

RHEINISCHE FRIEDRICH-WILHELMS-UNIVERSITÄT BONN  
INSTITUT FÜR INFORMATIK I



---

Elmar Langetepe  
**Online Motion Planning**

MA INF 1314

---

---

Sommersemester 2016  
Manuscript: Elmar Langetepe



# Contents

- 1 Labyrinths, grids and graphs** **3**
- 1.1 Shannons Mouse Algorithm . . . . . 3
- 1.2 Intuitive connection of labyrinths, grids and graphs . . . . . 4
- 1.3 A lower bound for online graph exploration . . . . . 4
- 1.4 Exploration of grid environments . . . . . 8
  - 1.4.1 Exploration of simple gridpolygons . . . . . 9
  - 1.4.2 Competitive ratio of SmartDFS . . . . . 20
  - 1.4.3 Exploration of general gridpolygons . . . . . 23
- 1.5 Constrained graph-exploration . . . . . 31
  - 1.5.1 Restricted graph-exploration with unknown depth . . . . . 36
  - 1.5.2 Mapping of an unknown graph . . . . . 39



# List of Figures

1.1	Shannons original mouse labyrinth. . . . .	4
1.2	An example of the execution of Shannons Algorithm. . . . .	4
1.3	Labyrinth, labyrinth-graph and gridgraph. . . . .	5
1.4	The agent return to $s$ . . . . .	6
1.5	The agent has visited $\ell + 1$ vertices in corridor 3. . . . .	6
1.6	A polygon $P$ and the gridpolygon $P_{\square}$ as a reasonable approximation. . . . .	8
1.7	Ist DFS optimal? . . . . .	9
1.8	The number of boundary edges $E$ in comparison to the number of cells $C$ is a measure for the existence of <i>fleshy</i> or <i>skinny</i> parts. . . . .	9
1.9	A lower bound construction for the exploration of simple gridpolygons. . . . .	10
1.10	First simple improvement of DFS. . . . .	11
1.11	Second improvement of DFS. . . . .	12
1.12	The $\ell$ -Offset of gridpolygon $P$ . . . . .	14
1.13	Decomposition at a split-cell. . . . .	15
1.14	Three types of components. . . . .	15
1.15	Special cases: No component of typ (III) exists. . . . .	16
1.16	Wave-Propagation. . . . .	19
1.17	SmartDFS is optimal in narrow passages. . . . .	20
1.18	A simple gridpolygon without narrow passages and no split-cell in the first layer has the property $E(P) \leq \frac{2}{3}C(P) + 6$ . After the first coil SmartDFS starts in the 1-Offset $P'$ . The return path to $c'$ from an arbitrary point in $P'$ is shorter than $\frac{1}{2}E(P)/2 - 2$ . . . . .	21
1.19	In a corridor of width 3 and with even lenght the bound $S(P) = \frac{4}{3}S_{\text{Opt}}(P) - 2$ holds. . . . .	21
1.20	A gridpolygon $P_i$ that is separated into components of type (I) or (II) at the split-cell. The rectangle $Q$ is always inside $P_i$ . . . . .	23
1.21	$2D$ -cells and $D \times D$ sub-cells. . . . .	23
1.22	Examples for (i) $2D$ -Spiral-STC and (ii) Spiral-STC. . . . .	25
1.23	(i) Double-sided edge, (ii) one-sided edge, (iii) locally disconnected $2D$ -cell. . . . .	25
1.24	Avoid horizontal edges with the Scan-STC. . . . .	26
1.25	Examplle for (i) $2D$ -Scan-STC, (ii) Scan-STC. . . . .	26
1.26	Estimating the double visits of sub-cells by STC locally. . . . .	27
1.27	Analysis of STC, all possible cases. . . . .	28
1.28	(i) Columns and the change of connectivity, (ii) Columns without changes, (iii) Difficult online situation. . . . .	29
1.29	(i) A Graph with $n$ vertices and with depth $r = 1$ , pure DFS would require a tether of length $n - 1$ . (ii) A graph of depth $n$ , BFS with a tether of length $n$ requires $\Omega(n^2)$ steps. . . . .	32
1.30	bDFS kann einige Knoten nicht erreichen. . . . .	32

---

1.31	The algorithm maintains a set of disjoint trees $\mathcal{T} = \{T_1, T_2, T_3\}$ and choose the tree $T_2$ with minimal distance $d_{G^*}(s, s_i)$ . After that the tree is pruned. Subtrees of distance 2 away from $s_2$ with vertices inside that have distance at least 4 from $s_2$ are cut-off. After that DFS starts on the rest of $T_2$ and starts bDFS on the incomplete vertices. Here some new graphs $G'$ will be explored and we build spanning trees $T'$ for them. Some trees in $\mathcal{T}$ get fully explored. $T_w$ and $T'$ are added to $\mathcal{T}$ , the tree $T_2$ is deleted. . . . .	33
1.32	A graph of depth $r = 6$ that cannot be explored by an accumulator of size $2r$ . . . . .	38
1.33	A graph with $n + 1 = 13$ vertices. A path of length $\frac{n}{2}$ visits a clique of size $\frac{n}{2} + 1$ . Any accumulator strategy with accumulator size $n + 2 + d$ requires $\Omega(n^3)$ steps. . . . .	38
1.34	Two different regular graphs of degree 3, an agent cannot distinguish them without a marker. . . . .	39

# List of Algorithms

1.1	Shannons Maus	3
1.2	DFS	11
1.3	DFS with optimal return trips	12
1.4	SmartDFS	13
1.5	Algorithm of Lee	19
1.6	2D-Spiral-STC	24
1.7	SpiralSTC	24
1.8	ScanSTC	27
1.9	boundedDFS	32
1.10	CFS	34





---

## Introduction

This lecture considers tasks for autonomous agents. In general, constructing autonomous machines is a very complex challenge and has many different engineering and scientific aspects, some of which are given in the following list.

- Electronic devices
- Mechanical devices
- Control/Process engineering
- Artificial Intelligence
- Softwareengineering
- ∴
- Plans: **Algorithmic/Motion planning**
- Full information (offline)/ **Incomplete information (online)**
- Input: Geometry of the Environment

As part of the algorithm track of the master program we will concentrate on the item *Algorithms*. That is, we concentrate on the description and analysis of efficient schedules for solving motion planning tasks for autonomous agents. Besides, we concentrate on problem definitions and models that take the geometry of the scene into account. In this sense the scientific aspects of this course are part of the scientific area called Computational Geometry. Furthermore we consider online problems, which means that the full information of the problem is not given in advance. The agent has to *move* around and collects more information.

We will mainly concentrate on the ground tasks of autonomous agents in unknown environments such as

- Searching for a goal,
- Exploration of an environment,
- Escaping from a labyrinth,

and we consider different abilities of the agents some of which are

- Continuous/discrete vision,
- Touch sensor/compass,
- Building a map/constant memory.

The first concern is that we construct correct algorithms which always fulfil the task. Second we concentrate on the efficiency of the corresponding strategy. We would like to analyse performance guarantees and would like to provide for formal proofs. The course is related to the undergraduate course on *Offline motion planning*. In the offline case the information for the task is fully given and we only have to compute the best path for the agent. The offline solution will be used as a comparison measure for the online case. This is a well known concept for online problems in general.



# Chapter 1

## Labyrinths, grids and graphs

In this section we first concentrate on discrete environments based on grid structures. For the grid structure we consider an agent that can move from one cell to a neighbouring cell with unit cost. We start with the task of searching for a goal in a very special grid environment. After that we ask for visiting all cells, which means that we would like to explore the environment. For this task the grid environment is only partially known, by a touch sensor the agent can only detect the neighbouring cells. The agent can build a map. Exploration and Searching are closely related. If we are searching for an unknown goal, it is clear that in the worst-case the whole environment has to be explored. The main difference is the performance of these *online* tasks. As a comparison measure we compare the length of the agent's path to the length of the optimal path under full information. Thus, in the case of searching for a goal, the comparison measure is the shortest path to the goal.

At the end of the section we turn over to the exploration task in general graphs under different additional conditions.

### 1.1 Shannons Mouse Algorithm

Historically the first online motion planning algorithm for an autonomous agent was designed by Claude Shannon [Sha52, Sha93] in 1950. He considered a  $5 \times 5$  cellular labyrinth, the inner walls of the labyrinth could be placed around arbitrary cells. In principle, he constructed a labyrinth based on a grid environment; see Figure 1.1.

The task of his electronical mouse was to find a target, i.e. the cheese, located on one of the fields of the grid. The target and the start of the mouse were located in the same *connected component* of the *grid labyrinth*. The electronical mouse was able to move from one cell to a neighbouring cell. Additionally, it could (electronically) mark any cell by a label  $N, E, S, W$  which indicates in which direction the mouse left the cell at the last visit. This label is updated after leaving the cell. With these abilities the following algorithm was designed.

---

**Algorithm 1.1** Shannons Maus

---

- Initialize any cell by the label  $N$  for 'North'.
  - While the goal has not been found:  
starting from the label direction, search for the first cell in clockwise order that can be visited. Change the label to the corresponding direction and move to this neighbouring cell.
- 

Sutherland [Sut69] has shown that:

**Theorem 1.1** *Shannon's Algorithms (Algorithm 1.1) is correct. For any labyrinth, any starting and any goal the agent will find the goal, if a path from the start to the goal exists.*



Figure 1.1: Shannons original mouse labyrinth.

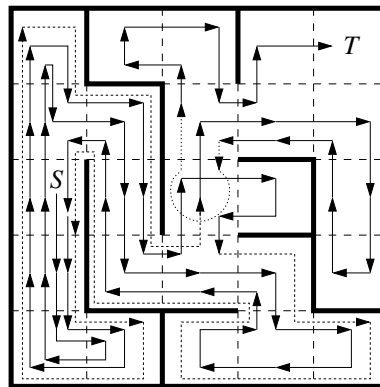


Figure 1.2: An example of the execution of Shannons Algorithm.

**Proof.** We omit the goal and show that any cell in the connected component of the start will be visited infinitely often. □

**Exercise 1** Formalize the above proof sketch!

As shown in Figure 1.2 the path of Shannons Mouse is not very efficient.

## 1.2 Intuitive connection of labyrinths, grids and graphs

For a human a labyrinth consists of corridors and connection points. In this sense the environment for Shannons task can be considered to be a labyrinth. Obviously any such labyrinth can be modeled by a planar graph.<sup>1</sup> More precisely the environment for Shannons task is a grid graph. Figure 1.3 shows the corresponding intuitive interpretations.

For any intuitive labyrinth there is a labyrinth-graph. On the other hand for any planar graph we can build some sort of labyrinth. This is not true for general graphs. For example the complete graph  $K_5$  has no planar representation and therefore a correspondance to a labyrinth does not exist.

## 1.3 A lower bound for online graph exploration

We consider the following model. Assume that a graph  $G = (V, E)$  is given. If the agent is located on a vertex it detects all neighbouring vertices. Let us assume that moving along an edge can be done with

<sup>1</sup>A graph, that has an intersection free representation in the plane.

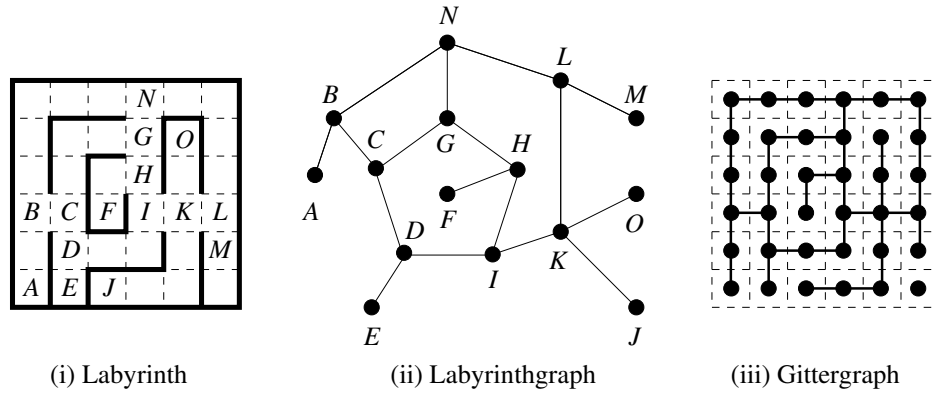


Figure 1.3: Labyrinth, labyrinth-graph and gridgraph.

unit cost. The task is to visit all edges and vertices and return to the start. The agent has the ability of building a map. If we apply a DFS (depth first search) for the edges we will move along any edges twice. DFS can run online. The best offline strategy has to visit any edge at least once. In this sense DFS is a 2-approximation.

The comparison and approximation between online and offline is represented by the following concept. A strategy that runs under incomplete information is denoted as an **Online-Strategy**. On the other hand an **Offline-Strategy** solves the same task with full information. In the above example the offline strategy is the shortest round trip that visits all edges of the graph.

The performance measure for Online-Algorithms is the so-called *competitive ratio*.

**Definition 1.2** (Sleator, Tarjan, 1995)

Let  $\Pi$  be a problem class and  $S$  be a strategy, that solves any instance  $P \in \Pi$ .

Let  $K_S(P)$  be the cost of  $S$  for solving  $P$ .

Let  $K_{opt}(P)$  be the cost of the optimal solution for  $P$ .

The strategy  $S$  is denoted to be  **$c$ -competitive**, if there are fixed constants  $c, \alpha > 0$ , so that for all  $P \in \Pi$

$$K_S(P) \leq c \cdot K_{opt}(P) + \alpha$$

holds.

The additive constant  $\alpha$  is often used for starting situations. For example if we are searching for a goal and have only two unknown options, the goal might be very close to the start, the unsuccessful step will lead to an arbitrarily large competitive ratio. This is not intended. Sometimes we can omit the additive constant, if we have additional assumptions. For example we can assume that the goal is at least distance 1 away from the start.

As already mentioned DFS on the edges visits any edge at most twice. There are graphs where the optimal offline solution also has to visit any edge twice. For such examples DFS is optimal with ratio 1. Now we are searching for a lower bound for the competitive ratio. That is, we would like to construct example such that any possible online strategy fails within a ratio of 2.

**Theorem 1.3** (Icking, Kamphans, Klein, Langetepe, 2000)

For the online-exploration of a graph  $G = (V, E)$  for visiting all edges and vertices of  $G$  there is always an arbitrarily large example such that any online strategy visits roughly twice as much edges in comparison to the optimal offline strategy. DFS always visit no more than twice as much edges against the optimum.

[IKKL00a]

**Proof.** The second part is clear because DFS visits exactly any edge twice. Any optimal strategy has to visit at least the edges.

The robot should explore a gridgraph and starts in a vertex  $s$ . Finally, the agent has to return to  $s$ . We construct an *open* corridor and offer two directions for the agent. At some moment in time the agent has explored  $\ell$  new vertices in the corridor. If this happens we let construct a conjunction at one end  $s'$  of the corridor. At this bifurcation two open corridors are build up which *run* back into the direction of  $s$ . If the agent proceeds one of the following events will happen.

1. The agent goes back to  $s$ .
2. The agent has visited more than  $\ell + 1$  edges in one of the new corridors.

Let  $\ell_1$  denote the length of the part of the starting open corridor into the opposite direction of  $s'$ . Let  $\ell_2$  and  $\ell_3$  denote the length of the second and third open corridor.

We analyse the edge visits  $|S_{ROB}|$  that an arbitrary strategy  $S_{ROB}$  has done so far.

1.  $|S_{ROB}| \geq 2\ell_1 + (\ell - \ell_1) + 2\ell_2 + 2\ell_3 + (\ell - \ell_1) = 2(\ell + \ell_2 + \ell_3)$ , see Figure 1.4. Now we close the corridors at the open ends. From now on the agent still requires  $|S_{OPT}| = 2(\ell + \ell_2 + \ell_3) + 6$  edge visits, where  $S_{OPT}$  is the optimal strategy if the situation was known from the beginning. Thus we have:  $|S_{ROB}| \geq 2|S_{OPT}| - 6$ .

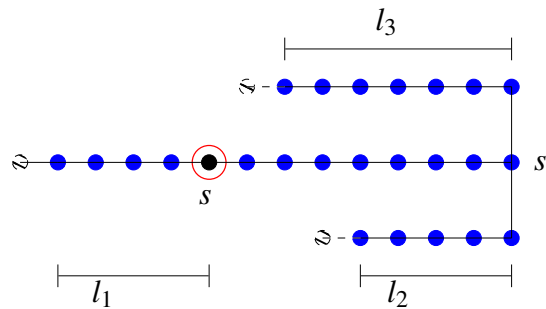


Figure 1.4: The agent return to  $s$ .

2. W.l.o.g. the agent has explored  $\ell + 1$ -ten vertices in corridor 3. We have  $|S_{ROB}| \geq 2\ell_1 + (\ell - \ell_1) + 2\ell_2 + (\ell + 1)$ . We connect corridor 3 with corridor 1 (see Figure 1.5) and close corridor 2. The agent still requires  $\ell + 1 + 2(\ell_2 + 1) + (\ell - \ell_1)$  edge visits; in total at least  $4\ell + 4\ell_2 + 4 = 4(\ell + \ell_2) + 4$  edge visits. From  $|S_{OPT}| = 2(\ell + 1) + 2(\ell_2 + 1) = 2(\ell + \ell_2) + 4$  we conclude  $|S_{ROB}| \geq 2|S_{OPT}| - 4 > 2|S_{OPT}| - 6$ .

