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# **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

Master's degree programme: Telematik

submitted to

**Graz University of Technology**

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Graz, November 2013

This document is set in Palatino, compiled with pdf $\text{\LaTeX}2\text{e}$  and Biber.

The  $\text{\LaTeX}$  template from Karl Voit is based on KOMA script and can be found online: <https://github.com/novoid/LaTeX-KOMA-template>

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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

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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.

better and longer introduction

## 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

*Is it possible to improve the performance of a genetic algorithm using the taguchi method?*



# 2 Foundations

## 2.1 ADAS/AD

probably  
talk about  
ADAS/AD  
here...

## 2.2 Genetic Algorithm

Pioneered by Holland, 1992, genetic algorithms (GAs) imitate the process of natural selection and the darwinian principle called "survival of the fittest". Genetic algorithms are a subclass of evolutionary algorithms and explore the solution space using a population of individuals (Mills, Filliben, and Haines, 2015). Each individual contains one chromosome, which serves as a candidate solution. Using genetic operations, the individuals mate among themselves and mutate independently.

The selection of an individual is determined by a fitness function, which defines the search problem. Each individual has a fitness value corresponding to the performance of its chromosome (Majumdar and Ghosh, 2015). Individuals that do poorly on the given problem die out, while individuals deemed "strong" propagate. Genetic algorithms optimize iteratively, with each iteration referred to as a generation. Chromosomes consist of individual genes, which form a solution to the search problem. Through successive generations using genetic operations, gene values and position will be optimized, resulting in progressively improved solutions (Srinivas and Patnaik, 1994).

Generational genetic algorithms, where the entire population is replaced each generation, will be utilized in this master's thesis. In contrast, steady

## 2 Foundations

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state genetic algorithms only replace a small fraction of the population at a time (Srinivas and Patnaik, 1994).

This search method on the basis of biological principles allows for a global and dispersed search through its population, which avoids various shortcomings of local search techniques (Grefenstette, 1986). Especially on challenging search spaces with multiple local optima, a GA is less prone to get stuck on a substandard solution (Katoch, Chauhan, and Kumar, 2021, Xia et al., 2019, Majumdar and Ghosh, 2015). Genetic algorithms offer advantages for complex optimization and non-deterministic polynomial (NP-hard) problems (Hussain and Muhammad, 2020).

*"GAs can find good solutions within a large, ill-defined search space, and can be readily adapted to a wide variety of search problems" (Mills, Filliben, and Haines, 2015)*

Grefenstette, 1986 further emphasizes its ability to outperform gradient techniques on difficult problems with high-dimensional, noisy or discontinuous fitness functions by efficiently exploiting a "relatively simple selection mechanism." The main advantage of a GA in tackling these problems comes from its capability to explore the search space using its entire population (Hussain and Muhammad, 2020). The "high implicit parallelism" makes genetic algorithms "numerically very efficient" (Marsili Libelli and Alba, 2000). A basic GA proposed by Holland, 1992 can be defined as follows:

```
1    simple_genetic_algorithm()
2    {
3        initialize_population();
4        evaluate_population();
5        for(int i = 0; i < num_of_generations; i++)
6        {
7            select_individuals_for_next_population();
8            perform_crossover();
9            perform_mutation();
10           evaluate_population();
11        }
12    }
```

The population is initialized randomly, and its individuals are subsequently evaluated using the fitness function. The following steps are iteratively repeated until some stopping criterium is triggered.

The individuals are chosen using a selection operation, which takes their fitness value under consideration. A crossover rate subsequently determines which individuals mate using a crossover function. This operation serves the purpose of exchanging information between the chromosomes. In order to add variation and diversity the resulting crossover offspring undergoes small changes using a mutation function. A mutation rate decides on which individuals this operation is applied. The newly generated population will then get evaluated for the next iteration. Different stopping methods like time limit, fitness limit, minimum convergence rate or maximum number of generations are available (Majumdar and Ghosh, 2015).

As discussed by Hussain and Muhammad, 2020, genetic algorithms suffer from an exploration vs exploitation dilemma. If the algorithm converges too quickly to a solution, most of the search space might not have yet been explored, thus increasing the probability of getting stuck in a local optimum. In contrast to that, a low convergence rate might be time consuming and result in inefficient utilization of computational resources.

*“Finding a balance between exploration and exploitation has been a difficult-to-achieve goal from the beginning.” (K. De Jong, 2007)*

Due to no real consensus on the best control parameter settings, their optimal selection proves to be difficult.

### 2.2.1 Chromosomes and Genes

The encoding of genes proved to be, besides the hyperparameter tuning, the main challenge of setting up a genetic algorithm pipeline. A simple and commonly used representation for genes is binary encoding. As suggested by its name, a gene can take on the form of 0 or 1. Further common encodings are octal and hexadecimal representations (Srinivas and Patnaik, 1994, Katoch, Chauhan, and Kumar, 2021).

explain genes  
and chromo-  
somes a bit  
more

Srinivas and Patnaik, 1994 suggest to use integer representations in case a optimization problem has real-valued continuous variables, where the objects are linearly mapped to integers defined in a specific range. It is possible to again represent these integers as binary encodings. Various other encodings are available, often very problem specific. For example tree encodings allows for genes to represent programming functions, leading to a subcategory of genetic algorithms called "genetic programming" (Katoch, Chauhan, and Kumar, 2021).

In case no encodings fits the given problem, a custom encoding can be used. Here the genetic operation crossover and mutation might have to be tailored accordingly.

### 2.2.2 Population Size

Controlling the population size, which defines the number of individuals per generation, has a direct impact on the performance of a genetic algorithm. Research seams to be in agreement, that a small population leads to less diverse individuals and might provide an insufficient sample size, which can lead to a premature convergence to a local optimum. A large population size will allow the GA to perform a more informed search. However computation time will suffer due to the larger number of individuals per generation and might lead to slower convergence to an optimum (Grefenstette, 1986, Katoch, Chauhan, and Kumar, 2021, K. De Jong, 2007).

Increasing the population size will increase the degree of parallelism in the genetic algorithm, as each individual represents one search point (Mills, Filliben, and Haines, 2015).

### 2.2.3 Selection

The selection operator chooses two parents for crossover and mutation operations until the list of offsprings has reached the desired population size. Individuals are selected based on their fitness value, ensuring their

## 2.2 Genetic Algorithm

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increased representation from generation to generation. Weak solutions will be discarded over time (Srinivas and Patnaik, 1994).

The selection algorithm needs to satisfy two requirements. On the one hand, high selection pressure will lead to decreased diversity in the population resulting in premature convergence (Katoch, Chauhan, and Kumar, 2021). The algorithm will subsequently behave more like a local search method, a hill-climber or a greedy algorithm (K. De Jong, 2007). The initial low average fitness value of the population will in combination with a few good performing individuals lead to them overtaking the population, drastically reducing diversity. On the other hand, a low selection pressure will struggle to converge to an optimum.

Different selection methods like Stochastic Uniform Remainder, Random Selection, Rank Selection, Roulette Wheel Selection and Tournament Selection exist (Majumdar and Ghosh, 2015). According to Hussain and Muhammad, 2020, who provide a extensive comparison of different selection techniques, the choice of selection methods highly affects the performance of the genetic algorithm.

Roulette wheel selection is a popular mechanism, where each individual corresponds to an area on a roulette wheel, based on its fitness value (Holland, 1992). Grefenstette, 1986 points out the scaling problem as its major drawback. As the algorithm progresses, its fitness variance to mean ratio becomes increasingly small, leading to low selection pressure. This problem can be mitigated by using ranks instead of the fitness value (Katoch, Chauhan, and Kumar, 2021). Individuals will be sorted based on their performance and their relative rank is inserted into the roulette wheel.

Tournament selection is a popular alternative to roulette wheel selection. First, a tournament size  $t$  has to be chosen. Until the desired number of offsprings is achieved,  $t$  individuals will be chosen at random from the population. Only from this smaller list, the best individual is selected. A popular tournament size is 2, larger sizes might enhance competition among individuals, however they can also has an impact on the diversity of the population (Hussain and Muhammad, 2020). In a comparison of different selection methods by Jinghui Zhong et al., 2005, tournament selection was deemed to be the most performant, converging more quickly then for example roulette wheel selection.

Elite selection is an additional method which can enhance a selection operation, proposed by K. A. De Jong, 1975. The best n individuals are automatically chosen to go into the next generation. Elite selection can help reduce variance, as the highest performing individuals can not be removed by randomness.

### 2.2.4 Crossover

The crossover operation serves as the mating process between two individuals (chosen from the population by the selection operation). Its resulting offspring is a combination of both parent genes. Grefenstette, 1986 highlights, that first, “it provides new points for further testing within the hyperplanes already represented in the population” and second, that “crossover introduces representatives of new hyperplanes into the population”.

Different types of crossovers can be implemented, the most simple approach being the single-point crossover. Here a point is chosen at random along the length of the chromosome. The two parents will swap their genetic information at this position, generating two new offsprings (Katoch, Chauhan, and Kumar, 2021). Extension are the two- or multipoint crossover operations, where chromosomes are divided by k segments, which will get exchanged. They eliminate a disadvantage of “the single-point crossover bias toward bits at the ends of strings” (Srinivas and Patnaik, 1994).

A second crossover operation is the uniform crossover where gene swaps between parents are randomly decided by a given probability, independent to the exchange of other genes (Katoch, Chauhan, and Kumar, 2021).

Srinivas and Patnaik, 1994 proposed the notion of positional and distributional bias. If a crossover operation has a positional bias, the probability of a bit to be swapped depends mainly on its position. Examples are the mentioned single and multipoint crossover operators. The uniform crossover can be found at the other end of the spectrum, as it has a maximal distributional bias, completely disregarding any positional information. Ignoring the gene positions results in a higher disruptiveness, which has potential drawbacks. However it will be more exploratory in homogeneous populations where k-point crossovers might struggle. Srinivas and Patnaik, 1994 suggest that

uniform crossover is more useful in small populations. Larger populations inherently are more diverse, making k-point crossover more suitable.

Various other crossover operations are analysed in paper [here](#).

Not only the crossover type has to be chosen, a crossover rate also needs a declaration. This value defines, how likely a single crossover operation is to be applied on the selected individuals. Choosing a suitable crossover probability is again influenced by the exploration-exploitation balance. Higher crossover rates will introduce new structures quickly into the population, however good sequences of genes might get disrupted. A low crossover rate will result in a low exploration which leads to stagnation (Grefenstette, 1986).

Find good research paper, maybe Katoch, Chauhan, and Kumar, 2021

### 2.2.5 Mutation

The mutation operation is applied after crossover as a means to maintain the diversity of the gene pool. Through small random changes, the variability of the population increases. Each individual can be exposed to mutation, irrespective of their fitness value. A mutation rate chooses the individuals to be mutated. An additional individual mutation rate selects which specific genes of the chosen chromosome are mutated (Srinivas and Patnaik, 1994). Depending on the gene encoding, the mutation operation might vary. In case of binary gene encoding, mutation will only flip certain bits. For custom encodings, the mutation operation needs to be tailored accordingly.

*"If the mutation is not considered during evolution, then there will be no new information available for evolution." (Katoch, Chauhan, and Kumar, 2021)*

The settings for mutation rates again needs to be chosen carefully. If they are set too high, the genetic algorithm might transform into random testing. In case they are set too low, the population will not be able to maintain diversity as no new genetic material is reintroduced. (Klampfl, Klück, and Wotawa, 2023, Grefenstette, 1986).

