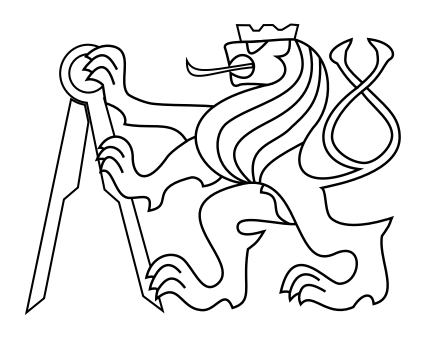
# CZECH TECHNICAL UNIVERSITY IN PRAGUE

Faculty of Electrical Engineering

# BACHELOR'S THESIS



Zdeněk Rozsypálek

Active 3D mapping using laser range finder with steerable measuring rays

Department of Cybernetics

Thesis supervisor: Ing. Tomáš Petříček

# Acknowledgements I would like to thank my adviser for his advice.

Abstract
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The abstract in English.

Abstrakt

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### 1 Introduction

Lidar sensors offer an accurate volumetric mapping of surrounding space. There is much utilization of volumetric space reconstructions in different fields. For example, lidar sensors are nowadays essential equipment for a large variety of autonomous vehicles. The sensor can help autonomous vehicles to orientate in an environment. One of the most significant issues which prevent broader implementation of these sensors is a relatively high price. Breakthrough in this field is solid-state lidars. These lidars do not have moving parts, and their price should be circa hundreds of dollars [1]. The major drawback of this sensor is a limited number of rays which can be sent over the certain time frame and only in chosen directions. Zimmermann et al. [2] proposed mapping agent which creates dense reconstructions from sparse measurements. They also proposed prioritized greedy planning for choosing directions of these rays. The objective of this thesis is to apply reinforcement learning (RL) methods to learn planning of the rays and contribute to methods of controlling these sensors. RL is a field of study based on concepts of behavioral psychology, especially the trial and error method, and has in recent years experienced a rapid development due to the growth of computational power and neural networks improvement. Richard Sutton has made a helpful summary of RL concepts in his book [3]. One of the biggest achievements was playing Atari games by a RL agent without any prior knowledge of the environment [4]. Soon after the RL agent, able to solve simple continuous problems such as balancing inverse pendulum on a cart, was introduced. Today state-of-the-art methods can solve complex environments with infinite action spaces. Although these methods reach great successes, they still suffer from lack of sample efficiency - they need for training many episodes before environment can be solved. This inefficiency makes creating agent controlling lidar very challenging, since training large neural networks is very time-consuming. The agent is divided into two parts - mapping and planning. The mapping part should create a best possible reconstruction from sparse measurements, while the planning part is focused on picking rays that will maximize reconstruction accuracy. Agents are trained using a publicly available dataset which contains drives of a car equipped with Velodyne lidar [5]. Theoretical background of RL is discussed in first part of this thesis. In the second part are methods from first part used to solve the Lidar-gym environment [6].

# 2 RL basics

Firstly, an environment where an agent can operate must be defined. The environment can be described as Markov decision process, where  $S_t \in \mathcal{S}$  is a state from a set of possible states  $\mathcal{S}$  in which environment is located in time t. An agent can usually observe the state of the environment and take action accordingly. An action is a probabilistic transition between states. Every action  $A_t \in \mathcal{A}$  moves the environment from  $S_t$  to  $S_{t+1}$ . The environment evaluates every action and returns appropriate reward  $R_t$  (figure 1). In RL set  $\mathcal{A}$  is often called action space and set  $\mathcal{S}$  observation space. The main goal of the agent is to find policy  $\pi$  which maximizes expected return. Return  $G_t$  is a sum of discounted future rewards [3].

$$G_t = \sum_{k=0}^{\infty} \gamma^k R_{t+k} \tag{1}$$

where  $\gamma \in [0,1]$  is discount factor. RL methods define how experiences from interacting with the environment will change the policy. The major issue is that maximizing immediate reward is often not an effective approach to maximize an expected sum of discounted rewards. This greedy policy can take the agent into a very disadvantageous state. Thus, the agent must take into account future states and rewards. This is done by value function  $V_p i(S_t)$  which assesses how advantageous is being in state  $S_t$  with policy  $\pi$ .