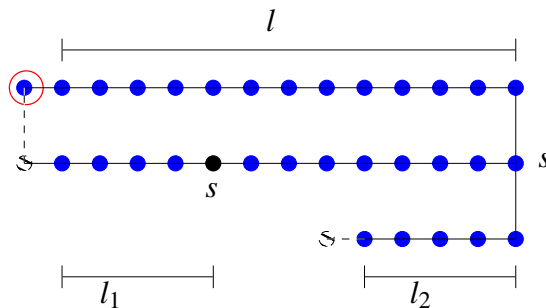


Figure 1.5: The agent has visited  $\ell + 1$  vertices in corridor 3.

We have  $|S_{ROB}|/|S_{OPT}| \geq 2 - 6/|S_{OPT}|$ . We also have  $|S_{OPT}| \geq 2(\ell + 1)$  and conclude  $2 - 6/|S_{OPT}| > 2 - 6/2\ell = 2 - 3/\ell$ . For arbitrary  $\delta > 0$  we choose  $\ell = \lceil 3/\delta \rceil$  and conclude  $|S_{ROB}|/|S_{OPT}| > 2 - \delta$ .  $\square$

**Remark 1.4** *There are always examples so that the optimal exploration tour visits any edge twice.*

**Corollary 1.5** *DFS for the Online-Edge-Exploration of general graphs is 2-competitive and optimal.*

**Exercise 2** *Show that the same competitive ratio holds, if the return to the starting point is not required.*

**Exercise 3** *Consider the problem of exploring the vertices (not the edges) of a graph. If the agent is located at a vertex it detects the outgoing edges but along non-visited edges it is not clear which vertex lies on the opposite side. Does DFS applied on the vertices result in a 2-approximation?*

## 1.4 Exploration of grid environments

Next we consider a simple discrete grid model. The agent runs inside a grid-environment. In contrast to Shannons the inner obstacles consist of full cells instead of single blocked edges.

We would like to design efficient strategies for such grid environments. First, we give a formal definition.

### Definition 1.6

- A **cell**  $c$  is a tuple  $(x, y) \in \mathbf{N}^2$ .
- Two cells  $c_1 = (x_1, y_1), c_2 = (x_2, y_2)$  are **adjacent**, if  $:\Leftrightarrow |x_1 - x_2| + |y_1 - y_2| = 1$ . For a single cell  $c$ , exact 4 cells are adjacent.
- Two cells  $c_1 = (x_1, y_1), c_2 = (x_2, y_2), c_1 \neq c_2$  are **diagonally adjacent**, if  $:\Leftrightarrow |x_1 - x_2| \leq 1 \wedge |y_1 - y_2| \leq 1$ . For a single cell  $c$ , exact 8 cells are diagonally adjacent.
- A **path**  $\pi(s, t)$  from cell  $s$  to cell  $t$  is a sequence of cells  $s = c_1, \dots, c_n = t$  such that  $c_i$  and  $c_{i+1}$  are adjacent for  $i = 1, \dots, n - 1$ .
- A **gridpolygon**  $P$  is a set of path-connected cells, i.e.,  $\forall c_i, c_j \in P : \exists \text{ path } \pi(c_i, c_j)$ , such that  $\pi(c_i, c_j) \in P$  verläuft.

The agent is equipped with a touch sensor so that the agent scans the adjacent cells and their nature (free cell or boundary cell) from its current position. Additionally, the agent has the capability of building a map. The task is to visit all cells of the gridpolygon and return to the start. This problem is NP-hard for known environments; see [IPS82]. We are looking for an efficient Online-Strategy. The agent can move within one step to an adjacent cell. For simplicity we count the number of movements.

The task is related to vacuum-cleaning or lawn-mowing. A cell represents the size of the tool, the tool should visit all cells of the environment. A general polygonal environment  $P$  can be approximated by a grid-polygon.

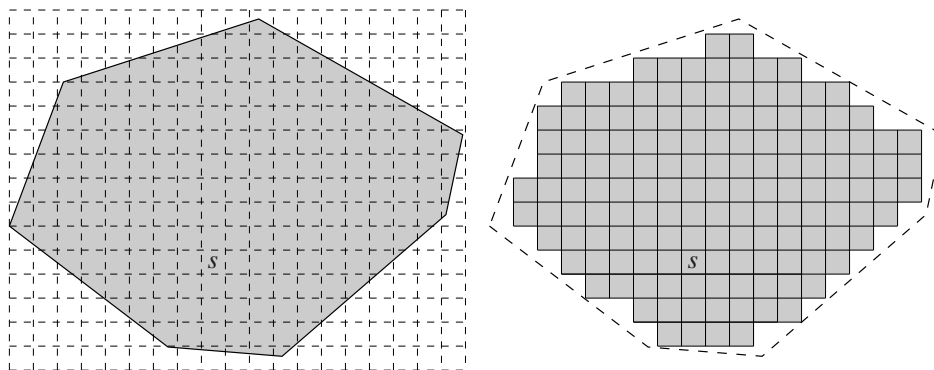


Figure 1.6: A polygon  $P$  and the gridpolygon  $P_{\square}$  as a reasonable approximation.

The starting position and orientation of the tool fixes the grid and all connected cells which are entirely inside  $P$  belong to the approximation  $P_{\square}$ ; see Figure 1.6. For any gridpolygon  $P'$  we use the following notation. Cells that do not belong to  $P'$  but are diagonally adjacent to a cell in  $P'$  are called boundary cells. The common edges of the boundary cells and cells of  $P'$  are the boundary edges. Let  $E(P')$  denote the number of boundary cells or  $E$  for short, if the context is clear. The number of cells is denoted by  $C(P')$  or  $C$  respectively.

From Theorem 1.3 we can already conclude a lower bound of 2 for the competitive ratio of this problem. On the other hand DFS on the cells finishes the task in  $2C - 2$  steps



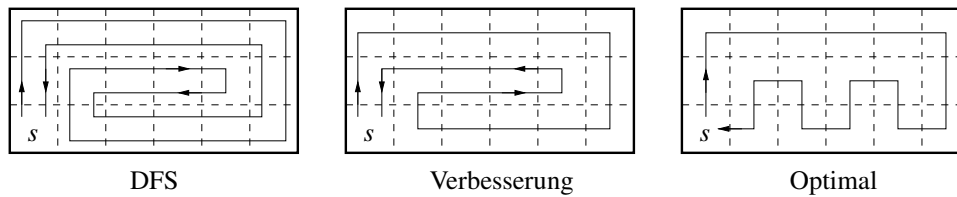


Figure 1.7: Ist DFS optimal?

**Exercise 4** Give a formal proof that for a gridpolygon  $P$  the DFS strategy on the cells requires exactly  $2C - 2$  steps for the exploration (with return to the start) of  $P$ .

But is DFS really the best strategy in general? For fleshy environments DFS obviously is not very efficient. Besides the lower bound construction makes use of corridors only. Compare Figure 1.7: After DFS has visited the *right* neighbour of  $s$  the environment is fully known and we can improve the strategy. It seems that even the optimal solution could be found in an online fashion in this example. On the other hand there are always *skinny* corridor-like environments where DFS is the best online strategy. Altogether, we require a case sensitive measure for the performance of an online strategy that relies on the existence of large areas. The existence of large *fleshy* areas depends on the relationship between the number of cells  $C$  and the number of (boundary) edges  $E$ . In Figure 1.7 the environment has 18 edges and 18 cells. In corridor-like environments we have  $\frac{1}{2}E \approx C$  in fleshy environments we have  $\frac{1}{2}E \ll C$ ; see also Figure 1.8.

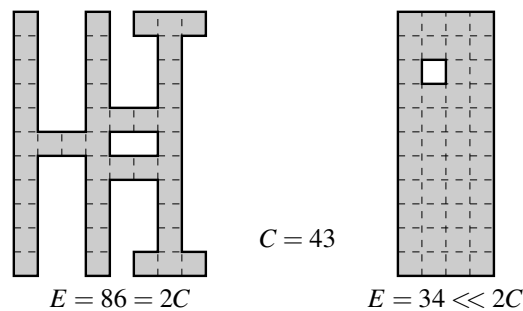


Figure 1.8: The number of boundary edges  $E$  in comparison to the number of cells  $C$  is a measure for the existence of *fleshy* or *skinny* parts.

### 1.4.1 Exploration of simple gridpolygons

We first consider *simple* gridpolygons  $P$  which do not have any *inner* boundary cell, i.e., also the set of all cells that do not belong to  $P$  are path connected.

Note that the lower bound of 2 is not given, because the lower bound construction in the previous section requires the existence of inner obstacles. We make use of a different construction.

**Theorem 1.7** Any online strategy for the exploration (with return to the start) of a simple gridpolygon  $P$  of  $C$  cells, requires at least  $\frac{1}{5}C$  steps for fulfilling the task.

**Proof.** We let the agent start in a corner as depicted in Figure 1.9(i) and successively extend the walls. Assume that the agent decides to move to the east first. By symmetry we apply the same arguments, if the agent moves to the south. For the second step the agent has two possibilities (moving backwards can be ignored). Either the strategy leaves the wall by a step to the south (see Figure 1.9(ii)) or the strategy follows the wall to the east (see Figure 1.9(iii)).

In the first case we close the polygon as shown in Figure 1.9(iv). For this small example the agent requires 8 steps whereas the optimal solution requires only 6 steps which gives a ratio of  $\frac{8}{6} \approx 1.3$ .

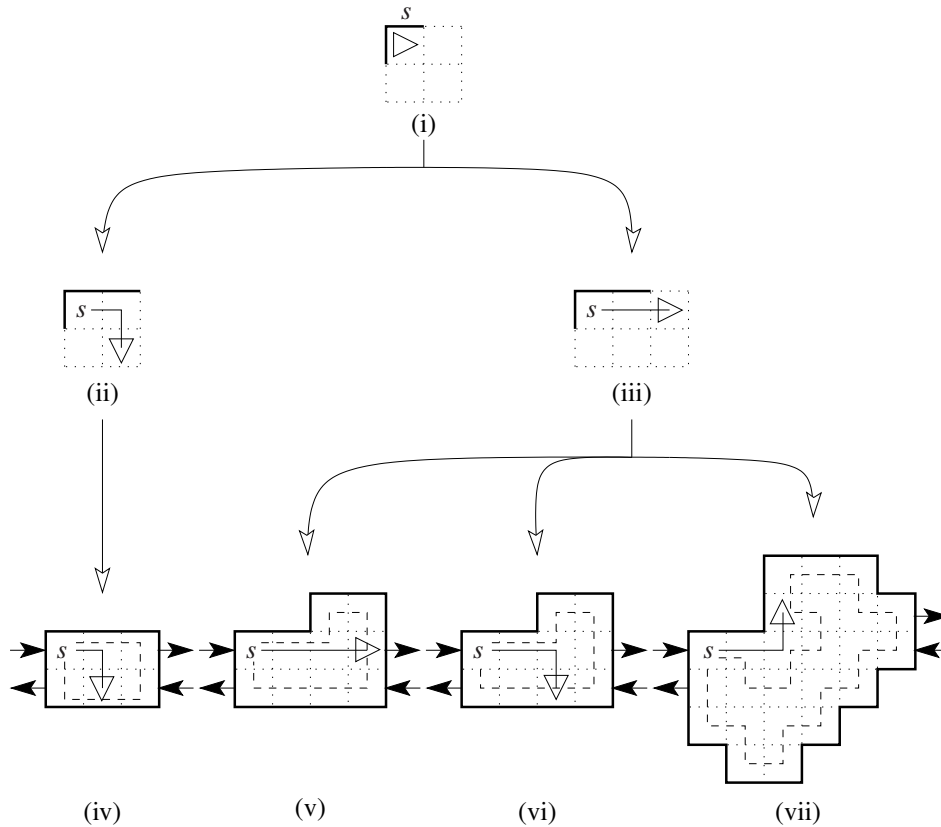


Figure 1.9: A lower bound construction for the exploration of simple gridpolygons.

In the second case we proceed as follows: If the robot leaves the wall (the wall runs upwards), we close the polygon as depicted in Figure 1.9(v) or (vi), respectively. In this small example the agent requires 12, respectively, whereas 10 steps are sufficient.

In the last and most interesting case the agent follows the wall upwards and we present the sophisticated polygon of Figure 1.9(vii). In the offline case an agent requires 24 steps. The online agent already made a mistake and can only finish the task within 24 steps. This can be shown by a tedious case distinction of all further movements. We made use of an implementation that simply checks all possibilities for the next 24 steps. There was no such path that finishes the task. For all cases we guarantee have a worst-case ratio of  $\frac{28}{24} = \frac{7}{6} \approx 1.16$ .

We use this scheme in order to present a lower bound construction of arbitrary size. Any block has an entrance and exit cell which are marked by corresponding arrows; see Figure 1.9(iv)–(vii). If an agent moves inside the next block, the game starts again. Since the arrows only point in east or west direction we take care that the concatenated construction results in a simple gridpolygon of arbitrary size. as required.  $\square$

Note that the arbitrary-size condition in the above proof is necessary. Assume that we can only construct such examples of fixed size  $D$ . This will not result in a lower bound on the competitive ratio. Any reasonable algorithm will explore the fixed environment with komeptitive ratio 1 since  $\alpha \gg D$  exists, with  $|S_{\text{ALG}}| \leq |S_{\text{OPT}}| + \alpha$ .

We consider the exploration of a simple gridpolygon by DFS and formalize the strategy; see Algorithm 1.2. The agent explores the polygon by the “Left-Hand-Rule”, i.e. the DFS preference is Left before Straight-On before Right. The current direction (North, West, East or South) is stored in the variable  $dir$ . The functions  $cw(dir)$ ,  $ccw(dir)$  and  $reverse(dir)$  result in the corresponding directions of a rotation by  $90^\circ$  in clockwise or counter-clockwise order or by a rotation of  $180^\circ$ , respectively. The predicate  $unexplored(dir)$  is true, if the adjacent cell in direction  $dir$  is a cell of the environment, which was not visited yet.

**Algorithm 1.2** DFS**DFS:**

Choose  $dir$ , such that  $reverse(dir)$  is a boundary cell;  
 ExploreCell( $dir$ );

**ExploreCell( $dir$ ):**

// Left-Hand-Rule:  
 ExploreStep(ccw( $dir$ ));  
 ExploreStep( $dir$ );  
 ExploreStep(cw( $dir$ ));

**ExploreStep( $dir$ ):**

**if** unexplored( $dir$ ) **then**  
   move( $dir$ );  
   ExploreCell( $dir$ );  
   move(reverse( $dir$ ));  
**end if**

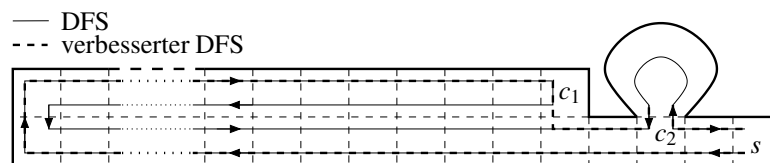


Figure 1.10: First simple improvement of DFS.

A first simple improvement for DFS is as follows:

If there are no unexplored adjacent cells around the current cell, move back along the shortest path (use all already explored cells) to the *last* cell, that still has an unexplored neighbouring cell.

Figure 1.10 sketches this idea: After visiting  $c_1$  the pure DFS will backtrack along the full corridor of width 2 and reach cell  $c_2$  where still something has to be explored. With our improvement we move directly from  $c_1$  to  $c_2$ . Note that for the shortest path we can only make use of the already visited cells. We have no further information about the environment.

By this argument we no longer use the step “move(reverse( $dir$ ))” in the procedure ExploreStep. After the execution of ExploreCell we can no longer conclude that the agent is on the same cell as before. Therefore we store the current position of the agent and use it as a parameter for any call of ExploreStep. The function unexplored( $base, dir$ ) gives “True”, if w.r.t. cell  $base$  there is an unexplored adjacent cell in direction  $dir$ . We re-formalize the behaviour as follows:

**Algorithm 1.3** DFS with optimal return trips**DFS:**

Choose  $dir$ , such that  $reverse(dir)$  is a boundary cell;  
 ExploreCell( $dir$ );  
 Move along the shortest path to the start;

**ExploreCell( $dir$ ):**

$base :=$  current position;  
 // Left-Hand-Rule:  
 ExploreStep( $base$ ,  $ccw(dir)$ );  
 ExploreStep( $base$ ,  $dir$ );  
 ExploreStep( $base$ ,  $cw(dir)$ );

**ExploreStep( $base$ ,  $dir$ ):**

**if** unexplored( $base$ ,  $dir$ ) **then**  
 Move along the shortest path  
 among all visited cells to  $base$ ;  
 move( $dir$ );  
 ExploreCell( $dir$ );  
**end if**

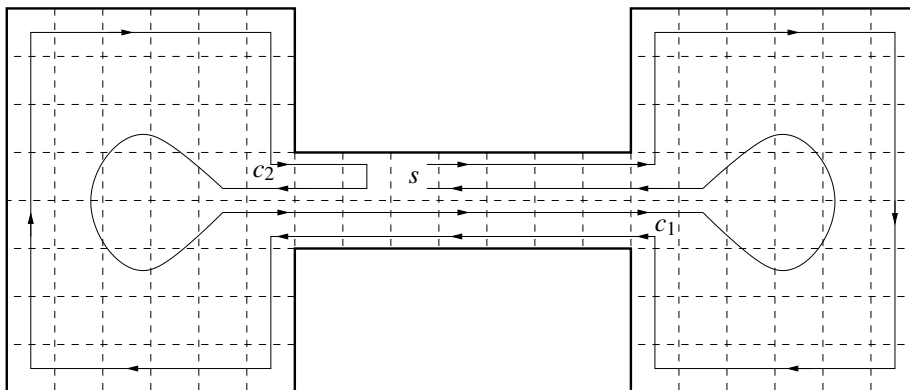


Figure 1.11: Second improvement of DFS.

