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## Animating a Sudoku solver

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## **Abstract**

**TODO:** Sudoku is a popular puzzle played all over the world. It consists of filling in a  $9 \times 9$  grid such that every row, column and  $3 \times 3$  sub-grids have different digits from 1 to 9. Solving the puzzle will make use of the all-different algorithm from Constraint Programming for which an implementation will be provided. Finally, the program will animate all the steps done by the algorithm.

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# Chapter 1

## Introduction

The Sudoku puzzle is a well known combinatorial problem where the goal is to fill in a partially completed  $9 \times 9$  grid in such a way that a single digit cannot appear twice in the same row, column or  $3 \times 3$  box. The problem is NP-complete and will be used as an example to introduce the concepts of the all-different algorithm used in Constraint Programming to solve many real-world combinatorial problems. Although the original  $9 \times 9$  Sudoku puzzle can be solved in a fraction of a second using a variety of algorithms, the power and effectiveness of the algorithm introduced in this paper can be seen when trying to solve a  $36 \times 36$  grid, where a solution is found in a matter of seconds, rather than days using a conventional approach. Students new to Constraint Programming will be shown a new way of approaching and solving real-world problems.

### 1.1 Aims

The aim of this project is to visualise how Constraint Programming techniques can be used to solve Sudoku puzzles. A Java application is developed to act as a teaching aid to be used by the lecturer to show students how the all-different algorithm works towards a solution. Sudoku puzzles represent an excellent choice to introduce the concept of the all-different constraint in a entertaining and engaging experience with the students. Students will be shown how to model the Sudoku puzzle as a Constraint Satisfaction Problem and get to the solution without resorting to guessing or bruteforcing. The constraint programming technique that will be used to solve the Sudoku puzzle is the same algorithm used worldwide to solve planning, scheduling, timetabling and supply chain management problems. The step by step visual representation of how the algorithm progresses towards a solution will help students better understand the new algorithm.

### 1.2 Background

The all-different algorithm introduced in 1994 by Jean-Charles Regin represents to this day a great achievement in Constraint Programming and other fields.

### 1.3 Motivation

The motivation of this project is to provide a tool that makes it easier for a lecturer to introduce the all-different algorithm to its students by providing a step by step view of how the algorithm progresses. Students will find

it easier to remember and understand the algorithm through the use of the visualisations. The tool based on the Sudoku puzzle is a great way to visualize an important technique used to solve Constraint Satisfaction Problems and will provide an engaging learning experience to the students.

## 1.4 Report Content

The rest of the report will provide information on the algorithms used and their implementation in the project.

- **Chapter 2** covers details about the Sudoku puzzle and Constraint Programming.
- **Chapter 3** introduces a powerful constraint of difference.
- **Chapter 4** explains the two algorithms needed to solve a constraint of difference.
- **Chapter 5** covers a presentation of the project solving a Sudoku puzzle.
- **Chapter 6** explains the implementation of the project.
- **Chapter 7** details the overall achievements of the project.

## Chapter 2

# Background

### 2.1 Sudoku

Sudoku, meaning *single number* in Japanese, is a logic-based, combinatorial puzzle that became mainstream in 1986. Today, many magazines and newspapers from all over the world include a Sudoku puzzle. The aim of the game is to place digits from 1 to 9 into a  $9 \times 9$  grid, such that the same digit may not appear twice in the same row, column or  $3 \times 3$  sub-squares of the puzzle. A Sudoku puzzle starts as a partially completed grid that has to be filled. Enough clues are provided such that the puzzle has a unique solution. An example of a relatively hard Sudoku puzzle is shown in Fig. 2.1. Variations of the puzzle include instances featuring  $n^2 \times n^2$  sized boards with  $n \times n$  sub-squares. Since the Sudoku puzzle is known to be NP-complete, this means that a general Sudoku requires significantly more computational resources as  $n$  increases.

		9				2	6	
2				8	4			
5			2					1
	6		1		9	7		
	9			7			4	
		7	4		3		9	
3					6			5
			5	3				8
	2	5				3		

**Fig. 2.1:** A Sudoku puzzle

<sup>1</sup> <sub>4</sub> 7 8	<sup>1</sup> <sub>4</sub> 7 8	9	<sup>3</sup> <sub>3</sub>	<sup>1</sup> <sub>5</sub>	<sup>1</sup> <sub>5</sub>	2	6	<sup>4</sup> <sub>7</sub>
2	<sup>1</sup> <sub>3</sub> 7	<sup>1</sup> <sub>3</sub> 6	<sup>6</sup> <sub>9</sub>	8	4	<sup>5</sup> <sub>9</sub> 7	<sup>5</sup> <sub>3</sub> 7	<sup>3</sup> <sub>9</sub>
5	<sup>3</sup> <sub>4</sub> 8	<sup>3</sup> <sub>4</sub> 8	<sup>3</sup> <sub>6</sub>	2	<sup>6</sup> <sub>9</sub>	<sup>7</sup> <sub>7</sub>	<sup>3</sup> <sub>8</sub>	1
<sup>4</sup> <sub>8</sub>	6	<sup>2</sup> <sub>3</sub> 4	<sup>2</sup> <sub>3</sub> 8	1	<sup>2</sup> <sub>5</sub>	9	7	<sup>2</sup> <sub>3</sub> 5
<sup>1</sup> <sub>8</sub>	9	<sup>1</sup> <sub>2</sub> 3	<sup>1</sup> <sub>2</sub> 8	<sup>6</sup> <sub>8</sub>	7	<sup>2</sup> <sub>5</sub> 8	<sup>1</sup> <sub>5</sub> 6	<sup>2</sup> <sub>3</sub> 4
<sup>1</sup> <sub>8</sub>	<sup>1</sup> <sub>5</sub> 8	7	4	<sup>2</sup> <sub>5</sub> 6	3	<sup>1</sup> <sub>5</sub> 6	9	<sup>2</sup> <sub>6</sub>
3	<sup>1</sup> <sub>4</sub> 7	<sup>1</sup> <sub>4</sub> 8	<sup>7</sup> <sub>8</sub> 9	<sup>1</sup> <sub>4</sub> 2	6	<sup>1</sup> <sub>4</sub> 9	<sup>1</sup> <sub>2</sub> 7	5
<sup>1</sup> <sub>4</sub> 7	<sup>1</sup> <sub>4</sub> 6	<sup>1</sup> <sub>4</sub> 7	5	3	<sup>1</sup> <sub>2</sub> 4	<sup>1</sup> <sub>4</sub> 6	<sup>1</sup> <sub>2</sub> 7	8
<sup>1</sup> <sub>4</sub> 7	<sup>1</sup> <sub>4</sub> 6	2	5	<sup>1</sup> <sub>4</sub> 7	<sup>1</sup> <sub>8</sub>	3	<sup>1</sup> <sub>7</sub>	<sup>4</sup> <sub>6</sub> 9

**Fig. 2.2:** The state of the puzzle after the propagation

One efficient way of solving Sudoku puzzles is by using techniques and algorithms used in Constraint Programming. The puzzle can be modelled as a Constraint Satisfaction Problem that has 81 variables and 27 all-different constraints, one for each of the 9 rows, column and  $3 \times 3$  sub-squares of the puzzle. Each constraint in the Sudoku puzzle represents a relationship between 9 variables that make up a row, column or  $3 \times 3$  sub-square. Variables that represent clues given in the initial puzzle have a domain that contains a single value, while the unknown variables have the set of digits from 1 to 9 as their domain of possible values. Multiple techniques are then used to get to a solution by removing values from the domain of the variables until each one of them is left with only a single value in their domains, representing the correct assignment to be used in the solution.



Propagation schemes are used to achieve consistency of the constraints. This means that for each variable that has been solved (has a single digit in their domain of possible values), one can remove that value from the domain of the variables that take part in the same constraints. In the case of the Sudoku puzzle, each variable is part of 3 all-different constraints, one for the row it takes part in, one for the column, and one for the  $3 \times 3$  sub-square. Propagation is done starting from the variables representing clues in the initial puzzle, and can trigger recursive calls once other variables have their domain reduced to a single value. The recursive calls stop when no more values can be removed from any of the variables' domains. Easy Sudoku puzzles can be solved by using this technique alone.

The Fig. 2.1 shows the state of the initial Sudoku puzzle after a recursive propagation is made. One can notice that the clues provided in the initial puzzle were enough to find out the correct assignment for two variables (circled in the figure). The rest of the variables had their domains reduced to keep the constraints consistent. Notice the first column that contains two variables that have  $\{1, 8\}$  as their domain. This is place where a human may resort to guessing, and where an intelligent algorithm should be used so one can find a solution to the puzzle. The all-different algorithm notices that values 1 and 8 should be assigned to the two variables at a later stage in the solution. This is enough information to decide that neither 1 nor 8 can be taken by the rest of the variables in the column. This will result in the variable that has  $\{4, 8\}$  as its domain to be assigned the value 4 and will inform the other variables in the same column that they cannot take the value 4. Since this is known, the variable that had  $\{1, 4, 7, 8\}$  as its domain will be forced to take the value 7 as the rest of the values were previously assigned to other variables. By applying the same all-different algorithm on rows, columns and  $3 \times 3$  sub-squares, one will eventually find the unique solution without having to resort to any guessings. Since the Sudoku puzzle is an NP-complete problem, there is no way to know beforehand which row, column or sub-square will solve the most variables. The advantage of the algorithm is that the search space for a solution is pruned by removing values that will never be part of the solution, making the search much faster.

## 2.2 Constraint Programming

Constraints can be found everywhere in our daily life, and represent mathematical abstractions of the dependencies in the physical world. More formally, a constraint is a logical relationship among a number of variables, each with its own domain. The constraint therefore imposes restrictions on the values the variables can take. It is good to note that the values of the variables are not always numeric, a heterogeneous constraint would require a word (string) to have a specific length (numeric). The constraints are declarative as they only specify a relationship that must hold between variables and do not provide an algorithm to enforce the relationship.

