

University of Padua

Information Engineering Department Master's Degree in Computer Engineering

Solutions to the Travelling Salesman Problem

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Chapter 1

Introduction

In this report I am going to describe, analyze and implement solutions for the Travelling Salesman Problem (hereafter referred to as TSP).

The TSP is a NP-hard problem formulated as follows:

Given a list of cities and the distances between them, find the shortest path that connects all the cities and return to the origin one.

It is used mainly in plain logistics, planning, and as a benchmark for testing optimization problems. However, it can be helpful in many other areas with a slight modification of the formula - such as in DNA sequencing, considering cities as DNA fragments, and astronomy, where they are stars.

Even though it is an NP-hard problem, instances with the dimensions of thousand or even millions of cities can be solved with great precision (around 1%) thanks to many heuristics and exact algorithms.

1.1 Brief history of the problem

The German handbook *Der Handlungsreisende* from 1832 was a guide used by salesman traveling through Germany and Switzerland. Albeit without any mathematical languages, it proves that people were starting to realize that optimal paths could save time and so it can be seen as the first example of TSP. The first TSP mathematical formula was made by Hmail and Kirkman in the XIX century, but it was in the 1930s that the TSP was implemented - mainly in Vienna and at Harvard University. An important leap forward was made in the 1950s, when G. Dantzig, D. R. Fulkerson, and S. M. Johnson expressed the problem as an integer linear program, even if they did not propose an algorithmic solution. They were able to devise the cutting plane method, and solved an instance with 49 nodes - by constructing a tour and proving that no other tour could be shorter. In the 1980s, Grötschel, Padberg, Rinaldi and others figured out instances with up to 2392 nodes, using both cutting planes and branch and bound. In 1991, Gerhard Reinelt published

the TSPLIB, a collection of benchmark instances of varying difficulty - which has been used for comparing results among many research groups. In the 1990s Applegate, Bixby, Chvátal, and Cook developed the Concorde TSP solver. Nowadays, this program can run even on mobile devices such as iPads and it has been used in many recent record solutions: in 2006, Cook and others computed the optimal tour for an instance of 85900 nodes given by a microchip layout problem and this is currently the largest solved TSPLIB instance. For many other instances with millions of cities, today's solutions are guaranteed to be within 2-3% of an optimal tour.

1.2 The mathematical formulation

In this introduction, I used a natural way to describe the problem using the notion of cities and distance travelled. But applying from a mathematical point of view, the TSP problem can be thought as graph G = (V, E) where $V = \{v_1, \ldots, v_n\}$ are the cities described in this introduction (from now on called nodes or vertices of the graph), and E in this case represents the path that connects one node to another and it can be described as $E \subseteq (V \times V)/\{\{i, j\} : i \in V\}\}$, is the set of edges of the graph.

nother fundamental aspects to be taken in consideration are the concept of distance and its mathematical counterpart, the cost. I assign to each edge a real number that will be used to give weight (or cost) to the path chosen.

Now that I have introduced the main concepts I can explain the TSP problem from a mathematical point of view:

Given a list of nodes and the distances between them, find the shortest hamiltonian circuit.

One of the most important formulations is the following one [1], even if it presents slight modifications:

$$x_{ij} = \begin{cases} 1 & \text{if the arc } (i,j) \in A \text{ is chosen in the optimal solution} \\ 0 & \text{otherwise} \end{cases}$$
 (1.1)

$$\min \sum_{(i,j)\in A} c_{ij} x_{ij} \tag{1.2a}$$

$$\sum_{(i,j)\delta^{-}(j)} x_{ij} = 1, \quad j \in V$$
(1.2b)

$$\sum_{(i,j)\delta^+(j)} x_{ij} = 1, \quad i \in V$$
(1.2c)

$$\sum_{e \in E_G(S)} x_e \le |S| - 1, \quad \forall S \subset V \quad , |S| \ge 2$$
(1.2d)

$$x_{ij} \ge 0 \text{ intero}, \quad (i,j) \in A$$
 (1.2e)

This model is not fully functional since it allows the presence of sub-tours in the solution. In order to avoid them, I am going to add some constraints to the model in the next sections.

Chapter 2

Compact models

In this section, I am going to explore the first method used to solve the TSP problem. In particular, I will describe the Miller, Tucker, and Zemlin model (known as MTZ model) and the Gavish and Graves model (known as GG model).