$$V_{\pi}(S_t) \doteq \mathbf{E}_{\pi}[G_t|S_t]. \tag{2}$$

Optimal policy  $\pi^*$  is then defined as

$$\pi^*(S_t) \doteq \max_{\pi} V_{\pi}(S_t), \tag{3}$$

for all  $S_t \in \mathcal{S}$ . In the past agents used big tables to estimate the value function. That is possible in environments with small action and observation spaces but is very memory consuming for larger environments and even impossible for continuous action or observation space. Therefore, modern methods use neural networks as function estimators.

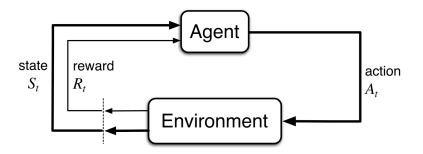


Figure 1: RL concept. Source - [3].

### 2.1 Temporal difference learning

Temporal difference (TD) learning combines the ideas of Monte Carlo methods and dynamic programming. It can learn directly from experience obtained by interactions with an environment without any prior knowledge of the said environment. TD learning is done by following assignment in each timestamp [3]

$$V(S_t) \leftarrow V(S_t) + \lambda [R_t + \gamma V(S_{t+1}) - V(S_t)] \tag{4}$$

where  $\lambda \in \mathbb{R}^+$  is step size.

# 2.2 Q-learning

Q-learning is a type of TD learning developed by Watkins [7]. The state value V from the previous subsection is replaced by Q value, which refers to a quality of action in a particular state instead of the quality of the state itself. When we rewrite TD learning (4) to Q-learning we get:

$$Q(S_t, A_t) \leftarrow Q(S_t, A_t) + \lambda [R_t + \gamma \max_{A_{t+1}} Q(S_{t+1}, A_{t+1}) - Q(S_t, A_t)].$$
 (5)

Our policy here is to take action with maximal Q value. That is called greedy policy. An obvious drawback of greedy policy is that it does not allow to explore the whole environment properly because an action with the highest Q value is always chosen. A solution to this problem is sometimes to take random action to explore the environment. This policy is often referred to as  $\epsilon$ -greedy policy.

### **Algorithm 1** $\epsilon$ -greedy policy in pseudocode

```
1: function ChooseAction
2: \epsilon \leftarrow \epsilon \cdot \epsilon_d
3: if \epsilon > \text{random} \in (0, 1) then
4: \text{action} \leftarrow \text{random} \in \mathcal{A}
5: else
6: \text{action} \leftarrow \max_{A_t} Q(S_t, A_t)
7: return action
8: end function
```

It is common to set  $\epsilon = 1$  at the beginning of the training and decay rate  $\epsilon_d$  close to one. The general idea behind this policy assumes that it is needed to explore an environment first and then exploit agents experience.

### 2.3 Prioritized experience replay

Experience replay is biologically inspired mechanism introduced by Schaul et al. [8] which stores all experiences (specifically:  $S_t$ ,  $A_t$ ,  $R_t$ ,  $S_{t+1}$ ) into a buffer and assigns priority to every experience. The main idea is that experiences with high TD should have higher priority. It is thus necessary to calculate priority p from TD error:

$$p = (|\delta_t| + \eta)^{\rho} \tag{6}$$

where  $\rho$  indicates how much we prefer experiences with higher priority and  $\eta \ll 1$  is a constant which helps to avoid priorities very close to zero. Considering a greedy selection would abandon experiences with low priority, a better approach is to choose experience  $i \in \mathcal{I}$  with probability:

$$P(i) = \frac{p_i}{\sum_{j \in \mathcal{I}} p_j},\tag{7}$$

where  $\mathcal{I}$  is set of all experiences in the buffer. It is possible now to sample a batch of experiences for training using this probability. It removes correlation in the observation sequence and improves sample efficiency of DQN. It is feasible to store all experiences in a buffer sorted by priority, but a more efficient implementation is a sum tree. That is a binary tree, where the value of each root is equal to the sum of its children values (see figure 2). Example of usage is in algorithm 0.