**insert papers**

### 2.2.6 Adaptive Control Parameters

Various literature suggests adaptively controlling selection methods, crossover rates and mutation rates over the duration of a genetic algorithm ( ... ). The main goal is to mitigate the previously mentioned exploration-exploitation problem. In the beginning, exploration is desired, thus low selection pressure as well as high crossover and mutation rates are required. However to the end of the genetic algorithms duration, convergence to an optimum is preferred, meaning more "elite" selection methods as well as less variation due to crossover and mutation (Srinivas and Patnaik, 1994).

Marsili Libelli and Alba, 2000 proposed a method, where different mutation rates are used depending on the fitness of the individual. They claim, that a higher mutation probability for low fitness population results in a more efficient search. Hussain and Muhammad, 2020 argue for a selection operation, where initially a low selection pressure is applied while increasing it to the end, stating that this approach will help mitigate both mentioned competing criteria, namely premature convergence as well as slow convergence.

Contrary, K. De Jong, 2007 is more critical on adaptive control parameters. Although adaptive control parameter might improve the performance of a genetic algorithm for specific problems, genetic algorithms are already quite robust in practice. Adding more tunable parameters will not make the task of the researcher easier.

*"The purist might argue that inventing feedback control procedures for EAs is a good example of over-engineering an already sophisticated adaptive system." (K. De Jong, 2007)*

K. De Jong, 2007 adds, that "after more than 30 years of experimenting with dynamic parameter setting strategies, is that, with one exception, none of them are used routinely in every day practice. The one exception are the strategies used by the ES community for mutation step size adaptation."

In order to keep the number of tunable parameters low, the approach implemented by this master's thesis will not utilize adaptive control parameters.

## 2.3 Behaviour Tree

A behaviour tree is a decision tree.

insert a short introduction to BT



# **3 Implementation**

## **3.1 Traffic Manager**

A custom developed Traffic Manger will be used to simulate an traffic environment, which can subsequently be controlled by the Genetic Algorithm. The Traffic Manager is not part of the master's thesis and will be regarded as a "blackbox", only receiving a brief introduction. Generally speaking, it will simulate traffic using a start scenario, in which the positions and types of actors (vehicles and pedestrians) are defined. A simulation always consist of at least one EGO vehicle. Additionally any number of NPCs can be added.

While the NPCs are only controlled by the Traffic Manager, the EGO vehicle can be either partly or completely supervised by an ADAS/AD function. The goal of the genetic algorithm is subsequently to test this function for safety, comfort etc... by controlling the NPCs in order to generate critical scenarios.

For all simulations done by this thesis, the Traffic Manager was set to 100Hz and the simulation duration set to 35 seconds.

### **3.1.1 Action Interface**

In order to control the behaviour of all actors inside the simulation, actions can be requested over the Action Interface, which is provided by the Traffic Manager. An action will initiate a certain behaviour from an actor and can

### 3 Implementation

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be set at any timestep<sup>1</sup>. Pedestrians and vehicles have a different set of possible actions. If no action is applied, the actor will behave in a normal manner inside the simulation. This means that the actor will follow along its given path until a new action changes its behaviour.

The following list shows all actions provided by the traffic manager which were available for the genetic algorithm at the time of this master's thesis.

- JunctionSelection
  - vehicle\_id: int, step: int, junction\_selection\_angle (radians): float
  - Vehicles will chose which direction to take at junctions based on this angle.
- LaneChange
  - vehicle\_id: int, step: int, direction: float, distance: float, delay (optional): float
  - Initiates a lane-change based on its given parameters.
- AbortLaneChange
  - vehicle\_id: int, step: int
  - Will abort a current lane-change.
- ModifyTargetVelocity
  - vehicle\_id: int, step: int, percentage: float
  - Modifies the internal target velocity of the vehicle by a percentage. If set to 0, the vehicle will stop.
- TurnHeading
  - pedestrian\_id: int, step: int
  - The pedestrian will turn 180 degrees and walk in the opposite direction
- CrossRoad
  - pedestrian\_id: int, step: int
  - The pedestrian will cross the road immediately.
- CrossAtCrosswalk

---

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

### 3.2 Genetic Algorithm

- pedestrian\_id: int, step: int
- The pedestrian will cross the road at the next crosswalk.

The genetic algorithm will control only NPCs, a behaviour tree is used for the EGO vehicle itself. Both methods use the Action Interface to control their respective actors. An overview is provided in figure 3.1.



Figure 3.1: Action Interface Structure

## 3.2 Genetic Algorithm

The genetic algorithm will search for sequences of actions that will result in the most interesting scenarios according to its cost function. For implementing the genetic algorithm, the Python library DEAP<sup>2</sup> was chosen. It is a popular tool in academia and allows for high customisability. The algorithm has full access over the setting of actions for all NPCs. Searching for sequences of actions, the GA tries to optimize a cost function.

A few default settings for the genetic algorithm were chosen. It was decided for example, that the genetic algorithm will be able to 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. If the GA decides to not set an action, "NoAction" will be used as a placeholder.

<sup>2</sup><https://deap.readthedocs.io/en/master>

### 3.2.1 Maximum Number of Generations

A fixed maximum number of generation is a commonly used stopping criteria. While more complex methods like a "adaptive convergence rate" might lead to better performance, it was decided that the additional complexity of having to choose a suitable convergence rate outweighs its potential benefits. Especially considering that a genetic algorithm already has a lot of different control parameters that need tuning.

Performance is always a big consideration in this thesis, thus a lower maximum number of generations is preferred. During testing, a generation size of 30 was almost always sufficient and will be used in all of the coming testing.

### 3.2.2 Encoding

When implementing a genetic algorithm, it is necessary to use an encoding that fits to the problem at hand. Here, each chromosome is required to include all actions of one whole simulation. Due to the deterministic nature of the Traffic Manager, it is sufficient to define a simulation by only the action sequence from a chromosome and the initial state defining start scenario.

The different encodings presented in section 2.2.1 were not directly applicable to the given problem. A custom encoding for both chromosomes and genes needed to be generated.

#### Chromosome

Two different encodings proved to be a good fit. Both use the position of a gene inside the chromosome to define the time step of an action. Which chromosome encoding is the most adequate will be discussed and tested in chapter 4.

### 3.2 Genetic Algorithm

**Time** The first encoding is called Time. Each gene corresponds to 1 time step (so 1 gene per every 0.5 seconds), and needs to contain all actions of all NPCs at this particular moment. A gene is a list of the size of the number of NPCs. Each object in this lists equals an action. The index of an action inside a gene corresponds to the NPC ID. For example, an action positioned in the gene at index 2, will be applied to NPC 2. If the gene is at position 6 in the chromosome, the action will be set at second 3.0 in the simulation. A visualization is seen in figure 3.2.



Figure 3.2: Time

Given the previously stated simulation time of 35 seconds, each chromosome has a length of  $35 * 2 = 70$  genes. Crossover can now only move all actions of a time step at once, modifications in the timeline between actions of the same time step will only happen due to a mutation operation.

**Time+NPC** The second encoding is called Time+NPC. Here, one gene corresponds to exactly one action. In order to not loose required information, the ID of each NPC now needs to be encoded into the genes position of its chromosome. One might visualize this as unfolding the matrix build by the genes and chromosome in encoding Time into a linear array. Each actors actions will be listed one after another. Now, each gene has a length of 1 and each chromosome has a length of  $35 * 2 * \text{number\_of\_actors}$ , which makes them much longer compared to the previous encoding. Figure 3.3 provides a visualization. In this example, a gene at position 13 corresponds to an action for NPC 2 at time step 3 (which is at 1.5 seconds in the simulation).

However for this encoding to make sense, the crossover operations OnePoint and TwoPoint had to be modified. For each NPC, these crossover opera-

provide better explanation for crossover

### 3 Implementation

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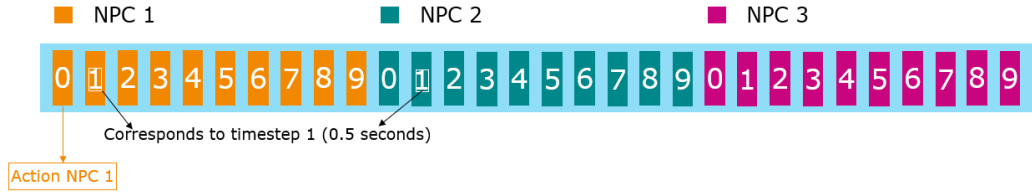


Figure 3.3: Time + NPC

tions are applied individually. If this was not the case, OnePoint crossover would introduce changes to actions of only one NPC. Because the different crossover points are independently chosen, the position of the crossing differs between the NPCs. This fact allows the crossover operation to break up actions at the same time step, which was not possible with the Time encoding.

#### Gene

Two different encodings for genes were implemented. A gene always consists of a list, which depending on the chromosome type either has a length of *number\_of\_actors* (in case of Time) or of length 1 (in case of Time+NPC). The following two encodings thus explain the type of object in these lists.

Bad explanation  
for integer en-  
coding, needs  
rewriting

ref appendix

**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. These ranges were assigned based on intuition and trial and error. In Appendix these probabilities can be viewed.

Actions have parameters split into different ranges, according to which values make sense. For example ModifyTargetVelocity has five different percentages assigned, namely 50, 70, 100, 130, 160. The range of integers assigned to each of these parts is different as well. A percentage setting of 100 for example has the largest integer range assigned.

### 3.2 Genetic Algorithm

Actions will be split into 1 or more parts that get an range of integer assigned. Combing these ranges results in a continuos list of integers. The encoding is visualized in 3.4

Encode as Integer	
0 = NoAction	4 = LaneChange(left, 3)
1 = JunctionSelection(straight)	5 = LaneChange(right, 3)
2 = JunctionSelection(left)	6 = AbortLaneChange()
3 = JunctionSelection(right)	7 ...

Figure 3.4: Integer

**Dictionary** The second encoding for genes corresponds exactly to the actual actions used in the simulations, requiring no translation. Each action is again selected based on the same probabilities as for the integer encoding. For actions without parameters, the different encoding will make no difference. However actions with parameters no longer are split into different discrete settings. Now, each parameter is chosen by a randomness function, which is individually selected per action. For example in case of the percentage parameter in ModifyTargetVelocity, the values are selected from a gaussian distribution with mu=100, sigma=25 and a range limit between 0 and 300. Figure 3.5 shows a visualization. Again, these probability functions with settings were assigned based on intuition as well as trial and error. Detailed information can be seen in Appendix .

Also bad explanation for dictionary encoding, needs rewriting

ref appendix

ref appendix

Compared to Integer encoding, this method allows for much higher granularity when selecting parameters. If the performance of a genetic algorithm differs between these settings will be investigated in chapter 4.

#### Encode as dictionary

```
{"type": "LaneChange", "direction": "left", "duration": 3}  
 {"type": "JunctionSelection", "junction_selection_angle": "straight"}  
 {"type": "AbortLaneChange"}  
 {"type": "ModifyTargetVelocity", "percentage": "133.4"}
```

Figure 3.5: Dictionary

#### 3.2.3 Cost Function

This master's thesis only utilized internal values from the Traffic Manager for the cost function. Namely if the ego vehicle has to initiate an emergency break. The emergency break values are booleans stored in a list called `result["ego_emergency_stop"]` which has a length of  $100 * simulation\_duration\_seconds$  (because of 100hz).

