UNIVERSITY OF PADUA

Department of Mathematics Computer Science Master Degree

Methods and Models for Combinatorial Optimization project

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1 Abstract

The project is composed by two parts:

- Implementing an Integer Linear Programming model with the Cplex API
- Implementing a meta-heuristic solution method

For the second part my choice of method is a genetic algorithm, details on implementation and decisions in section 3. Both solution methods will be applied to the following problem:

A company produces boards with holes used to build electric frames. Boards are positioned over a machines and a drill moves over the board, stops at the desired positions and makes the holes. Once a board is drilled, a new board is positioned and the process is iterated many times. Given the position of the holes on the board, the company asks us to determine the hole sequence that minimizes the total drilling time, taking into account that the time needed for making an hole is the same and constant for all the holes.

The two different approaches are then tested using some benchmark tests from literature and compared to spot differences in results (optimality, execution time, reliability, ...)

2 Integer Linear Programming Model

2.1 Network Flow Model Representation

The problem can be represented on a complete weighted graph G = (N, A) where N is the set of nodes (holes on the board) and A is the set of the arcs $(i, j), \forall i, j \in N$ (trajectory of the drill moving from hole i to hole j). To each arc will be associated a weight c_{ij} that represents the time needed to move from starting hole to destination.

Using this representation the problem is equivalent to determine the minimum weight hamiltonian cycle on G, so it is like the very popular Travelling Salesman Problem (TSP).

To solve the TSP problem we can formulate it as a network flow model on G. Given a starting node $0 \in N$, let |N| be the amount of its output flow, the problem can be solved by finding the path for which:

- Each node receives 1 unit of flow
- Each node is visited once
- The sum of costs in selected arcs is minimum

2.2 Variables and Constraints

The problem can then be represented like this:

SETS:

```
N = \text{the graph nodes (holes)}

A = \text{arcs in the form } (i, j), \forall i, j \in N \text{ (trajectories between holes)}
```

PARAMETERS:

```
c_{ij} = time taken by the drill to move from i to j, \forall (i,j) \in A
0 = starting hole, 0 \in N
```

2.2 Variables and ConstruMSEGER LINEAR PROGRAMMING MODEL

DECISION VARIABLES:

 $x_{ij} = \text{amount of the flow shipped from } i \text{ to } j, \forall (i, j) \in A$ $y_{ij} = 1 \text{ if arc } (i, j) \text{ ships some flow, 0 otherwise, } \forall (i, j) \in A$

OBJECTIVE FUNCTION:

$$\min \sum_{i,j|(i,j)\in A} c_{ij} \cdot y_{ij}$$

CONSTRAINTS:

$$\sum_{j|(0,j)\in A} x_{0j} = |N|$$

$$\sum_{i|(i,k)\in A} x_{ik} - \sum_{j|(k,j)\in A} x_{kj} = 1 \qquad \forall k \in N \setminus \{0\}$$

$$\sum_{j|(i,j)\in A} y_{ij} = 1 \qquad \forall i \in N$$

$$\sum_{i|(i,j)\in A} y_{ij} = 1 \qquad \forall j \in N$$

$$x_{ij} \le |N| \cdot y_{ij}$$
 $\forall (i,j) \in A$

$$x_{ij} \in \mathbb{Z}_+$$
 $\forall (i,j) \in A$

$$y_{ij} \in \{0, 1\} \qquad \forall (i, j) \in A$$

3 Genetic Algorithm

Genetic algorithms are a population based heuristic method that simulates nature's evolutionary system to evolve a starting population of random solutions to obtain very good (possibly optimal) solutions at the end of the process. To do this a random starting population is generated, then each solution is evaluated with a fitness function. After that some solutions, usually the best ones, are chosen to create offsprings that will replace worst solutions. This way the algorithm performs a sort of natural selection keeping the best solutions and improving them over generations.

3.1 Encoding

To represent a TSP solution the path representation has been used, it consists in a sequence of N nodes, where N is the total number of nodes, sorted from the starting one to the last visited on encoded path.

