodes, similarly to a balanced binary tree: All nodes are part of the root level of the binary tree. Half the nodes are promoted left, and half the nodes are promoted right, on each level. Hence on level \( i \) we have have \( 2^i \) lists (or, more symmetrically: rings) of about \( n/2^i \) objects. This is pretty much what we need for a nice hypercubic P2P architecture.

- One important goal in choosing a topology for a network is that it has a small diameter. The following theorem presents a lower bound for this.
Theorem 14.7. Every graph of maximum degree \( d > 2 \) and size \( n \) must have a diameter of at least \( \lceil \frac{(\log n)}{(\log(d-1))} \rceil - 2 \).

Proof. Suppose we have a graph \( G = (V,E) \) of maximum degree \( d \) and size \( n \). Start from any node \( v \in V \). In a first step at most \( d \) other nodes can be reached. In two steps at most \( d \cdot (d-1) \) additional nodes can be reached. Thus, in general, in at most \( k \) steps at most

\[
1 + \sum_{i=0}^{k-1} d \cdot (d-1)^i = 1 + d \cdot \frac{(d-1)^k - 1}{(d-1) - 1} \leq \frac{d \cdot (d-1)^k}{d-2}
\]

nodes (including \( v \)) can be reached. This has to be at least \( n \) to ensure that \( v \) can reach all other nodes in \( V \) within \( k \) steps. Hence,

\[
(d-1)^k \geq \frac{(d-2) \cdot n}{d} \iff k \geq \log_{d-1}((d-2) \cdot n/d).
\]

Since \( \log_{d-1}((d-2)/d) > -2 \) for all \( d > 2 \), this is true only if \( k > \lceil \frac{(\log n)}{(\log(d-1))} \rceil - 2 \).

Remarks:

- In other words, constant-degree hypercubic networks feature an asymptotically optimal diameter.

- There are a few other interesting graph classes, e.g., expander graphs (an expander graph is a sparse graph which has high connectivity properties, that is, from every not too large subset of nodes you are connected to a larger set of nodes), or small-world graphs (popular representations of social networks). At first sight hypercubic networks seem to be related to expanders and small-world graphs, but they are not.

14.4 DHT & Churn

As written earlier, a DHT essentially is a hypercubic structure with nodes having identifiers such that they span the ID space of the objects to be stored. We described the straightforward way how the ID space is mapped onto the peers for the hypercube. Other hypercubic structures may be more complicated: The butterfly network, for instance, may directly use the \( d+1 \) layers for replication, i.e., all the \( d+1 \) nodes with the same ID are responsible for the same hash prefix. For other hypercubic networks, e.g., the pancake graph (see exercises), assigning the object space to peer nodes may be more difficult.

In general a DHT has to withstand churn. Usually, peers are under control of individual users who turn their machines on or off at any time. Such peers join and leave the P2P system at high rates (“churn”), a problem that is not existent in orthodox distributed systems, hence P2P systems fundamentally differ from old-school distributed systems where it is assumed that the nodes in the system are relatively stable. In traditional distributed systems a single unavailable node is a minor disaster: all the other nodes have to get a consistent view of the system again, essentially they have to reach consensus which nodes are available.
In a P2P system there is usually so much churn that it is impossible to have a consistent view at any time.

Most P2P systems in the literature are analyzed against an adversary that can crash a fraction of random peers. After crashing a few peers the system is given sufficient time to recover again. However, this seems unrealistic. The scheme sketched in this section significantly differs from this in two major aspects. First, we assume that joins and leaves occur in a worst-case manner. We think of an adversary that can remove and add a bounded number of peers; it can choose which peers to crash and how peers join. We assume that a joining peer knows a peer which already belongs to the system. Second, the adversary does not have to wait until the system is recovered before it crashes the next batch of peers. Instead, the adversary can constantly crash peers, while the system is trying to stay alive. Indeed, the system is never fully repaired but always fully functional. In particular, the system is resilient against an adversary that continuously attacks the “weakest part” of the system. The adversary could for example insert a crawler into the P2P system, learn the topology of the system, and then repeatedly crash selected peers, in an attempt to partition the P2P network. The system counters such an adversary by continuously moving the remaining or newly joining peers towards the sparse areas.

Clearly, we cannot allow the adversary to have unbounded capabilities. In particular, in any constant time interval, the adversary can at most add and/or remove $O(\log n)$ peers, $n$ being the total number of peers currently in the system. This model covers an adversary which repeatedly takes down machines by a distributed denial of service attack, however only a logarithmic number of machines at each point in time. The algorithm relies on messages being delivered timely, in at most constant time between any pair of operational peers, i.e., the synchronous model. Using the trivial synchronizer this is not a problem. We only need bounded message delays in order to have a notion of time which is needed for the adversarial model. The duration of a round is then proportional to the propagation delay of the slowest message.