2.1 Basic model

The first model I took in consideration is a lightly modified version of the one presented in section 1.2. This formula still doesn't adopt the Sub-tour Elimination Constraint (SEC) since it is intended to be used jointly with other SECs and with other methods such as matheuristics.

This configuration gives an undirected complete graph G = (V, E). The formulation used in the code is the following:

$$\min \sum_{(i,j)\in A} c_{ij} x_{ij} \tag{2.1a}$$

$$\sum_{(i,j)\delta^{-}(j)} x_{ij} = 1, \quad j \in V$$
(2.1b)

$$\sum_{(i,j)\delta^+(j)} x_{ij} = 1, \quad i \in V$$
(2.1c)

$$x_{ij} \ge 0 \text{ intero}, \quad (i,j) \in A$$
 (2.1e)

It is the same model used in section 1.2 but in this case 1.2d is not included. In this implementation, the graph is symmetric. The arcs (i, j) and (j, i) have the same weight and then thus are represented with the same edge. In this way, the total number of variables are reduced too, considering the starting point not of n^2 , but only $\frac{n(n-1)}{2}$. The result using this model can be seen in figure 2.1.

Since sub-tours are present, this solution cannot be accepted. In the next sections SEC constraints will be added to the formula for avoiding this problem.

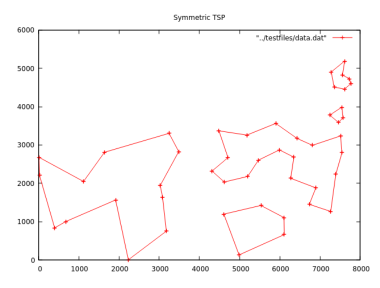


Figure 2.1: The image represent att48.tsp solved with the problem formulation showed in section 2.1

2.2 The Miller, Tucker, and Zemlin model

As said before, the basic model can create loops in the optimal solution A first constraint introduced to avoid this situation was described by Dantzig, Fulkerson, and Johnson. The constraint added was:

$$\sum_{i \in S} \sum_{j \in S} x_{ij} \le |S| - 1; \quad \forall S \subset V : \{1\} \notin S, \ |S| \ge 2, \ i \ne j$$
 (2.2)

This constraint limits the number of edges in the solution, so that no cycles are allowed in the result. Nonetheless, the usage of this method is infeasible even with a low number of nodes since the amount of constraints needed is exponential, $(O(n^2))$.

The model produced by Miller, Tucker, and Zemlin bypasses the exponential SECs by reducing their number to a simple polynomial. Differently from the basic model, the graph used in this case is asymmetrical: x_{ij} and x_{ji} can have different weights.

In the new formulation, a new variable called u_i is assigned to each node of the solution. This number represents an increasing sequence number in the optimal tour: starting from the second one $(u_2 = 0)$, each node will increase the value by 1 at each following node until we reach the end of the solution $(u_n = n - 2)$. The first node is considered special and its value is always set to 0.

This is the new constraint added to the basic model by MTZ:

$$u_i - u_j + nx_{ij} \le n - 1; \quad i, j \in \{2, \dots, n\}, i \ne j$$
 (2.3)

$$0 \le u_i \le n - 2; \quad integer \quad i \in V : i > 1 \tag{2.4}$$

The meaning of this formulation is the following: if the arc x_{ij} is selected, then the value of u_j is $u_j \ge u_i + 1$.

2.2.1 Implementation of the model

To express 2.3 as a CPLEX constraint I need to rewrite (Check) the inequation in a way called Big-M. This method expects a new variable called M, that allows the system to activate or deactivate the constraint in a simple way. The new constraint will be:

$$u_j \ge u_i + 1 - M(1 - x_{ij}) \tag{2.5}$$

With this approach, the constraint is strictly depending on the value of x_{ij} . If $x_{ij} = 1$ the constraint works like a normal one because the value of $M(1 - x_{ij})$ will be 0, so the meaning of 2.5 will be the same as the one expressed in the previous section. If $x_{ij} = 0$ the right-hand side of the inequation will be certainly negative, making the constraint deactivated and allowing u_j to take up any possible value from 0 to n-2. Between all the values that M can assume, the smallest one is surely n-1 due to the fact that, in the case of $X_{ij} = 0$, the constraint will be still useful even if u_i will reach his case limit of n-2.

