

AUTHOR, OLDDEGREE

Evaluation of a Genetic Algorithm on generating critical Scenarios in a Traffic Simulation

Master's Thesis

to achieve the university degree of Master of Science

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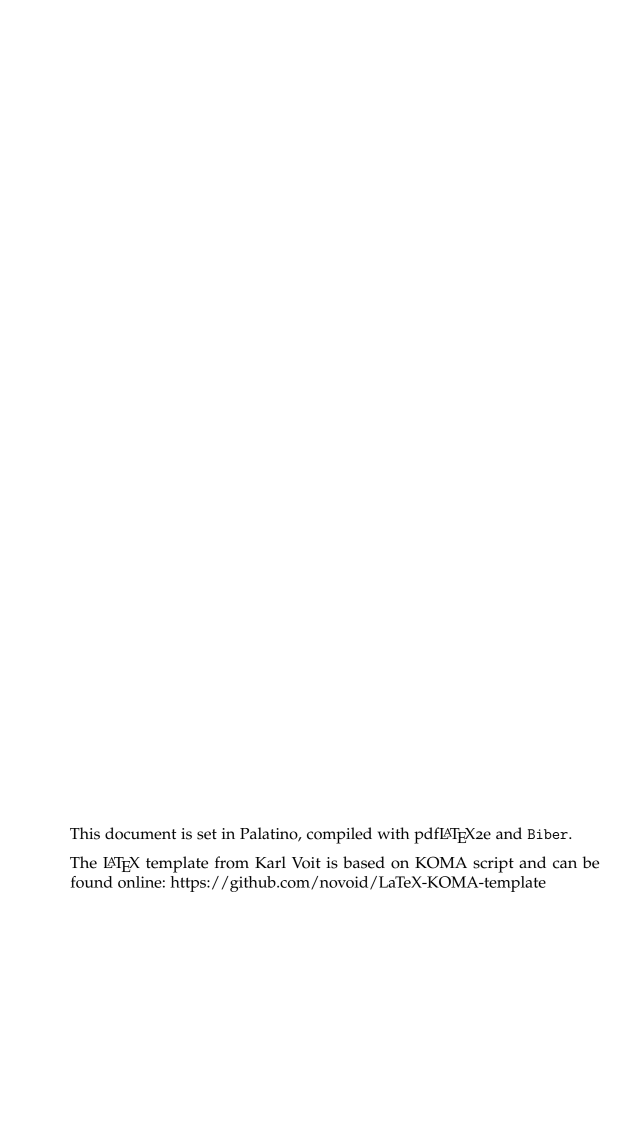
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Abstract

This is a placeholder for the abstract. It summarizes the whole thesis to give a very short overview. Usually, this the abstract is written when the whole thesis text is finished.

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1 Introduction

This Thesis will use a Genetic Algorithm in order to generate critical Driving Scenarios for testing ADAS/AD Functionality in vehicles. While generating these scenarios is the objective, the main task of the thesis will evolve around the implementation of the Genetic Algorithm as well as the Optimization of its Hyperparameter.

1.1 Research Questions

1.1.1 Research Question 1

Is a Genetic Algorithm suitable for generating critical driving scenarios compared to a random generation?

1.1.2 Research Question 2

Can hypertuning improve the performance of a Genetic Algorithm?

1.1.3 Research Question 3

Can a hypertuned Genetic Algorithm generalize on different start scenarios?

1.2 Shortcomings

This Master Thesis started with the developement of the Traffic Manger and thus progress was closely linked. Without a working simulations, no genetic alogirthmis could be tested. Due to time and performance constraints, it is not possible to test a full driving stack like autoware, as well as other professional ADAS/AD functions. In this Thesis, internal functions like Time-To-Collision and Emergency Braking will be optimized. The learned information on e.g. optimal hyperparameter settings can then be applied in further steps to test these functions. This will however not be tackled by this thesis.

Performance is also a problem and will lead to many shortcuts that need to be taken. There is a hughe number of possible compations of hyperparamter, so only a handful can be tested. In further chapers, these shortcuts will be explained and their relevancy will be dicussed.

2 Foundations

2.1 Genetic Algorithm

Genetic Algorithms are a popular search algorithm that utilizes the principle of Darwin. They have been used successfully in various areas. Some of their strengths are However we will also look at shortcomings, which mainly evolve around performance. We will have a look at its History and then discussing the most important parameters.

Define a vocabulary

The task of the Genetic Algorithm is to search for sequences of actions that will result in the most interesting Scenarios according to its cost function.

Genes are the building blocks of a GA

Usage of GA

dejong talks about dynamic param and why its not good

2.1.1 History

The GA was invented by....

2.1.2 Encoding

Binary, Hex,

cite what makes an encoding good: eg. simplicity,...

2.1.3 Different Hyperparameter

Hyperparamter have a huge influence on the performance of a Genetic Algorihm. They have an impact on the "convergin" ... It has been shown, that there is no universal hyperparamter set and that it needs to be optimized on a per "problem" basis.

Num of generations

The Number of Generation defines the duration of a GA. As long as the algorihtm has not converged,? For my testing, using a generation size of 40 was almost always sufficient, and will thus mostly be used.

Pop Size

Pop size will set the number of Individuals of a GA per Generation. The higher the pop size, the bigger the less change of premature converging. It will however also lead to a longer convergin time.

Selection

pros and cons of roulette vs Tournament Selection defines how which individuals are allowed to mate and move into the next generation.

tournament was chosen to be used for this works because of this paper (and also because of pros and cons list)

cite paper

Other ideas are evolve around having a flexible selection system debending on fitness

Crossover

Discuss all used crossover methods

Crossover is the mating process.

4

Mutation

Mutation is responsible for introducing new information into the gene pool.

Discuss individual mutation

Discuss all used mutation methods

Other

More to come....

2.2 Behavior Tree

A behavior tree is a decision tree.

insert a good introduction to BT

2.2.1 Usage for GA

Due to the fact, , that there is no full stack available for the EGO vehicle, a solution had to be found. In order to have the Genetic Algorithm controll only NPCs and not the EGO vehicle itselve, a behaviour tree is used. The behaviour tree is used to controll the EGO vehicle over the action interface provided by the Traffic Manager. This is the same as the Genetic Algorithm is doing.

insert ref to discussion

The behaviour tree will define which direction the EGO should take at junctions and it will realistically dodge obstacles intoduced by the Genetic Algorithm. The main goal of the BT is to make the EGO vehicle behave in a realistic way.

In a further chapter it will be discussed if a GA with controll of the EGO (i.e. no BT will be used) lead to better cost.

While the aim of the GA is to find the most optimal solution, considering the vastness of the hyperspace, this is unlikely. Rather, we want to find the "best" local minimas. Considering the contex of Automotive testing, it is not so much of importance to find "the best fail of the ADAS/AD System", rather its important to find "all" fails.

3 Implementation

3.1 Traffic Manager

The Genetic Algorithm will control the simulation of a custom developed Traffic Manager. This Traffic Manager was developed closely to fit the needs of the Genetic Algorithm. It, however is not part of this Thesis and will thus will only get a brief introduction. In general, it will simulate traffic starting from a predefined scenarios which defines the positions and types of vehicles and pedestrians (i.e. actors). A simulation always consist of at least one EGO vehicle. Additionally any number of NPCs can be used.

While the NPCs are only controlled by the Traffic Manager, the ego vehicle can be either partly or even completely controlled by an ADAS/AD Function. The stated goal is to test these functions for errors.