For example the path $0 \to 3 \to 1 \to 4 \to 2 \to 0$ is encoded as:

3.2 Fitness

As fitness concerns I took a different approach from usual genetic algorithms and considered the sum of costs in path as the fitness and I will try to minimize it. This choice was taken because other considered fitness functions had drawbacks, mainly because most of them were in the form upperBound-costs but upper bounds are not so strict and then a proportional choice based on fitness would yield a very similar probability for each individual (because maxFitness-minFitness is much lower than upperBound-anyFitness).

By setting the fitness function equal to the sum of path costs it it directly proportional to objective function resulting in a more correct reward based on fitness.

3.3 Select

The select operator is a little tricky because of the fact that solutions with lower fitness should have higher probabilities, so both Montecarlo and Linear ranking methods can't be used. N-Tournament is a valid alternative, but it was discarded because if a subset of k solutions is chosen, the k-1 worst solutions are never used for crossover and this reduces (slightly) diversification.

The final choice is a probability calculated based on position in an array of solutions sorted by fitness. The selected solution is the one with index

$$index = \lfloor N * r^2 \rfloor$$

where N is the total number of nodes and r is a random number $\in [0, 1)$

This way the solutions with lower cost are more likely to be chosen than the others, but each solution might be selected.

3.4 Crossover

The crossover operator takes a certain number of parents and creates an offspring that inherits parents properties. In the project was used the Order Crossover (OX), that from 2 parents creates one child, and works like this:

```
Parent 1: 8 4 7 <u>3 6 2 5 1</u> 9 0
Parent 2: 0 <del>1</del> <del>2</del> <del>3</del> 4 <del>5 6</del> 7 8 9
Child 1: 0 4 7 3 6 2 5 1 8 9
```

Figure 1: Example of Order Crossover (OX)

- 1. Select a random subsequence of consecutive alleles from parent 1. (underlined)
- 2. Drop the subsequence down to Child and mark out these alleles in Parent 2.

3. Starting on the right side of the subsequence, grab alleles from parent 2 and insert them in Child at the right edge of the subsequence. Since 8 is in that position in Parent 2, it is inserted into Child first at the right edge of the subsequence.

Notice that alleles 1, 2 and 3 are skipped because they are marked out and 4 is inserted into the 2nd spot in Child.

3.5 Mutation

Only one type of mutation is used and it is the 2-opt, an algorithm that takes a substring of the path representation and reverse it. The idea is to take a route that crosses over itself and reorder it so that it does not. This is good for the main purpose of mutations (prevent genetic drift) as well as for improving solutions.

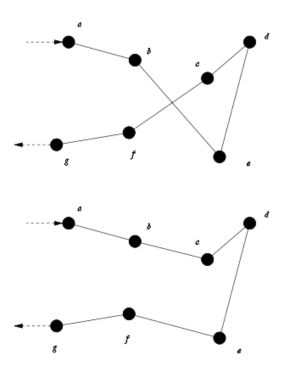


Figure 2: Example of 2-opt mutation

To control the mutation there is a parameter MUTATION_CHANCE that sets the probability that this mutation happens.

3.6 Generational Replacement

When a population advance to next generation we need to replace the old bad solutions with newly created offsprings. This is done by elitist, so only the best solutions are kept and all the others are replaced by new solutions. The percentage of survivors (and replaced solutions) is set by the parameter SURVIVAL_RATIO.

3.7 Improvements

After implementing all the elements of the algorithm and doing some basic tests I found out that the convergence was fast and after a few generations the solution was already quite good and it was rarely improved after. This means that algorithm was iterating through a lot of generation with very few improving solutions, so I decided to introduce random restarts reducing generations count and considering the best solution from all evaluated populations. This approach worked out well to find more reliably good solutions but slowed computation down. To make up for the loss in performance multithread has been used running genetic algorithm on many random starting populations in parallel. It is possible to control the amount of populations evaluated by setting the parameter POPULATIONS.

Another thing that has been tested is to modify the mutation chance at runtime based on the number of non-improving generations, but this didn't lead to significative improvement of solutions.