Applying the Big-M trick, the constraints can be written in the CPLEX environment. There are substantially 3 methods that can be implemented in the framework:

- the use of standard constraints: all the constraints wrote in 2.3 are directly saved into the problem at once. This lead to have $O(n^2)$ constraints active, making the optimization too large or even too expensive to solve;
- the use lazy constraints: as the name suggests, here the constraints are applied lazily. This means they are not always applied to the problem because CPLEX uses them only when necessary or not before needed. In doing so, a pool of constraints is created and every time an integer optimal solution is found, the violation of every constraint in the pool is checked. If one of them is infringed, it is added permanently to the instance. This method will hopefully make the problem smaller and faster to solve than the one created with the standard constraints;
- the use of indicator constraints: in the first two cases the Big-M trick is used to trigger a constraint when a particular variable assumes a predetermined value, but this method (Big-M) is not always preferable since it can behave in unstable ways. That is why a good implementation of 2.3 is the usage of the indicator constraints provided by the CPLEX API: this method automatically activates the constraint $u_j \geq u_i + 1$ when the x_{ij} assumes the user passed value.

2.3 The Gavish and Graves model

The second compact model implemented is the one proposed by Gavish and Graves (this model will be called GG from now on), based on the single commodity flow: the arcs are considered as pipes.

Like in the MTZ model, here too a new variable is introduced in the problem. It is called "Flow of the arc" and it is represented with the symbol y_{ij} . Compared with the model in section 2.2, the starting value of y_{ij} is n-1 and decrease by 1 at each following node in the optimal solution. To implement this proposition the following constraints are added to the instance:

$$y_{ij} \le (n-1)x_{ij} \tag{2.6}$$

$$\sum_{j \in V; j \neq 1} y_{1j} = n - 1 \tag{2.7}$$

$$\sum_{i,j \in V; i \neq j} y_{ij} - \sum_{j,k \in V; j \neq k} y_{jk} = 1$$
(2.8)

This formula finds an optimal solution without sub-tours. (vedere se aggiungere qualcos'altro ma non credo)

Chapter 3

Other sub-tour elimination methods

In chapter 2 I described models that cut out any possibility to have some sub-tours in the optimal solution. In this section I use particular techniques to remove the tours in a faster way in relation to the one in the previous chapter.

3.1 Benders method

This method is the simplest one presented in this chapter. The basic idea is the insertion of the SECs only when sub-tours are found. This way is quite different from the implementation of the lazy constraints of MTZ seen before: the constraints are not activated when one of them is violated, but they are manually added into the instance.

This method uses the basic model described in 2.1 and follows this algorithm:

Algorithm 1 Benders

```
Input: G = (V, E), c : E \to \Re^+
Output: z^* optimal solution
 1: instance \leftarrow *initializing basic model*
 2: successors, component← *initialize arrays*
 3: ncomp \leftarrow 99999
 4: while ncomp > 1 do
        z^* \leftarrow \text{CPXMIPOPT}
        successors, component, ncomp \leftarrow BUILD_SOLUTION(z^*)
 6:
        if ncomp > 1 then
 7:
           instance \leftarrow add violated constraints
 8:
        end if
 9:
10: end while
11: return z^*
```

This algorithm shows the simplicity of the Benders method. The first thing to do is building the basic model inside the instance of the CPLEX environment, afterwards the arrays that will contain the successors and the components are initialized. These two arrays are big enough to hold all the nodes, so they are n long. The most important

3.2 Callback method 9

function in this algorithm, excluded the optimization (CPXMIPOPT), is absolutely the function BUILD_SOLUTION, it fills the arrays successors and components following this idea: searches all the connected components inside the solution, then each successors[i] will contain the next node inside the connected component, and component[i] will enclose the index of the component of the node i.

Then the algorithm will check the number of the connected components, saved into ncomp, and if are present more than one loops it inserts into the instance a new constraint, for each component, following this formula:

$$\sum_{e(S)} x_e \le |S| - 1; \quad \forall S \subset \{2, \dots, n\}, \ |S| \ge 2$$
 (3.1)

where x_e are the edges contained inside the connected component S.

3.2 Callback method

In the previous section, the Benders method reaches the optimal solution of the problem through multiple calls of the CPLEX optimizer adding the sub-tour elimination constraints if the solution found is composed of various tours. The purpose of this section is to introduce a different approach than the earlier one: I will exploit the branch-and-cut technique through the use of the CPLEX callbacks.

The API provided by IBM's software grant the use of some callbacks during the optimization process. CPLEX provides a wide range of possibilities such as informational callbacks, query/diagnostic callbacks, and control callbacks. The first ones give the user additional information on the current optimization without affecting the performance or interfering with the solution search space. The second one access to more detailed information compared to the informational callbacks but can affect the overall performance of the problem resolution; the query/diagnostic callbacks are also incompatible with the dynamic search and deterministic parallel functions. The last ones are the one I am going to use and they allow the user to alter and customize how CPLEX performs the optimization.