For all simulations done by this thesis, the Traffic Manager was set to 100Hz.

3.1.1 Action Interface

To control the behaviour of the actors inside the simulation, actions can be requested over the "Action Interface" provided by the Traffic Manager. An action will request a certain behaviour from an actor. An action can be set to at any timestep for any actor¹. Pedestrians and vehicles have a different set of actions.

Insert graph of action interface

¹depending on the ADAS/AD function under test, the Action Interface might be disabled for the EGO vehicle

If no action is set, the actor will behave in a normal manner inside the simulation. This means that the actor will follow along its path until a new action changes its behaviour.

The following list are all actions provided by the traffic manager that were available for the genetic algorithm at the time of this master thesis.

• JunctionSelection

- Parameters: Vehicle ID: int, Junction_selection_angle: float
- Angle is set in radiant. Default value is o. Vehicles will chose which direction to take at a junction based on this angle.

LaneChange

- Parameters: Vehicle ID: int, ...
- Initiates a LaneChange based on its given parameters.

• AbortLaneChange

- Parameters: Vehicle ID: int, ...
- If a LaneChange is currently happening, it will get aborted.

• ModifyTargetVelocity

- Parameters: Vehicle ID: int, ...
- Modifies the interal Target Velocity of the Traffic Manager by a percentage. If it is for example o, the vehicle will stop.

TurnHeading

- Parameters: Pedestrian ID: int, ...
- The pedestrian will turn 180 degrees and walk in the oposite direction

• CrossRoad

- Parameters: Pedestrian ID: int, ...
- The pedestrian will cross the road immediately.

• CrossAtCrosswalk

- Parameters: Pedestrian ID: int, ...
- The pedestrian will cross the road at the next crosswalk.

All these actions are accessed by the Genetic Algorithm and the Behavior tree. The Behaviour tree sets only actions for the EGO vehicle, while the Genetic Algorithm will set all actions for the other actors in the simulation.

3.2 Genetic Algorithm

For implementing the Genetic Algorithm, DEAP was chosen. It is a popular tool for academia and allows for high customacibility. As has been stated in section 3.1.1, it has full access over setting actions for all NPCs. Using theses actions, it tries to optimize a cost function. This section aims to explain how the genetic algorithm was implemented and which different hyperparameters are variable. In chapter 4, the best hyperparameter combination will be generated, further analysis is done in 5.

generated, further analysis is done in 5.

A few default settings for the genetic algorithm had to be chosen. It was decided that the genetic algorithm will set an action per actor every 50 steps, which translates to 0.5 seconds (simulation runs at 100hz). In other words, every 50 steps of the simulation is 1 timestep for the genetic algorithm. The time of a simulation is always set to 35 seconds. Each genetic algorithm

reducing the amount of needed computations.

If the GA decides to not set an action for the integer, it sets "NoAction" as a placeholder.

will run for 30 generations. These to settings were chosen with the aim of

3.2.1 Encoding

When implementing a Genetic Algorithm, it is necessary to implement an encoding that fits to the problem. Each individual basically thus needs to include all actions that the genetic algorithm wants to apply. Different encodings presented in section 2.1.2, however none directly fitted to the problem presented. A custom encoding for both chromosomes and genes needed to be generated.

cite 3 examples

Chromosome

Each individual has 1 chomosome which consits of a list of genes, which has been explained in section. Starting out, 2 different encodings came to mind, in both cases, the genes position in the chromosome defined the time an action is set.

Time The first encoding is will be called "Time". Each gene corresponds to 1 timestep (so 1 gene per every 0.5 seconds). One gene has a list of the length of the number of all NPCs. This list is populated with actions. The index of an action in the list corresponds to the NPC id (index + 1 as the ego has id == 0, thus a start at 1 is needed). A visualization is seen in figure 3.1.

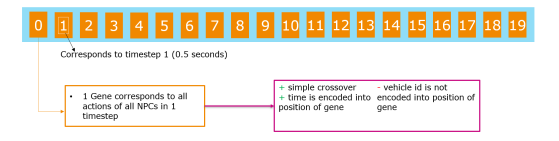


Figure 3.1: Time

Given the previously stated simulation time of 35 seconds, each chromosome has a length of 35 * 2 = 70 genes. Each gene consits of $number_of_actors$ actions. Crossover can thus only move all actions of a timestep at once, modifing between actions of the same timestep can only be done using mutation. If this is desired will be seen in the next chapters.

TimeNPC The second encoding has the name "TimeNPC", and is somewhat differently structured. Now, genes only hold 1 action, encoding now not only the timestep, but also the actor id in the position of the gene inside

ref

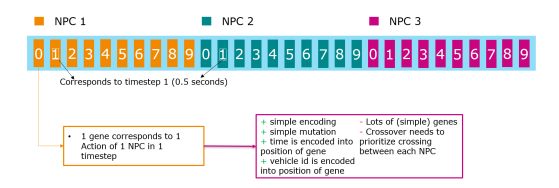


Figure 3.2: Time + NPC

the chromosome. Now, each actors actions will be listed one after another. This is visualized in figure 3.2.

Now, each gene has a length of 1 and each chromosome now has a length of $35 * 2 * number_of_actors$, which makes them much longer compared to the previous encoding. This now allows the crossover operation to modify only specific actions of one timesstep. Previously this was not possible.

However for this encoding to make sense, the crossover operations "One-Point" and "TwoPoint" had to be modified as follows. In an example of 10 NPCs, the operations will be executed for each NPC separately. Otherwise these two operations would have only had an effect on 1 or 2 different NPCs. For the reaiming NPCs, their actions would stay the same.

Gene

Two different encodings for genes were implemented as well. A gene always consits of a list, which depending on the chromosome type either has a length of $number_of_actors$ (In case ChromosomeEncoding == Time) or of length 1 (in case ChromosomeEncoding == TimeNPC). The following two encodings thus show the type of object, which is in these lists.

Integer The first encoding uses integer, which are translated into actions when the simulation is started. For each action, a range of integers is assigned, the larger the range, the more likely the action is chosen by the GA. Actions that have parameters are split into different ranges, according to which parameters make sense. For example ModifyTargetVelocity is split into five different parts, with different percentages, namely 50, 70, 100, 130, 160. The range of integers assigned to these parts is different. A percentage setting of 100 for example has the largest integer range assigned. In Appendix , the probabilty of an actions can be seen. In Appendix ... the probabilty per actions of the parameters can be viewed.

These ranges were assigned based on intuition and trial and error. The encoding is visualized in 3.3.

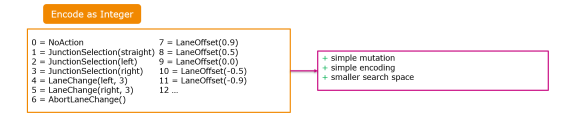


Figure 3.3: Integer

Dictionary The second encoding is much similar to the actual actions used in the simulations. Now, no translation is necessary anymore. During generation of the individuals, each action is again selected based on different probabilities assigned to actions, which again can be viewed in Appendix These probabilities are the same as for the integer encoding. However the difference is, in case an action has parameters that need to be chosen. For each parameter, a range and a randomness function was chosen. For example in case of the percenetage parameter in ModifyTargetVelocity, the values are selected from a GausDistribution, with mu= 100, sigma=25 and a range limit between 0 and 300.