In the remainder of this section, we give a sketch of the system: For simplicity, the basic structure of the P2P system is a hypercube. Each peer is part of a distinct hypercube node; each hypercube node consists of $\Theta(\log n)$ peers. Peers have connections to other peers of their hypercube node and to peers of the neighboring hypercube nodes. Because of churn, some of the peers have to change to another hypercube node such that up to constant factors, all hypercube nodes own the same number of peers at all times. If the total number of peers grows or shrinks above or below a certain threshold, the dimension of the hypercube is increased or decreased by one, respectively.

The balancing of peers among the hypercube nodes can be seen as a dynamic token distribution problem on the hypercube. Each node of the hypercube has a certain number of tokens, the goal is to distribute the tokens along the edges of the graph such that all nodes end up with the same or almost the same number of tokens. While tokens are moved around, an adversary constantly inserts and deletes tokens. See also Figure 14.7.

In summary, the P2P system builds on two basic components: i) an algorithm which performs the described dynamic token distribution and ii) an in-

\footnote{Having a logarithmic number of hypercube neighbor nodes, each with a logarithmic number of peers, means that each peer has $\Theta(\log^2 n)$ neighbor peers. However, with some additional bells and whistles one can achieve $\Theta(\log n)$ neighbor peers.}
14.5 STORAGE AND MULTICAST

Figure 14.7: A simulated 2-dimensional hypercube with four nodes, each consisting of several peers. Also, all the peers are either in the core or in the periphery of a node. All peers within the same node are completely connected to each other, and additionally, all peers of a node are connected to the core peers of the neighboring nodes. Only the core peers store data items, while the peripheral peers move between the nodes to balance biased adversarial changes.

formation aggregation algorithm which is used to estimate the number of peers in the system and to adapt the dimension of the hypercube accordingly:

**Theorem 14.8** (DHT with Churn). *We have a fully scalable, efficient P2P system which tolerates $O(\log n)$ worst-case joins and/or crashes per constant time interval. As in other P2P systems, peers have $O(\log n)$ neighbors, and the usual operations (e.g., search, insert) take time $O(\log n)$. 

**Remarks:**

- Indeed, handling churn is only a minimal requirement to make a P2P system work. Later studies proposed more elaborate architectures which can also handle other security issues, e.g., privacy or Byzantine attacks.

- It is surprising that unstructured (in fact, hybrid) P2P systems dominate structured P2P systems in the real world. One would think that structured P2P systems have advantages, in particular their efficient logarithmic data lookup. On the other hand, unstructured P2P networks are simpler, in particular in light of non-exact queries.

14.5 Storage and Multicast

As seen in the previous section, practical implementations often incorporate some non-rigid (flexible) part. In a system called Pastry, prefix-based overlay structures similar to hypercubes are used to implement a DHT. Peers maintain connections to other peers in the overlay according to the lengths of the shared prefixes of their respective identifiers, where each peer carries a $d$-bit peer identifier. Let $\beta$ denote the number of bits that can be fixed at a peer to route any message to an arbitrary destination. For $i = \{0, \beta, 2\beta, 3\beta, \ldots\}$, a peer chooses, if possible, $2^i - 1$ neighbors whose identifiers are equal in the $i$
most significant bits and differ in the subsequent $\beta$ bits by one of $2^\beta - 1$ possibilities. If peer identifiers are chosen uniformly at random, the length of the longest shared prefix is bounded by $O(\log n)$ in an overlay containing $n$ peers; thus, only $O(\log n(2^\beta - 1)/\beta)$ connections need to be maintained. Moreover, every peer reaches every other peer in $O(\frac{\log n}{\beta})$ hops by repetitively selecting the next hop to fix $\beta$ more bits toward the destination peer identifier, yielding a logarithmic overlay diameter.

The advantage of prefix-based over more rigid DHT structures is that there is a large choice of neighbors for most prefixes. Peers are no longer bound to connect to peers exactly matching a given identifier. Instead peers are enabled to connect to any peer matching a desired prefix, regardless of subsequent identifier bits. In particular, among half of all peers can be chosen for a shared prefix of length 0. The flexibility of such a neighbor policy allows the optimization of secondary criteria. Peers may favor peers with a low-latency and select multiple neighbors for the same prefix to gain resilience against churn. Regardless of the choice of neighbors, the overlay always remains connected with a bounded degree and diameter.

Such overlay structures are not limited to distributed storage. Instead, they are equally well suited for the distribution of content, such as multicasting of radio stations or television channels. In a basic multicasting scheme, a source with identifier 00...0 may forward new data blocks to two peers having identifiers starting with 0 and 1. They in turn forward the content to peers having identifiers starting with 00, 01, 10, and 11. The recursion finishes once all peers are reached. This basic scheme has the subtle shortcoming that data blocks may pass by multiple times at a single peer because a predecessor can match a prefix further down in its distribution branch.