In case an emergency break lasts longer than 3 seconds, a penalty will be applied. This penalty was introduced as it mitigated results, where the ego vehicle is constantly brake-checked<sup>3</sup> by the leading vehicle. The following code snipped shows the cost calculation:

explain code a bit

```
1 SEPS_PER_SECOND = 100  
2 # allow emergency breaks to last only 3 seconds  
3 MAX_DURATION = 3 * STEPS_PER_SECOND  
4  
5 cost = 0  
6 duration_counter = 0  
7 for i in range(len(result["ego_emergency_stop"])):  
8     if not result["ego_emergency_stop"][i]:  
9         # base cost in case of no current emergency break  
10        cost = cost + 1  
11        duration_counter = 0
```

<sup>3</sup>"the unsafe action of applying a car's brakes to dissuade a driver who is following too closely" according to <https://www.dictionary.com/e/slang/brake-check/>

```

12     else:
13         if duration_counter > MAX_DURATION:
14             # increase cost if emergency break max
15             # duration is exceeded
16             cost = cost + 10
17             duration_counter += 1
18
19     return cost

```

The resulting cost values however are not easy to interpret, thus these numbers are modified using equation 3.1.

$$\text{cumulated\_emergency\_break\_duration} = (3500 - \text{cost}) / 100 \quad (3.1)$$

The constant 3500 is chosen to account for the 100hz sampling rate and the duration of 35 seconds. The division by 100 transforms the value into seconds. This value will be used from now on instead. A higher cumulated emergency brake duration is preferable. It is important to consider, that this value might not exactly correspond to the actual cumulated emergency brake duration time from the simulation, due to the penalty which the cost function applies for emergency breaks that last longer than 3 seconds.

Optimizing for different cost functions, such as improving time to collision (TTC) would have been interesting as well. However at the time, no working TTC functionality was implemented. Consequently, only the emergency break cost function will be optimized by the genetic algorithm.

explanation

### 3.3 Behaviour Tree

The EGO vehicle will not be controlled by the genetic algorithm, rather by a behaviour tree. The general idea is to have an EGO vehicle moving in a relatable manner through the world. It will try to dodge standing or slow moving obstacles. Every 3 seconds the behaviour tree switches between left, straight and right for the junction selection, to produce more interesting

### 3 Implementation

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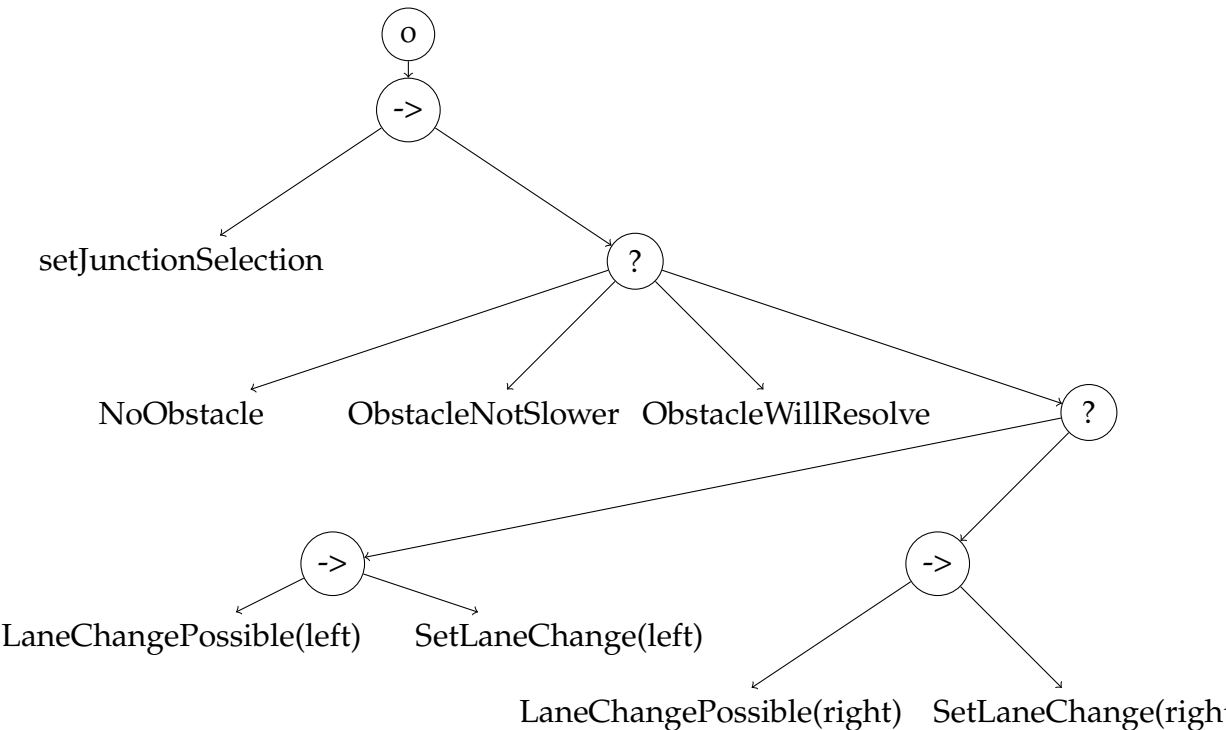


Figure 3.6: Used Behaviour Tree

scenarios, where the EGO vehicle is not always moving straight. Figure 3.3 shows the behaviour tree implemented.

better vis

While the behaviour tree has access to the same Action Interface (described in section 3.1.1) as the genetic algorithm, it needs a higher level of integration with the Traffic Manger. The genetic algorithm is only interested in the resulting emergency break values, the behaviour tree however needs access to internal functions during the simulation. Conditions like NoObstacle() or LaneChangePossible() execute internal Traffic Manager functions to get needed information on the world around the EGO vehicle.

detailed explanation on bt functions

# 4 Hyperparameter Tuning

The performance of a genetic algorithm is significantly influenced by its control parameters. Performant settings on a particular fitness landscape might not be appropriate on a different one (K. De Jong, 2007).

According to the "No Free Lunch Theorem", no single algorithm will outperform all other algorithms on a single class of problem (K. De Jong, 2007). K. De Jong, 2007 further states, that "the No Free Lunch results place an obligation on the EA practitioner to understand something about the particular properties of the problems that (s)he is trying to solve that relate to particular choices of EA parameter settings." A human-in-the-loop approach is needed to fine-tune parameter settings to a particular problem.

explain No Free Lunch Theorem better

This chapter will focus on tuning the genetic algorithm to perform well on the given cost function. First, a choice on the optimal population size is made. Afterwards a Taguchi method is used for tuning the remaining hyperparameter.

## 4.1 Simulation Setup

Two workstations were available for running all of the following simulations. The first workstation had an Intel Core i7-9700K and a GeForce RTX 2070 SUPER with 32 GB of RAM. The second workstation had an Intel Core i7-6850K as well as two Nvidia GeForce GTX 1080 also with 32 GB of RAM. On both systems, Kubuntu 20.04 LTS was the operating system.

As has been defined in chapter 3, each genetic algorithm will run for 30 generations. Additionally 1 simulation will have a duration of 35 seconds. On a single workstation, it takes approximately 3 hours and 50 minutes

## 4 Hyperparameter Tuning

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not sure where to show the start scenarios, they are currently in the appendix

to complete one genetic algorithm with a population size of 96. This time indication is however influenced by the number of actors.

Town 10 from the driving simulator Carla<sup>1</sup> will be used as the map for all experiments. It has an adequate size, yet is not too big and allows for interesting manoeuvres.

In order to reduce the required number of tests, only one starting scenario was used for all of the tuning of the control parameters. Start scenario 1 can be seen in appendix 6.1.1. In chapter 5, the performance of the tuned genetic algorithm on different start scenarios will be investigated.

### 4.2 Population

Finding a suitable population size is of high importance to a genetic algorithm, especially considering the limited processing resources available. On one hand, a population that is too small might result in less diverse runs of the genetic algorithm, on the other hand, if the population size is too high, the simulations will become too costly (see section 2.2).

In order to test for the best population size, other hyperparameters first have to be fixated. While reviewing the literature, trends of general settings for genetic algorithms can be found. Grefenstette, 1986 suggests that a range of control parameter will already lead to acceptable performance, yet optimal performance needs tuning. K. De Jong, 2007 complements these findings, adding that the "sweet spot" for control parameters of genetic algorithms is reasonably large and easy to find. A default set of static parameter values is generally speaking sufficient.

Following this advice, the most suitable population parameters will now be evaluated by fixating the remaining hyperparameters to a small range of suggested values from the literature.

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<sup>1</sup>[https://carla.readthedocs.io/en/latest/map\\_town10/](https://carla.readthedocs.io/en/latest/map_town10/)

### 4.2.1 Suggested Hyperparameter from the Literature

After reviewing various literature regarding control parameter of genetic algorithms, no clear consensus emerged. Mills, Filliben, and Haines, 2015 come to a similar conclusion, mentioning the inconsistencies between findings during their literature review and highlighting the conflicting evidence regarding "key GA control parameter".

Table 4.1 aims to provide a short, though not exhaustive, overview on different control parameter settings used in the literature. This compilation does not claim to cover the entire scope of available research in this domain, rather it served as a focused effort to identify usable hyperparameters.

Table 4.1: Summary of Genetic Algorithm Hyperparameters

Parameter Set	Pop	Cross	Mut	Sel	Elite
K. A. De Jong, 1975	50	0.6	0.001	?	1
Mills, Filliben, and Haines, 2015	200	?	Adaptive	SUS	8%
Grefenstette, 1986	30	0.95	0.01	?	1
Grefenstette, 1986	80	0.45	0.01	?	1
Almanee et al., 2021	50	0.8	0.2	?	?
Srinivas and Patnaik, 1994	30-100	0.9	0.01	?	?
Fazal et al., 2005	50	0.5	?	Tourn	5
Dao, Abhary, and Marian, 2016	200	0.7	?	Roul	1
Naruka et al., 2019	200	0.4	?	Roul	10
Jinghui Zhong et al., 2005	50-250	0.1-0.9	0.05-0.25	?	?

Generally speaking, a low mutation rate is recommended. For example Grefenstette, 1986 suggest poor performance using a rate over 0.05. 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 found higher mutation rates for their application to be more suitable. Srinivas and Patnaik, 1994 differentiates between higher and lower population numbers, claiming that a smaller population needs higher mutation rates in order to maintain a sufficient diversity.

analyze other control parameters a bit more

### 4.2.2 Comparison of Population Size

Based on the described research, population sizes of 32, 48, 64 and 96 will be compared. Crossover rates are set to 0.8 and 0.6. For mutation, 0.01 and 0.2 will be discussed. Further, tournament selection of 2 and 4 is used. Individual mutation probability will stay at 0.1. Chromosome encoding is set to Time and gene encoding is set to Integer. Each run will be executed 5 times to reduce randomness and to make the results more robust. Each simulation will last for 30 generations. A list of all settings with the mean over 5 repetitions per population size can be seen in table 4.2.2.

Settings	Code	32	48	64	96
C: 0.6, M: 0.01, TS: 2	A	4.49	4.84	6.49	6.29
C: 0.6, M: 0.01, TS: 4	B	3.89	4.79	4.21	5.63
C: 0.6, M: 0.20, TS: 2	C	4.38	4.90	4.98	6.69
C: 0.6, M: 0.20, TS: 4	D	4.80	5.33	6.09	6.50
C: 0.8, M: 0.01, TS: 2	E	4.37	6.08	5.29	5.84
C: 0.8, M: 0.01, TS: 4	F	4.48	4.51	4.46	6.03
C: 0.8, M: 0.20, TS: 2	G	4.01	5.60	5.41	6.31
C: 0.8, M: 0.20, TS: 4	H	4.42	4.95	7.06	6.91

Table 4.2: results from testing population size - mean of 5 repetitions

In figure 4.1, the results per population are plotted. The line corresponds to the mean, while the bars show the spread (min to max) of all 5 repetitions.

