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Minimum Weight Vertex Cover Problem

ARTIFICIAL INTELLIGENCE PROJECT

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Introduction

1.1 Problem

Given a problem instance (G, ω) , where G is a undirected graph G(V, E) and $\omega : V \to \mathbb{R}^+$ a function that associates a positive weight value $\omega(v)$ to each vertex $v \in V$, the Minimum Weight Vertex Cover can formally be defined as follows:

$$\mathbf{minimize} \quad \omega(S) = \sum_{v \in S} \omega(v), \quad S \in V$$

such that $\forall (v_i, v_j) \in E, v_i \in S \lor v_j \in S$.

Note that the MWVC is a NP-complete problem.

1.2 Proposed Solution and Motivation

This project implements a solution to the MWVC problem using a Genetic Algorithm (GA) with one-point crossover and k-tournament selection.

The chosen algorithm facilitates parallel computing compared to Tabu Search, as well as allowing to reduce the risk of the local optima stagnation. Also, Branch-and-Bound has limitations concerning the problem size, while Genetic Algorithms are suitable for NP-Hard problems such as MWVC.

Regarding the chosen Genetic Algorithm variation, even though it is not immediate to state whether one-point crossover will achieve better results over uniform crossover without prior testing, it has been widely proved that k-tournament selection outperforms roulette wheel in most scenarios [1].

Genetic Algorithms

2.1 Behaviour and Structure

Genetic Algorithms (GAs) are a class of evolutionary algorithms inspired by the principles of natural selection. They operate by iteratively evolving a population of potential solutions towards an optimal or near-optimal state. The process unfolds as follows:

Initialization An initial population of candidate solutions is randomly generated.

Evaluation Each individual in the population is evaluated based on a predefined fitness function, which quantifies its quality or suitability with respect to the problem at hand.

Selection A subset of individuals are selected from the populations. The chosen individuals are selected as parents of the following population. Selection is based on fitness, with fitter individuals having a higher probability of being chosen.

Crossover Using crossover, selected parents genetic material are combined to create offspring.

Mutation With a certain probability, random mutations are introduced into the offspring. This is called mutation and it helps to maintain diversity within the population and prevents premature convergence towards suboptimal solutions.

Replacement This newly generated population replaces the old one.

The cycles of evaluation, selection, crossover, mutation, and replacement are repeated until a halting criteria is satisfied.

2.2 k-Tournament Selection

The selection algorithm for the proposed solution is *k-tournament*. This algorithm involves running several "tournaments" among k individuals chosen at random from the population, until the desired amount of population is reached. Each tournament selects the best amongst the k selected individuals.

The tournament size k can be adjusted to balance exploration and exploitation. Smaller k introduces more diversity, while larger k focuses more on exploiting fittest individuals. Given a population of n individuals:

k=1: selection is random, there is no preference based on fitness

k=n: the fittest individuals are always selected

Algorithm 2.1: k-tournament selection 1 function k-tournament(population, k, n) new_population = [] while new_population.size < n</pre> 3 selected = []while selected.size < k</pre> individual = random element from population selected.push(individual) end new_population.push(best(selected)) end 10 11 return new_population 12 end

2.3 Single-Point Crossover

The simplest form of crossover is the single-point crossover, where a random crossover point is selected and the genetic material is exchanged between the parents at that point.

2.4 Bit-Flip Mutation

Mutation is a genetic operator that introduces random changes in the offspring. The simplest form of mutation is the bit-flip mutation, where a bit has a probability p of being flipped.

Implementation

3.1 DEAP Framework

The chosen framework for the implementation is DEAP [2]. DEAP (Distributed Evolutionary Algorithms in Python) is a Python library that excels at rapid prototyping and testing of ideas, making the tool ideal for the project.

3.2 Gene Representation

The gene representation is a binary string, where each bit represents the presence of a vertex in the solution. The length of the string is equal to the number of vertices in the graph. The weight of the vertex is stored in a separate list, where the index corresponds to the vertex index.