Constraint Programming is the area of Computing Science that solves problems by specifying a list of constraints and then finding the solutions that satisfy all the constraints. A Constraint Satisfaction Problem (CSP) can be defined as a triple  $\mathcal{P} = \langle X, D, C \rangle$ , where  $X$  represents a set of  $n$  variables  $X = \langle x_1, x_2, \dots, x_n \rangle$ ,  $D$  represents the  $n$  domains associated to the  $n$  variables  $D = \langle D_1, D_2, \dots, D_n \rangle$  such that  $x_i \in D_i$ , and finally  $C$  represents a set of  $t$  constraints  $C = \langle C_1, C_2, \dots, C_t \rangle$ . A constraint  $C_i$  restricts the values one or more variables can simultaneously take.

A solution for a CSP  $\mathcal{P}$  is a set of values  $S = \langle s_1, s_2, \dots, s_n \rangle$ , such that each variable gets assigned a value from its domain and all the constraints are satisfied at once. The set of all the solutions to the problem is noted as  $sol(\mathcal{P})$  for a CSP that has multiple solutions. One may also be interested in finding an optimal solution to the problem, one that for example minimizes the path of a traveling salesman. When  $sol(\mathcal{P})$  is equal to the empty set, it means that the original CSP is unsatisfiable. This simple representation can be used to model complex real world problems such as planning, scheduling, timetabling, supply chain management and more.

There are two strategies for solving CSPs: inference and search. If all the variables in the problem have finite domains, this means the search space for a solution is finite and represented by all the possible combinations

of the values of the variables. Although one can, in theory, enumerate all the possible combinations, **TODO WE:**we use inference and search to reduce the search space. Inference techniques remove large subspaces of the search tree through local constraint propagation on the basis that no solutions may be found using a particular value from the domain of a variable. Search techniques, such as backtracking, are used to systematically explore the search space, usually with the use of some heuristics, and reduce the search space whenever a single failure is found. Both strategies are frequently used together and work because a solution to a CSP must have all its constraints satisfied, therefore a local inconsistency on a subset of the variables will not result in a solution.

Backtracking search algorithms look for solutions to a CSP by traversing the search tree in a depth-first search manner. At each node of the search tree, a variable is assigned a value out of its domain, and the node gets extended with branches for the remaining values in the variables domain that need to be considered later, as they could be part of a solution. For example, consider a simple CSP problem with three variable  $x$ ,  $y$  and  $z$ . Suppose that the search algorithm already assigned a value to the variable  $x$  out of its domain, and it reaches a node representing the variable  $y$  with the domain  $D_y = \{1, 2, 3\}$ . The backtracking search algorithm will generate three branches, each one of them representing the variable  $y$  getting instantiated to a value from its domain. For every value assigned to the variable  $y$ , all the constraints tied to this variable and the previously visited variables are checked to see if they still hold. When the constraints are not satisfied by a selected value for a variable, the algorithm proceeds in assigning and testing the rest of the values in the domain of  $y$ . Once all the values from the domain of the variable  $y$  have been tried, the algorithm will backtrack to the previous variable,  $x$  in this case, and assign it the next value from its domain. Note that every time backtracking occurs, subtrees of the search tree will no longer be generated and visited since they will not contain any solutions to the CSP. In the problem mentioned above, the subtree for the values of  $z$  is no longer generated and tested as the partial solution made out of some specific values for  $x$  and  $y$  already violated the constraints of the problem. A complete solution is found once all the variables are assigned values that are allowed by the constraints. If the user is interested in only one solution to the problem, the search will terminate, otherwise the algorithm will continue to explore the search tree for other solutions. In the case the problem has no solutions, the algorithm will terminate after visiting all the possibilities in the search tree.

The efficiency of the backtracking search algorithm can be improved by performing constraint propagation to maintain consistency between the values of the variables and their associated constraints. Every time a variable is assigned a value from its domain, constraint propagation is made on the constraints the variable was part of. Propagation will check all the variables that were in a constraining relationship with the original variable and will remove values from their domain that are inconsistent with the problem requirements. By reducing the domains of the variables, the search tree is pruned and the efficiency of the search increases. The domain of a variable may be reduced to the empty set after propagation, which means that there is no value left for that variable that will satisfy the constraints and therefore backtracking should be initiated. On other occasions, the domain of a variable will be reduced to a single value, which means that the value for the variable is now known and removing the need to search through values that would not lead to a solution. The backtracking search algorithm will resume when the recursive propagation results in no changes to the domains of the variables. Information from constraint propagation is usually incorporated in the search algorithm in form of variable and value ordering heuristics that improve the efficiency of the search even more.

Variable and value ordering heuristics are ways to improve the backtracking search algorithm by prioritizing the instantiation of some variables and by choosing some values from their domain first. Ordering heuristics are essential to effectively solve hard combinatorial problems. The most popular heuristic is the fail-first approach, something Haralick and Elliot described as To succeed, try first where you are most likely to fail. [5]. The principle is to prioritize the instantiation of variables with the smallest domains of values. The assumption behind this is that branches in the search tree that have no solutions will be discovered earlier and pruned to bring a noticeable increase in efficiency later in the search. Consider the worst-case scenario when performing a backtracking search on a problem that has no solutions, the search tree will be much smaller as earlier failure is encouraged, making it faster to prove that a problem has no solutions. In the case of problems that start off with equally sized domains, the fail-first principle can still be applied by prioritizing the instantiation of the variables

that take part in many constraints. Once a variable was selected for instantiation, some values from its domain could lead to a solution faster than others. The order values are chosen for assignment to the variables is only relevant when trying to find a single solution to the problem. The reason behind this is that trying to find all the solutions to a problem or proving that there are no solutions would have to traverse all the search space anyway. A heuristic to prioritize values from a domain is to assess the likelihood each value has in getting closer to a solution, a choice that will have minimum impact on the domain of other variables after propagation, therefore increasing the chances of finding a solution faster.

Some solutions to a problem are better than others. Consider the Traveling Salesman Problem (TSP), where the shortest possible path between some cities should be found. The problem is NP-hard as the running time to find the solution increases exponentially with the number of cities. The general approach to the optimization problem is to introduce a constraint that minimizes a variable representing an objective, the length of the path in the case of the TSP. Finding an optimal solution is often hard, and proving optimality of a solution might be impossible for complex combinatorial problems.

Constraint Programming can be regarded as a form of declarative programming, where a user specifies a problem in terms of its variables, their domains and the constraints that should be satisfied. The computer then solves the given problem using a range of techniques that are general approaches to problem solving which can be applied to all the problems that can be modeled as constraint satisfaction problems.

The original exercise of finding an algorithm to solve a specific problem is now turned into an exercise of finding the best way to model the problem as a CSP, which can later be solved by the computer. The model can make the difference between efficient solving of a problem or a problem that cannot be solved due to its combinatorial complexity. There are usually many ways in which one can model the same problem, but the efficiency of the solution could be different. Consider a problem in which a number of  $g$  guests should be offered a seat at one of the  $t$  tables, with the constraint that couples should sit together. One could decide to represent each of the  $g$  guests as a variable associated with an initial domain representing a listing of the  $t$  tables, from which a choice for the table should be made based on the constraints. Another way to model the same problem is to have a number of  $t$  variables representing the tables, each with an initial domain representing a listing of all the guests. The search for a solution would reduce the domains of the variables in both models until there is only a single table in the domains of the variables (for the first model where guests are variables), or in the case of the second model, until all the variables representing tables have the cardinality of their domains reduced to their sitting capacity. Although both models are logically equivalent, one may perform worse than the other because of how its represented in the memory of the computer.

When modeling a problem, it is important to avoid symmetries in the form of equivalent solutions. Symmetries represent waste computational resources during the backtracking search that is used to explore symmetrical assignments. One should avoid such symmetries by adding new constraints that will filter out the symmetrical solutions. Such constraints are problem specific. If it is necessary to find all the solutions to a problem, including the equivalent ones, one can easily perform permutations of the variables outside of the search space once the main solutions are found, thus avoiding exploring similar branches in the search tree. In the example problem mentioned above, there are symmetries in the solutions, as the same group of guests will be seated at different tables throughout the solutions. **TODO WE:**As we know tables are the same in real life and can be easily swapped without violating the constraints, such computations should not be made to improve efficiency of the search.

Real world problems are often large and over-constrained, which may raise difficulties in modeling them. Most of the time there are conflicting objectives such as trying to maximize the efficiency of a production facility, while minimizing the operational costs. Tradeoffs between the constraints should be made to come up with a solution to the problem. Problem decomposition should also used to break a large real world scenario into smaller problems that are easier to solve. Finally, one should consider which model is easier to upgrade when more constraints should be added.

## Chapter 3

# The All-Different Constraint

### 3.1 Review of Jean-Charles Regin's paper

This section will introduce the reader to Jean-Charles Regin's paper published in 1994, which describes a filtering algorithm for constraints of difference used in solving CSPs. As mentioned before, because the search for all the solutions to a CSP is NP-complete, there is a justified need to prune the search space before both before and during the search for the solutions.

A constraint of difference is defined as a relationship on a subset of the variables of a problem for which the values that are assigned to them should be all different. Many real world problems require such constraints. The paper starts by recognizing the need for a powerful n-ary constraint of difference, as binary constraint of difference cannot efficiently prune the search space of values for variables that will never be part of a solution, and for most instances it does not delete any value. Before the publishing of the paper, the common practice to represent n-ary constraint of difference was to model it as many binary constraints, one for each pair of variables in the n-ary constraint.

Consider a CSP problem with three variables  $x$ ,  $y$  and  $z$  with their domains  $D_x = \{1, 2\}$ ,  $D_y = \{1, 2\}$  and  $D_z = \{1, 2, 3\}$  with a constraint that solutions should have different values assigned to the variables. The first way one could model such a problem is by introducing 3 binary constraints of difference for all the pairs of the variables:  $\{x, y\}$ ,  $\{x, z\}$  and  $\{y, z\}$ . The second way, is to use a 3-ary constraint of difference between the variables  $\{x, y, z\}$ . The difference between the two approaches is that a propagation of the constraints for the binary representation will not reduce the domains of the variables. On the other hand, the model using a 3-ary constraint will successfully remove the value 1 and 2 from variable  $z$ , as they will already be used by  $x$  and  $y$ .