Again, these probability functions with settings were assigned based on intuition as well as trial and error. Detailed information can be seen in Appendix...

Figure 3.4 shows a visualization.

```
Examples:
{"type": "LaneChange", "direction": "left", "duration": 3}
{"type": "JunctionSelection", "junction_selection_angle": "straight"}
{"type": "AbortLaneChange"}
{"type": "LaneOffset", "offset_percentage": "0.6"}

+ maps exactly to problem - larger search space + more control over parameters of actions lengths
```

Figure 3.4: Dictionary

3.2.2 Cost Function

Cost function is a bit difficult, as we are only using internal values. No ADAS/AD system is tested and we thus have to work with what we got. This is the code of the cost function:

```
1 SEPS_PER_SECOND = 100
2 # allow emergency breaks to last only 3 seconds
 MAX_DURATION = 3 * STEPS_PER_SECOND
3
4
5 cost = 0
6 duration_counter = 0
  for i in range(len(result["ego_emergency_stop"])):
8
      if not result["ego_emergency_stop"][i]:
          # base cost for no current emergency break
9
          cost = cost + 1
10
          duration_counter = 0
11
      else:
12
          if duration_counter > MAX_DURATION:
13
               # increase cost if emergency break max
14
                  duration is exceeded
               cost = cost + 10
15
          duration_counter += 1
16
17 return cost
```

result["ego_emergency_stop"] is a list with the length $100*simulation_duration_seconds$ (because 100hz). It contains a boolean per step, if the EGO vehicle has initiated an emergency stop.

Ref florian

It would have been interesting to not only test for emergency stops (which will make the NPCs try to get the EGO to hard break often) but also improve time to collison (TTC), as was done by . However by the time of starting the testing, no working TTC functionality was implemented. Thus, only the emergency break cost function is used by the GA to be optimized.

3.3 Behavior Tree

The Behavior Tree will controll the ego vehicle

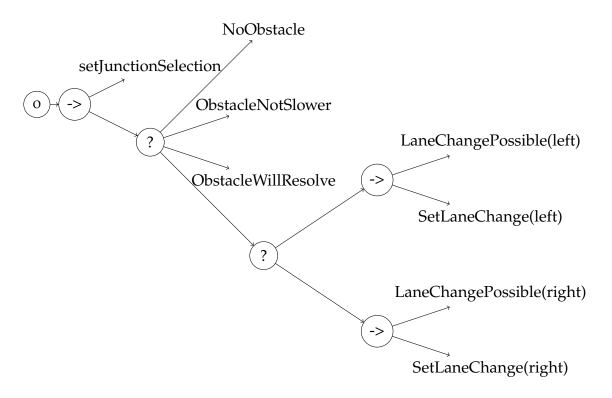


Figure 3.5: Used Behaviour Tree

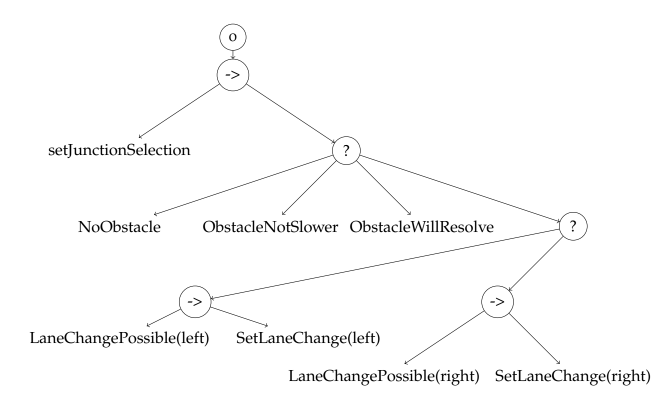


Figure 3.6: Used Behaviour Tree

4 Hyperparameter Tuning

In this chapter, we will incrementally move to an optimized Genetic Algorithm

4.1 No Free Lunch Theorem

No Free Lunch Theorem: The best hyperparameter settings of a Genetic Algorithm are very problem specific. K. De Jong, 2007, Dao, Abhary, and Marian, 2016

More ref

4.2 Map and Starting Scenario

The map is Town10 from Carla. It was chosen, because 1. its roads are self contained, 2. its not too big, yet still complex and 3. its supported by Carla and thus visualization looks better.

The Starting Scenario defines the number and type of all actors as well as their position. It needs to be created manually. Changing the scenario will have a great impact on the Genetic Algorithms performance. For time and complexity reasons, it was thus decided to first stick with one scenario and do all hyperparameter testing there. And finally test the performance for a handfull different scenarios.

4.3 Population

The number of Individuals is of high importance to a genetic algorithm, as has been explained in section 2.1. Especially considering the limed processing resources available, a suitable population size has to be found. On one hand, a population that is too low might result in less diverse runs of the genetic algorithm, on the other hand, if population is too high, the simulations will become too costly. Considering these points, the first step of the hyper parameter tuning was to find a suitable population size. In the next chapter 4.4, we will aim to improve the hyperparameter using a more robust approach.

In order to test for the best population size, the other hyperparameters have to be assumed using an educated guess. While reviewing the literature, trends of general settings for genetic algorithms can be found. However Mills, Filliben, and Haines, 2015 highlight the inconsistencies between findings, stating to have "uncovered conflicting opinions and evidence regarding key GA control parameters".

However Grefenstette, 1986 suggests, that "while it is possible to optimize GA control parameters, very good performance can be obtained with a range of GA control parameter settings." This is also complimented by findings from K. De Jong, 2007: "The key insight from such studies is the robustness of EAs with respect to their parameter settings. Getting "in the ball park" is generally sufficient for good EA performance. Stated another way, the EA parameter "sweet spot" is reasonably large and easy to find [18]. As a consequence most EAs today come with a default set of static parameter values that have been found to be quite robust in practice."

Chosing the right selection method is complicated as well, as discuees by K. De Jong, 2007: "One source of difficulty here is that selection pressure is not as easy to "parameterize" as population size. We have a number of families of selection procedures (e.g, tournament selection, truncation selection, fitness-proportional selection, etc.) to choose from and a considerable body of literature analyzing their differences (see, for example, [19] or [15]), but deciding which family to choose or even which member of a parameterized family is still quite difficult, particularly because of the interacting effects with population size [13]."

Looking at the literature might lead to hyperparameters are used that at least sufficient enough, to get an idea which range for population size is suitable. We will now look at different concrete hyperparameter suggestions from the literature.

4.3.1 Suggested hyperparameter from the literature

In an often cited thesis by K. A. De Jong, 1975, the following parameters have been suggested: GA(50, 0.6, 0.001, 1.0, 7, E) These suggested parameters have been used successfully by various different genetic algorithms Grefenstette, 1986.

An extensive study by Mills, Filliben, and Haines, 2015 which that took over "over 60 numerical optimization problems." into consideration found that "the most effective level settings found for each factor: population size = 200, selection method = SUS, elite selection percentage = 8%, reboot proportion = 0.4, number of crossover points = 3, mutation rate = adaptive and precision scaling = 1/2 as fine as specified by the user."