1	1	0	1	0	0	1	0	1	1	
---	---	---	---	---	---	---	---	---	---	--

Figure 3.1: Example of a MWVC gene representation

3.3 Fitness Function

The fitness function for the Minimum Weight Vertex Cover problem calculates the weight of the solution by summing the weights of the vertices present in the solution. Additionally, it imposes a penalty for each edge that is not covered by the vertices in the solution, ensuring that the fitness value is higher for incomplete solutions. The penalty function (or a similar alternative) is mandatory since we're using a direct gene representation that could create invalid configurations of uncovered vertices.

```
for edge in graph.edges
if individual[edge.from] == 0 and individual[edge.to] == 0

f = f + penalty
end
end
return f
send
```

The objective function is the same but with a penalty of zero, and it returns NULL if the solution is invalid (i.e. there are uncovered edges).

3.4 Performance Metrics

In order to evaluate the performance of the algorithm, it is necessary to define a set of metrics to evaluate the quality of the solution and the performance of the algorithm.

The most important one is the fitness, as well as the objective (that does not include the penalties). Another metric to optimize is the number of evaluations, as it is a direct measure of the computational cost of the algorithm.

The execution time is not considered as a metric as it is highly dependent on the hardware and software environment.

It would be useful to include an optimality metric representing the difference between the best known solution and the solution found by the algorithm, but this information is not easily computable as the problem is NP-hard.

3.5 Parameters

Genetic Algorithms have a set of parameters that need to be tuned in order to achieve the best performance.

The following is the list of the meta-parameters configurable in the implementation:

- Population size (POPULATION_SIZE): the number of individuals in the population.
- Crossover probability (CXPB): the probability of crossover.
- Mutation probability (MUTPB): the probability of mutation.
- **k-tournament selection size** (K_TOURNAMENT): the size k of the tournament in the k-tournament selection.
- Number of generations (NUMBER_OF_GENERATIONS): the number of generations the algorithm will run.

The problem of finding the best parameters is an optimization problem itself, and it is possible to use a meta-heuristic algorithm to find the best parameters, such as a Genetic Algorithm [3]. This is out of the scope of this project even though it could be an interesting future development.

3.6 Benchmarking Flow

This paragraph describes the flow of the benchmarking process. The goal of the benchmarking process is to test the algorithm with a fixed parameters combination over different graph instances to extract the metrics and evaluate the performance of the algorithm. The parameters are chosen based on the literature and are discusse in the chapter 4.

The process is divided into the following steps:

- 1. Load the graphs: the problem instances for testing are loaded from the files.
- 2. Compute graphs properties: for each graph a set of properties are computed, such as the number of vertices, the number of edges and the density.
- 3. Create the DEAP structures: the DEAP structures are created, including the individuals, the population and the evaluation function.
- 4. Run the algorithm: the algorithm is executed with the set parameters over the graph instances.
- 5. Save the results: the results are saved to a multiple files in JSON format for further analysis.
- 6. **Analyze the results**: the results are analyzed to extract the metrics and evaluate the performance of the algorithm.

Every step, except the last one, is executed within a Python script with configurable parameters. The last step is performed using a Jupyter Notebook for a faster and more interactive analysis.

The benchmarking process is repeated iteratively over different parameters combinations to find the best configuration for the algorithm.

Results

4.1 Introduction

As previously stated, this chapter describes the process of tuning the parameters of the genetic algorithm and the results obtained from the experiments.

4.2 Iteration 1: Default Parameters

This first iterations consists of the most basic configuration of the genetic algorithm with literature defaults [4].

Parameter	Value
POPULATION_SIZE	100
CXPB	0.5
MUTPB	0.2
K_TOURNAMENT	3
NUMBER_OF_GENERATIONS	1 000 000

Table 4.1: Parameters for Iteration 1

After grouping and plotting the results by the problem class it's immediately clear that the genetic algorithm converges depending on the problem size (amount of vertices). This will be useful for later when limiting the number of generations. Since we still need to try different configurations, we will change the number of generations for last.