A higher spread in the results 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 was deemed to be more important to keep the variation low.

explain decision  
a bit more

## 4.3 Design of Experiment

This section aims to find optimal settings for the remaining control parameters of the given problem. Following the conclusion from the previous

### 4.3 Design of Experiment

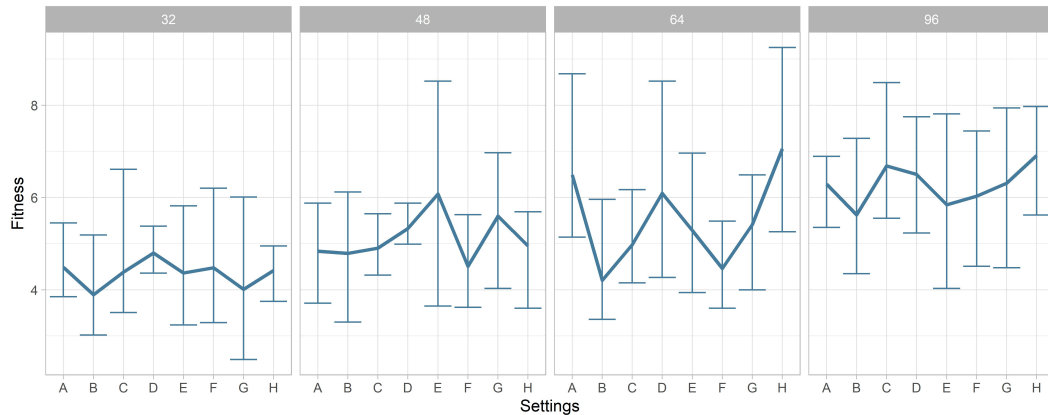


Figure 4.1: mean and error bars per population size

section 4.2, a population size of 96 will be fixated.

In order to tune the control parameter of the genetic algorithm, various different strategies can be used. Using automated approaches like "grid search", "bayesian optimization", "simulated annealing" or "hyperband" might lead to good results with minimal effort (tuning of hyperparameter for these search algorithms is still needed), however they each require a high number of runs (K. De Jong, 2007). [K. De Jong, 2007 even suggest using a second, higher level genetic algorithm for the optimization process.](#)

find more references

Due to performance considerations, many optimization methods did not fit the requirements. Executing one run for 30 generations currently takes around 3:50 hours. The high variance, between runs requires a certain number of repetitions for each setting. 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 budget. For this section, 8 repetitions were used per setting, which makes one evaluation last 30 hours.

A different approach is called design of experiment (DOE), also known as statistically designed experiments. DOE tries to find the cause-and-effect relationship between the factors and the output of experiments. It uses factorial design where the variables in an experiment are named 'factors'. Each factor consists of at least two settings, with the actual number of

## 4 Hyperparameter Tuning

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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.

*"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. Using ANOVA (Analysis of Variance) it is possible to identify the significance of each factor and interaction, which enables their ranking. More details on these analysis tools will be provided in section 4.3.3.

A full factorial design will test over all possible combinations of the manually selected factor levels. Looking at the proposed factors in table 4.3.1, 1024 runs<sup>2</sup> are required, which not feasible performance wise. A full factorial design has the drawback, that as the number of 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 used for estimating higher-order interactions, which are in most cases insignificant.

Add Short explanation

Fractional factorial experiments require less runs ...

*"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)*

### 4.3.1 Taguchi Method

Various improvements to design of experiment have been put forward by Dr. Genichi Taguchi, such as reducing the influence of uncontrollable (noise)

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<sup>2</sup>number of runs calculated using: <https://datatab.net/statistics-calculator/design-of-experiments>

### 4.3 Design of Experiment

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factors on processes and products and reducing variability (Roy, 1990). This master's thesis will not discuss all of Taguchi's proposed considerations, for more detail Roy, 1990 as well as Yang and El-Haik, 2009 are recommended.

Taguchi's proposed experimental design is a fractional factorial design, that requires significantly less runs. While different fractional factorial designs are available, Taguchi was chosen, because he provides a simple and easy to follow procedure which requires only a minimal number of runs.

*"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 non-existent. 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)*

Using Taguchi design for finding optimal hyperparameter of a genetic algorithm has been successfully performed by Dao, Abhary, and Marian, 2016 as well as Naruka et al., 2019.

Taguchi predefined a number of different orthogonal arrays where each row contains the specific levels (i.e the settings) of one experiment, while the columns correspond the factors (Hamzaçebi, 2021). Taguchi's orthogonal arrays "represent the smallest fractional factorials" (Roy, 1990). Only a fraction of combinations needs to be tested, drastically reducing computational needs.

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 a much smaller amount of simulation runs, while the full factorial experiments "might not provide appreciably more useful information" (Roy, 1990).

**Selection of an Orthogonal Array** When choosing a suitable Taguchi orthogonal array, various factors have to be taken into account. According to Yang and El-Haik, 2009, a three step procedure needs to be followed:

explain cons  
(e.g. higher order interactions are not possible)  
and orthogonality

## 4 Hyperparameter Tuning

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1. Calculate the total degree of freedom (DOF).
2. Base on the following two rules a 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 design of experiment, 7 factors (3 factors of level 4 and 4 factors of level 2) have been selected. Which factors and levels to choose was done based on experience gained on section 4.2. In table 4.3.1, every factor with the 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
CrossoverRate	B	0.2	0.5	0.8	0.9
MutationRate	C	0.01	0.1	0.3	0.5
ChromosomeType	D	Time	Time+NPC	-	-
GeneType	E	Int	Dict	-	-
TournamentSize	F	2	4	-	-
IndMutationRate	G	0.1	0.5	-	-

Table 4.3: List of control parameters (factors) with correspond settings (levels)

It is important to state, that Taguchi allows to only test for possible (pre determined) two-level interactions, evaluating higher level interactions is not possible (Yang and El-Haik, 2009). Analysing interactions comes at the cost of degrees of freedom. An interaction between ChromosomeType and GeneType might be of interest and will thus be chosen.

### 4.3.2 Selection of a Suitable Standard Orthogonal Array

The total degree of freedom can be 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 interaction ChromosomeType-GeneType:

$$DOF = 1 + 3 * (4 - 1) + 4 * (2 - 1) + 1 * (2 - 1) * (2 - 1) \quad (4.1) \\ = 15$$

A  $L_{16}$  array seems suitable to accommodate the required 15 DOF, which can be seen in table 4.3.2.

4 level factors need additional space which will be generated using column merging, while the interaction will need to be assigned as well. Either an interaction table or linear graphs of the  $L_{16}$  array can be used for both column merging and interaction assignment (NazanDanacioglu, 2005). Both illustrate the interaction relationships in the orthogonal array (Yang and El-Haik, 2009). The linear graph is more straight forward and will be the chosen approach. While there are multiple linear graphs for the  $L_{16}$  array, 4.3.2 describes the graph which best fits the requirements from table 4.3.1. If no suitable graph is found, they can be modified using rules described by NazanDanacioglu, 2005.

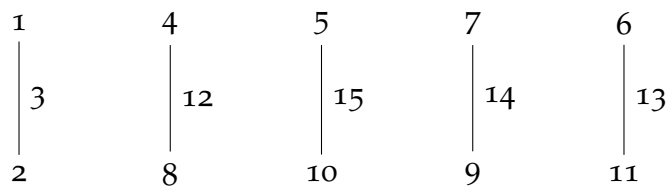


Figure 4.2: Linear Graph of  $L_{16}(2^{15})$  taken from Yang and El-Haik, 2009

## 4 Hyperparameter Tuning

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NO.	$L_{16}(2^{15})$														
	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

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

In a Taguchi linear graph, the nodes as well as the connections both represent columns in the orthogonal array. The interaction between two nodes is described by the connecting line (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 array only fits 2 level factors. Column merging, it is possible to extend columns to accommodate 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. Column merging needs the to be merged columns to be part of an interaction group (Yang and El-Haik, 2009). The available interaction groups are visualized by the linear graph in figure 4.3.2.

### 4.3 Design of Experiment

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3 2-level interaction columns need to be selected first. One column is discarded, the remain two columns are merged using the rules in tabular 4.5.

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

Table 4.5: 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 have to be merged in total.

**Assigning Interactions** Interactions between two-level factors can be assigned using the linear graph as well. An interaction between ChromosomeType and GeneType seems possible, thus D and E will be assigned to connected nodes in the linear graph. The resulting graph can be seen in 4.3.2. An interaction between F and G can not be investigated, as the chosen orthogonal array is not able to fit the additional 1 degree of freedom (see equation 4.1).

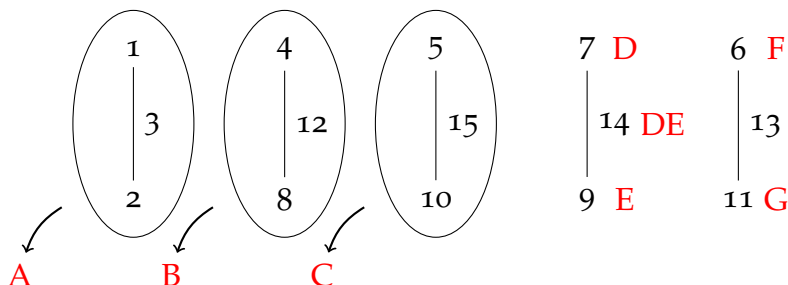


Figure 4.3: Modified Linear Graph to fit our needs

Combining columns 1 2 3 to A, 4 8 12 to B and 5 10 15 to C using rules defined by table 4.5 is done in 4.6.

## 4 Hyperparameter Tuning

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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		

Table 4.6: Building 4 Level columns from 2 Level columns

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 4.7. This combination table will subsequently be used as settings for the simulation runs.

### 4.3.3 Result Analysis

The final table 4.7 will 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 the table from 4.3.1. Every setting will be repeated 8 times to reduce randomness and gain information about variance. Running the genetic algorithm with these 16 different settings each repeated 8 times took 10 days on the two in section 4.1 described workstations. The results are found in the appendix at 6.1.1.

### 4.3 Design of Experiment

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NO.	A	B	C	D	E	F	G	DE
1	1	1	1	1	1	1	1	1
2	1	2	2	1	2	1	2	2
3	1	3	3	2	1	2	1	2
4	1	4	4	2	2	2	2	1
5	2	1	2	2	1	2	2	2
6	2	2	1	2	2	2	1	1
7	2	3	4	1	1	1	2	1
8	2	4	3	1	2	1	1	2
9	3	1	3	2	2	1	2	1
10	3	2	4	2	1	1	1	2
11	3	3	1	1	2	2	2	2
12	3	4	2	1	1	2	1	1
13	4	1	4	1	2	2	1	2
14	4	2	3	1	1	2	2	1
15	4	3	2	2	2	1	1	1
16	4	4	1	2	1	1	2	2

Table 4.7: Final version of used Taguchi orthogonal array

## ANOVA

ANOVA analysis (analysis of variance) will provide information on the magnitude of contribution of the main effects and interactions. The calculation of ANOVA on a Taguchi experiment is the same as for a classical design of experiment (Yang and El-Haik, 2009). Calculating ANOVA is done using the programming language R<sup>3</sup> and the result can be seen in table 4.8.