Grefenstette, 1986 claim that GA(30, 0.95, 0.01, 1.0, 1, E) and GA(80, 0.45, 0.01, 0.9, 1, P) produced the best results. They also advised against, a mutation rate of over 0.05, suggesting poor performance. Using a low mutation rate is also suggested by Whitley, 1994 and Jinghui Zhong et al., 2005. On the other hand, Boyabatli and Sabuncuoglu, 2004 state, that "Controversial to existing literature on GA, our computational results reveal that in the case of a dominant set of decision variable the crossover operator does not have a significant impact on the performance measures, whereas high mutation rates are more suitable for GA applications." Other paper also find a relatively high mutation rate useful. Almanee et al., 2021 uses genetic algorithms in a similar domain as this thesis. There, a Population of 50, crossover of 0.8 and mut of 0.2 was used. These used params are the same as the default params from deap (pop = 50 CXPB, MUTPB, NGEN = 0.5, 0.2, 4).

Srinivas and Patnaik, 1994 state, that for a higher population, cross: 0.6, mut: 0.001 and pop: 100 is a good starting point, while a lower population

Use best values also from:
Using genetic algorithms for automating automated lane-keeping system testing

Talk about rules (e.g. 1/n for mut rate...) look at: Parameter selection in genetic algorithms

cite https://deap.readtheo needs higher crossover and mutation rates like this cross: 0.9, mut: 0.01, pop: 30

Fazal et al., 2005 recommends a population size of 50, a scattered crossover function with a crossover probability of 0.5. The used selection function was tournament selection. Elite count was set to 5.

Dao, Abhary, and Marian, 2016 suggests a population size of 200, two point crossover with a crossover probability of 0.7. A Gausian Mutation Function as well as roulette selection and elite count set to 1.

A population size of 200 and roulette selection is used by Assistant Professor, Amity University, Jaipur, Rajasthan, India et al., 2019. Further, the elite count is set to 10. A heuristic crossover function with a crossover probability of 0.4 is also used.

4.3.2 results

This now leads to a difficult decision in choosing the right parameters. Based on the extensive research, we will compare population size of 32, 48, 64 and 96. We will compare the different crossover rates: 0.8 and 0.6. For mutation, 0.01 and 0.2 will be discussed. Further we will use tournament selection with 2 and 4. Each run will be executed 5 times to get rid of randomness and to make the results more robuts. We will run each simulation for 40 Generations.

In figure 4.2, the results per population are plotted. The line is corresponds to the mean, while the bars show the spread (min to max) of all 5 repetitions.

A high spread can be seen when looking at small population sizes. Considering these findings, a population size of 96 was chosen. While such a high value will result in a performance impact, it is important to keep the variation low.

Comparison of Population Size - mean											
Settings	Code	32	48	64	96						
C: 0.6, M: 0.01, TS: 2	A	3051	3016	2851	2871						
C: 0.6, M: 0.01, TS: 4	В	3111	3021	3079	2937						
C: 0.6, M: 0.2, TS: 2	C	3062	3010	3002	2831						
C: 0.6, M: 0.2, TS: 4	D	3020	2967	2891	2850						
C: 0.8, M: 0.01, TS: 2	E	3063	2892	2971	2916						
C: 0.8, M: 0.01, TS: 4	F	3052	3049	3054	2897						
C: 0.8, M: 0.2, TS: 2	G	3099	2940	2959	2869						
C: o.8, M: o.2, TS: 4	H	3058	3005	2794	2809						

Figure 4.1: List Settings per Population Size

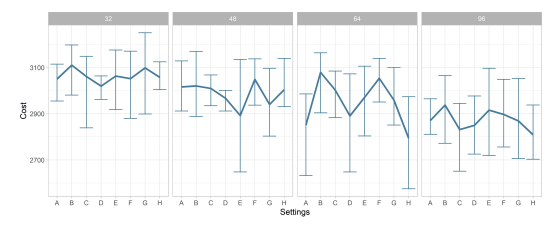


Figure 4.2: mean and error bars per population

4.4 Design of Experiment

Following the conclusion from the previous section 4.3, a population size of 96 will be used. Executing one run for 30 generations currently takes around 3:50 hours. Although two different workstations were available, the time required to execute the needed number of runs for these automated tests would exceed the available time budged. This is without considering a minimum required number of repetitions to remove randomness in the results.

In order to tune the hyperparameter of the genetic algorithm, various different strategies can be used. Using automated hyperparameter tuning approaches like "Grid Search", "Bayesian Optimization, "Simmulated Annealing or "Hyperband" might lead to good results with minimal effort (tuning hyperparameter of these search algorithms is still needed), however they require a high number of runs, which is not feasable.

find references

A different approach called "design of experiment" (DOE), also known as statistically designed experiments. DOE tries to find the causeand-effect relationship between the factors and the output of experiments. It uses factorial design where each experiment has factors, of which each consists of at least two settings, with the actual number of settings being called "levels" (Yang and El-Haik, 2009). Design of experiment needs manual expertise to define which factors are possibly of importance and which settings each factor should have, this is a drawback compared to automatic hyperparameter tuning.

"If the range of variable is too small, then we may miss lots of useful information. If the range is too large, then the extreme values might give infeasible experimental runs." (Yang and El-Haik, 2009)

Afterwards, main effects and interactions can be calculated to find the best settings per factor. It provides a graphical representation of these relationship by using interaction as well as main-effects charts. Using ANOVA (Analysis of Variance) it is possible to identify the significance of each factor and interaction, which enables the ranking of these factors. More details on these analysis tools will be provided in section 4.4.3.

A full factorial design will test over all possible combinations of the manually selected factor levels. Looking at the proposed factors in table 4.4.1, we would require 1024 runs¹, which not not feasible performance wise. A full factorial design has the drawback, that as the number if factors k gets increased, the number of needed experimental runs increases exponentially, thus resulting in lengthy experiments. Yang and El-Haik, 2009 state, that most of the results obtained by testing over all combinations are only

¹number of runs calculated using: https://datatab.net/statistics-calculator/design-of-experiments

used for estimating higher-order interactions, which are in most cases insignificant.

"Techniques such as fractional (or partial) factorial experiments are used to simplify the experiment. Fractional factorial experiments investigate only a fraction of all possible combinations. This approach saves considerable time and money but requires rigorous mathematical treatment, both in the design of the experiment and in the analysis of the results. (Roy, 1990)"

4.4.1 Taguchi Design

Various improvements to Design of experiment have been but forward by Dr. Genichi Taguchi, such as reducing the influence of uncontrollable (noise) factors on processes and products and reducing variability. Some of these methods evolve around Signal-to-noise (S/N) analysis and utilizing cost functions to "express predicted improvements from DOE results in terms of expected cost saving" (Roy, 1990). This master thesis will not discuss all of Taguchi's proposed considerations, for more detail Roy, 1990 as well as Yang and El-Haik, 2009 is highly recommended.

Using a Taguchi design for evaluating the best hyperparamter has been successfully performed by Dao, Abhary, and Marian, 2016 as well as Assistant Professor, Amity University, Jaipur, Rajasthan, India et al., 2019.

"There are many similarities between "regular" experimental design and Taguchi's experimental design. However, in a Taguchi experiment, only the main effects and two-factor interactions are considered. Higher-order interactions are assumed to be nonexistent. In addition, experimenters are asked to identify which interactions might be significant before conducting the experiment, through their knowledge of the subject matter." (Yang and El-Haik, 2009)

This masters thesis will mainly utilizes Taguchis orthogonal arrays (OAs), "which represent the smallest fractional factorials and are used for most common experiment designs." (Roy, 1990). This means, that only a fraction of combinations needs to be tested which drastically improves performance.

Each row of these matrices contains the factors of one experiment, while the columns correspond the factors Hamzaçebi, 2021.