---

**Algorithm 1.4** SmartDFS

---

**SmartDFS:**

Choose direction  $dir$ , such that  $reverse(dir)$  is a boundary cell;  
ExploreCell( $dir$ );  
Move along the shortest path to the start;

**ExploreCell( $dir$ ):**

Mark the cell with its layernumber;  
 $base :=$  current Position;  
**if** not SplitCell( $base$ ) **then**  
    // Left-Hand-Rule:  
    ExploreStep( $base$ ,  $ccw(dir)$ );  
    ExploreStep( $base$ ,  $dir$ );  
    ExploreStep( $base$ ,  $cw(dir)$ );  
**else**  
    // Choose different order:  
    Calculate the type of the components by the layernumbers  
    of the surrounding cells;  
    **if** No component of typ (III) exists **then**  
        Move one step by the Right-Hand-Rule;  
    **else**  
        Visit the component of type (III) last.  
    **end if**  
**end if**

**ExploreStep( $base$ ,  $dir$ ):**

**if** unexplored( $base$ ,  $dir$ ) **then**  
    Move along the shortest path along  
    the visited cells to  $base$ ;  
    move( $dir$ );  
    ExploreCell( $dir$ );  
**end if**

---

For a second kind of improvement we consider the gridpolygon Figure 1.11. In this example the current DFS variant fully surrounds the polygon. Finally the agent has to move back from  $c_2$  to  $c_1$  so that the corridor of width 2 is visited almost 4 times. Obviously it would be better to first fully explore the component at  $c_1$  move to the other component at  $c_2$  and finally move back to the start. In this case the critical corridor will be visited only once. So, if the exploration splits the polygon into components that have to be considered, we have to take care which component should be visited first.

A cell (like the cell  $c_1$ ) where the remaining polygon definitely splits into different parts is called a **split-cell**. At the first visit of split-cell  $c_1$  in Figure 1.11 it seems to be better to not apply the Left-Hand preference. This depends on the location of the starting point, because we have to move back at the end. The idea can be formulated as follows.

If the unexplored part of the polygon definitely is splitted into different components (i.e., the graph of unexplored cells is splitted into different components), try to visit the unexplored part that does not contain the starting point.

This idea leads to the Algorithm 1.4 (SmartDFS). It remains to decide, which component actually *contains* the starting point. For this we introduce some notions. Until the first split happens we apply the Left-Hand-Rule and successively explore the polygon layer by layer from the outer boundary to the inner parts. We require a formal definition of the layers.

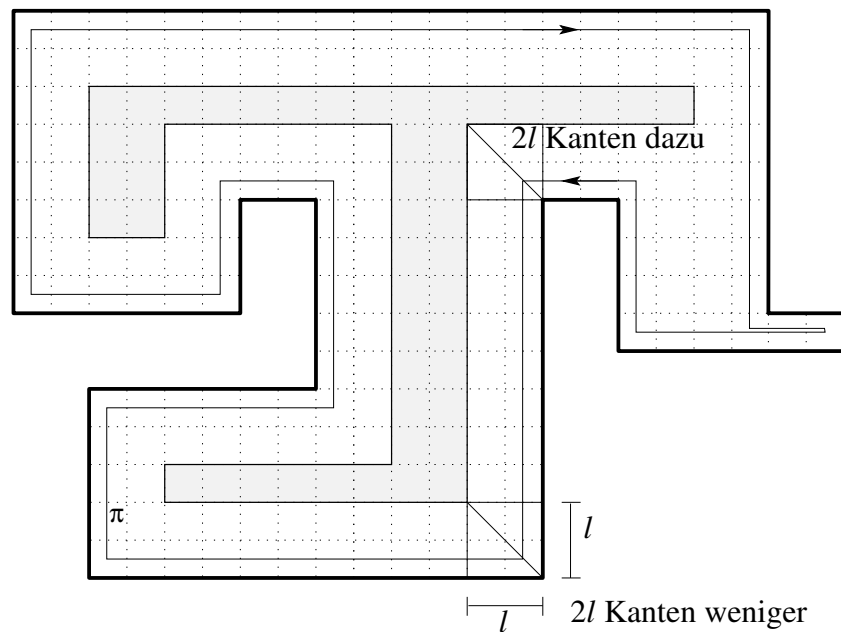


Figure 1.12: The  $l$ -Offset of gridpolygon  $P$ .

**Definition 1.8** Let  $P$  be a (simple) gridpolygon. The cells of  $P$  that share a boundary edge belong to the first layer, the **1-Layer** of  $P$ . The gridpolygon that stems from  $P$  without the 1-Layer is called the **1-Offset** of  $P$ . Recursively, the **2-Layer** of  $P$ , is the **1-Layer** of the **1-Offset** of  $P$  and the **2-Offset** of  $P$  is the **1-Offset** of the **1-Offset** of  $P$  and so on.

Note that the  $l$ -Offset of a gridpolygon need not be connected and finally the Offsets will decrease to an empty polygon. The definition is totally independent from any strategy. Fortunately, during the execution of SmartDFS on a simple gridpolygon, we can successively mark and store the layers for any visited cell. The  $l$ -Offset has an interesting property.

**Lemma 1.9** The non-empty  $l$ -Offset of a simple gridpolygon  $P$  has at least  $8l$  edges less than  $P$ .

**Proof.** We surround the boundary of the gridpolygon in clockwise order and visit all boundary edges along this path. Let us assume that the offset remains a single component. For a left turn the  $\ell$ -Offset  $2\ell$  loses  $2\ell$  edges for a right turn the  $\ell$ -Offset  $2\ell$  wins  $2\ell$  edges. We can show that there are 4 more right turns than left turns. So the  $\ell$ -Offset has at least  $8\ell$  edges less than  $P$ . Even more edges will be cancelled, if the polygon fell into pieces.  $\square$

**Exercise 5** Show that for any surrounding of the boundary of a simple gridpolygon in clockwise order there are 4 more right turns than left turns. Make use of induction.

**Exercise 6** Show that in the above proof the non-empty  $\ell$ -Offset will lose even more edges, if it consists of more than one connected component. Show the statement for the 1-Offset.

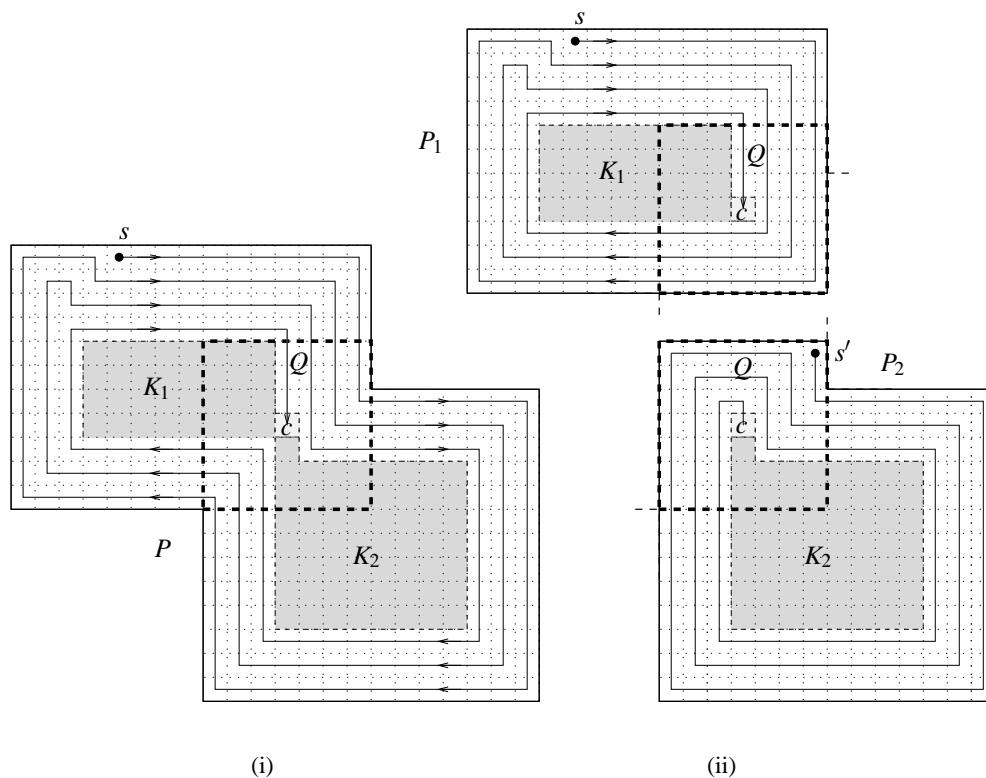


Figure 1.13: Decomposition at a split-cell.

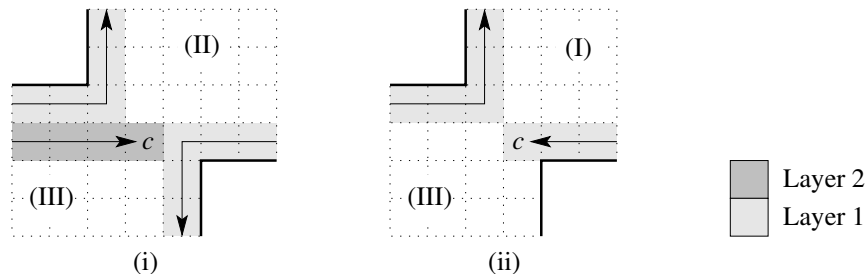


Figure 1.14: Three types of components.

We consider Figure 1.13(i): In the 4. Layer for the first time a split-cell  $c$  occurs. Now we decompose the polygon into different components<sup>2</sup>:

<sup>2</sup>Let  $A \dot{\cup} B$  denote the **disjoint union**  $A \dot{\cup} B = A \cup B$  mit  $A \cap B = \emptyset$ .

$$P = K_1 \dot{\cup} K_2 \dot{\cup} \{ \text{visited cells of } P \},$$

where  $K_1$  denotes the component that was visited last. SmartDFS recursively works on  $K_2$ , returns to  $c$  and proceeds with  $K_1$ .

By the layernumbers we would like to avoid the situation of Figure 1.11. We will find the split-cell in layer  $\ell$ , which gives three types of components; see Figure 1.14:

- (I) Component  $K_i$  is *fully* surrounded by layer  $\ell$ .
- (II) Component  $K_i$  is *not* surrounded by layer  $\ell$  (may be touched by the split-cell only).
- (III) Component  $K_i$  is *partly* surrounded by layer  $\ell$  (not only touched by the split cell).

Obviously, if a split-cell occurs, we should visit the component of type (III) last because the starting point lies in the outer layers of this component.

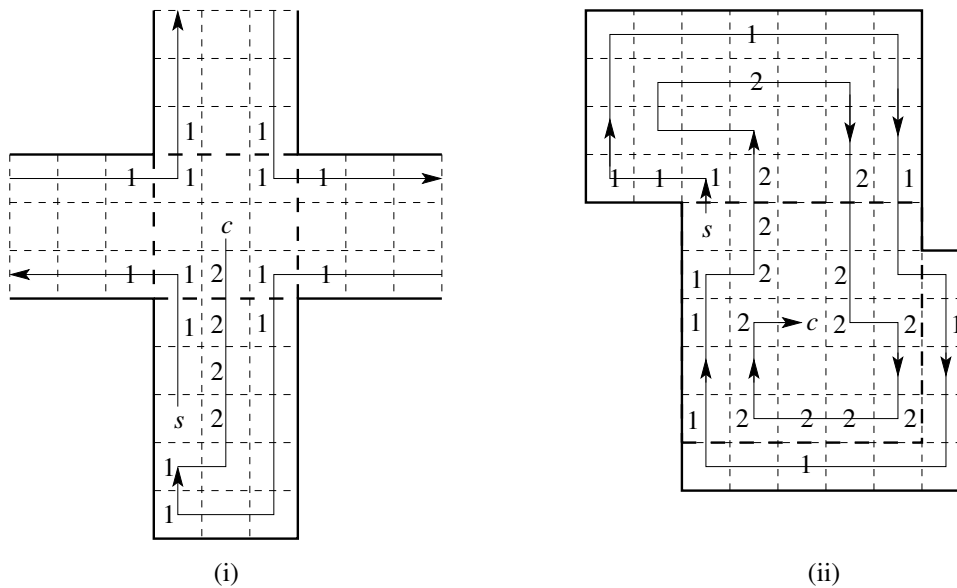


Figure 1.15: Special cases: No component of type (III) exists.

There are some situations where a component of type (III) does not exist. For example if the split-cell is the first cell on the next layer, or the component of the starting point was just explored (efficiently). More precisely:

- (a) The component with the starting point on its layer was just fully explored in the current layer; see Figure 1.15(i). In this case the order of visiting the remaining components is not critical, we can choose an arbitrary order. This example also shows that at a split-cell more than two components has to be visited. We simply apply one the next step by changing to the Right-Hand-Rule.
- (b) Two components have been fully surrounded, because at the split-cell we change from layer  $\ell$  to  $\ell + 1$ ; see Figure 1.15(ii). In all other cases at least one additional visited cell is marked with layer number of the split-cell. We can conclude that layer  $\ell$  was closed with the split-cell. This means that the starting point is not part of the layer of the component where the agent currently comes from. Because the agent normally moves by the Left-Hand-Rule, it suffices to apply the Right-Hand-Rule in this case also.

Altogether in both cases we simply apply the Right-Hand-Rule for a single step.

For the overall analysis at a split-cell we consider two polygons  $P_1$  and  $P_2$  as depicted in Figure 1.13(i). Here we detect the component of type (III).  $K_2$  is a component of type (II). Let  $Q$  be a



rectangle of edge length (width or height)  $2q + 1$  around the split cell  $c$  so that

$$q := \begin{cases} \ell, & \text{if } K_2 \text{ has type (I)} \\ \ell - 1, & \text{if } K_2 \text{ has type (II)} \end{cases}.$$

Now choose  $P_2$  so that  $K_2 \cup \{c\}$  is the  $q$ -Offset of  $K_2 \cup \{c\}$ . The idea is that the rectangle  $Q$  will be *added* so that  $P_2$  has the desired form. Now let  $P_1 := ((P \setminus P_2) \cup Q) \cap P$ , comp. Figure 1.13. The intersection with  $P$  is necessary, since there are cases where  $Q$  does not totally fit into  $P$ . We would like to apply arguments recursively for  $P_2$  and  $P_1$ . Let us consider them separately as shown in Figure 1.13(ii). We have chosen  $P_1, P_2$  and  $Q$  in a way so that the paths in  $P_1 \setminus Q$  and  $P_2 \setminus Q$  did not change w.r.t. the paths already performed for  $P^3$ . The already performed paths that lead in  $P$  from  $P_1$  to  $P_2$  and from  $P_2$  to  $P_1$  will be used and adapted so that the paths outside  $Q$  will not change; see Figure 1.13. We can consider  $P_1$  and  $P_2$  separately.

We know that any cell has to be visited at least once. Therefore we count the number of steps  $S(P)$  for polygons  $P$  as follows. It is the sum of the cells,  $C(P)$ , of  $P$  plus the extra cost  $excess(P)$  for the overall *path* length.

$$S(P) := C(P) + excess(P).$$

The following Lemma gives an estimate for the extra cost w.r.t. the above decomposition around a split-cell.

**Lemma 1.10** *Let  $P$  be a gridpolygon,  $c$  a split-cell, so that two remaining components  $K_1$  and  $K_2$  has to be considered. Assume that  $K_2$  is visited first. We conclude:*

$$excess(P) \leq excess(P_1) + excess(K_2 \cup \{c\}) + 1.$$

**Proof.** The agent is located at cell  $c$  and decides to explore  $K_2 \cup \{c\}$  starting from  $c$  and return to  $c$ . This gives additional cost at most  $excess(K_2 \cup \{c\})$ , note that the part  $P_2 \setminus (K_2 \cup \{c\})$  can only help for the return path. Because  $c$  was already visited, we count one additional item for the excess of visited cells. After that we proceed with the exploration of  $P_1$  and require  $excess(P_1)$  for this part.  $\square$

For the full analysis of SmartDFS we have to prove some structural properties:

**Lemma 1.11** *The shortest path between to cells  $s$  and  $t$  in a simple gridpolygon  $P$  with  $E(P)$  boundary edges consists of at most  $\frac{1}{2}E(P) - 2$  cells.*

**Proof.** W.l.o.g. we assume that  $s$  and  $t$  are in the first layer, otherwise we can choose different  $s$  or  $t$  whose shortest path is even a bit longer. Consider the path,  $\pi_L$ , in clockwise order in the first layer from  $s$  to  $t$  and the path,  $\pi_R$ , in counter-clockwise order in the first layer from  $s$  to  $t$ . Connecting  $\pi_L$  and  $\pi_R$  gives a full roundtrip. As in the proof of Lemma 1.9 counting the edges gives 4 more edges than cells which gives

$$|\pi_R| + |\pi_L| \leq E(P) - 4$$

visited cells.