The sum of squares of a model tell how much of the total variation is explained by the model. The variance is broken down into the factors A-G along with the interaction D:E. The residual sum of squares shows the difference between the models prediction versus what was actually observed (i.e the variance that can not be explained by the model) (A. P. Field, Miles, and Z. Field, 2012).

The F ratio (or value) measures the ratio of variance explained by the factor

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<sup>3</sup><https://www.r-project.org/>

## 4 Hyperparameter Tuning

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	Df	Sum Sq	Mean Sq	F value	Pr(>F)
A	3	23.89	7.96	6.59	0.0004
B	3	5.00	1.67	1.38	0.2532
C	3	34.32	11.44	9.46	0.0000
D	1	3.88	3.88	3.21	0.0759
E	1	0.35	0.35	0.29	0.5912
F	1	18.91	18.91	15.64	0.0001
G	1	6.98	6.98	5.77	0.0179
D:E	1	4.10	4.10	3.40	0.0680
Residuals	113	136.60	1.21		

Table 4.8: ANOVA results

and the variation explained by the error term (A. P. Field, Miles, and Z. Field, 2012). Simply speaking, how good is the model versus how bad is the model.

Finally, the p value (in column labelled Pr(>F)) shows how likely the size of the given F ratio is obtained in case there is no effect on the results. Commonly, if p is smaller than 0.05, the effect can be viewed as statistically significant (A. P. Field, Miles, and Z. Field, 2012).

The higher number of DOF can be explained with the number of repetitions and can be calculated with the equation 4.2 (taken from Roy, 1990).

$$\begin{aligned}
 DOF &= totalNumberOfResults - 1 \\
 &= numberTrials * numberRepetitions - 1 \\
 &= 16 * 8 - 1 = 127
 \end{aligned} \tag{4.2}$$

The multiple R-squared value of the model is 0.416 while the adjusted R-squared value is 0.344. Multiple R-squared gives a measure on how much variability in the outcome is explained by the predictors (A. P. Field, Miles, and Z. Field, 2012). Having only 41.6% does not seem optimal. A. P. Field, Miles, and Z. Field, 2012 state further that a model which generalizes well has an adjusted R-squared value that is similar to multiple R-squared, which is also not the case in this scenario. The high error might possibly be

### 4.3 Design of Experiment

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explained by the huge search space in the scenario. Increasing the population and number of generations might lead to improvements, however at the cost of computational time. If the model, having this much of an error, will perform well compared to either a genetic algorithm build from values from the literature or compared to random search will be evaluated in section 5.

Looking at the factors as well as the interaction, significant main effects can be seen. The highest F value has the main effect F (TournamentSize). The effect is significant with  $F(1, 112) = 15.64$ ,  $p < 0.001$ . Next, the effect C (MutationRate) is significant with  $F(3, 112) = 9.46$ ,  $p < 0.001$ . A (CrossoverType) is also significant  $F(3, 112) = 6.59$ ,  $p < 0.001$ . Finally G (IndependentMutationRate) is significant with  $F(1, 112) = 5.77$ ,  $p < 0.05$ . D (ChromosomeType) and the interaction D:E will be mentioned as well with  $F(1, 112) = 3.21$ ,  $p < 0.1$  and  $F(1, 112) = 3.4$ ,  $p < 0.1$  respectively. There is not enough evidence that suggests significant main effects for the factors B (CrossoverRate) and E (GeneType).

Especially the fact, that the CrossoverRate does not show significant effects is surprising, which is not supported by the literature (todo: find references). The low influence of GeneType however might be explained by the fact, that it does not have a huge impact on the action selected apart from more granularity of the parameters when Dictionary encoding is used.

To look at the percentage contribution of each factor the formulas 4.3 and 4.4 (gathered by Yang and El-Haik, 2009) are used.

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

Example calculation for factor A.

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

The percentage contribution of all factors is plotted in 4.3.3 and shows the high error of the model.

## 4 Hyperparameter Tuning

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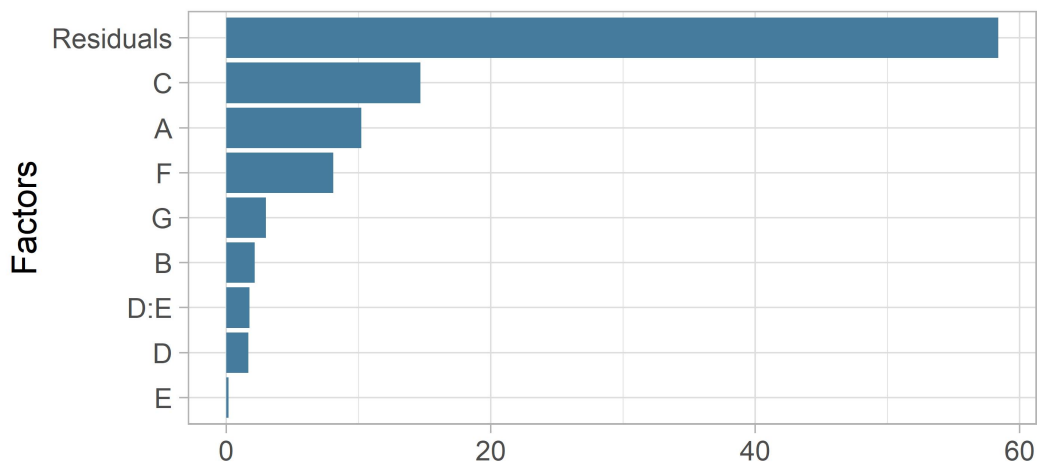


Figure 4.4: Percentage Contribution

### Main-effects and interaction chart

Identifying the optimal conditions needs analysis of the main effects per factor. They allow to predict the levels, that lead to the best result (Roy, 1990).

*"The main-effects chart is a plot of average responses at different levels of a factor versus the factor levels." (Yang and El-Haik, 2009)*

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

The resulting main-effect charts can be seen in figure 4.3.3.

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 highest value).

If interactions exist, they might have an influence on the best settings and need further investigation (Yang and El-Haik, 2009). To investigate the previously defined interaction an interaction graph can be used. The calculation is similar to calculating main effects and shown in figure 4.3.3.

Show calculation example for D

Show calculation example for DE

### 4.3 Design of Experiment

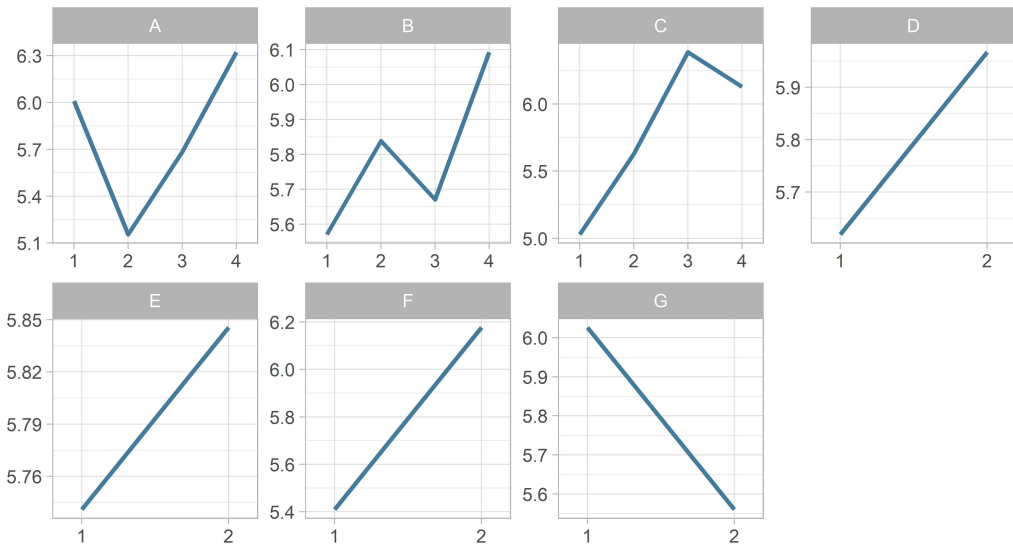


Figure 4.5: Main Effects

The crossing of lines indicates, that an interaction between the two factors might exist (A. P. Field, Miles, and Z. Field, 2012). 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).

#### 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 according the main effects is the following: A<sub>4</sub>, B<sub>4</sub>, C<sub>3</sub>, D<sub>2</sub>, E<sub>2</sub>, F<sub>2</sub>, G<sub>1</sub>.

Analysing the ANOVA table, the interaction D:E however seems to have significance, especially compared to E. According to (A. P. Field, Miles, and Z. Field, 2012), it makes little sense to further interpret main effects if the interaction effect is significant (which it almost is with  $p < 0.1$ ), thus this interaction will be integrated.

## 4 Hyperparameter Tuning

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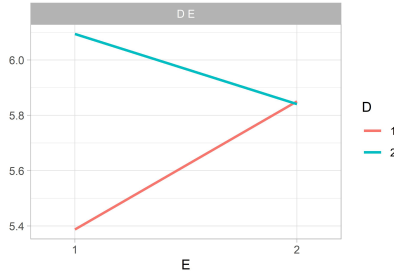


Figure 4.6: Test of interactions

The test of interaction in figure 4.3.3 suggest D<sub>2</sub> and E<sub>1</sub> as the best combination. This is optimal, as D<sub>2</sub> is also recommended by the main effects. While E<sub>1</sub> does not correspond to its main effect, the low F value suggests low significance for E anyway. Concluding this line of thought, the combination A<sub>4</sub>, B<sub>4</sub>, C<sub>3</sub>, D<sub>2</sub>, E<sub>1</sub>, F<sub>2</sub>, G<sub>1</sub> seems to be optimal.

**Optimum performance calculation** To calculate the predicted results of an algorithm with these settings, optimal performance calculation can be used. Equation 4.5 only uses the optimal main effects while equation 4.6 has the interaction D:E applied. The calculation formulas are provided by Roy, 1990.

$$\begin{aligned}
 Y_{opt} &= \bar{T} + (\bar{A}_4 - \bar{T}) + (\bar{B}_4 - \bar{T}) + (\bar{C}_3 - \bar{T}) + (\bar{D}_2 - \bar{T}) + \\
 &\quad (\bar{E}_2 - \bar{T}) + (\bar{F}_2 - \bar{T}) + (\bar{G}_1 - \bar{T}) \\
 &= 8.06
 \end{aligned} \tag{4.5}$$

$$\begin{aligned}
 Y_{opt} &= \bar{T} + (\bar{A}_4 - \bar{T}) + (\bar{B}_4 - \bar{T}) + (\bar{C}_3 - \bar{T}) + (\bar{D}_2 - \bar{T}) + \\
 &\quad (\bar{E}_1 - \bar{T}) + ([\bar{D}x\bar{E}]_2 - \bar{T}) + (\bar{F}_2 - \bar{T}) + (\bar{G}_1 - \bar{T}) \\
 &= 8.13
 \end{aligned} \tag{4.6}$$

Utilizing the interaction D:E, the performance estimation improves from 8.06 to 8.13. Finally the optimized settings are as follows: CrossoverType:

Uniform 0.5, CrossoverProbability: 0.9, MutationProbability: 0.3, ChromosomeType: Time+NPC, GeneType: Integer, TournamentSize: 4 and IndividualMutationProbability: 0.1.

#### signal-to-noise (S/N)

Talk about high randomness in the recommended settings

As previously discussed, the ANOVA model has a high error, 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.