Different orthogonal arrays have been proposed by Taguchi. The researcher has the responsibility to select an array based on the individual needs (Hamzaçebi, 2021). Using these orthogonal arrays instead of full factorial experiments will lead to needing a much smaller amount of simulation runs (in our case only 16 compared to 1024), while the latter "might not provide appreciably more useful information" Roy, 1990.

Definition orthogonal array An orthogonal array has multiple properties:

As has been stated, probably the biggest drawback of using Taguchi orthogonal arrays is on the one hand to increased manual labour and on the other hand the fact, that higher order interactions ignored.

Roy, 1990 explains why this might not be a big problem: "Generally speaking, OA experiments work well when there is minimal interaction among factors; that is, the factor influences on the measured quality objectives are independent of each other and are linear. In other words, when the outcome is directly proportional to the linear combination of individual factor main effects, OA design identifies the optimum condition and estimates performance at this condition accurately. If, however, the factors interact with each other and influence the outcome, there is still a good chance that the optimum condition will be identified accurately, but the estimate of performance at the optimum can be significantly off. The degree of inaccuracy in performance estimates will depend on the degree of complexity of interactions among all the factors."

This is complimented by Yang and El-Haik, 2009, who states, that: "During many years of applications of factorial design, people have found that higher-order interaction effects (i.e., interaction effects involving three or more factors) are very seldom significant. In most experimental case studies, only some main effects and two-factor interactions are significant."

Selection of orthogonal array When choosing a suitable Taguchi orthogonal array, we need to take various factors into account, which can make the

process tricky. According to Yang and El-Haik, 2009, we will have to follow a three step procedure:

- 1. Calculate the total degree of freedom (DOF).
- 2. Following two rules, standard orthogonal array should be selected:
 - a) Total DOF need to be smaller than the number of runs provided by the orthogonal array.
 - b) All required factor level combinations need to be accommodated by the orthogonal array.
- 3. Factors have to be assigned using these rules:
 - a) In case the factor level does not fit into the orthogonal array, methods such as column merging and dummy level can be used to modify the original array.
 - b) Using the linear graph and interaction table, interactions can be defined.
 - c) In case some columns are not assigned, its possible to keep these columns empty.

For this genetic algorithm, 7 factors (3 Factors of Level 4 and 4 Factors of Level 2) have been selected. Which factors to choose and with which level was done based on experience gained on section 4.3. When selecting levels, it is important to have them "as far away from either side of the current working condition as possible."(Roy, 1990) In table 4.4.1, every factor with corresponding levels has been listed,

Factors	Code	Level 1	Level 2	Level 3	Level 4
CrossoverType	A	one point	two point	uniform 0.1	uniform 0.5
CrossoverProp	В	0.2	0.5	0.8	0.9
MutationProp	C	0.01	0.1	0.3	0.5
ChromosomeType	D	Time	Time+NPC	-	-
GeneType	Е	int	dict	-	-
TournamentSize	F	2	4	-	_
IndMutationProp	G	0.1	0.5	-	_

Figure 4.3: List of Hyperparamters (Factors) matched to a Code and defined settings (Levels)

Using this table, we will now find the best standard orthogonal array in section 4.4.2. Before doing so, it is important to state, that Taguchi allows to test for possible (pre determined) two-level interactions (Yang and El-Haik, 2009). Analysing interactions comes at a cost of Degrees of freedom. If we look at the table, an interaction between ChromosomeType and GeneType might be of interest. Using the power of hindsight, we know, that a second two factor interaction is possible within our chosen array, thus we will have a look at the interaction between Tournament Size and IndMutationPropability as well.

4.4.2 Selection of a suitable standart orthogonal array

The total degree of freedom can be quickly calculated using the rules provided by Yang and El-Haik, 2009:

- 1. 1 DOF is always used for the overall mean.
- 2. Each factor has a DOF of NumberOfLevels 1.
- 3. Two-factor interactions use this equation to calculate DOF: $(n_{factor1} 1)(n_{factor2} 1)$ where n = number of levels.

This leads to the following calculation for the needed 3 Factors of Level 4 and 4 Factors of Level 2 as well as the two interactions between ChromosomeType-GeneType and TournamentSize-IndMutationProp:

$$DOF = 1 + 3 * (3 - 1) + 4 * (2 - 1) + 2 * (2 - 1) * (2 - 1)$$

$$= 13$$
(4.1)

A L_{16} array seems suitable to accommodate the required 13 DOF, which can be seen in 4.4.2.

This graph now needs to be fitted and modified to accommodate the needed factors. 4 Level Factors need additional space which will be generated using column merging, while interactions will need to be assigned as well. For this, either an interaction table or linear graphs of the L_{16} array can be used (NazanDanacioğlu, 2005). The linear graph approach is straight forward

		$L_{16}(2^{15})$													
NO.	1	2	3	4	5	6	7	8	9	10	11	12	13	14	15
1	1	1	1	1	1	1	1	1	1	1	1	1	1	1	1
2	1	1	1	1	1	1	1	2	2	2	2	2	2	2	2
3	1	1	1	2	2	2	2	1	1	1	1	2	2	2	2
4	1	1	1	2	2	2	2	2	2	2	2	1	1	1	1
5	1	2	1	1	1	2	2	1	1	2	2	1	1	2	2
6	1	2	2	1	1	2	2	2	2	1	1	2	2	1	1
7	1	2	2	2	2	1	1	1	1	2	2	2	2	1	1
8	1	2	2	2	2	1	1	2	2	1	1	1	1	2	2
9	2	1	2	1	2	1	2	1	2	1	2	1	2	1	2
10	2	1	2	1	2	1	2	2	1	2	1	2	1	2	1
11	2	1	2	2	1	2	1	1	2	1	2	2	1	2	1
12	2	1	2	2	1	2	1	2	1	2	1	1	2	1	2
13	2	2	1	1	2	2	1	1	2	2	1	1	2	2	1
14	2	2	1	1	2	2	1	2	1	1	2	2	1	1	2
15	2	2	1	2	1	1	2	1	2	2	1	2	1	1	2
16	2	2	1	2	1	1	2	2	1	1	2	1	2	2	1

Figure 4.4: $L_{16}(2^{15})$ Taguchi ortohogonal array taken from Roy, 1990

and will be selected. While there are multiple linear graphs for L_{16} array, 4.4.2 describes the graph which best fits the requirements from table 4.4.1. If no graph with the perfect fit is found, theses graphs can be modified as well, using rules described by NazanDanacioğlu, 2005.

"In each of Taguchi's orthogonal arrays, there are one or more accompanying linear graphs. A linear graph is used to illustrate the interaction relationships in the orthogonal array." Yang and El-Haik, 2009

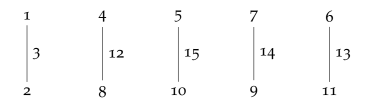


Figure 4.5: Linear Graph of $L_{16}(2^{15})$ taken from Yang and El-Haik, 2009

In a Taguchi linear graph, the nodes as well as the connections both represent columns in the orthogonal array. An interaction between to columns that are represented as nodes "comes out to" to the connecting line column Taguchi et al., 2005. This is useful for both analysing interactions between columns as well as combining (merging) interacting columns in case a higher factor is needed.

Column Merging A, B and C are both 4 level factors. The currently selected orthogonal only fits 2 level factors. Using column merging, it is possible to extend columns to accomodate higher order levels.