In the worst case both path have the same length, which gives  $|\pi(s, t)| = |\pi_R| = |\pi_L|$ , and  $2|\pi(s, t)| \leq E(P) - 4 \Rightarrow |\pi(s, t)| \leq \frac{1}{2}E(P) - 2$ .  $\square$

**Lemma 1.12** *Let  $P$  be a gridpolygon and let  $c$  be a split-cell. Define  $P_1, P_2$  and  $Q$  as above. For the number of edges we have:*

$$E(P_1) + E(P_2) = E(P) + E(Q).$$

---

<sup>3</sup>For the uniqueness of this decomposition into  $P_1$  and  $P_2$  we remark that  $P_1$  and  $P_2$  are connected, respectively and  $P \cup Q = P_1 \cup P_2$  and  $P_1 \cap P_2 \subseteq Q$  holds.

**Proof.** For arbitrary gridpolygons  $P_1$  and  $P_2$  we conclude

$$E(P_1) + E(P_2) = E(P_1 \cup P_2) + E(P_1 \cap P_2).$$

Let  $Q' := P_1 \cap P_2$ , we have:

$$\begin{aligned} E(P_1) + E(P_2) &= E(P_1 \cap P_2) + E(P_1 \cup P_2) \\ &= E(Q') + E(P \cup Q) \\ &= E(Q') + E(P) + E(Q) - E(P \cap Q) \\ &= E(P) + E(Q), \text{ since } Q' = P \cap Q \end{aligned}$$

□

**Exercise 7** Show that for arbitrary two gridpolygons  $P_1$  and  $P_2$  we have  $E(P_1) + E(P_2) = E(P_1 \cup P_2) + E(P_1 \cap P_2)$ .

Using all these arguments we can show:

**Theorem 1.13** (Icking, Kamphans, Klein, Langetepe, 2000)

For a simple gridpolygon  $P$  with  $C$  cells and  $E$  boundary edges the strategy SmartDFS required no more than

$$C + \frac{1}{2}E - 3$$

for the exploration of  $P$  (with return to the start). This bound will be attained exactly in some environments. [IKKL00b]

**Proof.** By the above arguments it suffices to show  $excess(P) \leq \frac{1}{2}E - 3$ . We give a proof by induction on the number of components.

**Induction base:**

Assume that there is no split-cell. For the exploration of a single component, SmartDFS visits all cells exactly once and return to the start. For visiting all cells we require  $C - 1$  steps. Now the excess is the shortest path back. By Lemma 1.11  $\frac{1}{2}E - 2$  steps suffices which gives the conclusion

**Induction step:**

Consider the (first) decomposition at a split-cell  $c$ . Let  $K_1, K_2, P_1, P_2, Q$  be defined as above, assume that  $K_2$  is visited last. We have:

$$\begin{aligned} excess(P) &\leq excess(P_1) + excess(K_2 \cup \{c\}) + 1 \text{ (Lemma 1.10)} \\ &\leq_{(1.A)} \frac{1}{2}E(P_1) - 3 + \frac{1}{2} \underbrace{E(K_2 \cup \{c\})}_{\leq E(P_2) - 8q} - 3 + 1 \text{ (Lemma 1.9)} \\ &\leq \frac{1}{2} \left[ \underbrace{E(P_1) + E(P_2)}_{\leq E(P) + 4(2q+1)} \right] - 4q - 5 \text{ (1.12, Def. of } Q) \\ &\leq \frac{1}{2}E(P) - 3 \end{aligned}$$

□

A Java-Applet for the Simulation of SmartDFS and different strategies can be found at:

<http://www.geometrylab.de/>

Finally, we would like to show, how to compute the offline shortest paths in gridpolygons. Of course the Dijkstra algorithm can also be applied on the gridgraph, but this algorithm does not use the grid structure directly. As an alternative we apply Algorithm 1.5 (C. Y. Lee, 1961, [Lee61]), the running time is only linear in the number of overall cells. The algorithm simulates a wave propagation starting from the goal. Any cell will be marked with a label indicating the distance to the goal. Obstacles *slow down* the propagation a bit; see Figure 1.16. When the wave reaches the starting point  $s$ , we are done with the first phase. For computing the path we start at  $s$  and move along cells with strictly decreasing labels. Obviously, the shortest path need not be unique.

---

**Algorithm 1.5** Algorithm of Lee
 

---

 Shortest path from  $s$  to  $t$  in a gridpolygon
 

---

```

Datastructure: Queue  $Q$ 
// Initialise
 $Q.InsertItem(t)$ ;
Mark  $t$  with label 0;
// Wave propagation:
loop
   $c := Q.RemoveItem()$ ;
  for all Cells  $x$  such that  $x$  is adjacent to  $c$  and  $x$  is not marked do
    Mark  $x$  with the label of  $label(c) + 1$ ;
     $Q.InsertItem(x)$ ;
    if  $x = s$  then break loop;
  end for
end loop
// Backtrace:
Move along cells with strongly decreasing labels from  $s$  to  $t$ .

```

---

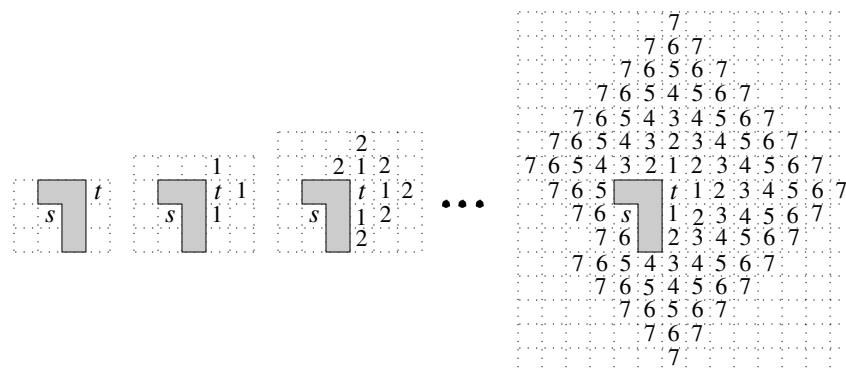


Figure 1.16: Wave-Propagation.

### 1.4.2 Competitive ratio of SmartDFS

The corridor of width 3, see Figure 1.7, indicates that the competitive ratio of SmartDFS should be better than 2. SmartDFS runs 4 times though the corridor whereas the shortest path visits any cell only once. This gives roughly a ratio of  $\frac{4}{3}$ . We will show that this is the worst-case for SmartDFS. The gap between  $e^{\frac{7}{6}}$  and  $\frac{4}{3}$  is small.

For the analysis we first give a precise definition of the structure of parts of gridpolygons which will be explored in an optimal fashion. The SmartDFS Strategy does not make any detours within these passages.

For a *corridor* of widths 1 this is obviously true. But also corridors of width 2 will be passed optimally, since SmartDFS runs forth and back along different tracks; see Figure 1.17. We give a formal definition of the *narrow passages*.

**Definition 1.14** The set of cells that can be deleted such that the layernumber of the remaining cells do not change are called narrow passages of  $P$ .

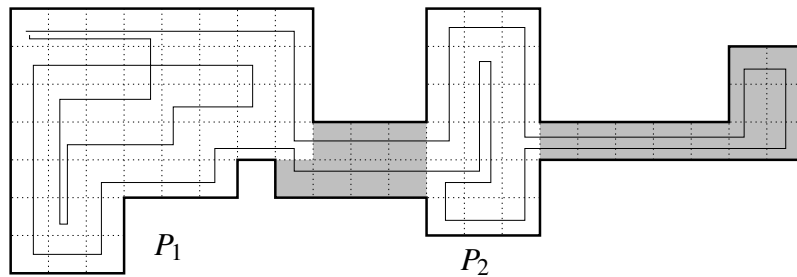


Figure 1.17: SmartDFS is optimal in narrow passages.

SmartDFS passes narrow passages optimally since they allow an optimal forth and back pass-through. There are no additional detours at the entrance and exit of a narrow passage because they consist of cells in the first layer. They can be considered as *gates*. The entrance and exit is always precisely determined.

The idea is to consider polygons without narrow passages first. There is a fixed relationship between edges and cells.

**Lemma 1.15** Let  $P$  be a simple gridpolygon without narrow passages and without a split-cell in the first layer. We have

$$E(P) \leq \frac{2}{3}C(P) + 6.$$

**Proof.** A  $3 \times 3$  gridpolygon has precisely this property,  $C(P) = 9$  und  $E(P) = 12$ . Any gridpolygon with the above conditions can be reduced by successively removing columns or rows such that in each step the property remains true and such that always at least 3 cells and at most 2 edges will be removed. This is an exercise below.

Starting backwards from the property  $E(P_0) = \frac{2}{3}C(P_0) + 6$  we will maintain the bound  $E(P_i) \leq \frac{2}{3}C(P_i) + 6$  since we add at least 3 cells and add at most 2 edges. Finally,  $E(P) \leq \frac{2}{3}C(P) + 6$  holds.  $\square$

First, we show that the overall number of exploration steps of SmartDFS decreases for the given class of polygons.

**Lemma 1.16** A simple gridpolygon  $P$  with  $E(P)$  edges and  $C(P)$  cells, without narrow passages and without a split-cell in the first layer will be explored by SmartDFS with no more than  $S(P) \leq C(P) + \frac{1}{2}E(P) - 5$  steps.



$$\begin{aligned}
&\leq C(P_i) + \frac{1}{2} \left( \frac{2}{3} C(P_i) + 6 \right) - 5 \\
&= \frac{4}{3} C(P_i) - 2.
\end{aligned}$$

**Induktion-Step:** If there is no split-cell in the first layer we can apply the same arguments as above. Therefore, we assume that the first split occurs in the first layer. Two cases can occur as depicted in Figure 1.20.

In the first case the component of type (II) was not visited before and we define  $Q := \{c\}$ . The second case occurs, if the split-cell  $c$  is diagonally adjacent to a cell  $c'$ ; compare Figure 1.20(ii), (iii) and (iv). We build the smallest rectangle  $Q$  that contains  $c$  and  $c'$ . In case (ii) and (iii)  $Q$  is a square of size 4. In case (iv) by simple adjacency  $Q$  is a rectangle and  $|Q| = 2$ .

Analogously to the proof of Theorem 1.13 we split the polygons into parts  $P'$  and  $P''$  both containing  $Q$ .

Here  $P''$  is of type (I) or (II) and  $P'$  the remaining polygon. das Polygon der Komponenten vom Typ (I) oder (II) und  $P'$  das andere.

For  $|Q| = 1$  (see Figure 1.20(i)) we have  $S(P_i) = S(P') + S(P'')$  and  $C(P_i) = C(P') + C(P'') - 1$ . We apply the induction hypothesis on  $P'$  and  $P''$  (they have one split-cell less) and obtain:

$$\begin{aligned}
S(P_i) &= S(P') + S(P'') \\
&\leq \frac{4}{3} C(P') - 2 + \frac{4}{3} C(P'') - 2 \\
&\leq \frac{4}{3} C(P_i) + \frac{4}{3} - 4 < \frac{4}{3} C(P_i) - 2.
\end{aligned}$$

For  $|Q| = 4$  we argue that by the union we will save some steps that will occur for the separate explorations. We consider  $P'$  and  $P''$  separately, first. The movements from  $c'$  to  $c$  (and  $c$  to  $c'$ ) count in both polygons. For the complete  $P_i$  the path from  $c'$  to  $c$  (and  $c$  to  $c'$ ) are given either  $P'$  or in  $P''$ , this means that we save  $4 = |Q|$  steps.

We have  $S(P_i) = S(P') + S(P'') - 4$  and  $C(P_i) = C(P') + C(P'') - 4$ . By induction hypothesis for  $P'$  and  $P''$  we conclude:

$$\begin{aligned}
S(P_i) &= S(P') + S(P'') - 4 \\
&\leq \frac{4}{3} C(P') + \frac{4}{3} C(P'') - 8 \\
&= \frac{4}{3} (C(P') + C(P'') - 4) - \frac{8}{3} \\
&< \frac{4}{3} C(P_i) - 2.
\end{aligned}$$

The case  $|Q| = 2$  is left as an exercise.

Altogether an optimal strategy requires  $\geq C(P_i)$  steps or  $\geq C(P)$  in total and we have a competitive ratio of  $\frac{4}{3}$ .  $\square$

**Exercise 8** Analyse the remaining case  $|Q| = 2$  in the above proof.

If we compare the result to Theorem 1.7 there is a gap of size  $\frac{1}{6}$  between  $\frac{7}{6}$  and  $\frac{4}{3}$ . Recently, both parts have been improved. There is a lower bound of  $\frac{20}{17}$  and an upper bound of  $\frac{5}{4}$  shown by Kolenderska et. al 2010. In principle the strategy is a local improvement of SmartDFS and the lower bound is an extension of our construction. The result comes along with a tedious case analysis.

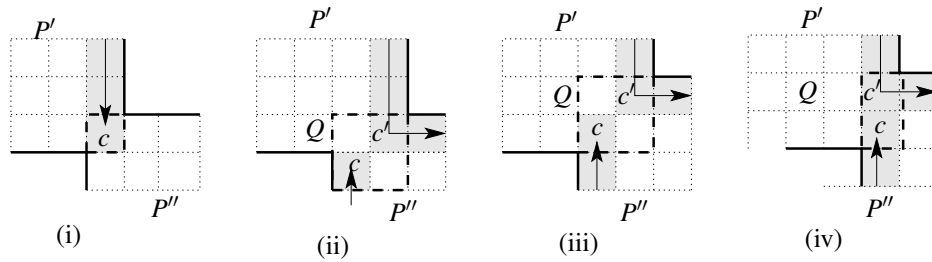


Figure 1.20: A gridpolygon  $P_i$  that is separated into components of type (I) or (II) at the split-cell. The rectangle  $Q$  is always inside  $P_i$ .

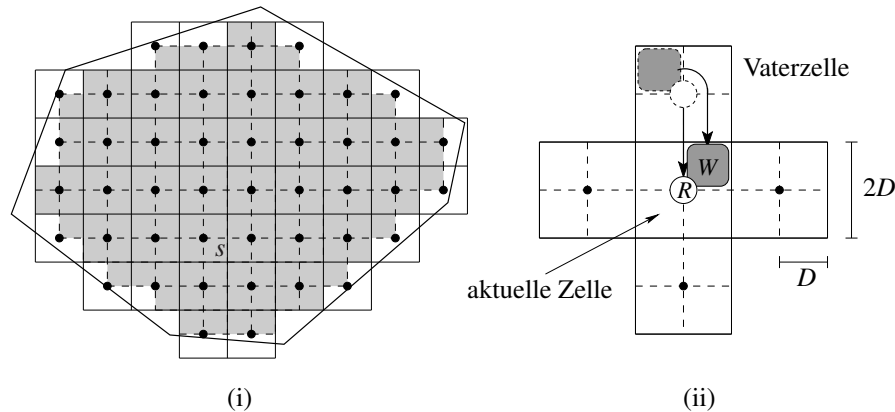


Figure 1.21:  $2D$ -cells and  $D \times D$  sub-cells.

### 1.4.3 Exploration of general gridpolygons

For the more general exploration of gridpolygons we first slightly change the model<sup>4</sup>: We consider an agent that is located at the center of 4 cells of size  $D \times D$ . The tool for the exploration still has size  $D \times D$  as before and moves freely around the agent. More precisely, we consider 4 **sub-cells** of size  $D \times D$  and unify them to a  $2D$ -cell<sup>5</sup>; see Figure 1.21(i).

It can happen that for the  $2D$ -cell, not all sub-cells belong to the initial gridpolygon, since some of the sub-cells simply belong to the boundary. Such  $2D$ -cell are denoted as **partially occupied cells**.

In Figure 1.21(i) all cells intersected by the original polygonal segments are partially occupied (compare also [reffigfigOnline/PolyToGrid](#) on page 8). The agent is always located in the center of the  $2D$ -cell. Analogously to the SmartDFS model, the agents scans the four adjacent  $2D$ -cells. The tool moves freely around the agent, we would like the count the number of steps of the tool; see also Figure 1.21(ii).

The current cell of the agent is denoted as *current cell*. The *parent cell* of the agent is the cell where he is actually coming from. In the beginning we initially an arbitrary adjacent  $2D$ -cell as the parent cell.

The strategy “Spanning-Tree-Covering” (STC) constructs a spanning tree for all connected  $2D$ -cells that are also not occupied. The tool moves along the spanning tree by the Left-Hand-Rule. The construction can be done fully online. The  $2D$ -cells are detected by the Right-Hand-Rule. Obviously by this approach the tool exactly visits any cell at most once by following the spanning tree. Figure 1.22(i) shows an example for the efficient exploration of all non-occupied cells by  $2D$ -Spiral-STC. As mentioned before, for the start we can choose an arbitrary parent cell.

The disadvantage of  $2D$ -Spiral-STC is, that we do not visit sub-cells by that tool which actually lie in the connected component of the sub-cells. Now we relax the behaviour of  $2D$ -Spiral-STC. The strategy

<sup>4</sup>We will see later that the change was only done for the reason of a convenient analysis and description.

<sup>5</sup>In the following a cell always denotes a  $2D$ -cell.