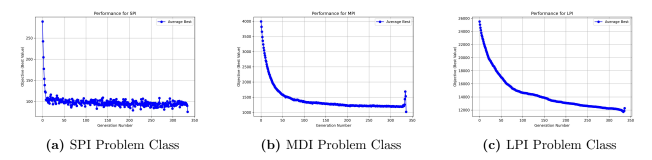


Figure 4.1: Objective function value by generation for each problem class (Iteration 1)

The results for the initial iteration are as follows.

Problem Class	Test Instance	Average Objective	Average Evaluations
LPI	10000	12317.3	20000.0
MPI	3000	1027.2	20000.0
MPI	2000	223.9	20000.0
MPI	750	2773.6	20000.0
MPI	500	903.5	20000.0
SPI	150	113.7	20000.0
SPI	120	92.2	20000.0
SPI	60	178.3	20000.0

Table 4.2: Results for Iteration 1

4.3 Iteration 2: Population Size

The first parameter to be tuned is the population size. The test is initially performed using a population size of 50, 75, 100, 150, 200 and 400. The results are compared using the average objective function value since the average number of evaluations would still be capped at 20000.

After performing the tests, it looks like the population size of 150 is the best for every test instance. The improvement is not significant, but it's still better than the other population sizes. Other than the original population sizes, different ones have been tested as well, but none of these have shown any improvement.

Pop. size	50	7 5	100	150	200	400
Avg obj	2229.7875	2196.2875	2203.7125	2126.1125	2259.25	2156.4375

Table 4.3: Average Objective for different Population Sizes

This iteration keeps the parameters the same as the precedent iteration, except for the population size which is now set to 150.

Parameter	\mathbf{V} alue
POPULATION_SIZE	150
CXPB	0.5
MUTPB	0.2
K_TOURNAMENT	3
NUMBER_OF_GENERATIONS	1 000 000

Table 4.4: Parameters for Iteration 2

The results are as follows.

Problem Class	Test Instance	Average Objective	Average Evaluations
LPI	10000	12321.9	20000.0
MPI	3000	1037.4	20000.0
MPI	2000	217.5	20000.0
MPI	750	2683.9	20000.0
MPI	500	850.6	20000.0
SPI	150	116.8	20000.0
SPI	120	84.0	20000.0
SPI	60	177.0	20000.0

Table 4.5: Results for Iteration 2

4.4 Iteration 3: Crossover and Mutation Probabilities

The next parameters to be tuned are the crossover and mutation probabilities. Since these two parameters are closely related, they will be tested together. The test is performed using a grid search with the following values: 0.01, 0.05 and 0.1 for mutation probability and 0.3, 0.5 and 0.7 for crossover probability. A bash script is used to run all the tests at once.

The results show that the best combinations are 0.1 for mutation probability and 0.3 and 0.5 for crossover probability. The first combination is the best for the LPI problem class, while the second one performs better on the other two classes. After making these discoveries, the tests are run again with a finer grid search around these values.

The final choice consists of 0.4 and 0.25 for crossover and mutation probability respectively.

Parameter	Value
POPULATION_SIZE	150
CXPB	0.4
MUTPB	0.25
K_TOURNAMENT	3
NUMBER_OF_GENERATIONS	1 000 000

Table 4.6: Parameters for Iteration 3

The randomness factor has an impact on the average objective function value, even though ten runs are performed for each test instance. The final results are as follows.

Problem Class	Test Instance	Average Objective	Average Evaluations
LPI	10000	12093.5	20000.0
MPI	3000	1007.8	20000.0
MPI	2000	205.4	20000.0
MPI	750	2656.4	20000.0
MPI	500	807.8	20000.0
SPI	150	118.4	20000.0
SPI	120	85.5	20000.0
SPI	60	172.7	20000.0

Table 4.7: Results for Iteration 3

4.5 Iteration 4: Tournament Size

The next parameter to be tuned is the tournament size. The test is performed with the following values: 2, 3, 4, 5.

The results show that the best tournament size is 2. Even tough the tournament size of 3 looks better for some test instances, the average objective function value is better for the tournament size of 2. Another reason to choose the tournament size of 2 is that it allows for a faster convergence.

Parameter	Value
POPULATION_SIZE	150
CXPB	0.4
MUTPB	0.25
K_TOURNAMENT	2
NUMBER_OF_GENERATIONS	1 000 000

Table 4.8: Parameters for Iteration 4

The average objective value for each test instance is as follows.