As calculated in 4.1, a four-level column requires three degrees of freedom, thus three two-level columns need to be merged. For column merging, it is required, that the to be merged columns are part of an interaction group (Yang and El-Haik, 2009).

So, 3 interaction 2-level columns need to first be selected. One column is discarded, the remain two columns need to be merged using the rules in tabular 4.6.

OLD C	COLUMN		NEW COLUMN
1	1	->	1
1	2	->	2
2	1	->	3
2	2	->	4

Figure 4.6: Rules taken from Roy, 1990

The four-level factor can then be assigned to this newly generated column. Because three four-level factors are needed for the current experiment, nine two-level columns need to be merged in total.

Assigning Interactions Interactions between two-level factors can be assigned using the linear graph as well. Here, select two connected nodes. The column describing their connection will subsequently contain the interaction (Taguchi et al., 2005).

An interaction between ChromosomeType and GeneType seems possible, thus D and E will be assigned to connected nodes in the linear graph. As we still have some unused space in the graph, we will also look at the interaction of TournamentSize and IndMutationProp (F and G). The resulting graph can be seen in 4.4.2.

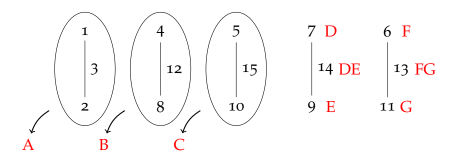


Figure 4.7: Modified Linear Graph to fit our needs

Combining columns 1 2 3 to A, 4 8 12 to B and 5 10 15 to C using rules defined by table 4.6 is done in 4.8.

Removing the old and inserting the new columns in the table and transcoding 7 to D, 9 to E, 14 to DE, 6 to F, 11 to G and 13 to FG results in the final table 4.9. This combinations table will subsequently be used as settings for the simulation runs.

4.4.3 Analysing the results

Table 4.9 can now can be used for running all the needed testcases (the interaction columns can be ignored until the evaluation). Transcoding all factors and levels to get the corresponding setting can be done using in the table from 4.4.1. We will repeat every setting 8 times to reduce randomness and gain information about variance. Running the Genetic Algorithm using these 16 different settings each repeated 8 times took 10 days on the two previously described workstations.

ref to section

The results are found in the appendix at 6.2.

NO.	1 2 3	4 8 12	5 10 15
1	1 1 > 1	1 1 > 1	1 1 > 1
2	1 1 > 1	1 2 > 2	1 2 > 2
3	1 1 > 1	2 1 > 3	2 1 > 3
4	1 1 > 1	2-2 > 4	2-2 > 4
5	1 2 > 2	1 1 > 1	1 2 > 2
6	1 2 > 2	1 2 > 2	1 1 > 1
7	1 2 > 2	2 1 > 3	2-2 > 4
8	1 2 > 2	2-2 > 4	2 1 > 3
9	2 1 > 3	1 1 > 1	2 1 > 3
10	2 1 > 3	1 2 > 2	2-2 > 4
11	2 1 > 3	2 1 > 3	1 1 > 1
12	2 2 > 3	2 2 > 4	1 2 > 2
13	2-2 > 4	1 1 > 1	2-2 > 4
14	2-2 > 4	1 2 > 2	2 1 > 3
15	2-2 > 4	2 1 > 3	1 2 > 2
16	2-2 > 4	2 2 > 4	1 1 > 1

Figure 4.8: Building 4 Level columns from 2 Level columns

Main-effects and interaction chart

Identifying the optimal conditions is done by analyzing the main effects per factor. Using them, it is possible to predict the factors, that lead to the best result Roy, 1990.

Yang and El-Haik, 2009 explains them well: "The main-effects chart is a plot of average responses at different levels of a factor versus the factor levels"

So, for every factor, sum up the mean of all results per level, then divide by the number of runs per level.

Example for D

The resulting main-effect charts can be seen here:

In case there is no interaction, the optimal setting is easily determined by using the main effects chart. Go over every factor in the chart and use the best level (in case of this experiment, the level with the lowest cost value).

NO.	A	В	С	D	Е	F	G	FG	DE
1	1	1	1	1	1	1	1	1	1
2	1	2	2	1	2	1	2	2	2
2 3 4 5 6	1	3	3	2	1	2	1	2	2
4	1	4	4	2	2	2	2	1	1
5	2	1	2	2	1	2	2	1	2
6	2	2	1	2	2	2	1	2	1
7 8	2	3	4	1	1	1	2	2	1
8	2	4	3	1	2	1	1	1	2
9	3	1	3	2	2	1	2	2	1
10	3	2	4	2	1	1	1	1	2 2
11	3	3	1	1	2	2	2	1	2
12	3	4	2	1	1	2	1	2	1
13	4	1	4	1	2	2	1	2	2
14	4	2	3	1	1	2	2	1	1
15	4	3	2	2	2	1	1	1	1
16	4	4	1	2	1	1	2	2	2

Figure 4.9: Final version of used Taguchi orthogonal array

If interactions exist, they might have an influence on the best settings and need to be investigated (Yang and El-Haik, 2009).

To investigate previously defined interactions, a test of interactions can be used. Their calculation is similar to calculating main effects.

Example for DE

If lines cross, an interaction between the two factors exists. The more parallel the lines are, the less likely an interaction. Magnitude of the angle between the lines corresponds to the degree of interaction presence, according to Roy, 1990.

ANOVA

Before choosing the best settings, ANOVA analysis (analysis of variance) should be performed on the results. Among other things, this will provide information on the magnitude of contribution of each main effects and

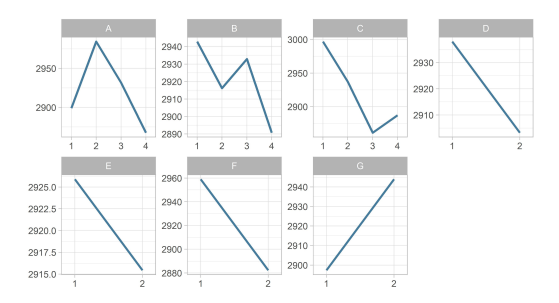


Figure 4.10: Main Effects

Summary after reading "Introduction into R"

interactions. The calculation of ANOVA is the same as for a classical design of experiment, according to Yang and El-Haik, 2009.

"In analysis of variance, mean squares are used in the F test to see if the corresponding effect is statistically significant." Yang and El-Haik, 2009 "F ratio is a better measure for relative performance" Yang and El-Haik, 2009

". The most commonly used criterion is to compare the p value with 0.05, or 5%, if p value is less than 0.05, then that effect is significant." Yang and El-Haik, 2009

"The variance ratio, commonly called the F statistic, is the ratio of variance due to the effect of a factor and variance due to the error term. (The F statistic is named after Sir Ronald A. Fisher.) This ratio is used to measure the significance of the factor under investigation with respect to the variance of all of the factors included in the error term. The F value obtained in the analysis is compared with a value from standard F-tables for a given statistical level of significance." Roy, 1990.

Due to our number of repetitions, the number of DOF increases according

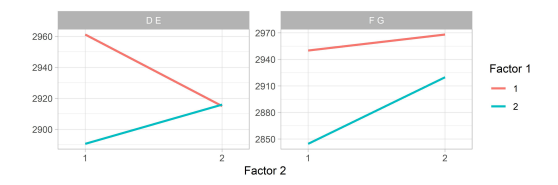


Figure 4.11: Test of interactions

to the following equation (taken from Roy, 1990):

$$DOF = totalNumberOfResults - 1$$

= $numberOfTrials * numberOfRepetitions - 1$ (4.2)
= $16 * 8 - 1 = 127$

Calculating ANOVA can be done simply be done using R, which will result in table 4.1.