*"use of the S/N ratio offers an objective way to look at the two characteristics (consistency and average value) together." (Roy, 1990)*

S/N is calculated in two steps, according to Roy, 1990. First the mean square deviation (MSD) is needed. Depending on the quality characteristic, a different equation has to be chosen. In this case, a higher result is better, thus equation 4.7 is used for each repetition, with  $y$  being a result and  $n$  the number of repetitions.

$$MSD = (1/y_1^2 + 1/y_2^2 + 1/y_3^2 + \dots)/n \quad (4.7)$$

S/N is further calculated using equation 4.8.

$$S/N = -10\log_{10}(MSD) \quad (4.8)$$

Generating main effects and ANOVA is now done using S/N instead. As a side note, only 15 DOF for the ANOVA analysis are available, as repetitions per run get merged into 1 result (in contrast to 4.2), which can be seen in equation 4.9.

$$\begin{aligned}
 DOF &= totalNumberOfResults - 1 \\
 &= numberOfTrials * 1 - 1 \\
 &= 16 - 1 = 15
 \end{aligned} \tag{4.9}$$

The resulting ANOVA table can be seen in 4.9.

	Df	Sum Sq	Mean Sq	F value	Pr(>F)
A	3	5.14	1.71	3.54	0.3682
B	3	2.04	0.68	1.40	0.5397
C	3	10.59	3.53	7.28	0.2644
D	1	1.29	1.29	2.67	0.3498
E	1	0.39	0.39	0.81	0.5329
F	1	4.43	4.43	9.15	0.2033
G	1	2.28	2.28	4.70	0.2751
D:E	1	0.92	0.92	1.89	0.4005
Residuals	1	0.48	0.48		

Table 4.9: S/N ANOVA results

When looking at the p values of this table, it is obvious that no factor can discard the null-hypothesis, which states that a factor has no significant effect. Considering this fact, it was deemed to not be necessary to perform further investigations. The often recommend Signal to Noise ratio does not seem to suitable for this experiment, thus the previously calculated settings in paragraph 4.3.3 will not be adjusted.

## Elite Selection

Although the optimal hyperparameter setting are discussed in chapter 5, a problem was obvious when analysing results of a GA using the settings from 4.3.3. Setbacks in the optimal cost between two generations happen frequently, likely due to the high crossover and mutation rates. In order to mitigate this problem, it was decided to implement elite selection with a size of 2. Per generation, the two best individuals are now copied into the

### 4.3 Design of Experiment

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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 4.3.3 compares both no elite selection with an elite selection of 2. For a few selected repetitions, the best individual cost per generation is plotted.

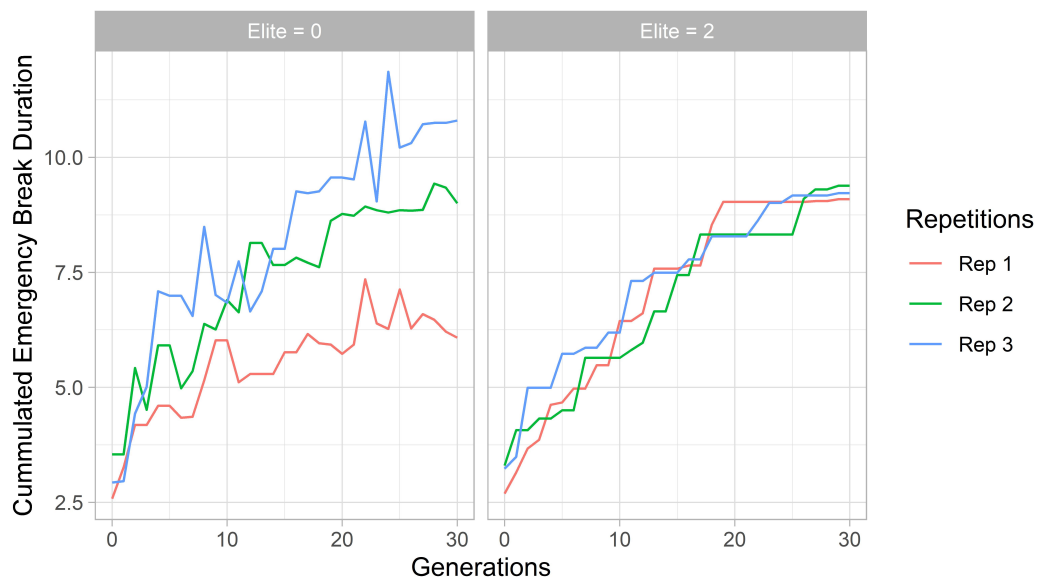


Figure 4.7: Elite Selection Comparison

For analysing the statistical significance in the differences in mean between the a genetic algorithm using elite selection of 2 vs no elite selection, a t-test can be used. In order to ignore possible violation of the assumption of homogeneity, a robust welch t-test is applied which adjusts the DOF accordingly (A. P. Field, Miles, and Z. Field, 2012).

On average, using elite improved the performance ( $M = 8.52$ ,  $SE = ^*$ ), compared to using no elite ( $M = 7.92$ ,  $SE = ^*$ ). This difference was not significant  $t(14.21) = 0.96$ ,  $p > 0.05$ ; however, it did represent a small-sized effect  $r = 0.25$ . A comparison of both settings each repeated 10 times can be seen in figure 4.3.3.

Due to the existence of the mentioned small effect, it is concluded that the

## 4 Hyperparameter Tuning

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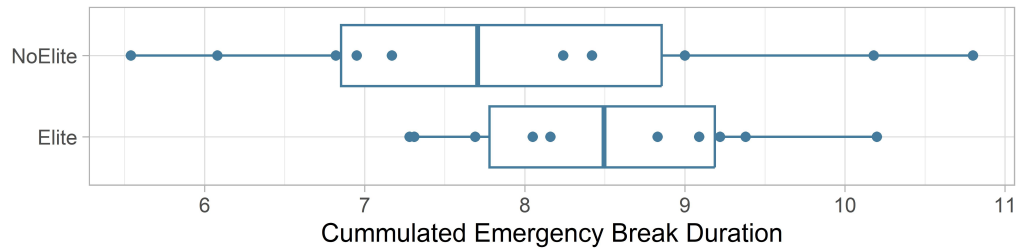


Figure 4.8: Comparison Elite Selection vs No Elite Selection

slightly modified version of the settings from 4.3.3 will be used in chapter 5, where elite selection is set to 2.

Find paper

**Effect Size** According to A. P. Field, Miles, and Z. Field, 2012 effect sizes provide an objective measure on the importance of an effect, where 0 means no effect and 1 means a perfect effect. They allow for a standardized measure and are not affected by sample size. He further recommends to use the widely used suggestions made by Cohen (1988, 1992) on defining between a large or small effect:

- $r = .10$  (small effect): The effect explains 1% of the variance.
- $r = .30$  (medium effect): The effect explains 9% of the variance.
- $r = .50$  (large effect): The effect explains 25% of the variance.

Effect sizes will be further used to compare different algorithms in chapter 5 as well and are calculated for a t test using equation 4.10.

$$r = \sqrt{\frac{t^2}{t^2 + DOF}} \quad (4.10)$$

# 5 Evaluation

This chapter will compare the GA settings proposed in 4.3.3 with the best settings found in 4.2 as well as with random search. The three different algorithms that are going to be compared are as follows:

- **Default Genetic Algorithm** - using the best settings found in 4.2
  - CrossoverType: Two-Point, CrossoverRate: 0.8, MutationRate: 0.20, TournamentSize: 4, ChromosomeType: Time, GeneType: Integer, IndividualMutationRate: 0.1 and EliteSelection: 0.
  - The algorithm will be run 10 times. The population of 96 needs to be simulated 31 times per run (30 generations + initialization), which leads to  $96 * 31 * 10 = 29,760$  simulations.
- **Optimized Genetic Algorithm** - using the recommended settings found in 4.3.3
  - CrossoverType: Uniform 0.5, CrossoverRate: 0.9, MutationRate: 0.3, ChromosomeType: Time+NPC, GeneType: Integer, TournamentSize: 4 and IndividualMutationRate: 0.1 and EliteSelection: 2.
  - The algorithm will be run 10 times. The population of 96 needs to be simulated 31 times per run (30 generations + initialization), which leads to  $96 * 31 * 10 = 29,760$  simulations.
- **Random Search**
  - For randomly choosing actions, the same dict probabilities from [appendix](#) are used.
  - The algorithm will be run 10 times each with 2,976 simulations, always taking the maximum value as the result. 29,760 simulations were executed in total.

ref appendix

## 5 Evaluation

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The evaluation is done over 4 different start scenarios which can be found in the appendix at 6.1.1

### 5.1 Start Scenario 1

First, a comparison of the three algorithms on start scenario 1 was made. This is the same start scenario, that was used in chapter 4. The results are shown in figure 5.1.

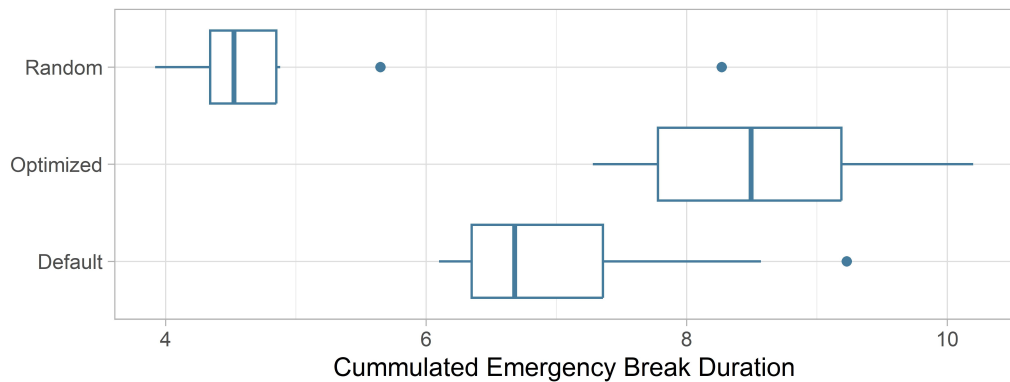


Figure 5.1: Start Scenario 1: Default GA vs Optimized GA vs Random Search

Analysing the graph, the Optimized GA clearly outperformed the Default GA as well as Random Search.

Welches t-test shows that on average, greater fitness is achieved by using Optimized GA ( $M = 8.52$ ,  $SE = ?$ ) than from using Default GA ( $M = 7.09$ ,  $SE = ?$ ). This difference was significant  $t(17.87) = 3.15$ ,  $p < .01$ . It did represent a large effect  $r = 0.60$ . Compared to Random ( $M = 4.943$ ,  $SE = ?$ ), the Optimized GA has even higher dominance, with a significant difference  $t(16.9) = 7.12$ ,  $p < .001$  and a large effect  $r = 0.87$ .

The better performance of the Optimized GA was however unsurprising, as it was specifically trained on the used start scenario.

## 5.2 Start Scenario 2

To further analyse both genetic algorithms, a look at their performance over the generations next to their diversity chart is of interest. Figure 5.1 plots the mean over 10 repetitions, the outline show the first and third quantiles.