Jean-Charles Regin goes on to describe a generalized constraint of difference to take advantage of the improved pruning of the search space. The aim is to achieve arc-consistency across all the variables in the CSP. A variable  $x$  is said to be arc-consistent with the other variables participating in the same n-ary constraint if for all the values in  $D_x$ , there exists admissible values in the domains of the other variables such that the constraint holds. For the purpose of this exercise, the aim is to achieve diff-arc-consistency, which is concerned with achieving arc-consistency across the constraints of difference.

A constraint of difference  $C$  can be represented by its value graph. The value graph of  $C$  is a bipartite graph where the variables in the constraint are grouped in the left partition of the graph, while the right partition represents a union of all the values from the domains of the variables. Edges between the two partitions of the graph connect the variables on the left hand side with values from their domains.

Matches in the value graph of  $C$  are a subset of the edges in the graph, where no two edges share a common

vertex. A maximum match is when all the vertices in the left partition have an edge to a vertex from the right partition, or more specifically, all the variables in the left partition are assigned a value from the right one. After finding maximum matchings for all the value graphs associated to the constraints of difference in a given CSP, the values assigned to the variables do not break the diff-arc-consistency for the problem. Since each variable is assigned a value from its domain represented by an edge in the graph, and no two variables are assigned the same value, the diff-arc-consistency for the problem still holds. Knowing this property, an algorithm is built to filter the domains of the variables by removing values that would never be part of a maximum matching, therefore breaking the constraint and not leading to any solutions to the problem. It is good to note that by using the notation mentioned earlier, all the constraints of difference can be represented by their value graphs that have a space complexity in  $O(pd)$ , where  $p$  is the arity of the constraint and  $d$  is the maximal cardinality of the domains for the variables in the constraint. Computing maximum matchings in a bipartite graph  $G = (X, Y, E)$  with  $m$  edges, can be done using Hopcroft and Karp's (1973) algorithm which has a complexity of  $O(\sqrt{d} \cdot m)$  or Ford-Fulkerson (TODO) algorithm with a complexity  $O(VE)$ .

To begin reducing the domains of the variables, more definitions regarding matches are introduced. First, a vertex is *matched* if there is an edge connecting it to a vertex in another partition of the graph. Vertices that are not matched are said to be *free*. Alternating paths represent paths made of edges that are alternatively matches and free and have a length corresponding to the number of edges in the path. Vital edges are edges that are part of all the maximum matchings in the value graph of a constraint.

Berge's property [1] is used to find the edges that correspond to no maximum matchings and can therefore be deleted. According to the property, such edges are not vital and when starting from an arbitrary maximum matching, no even alternating paths can be found to include that edge. The algorithm starts by computing a maximum matching for the value graph of a constraint of difference. After this, an algorithm such as the one proposed by Tarjan is used to find the strongly connected components in the graph. This means that edges in such a component will belong to even alternating paths, as the graph is bipartite and doesn't have any odd cycles. The step of finding strongly connected components is necessary as this means that every edge connecting vertices from two strongly connected components can be deleted as it will never be part of an even alternating path. The deletion of an edge in the value graph of the constraint represents the reduction of a variable's domain and is done for reasons independent from the constraint. The complexity of this algorithm proposed by Jean-Charles Regin is of  $O(p^2 \cdot d^2)$  for one constraint of difference.

Reducing the domain of a variable by deleting edges in the value graph of a constraint can trigger modifications for other constraints involving the same variable. If a variable is part of other constraints of difference, the author shows a more efficient propagation approach than simply repeating the same algorithm. The filtering algorithm for the second constraint builds on the information from the previous step by trying to remove the same edges in the value graph of the new constraint as long as they are not vital and a new maximum matching can be found.

The paper goes on to solve the zebra problem, a famous logic puzzle. The author models the problem in different ways to prove that his approach can prune the search space of the problem more when compared to the previous methods used in the industry. One representation of the problem uses binary constraints for each pair of variables involved in the constraint of difference and another representation uses the  $n$ -ary constraint of difference introduced by the author.

The results of his experiment show that the search space of the problem gets pruned more using the approach proposed in the paper, making the search for a solution to the problem more efficient. The same method was successfully used to solve the subgraph isomorphism problem.

## 3.2 Usage example

To demonstrate the usage and effects of the all-different constraint, **TODO WE:**we are going to use a simple example. The CSP problem will be modeled in Java and will make use of Choco3, a powerful free and open-source Java library used to solve Constraint Satisfaction Problems. The implementation of the all-different constraint in the Choco3 library is based on Jean-Charles Regin’s paper[7] reviewed above.

The problem has three variables  $x, y, z$  with the following initial domains:  $x = \{1, 2\}$ ,  $y = \{1, 2\}$  and  $z = \{2, 3\}$ . The constraint that should be satisfied by all the solutions of the problem is that  $x, y$  and  $z$  should all be different. The listing Section 3.2 shows how the problem is modeled and implemented using the Choco3 library.

```
1 Solver solver = new Solver("AllDifferentExample");
2
3 IntVar x = VF.enumerated("x", new int[]{1, 2}, solver);
4 IntVar y = VF.enumerated("y", new int[]{1, 2}, solver);
5 IntVar z = VF.enumerated("z", new int[]{2, 3}, solver);
6
7 solver.post(ICF.alldifferent(new IntVar[]{x, y, z}));
8
9 try{
10     solver.propagate();
11 } catch (ContradictionException e){
12     // Handle the exception
13 }
14
15 System.out.println(x + "\n" + y + "\n" + z);
16
17 if(solver.findSolution()){
18     do{
19         int x_val = x.getValue();
20         int y_val = y.getValue();
21         int z_val = z.getValue();
22
23         System.out.println(x_val + ";" + y_val + ";" + z_val);
24     } while(solver.nextSolution());
25 }
```

**Listing 3.1:** Usage of the all-different constraint in Choco3

Line 1 defines a Choco3 solver object, that will store the variables, their domains and constraints associated to them. Lines 3 to 5 define the three integer variables in the problem and their associated initial domains. The reference to the *solver* object makes sure the variables are added to the model. Line 7 introduces the *all – different* constraint between the three variables. The constraint is then posted to the model and the CSP is now fully modeled.

Before trying to find a solution, a call to the *propagate* method is made on the model on line 10. The propagation algorithm will make all the constraints consistent by looping thorough all the constraints in the model (only one in this case) with the aim of reducing the domains of the variables by removing values that will never part of a solution. When a value is removed from the domain of a variable, this might trigger a recursive propagation call on all its dependant variables to make the constraints consistent. The recursive propagation calls end when the problem is in a consistent state. It is good to note that propagation will preserve all the original solutions to the problem, while greatly reducing the search space.

The *ContradictionException* on line 11 is an exception that is raised by Choco3 whenever a variable is left with an empty domain or is instantiated to a value that is out of its initial domain. This usually happens in overconstrained problems, where there are no values in the domains of the variables that satisfy all the constraints at once. In such cases, there would be no valid solutions to the problem. This is not the case with the example showcased here, as there are three variables with three different values in total.

Once the propagation is finished, a call is made on line 15 to output the updated domains of the variables. The updated domains after the call to the *propagation* method are as follows:  $x = \{1, 2\}$ ,  $y = \{1, 2\}$  and  $z = \{3\}$ . The variable  $z$  lost the value 2 from its domain, while to the domains of  $x$  and  $y$  stayed the same. There are two reasons behind removing the value 2 from the domain of variable  $z$ . Firstly, there would no solutions with  $z$  taking the value of 2 as this would leave either  $x$  or  $y$  with an empty domain after one of them takes the value of 1. Secondly, two variables ( $x$  and  $y$ ) have the same domain of  $\{1, 2\}$ , therefore a solution that is consistent with the all-different constraint will have  $x$  taking the value of 1 and  $y$  taking the value of 2 (or vice-versa), making it impossible for  $z$  to take the value of 2. Since **TODO WE:**we know that the values 1 and 2 will be attributed to  $x$  and  $y$ , the value 2 is removed from the domain of  $z$  to reflect this change and reduce the search space.

Lines 17 to 25 will iterate over the two solutions of the problem and print the values for the variables. The solutions to the problem are  $S_1 = \{1, 2, 3\}$  and  $S_2 = \{2, 1, 3\}$  and they both contain values for the three variables that satisfy the all-different constraint.

## Chapter 4

# Algorithms for All-Different

### 4.1 Ford Fulkerson

A flow represents a directed graph with special nodes called the source and the sink. Source nodes have outgoing edges to inner nodes, while sink nodes have incoming edges from inner nodes of the graph. The problem requires sending flows, for example water through a pipe, from the source to the sink, while conserving the equilibrium.

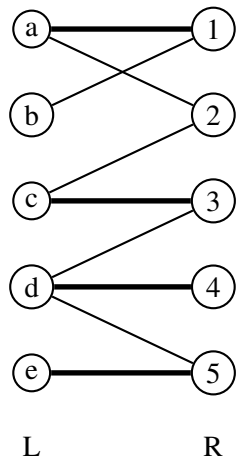
Flows need to get from the source to the sink by following the directed edges that often have different maximum capacity constraints, such as the diameter of a water pipe. Given a flow  $f$  in graph  $G$ , **TODO WE:**we build a residual directed graph  $G_f$ . Forward edges in the graph haven't reached their maximum capacity, while backward edges reached their upper bound.

Ford-Fulkerson algorithm uses the residual graph to achieve a maximum flow in the graph. The algorithm looks for augmenting paths from source  $S$  to sink  $T$  in a breadth first-search manner. Once a path  $p$  is found, **TODO WE:**we send the weight that will make the bottleneck edge in  $p$  fill its maximum capacity. Then, **TODO WE:**we add (subtract) the weight of each forward (backward) edge in  $p$ . Forward edges that reach their maximum capacity turn into backward edges. This results in flow  $f$  that has a greater value than before. Berge's theorem guarantees that once all the augmenting paths have been successively visited in the algorithm, the end result is a maximum flow of the graph [1].