	Df	Sum Sq	Mean Sq	F value	Pr(>F)
A	3	238901.41	79633.80	6.66	0.0004
В	3	49972.09	16657.36	1.39	0.2488
C	3	343169.03	114389.68	9.56	0.0000
D	1	38781.12	38781.12	3.24	0.0745
E	1	3507.03	3507.03	0.29	0.5893
F	1	189112.50	189112.50	15.81	0.0001
G	1	69751.13	69751.13	5.83	0.0174
D:E	1	41041.12	41041.12	3.43	0.0666
F:G	1	26277.78	26277.78	2.20	0.1411
Residuals	112	1339693.00	11961.54		

Table 4.1: ANOVA results

A, C, F and G have a relatively high F value, which suggests high influence on the model. ____

explain using R book The Multiple R-squared: 0.4275, Adjusted R-squared: 0.3509 ... both are bad.

We can also look at the percentage contribution of each factor, using the formula gathered by Yang and El-Haik, 2009:

$$SS_T = SS_A + SS_B + SS_C + \dots + SS_{error}$$
 (4.3)

$$contribution_A = SS_A/SS_T * 100 (4.4)$$

The percentage contribution is plotted in 4.4.3 (Sum of all factor contributions == Multiple R-squared in theory)

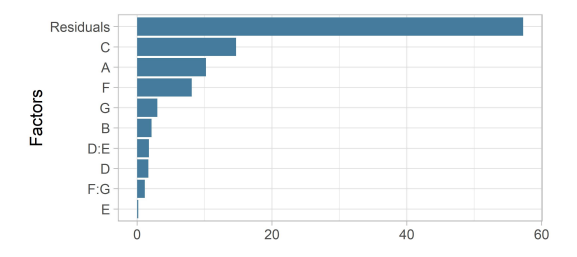


Figure 4.12: Percentage Contribution

We can clearly see a high contribution of the residuals (error), which is concerning.

Selection of optimal setting

When choosing the optimal setting, the first step is to look at the best main effects combination. For this experiment, the best combination would be the following: A4, B4, C3, D2, E2, F2, G1. Looking the ANOVA table, the interaction D:E has seams to have significance, especially compared to E. We will try to integrate this interaction. Compared to D:E, the second interaction (F:G) has a lower influence. This is also the case when looking at the individual factors F and G. The interaction F:G will thus not further be discussed.

The test of interaction in figure 4.4.3 suggest D2 and E1 as the best combination. This is optimal, as D2 is also suggested by the main effects. E1 is different to the suggested main effects, however its low F value in the anova table suggests low significance for using E2. Concluding this line of thought, the combination A4, B4, C3, D2, E1, F2, G1 looks to be optimal.

Optimum performance calculation Using optimal performance calculation 1. using only main effects or 2. using main effects with applied interaction can be applied. The equation provided by Roy, 1990.

Better bars

$$Y_{opt} = \bar{T} + (\bar{A}_4 - \bar{T}) + (\bar{B}_4 - \bar{T}) + (\bar{C}_3 - \bar{T}) + (\bar{D}_2 - \bar{T}) + (\bar{E}_2 - \bar{T}) + (\bar{F}_2 - \bar{T}) + (\bar{G}_1 - \bar{T})$$

$$= 2693.984$$
(4.5)

$$Y_{opt} = \bar{T} + (\bar{A}_4 - \bar{T}) + (\bar{B}_4 - \bar{T}) + (\bar{C}_3 - \bar{T}) + (\bar{D}_2 - \bar{T}) + (\bar{E}_1 - \bar{T}) + ([\bar{D}\bar{x}E]_2 - \bar{T}) + (\bar{F}_2 - \bar{T}) + (\bar{G}_1 - \bar{T}) + (\bar{G$$

Using the interaction D:E, the performance estimation improves from 2693.984 to 2686.547. Considering this result, interaction will be taken into account and the optimized settings are as follows: CrossoverType: Uniform 0.5, CrossoverPropability: 0.9, MutationPropability: 0.3, ChromosomeType: Time+NPC, GeneType: integer encoding, TournamentSize: 4 and Individual-MutationPropability: 0.1.

signal-to-noise (S/N) As previously discussed when looking at the anova model, the error is very high, which suggests high randomness. Taguchi recommends using signal-to-noise (S/N) ratio to reduce the variability, as using only the mean of the results does not take the variation into account (Roy, 1990). The greater the signal-to-noise ratio, the smaller the variance. Roy, 1990 further states, that the "use of the S/N ratio offers an objective way to look at the two characteristics (consistency and average value) together."

When using S/N, todo: talk about equation.

Afterwards, generating main effects and anova table is the same as using the mean. However the DOF calcualtion done in equation 4.2 is no longer valid, as repetitions get combined to 1 value per run. So, the DOF for anova changes to the following equation in 4.7 according to Roy, 1990.

$$DOF = totalNumberOfResults - 1$$

= $numberOfTrials * 1 - 1$
= $16 - 1 = 15$ (4.7)

Find out why 15 DOF is not enough

This is not enough residuals for generating the F value in anova. Thus in order to reduce the current DOF, the anova table was generated without the interaction F:G considered, which can be seen in 4.2.

	Df	Sum Sq	Mean Sq	F value	Pr(>F)
A	3	0.26	0.09	3.01	0.3953
В	3	0.05	0.02	0.60	0.7139
C	3	0.38	0.13	4.44	0.3326
D	1	0.04	0.04	1.48	0.4378
E	1	0.00	0.00	0.12	0.7845
F	1	0.21	0.21	7.35	0.2250
G	1	0.08	0.08	2.80	0.3429
D:E	1	0.04	0.04	1.56	0.4296
Residuals	1	0.03	0.03		

Table 4.2: S/N ANOVA results

When looking at the p values of this table, it is very obvious that no factor can discard the null - hypothesis, which states that a factor has no significant effect. Considering this, it was deemed to be not necessary to perform further investigations. Somehow argument, that having less variablity is only to an extend important. It is always possible to restart a simulation, if the variability is not too large, it is important that the results have a good mean overall. Less variability is in producing products much more important.

Elite Although the optimal hyperparameter setting will be discussed in chapter 5, a problem was obvious when analyzing a run using the optimzed GA. Figure 4.4.3 shows for a few selected repetitions the best individual cost per generation.

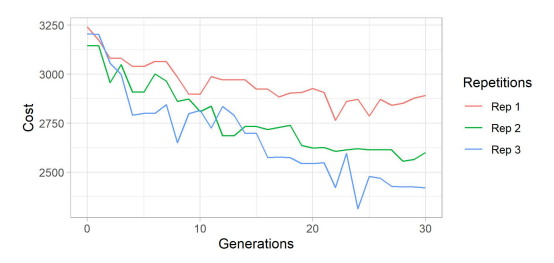


Figure 4.13: Genetic Algorithm without Elite

The lines show that setbacks in the optimal cost between two generations happens frequenctly. In order to mitigate this problem, it was decided to implement elite selection with a size of 2. This means, that per generation, the two best individuals are copied into the next generation without modifications, which makes worse performance between generations not possible. It is important to note, that the two best individuals can still be selected by tournament selection for modification, its just that a copy of them is saved. Figure ?? shows the effect of these changes.