---

**Algorithm 1.6** 2D-Spiral-STC

---

**2DSPSTC**( *parent*, *current* ):Mark *current* as visited.**while** *current* has unvisited neighbour cell **do**

- From *parent* search in ccw order for a neighbouring cell *free*, which is not marked as visited and is not partially occupied.
- Build the spanning tree edge from *current* to *free*.
- Move the tool by Left-Hand-Rule along the spanning tree edge to the first sub-cell of *free*.
- Call 2DSPSTC( *current*, *free* ).

**end while****if** *current*  $\neq$  *s* **then**

- Move by the Left-Hand-Rule along the spanning tree edge back from *current* to the first sub-cell of *parent*.

**end if**

---

---

**Algorithm 1.7** SpiralSTC

---

**SPSTC**( *parent*, *current* ):Mark *current* as visited.**while** *current* has unvisited neighbour cell **do**

- From *parent* search in ccw order for the first neighbouring cell *free*.
- Build a spanning tree edges from *current* to *free*.
- Move the tool along the spanning tree edge to the first sub-cell of *free*. The movement depends on the local situation. For double-sided edges the tool moves by Left-Hand-Rule along the edge. For single-sided edges the tool might change to the other (left) side of the spanning tree edge in order to avoid an occupied sub-cell for reaching the corresponding sub-cell.
- Call SPSTC( *current*, *free* ).

**end while****if** *current*  $\neq$  *s* **then**

- Move along the spanning tree edge back from *current* to the first possible sub-cell of *parent*. The movement depends on the type of the edge, as mentioned above.

**end if**

---



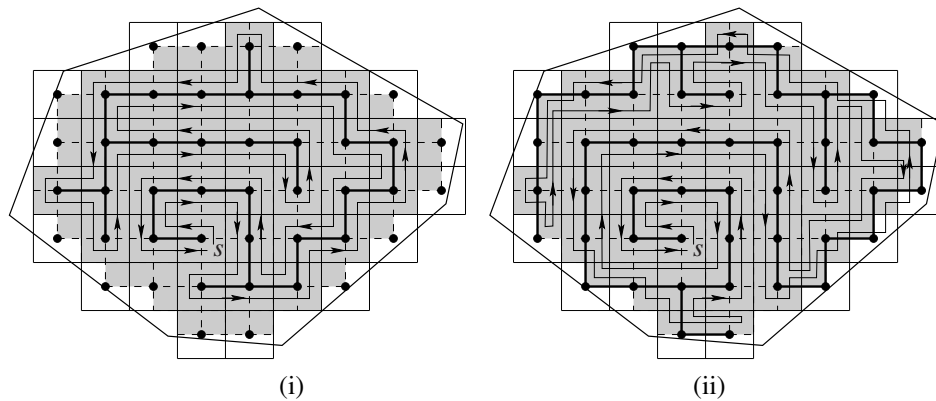


Figure 1.22: Examples for (i) 2D-Spiral-STC and (ii) Spiral-STC.

Spiral-STC (Algorithm 1.7) also constructs a spanning tree in an online fashion. But we also insert a corresponding edge if a partially occupied 2D-cell contains sub-cells that are still reachable by the tool. In this case the tool cannot always move the the Left-Hand-Rule along the spanning tree edge. The tool has to avoid occupied sub-cells and visits some sub-cells more than once. For systematically analysing the corresponding additional sub-cell visits of the tool we make use of the following notion:

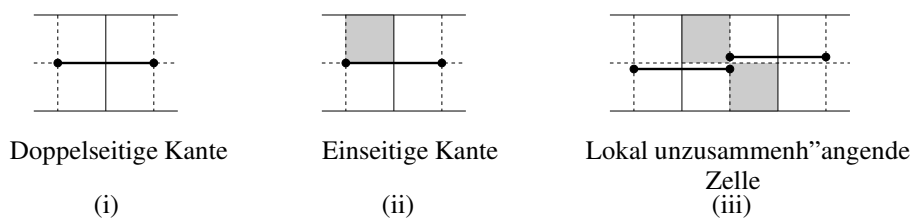


Figure 1.23: (i) Double-sided edge, (ii) one-sided edge, (iii) locally disconnected 2D-cell.

**Definition 1.18** A spanning tree edge constructed by STC in a gridpolygon  $P$  is denoted as

- (i) **double-sided edge**, if all adjacent sub-cells belong to the gridpolygon  $P$  (Figure 1.23(i)),
- (ii) **single-sided edge**, if at least one of the adjacent sub-cells is a boundary sub-cell of  $P$  (Figure 1.23(ii)).

Double-sided edges are handled in the same way as in the 2D-Spiral-STC strategy. Single-sided edges impose a detour for the tool, some sub-cells will be visited more than once since the tool changes to the other side of the spanning tree edge. For the analysis we will consider the corresponding cases systematically. A special case occurs, if the situation imposes two spanning tree edges for the same cell from different directions. The cell is locally disconnected in this case; see Figure 1.23(iii). This 2D-cell will be visited twice from different directions. For simplicity we internally double the corresponding vertex and the spanning tree has exactly one incoming edge for any vertex. For the analysis we have to take care that we count the cell only once. An example of the execution of Spiral-STC is shown in Figure 1.22(ii).

By the preference rule for the 2D-cells the Spiral-STC constructs spanning trees with many windings. This is not always intended, especially for lawn-mowing or vacuum-cleaning a tool should try to avoid so many turns. The number of turns might also be part of the cost model. The Scan-STC variant has a fixed given preference for vertical or horizontal edges. We would like to make local decision for the construction of spanning tree edges. In our examples we prefer a vertical scan of the gridpolygon. For this we extend the sensor model and allow to have information about all diagonally adjacent 2D-cells of a current cell.

The idea is that the construction of a horizontal edge will be postponed, if it is clear that we can also reach the 2D-cell by another vertical spanning tree edge. To keep the rule simple we only look ahead as

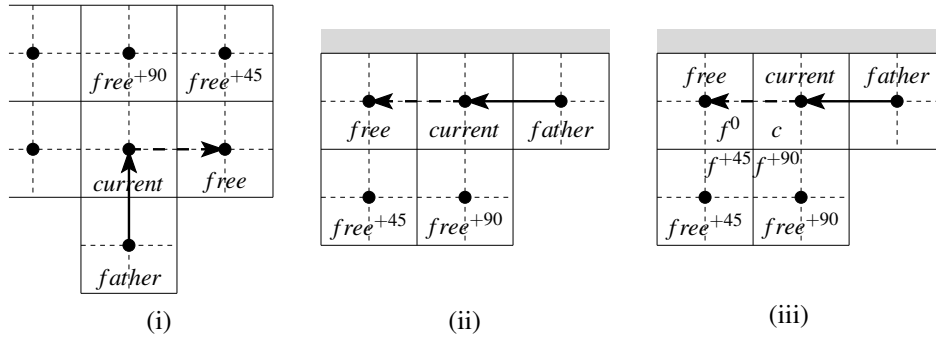


Figure 1.24: Avoid horizontal edges with the Scan-STC.

indicated in Figure 1.24 (i) and (ii). Here we currently would like to build a horizontal edge. The agent is located at cell *current* and is looking (in ccw order starting from *father*) for the first free cell *free*. If there is a counterclockwise path from *free* over  $free^{+45}$  and  $free^{+90}$  back to the current cell, we change the preference and build a spanning tree edge to  $free^{+90}$ . Here  $free^{+45}$  lies on the same row as *free* and is the next cell in ccw order from *free*.  $free^{+90}$  is the next cell in ccw order from  $free^{+45}$  in the same column as *current*.

If the full turn exists, the cell *free* will also be reached from  $free^{+45}$  by a vertical edge and  $free^{+45}$  can be reached from  $free^{+90}$ . Note that we have extended the sensor model in this case and also have information about diagonally adjacent edges.

Analogously, we can also consider partially occupied 2D-cells and apply the same idea. For the corresponding avoidance rule we consider the sub-cells  $c$ ,  $f^0$ ,  $f^{+45}$  and  $f^{+90}$  instead of the cells *current*, *free*,  $free^{+45}$  and  $free^{+90}$ ; see Figure 1.24(iii).

By the above idea we could define a strategy 2D-Scan-STC that corresponds to 2D-Spiral-STC. We skip this step and directly define a Scan-STC Algorithm that makes use of the sub-cells  $c$ ,  $f^0$ ,  $f^{+45}$  and  $f^{+90}$  by the same arguments. If  $f^{+45}$  and  $f^{+90}$  are also free, we will reach  $f^0$  from  $f^{+45}$  and in turn  $f^{+45}$  from  $f^{+90}$ . Algorithm 1.8 summarizes this behaviour.

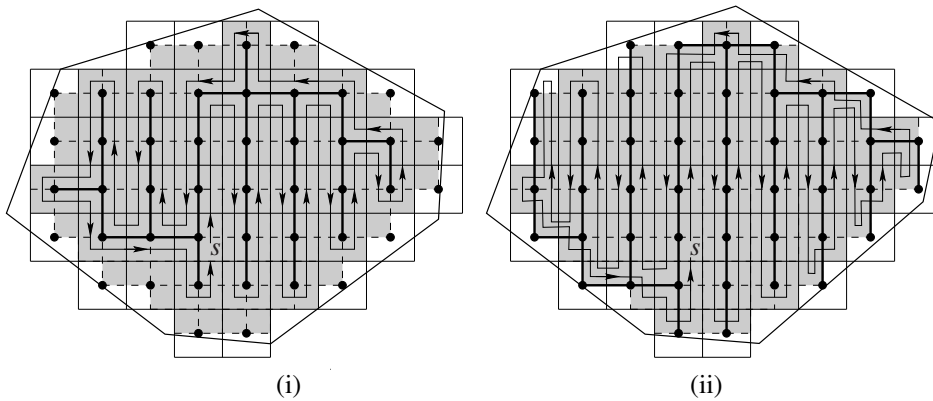


Figure 1.25: Example for (i) 2D-Scan-STC, (ii) Scan-STC.

**Algorithm 1.8** ScanSTCSCSTC(*parent*, *current*):Mark *current* as visited.**while** *current* has unvisited neighbouring cell **do**

- From *parent* search in ccw order for the first non-visited neighbouring cell *free*.

- **if** Spanning tree edge from *current* to *free* is horizontal and sub-cells  $f^{+45}$  and  $f^{+90}$  are free **then**  
 $free := free^{+90}$ .

**end if**

- Build a spanning tree edge from *current* to *free*.

- Move the tool along the spanning tree edge to the first sub-cell of *free*. The movement depends on the local situation. For double-sided edges the tool moves by Left-Hand-Rule along the edge. For single-sided edges the tool might change to the other (left) side of the spanning tree edge in order to avoid an occupied sub-cell and reach the corresponding sub-cell.

- Call SCSTC(*current*, *free*) auf.

**end while****if** *current*  $\neq s$  **then**

- Move along the spanning tree edge from *current* back to the first possible sub-cell of *parent*. The movement depends on the type of the edge, as mentioned above.

**end if****Theorem 1.19** (Gabriely, Rimon, 2000)

Let  $P$  be a gridpolygon with  $C$  sub-cells. Let  $K$  be the number of all sub-cells, which are diagonally adjacent to an occupied (boundary) sub-cell<sup>6</sup>. The gridpolygons  $P$  will be explored by Spiral-STC and Scan-STC in time  $O(C)$  and space  $O(C)$ . There are no more than

$$S \leq C + K$$

exploration steps,  $S$ , for the tool.

[GR03]

**Proof.****Correctness:**

Both algorithms construct a spanning tree by DFS such that any 2D-cell which has reachable  $D$  sub-cells will be visited. The tool moves along the spanning tree on both sides – as long as the path is not blocked – and visits all sub-cells that are *touched* by the spanning tree.

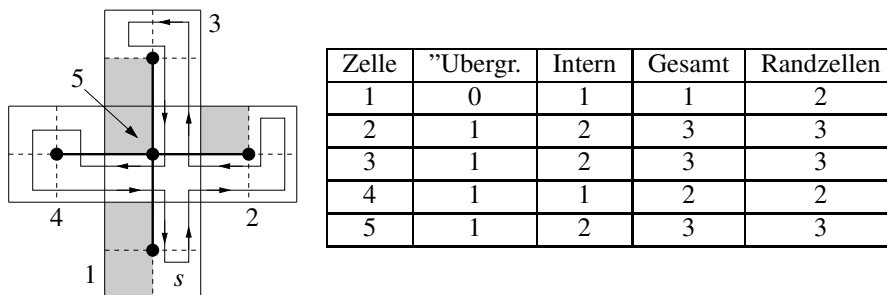


Figure 1.26: Estimating the double visits of sub-cells by STC locally.

**Path length:**

The number of steps for the tool is essentially the sum of the visited sub-cells  $C$ . If the tool changes to the left side of a spanning tree a detour has to be made and some sub-cells will be visited more than once. Beyond  $C$  we simply count the number of sub-cells that are visited more than once and locally charge the sub-cells of a 2D-cell for these visits.

<sup>6</sup> $K$  can be estimated by the number of sub-cells in the first layer of  $P$ .

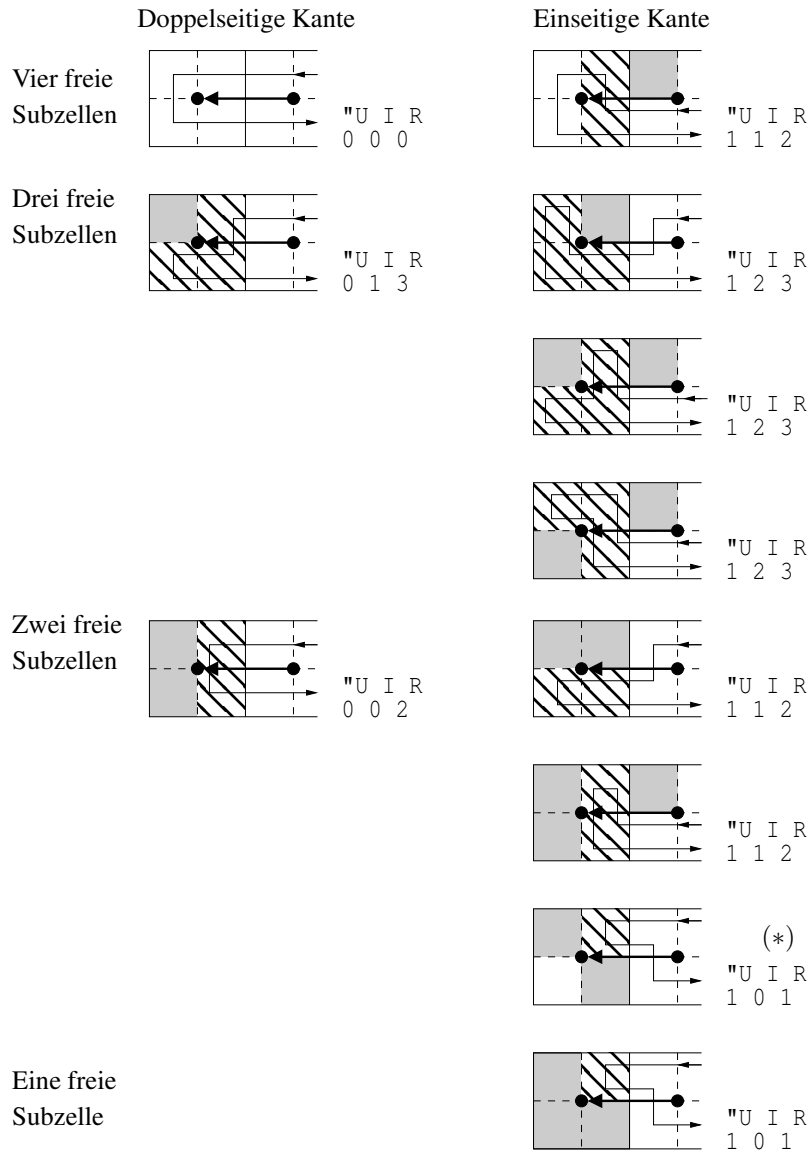


Figure 1.27: Analysis of STC, all possible cases.

We differentiate between *inner* double visits and *intra* double visits. The latter one occur during the movement inside a 2D-cell if a sub-cell is visited again. The former one occur if we leave a 2D cell  $c$  along the spanning tree to a neighbouring cell and the corresponding sub-cell was visited before. For this double visit we also charge the 2D cell  $c$ , since it was responsible for the detour.

Any 2D-cell  $c$  is visited for the first time by an incoming spanning tree edge. The inner-cell double visit will occur only if the cell  $c$  is left again along this edge. Figure 1.26 shows an example for counting inner and intro double visits. For cell 1 sub-cell  $s$  is visited twice, an intra double visit. The sub-cell above  $s$  is also visited twice, but by the movement back for 5 to 1 along the spanning tree edge. Therefore 2D-cell 5 is charged for this by an inner double visit.

The table of 2D-cells Figure 1.26 shows the number of inner and intra double visits for any 2D-cell. We charge the 2D-cells for these double visits. On the other hand, for any 2D-cell we also count the number of sub-cells that are diagonally adjacent to a boundary sub-cell. The corresponding boundary sub-cell need not lie inside the 2D-cell itself. Surprisingly, the sum of inner and intra double visits does never exceed the number sub-cells with diagonally adjacent neighbours. This is also given in the table of Figure 1.26.