During the optimization process, CPLEX will find numerous possible solutions. Suppose that during operation a candidate solution x^* is found. Then if the cost of this new result is better than the anterior one the software will update the current best solution with the last found. Otherwise, the candidate is considered infeasible and is rejected by the system.

The CPXCALLBACKSETFUNC method will set my custom callback that will be used every time a candidate solution is found. The algorithm adopt the same method explained in section 3.1 (BUILD_SOLUTION). In fact inside the callback if more than one connected component is found a new SEC is added to the instance of the problem and the candidate

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solution is rejected (through the function CPXCALLBACKREJECTCANDIDATE).

The real difference between this implementation of the branch-and-cut and the Benders method is that while the last one needs the iteration of various optimizations processes this one rejects the possible solution aforehand it is provided to the user. The algorithm used is the following:

Algorithm 2 Callback method

```
Input: G = (V, E), c : E \to \Re^+
Output: z^* optimal solution
 1: procedure MAIN(G = (V, E), c : E \to \Re^+)
       instance \leftarrow *initializing basic model*
 2:
       instance \leftarrow *instance \cup custom callback*
 3:
       z^* \leftarrow \text{CPXMIPOPT}
 4:
       return z^*
 5:
 6: end procedure
 7: procedure CUSTOMCALLBACK(z)
       ncomp \leftarrow BUILD\_SOLUTION(z)
 8:
       if ncomp > 1 then
 9:
           *Add SEC and reject candidate solution*
10:
       end if
11:
       return
12:
13: end procedure
```

3.2.1 Callback on the fractional solutions

In the previous section, the callback was called on each integer candidate solution. This sector will describe the use of the callback even on the fractional solution.

The object of this idea is to save computational time by adding new SECs that are probably common in the decision tree. The procedure defined in this phase is the same in section 3.2, except that this callback is applied in the continuous relaxation of the problem. To implement this kind of operation I used an external library called Concorde [2]. This library is written in ANSI C and is one of the most powerful tool on the market: it can solve really large instances of the TSP problem to the optimal solution.

When the callback is called I use the library mentioned above to compute which SEC I require to exclude the fractional solution from the decision tree.

Chapter 4

Non optimal solutions

In this section I am going to explore a new branch of the TSP resolution. Now the discovery of an optimal solution is no longer important. This can be achieved through the use of the matheuristic and heuristic to find a solution, with a great approximation to the optimal one, even with instances including millions of nodes. In this chapter I am going to cover different approach such as: matheuristic methods, heuristic and metaheuristic algoritms.

4.1 Matheuristics

As aforementioned in this chapter I am going to explore how big instances can be solved when the classical methods, such as MTZ and GG, would take too long to be solved to the optimal value. This section will explore the concept of matheuristic that combine mathematical programming and heuristics techniques. "The objective of a heuristic is to produce a solution in a reasonable time frame that is good enough for solving the problem at hand. This solution may not be the best of all the solutions to this problem, or it may simply approximate the exact solution. But it is still valuable because finding it does not require a prohibitively long time" [3].

The two techniques that I will explore are the hard-fixing and the soft-fixing.

4.1.1 Hard-fixing

The main purpose of this method is to try to reduce the search space by cutting down the complexity of the optimization. For achieving this an initial feasible solution is needed, obtained by any approach described in this report.

The central thought of hard-fixing is to obtain an easier problem by previously setting some variables of the solution passed to the method - and so by having fewer variables to compute. The variables that are going to be fixed are x_{ij} , $i, j \in V$. This task is done by settling the values to 1, so the edge will be surely part of the solution. A valid method

4.1 Matheuristics

to choose which path is chosen to be fixed is to link each edge to a probability 0 , then setting randomly - with the probability picked - the value to 1.

Hereupon the problem will have presumably p * |E| variables fixed and the instance can be solved with a lower effort, this will bring to a hopefully better solution. Although the choice of using a probability is a good pick, any other method can be used to block the edges.

The performance of this technique is strictly related to the choice of the first feasible solution, if it is not good enough this approach will get stuck on some non-optimal solution. A good fix is to lower down p whenever the solution is not improved for a predetermined amount of time.