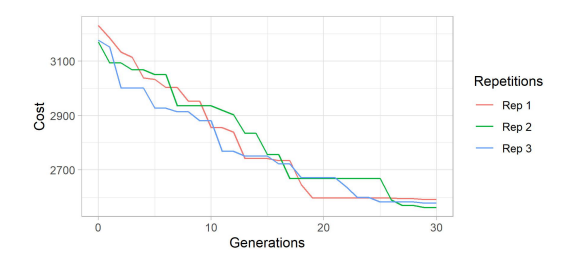


Figure 4.14: Genetic Algorithm with Elite

Comparing the 10 repetitions also provides a clear picture in figure 4.4.3. It is thus concluded that the slightly modified version of the optimized Ga now using Elite of 2 will be used for chapter 5.

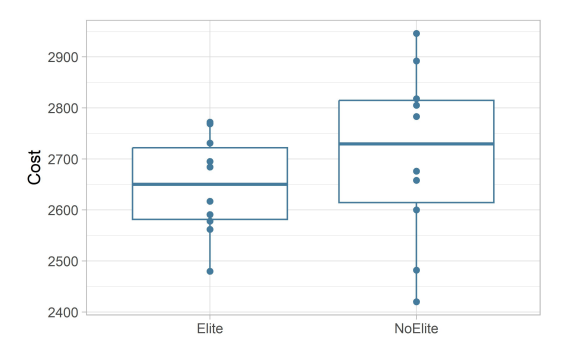


Figure 4.15: Comparison Elite vs No Elite

5 Evaluation

in this chapter, we evaluate and compare various different settings

5.1 Comparison with random and default ga Values

Compare with results using boxplot, also compare (average) diversity

Different Algorithms: 1. Optimized GA 2. Best GA setting from Population
3. Random

5.2 Generalization on different start scenarios

scenario 1: default : 9v 5p scenario 2: 9v 5p scenario 3: 5v 3p scenario 4: 18v 10p

6 Conclusion

6.1 Future Work

6.1.1 Oracles

While not implemented here, Oracles are needed in order to get a list of good scenarios.

6.2 Final words

Appendix

Appendix A.

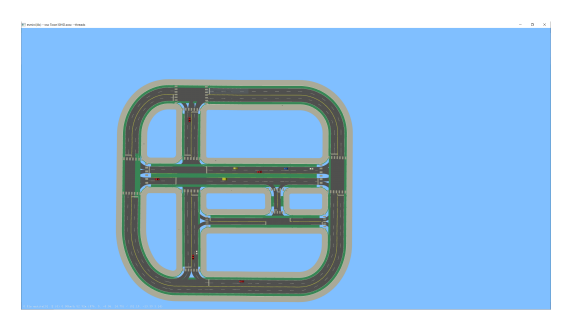


Figure 1: Start scenario 1

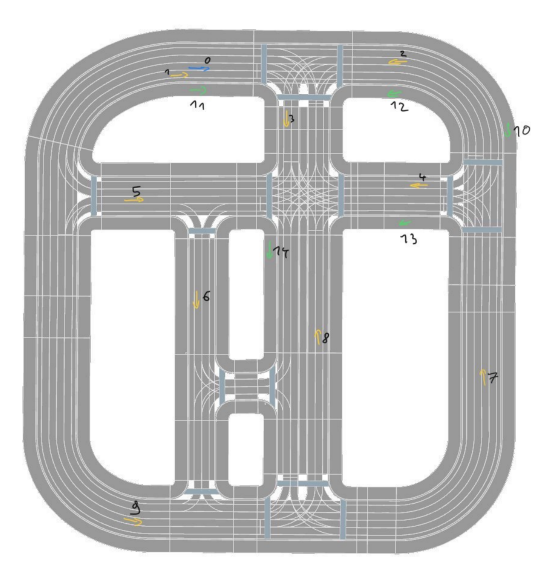


Figure 2: Start scenario 2

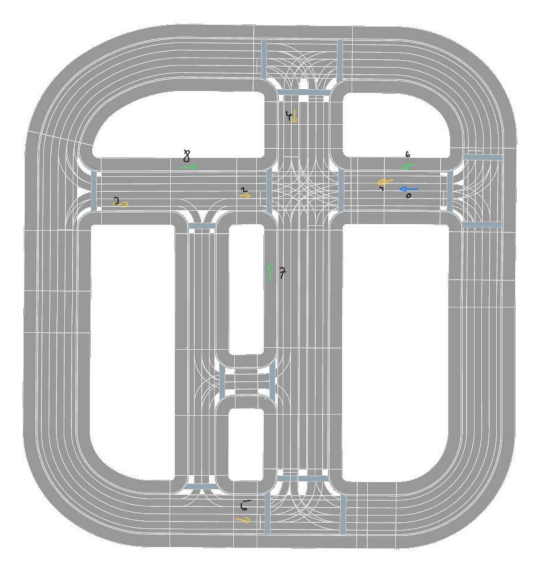


Figure 3: Start scenario 3

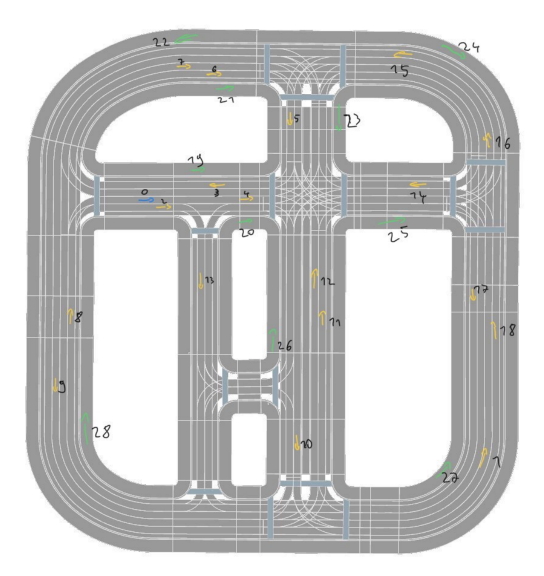


Figure 4: Start scenario 4

Appendix B.

NO.	rep1	rep2	rep3	rep4	rep5	rep6	rep7	rep8
1	3124	3110	3025	3077	3068	2925	3106	3105
2	2694	2980	3025	2996	3037	2921	3068	2900
3	2638	2711	2624	2856	2623	2832	2904	2778
4	2735	2805	2851	2965	2876	2703	2848	2858
5	3074	2955	3045	3080	3120	2971	2895	2798
6	2974	2979	2929	2941	2952	2936	3139	2953
7	3108	3099	3049	3020	3092	3057	3090	2895
8	2840	2931	2921	2921	2957	2997	2889	2895
9	3007	2995	2983	3009	2847	2996	2734	2927
10	2916	2828	3013	2787	2818	2926	3034	2822
11	3007	2879	3090	3033	2906	2981	3109	3104
12	2946	3016	2790	2917	2904	2983	2898	2606
13	2378	2712	2906	2800	2912	2795	2860	2834
14	2895	2760	2750	2849	2542	2997	2965	2991
15	2842	3065	3050	2779	2862	2923	2955	2892
16	3065	2834	2643	3056	3051	3011	2828	2963

Figure 1: List of results

Appendix C.

Insert information of Gene action probabitlities

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