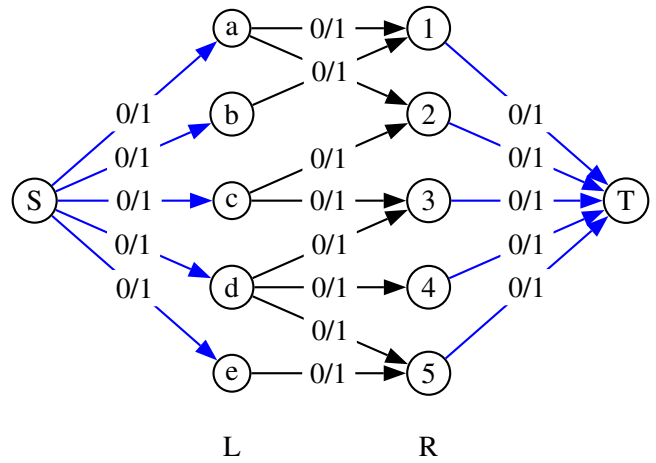
The maximum matching problem is a special case of the maximum flow problem. For the purpose of our example, there will be only one source  $S$  that has outgoing edges to the nodes in the left partition  $L$  of the bipartite graph and one sink  $T$  with incoming edges from the right partition  $R$  of the graph. Nodes in  $L$  are connected through edges to nodes in  $R$ . A matching **TODO**. A maximum matching **TODO**. To find a maximum matching for a Graph  $G$ , **TODO WE:**we add a source and sink nodes and set all edges to a maximum capacity of one. **TODO WE:**We then run the Ford-Fulkerson algorithm on the graph, by finding all the augmenting paths from free nodes in  $L$  to free nodes in  $R$ .

Having a maximum matching for our graph, the next step in the all-different algorithm is to turn the previously undirected graph, into a directed one. The matches resulted from the Ford Fulkerson algorithm are assigned a direction from the 1 to 9 values to the corresponding cells of the Sudoku row. The edges that remain unused in the matching are given a direction from left to right (i.e. from the Sudoku cells of the row to the 1-9 values).

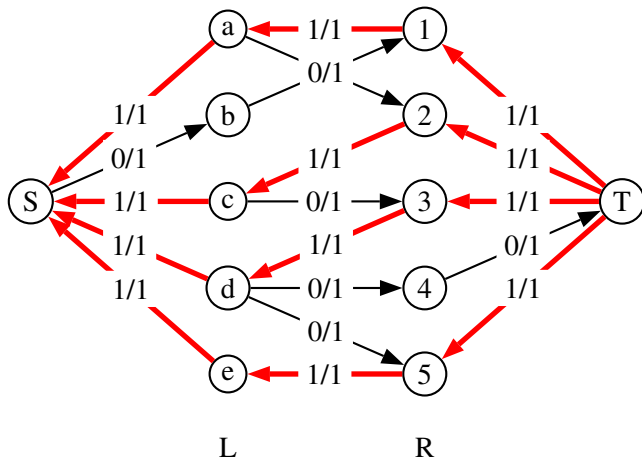




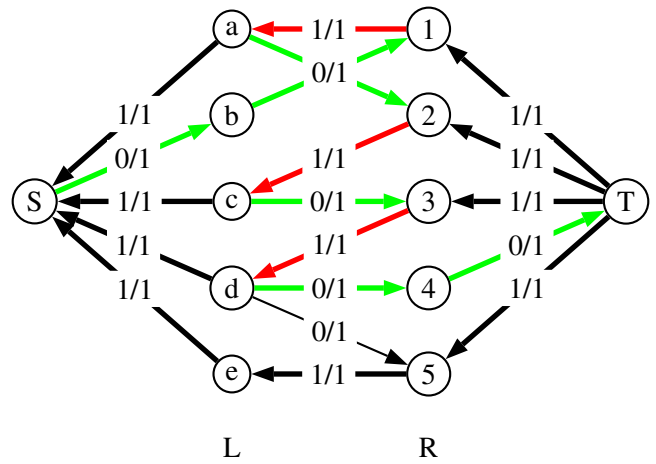
**Fig. 4.1:** Initial graph



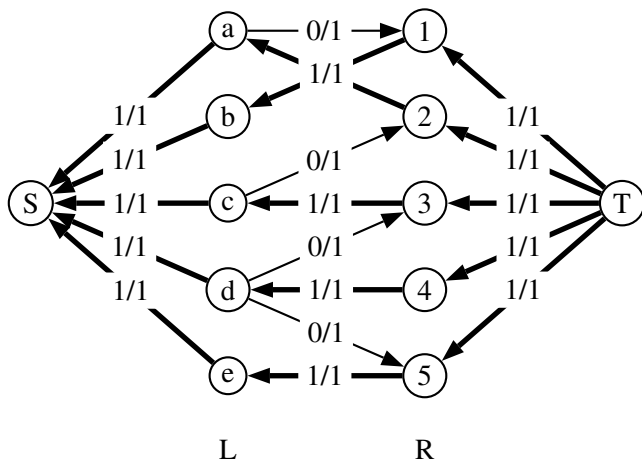
**Fig. 4.2:** S&T nodes added



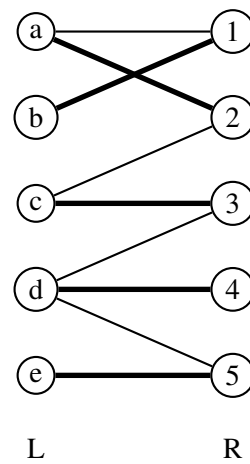
**Fig. 4.3:** Greedy match



**Fig. 4.4:** Longer augmenting path



**Fig. 4.5:** New matching



**Fig. 4.6:** Maximum matching

**Fig. 4.7:** FordFulkerson algorithm

---

**Algorithm 1: Ford Fulkerson**

---

```
1 void FordFulkerson(Graph G)
2 begin
3   Global int capacity[][]  $\leftarrow$  getCapacity(G)
4   Global int n  $\leftarrow$   $|V(G)|$ 
5   Global int source  $\leftarrow$  0
6   Global int sink  $\leftarrow$  n - 1
7   Global pred  $\leftarrow$  new int[n]
8   Global visited  $\leftarrow$  new boolean[n]
9   Global Q  $\leftarrow$  new Queue()
10  while bfs(G) do
11     $\lfloor$  update()

12 boolean bfs(Graph G)
13 begin
14   clear(Q)
15   fill(pred, -1)
16   fill(visited, false)
17   enqueue(source, Q)
18   visited[source]  $\leftarrow$  true
19   while ( $\neg$ isEmpty(Q)) do
20     int v  $\leftarrow$  dequeue(Q)
21     if v = sink then return true
22     for w  $\leftarrow$  0 to n do
23       if  $\neg$ visited[w] and capacity[v][w] > 0 then
24          $\lfloor$  pred[w]  $\leftarrow$  v
25          $\lfloor$  enqueue(w, Q)
26          $\lfloor$  visited[w]  $\leftarrow$  true
27   return false

28 void update()
29 begin
30   int f  $\leftarrow$  minCost()
31   int v  $\leftarrow$  sink
32   while pred[v]  $\neq$  -1 do
33     int u  $\leftarrow$  pred[v]
34     capacity[u][v]  $\leftarrow$  capacity[u][v] - f
35     capacity[v][u]  $\leftarrow$  capacity[v][u] + f
36     v  $\leftarrow$  u

37 int minCost()
38 begin
39   int minCost  $\leftarrow$   $\infty$ 
40   int v  $\leftarrow$  sink
41   while pred[v]  $\neq$  -1 do
42      $\lfloor$  minCost  $\leftarrow$  minimum(minCost, capacity[pred[v]][v])
43      $\lfloor$  v  $\leftarrow$  pred[v]
44   return minCost
```

---

Ford Fulkerson's algorithm for finding a maximum flow in a graph is shown in algorithm 1. [3]. The *Ford Fulkerson* procedure on line 1 takes a graph  $G$  as an argument and represents the start of the algorithm. The graph passed as an argument holds information about the capacity of each edge. Line 3 stores the capacity of the graph  $G$  in a global two-dimensional integer array. The capacity of an edge between vertices  $u, v$  is stored in  $capacity[u][v]$ . Line 4 declares an integer  $n$  used to store the number of vertices in the given graph. Lines 5 and 6 declare two integers called *source* and *sink* representing references to the first and the last vertices in the given graph.

In line 7 **TODO WE:**we declare a vertex-indexed array *pred* that is used to store references to the previous vertex discovered in the breadth-first search tree. Line 8 declares a vertex-indexed array *visited* used to keep track if a vertex has already been visited in the breadth-first search procedure. Line 9 declares a queue of integers  $Q$  used to store vertices in the order they are visited.

The algorithm consists of repeated calls to the breadth-first search procedure (line 10). In line 11, **TODO WE:**we call the update procedure to update the capacities of the edges after finding augmenting paths.

The *breadth-first search* procedure starts at line 12 and takes a graph  $G$  as an argument. In lines 14 to 16, the queue  $Q$ , *pred* and *visited* array are reset to have default values. The breadth-first search calls always start from the first vertex, representing the source. This vertex is enqueued on queue  $Q$  in line 17 and is marked as visited in line 18.

Lines 19 to 26 contain a loop performing changes on the queue  $Q$  that is used to store an augmenting path. The loop starts on line 20 by dequeuing a vertex from the queue  $Q$  and storing it in  $v$ . Line 21 forces the breadth-first search procedure to return *true* when **TODO WE:**we reach the last vertex in the graph, the sink. Lines 22 to 26 contain a loop over all the vertices  $w$  in the graph. In line 23 **TODO WE:**we check to see if the discovered vertex  $w$  hasn't been already visited and **TODO WE:**we consider it if it still has available capacity. Line 24 stores the index value of  $w$ 's predecessor in the augmenting path. Line 25 enqueues the discovered vertex  $w$  on the queue  $Q$  and marks it as visited in line 26. Finally, the breadth-first search procedure returns *false* when no more augmenting paths are found.