Algorithm 3 Hard-fixing

```
Input: G = (V, E), c : E \to \Re^+, qlobal\_timelimit, iteration\_timelimit
Output: z hopefully good solution
 1: instance \leftarrow *initializing model (any of this report)*
 2: z \leftarrow \text{CPXMIPOPT} *with nodelimit 0*
 3: p \leftarrow 0.9
 4: i \leftarrow 0
 5: while time_elapsed < global_time_limit do
        if time_remaining > iteration_timelimit then
 7:
            instance \leftarrow *set timelimit to iteration_timelimit*
        else
 8:
            instance \leftarrow *set timelimit to time\_remaining*
 9:
10:
        end if
        instance \leftarrow *hard-fixing with probability p^*
11:
        z_{new} \leftarrow \text{CPXMIPOPT}
12:
        if cost(z_{new}) < cost(z) then
13:
14:
            z \leftarrow z_{new}
15:
            i \leftarrow 0
16:
        else
            i \leftarrow i + 1
17:
        end if
18:
        if i = 10 then
19:
20:
            p \leftarrow p - 0.1
21:
        end if
22:
        instance \leftarrow *remove hard-fixing*
23: end while
24: return z
```

In this algorithm, there are some variables never mentioned before: the global timelimit and the iteration timelimit. As aforementioned the hard-fixing is a technique that aims to obtain a good solution in a short amount of time. So the parameters described above are necessary to establish - as the name suggests - the timespan in which the optimization is solved: global_timelimit is the total time reserved to the solver, interation_timelimit is the duration of each optimization in which the hard-fixing is applied. 4.1 Matheuristics

In the algorithm, the initial solution is provided by the solver, limited in the depth of its analysis. Then algorithm 3 starts to iterate the main process until the global_timelimit is reached. During this phase the optimization is called several times, in each of which the instance is solved blocking some variables to 1.

4.1.2 Soft-fixing

The technique described is called Local Branching. Since it is a slight modification of the hard-fixing it is also called soft-fixing.

In section 4.1.1 the value of the variables is fixed in a manual way using a probability system. In local branching this operation is performed by adding a new constraint that forces the instance itself to block a predetermined number of variables, giving a degree of freedom to the solver on which variable to fix and which not.

The main idea of soft-fixing is the Hamming distance: "it measures the minimum number of substitutions required to change one string into the other" [4]. This description can be applied also to vectors. So given two vectors x and \tilde{x} in $\{0,1\}$ the hamming distance is the number of different bits they have. It can be described in this way:

$$H(x,\tilde{x}) = \sum_{j:\tilde{x}_j=1} (1 - x_j) + \sum_{j:\tilde{x}_j=0} x_j$$
 (4.1)

Considering that the output solution of the solver is a vector in $\{0,1\}$ it is possible to insert into the instance a new constraint that limits the hamming distance between the old solution and the new one. Since the number of edges active in each solution is forced to be n = |V| and therefore the Hamming distance is computed one the differences in the bits equal to 1, it is possible to reduce 4.1 to:

$$H(x,\tilde{x}) = \sum_{j:\tilde{x}_j=1} (1 - x_j) = n - \sum_{j:\tilde{x}_j=1} x_j$$
 (4.2)

I recall that the purpose of this method is to limit the diversity of two solutions it is possible to add a new variable k. This will be the value that will limit the Hamming distance. Through elementary math this formulation is built:

$$H(x, \tilde{x}) \le k \Rightarrow n - \sum_{j: \tilde{x}_j = 1} x_j \le k \Rightarrow \sum_{j: \tilde{x}_j = 1} x_j \ge n - k$$
 (4.3)

The effect of 4.3 is to narrow the next iteration of the optimization to a k-neighborhood of the previous one. As the method described in section 4.1.1 the value of k can vary if for a predetermined number of times the optimization doesn't improve the value.

As it is evident the algorithm 3 and 4 are almost identical. The unique change is upon the variable used and the constraint added to the instance. In this implementation, the 4.1 Matheuristics

Algorithm 4 Soft-fixing

```
Input: G = (V, E), c : E \to \Re^+, global\_timelimit, iteration\_timelimit
Output: z hopefully good solution
 1: instance \leftarrow *initializing model (any of this report)*
 2: z \leftarrow \text{CPXMIPOPT} *with nodelimit 0*
 3: k \leftarrow 2
 4: i \leftarrow 0
 5: while time\_elapsed < global\_time\_limit do
        if time_remaining > iteration_timelimit then
             instance \leftarrow *set timelimit to iteration_timelimit*
 7:
 8:
        else
             instance \leftarrow *set timelimit to time\_remaining*
 9:
10:
        instance \leftarrow *soft-fixing using a k-neighborhood*
11:
        z_{new} \leftarrow \text{CPXMIPOPT}
12:
        if cost(z_{new}) < cost(z) then
13:
14:
             z \leftarrow z_{new}
15:
            i \leftarrow 0
16:
        else
             i \leftarrow i + 1
17:
        end if
18:
        if i = 10 then
19:
             k \leftarrow k + 1
20:
        end if
21:
        instance \leftarrow *remove soft-fixing*
23: end while
24: return z
```

value of k is implemented by 1 every time the solution is not improved for 10 times.