For a full systematic proof we refer to Figure 1.27. Any 2D-cell  $c$  is visited by some spanning tree

edge for the first time and the inner double visits can only occur on this edge. Therefore it is sufficient to consider the  $2D$ -cell without other outgoing spanning tree edges. For any intra detours only sub-cells of the current cell are responsible. For the inner detour only the parent cell was responsible.

We distinguish between double sided and single sided edges and between the number of boundary sub-cells inside the corresponding  $2D$ -cell  $c$ . We always count inner and intra double visits and compare the sum to the number of sub-cells adjacent to boundary sub-cells.

For all reasonable cases the sum of double visits is always covered locally by the number sub-cells adjacent to boundary sub-cells. The case marked with (\*) is a bit tricky. The corresponding  $2D$  cell might also be visited by another spanning tree edge. This is not critical because there is only 1 double visit in this case for each sub-case. They can be handled separately.

### Running time and space requirement

The tool performs at most  $C + K \leq 2C$  steps. Any movement is computed locally in  $O(1)$  time. The corresponding overall information required does not exceed  $O(C)$ .  $\square$

Finally, we consider the Scan-variants of the STC-Algorithms. We would like to give a rough estimate for the efficiency in avoiding horizontal edges by  $2D$ -Scan-STC.

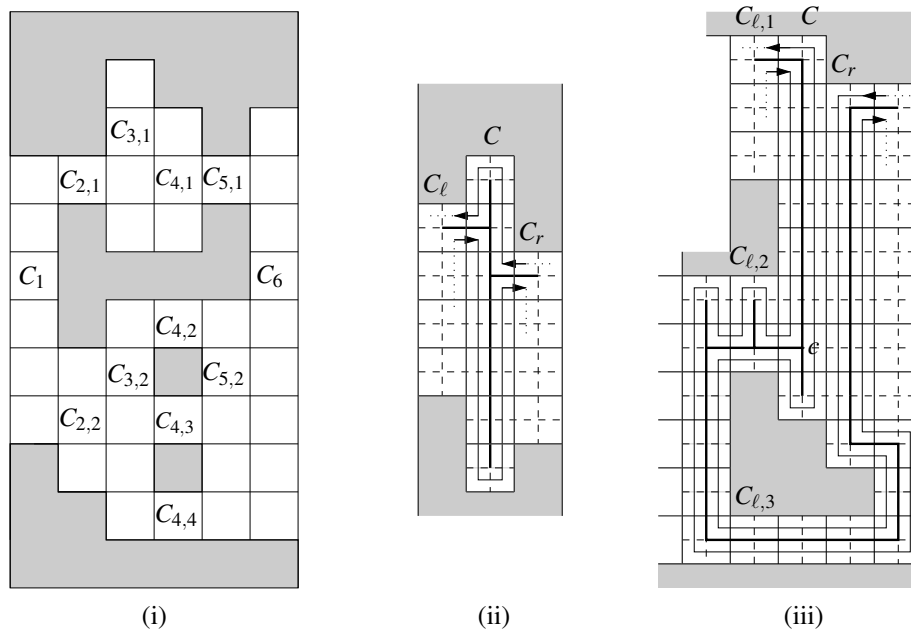


Figure 1.28: (i) Columns and the change of connectivity, (ii) Columns without changes, (iii) Difficult online situation.

We consider columns of the gridpolygon and from left to right we count the change of the connectivity from a column to its neighbour on the right. For example on Figure 1.28(i) there is a numbering of the columns and the number of different vertical components of the columns. From left to right we sum up all differences in the number of components of a column to its neighbour. In Figure 1.28(i) column  $C_1$  has one component and in column  $C_2$  this component split into two components  $C_{2,1}$  and  $C_{2,2}$ . This gives a difference of 1. The components  $C_{5,1}$  and  $C_{5,2}$  of column  $C_5$  run together in  $C_6$  to a single component. This also is a change of 1 in the difference. Of course also many parts might be involved. We count the changes of any component separately. Let  $Z$  denote the sum of all these local changes.

The number  $Z$  is a measure for the additional horizontal edges of the spanning tree of Scan-STC against an optimal number of spanning tree edges:

### Theorem 1.20 (Gabriely, Rimon, 2000)

Let  $P$  be a gridpolygon. Let  $H_{\text{Opt}}$  denote the minimal number of spanning tree edges among all  $2D$  spanning trees of  $P$ . Let  $Z$  be the above number of connectivity changes for the columns of the  $2D$ -cells.

*2D-Scan-STC constructs a spanning tree with at most*

$$H_{\text{STC}} \leq H_{\text{Opt}} + Z + 1$$

*horizontal edges.*

[GR03]

**Proof.**(Sketch)

If there is no change in a  $2D$  column, the optimal spanning tree and  $2D$ -Scan-STC will visit and leave the column only once; compare Figure 1.28(ii). The main problem is that by  $2D$ -Scan-STC a connected component of a column will be left by the spanning tree to the same side more than once. This can only happen, if there are changes in the connectivity; see Figure 1.28(iii).  $\square$

**Concluding remarks**

Arkin, Fekete and Mitchell gave some approximation results for the offline exploration of gridpolygons; see [AFM00]. Betke, Rivest und Singh considered a variant of the exploration problem. They introduced the following piecemeal-condition: The agent has to explore an environment with rectangular obstacles and has to return to the start from time to time (charging an accumulator); see [BRS94]. A strategy for this problem for general grid-environments stems from Albers, Kursawe und Schuierer [AKS02].

## 1.5 Constrained graph-exploration

We consider the problem of the exploration of an unknown graph  $G = (V, E)$  starting from some fixed vertex  $s \in V$ . This means that we would like to visit all edges and vertices of  $G$ . First, we consider unit-weights which means that any visit of an edge has cost 1. Different from the previous section we consider a constrained version of the exploration, due to the following practical models. Let  $r$  denote the radius or depth of the graph w.r.t.  $s$ . I.e.,  $r$  is the maximal length of a shortest path from  $s$  to some vertex  $v \in V$ . Let us first assume that  $r$  is known, but not the graph itself.

1. The agent is bounded by a tether of length  $\ell = (1 + \alpha)r$  (for example a cable constraint).
2. The agent has to return to the start after any  $2(1 + \alpha)r$  steps (for example an accumulator has to be recharged).
3. A large graph should be explored up to a given fixed depth  $d$  (for example for searching a close by target).

The above third variant will be applied to a searching heuristic with increasing depth, later. First, we show some simple simulation results. If an algorithm for the tether variant is known, one can also establish an accumulator strategy with some extra cost.

**Lemma 1.21** *Given a tether variant strategy with tether length  $l = (1 + \alpha)r$  and overall cost  $T$ . For any  $\beta > \alpha$  there is an accumulator-strategy with cost  $\frac{1+\beta}{\beta-\alpha}T$*

**Proof.** We design the accumulator strategy by following the tether strategy. After any  $2(\beta - \alpha)r$  steps we move back from the current vertex  $v$  to the start, recharge the agent and move back to  $v$ . Then we proceed with the next step of length  $2(\beta - \alpha)r$  of the tether strategy path. In the tether strategy for any vertex  $v$ , we are never more than  $(1 + \alpha)r$  away from the start. That is  $2(\beta - \alpha)r + 2(1 + \alpha)r = 2(1 + \beta)r$  always result in correct loops. The strategy is correct.

On the other hand, we have cost  $T$  for following the tether path and additional cost for moving back and force. We move back at most  $\frac{T}{2(\beta-\alpha)r}$  times and require  $2(1 + \alpha)r$  steps for any movement. This gives total cost:

$$T + \frac{T}{2(\beta - \alpha)r} \cdot 2(1 + \alpha)r = T \frac{\beta - \alpha + 1 + \alpha}{\beta - \alpha} = \frac{1 + \beta}{\beta - \alpha} T.$$

□

**Exercise 9** *Given an accumulator strategy  $S$  with accumulator size  $2(1 + \beta)r$  and overall cost  $T$ . For a given  $\alpha > \beta$  design an efficient tether strategy that makes use of  $S$  so that the cost of the tether strategy is  $f(\alpha, \beta) \cdot T$ . Determine  $f(\alpha, \beta)$  precisely.*

We can also consider the Offline-variant of the problem. In this case the graph is fully known. To the best of our knowledge the complexity of the Offline-variant (computing the best strategy) is not known. Since in the case that the tether is very long, the Hamiltonian-path problem appears to be part of the problem, the problem is assumed to be NP-hard.

If the optimal Offline-strategy is not known, we can design an Offline-strategy that approximates the optimal strategy. We consider the accumulation variant and assume that the accumulator has size  $4r$ .

**Lemma 1.22** *Let us assume that an accumulator of size  $4r$  is given. There is a simple Offline algorithm that explores a graph of depth  $r$  with no more than  $6|E|$  steps.*

**Proof.** We consider the DFS walk among the edges of the graph which requires  $2|E|$  steps. Now we split this overall path into pieces of size  $2r$ . Similarly to the simulation in the proof above we successively move to the start vertices of these subpaths, follow the DFS path for  $2r$  steps and return to the start after that. In total the accumulator of size  $4r$  is sufficient. Moving along the DFS path gives  $2|E|$  steps. There are no more than  $\lceil \frac{2|E|}{2r} \rceil$  sub-paths that require no more than  $\lceil \frac{|E|}{r} \rceil 2r$  steps in total. We have  $\lceil \frac{|E|}{r} \rceil 2r \leq \left(\frac{|E|}{r} + 1\right) 2r \leq 2|E| + 2r$  which shows that  $4|E| + 2r \leq 6|E|$  is sufficient.  $\square$

From now on we consider only the tether variant, for the accumulation variant similar results can be shown. A first simple idea is to take the tether length for the DFS walk into account.

Just performing DFS is not always possible. A BFS approach is always possible but results in too many exploration steps; see Figure 1.29. Therefore we apply DFS with the tether restriction as given in Algorithm 1.9. There is a backtracking step, if the tether is fully exhausted. We call this algorithm bDFS for bounded DFS.

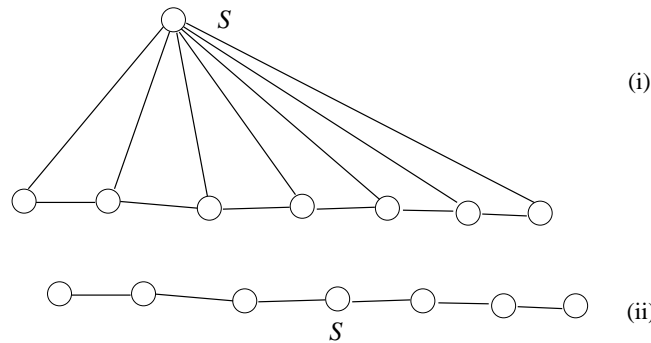


Figure 1.29: (i) A Graph with  $n$  vertices and with depth  $r = 1$ , pure DFS would require a tether of length  $n - 1$ . (ii) A graph of depth  $n$ , BFS with a tether of length  $n$  requires  $\Omega(n^2)$  steps.

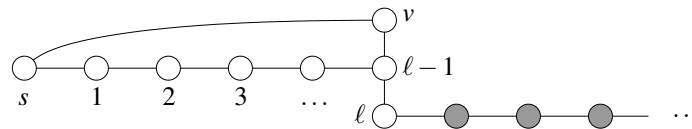


Figure 1.30: bDFS kann einige Knoten nicht erreichen.

---

#### Algorithm 1.9 boundedDFS

---

**bDFS**(  $v, \ell$  ):

**if**  $(\ell = 0) \vee$  (no adjacent non-explored edges) **then**

    RETURN

**end if**

**for all** non-explored edge  $(v, w) \in E$  **do**

    Move from  $v$  to  $w$  along  $(v, w)$ .

    Mark  $(v, w)$  as *explored*.

    bDFS( $w, \ell - 1$ ).

    Move back from  $w$  to  $v$  along  $(v, w)$ .

**end for**

---

In general bDFS is not sufficient for the full exploration of a graph. For example in Figure 1.30 we have the problem that the dark-colored vertices cannot be reached, if the algorithm first chooses the path along the vertices  $1, 2, \dots, \ell - 1$ , visits vertex  $l, v$  and  $s$  and winds back to the start  $s$ . The path from  $s$  over  $v$  is short enough but will not be considered by bDFS.



Therefore we would like to call bDFS from different sources. The aim is to achieve a constant competitive algorithm. In Algorithm 1.10 we maintain a set of (edge) disjoint trees  $\mathcal{T} = \{T_1, T_2, \dots, T_k\}$  with root vertices  $s_1, s_2, \dots, s_k$ , respectively. The trees still contain incomplete vertices where not all adjacent edges have been visited. We choose a tree  $T_i$  with the minimal distance from  $s$  to root  $s_i$  among all trees of  $\mathcal{T}$ . From this tree we prune subtrees  $T_{w_j}$  with root vertices  $w_j$ , so that  $w_j$  is a certain distance ( $\text{minDist} = \frac{\alpha r}{4}$ ) away from  $s$  and  $T_{w_j}$  has a certain minimal depth (determined over  $\text{minDepth} = \frac{\alpha r}{2}$ ). Those trees will be inserted into  $\mathcal{T}$ . The pruning forces the trees of  $\mathcal{T}$  to have a minimum size, it is still worth visiting them.

After pruning, the rest of  $T_i$  will be explored by DFS and if an incomplete vertex will be found, we start bDFS with the current remaining tether length for the exploration of *new* edges. The newly explored edges and vertices build a graph  $G'$ . If  $G'$  has incomplete vertices, we construct a spanning tree  $T'$  with a root vertex  $s'$ , where  $s'$  is the vertex in  $T'$  closest to  $s$  in the current overall explored graph  $G^*$ .  $T'$  will be inserted into  $\mathcal{T}$ . After the overall DFS (and bDFS) walk in  $T_i$  we delete all trees of  $\mathcal{T}$  that are now fully explored. Some of the trees in  $\mathcal{T}$  might have common vertices. We merge those trees and build a new spanning tree for them with a new root vertex.

A scheme of the algorithm is shown in Figure 1.31. We have done the prune step by values  $(2, 4)$ . Otherwise, we have to build very large example graphs.

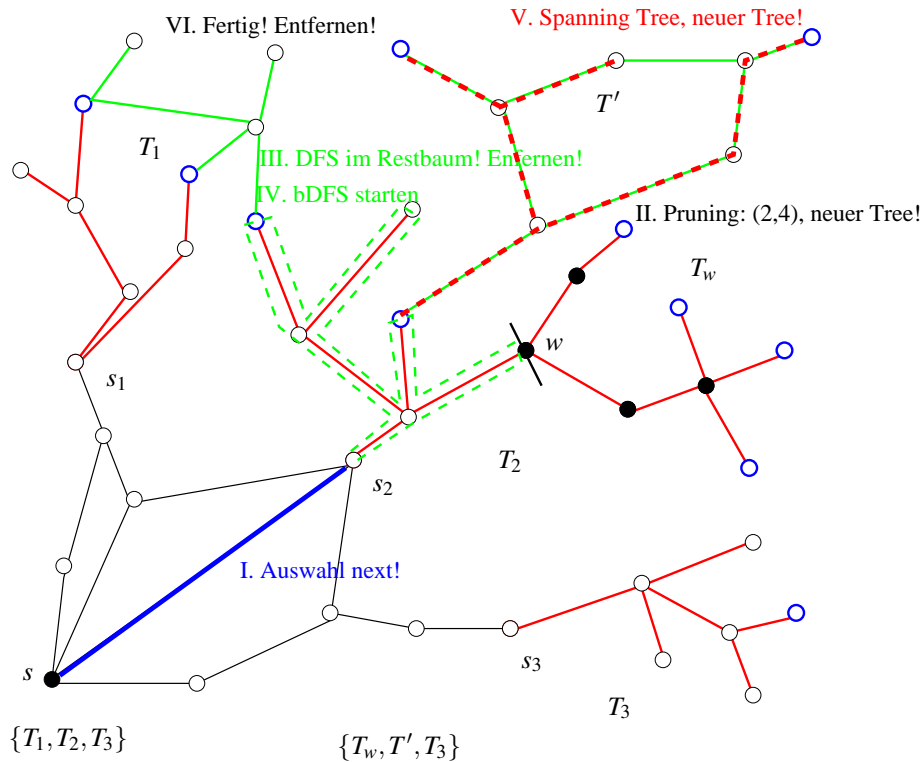


Figure 1.31: The algorithm maintains a set of disjoint trees  $\mathcal{T} = \{T_1, T_2, T_3\}$  and choose the tree  $T_2$  with minimal distance  $d_{G^*}(s, s_i)$ . After that the tree is pruned. Subtrees of distance 2 away from  $s_2$  with vertices inside that have distance at least 4 from  $s_2$  are cut-off. After that DFS starts on the rest of  $T_2$  and starts bDFS on the incomplete vertices. Here some new graphs  $G'$  will be explored and we build spanning trees  $T'$  for them. Some trees in  $\mathcal{T}$  get fully explored.  $T_w$  and  $T'$  are added to  $\mathcal{T}$ , the tree  $T_2$  is deleted.

In the following let  $d_{G'}(v, w)$  denote the distance between vertices  $v$  and  $w$  in the subgraph or tree  $G'$ .  $G^* = (V^*, E^*)$  denotes the currently known part of  $G$ .