The subsequent multicasting scheme $\mathcal{M}$ avoids this problem by modifying the topology and using a different routing scheme. For simplicity, the neighbor selection policy is presented for the case $\beta = 1$. In order to use $\mathcal{M}$, the peers must store links to a different set of neighbors. A peer $v$ with the identifier $b_v^0 ... b_v^{d-1}$ stores links to peers whose identifiers start with $b_v^0 b_v^1 ... b_v^{i-1} b_v^i b_v^{i+1}$ and $b_v^0 b_v^1 ... b_v^{i-1} \overline{b_v^i} b_v^{i+1}$ for all $i \in \{0, ..., d - 2\}$. For example, the peer with the identifier 0000 has to maintain connections to peers whose identifiers start with the prefixes 10, 11, 010, 011, 0010, and 0011. Pseudo-code for the algorithm is given in Algorithm 56.

**Algorithm 56 $\mathcal{M}$: forward($\pi, v_c$) at peer $v$.**

1: $\mathcal{S} := \{v' \in \mathcal{N}_v : \ell(v', v) \geq \pi + 1\}$
2: choose $v_1 \in \mathcal{S}$: $\ell(v_1, v) \leq \ell(\tilde{v}, v)$ $\forall \tilde{v} \in \mathcal{S}$
3: if $v_1 \neq \emptyset$ then forward($\ell(v_1, v), v$) to $v_1$ fi
4: if $v_c \neq \emptyset$ then
5: choose $v_2 \in \mathcal{N}_v$: $\ell(v_2, v_c) = \pi + 1$
6: if $v_2 = \emptyset$ then $v_2 :=$ getNext($v$) from $v_c$ fi
7: if $v_2 \neq \emptyset$ then forward($\ell(v_2, v_c), v_c$) to $v_2$ fi
8: else
9: choose $v_2 \in \mathcal{N}_v$: $\ell(v_2, v) = \pi$
10: if $v_2 \neq \emptyset$ then forward($\pi + 1, v_c$) to $v_2$ fi
11: fi
The parameters are the length \( \pi \) of the prefix that is not to be modified and at most one critical predecessor \( v_c \). If \( \beta = 1 \), any node \( v \) tries to forward the data block to two peers \( v_1 \) and \( v_2 \). The procedure is called at the source \( v_0 \) with arguments \( \pi := 0 \) and \( v_c := \emptyset \), resulting in the two messages \( \text{forward}(1, v_0) \) to \( v_1 \) and \( \text{forward}(1, \emptyset) \) to \( v_2 \). The peer \( v_1 \) is chosen locally such that the prefix its identifier shares with the identifier of \( v \) is the shortest among all those whose shared prefix length is at least \( \pi + 1 \). This value \( \ell(v_1, v) \) and \( v \) itself are the parameters included in the forward message to peer \( v_1 \), if such a peer exists.

The second peer is chosen similarly, but with respect to \( v_c \) and not \( v \) itself. If no suitable peer is found in the routing table, the peer \( v_c \) is queried for a candidate using the subroutine \( \text{getNext} \) which is described in Algorithm 57. This step is required because node \( v \) cannot deduce from its routing table whether a peer \( v_2 \) with the property \( \ell(v_2, v_c) \geq \pi + 1 \) exists. In the special case when \( v_c = \emptyset \), \( v_2 \) is chosen locally, if possible, such that \( \ell(v_2, v) = \pi \). In Figure 14.8, a sample spanning tree resulting from the execution of \( M \) is depicted.

**Algorithm 57 get
\[ \text{getNext}(v_s) \text{ at peer } v \]

1. \( S := \{ v' \in \mathcal{N}_v \mid \ell(v', v) > \ell(v_s, v) \} \)
2. choose \( v_r \in S : \ell(v_r, v) \leq \ell(\tilde{v}, v) \forall \tilde{v} \in S \)
3. send \( v_r \) to \( v_s \)

Figure 14.8: The spanning tree induced by a forward message initiated at peer \( v_0 \) is shown. The fixed prefix is underlined at each peer, whereas prefixes in bold print indicate that the parent peer has been constrained to forward the packet to peers with these prefixes.

The presented multicasting scheme \( M \) has the property that, at least in a static setting, wherein peers neither join nor leave the overlay, all peers can be reached and each peer receives a data block exactly once as summarized by the following theorem:

**Theorem 14.9.** In a static overlay, algorithm \( M \) has the following properties:

(a) It does not induce any duplicate messages (loop-free), and

(b) all peers are reached (complete).
Remarks:

- The multicast scheme $\mathcal{M}$ benefits from the same overlay properties as DHTs; there is a bounded diameter and peer degree. Peers can maintain backup neighbors and favor low-latency, high-bandwidth peers as neighbors. Most importantly, intermediate peers have the possibility to choose among multiple (backup) neighbors to forward incoming data blocks. This, in turn, allows peers to quickly adapt to changing network conditions such as churn and congestion. It is not necessary to rebuild the overlay structure after failures. In doing so, a system can gain both robustness and efficiency.