4.2 Heuristics

In this section the concept of heuristic is explored in some of algorithms used to solve the TSP problem. In particular a set of methods will be presented such as greedy algorithm, extra mileage, and 2-opt optimization.

4.2.1 Greedy algorithm

This type of method is the easiest to understand and implement. This heuristic has its base on the concept to construct the shortest path by connecting the nearest nodes.

This algorithm starts from an arbitrary node and chooses its following node by selecting the nearest one from the ones that are not already in the path. This greedy approach is also called Nearest Neighbor.

The main side effect it has is that the shortest path is easily missed since the last nodes - apart from special cases - are far from each other. In this way, the optimal solution is surely not selected. The algorithm is the following:

Algorithm 5 Greedy

```
Input: G = (V, E), c : E \to \Re^+
Output: z hopefully good solution
 1: best\_cost \leftarrow +\infty
 2: z \leftarrow \text{*empty*}
 3: for start\_node \leftarrow 1 to n do
        current\_node \leftarrow start\_node
 4:
         while *each node is visited* do
 5:
             candidate\_node \leftarrow -1
 6:
             for i \leftarrow 1 to n do
 7:
                 *Finds the nearest node and marks it as visited*
 8:
                 candidate\_node \leftarrow *nearest node*
 9:
            end for
10:
             *Saves candidate_node as next node*
11:
             current\_node \leftarrow candidate\_node
12:
         end while
13:
        if cost(z_{curr}) < best\_cost then
14:
             best\_cost \leftarrow cost(z_{curr})
15:
16:
             z \leftarrow z_{curr}
17:
        end if
18: end for
19: return z
```

This algorithm describes exactly the behavior of the greedy algorithm. In my implementation, each node is tested as starting node to obtain the best nearest neighbor

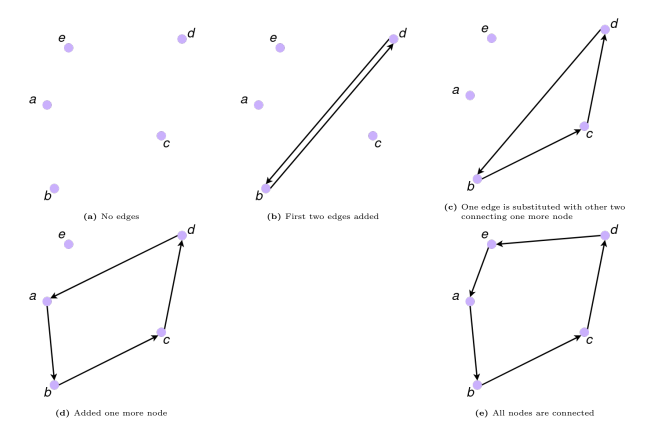


Figure 4.1: In this image we can see the process done by extra-mileage to find the solution.

solution. The classic implementation has a complexity of $O(n^2)$ since each node has to search the nearest node all over the graph. In particular 5 has a complexity of $O(n^3)$ because of the aforementioned method to select the best starting node.

This complexity is quite low but with some instances with a great number of nodes can be slower than expected.

4.2.2 Extra mileage approach

This approach is more complex than the previous one but in favor of a better solution. The main idea behind this algorithm is very simple and it starts from an empty solution and provides a good solution.

It starts with connecting with a cycle the farthest nodes in the instance, this will be the beginning of the solution. Then the closest node among the active edges is selected and is added to the solution. This action is performed by replacing the edge with two more that allow the new node to enter the solution.

The process can be seen in figure 4.1 where it is performed with a bunch of nodes.

The method with the next node to be inserted is chosen is the triangle inequality. Actually, with the replacing phase, some cost is added to the solution. Imagine taking into consideration three nodes (x, y, z), where x and y are already in the solution and z

wants to join them. The mathematical operation to use is the following:

$$\Delta(x, y, w) = c_{xw} + c_{yw} - c_{xy} \tag{4.4}$$

This value will always be positive thanks to the aforementioned triangle inequality which states that the sum of the lengths of any two sides must be greater than or equal to the length of the remaining side.