### Algorithm 2 Retrieve node from sum tree in pseudocode

- 1: function GetChild(parent, value)
- 2: **if** parent.left is None **then return** parent
- 3: **if** value < parent.left.value **then**
- 4: **return** GetChild(parent.left, value)
- 5: else
- 6: **return** GetChild(parent.right, value parent.left.value)
- 7: end function

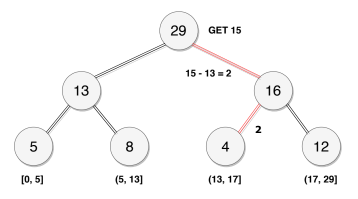


Figure 2: Simple example of sum tree.

# 3 Deep neural networks in RL

As was stated in the previous chapter, tabular methods are very inefficient in large environments. In these instances, it is possible to use deep neural networks which can replace tables. Deep Q networks (DQN) proposed by Googles Deepmind [4] outperformed all previous RL algorithms in playing Atari games. With neural networks also grew the popularity of policy gradient methods where function estimator outputs an action instead of Q values. Note that the most of these methods are general and not necessarily tied to neural networks.

### 3.1 Deep Q network

The neural network takes current state as input and outputs Q value for each possible action. The network is trained using gradients of Q-value in the current state with respect to trainable weights  $\theta$  of our neural network.

$$\delta_t = R_t + \gamma \max_{A_{t+1}} Q^{\theta}(S_{t+1}, A_{t+1}) - Q^{\theta}(S_t, A_t)$$
(8)

$$\theta_{t+1} = \theta_t + \lambda \delta_t \nabla_\theta Q^\theta(S_t, A_t). \tag{9}$$

Gradients are updated in proportion to TD  $\delta_t$ . Unfortunately, this simple DQN agent suffers from a lack of sample efficiency and does not converge well. There are many techniques which can help DQNs to achieve satisfying results.

# 3.2 Target network

Target network is a technique which improves convergence of DQN learning [4]. It uses two neural nets instead of one. The first is trained online network on a batch of data and the second target network is used for predictions during training. After the completion of training on a batch of data, the target network is updated.

$$\theta^{-} = \tau \theta + (1 - \tau)\theta^{-},\tag{10}$$

where  $\theta^-$  is set of trainable weights of the target network,  $\theta$  indicates online network weights and  $\tau \ll 1$  is constant. TD  $\delta$  is now calculated using target network:

$$\delta_t = R_t + \gamma \max_{A_{t+1}} Q^{\theta^-}(S_{t+1}, A_{t+1}) - Q^{\theta}(S_t, A_t). \tag{11}$$

Target network stabilizes training since predicting network does not change after each training step.

# 3.3 Double Q-learning

Classic Q-learning algorithm tends to overestimate actions under certain conditions. Hasselt et al. propose the idea of Double Q-learning which decompose the max operation into action selection and action evaluation [9]. TD is then computed by the following equation.

$$\delta = R_t + \gamma Q^{\theta^-}(S_{t+1}, \underset{A_{t+1}}{\operatorname{argmax}} Q^{\theta}(S_{t+1}, A_{t+1})) - Q^{\theta}(S_t, A_t).$$
(12)

Double DQN outperforms DQN in terms of value accuracy and in terms of policy quality.

# 4 Policy gradient

By this section, the goal of the neural network was predicting values by which the policy was determined. In policy gradient method neural network approximates the policy itself.

$$\theta_{t+1} = \theta_t + \lambda \widehat{\nabla J(\theta_t)} \tag{13}$$

where J is performance measure with respect to our neural network parameters and  $\nabla J(\theta_t)$  is stochastic estimate which approximates gradient of the performance measure. In other words, this method is basically doing stochastic gradient ascent of J with respect to  $\theta$  [10]. Policy gradient methods are outperforming DQNs, especially in continuous action spaces, because their output is directly continuous action instead of Q-value for every possible action.