The *update* procedure at line 28 is used to update the capacities of the edges in the graph to reflect the updated flow. Line 30 declares an integer  $f$  used to store the amount of flow **TODO WE:**we are adding along the found path. The value for  $f$  is calculated in the *minCost* procedure at line 37. Line 31 declares an integer  $v$ , initially representing the sink which is the last vertex in the graph. Lines 32 to 35 contain a loop that updates the capacities of the edges in the augmenting path. Line 33 uses the integer  $u$  to store the index of a vertex that is the predecessor of vertex  $v$  along the path. Lines 34 and 35 updates the flow along the edge from  $u$  to  $v$  and  $v$  to  $u$  by the found difference. Finally,  $v$  is updated on line 36 to take the value of  $u$  such that the while loop continues to change the capacities of the remaining edges along the path.

The *minCost* procedure at line 37 is used to find the amount of flow to be sent along the new augmenting path from the sink to the source. The value is equal to the minimum capacity that is still available along the edges of the path. Line 39 declares an integer *minCost* used to represent the current value of the minimum cost, initialized to a big value. Line 40 declares an integer variable  $v$  initially used to store a reference to the sink vertex. Lines 41 to 43 contain a loop over the edges along the path, starting from the sink toward the source. Line 42 updates the value of *minCost* if the current edge in the loop has a smaller capacity left than what was previously discovered. The *minimum* procedure returns the minimum of two given integers. Line 43 updates the value of integer  $v$  to its predecessor so the loop can continue along the path until it reaches the *source*. At the end of the *minCost* procedure, **TODO WE:**we return the value of the currently found *minCost* variable.

## 4.2 Tarjan algorithm

A strongly connected component *SCC* of a directed graph  $G$  is a maximal set of vertices such that every vertex is reachable from every other vertex by following the directed edges.

There are four kinds of edges in a depth-first traversal: tree edges, forward edges, back edges and cross edges. Tree edges are edges between a vertex and vertices that were not previously visited. Forward edges are from ancestor vertices to descendant vertices that were already visited during the search. Back edges link descendant vertices back to already visited ancestor vertices. Lastly, cross edges represent edges between two vertices that are neither an ancestor nor a descendant for each other, such as the edges between different components of the graph.

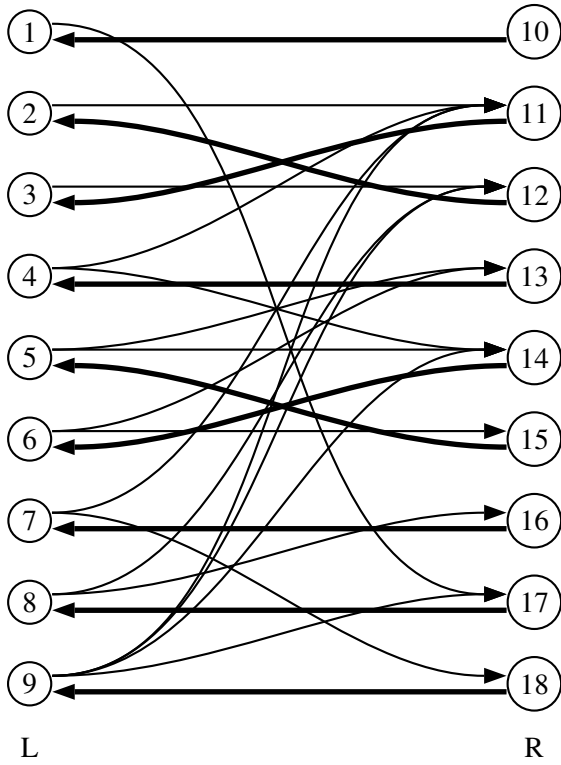
For each vertex in a graph  $G$ , the algorithm keeps track of two properties. The first property **TODO WE:**we remember is the *index* of the node, its consecutive order of discovery during the depth-first search, a value that will remain constant throughout the algorithm. The second property associated with all the nodes is named *low* and keeps track of the lowest (oldest) ancestor reachable from their position in the graph. As each node of the graph gets discovered in a depth-first search manner, the initial value that *low* gets assigned is the same as the *order* the node was discovered. The value for *low* may get changed during the search if an ancestor for the node is discovered by following a back edge. After the recursive depth-first search finishes visiting all of a nodes neighbours, the nodes value for *low* gets updated to the lowest index of its oldest reachable ancestor. If the value for *low* of a node stays the same even after visiting all of its neighbours, then it means it is the root of a strongly connected component or its an individual vertex that will be a component on its own. [10]

For the purpose of finding SCCs, every time a node is discovered in the search, the node is put on a stack  $S$ . After returning from the recursive DFS calls on all of the adjacent neighbours of vertex, if both *index* and *low* have the same value, it means **TODO WE:**weve finished discovering a SCC. The vertices in the newly discovered strongly connected component will be popped from the stack  $S$  until **TODO WE:**we reach the current vertex representing the root of the SCC. Vertices that are part of other SCCs are still left on the stack as they have different roots represented by the value of *low*.

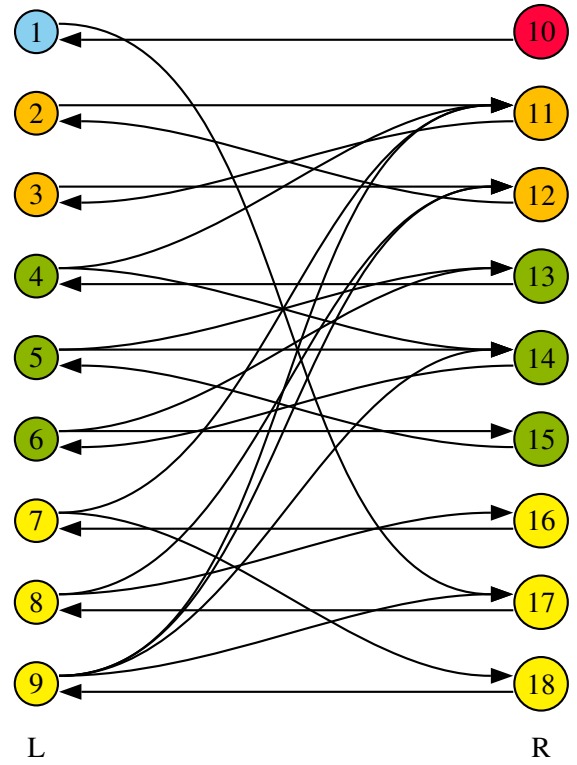
The Fig. 4.8 shows a bipartite directed graph with 18 vertices. There are 9 backward edges that represent the maximum matching found using the Ford-Fulkerson algorithm in Section 4.1. The rest of the possible edges are assigned a forward direction. This graph is similar to the graphs that will be generated to represent a row, column or one of the nine sub-grids of the Sudoku puzzle. For example, the left partition of the graph will contain the nine variables of a Sudoku row, and the right partition will contain the digits from 1 to 9. The edges will then link each variable of the row to values that are still available in their domain.

The Table 4.1 contains the output after running the Tarjans algorithm on the graph in Fig. 4.8. As previously discussed, vertices are visited in a depth-first search manner, in this case starting from the first vertex. Each node has two properties associated to it: its order of discovery or *index*, and its oldest reachable ancestor. Initially, both properties take the value of the index, but as the algorithm progresses the values for *low* may become lower after visiting all of the neighbours of the node and finding the oldest (lowest) reachable ancestor. All vertices are pushed on a stack in the order they are visited. Once a back edge is found in the graph, vertices that are currently on the stack get popped including the root of the newly discovered SCC. Note that vertices may still remain on the stack as they will be part of different components that have other roots. As each vertex is made part of a component, its value for *low* gets updated to a large value to mark it as already part of a component such that it will not become the root of another component in the future.

The graph in Fig. 4.9 shows the 5 strongly connected components identified after running Tarjans algorithm on the initial graph in Fig. 4.8. Vertices that have the same background are part of the same component; meaning that one vertex can reach every other vertex in the component by following the directed edges. Note that some vertices make a component on their own as there is no way to reach them again in a cycle.



**Fig. 4.8:** An initial graph



**Fig. 4.9:** The components of the graph

node	index	low	stack	new component	color
1	1	1	[1]		blue
17	2	2	[1, 17]		yellow
8	3	2	[1, 17, 8]		yellow
12	4	4	[1, 17, 8, 12]		orange
2	5	4	[1, 17, 8, 12, 2]		orange
11	6	4	[1, 17, 8, 12, 2, 11]		orange
3	7	4	[1, 17, 8, 12, 2, 11, 3]		orange
			[1, 17, 8]	[3, 11, 2, 12]	
16	8	2	[1, 17, 8, 16]		yellow
7	9	2	[1, 17, 8, 16, 7]		yellow
18	10	2	[1, 17, 8, 16, 7, 18]		yellow
9	11	2	[1, 17, 8, 16, 7, 18, 9]		yellow
14	12	12	[1, 17, 8, 16, 7, 18, 9, 14]		green
6	13	12	[1, 17, 8, 16, 7, 18, 9, 14, 6]		green
13	14	12	[1, 17, 8, 16, 7, 18, 9, 14, 6, 13]		green
4	15	12	[1, 17, 8, 16, 7, 18, 9, 14, 6, 13, 4]		green
15	16	12	[1, 17, 8, 16, 7, 18, 9, 14, 6, 13, 4, 15]		green
5	17	12	[1, 17, 8, 16, 7, 18, 9, 14, 6, 13, 4, 15, 5]		green
			[1, 17, 8, 16, 7, 18, 9]	[5, 15, 4, 13, 6, 14]	
			[1]	[9, 18, 7, 16, 8, 17]	
			[]	[1]	
10	18	18	[10]		red
			[]	[10]	

**Table 4.1:** The depth-first search performed in Tarjan's algorithm

---

**Algorithm 2:** Tarjan's strongly connected components

---

```
1 void Tarjan(Graph G)
2 begin
3   Global int index  $\leftarrow$  0
4   Global int components  $\leftarrow$  0
5   Global int n  $\leftarrow$  |V(G)|
6   Global S  $\leftarrow$  new Stack()
7   Global stacked  $\leftarrow$  new boolean[n]
8   Global id  $\leftarrow$  new int[n]
9   Global low  $\leftarrow$  new int[n]
10  for u  $\in$  V(G) do
11    if  $\neg$ stacked[u] then dfs(u, G)

12 void dfs(int u, Graph G)
13 begin
14   push(u, S)
15   stacked[u]  $\leftarrow$  true
16   low[u]  $\leftarrow$  index
17   index  $\leftarrow$  index + 1
18   int min  $\leftarrow$  low[u]
19   for v  $\in$  N(u, G) do
20     if  $\neg$ stacked[v] then dfs(v, G)
21     min  $\leftarrow$  minimum(low[v], min)
22   low[u]  $\leftarrow$  minimum(low[u], min)
23   integer v
24   repeat
25     v  $\leftarrow$  pop(S)
26     id[v]  $\leftarrow$  components
27     low[v]  $\leftarrow$  n
28   until v  $\neq$  u
29   components  $\leftarrow$  components + 1
```

---

Tarjan's algorithm for finding strongly connected components is shown in algorithm 2. The *Tarjan* procedure in line 1 takes a Graph  $G$  as an argument and represents the start of the algorithm. The global integer *index* declared in line 3 stores the current order in the depth-first search, and gets incremented each time a vertex is discovered. The integer *components* in line 4 keeps track of how many components were identified in the given graph. The integer  $n$  in line 5 represents the number of vertices in the given graph.