The algorithm used is the following:

```
Algorithm 6 Extra mileage
```

```
Input: G = (V, E), c : E \to \Re^+
Output: z hopefully good solution

1: x, y \leftarrow \text{*finds the farthest nodes*}

2: z \leftarrow \text{Add}(x, y)

3: while |z| < n do

4: (x, y, w) \leftarrow \operatorname{argmin}(\Delta(x, y, w) : x, y \in z, w \notin z)

5: z \leftarrow \text{Add}(w)

6: end while

7: return z
```

4.2.3 2-opt refining

This section will analyze one refining technique. The target of this approach is to take an existing solution x and try to take it closer to the optimum one.

A method to do so is represented by the k-opt refining that consists on rearrange k edges in order to obtain a new solution with a lower cost. In particular, this section it is describes the 2-opt refining.

To apply this approach effectively it is applied starting from a solution obtained with a heuristic approach such as the ones described in section 4.2.1 and 4.2.2. The idea of this algorithm is to remove all the crossing edges and insert new edges that will decrease the cost of the final solution.

To do so it is used the triangle inequality, is used to find the couple of edges that allow the best improvement of the solution. In this implementation the operation is done across four nodes: i, succ[i], j, and succ[j]. Since the tour is asymmetric it has a direction, succ[i] and succ[j] are respectively the following nodes in the path of the nodes i and j.

$$\Delta(i,j) = (c_{i,succ[i]} + c_{j,succ[j]}) - (c_{i,j} + c_{succ[i],succ[j]}) \quad i,j \in V; i \neq succ[j]; j \neq succ[i] \quad (4.5)$$

In 4.5 in is described the correct way to compute the delta of the method. Greater the value, better the improving.

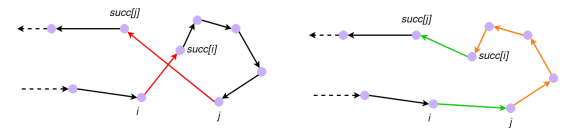


Figure 4.2: The image represent the replacement of two edges during the 2-opt refining.

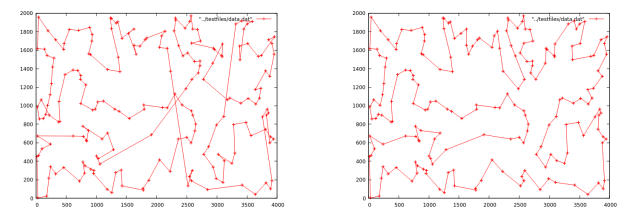


Figure 4.3: The improvement obtained applying to a greedy algorithm (left) the 2-opt refining approach (right).

In figure 4.2 it is possible to notice the replacement of two arcs. A detail to give attention to is that the path from i and succ[i] changes direction after the substitution of the old edges, this is done to maintain the solution integrity. In particular, this optimization is performed until no more $\Delta(i,j)$ positive are found, in that case, the solution is no more improvable by the 2-opt algorithm.

```
Algorithm 7 2-opt refining
```

```
Input: G = (V, E), c : E \to \Re^+
Output: z hopefully good solution
 1: z \leftarrow *find feasible solution with an algorithm*
 2: flag \leftarrow false
 3: while flag == false do
         flag \leftarrow true
 4:
         (i, j) \leftarrow \operatorname{argmax}(\Delta(i, j) : i, j \in V)
 5:
         if \Delta(i,j) > 0 then
 6:
             flag \leftarrow false
 7:
             z \leftarrow *edge replacement*
 8:
         end if
10: end while
11: return z
```

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4.3 Metaheuristics

While the heuristic approach seen before was strictly specific to the TSP problem, the concept of metaheuristic is quite different. Their procedure is problem-independent and does not take advantage of any specificity of the problem. Generally, these techniques are not greedy and can accept a temporary deterioration of the solution in order to have a bigger search space in which to find a better solution.

In this section, some of them will be used such as the variable neighborhood search and the tabu search.

4.3.1 Variable Neighborhood Search

The variable neighborhood search (VNS) is the first metaheuristic approach presented. Its main purpose is to try to escape from a local optimal of the solution aiming to find the optimal solution of the problem.