- In contrast, for more rigid data structures, such as trees, data blocks are forced to travel along fixed data paths, rendering them susceptible to any kind of failure.

- Conversely, unstructured and more random overlay networks lack the structure to immediately forward incoming data blocks. Instead, such systems have to rely on the exchange of periodic notifications about available data blocks and requests and responses for the download of missing blocks, significantly increasing distribution delays. Furthermore, the lack of structure makes it hard to maintain connectivity among all peers. If the neighbor selection is not truly random, but based on other criteria such as latency and bandwidth, clusters may form that disconnect themselves from the remaining overlay.

There is a variety of further flavors and optimizations for prefix-based overlay structures. For example, peers have a logarithmic number of neighbors in the presented structure. For 100,000 and more peers, peers have at least 20 neighbors. Selecting a backup neighbor doubles the number of neighbors to 40. Using $\mathcal{M}$ further doubles their number to 80. A large number of neighbors accrues substantial maintenance costs. The subsequent variation limits the number of neighbors with a slight adjustment of the overlay structure. It organizes peers into disjoint groups $G_0, G_1, \ldots, G_m$ of about equal size. The introduction of groups is motivated by the fact that they will enable peers to have neighboring connections for a subset of all shared prefixes while maintaining the favorable overlay properties. The source, feeding blocks into the overlay, joins group $G_0$. The other peers randomly join groups. Let $g(v)$ denote the function that assigns each peer $v$ to a group, i.e., $v \in G_{g(v)}$.

Peers select neighboring peers based not solely on shared prefixes but also on group membership. A peer $v$ with the identifier $b_v^0 \ldots b_v^{d-1}$ stores links to neighboring peers whose identifiers start with $b_v^0 b_v^1 \ldots b_{i-1}^i b_i^j$ and belong to group $g(v) + 1 \mod m$ for all $i \in \{g(v), g(v) + m, g(v) + 2m, g(v) + 3m, \ldots\}$. Furthermore, let $f$ denote the first index $i$ where no such peer exists. As fallback, peer $v$ stores further links to peers from arbitrary groups whose identifiers start with $b_v^0 b_v^1 \ldots b_{k-1}^k b_k^j$ for all $k \geq f - m + 1$. The fallback connections allow a peer to revert to the regular overlay structure for the longest shared prefixes where only few peers exist.

As an example, a scenario with $m = 4$ groups is considered. A peer with identifier 00...0 belonging to group $G_2$ has to maintain connections to peers from group $G_3$ that share the prefixes 001, 0000001, 00000000001, etc. In an
overlay with 100 peers, the peer is unlikely to find a neighbor for a prefix length larger than \( \log(100) \), such as prefix 00000000001. Instead, he further maintains fallback connections to peers from arbitrary groups having identifiers starting with the prefixes 000000001, 000000001, 000000001, etc. (if such peers exist).

Remarks:

- By applying the presented grouping mechanism, the total number of neighbors is reduced to \( \frac{2 \log n}{m} + c \) with constant \( c \) for fallback connections. (Note that peers have both outgoing neighbors to the next group and incoming neighbors from the previous group, doubling the number of neighbors.)

- Setting the number of groups \( m = \log n \) gives a constant number of neighbors regardless of the overlay size.

Chapter Notes

The paper of Plaxton, Rajaraman, and Richa [PRR97] laid out a blueprint for many so-called structured P2P architecture proposals, such as Chord [SMK+01], CAN [RFH+01], Pastry [RD01], Viceroy [MNR02], Kademlia [MM02], Koorde [KK03], SkipGraph [AS03], SkipNet [HJS+03], or Tapestry [ZHS+04]. Also the paper of Plaxton et. al. was standing on the shoulders of giants. Some of its eminent precursors are: linear and consistent hashing [KLL+97], locating shared objects [AP90, AP91], compact routing [SK85, PU88], and even earlier: hypercubic networks, e.g. [AJ75, Wit81, GS81, BA84].

Furthermore, the techniques in use for prefix-based overlay structures are related to a proposal called LAND, a locality-aware distributed hash table proposed by Abraham et al. [AMD04].

More recently, a lot of P2P research focussed on security aspects, describing for instance attacks [LMSW06, SENB07, Lar07], and provable countermeasures [KSW05, AS09, BSS09]. There are several recommendable introductory books on P2P computing, e.g. [SW05, SG05, MS07, KW08, BYL08].

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Bibliography


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