In line 6 **TODO WE:**we introduce a Stack data structure that will hold all the vertices that are part of the same strongly connected component. Line 7 contains a declaration for the *stacked* vertex-indexed array of booleans used to keep track if a vertex is present or not on the stack  $S$ .

An integer vertex-indexed array *id* is declared in line 8 to store the id of the strongly connected component of each vertex. Line 9 declares a vertex-indexed array *low* used to store the topmost reachable vertex in the depth-first search tree through a back edge. Lines 10 and 11 contain a for loop that calls a procedure for performing depth first search on every vertex in the given graph that is not currently stacked.

Line 12 declares a recursive procedure for performing a depth-first search starting from vertex  $u$  in graph  $G$ . The procedure starts by pushing the vertex  $u$  on the stack  $S$  (line 14) and then updating the value in the *stacked* array (line 15) to reflect the change. Line 16 sets the vertex's  $v$  value for *low* to *index* as every vertex assumes to have

no ancestors until a back edge to one is found. In line 17, the value for *index* is incremented by one to keep count of the current number of discovered vertices. Line 18 declares an integer *min* used to hold information about the oldest, lowest ancestor that will be discovered later in the search and is initialized to take the same value of the current vertex.

Lines 19 to 21 loop through every vertex *v* that is a neighbour of vertex *u*. The procedure  $N(u, G)$  on line 19 returns a list of neighbouring vertices of a given vertex, not including the given vertex. If the discovered neighbour *v* is not currently on the stack *S*, then **TODO WE:**we proceed to make a recursive call to the depth-first search function on the found vertex.

After the recursive call finishes processing the sub-tree of the neighbouring vertex *v*, *min* gets updated on line 21 to store the topmost reachable vertex that could be the same, or even higher in the tree if a higher back-edge from *v* was discovered.

Line 22 updates the topmost reachable vertex of vertex *u* to reflect any changes after processing all the neighbouring vertices.

Line 23 declares an integer *v* used to store a vertex that for the repeat-until loop in lines 24 to 28. The loop begins by popping vertices out of the stack *s* on line 25. Each vertex *v* found on the stack is assigned the current component id. Line 26 assigns the last leaf node of the graph as a topmost reachable vertex of vertex *v* in order to mark it as already part of a component. The repeat-until loop in line 24 to 28 is runs until reaching the root of the component.

The depth-first search procedure finishes on line 29 by updating the total number of strongly connected components discovered.

### 4.3 Deletion of edges from the graph

The final step in the all-different algorithm uses the knowledge gained after computing a maximum matching and finding the strongly connected components in the graph. For this step, every edge that does not belong to any matching covering the vertices in the left partition of the graph will be deleted. The algorithm shown in ?? 3 is a slight adaptation of the fourth and final step in Jean-Charles Regin’s paper [7] describing the all-different constraint. The algorithm starts by computing a matching using the Ford Fulkerson algorithm. Having the match, a breadth-first search is made on the free vertices in the graph and traversed edges are marked as *used*. Tarjan’s algorithm is then used to find the strongly connected components in the graph. Each edge connecting two vertices in the same component is marked as *used*. Finally, all the edges in the graph that are *unused* and are not part of the matching will be deleted from the graph. The *unused* edges represent edges that are not part of any alternating path of even length, and therefore belong to no possible matching in the value graph of the constraint. The removed edges in the graph correspond to values getting removed from the domains of the variables, and represents the way the search spaced is pruned making the search for a solution more efficient.

18	23	23	245	456	456	279	378	23589
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**Fig. 4.10:** A Sudoku row used to illustrate the all-different algorithm

---

**Algorithm 3:** Remove edges from the value graph of a constraint of difference

---

*RemoveEdgesFromGraph*(**Graph**  $G$ )

// *RemovedEdges* is the set of edges removed from  $G$

//  $M(G)$  is a matching over  $G$  using the Ford-Fulkerson algorithm

// The function returns *RemovedEdges*

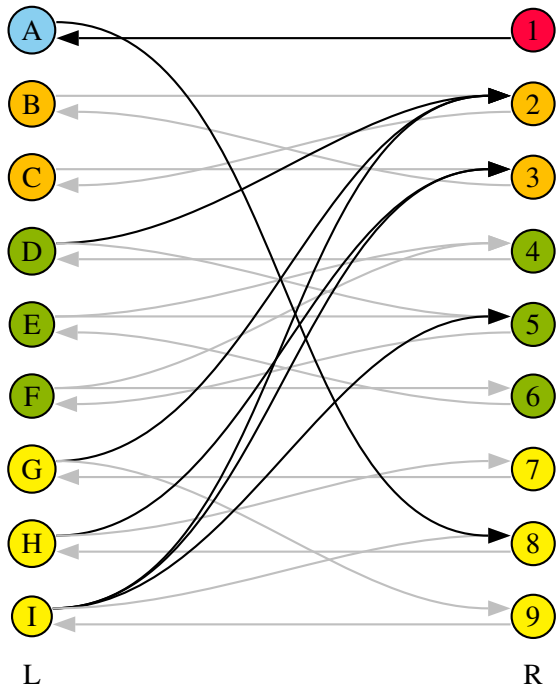
**begin**

```
1  Mark all the edges in graph  $G$  as unused
2  Initialize RemovedEdges to the empty set
3  Compute a matching over  $G$  using the FordFulkerson algorithm
4  Perform a breadth-first search starting from the free vertices and mark all visited edges as used
5  Compute the strongly connected components of graph  $G$  using Tarjan's algorithm
6  Mark each edge that connects two vertices in the same component as used
7  for each edge  $e$  in graph  $G$  that is unused do
    if  $e \in M(G)$  then
      | mark  $e$  as vital
    else
      |  $RemovedEdges \leftarrow RemovedEdges \cup \{e\}$ 
      | Remove  $e$  from  $G$ 
  return RemovedEdges
```

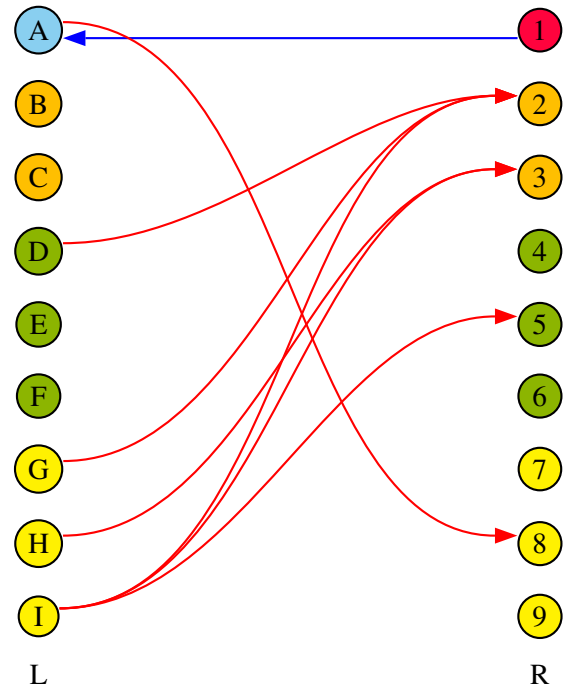
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The recently described last step of the all-different algorithm is the general implementation of the all-different constraint. The properties of the all-different constraints used in the Sudoku puzzle make the last step of the all-different algorithm much easier. It is good to notice that the chosen value graph used in this paper has 18 vertices in two partitions. The Sudoku row shown in Fig. 4.10 corresponds to the value graph that was used in this paper and was chosen as an example to illustrate the effects of running the all-different algorithm on the 9 variables involved in the constraint. The value graph of the constraint is shown in Fig. 4.11. The variables of the Sudoku row in the left  $L$  partition of the value graph and have names from A to I. Each variable has edges to vertices in the right  $R$  partition of the graph representing digits in the domain of the variable. As all the constraints of difference in the Sudoku puzzle have 9 variables taking at most 9 possible values, the two partitions of the graph will always be balanced. This property makes the last step of the all-different constraint a straightforward process in the case of the Sudoku puzzle. This alternative algorithm for the last step in the all-different constraint considers all the edges that connect two vertices from different components of the value graph. Edges of interest are highlighted in Fig. 4.11. The direction of the edges indicate two outcomes and are coloured in blue and red as shown in Fig. 4.12. Forward edges (coloured in red) indicate that the digits from the right  $R$  partition should be removed from the domain of the variables in the  $L$  partition as they will never be part of an even alternating path and therefore can't be part of a solution where the constraint holds. Backward edges (coloured in blue) indicate finding the unique assignment to a variable. In the case shown in Fig. 4.12, the first variable  $A$  of a selected Sudoku row will certainly take the value of 1 as all other values got removed from its domain.