4 Tests

Tests have been taken from the ones used in laboratories and from TSPLIB, a library of sample instances for the TSP from various sources and of various types. A total of 13 instances, ranging in size from 12 to 100, have been tested 10 times for each method. Higher size tests should have been included but ILP solver already starts to slow at 100 nodes and the time to test was not so much.

4.1 ILP

Results obtained testing the ILP solver gives expected results, as it always finds optimal solution, and execution times grow with problem size. Average solution times on 10 runs are displayed in Table 1.

Test	Size	Avg Time (s)
tsp12.dat	12	0,133
tsp15.dat	15	0,062
tsp17.dat	17	0,162
tsp26.dat	26	0,298
tsp29a.dat	29	0,633
tsp29b.dat	29	0,668
tsp42.dat	42	2,462
tsp48a.dat	48	8,371
tsp48b.dat	48	10,357
tsp48c.dat	48	2,275
tsp60.dat	60	27,066
tsp100a.dat	100	101,527
tsp100b.dat	100	128,637

Table 1: ILP tests results

We can see that while solving small instances is very efficient, bigger ones require more time and going from size 60 to 100 (5/3 ratio) it gets way slower

(more than 5 times), and the trend continue growing in size. This is due to exponential growth of the solution space the solver has to explore to find and prove optimality.

4.2 Genetic Algorithm

Genetic algorithms, due to their random nature, give different results at each execution, so an average of 10 runs is considered. Obtained solutions and computation time are also greatly related to set parameters, which are reported for each instance. Parameters don't differ so much, but more generations and bigger populations are used for bigger instances, and to keep computational times low populations evaluated are reduced. Results obtained with specified tests are reported in Table 2.

Test	Size	Average	Avg. opt.	Opt.	Parameters		
Test		time	distance	found	Populations	Pop. size	Generations
tsp12.dat	12	0,405	0,00%	10/10	5	500	200
tsp15.dat	15	0,401	0,00%	10/10	5	500	200
tsp17.dat	17	0,502	0,02%	9/10	5	500	250
tsp26.dat	26	1,037	0,36%	9/10	10	500	250
tsp29a.dat	29	1,038	0,68%	2/10	10	500	250
tsp29b.dat	29	1,049	0,40%	3/10	10	500	250
tsp42.dat	42	1,246	0,75%	2/10	10	500	300
tsp48a.dat	48	1,779	2,30%	0/10	7	1000	300
tsp48b.dat	48	1,777	1,65%	0/10	7	1000	300
tsp48c.dat	48	1,859	2,91%	0/10	7	1000	300
tsp60.dat	60	1,802	3,19%	0/10	7	1000	300
tsp100a.dat	100	3,053	7,46%	0/10	7	1000	500
tsp100b.dat	100	3,083	6,33%	0/10	7	1000	500

Table 2: Genetic algorithm tests results

Note that computation time is highly dependent on set parameters, so if time is less important parameters could be increased to get better solutions. A

4.3 Tests details 4 TESTS

direct consequence is that with genetic algorithms it's possible to get the desired precision/efficiency ratio. However as GAs don't provide an optimality proof, if an optimal solution is found, all remaining computation is executed anyway.

4.3 Tests details

Here have been shown the most important results, all recorded data can be found in this spreadsheet:

 $https://docs.google.com/spreadsheets/d/1wJ98av-soxw_okteJ7X8zMJUfesh_16ECjA9uRbP8Hg/edit?usp=sharing$

5 Conclusions

TODO

A Bibliography

- Project reference 1/2: http://www.math.unipd.it/luigi/courses/metm odoc/z01.eserc.lab.01.en.pdf
- Project reference 2/2: http://www.math.unipd.it/luigi/courses/metm odoc/z02.eserc.lab.01.en.pdf
- Genetic algorithms for TSP: https://www.iccl.inf.tu-dresden.de/w/images/b/b7/GA_for_TSP.pdf
- TSPLIB (test source): http://www.comopt.ifi.uni-heidelberg.de/software/TSPLIB95/