The algorithm makes us of the following subdivision of vertices:

**non-explored** a vertex, which was never been visited before.

**incomplete** a vertex already visited before but some of the adjacent edges are still non-explored.

---

**Algorithm 1.10** CFS

---

**CFS**(  $s, r, \alpha$  ) $\mathcal{T} := \{ \{s\} \}$ .**repeat** $T_i :=$  closest subtree of  $\mathcal{T}$  to  $s$  in  $G^*$ . $s_i :=$  vertex of  $T_i$  closest to  $s$  in  $G^*$ . $(T_i, \mathcal{T}_i) := \text{prune}( T_i, s_i, \frac{\alpha r}{4}, \frac{\alpha r}{2} )$ . $\mathcal{T} := \mathcal{T} \setminus \{T_i\} \cup \mathcal{T}_i$ .**explore**(  $\mathcal{T}, T_i, s_i, (1 + \alpha)r$  ).Delete fully explored trees from  $\mathcal{T}$ .Merge the trees of  $\mathcal{T}$  with common vertices.Define a root vertex closest to  $s$  in  $G^*$ .**until**  $\mathcal{T} = \emptyset$ **prune**(  $T, v, \text{minDist}, \text{minDepth}$  ) $v :=$  Wurzel von  $T$ . $\mathcal{T}_i := \emptyset$ .**for all**  $w \in T$  with  $d_T(v, w) = \text{minDist}$  **do** $T_w :=$  subtree of  $T$  with root  $w$ .**if** maximale Distanz between  $v$  and a vertex in  $T_w > \text{minDepth}$  **then**// Cut-Off  $T_w$  from  $T$  ab: $T := T \setminus T_w$ . $\mathcal{T}_i := \mathcal{T}_i \cup \{T_w\}$ .**end if****end for**RETURN (  $T, \mathcal{T}_i$  )**explore**(  $\mathcal{T}, T, s_i, \ell$  )Move from  $s$  to  $s_i$  along shortest path in  $G^*$ .Explore  $T$  by DFS. If incomplete vertex occurs, do: $\ell' :=$  remaining tether length.bDFS(  $v, \ell'$  ). $E' :=$  set of newly explored edges. $V' :=$  set of vertices of  $E'$ .Calculate spanning tree  $T'$  for  $G' = (V', E')$ .Define root vertex of  $T'$  closest to  $s$  in  $G^*$  $\mathcal{T} := \mathcal{T} \cup \{T'\}$ .Move back from  $s_i$  to  $s$ .

---

**explored** a vertex, that was visited and all adjacent edges have been explored.

Additionally, for the bDFS walk we mark the edges as 'non-explored' or 'explored'.

**Lemma 1.23** *The following properties hold during the execution of the CFS-Algorithm:*

- (i) Any incomplete vertex belongs to a tree in  $\mathcal{T}$ .
- (ii) Until  $G^* \neq G$ , there is always an incomplete vertex  $v \in V^*$  so that  $d_{G^*}(s, v) \leq r$ .
- (iii) For any chosen root vertex  $s_i$ :  $d_{G^*}(s, s_i) \leq r$ .
- (iv) After pruning  $T_i$  is fully explored by DFS. All trees  $T \in \mathcal{T}$  have size  $|T| \geq \frac{\alpha r}{4}$ .
- (v) All trees  $T \in \mathcal{T}$  are disjoint (w.r.t. edges).

**Proof.**

- (i) Follows directly from the construction of the trees by bDFS and Pruning. No incomplete vertex is missing.
- (ii) Assume that for all  $v \in V^*$  we have  $d_{G^*}(s, v) > r$  and let  $v$  be an incomplete vertex of  $V^*$ . In  $G$  there is a shortest path  $P(s, v)$  from  $s$  to  $v$  with length  $\leq r$ . Along  $P(s, v)$  there is a first vertex  $w$  that does not belong to  $G^*$ . Thus its predecessor  $w'$  along  $P(s, v)$  belongs to  $V^*$  and is incomplete. We have  $d_{G^*}(s, w') \leq r$ .
- (iii) Follows from (ii), the root of a corresponding tree  $T$  is always the vertex of  $T$  closest to  $s$ .
- (iv) We show the property by successively considering the upcoming trees. Or by induction on the number of pruning steps. In the beginning the algorithm starts with bDFS at the root  $s$ . Either, the graph will be fully explored and we are done, or bDFS have exhausted the tether of length  $(1 + \alpha)r$  and have visited more than  $(1 + \alpha)r$  edges. The single spanning tree  $T$  has size  $|T| \geq (1 + \alpha)r > \frac{\alpha r}{4}$ . Let us assume that the condition holds for the trees inside  $\mathcal{T}$  and the next pruning step happens. Now by the next iteration we are choosing tree  $T_i$  with root  $s_i$  closest to  $s$  among all trees in  $\mathcal{T}$ . After that we prune  $T_i$ . The rest of  $T_i$  has still size  $|T_i| \geq \frac{\alpha r}{4}$  since we cut off subtrees  $T_w$  with distance  $\geq \frac{\alpha r}{2}$  away from  $s_i$ . For a corresponding subtree  $T_w$  we conclude  $|T_w| \geq \frac{\alpha r}{2} - \frac{\alpha r}{4} = \frac{\alpha r}{4}$  since there is a vertex inside  $T_w$  that is at least distance  $\frac{\alpha r}{2}$  away from  $s$ . Now consider the remaining DFS/bDFS combination on (the rest of)  $T_i$ . The distance from  $s$  to  $s_i$  is at most  $r$ . Any incomplete vertex in the current  $T_i$  has at most distance  $\frac{\alpha r}{2}$  from  $s_i$  otherwise this vertex would be part of a tree  $T_w$  that has to be considered in the pruning step. This means that at any incomplete vertex there is a rest tether of length  $\frac{\alpha r}{2}$  which can be used for the bDFS part. If the exploration results in another spanning tree  $T'$  with incomplete vertices, this tree has size at least  $\frac{\alpha r}{2}$ . Finally fully explored trees are deleted from  $\mathcal{T}$  which is not critical. Additionally, some other trees might be merged and still have incomplete vertices. These trees only grow. □

Finally, we show:

**Theorem 1.24** (Duncan, Kobourov, Kumar, 2001/2006)

*The CFS-Algorithm for the constrained graph-exploration of an unknown graph with known depth is  $(4 + \frac{8}{\alpha})$ -competitive. [DKK06, DKK01]*

**Proof.** We split the cost for any appearing subtree  $T_R$ . Let  $K_1(T_R)$  denote the cost for moving from  $s$  to  $s_i$  in  $G^*$ . Let  $K_2(T_R)$  denote the cost of DFS for  $T_R$  and let  $K_3(T_R)$  denote the cost for the bDFS exploration done for the incomplete vertices starting at  $T_R$ . The trees are edge disjoint.

The total cost is a sum of the cost for any  $T_R$ . We have

$$\sum_{T_R} K_3(T_R) \leq 2 \cdot |E|, \text{ since bDFS only visits non-explored edges (twice).}$$

$$\sum_{T_R} K_2(T_R) = \sum_{T_R} 2 \cdot |T_R| \leq 2 \cdot |E|, \text{ the cost for all DFS walks.}$$

For  $K_1(T_R)$  we have  $K_1(T_R) = 2 \cdot d_{G^*}(s, s_i) \leq 2r$ . The complexity of any  $T_R$  is at least  $\frac{\alpha r}{4}$  which gives  $|T_R| \geq \frac{\alpha r}{4}$  for the number of edges. We conclude  $r \leq \frac{4|T_R|}{\alpha}$  and

$$\sum_{T_R} K_1(T_R) \leq \sum_{T_R} 2r \leq \frac{8}{\alpha} \sum_{T_R} |T_R| \leq \frac{8}{\alpha} |E|$$

Altogether, the algorithm makes  $(4 + \frac{8}{\alpha})|E|$  step whereas any optimal algorithm visits at least any edge once.  $\square$

In general we assume that  $\alpha$  is a small constant with  $0 < \alpha < 1$ . The above proof works for any  $\alpha > 0$ . The cost of the algorithm for known depth  $r$  are within  $O(|E|/\alpha)$ . More precisely we can show that actually  $O(|E| + |V|/\alpha)$  steps are made. For this we have a closer look at the cost. bDFS work on the edges only. The DFS walk work on trees where the number of vertices is the same as the number of edges. Some of these vertices appear in two trees, so by a factor of 2 we are on the save side. The movements from  $s$  to  $s_i$  are analysed over the size of spanning trees, where vertices and edges are also the same.

The cost  $K_1(T_R)$  and  $K_2(T_R)$  sum up to  $(2 + \frac{8}{\alpha})2|V|$ .

Altogether there is an  $\Theta(|E| + |V|/\alpha)$  algorithm for the exploration of arbitrary graphs.

**Corollary 1.25** *The CFS-Algorithm for the constrained graph-exploration of an unknown graph with known depth has optimal exploration cost  $\Theta(|E| + |V|/\alpha)$ .*

Now we have some possibilities for extensions. First, we assume that the depth of the graph is unknown in the beginning. Next we would like to consider weighted edges.

### 1.5.1 Restricted graph-exploration with unknown depth

Let us now assume that the radius, say  $R$ , of the unknown graph  $G$  is not known. From a practical point of view, spending some cable is costly and we would like to extend the tether only if it is necessary. A first simple idea is that we guess the depth, say  $r$ , and successively double its length until the algorithm finally explores the whole graph. Obviously, the repeated application of the CFS-algorithm runs in  $O(\log r|E|)$  step. As shown above we can also refine the analysis of this approach. For any bDFS step we make use of the already visited edges and directly *jump* to incomplete vertices (now with larger tether length). Therefore the bDFS steps are still subsumed by  $2|E|$  steps. But we still have to take the movements to the roots of the trees into account as well as the DFS movements on the new subtrees. Therefore we have the following result.

**Corollary 1.26** *Applying the CFS-Algorithmus by successively doubling the current depth  $r$  gives an algorithm that explores an unknown graph  $G$  with unknown depth  $R$  with  $\Theta(|E| + (\log R)|V|/\alpha)$  steps.*

We will now show that we can get rid of the log-factor by successively adjusting  $r$  appropriately. We only exchange two calls in the main procedure. In principle, instead of the known value  $r$  we successively make use of  $r := d_{G^*}(s, s_i)$ , which is the smallest distance from  $s$  to one of the roots of the trees in  $\mathcal{T}$ .

More precisely, we exchange  $\text{prune}(T_i, s_i, \frac{\alpha r}{4}, \frac{\alpha r}{2})$  by  $\text{prune}(T_i, s_i, \frac{\alpha d_{G^*}(s, s_i)}{4}, \frac{9\alpha d_{G^*}(s, s_i)}{16})$  and  $\text{explore}(\mathcal{T}, T_i, s_i, (1 + \alpha)r)$  by  $\text{explore}(\mathcal{T}, T_i, s_i, (1 + \alpha)d_{G^*}(s, s_i))$ . This means that the pruning-step is done with the values  $\frac{\alpha d_{G^*}(s, s_i)}{4}$  and  $\frac{9\alpha d_{G^*}(s, s_i)}{16}$  and the explore-step is done with tether length  $(1 + \alpha)d_{G^*}(s, s_i)$ .

In the beginning we have  $d_{G^*}(s, s_i) = 0$ , therefore we make use of some fixed constant  $c$  in the beginning and use  $r := \max(d_{G^*}(s, s_i), c)$ . Let  $d_{G^*}(s, T)$  denote the shortest distance from  $s$  to some vertex in  $T$  inside  $G^*$ .

**Lemma 1.27** *For the CFS-Algorithmus with unknown depth  $R$  we have the following properties:*

- (i) *Any incomplete vertex belongs to a tree in  $\mathcal{T}$ .*
- (ii) *There is always an incomplete vertex  $v \in V^*$  with  $d_{G^*}(s, v) \leq r$ , until  $G^* \neq G$ .*
- (iii) *For the closest root  $s_i$  we have:  $d_{G^*}(s, s_i) \leq r$ .*
- (iv) *For all trees  $T \in \mathcal{T}$  we have  $|T| \geq \frac{\max(d_{G^*}(s, T), c)\alpha}{4}$ . After pruning the remaining tree will be fully explored by DFS.*
- (v) *All trees ever considered in  $\mathcal{T}$  are (edge) disjoint.*

**Proof.** For the proof of (i),(ii),(iii) and (v) we apply the same arguments as in the proof of Lemma 1.23. It remains to show that (iv) holds. The main difference is that the size of a tree  $T$  is directly correlated to the distance from  $s$  to  $T$ , this is different from the previous argumentation.

Let us first show that the remaining tree  $T_i$  (after pruning) will be fully explored by DFS. For any vertex  $v$  in  $T_i$  we have  $d_{T_i}(s_i, v) \leq \frac{9d_{G^*}(s, s_i)\alpha}{16}$ , otherwise  $v$  has been cut of by pruning. Thus we have

$$(1 + \alpha)d_{G^*}(s, s_i) - d_{G^*}(s, s_i) - d_{T_i}(s_i, v) \geq \frac{7d_{G^*}(s, s_i)\alpha}{16},$$

which shows that the tether is long enough  $T_i$  will be fully explored by DFS.

By induction over the number of pruning steps we will finally show:  $\forall T \in \mathcal{T} : |T| \geq \frac{\max(d_{G^*}(s, T), c)\alpha}{4}$ .

In the beginning we apply bDFS from the start with tether length  $c$ . Either we explore the whole graph or we have  $|T| \geq (1 + \alpha)c > \frac{\alpha c}{4}$  for the resulting spanning tree  $T$ . For simplicity we assume  $d_{G^*}(s, T_i) > c$  from now on.

We would like to show that for any tree  $T_w$ , resulting from the pruning of some  $T_i$ , we have  $|T_w| \geq \frac{d_{G^*}(s, T_w)\alpha}{4}$ . Also the remaining tree  $T_i$  has this property.

For the remaining tree  $T_i$  (after pruning), we conclude  $d_{G^*}(s, T_i) = d_{G^*}(s, s_i)$  and pruning guarantees  $|T| \geq \frac{d_{G^*}(s, T)\alpha}{4}$ . For a tree  $T_w$  pruned from  $T_i$  we have:  $|T_w| \geq \frac{9d_{G^*}(s, s_i)\alpha}{16} - \frac{d_{G^*}(s, s_i)\alpha}{4} = 5\frac{d_{G^*}(s, s_i)\alpha}{16}$  by the pruning values. Additionally, we have  $d_{G^*}(s, T_w) \leq d_{G^*}(s, s_i) + d_{G^*}(s_i, w) = (1 + \frac{\alpha}{4})d_{G^*}(s, s_i)$ , since the root  $w$  of  $T_w$  is exactly  $\frac{\alpha d_{G^*}(s, s_i)}{4}$  steps away from  $s$ . Für  $0 < \alpha < 1$  we conclude:  $d_{G^*}(s, T_w) < \frac{5d_{G^*}(s, s_i)}{4}$  and together with the above inequality we have  $|T_w| > \frac{d_{G^*}(s, T_w)\alpha}{4}$ .

Finally, we have to analyse the emerging spanning trees  $T_v$ , which will be constructed from the bDFS steps starting during the DFS walk in  $T_i$ . Such a tree  $T_v$  starts at some incomplete vertex  $v$  in  $T_i$ . We have  $d_{G^*}(s_i, v) \leq \frac{9\alpha d_{G^*}(s, s_i)}{16}$ , otherwise  $v$  would have been pruned and could not be a leaf of the rest of  $T_i$  any more. Thus we have  $d_{G^*}(s, T_v) \leq d_{G^*}(s, s_i) + d_{G^*}(s_i, v) < \frac{25d_{G^*}(s, s_i)}{16}$  or  $d_{G^*}(s, s_i) > \frac{16d_{G^*}(s, T_v)}{25}$ . If  $T_v$  is fully explored, we are done, since the tree will be deleted. Assume that  $T_v$  still has incomplete vertices. As mentioned above we have  $d_T(s_i, v) \leq \frac{9\alpha d_{G^*}(s, s_i)}{16}$ . Starting from  $v$  there was a remaining tether length of  $\frac{7\alpha d_{G^*}(s, s_i)}{16}$  for the construction of the incomplete  $T_v$ , which gives  $|T_v| \geq \frac{7\alpha d_{G^*}(s, s_i)}{16}$ . Application of  $d_{G^*}(s, s_i) > \frac{16d_{G^*}(s, T_v)}{25}$  gives  $|T_v| > \frac{7\alpha d_{G^*}(s, T_v)}{25} > \frac{d_{G^*}(s, T_v)\alpha}{4}$ . Either we have explored everything behind  $v$  or the spanning tree  $T_v$  has size  $|T_v| > \frac{d_{G^*}(s, T_v)\alpha}{4}$ .

We have considered any emerging  $T \in \mathcal{T}$ ! □

**Theorem 1.28** (Duncan, Kobourov, Kumar, 2001/2006)

*Applying the CFS-Algorithm with the adjustments above results in a correct restricted graph-exploration of an unknown graph with unknown depth. The algorithm is  $(4 + \frac{8}{\alpha})$ -competitive. [DKK06, DKK01]*

**Proof.** We apply the same analysis as in the proof of Theorem 1.24. For the analysis of the movements from  $s$  to the roots of the trees we make use of the correlation  $|T_R| > \frac{d_{G^*}(s, T_R)\alpha}{4}$ . □

For the number of steps we can also refine the analysis, analogously.