**Fig. 4.11:** Edges connecting vertices in separate components



**Fig. 4.12:** Deletions (red) and assignments (blue) for the variables

18	23	23	245	456	456	279	378	23589
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**Fig. 4.13:** Removal of values from the domain (in red), and assignments (in blue) in a Sudoku row

The Fig. 4.13 shows the final state of the Sudoku row after the last step in the all-different algorithm. Values coloured in red correspond to edges that got removed from the value graph of the constraint. Values coloured in blue represent the unique value that is assigned to a variable as all other values from the domain got removed.

## Chapter 5

# AllDifferent Demonstration

This chapter provides a demonstration of the Java application developed to solve Sudoku puzzles using Constraint Programming. The Graphical User Interface of the application is split into two parts, the left-hand side showing a Sudoku puzzle, while the right-hand side shows the value graph of an all-different constraint. Each time a row, column or  $3 \times 3$  sub-square is selected by the user, the corresponding 9 variables are highlighted in the Sudoku puzzle. The steps in the all-different algorithm perform changes on the graph of the selected constraint of difference.

Every Constraint Satisfaction Problem (CSP) has variables with domains associated to them and constraints that should hold for a solution. For the purpose of this project, there are 81 variables representing the  $9 \times 9$  cells in the Sudoku grid, each with an initial domain of integers from 1 to 9 as shown in Fig. 5.1. The aim is to reduce the domain of all the variables to a single value, the correct value that is part of the solution. As all initial Sudoku puzzles start with a partially completed grid, values that represent clues are reflected as variables that have an initial domain of a single value, the known digit. Such variables keep their initial assignment throughout the algorithm, as no other values are present in their domain.

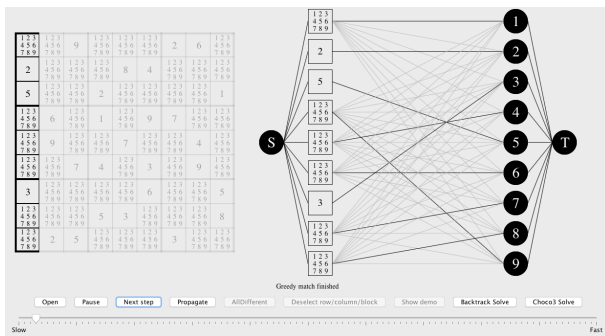
The constraints for a Sudoku puzzle can be expressed in a natural language as such: digits may not appear twice in the same row, column or  $3 \times 3$  sub-squares. The rules of the puzzle highlight the need of thinking in terms of 9 rows, 9 columns and 9 sub-squares, each of them having digits from 1 to 9 with no repetition. The CSP shown in the application has 27 all-different constraints, one for each of the 9 rows, columns and sub-squares. Each constraint contains 9 variables that should take different values. As each variable is part of a row, column and sub-square, it is therefore involved in 3 constraints of difference at the same time.

One gets to the solution by running the all-different algorithm on rows, columns and sub-squares to remove values from the domain of the variables. Initially, the known values given at the start of the puzzle will be removed from the domains of possible values in the variables that are part of the same row, column or subregion. As the algorithm progresses, it will eventually remove enough values from the domain of a variable until a single value is left, representing the correct answer for that position. Once a value is found, there is a waterfall-like effect across the puzzle as the knowledge is propagated across the grid. The propagation takes place across all the variables that are in the same row, column and sub-square with the one just solved. This keeps the 3 constraints of difference in a consistent state, as all the variables will have remaining domains not containing any already solved values. Initially, the propagation mechanism is disabled in the application such that all changes to the puzzle are made by the all-different algorithm that detects such inconsistencies and removes them. Propagation after assignments can be enabled from the Settings menu.

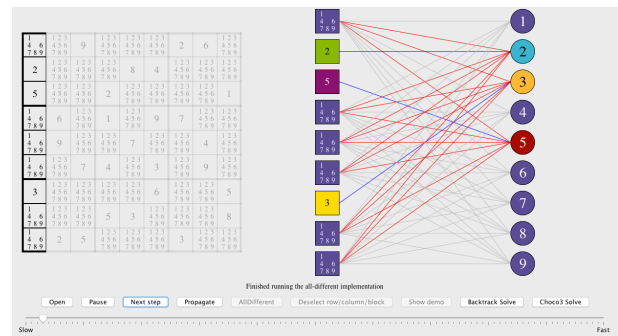
In order to test and show that the all-different algorithm is implemented correctly and performs the right decisions, a demonstration is hard coded consisting of running the all-different algorithm on a sequence of 5 rows, columns and sub-squares. The step by step demonstration can be run using the *Show demo* button and can be

paused at any time. Alternatively, the same sequence can be followed by manually selecting the rows, columns and sub-squares mentioned in Chapter B.

There are a number of operations performed by the all-different algorithm on the value graph of a selected constraint of difference. All operations on the graph are shown in a step by step fashion in the right-hand side of the screen. First, a greedy match is made on the graph to associate most of the variables with a value. The second step is to compute a maximum matching for the 9 variables in the left partition of the graph. The maximum matching is computed using the Ford Fulkerson algorithm introduced in Section 4.1. Fig. 5.1 shows the maximum matching for the constraint of difference in the 1st column of the puzzle. The third step uses information from the maximum matching to find the strongly connected components in the value graph of the constraint, by using Tarjan's algorithm introduced in Section 4.2. Fig. 5.2 shows vertices belonging to the same component having the same background colour. Finally, the edges connecting vertices that belong to different components get colored in red or blue based on their direction. Red edges connect a variable with a value that was removed from its domain during the algorithm. Blue edges highlight the single value a variable is forced to take, as it is the only option left in its domain. The state of the 1st column after running the all-different algorithm is shown in Fig. 5.2. Notice how the all-different algorithm performs changes only to the selected 9 variables inside a row, column or  $3 \times 3$  sub-square.



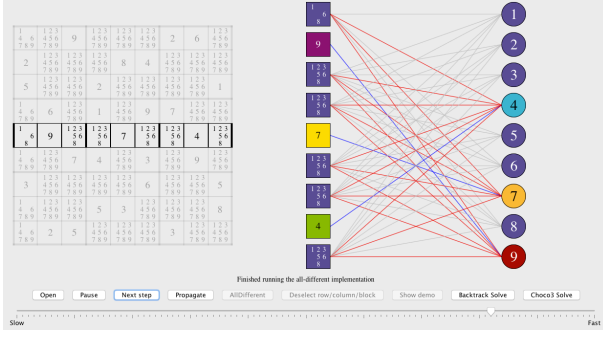
**Fig. 5.1:** 1st column after finding a maximum matching



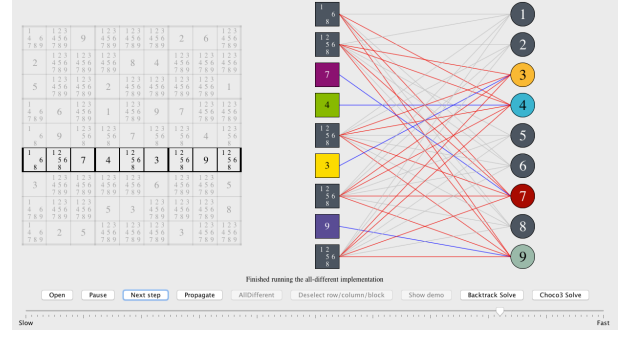
**Fig. 5.2:** 1st column after running the all-different algorithm

The same steps are followed for the 5th row (Fig. 5.3), the 6th row (Fig. 5.4) and the 4th  $3 \times 3$  sub-grid (Fig. 5.5). As seen in the figures mentioned before, all the previous runs of the all-different algorithm removed values from the domain of the variables that represented clues in the initial puzzle. Notice how in Fig. 5.5 there are two variables in the 1st column of the puzzle, each having a domain of  $\{1, 8\}$ . One doesn't know which of the two variables will take which value, but since the two values will have to be used by the two variables, this means both values should be removed from the other variables in the same constraint. This detail is spotted by the all-different algorithm that decides to remove the two values from the domains of the adjacent variables. The last selection in this demonstration calls the all-different algorithm on the 1st column of the puzzle once again. Fig. 5.6 shows the state of the 1st column after running the all-different algorithm the second time. Notice how only two variables in the 1st column have the values 1 or 8 in them, compared to last time. It is good to mention once again that the all-different algorithm doesn't make guesses, instead it uses knowledge about the domains of the variables to remove values that will never be part of a solution.

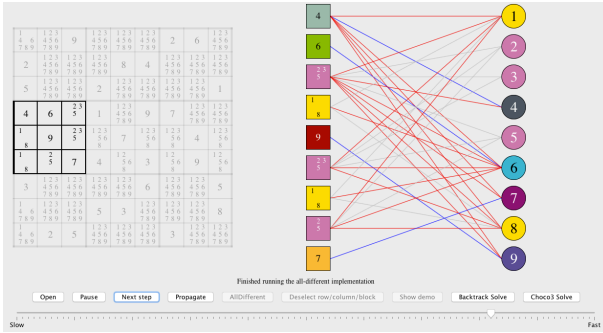
The algorithm is intelligent in a sense that it mimics the human thinking when solving a Sudoku puzzle. At some stages of the game there is enough information to know that, for example, two particular values will fill two cells, but no information about which value corresponds to which cell. The hardness of a Sudoku puzzle increases with the number of such dilemmas, and the number of variables having the same values in their domain. By using the all-different algorithm, one can ensure the progress towards a solution, despite not knowing which value goes where. It is good to note that although humans may choose a guessing approach to continue towards a solution, the all-different algorithm performs no guessing and works only with the information about the current state of the puzzle.