Problem Class	Test Instance	Average Objective	Average Evaluations
LPI	10000	11693.3	20000.0
MPI	3000	905.1	20000.0
MPI	2000	228.3	20000.0
MPI	750	2657.8	20000.0
MPI	500	815.7	20000.0
SPI	150	113.8	20000.0
SPI	120	85.0	20000.0
SPI	60	177.7	20000.0

Table 4.9: Results for Iteration 4

4.6 Iteration 5: Number of Generations

The number of generations is by far the most important parameter to be tuned, as well as the hardest to tune. By now, the genetic algorithm has been tuned to the best of its abilities, and the number of generations was not considered in the previous iterations. This iteration will improve considerably the efficiency of the genetic algorithm, with the downside of getting a slightly less accurate solution. The best way to tune this parameter is to plot the average objective function value by generation and see when the genetic algorithm converges.

At first glance, the plots show how the problem size doesn't look to depend at all on the number of generations. Figure 4.2 shows this behavior. Binding the number of generations to the problem size is not a good idea.

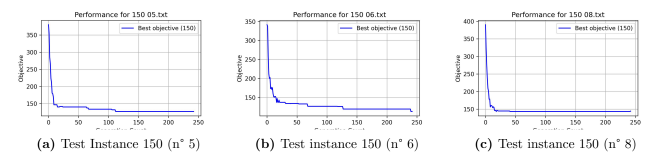


Figure 4.2: Objective function value by generation for different test instances

The remaining option is to dynamically set the number of generations depending on a dynamic convergence criteria. The chosen criteria is to stop the genetic algorithm when the average objective function value doesn't improve for 100 generations.

There could be better criteria, but this one was chosen for its simplicity. This still halves the number of generations for SPI test instances and leaves room for improvement in case one decides to increase the evaluation limit.

Parameter	Value
POPULATION_SIZE	150
CXPB	0.4
MUTPB	0.25
K_TOURNAMENT	2
NUMBER_OF_GENERATIONS	dynamic

Table 4.10: Parameters for Iteration 5

The final results are as follows.

Problem Class	Test Instance	Average Objective	Average Evaluations
LPI	10000	11426.8	20000.0
MPI	3000	917.4	20000.0
MPI	2000	226.9	19335.8
MPI	750	2479.1	20000.0
MPI	500	834.7	19459.4
SPI	150	113.0	11900.9
SPI	120	83.5	10097.8
SPI	60	176.8	12075.9

Table 4.11: Results for Iteration 5

Conclusion

The analysis shows how an iterative process can be a solution to find the best parameters for a genetic algorithm implementation. I found it hard to find a linear path to the solution, but the iterative process helped me to find the best parameters for the genetic algorithm. I learned how this kind of tuning process is very difficult to standardize due to the uniqueness of each problem. I am pround and happy of the results I got, especially regarding the framework created to test and then analyze the results. I think this was the key to succeed in these kinds of experimental projects.

The problem task suggested to run each test ten times and then perform the average of the results. I think this is a good approach to reduce the noise in the results, but I feel that for this algorithm implementation it wasn't enought. It looked like different executions gave too different results, and I think that running the tests more times could have helped to reduce the noise even more, allowing for more accurate results.

Despite the obtained results, I believe it would be interesting to approach the problem using a metaheuristic algorithm, such as the genetic algorithm itself, to solve the problem of finding the best parameters for the genetic algorithm. The iterative process is slow and requires a lot of manual work, while a metaheuristic algorithm could be more challenging and fun.

I really enjoyed working on this project, I think I learned a lot about genetic algorithms and how to tune them. The task problem was clear and having the test instances ready saved me plenty of time. This was my first experimental research project, and the freedom to choose any language and technology I believe was the essence of the project, even though it was a bit challenging to find the best tools to use and I was a little bit lost in the beginning. When looking for literature resources, I found the No Free Lunch Theorem [5], which left me slightly unsettled but gave me a different approach to the problem. I hope to have the opportunity to work on more projects like this in the future.

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