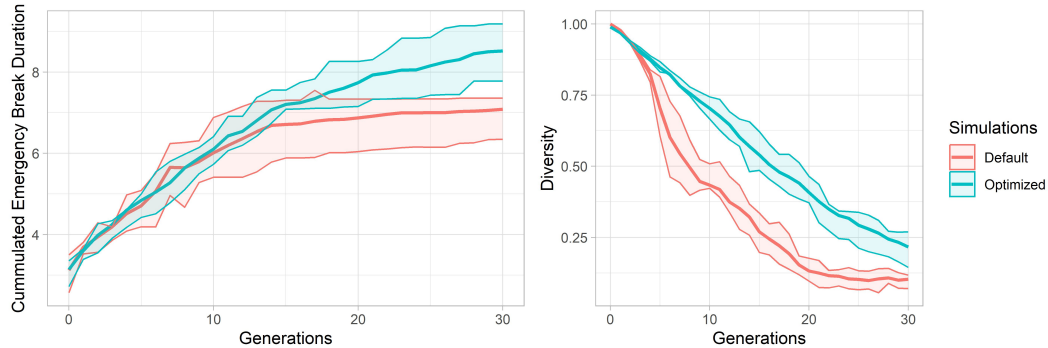


Figure 5.2: Start Scenario 1: Comparison of GAs

After generation 10, the rate of improved fitness of the Default GA drops compared to the Optimized GA. A combined early sharp decline in the diversity, suggests a connection. The Optimized GA shows to hold the diversity in the population longer, its rate of convergence is linear. Its high crossover and mutation rates seem to help with the improved diversity. The graph also shows, that a higher number of generations might not be useful for improved performance, as after 30 generations, the Optimized GA does not seem to hold enough diversity to continue with adequate performance gains.

## 5.2 Start Scenario 2

9 Vehicles and 5 pedestrians. same number as start scenario 1.

simulations have not yet finished

## 5.3 Start Scenario 3

Start scenario 3 is described in more detail in Appendix 6.1.1. 5 vehicles with 3 pedestrians are initialized, resulting in a simulation with only a small

## 5 Evaluation

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number of NPCs.

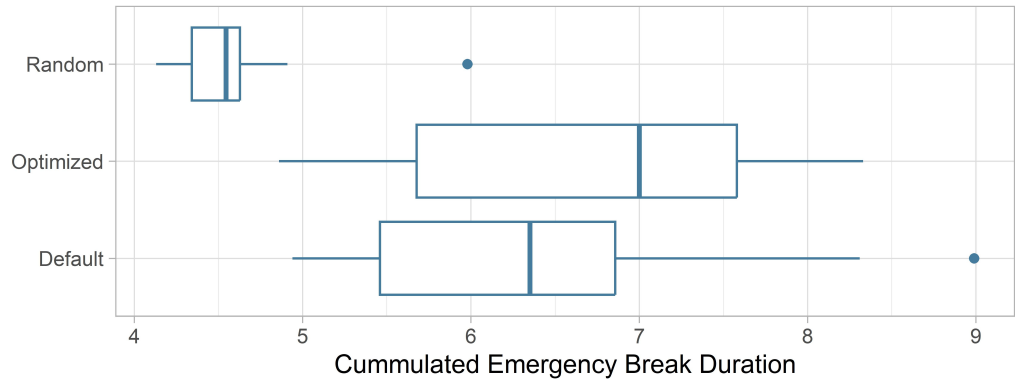


Figure 5.3: Start Scenario 3: Default GA vs Optimized GA vs Random Search

Looking at the graph, the Optimized GA again outperforms the Random Search, however there seems to be only marginal improvements compared to Default GA.

A t-test confirms these findings. While on average, greater fitness is achieved by using Optimized GA ( $M = 6.66$ ,  $SE = ?$ ) than from using Default GA ( $M = 6.49$ ,  $SE = ?$ ). This difference was however not significant  $t(17.83) = 0.29$ ,  $p > .05$  and an effect size of  $r = 0.07$ . Verifying the better performance of the Optimized GA compared to Random Search ( $M = 4.619$ ,  $SE = ?$ ) shows a significant difference  $t(12.4) = 4.9$ ,  $p < .001$  and a large effect size of  $r = 0.81$ .

To further analyse both genetic algorithms, their performance over the generations next to their diversity chart is again shown in figure 5.3. The mean over 10 repetitions is plotted, the outline show the first and third quantiles.

The rate of improved fitness of the Default GA now drops very similar to the Optimized GA. The early decline in the diversity is also not as pronounced as in the previous comparisons. While the optimized GA shows to still hold the diversity in the population longer, this only has a minimal impact on its average fitness. The similarity in performance might be explained by the smaller search space due to the smaller number of NPCs. Here,

## 5.4 Start Scenario 4

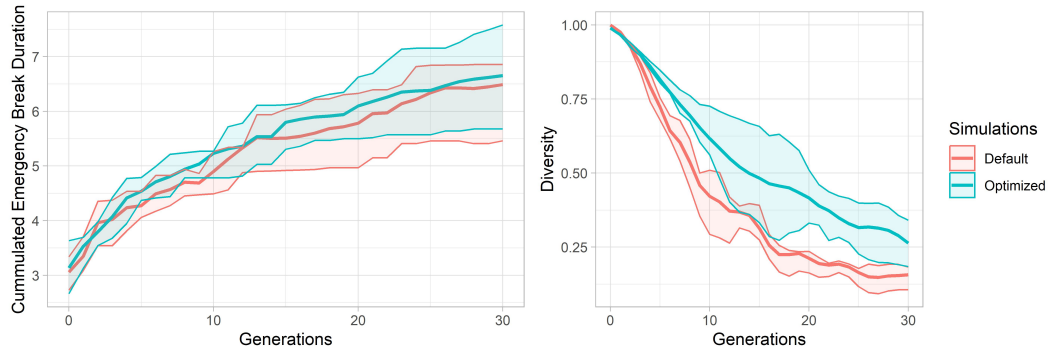


Figure 5.4: Start Scenario 3: Comparison of GAs

high mutation and crossover rates might not have the previous pronounced advantage.

## 5.4 Start Scenario 4

Start scenario 4 can be seen in Appendix at 6.1.1. 18 vehicles with 10 pedestrians are initialized, resulting in the simulation with the most NPCs.

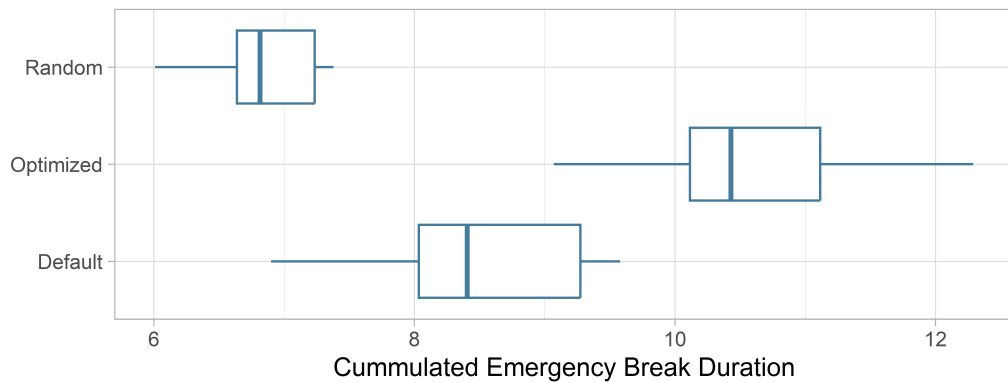


Figure 5.5: Start Scenario 4: Default GA vs Optimized GA vs Random Search

Figure 5.4 shows the Optimized GA clearly outperforming the Default GA as well as Random Search. The Optimized GA ( $M = 10.60$ ,  $SE = ?$ ) has

## 5 Evaluation

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significantly greater fitness than the Default GA ( $M = 8.46$ ,  $SE = ?$ ) with  $t(17.98) = 5.30$ ,  $p < .001$  and a large effect  $r = 0.78$ . Compared to Random Search ( $M = 6.86$ ,  $SE = ?$ ), the greater fitness of the Optimized GAs is significant with  $t(11.71) = 12.7$ ,  $p < .001$  and a large effect  $r = 0.96$ .

Both genetic algorithms performance over the generations next to their diversity chart is compared in figure 5.4.

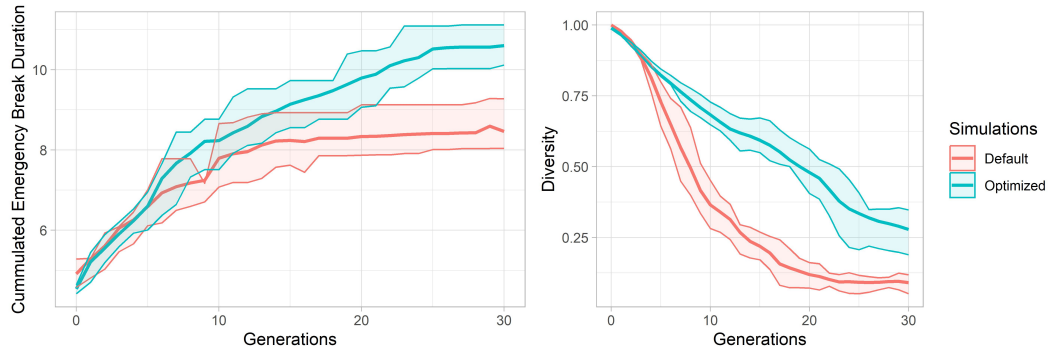


Figure 5.6: Start Scenario 4: Comparison of GAs

Already at generation 5, the average rate of improved fitness of the Default GA drops compared to the Optimized GA. This is accompanied by an early sharp decline in the diversity. The optimized GA again shows be able to hold the diversity much longer at a high level, its rate of diversity drop is linear. The results underline the statement made in section 5.3, which claims, that the Optimized GA excels at scenarios with lots of NPCs, where a vast search space is given.

# 6 Conclusion

The in this master's thesis proposed settings for a genetic algorithm show good performance on the given search problem.

write better and longer conclusion

The results of chapter 5 prove, that the optimized settings GA shows significant improvements in most start scenarios compared to general settings from the literature as well as compared to random search. Only in the case of a small number of NPCs, the performance seems to be on par with a genetic algorithm utilizing settings from the literature.

Compared to random testing, research question 1 definitely holds true. The optimized genetic algorithm will drastically improve performance when using the given cost function.

The second research question can be answered as well with yes. Taguchi orthogonal testing only needs a minimal amount of different experiments while leading to impressive performance improvements compared to using control parameters suggested by the literature.

## 6.1 Future Work

### 6.1.1 Oracles

Using oracle functions to select interesting scenarios during the runtime of a genetic algorithm was already done by Almanee et al., 2021. Different oracle functions test for different thresholds in the simulation. Examples are 'overSpeedLimit', 'criticalLaneChange', 'criticalComfortLevel'. These functions will test each individual in the population. In case one or multiple functions return true, the individual scenario will be saved and automatically

add Responsibility-Sensitive Safety (RSS)

Better introduction

## 6 Conclusion

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categorized according to the types of thresholds. The genetic algorithm remains unaffected.

Oracles have the potential to extract a multiple different scenarios from only one genetic algorithm run. The automatic categorization drastically improves the usefulness of this testing approach. Otherwise each result from the genetic algorithm needs to be evaluated manually for finding errors in the ADAS/AD function.