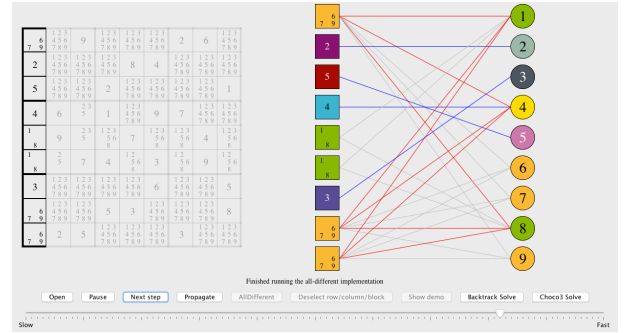
**Fig. 5.3:** 5th row after running the all-different algorithm



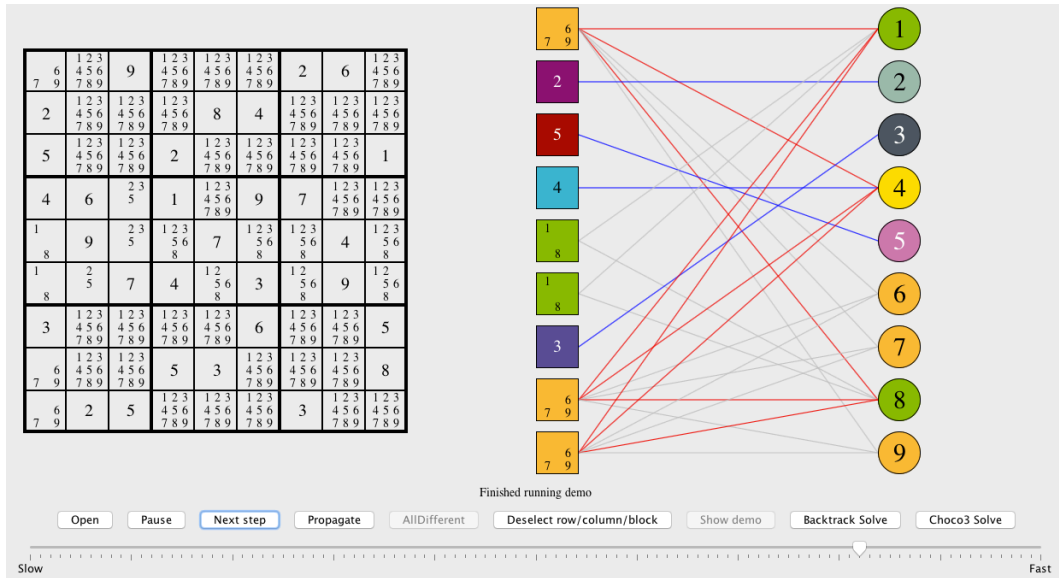
**Fig. 5.4:** 6th row after running the all-different algorithm



**Fig. 5.5:** 4th  $3 \times 3$  sub-grid after running the all-different algorithm



**Fig. 5.6:** 1st column after running the all-different algorithm



**Fig. 5.7:** State of the program after running the demo

The end result of following the 5 steps in the demonstration can be seen in Fig. 5.7. In the 1st column there are two cells with a domain of 1, 8. Although, one doesn't know yet which cell takes which value, the human and the all-different algorithm is sure that the digits 1, 8 will be distributed to the two particular cells. The last step illustrates the all-different algorithm decision to remove the values 1, 8 from the domains of the rest of the cells in the selection.

## Chapter 6

# Implementation

One could create 36 not equals constraints between the 9 variables of each row, column, subregions times 27 such regions for a total of 972 constraints between two variables. This would be highly inefficient, hence the need for a better approach.

The Graphical User Interface (GUI) is inside a `JLayeredPane` which is a special swing component that allows components inside it to overlap according to their specified depth. In my implementation, the backmost layer contains a `JPanel` responsible for the drawn edges in the graph. The layer above contains the rest of the GUI represented by the Sudoku puzzle on the left hand side of the screen, and the vertices of the graph on the right hand side of the screen. By using a layered panel, the edges of the graph can be drawn behind the vertices.

The Sudoku Grid present on the left hand side of the screen is made up of 9 by 9 Sudoku Cells positioned at specific locations. When a new Sudoku Cell instance is made, its *i*th row and *j*th column number are passed as arguments to the constructor. Using the information related to its location in the grid, it is now possible to render the border of each cell. Initially all the top, left, bottom and right borders have a width of 1px. To represent the 3 by 3 sub-grids, specific borders are set to have a width of 2px. In the end, the 9 by 9 grid is given an overall thick border by specifying a width of 4px to the cells that are in the first and last columns and rows. All Sudoku Cells contain a `JLabel` with text retrieved from the Model denoting which possible values are still available for that particular cell.

Not visible to the human eye, a new Sudoku grid instance is drawn on top of the existing one, this time with all borders set to a width of 1px. The Sudoku Cells inside this second grid contain the same information about the state of the puzzle retrieved from the Model. Additional information such as a custom background will be applied to these cells as the algorithm progresses. Cells inside the second grid become visible once the user makes a selection. Once visible, the cells inside the selection are animated into their position inside the graph on the right hand side of the screen. When the user performs deselection, the nice cells are animated back into their original position in the grid and become invisible again.

Particular attention was given to make the software thread safe. It is crucial that the application logic doesn't make the Graphical User Interface unresponsive. To achieve this, the program will run on two threads, one main thread responsible with all the processing need to be done by the program, and the Event Dispatching Thread.

The Event Dispatching Thread (EDT) is a background thread used to invoke Swing methods that change the GUI and listen to events associated to components. Most Swing components are not thread safe, therefore bugs such as race conditions could unexpectedly arise if it was to perform the actions from normal threads. The EDT acts as a queue of events that are performed sequentially.

Throughout the application, Timer objects are used to schedule repeated actions such as the fading out of the

components or movement of the cells. The model contains a two-dimensional ArrayList of Timer objects that acts as a queue of animations. Timers on the same level start running at the same time until they all finish. Once all the timers on a level finish, the next level in the two-dimensional ArrayList is started, allowing us to queue a movement of a row after all the cells finish fading out.

## Chapter 7

# Conclusion

### 7.1 Summary

The aim of this project was to develop a Java application to be used during the Constraint Programming lectures to introduce students to the all-different algorithm. This paper started by introducing general concepts from Constraint Programming, a programming paradigm that originated from Artificial Intelligence. A chapter is dedicated to reviewing Jean-Charles Regin's paper [7] from 1994 describing the all-different constraint. The algorithm is a sequence of steps that perform operations such as finding a maximum matching and the strongly connected components in the value graph of a constraint of difference. The Ford-Fulkerson algorithm was used in this project to compute the maximum matching, and Tarjan's algorithm to find the strongly connected components in the graph. The Sudoku puzzle was chosen to illustrate the steps in the all-different algorithm. A Java program was developed to model the problem and provide a Graphical User Interface showing the puzzle and the value graphs of the all-different constraints. A description of the algorithms used and how the project was implemented in Java is provided in the paper.

### 7.2 Future Work

The program developed in this project implements a model for the popular Sudoku puzzles with a  $9 \times 9$  grid. There are multiple techniques and algorithms to solve such a puzzle in approximately the same time using a regular CPU. It is therefore difficult to accurately measure the performance of the algorithms since searching for a solution takes hundreds of milliseconds. The benefits of using Constraint Programming to solve such problems, using techniques such as the all-different algorithm presented in this paper, starts to become observable when trying to solve a bigger Sudoku grid, such as one of size  $36 \times 36$ . The performance in such a case is a second compared to more than two days when using less efficient algorithms. Future work could develop models and a Graphical User Interface for general  $n^2 \times n^2$  sized boards, along with a way to compare the performance of different algorithms.

Another interesting avenue for future work is to implement and make use of different heuristics that may prove to increase the performance of the program when trying to find a solution. The project uses human judgements as its heuristic, as one has to select which row, column or  $3 \times 3$  sub-grid to solve next. An example of such a heuristic is one that will check which selection has variables with many values in their domains, as there may have values that are more likely to fail by breaking a constraint.

### 7.3 Personal Reflection

As a Joint Honours student which studies Computing Science and Business Management, I have come to appreciate the importance of an efficient algorithm over an expensive hardware upgrade. Powerful algorithms such as the one in this paper can offer a great competitive advantage in business environments that deal with hard combinatorial problems on a daily basis. Such an example would be the scheduling of the deliveries that need to be made the next day, where an optimal solution should be found in a matter of hours. In the end I would like to quote Robert Tarjan, a Turing Award winner, on the characteristics of a good algorithm: “It is one thing to have an algorithm that is theoretically good. It is another to have an algorithm that is simple enough that someone would want to program it and use it in practice.” [11].

### 7.4 Acknowledgements

I would like to thank my supervisor, Dr. Patrick Prosser, and Ciaran McCreesh, for all their guidance and feedback provided throughout this project. [2] [4] [8] [9] [6] TODO



# **Appendices**

## Appendix A

# Running the Program

Running the program from the command line is done after compiling the source code as follows:

```
> javac *.java  
> java Sudoku
```

This will open the application loaded with the Sudoku puzzle */puzzles/lockedset.txt*.

Detailed debug output in the console can be activated using:

```
> java Sudoku debug
```

NOTE that the Choco3 library found in *lib/choco-solver-3.3.1-with-dependencies.jar* has to be added to the CLASSPATH environment variable as follows:

```
> export CLASSPATH=$CLASSPATH:/path/to/the/lib/choco-solver-3.3.1-  
with-dependencies.jar
```

## Appendix B

### Proof of concept

The following sequence of steps will provide a visual proof of the all-different algorithm. The following figures capture the user running the all-different algorithm on 5 predetermined rows, columns or  $3 \times 3$  sub-grids. For demonstration purposes, the Sudoku instance used is */puzzles/lockedset.txt*.

The sequence is as follows:

```
> Open lockedset.txt Sudoku instance
>
> Select the 1st column
> Run the all-different algorithm
> Deselect the column
>
> Select the 5th row
> Run the all-different algorithm
> Deselect the row
>
> Select the 6th row
> Run the all-different algorithm
> Deselect the row
>
> Select the 4th 3x3 sub-grid
> Run the all-different algorithm
> Deselect the sub-grid
>
> Select the 1st column
> Run the all-different algorithm
> Deselect the 1st column
```

Alternatively, the user can press the *ShowDemo* button to run the same steps.

Detailed description about why the last state of the program is a proof that the algorithm works as expected is provided in Chapter 5.

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