The instance of the problem can be seen as a function with its local minima, local maxima, and so on. During the utilization of some heuristic techniques, it is possible that the process remains stuck in a sub-optimal area. The way in which it tries to escape from this region is done by changing the *neighborhood* of the solution: some data are modified and then a new minimum is searched.

In this report, the adjustment of the neighborhood is performed by a perturbation phase in which is applied a k-opt kick. This change is performed by selecting randomly a k set of edges and then replacing them with others in order to obtain a poor solution. In particular, the procedure done is the following: a 2-opt refining is applied to a reference solution, once the minimum is reached a kick is given to the solution to worsening the solution. Then the 2-opt refining approach is used again to find a new minimum. In my implementation are implemented three perturbations: 3, 5, and 7.

4.3.2 Tabu search

The tabu search is the second metaheuristic presented in this report. It is a method to improve the local search and avoid falling back in a previous local minimum. To do that the concept of tabu is introduced: a set of banned solutions prevents the algorithm to return an already visited result.

In this implementation of the tabu search the refining phase is performed by a 2-opt move as it happened in section 4.3.1, then the method to escape from this minimum is to perform a kick like as the previous section. The main difference is that during the VNS is possible to fall again and again in the previous solution, using the tabu search is not possible.

The worsening move is performed by swapping two non-consecutive edges, then they

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Algorithm 8 VNS

```
Input: G = (V, E), c : E \to \Re^+, k, qlobal\_timelimit
Output: z hopefully good solution
 1: z_{curr} \leftarrow z \leftarrow \text{*built solution using an heuristc*}
 2: best\_cost \leftarrow cost(z)
 3: cycles \leftarrow 0
 4: while time_elapsed < global_timelimit do
         z_{curr} \leftarrow 2\text{-}\mathrm{OPT}(z_{curr})
         if cost(z_{curr}) < best\_cost then
 6:
 7:
              z \leftarrow z_{curr}
              best\_cost \leftarrow cost(z_{curr})
 8:
         end if
 9:
10:
         *Bigger the value of cycles bigger the kick*
         z_{curr} \leftarrow \text{K-KICK}(cycles)
11:
12:
         cycles \leftarrow cycles+1
13: end while
14: return z
```

are declared as tabu: they cannot be swapped for a while. This process precludes the possibility of falling into a cycle of deterioration-recover, where to final solution is always the same. This algorithm will continuously worsen the solution until a non-tabu move is performed.

The way in which this method is implemented is the following: each time a worsening action is performed the nodes selected are declared as tabu and the edges connected with them cannot be changed by any improving moves. If each time this action has been performed the list of tabu nodes becomes so large that no more moves are allowed. To avoid this case it is introduced a new variable called *tenure*, its main idea is to limit how many times a tabu is valid.

To do that we declare a node tabu (for example i) by inserting into an array the iteration number h, so $tabu_nodes[i] = h$. This constraint will last for a tenure number of times, following this rule:

$$iteration_number - tabu[i] \le tenure$$
 (4.6)

A good choice of *tenure* is crucial to allow the algorithm to perform adequately. The best decision is to make it variable regarding the number of iterations already completed. In algorithm 9 can be seen the proceeding of this method.

4.3.3 Genetic algorithms

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Algorithm 9 Tabu search

```
Input: G = (V, E), c : E \to \Re^+, k, global\_timelimit
Output: z hopefully good solution
 1: z_{curr} \leftarrow z \leftarrow \text{*built solution using an heuristc*}
 2: best\_cost \leftarrow cost(z)
 3: iteration\_counter \leftarrow 1
 4: while time_elapsed < global_timelimit do
         *Performs a 2-opt refining applying the constaraint described in section 4.3.2. The
    iteration counter is increased each time a move is performed. If no moves are allowed
    this method does nothing*
        z_{new} \leftarrow \text{TABU-2-OPT}(z_{curr}, tabu, tenure, iteration\_counter)
 6:
 7:
        if cost(z_{curr}) < best\_cost then
 8:
             z \leftarrow z_{curr}
 9:
             best\_cost \leftarrow cost(z_{curr})
        end if
10:
        first\_node \leftarrow RANDOM(|V|)
11:
        second\_node \leftarrow RANDOM(|V|)
12:
        tabu[first\_node] \leftarrow iteration\_counter
13:
        tabu[second\_node] \leftarrow iteration\_counter
14:
        z_{curr} \leftarrow 2\text{-KICK}(first\_node, second\_node)
15:
        iteration\_counter \leftarrow iteration\_counter + 1
17: end while
18: return z
```

References

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