# **Appendix**



# Appendix A.

Better images



Figure 1: Start scenario 1

## Appendix A.

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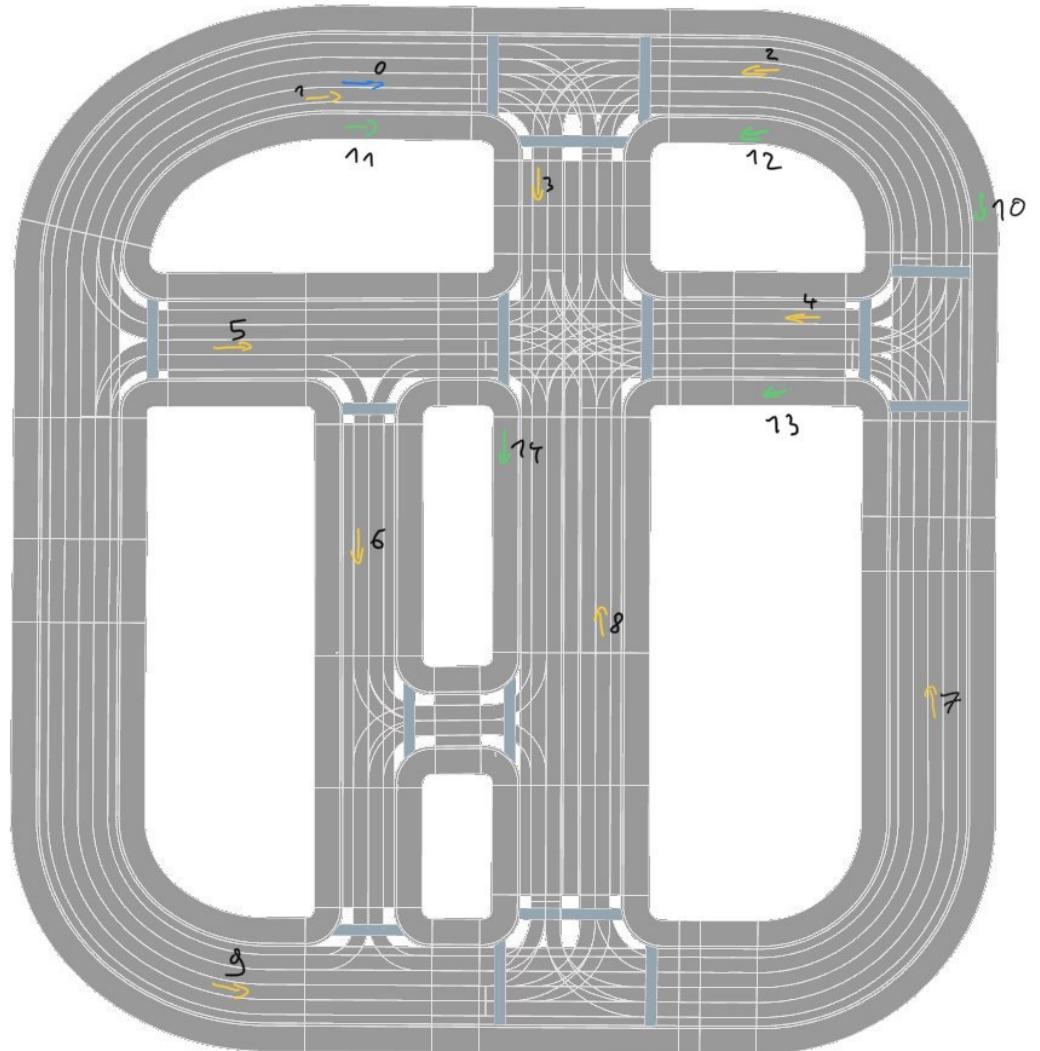


Figure 2: Start scenario 2

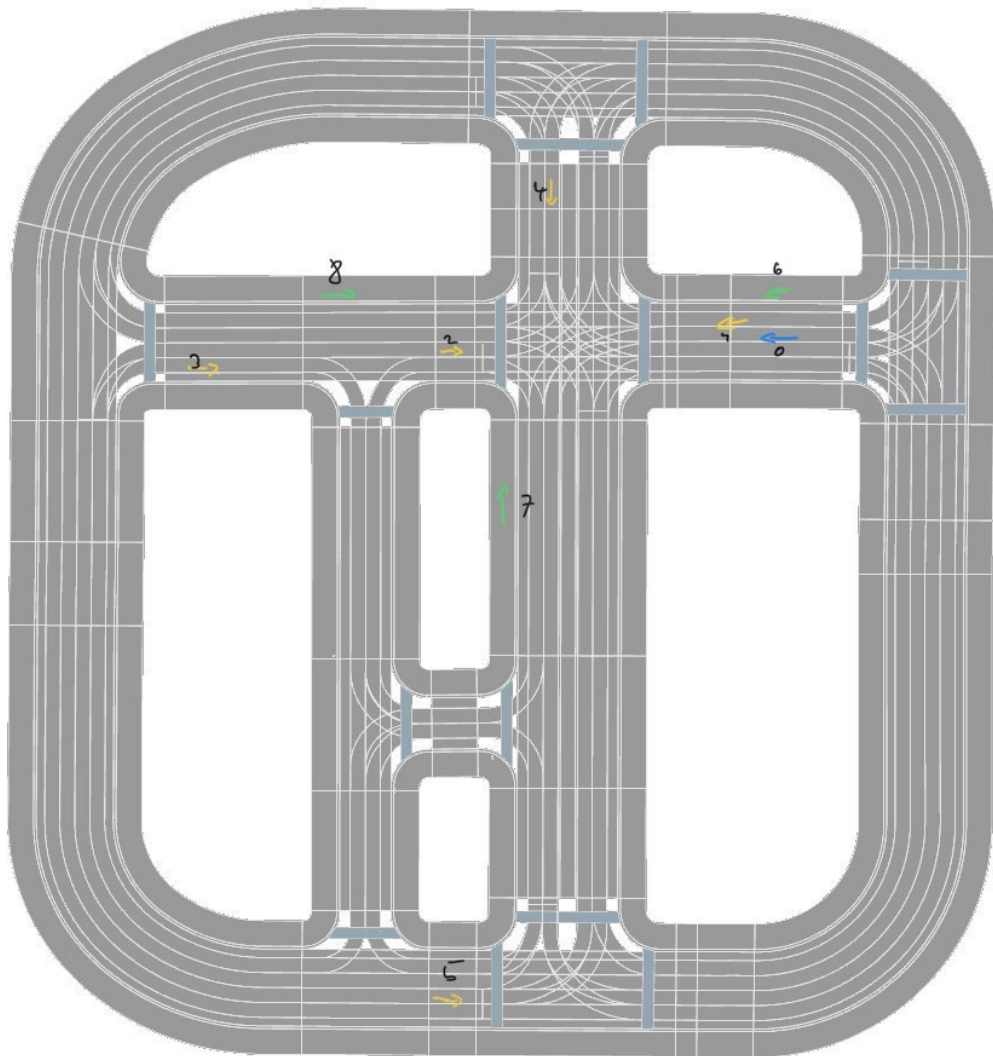


Figure 3: Start scenario 3

## Appendix A.

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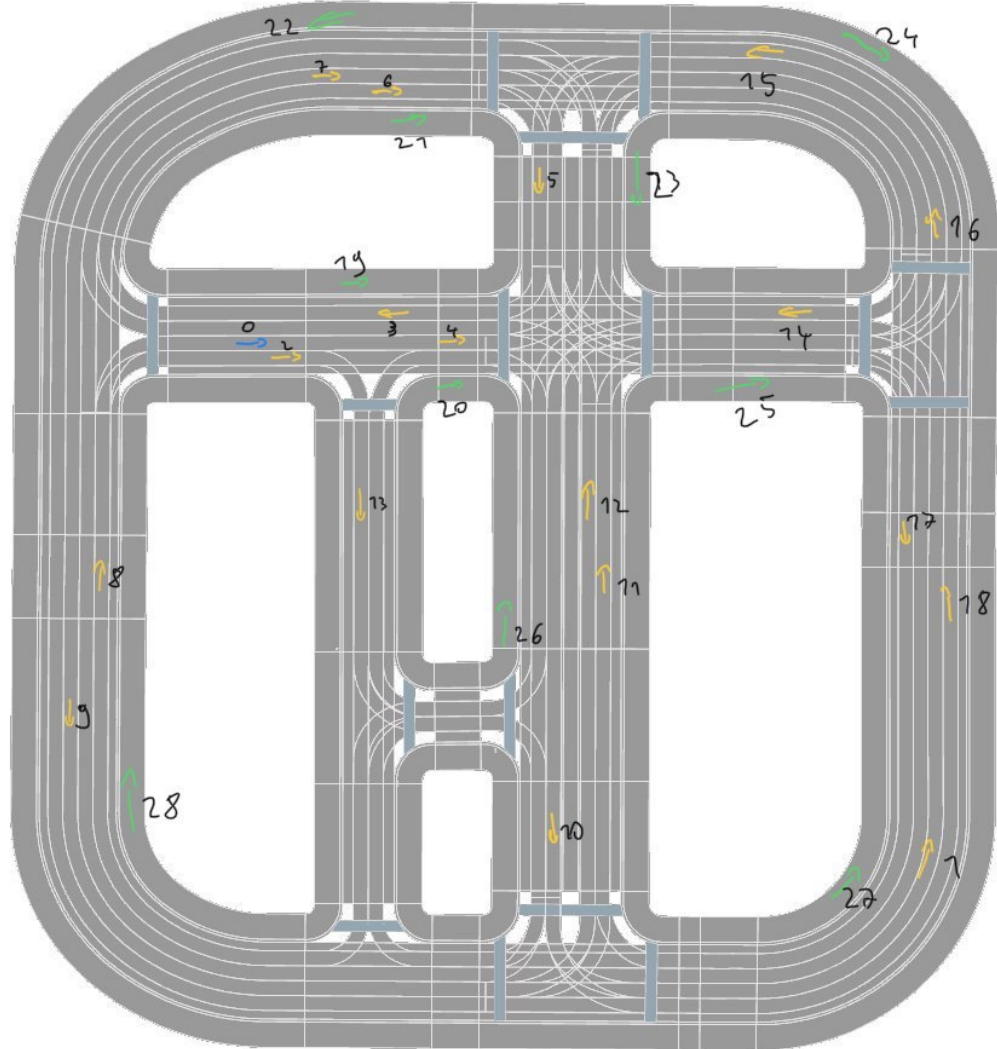


Figure 4: Start scenario 4

## Appendix B.

NO.	rep1	rep2	rep3	rep4	rep5	rep6	rep7	rep8
1	3.76	3.90	4.75	4.23	4.32	5.75	3.94	3.95
2	8.06	5.20	4.75	5.04	4.63	5.79	4.32	6.00
3	8.62	7.89	8.76	6.44	8.77	6.68	5.96	7.22
4	7.65	6.95	6.49	5.35	6.24	7.97	6.52	6.42
5	4.26	5.45	4.55	4.20	3.80	5.29	6.05	7.02
6	5.26	5.21	5.71	5.59	5.48	5.64	3.61	5.47
7	3.92	4.01	4.51	4.80	4.08	4.43	4.10	6.05
8	6.60	5.69	5.79	5.79	5.43	5.03	6.11	6.05
9	4.93	5.05	5.17	4.91	6.53	5.04	7.66	5.73
10	5.84	6.72	4.87	7.13	6.82	5.74	4.66	6.78
11	4.93	6.21	4.10	4.67	5.94	5.19	3.91	3.96
12	5.54	4.84	7.10	5.83	5.96	5.17	6.02	8.94
13	11.22	7.88	5.94	7.00	5.88	7.05	6.40	6.66
14	6.05	7.40	7.50	6.51	9.58	5.03	5.35	5.09
15	6.58	4.35	4.50	7.21	6.38	5.77	5.45	6.08
16	4.35	6.66	8.57	4.44	4.49	4.89	6.72	5.37

Figure 1: Taguchi experiment results

insert all results  
from evaluation



# Appendix C.

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Insert information of Gene action probabilities



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