**Corollary 1.29** *The above CFS-Algorithm for the restricted exploration of an unknown graph with unknown depth requires  $\Theta(|E| + |V|/\alpha)$  exploration steps, which is optimal.*

Finally, we would like to argue that the usage of a look-ahead of  $\alpha r$  is necessary for attaining linear optimal exploration cost (i.e., in comparison to  $|E|$  and  $|V|$ ). This can be shown for the accumulator variant as follows. First, it is clear that an accumulator of size  $2r$  is not sufficient for exploring all edges. The graph in Figure 1.32 has depth 6, but exploring all edges requires an accumulator of size 13.

This means that an accumulator size  $2r + 1$  is necessary. We show that an accumulator of size  $2r + d$  for constant  $d$  is not sufficient in the sense of performing no more than  $C \cdot |E|$  exploration steps.

**Lemma 1.30** *For the accumulator variant with accumulator size  $2r + d$  for constant  $d$ , there are examples do that any algorithm attains at least  $\Omega(|E|^{\frac{3}{2}})$  exploration steps.*

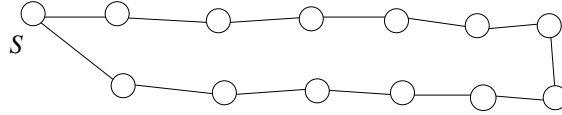


Figure 1.32: A graph of depth  $r = 6$  that cannot be explored by an accumulator of size  $2r$ .

**Proof.** We consider the following example as given in Figure 1.33. Starting from  $s$  there is a path of length  $\frac{n}{2}$  that visits a clique of size  $\frac{n}{2} + 1$ . Moving forth and back along the path requires  $n$  steps, the depth of the graph is  $\frac{n}{2} + 1$ . Exploration with accumulator size  $n + 2 + d$  means that we have to visit the clique  $\Omega\left(\frac{n^2}{d}\right)$  times since the clique has  $\Omega(n^2)$  edges. This gives  $\Omega\left(\frac{n^2}{d} \cdot n\right) = \Omega(n^3)$  exploration steps. The statement follows from  $|E| \in \Theta(n^2)$ .  $\square$

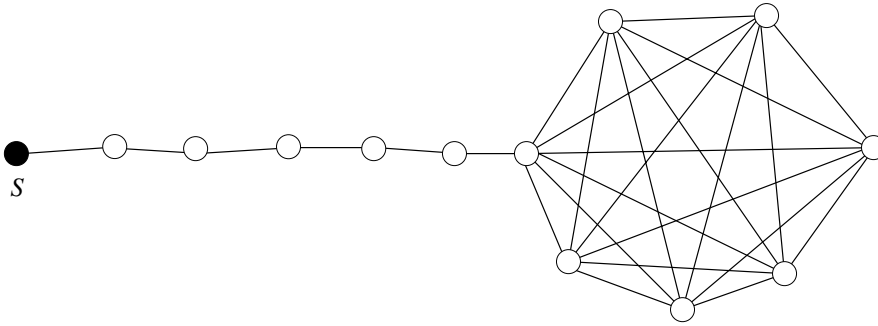


Figure 1.33: A graph with  $n + 1 = 13$  vertices. A path of length  $\frac{n}{2}$  visits a clique of size  $\frac{n}{2} + 1$ . Any accumulator strategy with accumulator size  $n + 2 + d$  requires  $\Omega(n^3)$  steps.

With a similar argument we conclude that an sub-linear extension of the accumulator, i.e., size  $2r + o(r)$ , is not sufficient for attaining a linear cost strategy. Let us briefly repeat the small-o notation. For real valued functions or series  $f$  and  $g$  we define  $f \in o(g)$ , if and only if  $\lim_{r \rightarrow \infty} \frac{f(r)}{g(r)} \rightarrow 0$  holds. Therefore we conclude  $r \in o(r^2)$ ,  $c \in o(r)$  for any constant  $c$  and also  $\frac{1}{r} \in o(1)$ . By the above arguments and example we can show that  $\Omega\left(\frac{n^3}{f(n)}\right)$  exploration steps are necessary for an accumulator of size  $n + 2 + f(n)$ . For  $f(n) = n^{1-\epsilon}$  (this means  $f \in o(n)$ ) we have to perform  $\Omega(|E|^{1+\epsilon})$  exploration steps.

Note, that for the tether variant up to our knowledge there is no such statement that a tether of length  $r + o(r)$  is necessary for attaining  $O(|E|)$  exploration cost.

We have shown that we can explore any graph (online and offline) with at most  $\Theta(|V| + |E|)$  exploration steps. These are the pure cost for the motion of the agent. In the literature this is also denoted as the *mechanical cost*; see also [DJMW91]. Besides, there are also some *computational cost*, for the planning and preparation of the strategy.

For example the computational cost of the CFS-Algorithm have to be analysed for the following tasks:

- Build the spanning trees
- Update the shortest paths to the trees of  $\mathcal{T}$
- Merge the trees
- Detect fully explored trees
- Prune a tree
- Maintain the list  $\mathcal{T}$
- Apply DFS/bDFS

For unit-length edges some of the above tasks can be done very efficiently. The overall approach can be easily extended to weighted graphs (positive edge weights).

**Exercise 10** Analyse the computational cost for the CFS-Algorithm in  $O$ -notation for  $|E|$  and/or  $|V|$ .

**Exercise 11** Show that the CFS-Algorithm approach also works for graphs with positive edges weights. How do we have to adjust the CFS-Algorithm?

### 1.5.2 Mapping of an unknown graph

Finally, in this section we would like to show the influence of different capabilities of the agent. Up to now we assumed that an already visited vertex or edge will be recognized at the next visit. This means that we have marked any visited edge and vertex.

Let us now assume that the agent cannot mark parts of the environment. We do not have any *landmarks*. We still assume that we have enough storage for constructing the sub-graph detected so far.

The following model is taken from Dudek et al.; see [DJMW91]. The agent has no orientation and no compass. At any vertex the outgoing edges are presented in the same order. This order need not represent a planar embedding. If the agent visits the vertex from different incoming edges, the order will be consistent. This means that there is a fixed cyclic order, the relative presentation of the order stems from the edge where the agent currently comes from. Figure ?? shows an example of a relative order. By this order, the agent knows where he was coming from and can also return to this vertex. Since the storage is not limited, it is possible to remember a return path. Let us for example assume that the agent visits vertex  $v_2$  by edge  $e_1$  and then visits the second edge  $e_3$  in ccw-order from  $e_1$ . If the agent moves back along  $e_3$  to  $v_2$ , it already knows that it was recently coming from the first edge in ccw order, which is  $e_1$ . The agent can make use of this return path. If the agent visits a vertex in a forward step, it has no idea which of the vertices the visited vertex actually is.

Is it possible to build a map of the graph and to locate oneself inside the graph? The offline input is a triple  $G = (V, E, S)$ , where by  $S$  for any vertex the cyclic local order of the edges is given.

First, it is easy to see that without further capabilities, one can not fully detect a given graph. Figure 1.34 shows two different regular graphs of fixed degree 3. For an agent the information on any vertex is exactly the same. It is not possible to distinguish between the two variants. At least one marker is necessary.

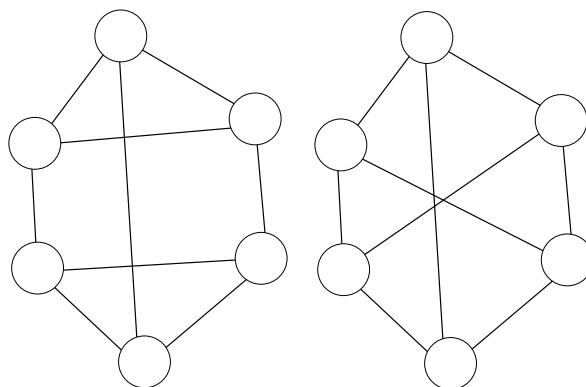


Figure 1.34: Two different regular graphs of degree 3, an agent cannot distinguish them without a marker.

**Corollary 1.31** Let  $G = (E, V, S)$  be a graph with local cyclic edge order- Without a marker an online agent cannot build a correct map of the graph.

**Exercise 12** Give a formal argument that the graphs in Figure 1.34 are different. Which class of graphs can be correctly detected by an online agent without a marker?

A single marker (or pebble) is sufficient as shown by Dudek et al. [DJMW91]. We describe the corresponding *Marker-Algorithm*. The algorithm maintains the current known graph  $G^*$  and a list  $L$  of non-determined (seen but not correctly detected) edges. In the beginning the starting vertex is known and its outgoing edges belong to  $L$ . They are given in the cyclic order.

In the main step, the algorithm choose an edge  $e$  of  $L$  starting at a detected vertex  $b$  and moves to a vertex  $u$  along the edges  $e = (b, u)$ . Now the agent sets the marker on  $u$ , moves back to  $b$  along  $e$  and searches for the pebble in  $G^*$ .

**Case 1:** The pebble was not found in  $G^*$ . In this case we add the edge  $e = (b, u)$  to  $G^*$  w.r.t. the cyclic order. All outgoing edges of  $u$  different from  $e$  will be inserted into the list  $L$  of non-determined edges.

**Case 2:** The marker has been found at some vertex  $v \in G^*$ . If there is more than one non-determined outgoing edges at  $v = u$ , we cannot precisely detect  $e$ . Therefore we take the marker, move back to  $b$ , place the pebble there, move back to  $v$  again and successively check the non-determined edges. Finally, we will detect the edge  $e$  and add it to  $G^*$  by the local order.

The above algorithm is simple and correct. By construction in any step an additional edge will be correctly detected. The number of exploration steps is restricted by  $O(|E| \times |V|)$  and the same holds for the computational cost. We assume that the graph is not a multigraph and has no loop edges  $(v, v)$ . Besides, we assumed that any edge has unit-length.

**Theorem 1.32** (Dudek, Jenkin, Milios, Wilkes, 1991)

Let  $G = (E, V, S)$  be a graph with given cyclic local order of the edges. By the use of one marker it is possible to fully detect the structure of the graph by online navigation with  $O(|E| \times |V|)$  exploration steps and also overall  $O(|E| \times |V|)$  computational cost.

**Proof.**

Let  $G^* = (V^*, E^*, S^*)$  be the current graph during the execution of the Marker-Algorithm. Setting the marker has cost  $O(1)$ , searching for the marker in  $G^*$  can be done by DFS by  $O(|V^*|)$  steps. Moving back and force along a path can be done in  $O(|V^*|)$  steps as well. The traversal cost are considered for any edge, which gives  $O(|E| \times |V|)$  steps in total.

For unit-edge length the computational cost are precisely the same for any edges we have to compute the shortest paths between two vertices. The effort is bounded by  $O(|V^*|)$ . This gives  $O(|E| \times |V|)$ .  $\square$

**Exercise 13** Explain why the cyclic order of the edges is necessary for the above Marker-Algorithm. Where is it used during the execution of the algorithm?

**Exercise 14** Analyse the mechanical and the computational cost of the marker algorithm for graphs with positive edge weights.



# Bibliography

- [AFM00] E. M. Arkin, S. P. Fekete, and J. S. B. Mitchell. Approximation algorithms for lawn mowing and milling. *Comput. Geom. Theory Appl.*, 17:25–50, 2000.
- [AKS02] Susanne Albers, Klaus Kursawe, and Sven Schuierer. Exploring unknown environments with obstacles. *Algorithmica*, 32:123–143, 2002.
- [BRS94] Margrit Betke, Ronald L. Rivest, and Mona Singh. Piecemeal learning of an unknown environment. Technical Report A.I. Memo No. 1474, Massachusetts Institute of Technology, March 1994.
- [DJMW91] G. Dudek, M. Jenkin, E. Milios, and D. Wilkes. Robotic exploration as graph construction. *Transactions on Robotics and Automation*, 7:859–865, 1991.
- [DKK01] Christian A. Duncan, Stephen G. Kobourov, and V. S. Anil Kumar. Optimal constrained graph exploration. In *Proc. 12th ACM-SIAM Symp. Discr. Algo.*, pages 307–314, 2001.
- [DKK06] Christian A. Duncan, Stephen G. Kobourov, and V. S. Anil Kumar. Optimal constrained graph exploration. *ACM Trans. Algor.*, 2:380–402, 2006.
- [GR03] Yoav Gabriely and Elon Rimon. Competitive on-line coverage of grid environments by a mobile robot. *Comput. Geom. Theory Appl.*, 24:197–224, 2003.
- [IKKL00a] Christian Icking, Thomas Kamphans, Rolf Klein, and Elmar Langetepe. Exploring an unknown cellular environment. In *Abstracts 16th European Workshop Comput. Geom.*, pages 140–143. Ben-Gurion University of the Negev, 2000.
- [IKKL00b] Christian Icking, Thomas Kamphans, Rolf Klein, and Elmar Langetepe. Exploring an unknown cellular environment. Unpublished Manuscript, FernUniversität Hagen, 2000.
- [IKKL05] Christian Icking, Tom Kamphans, Rolf Klein, and Elmar Langetepe. Exploring simple grid polygons. In *11th Internat. Comput. Combin. Conf.*, volume 3595 of *Lecture Notes Comput. Sci.*, pages 524–533. Springer, 2005.
- [IPS82] A. Itai, C. H. Papadimitriou, and J. L. Szwarcfiter. Hamilton paths in grid graphs. *SIAM J. Comput.*, 11:676–686, 1982.
- [Lee61] C. Y. Lee. An algorithm for path connections and its application. *IRE Trans. on Electronic Computers*, EC-10:346–365, 1961.
- [Sha52] Claude E. Shannon. Presentation of a maze solving machine. In H. von Foerster, M. Mead, and H. L. Teuber, editors, *Cybernetics: Circular, Causal and Feedback Mechanisms in Biological and Social Systems, Transactions Eighth Conference, 1951*, pages 169–181, New York, 1952. Josiah Macy Jr. Foundation. Reprint in [Sha93].
- [Sha93] Claude E. Shannon. Presentation of a maze solving machine. In Neil J. A. Sloane and Aaron D. Wyner, editors, *Claude Shannon: Collected Papers*, volume PC-03319. IEEE Press, 1993.

- [Sut69] Ivan E. Sutherland. A method for solving arbitrary wall mazes by computer. *IEEE Trans. on Computers*, 18(12):1092–1097, 1969.

# Index

$\dot{\cup}$ .....	<i>see</i> disjoint union	<b>G</b>	
1-Layer .....	<b>14</b>	<i>Gabriely</i> .....	27, 29
1-Offset .....	<b>14</b>	grid-environment .....	8
2-Layer .....	<b>14</b>	gridpolygon .....	<b>8</b> , 30
2-Offset .....	<b>14</b>	<b>I</b>	
lower bound .....	5	<i>Icking</i> .....	5, 18, 21
<b>A</b>		<i>Itai</i> .....	8
accumulator strategy .....	31	<b>J</b>	
adjacent .....	<b>8</b>	Java-Applet .....	18
<i>Albers</i> .....	30	<i>Jenkin</i> .....	40
approximation .....	30	<b>K</b>	
<i>Arkin</i> .....	30	<i>Kamphans</i> .....	5, 18, 21
<b>B</b>		<i>Klein</i> .....	5, 18, 21
Backtrace .....	19	<i>Kobourov</i> .....	35, 37
<i>Betke</i> .....	30	<i>Kumar</i> .....	35, 37
<b>C</b>		<i>Kursawe</i> .....	30
cell .....	<b>8</b>	<b>L</b>	
columns .....	29	<i>Langetepe</i> .....	5, 18, 21
competitive .....	35, 37	Layer .....	15
constrained .....	31	layer .....	27
Constraint graph-exploration .....	31	<i>Lee</i> .....	19
<b>D</b>		Left-Hand-Rule .....	10–13
DFS .....	8, 11	Lower Bound .....	9
diagonally adjacent .....	<b>8</b> , 27	lower bound .....	8
<i>Dijkstra</i> .....	19	<b>M</b>	
disjoint union .....	<b>15</b>	<i>Milios</i> .....	40
<i>Dudek</i> .....	40	<i>Mitchell</i> .....	30
<i>Duncan</i> .....	35, 37	<b>N</b>	
<b>F</b>		narrow passages .....	20
<i>Fekete</i> .....	30	NP-hard .....	8
		<b>O</b>	
		Offline-Strategy .....	<b>5</b>
		Online-Strategy .....	<b>5</b>
		Online-Strategy .....	8

**P**


---

<i>Papadimitriou</i> .....	8
partially occupied cells .....	<b>23</b>
path .....	<b>8</b>
piecemeal-condition .....	30

**Q**


---

Queue .....	19
-------------	----

**R**


---

<i>Rimon</i> .....	27, 29
<i>Rivest</i> .....	30

**S**


---

<i>Schuieler</i> .....	30
<i>Shannon</i> .....	3
<i>Singh</i> .....	30
<i>Sleator</i> .....	5
SmartDFS .....	13, 14
spanning tree .....	23
Spanning-Tree-Covering .....	23
split-cell .....	<b>14</b>
sub-cells .....	<b>23</b>
<i>Sutherland</i> .....	3
<i>Szwarcfiter</i> .....	8

**T**


---

<i>Tarjan</i> .....	5
tether strategy .....	31
tool .....	23
touch sensor .....	8

**W**


---

Wave propagation .....	19
<i>Wilkes</i> .....	40