### 4.1 Actor-Critic

Thanks to predicting action directly, we gain the possibility to predict in continuous action space, but the Q-value which assessed the quality of action in a certain state has been lost. That is why the Actor-Critic framework was created. It uses two separate neural networks - actor which predicts action and critic which assesses action advantage. Concept is visualised in the figure 3.

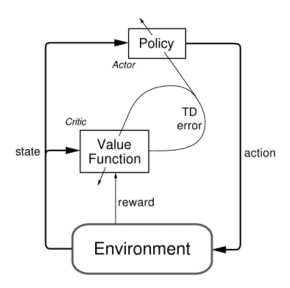


Figure 3: Actor-Critic framework. Source - [3].

Consider critic using Q-values for the update.  $\theta$  and  $\omega$  denote trainable weights of actor and critic, respectively. Critic update is similar to DQN:

$$\delta_t = r_t + \gamma Q^{\omega}(S_{t+1}, \mu^{\theta}(S_{t+1})) - Q^{\omega}(S_t, A_t)$$
(14)

$$\omega_{t+1} = \omega_t + \lambda \delta_t \nabla_\omega Q^\omega(S_t, A_t). \tag{15}$$

Note that instead of  $A_{t+1}$  is now used function  $\mu^{\theta}(S)$ , which is an action estimate by actor neural network. Actor update rule is not so straightforward.

$$\theta_{t+1} = \theta_t + \lambda \nabla_\theta \mu^\theta(S_t) \nabla_a Q^\omega(S_t, A_t)|_{a = \mu^\theta(S_t)}. \tag{16}$$

This equation uses chain rule for derivatives to obtain the gradient of Q-values with respect to trainable weights  $\theta$ . Namely:

$$\frac{\partial Q^{\omega}(S_t, A_t)}{\partial \theta} = \frac{\partial Q^{\omega}(S_t, A_t)}{\partial A_t} \frac{\partial A_t}{\partial \theta}.$$
 (17)

Actor neural network is updated by gradients which change action output to maximize Q-value of the critic [11]. There are other approaches, which doesn't use Q-value as critic assessment, but they rather use so-called advantage [12]. These methods are beyond the scope of this thesis.

### 4.2 Stochastic Actor-Critic

A stochastic Actor-Critic method is frequently used approach. In this method actor outputs parameters of distribution and action itself is sampled within the parameterized distribution. It is standard to use normal distribution and predict the mean and variance of action. The biggest advantage of the normal distribution is that it can be adjusted to use of backpropagation [13]. Another benefit of the stochastic actor is that it does not need any other techniques for action space exploration.

### 4.2.1 Beta distribution

On the other hand, obvious drawback of the normal distribution is that there is always some small probability of sampling an outlier. There is also an issue for bounded action space. When mean value of the normal distribution is close to the boundary, an agent can experience not negligible bias. A solution for both problems is to use beta distribution as a stochastic policy [14]. The beta distribution is defined by the following function:

$$f(x;\alpha,\beta) = \frac{\Gamma(\alpha+\beta)}{\Gamma(\alpha)\Gamma(\beta)} x^{\alpha-1} (1-x)^{\beta-1},$$
(18)

where  $\alpha, \beta \in R_0^+$  are distribution parameters and  $x \in [0, 1]$ .  $\Gamma$  is Euler's gamma function, which extends factorial into the set of real numbers. Beta distribution is shown in the figure 4. The biggest advantage is that Beta distribution is bounded by definition and does not need additional clipping.

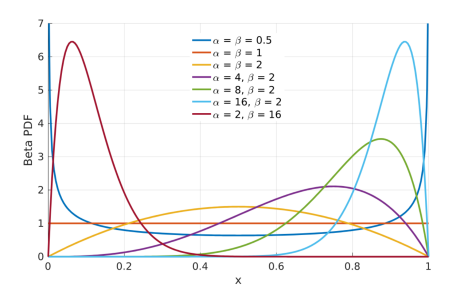


Figure 4: Probability density of beta distribution. Source - [14].

For reinforcement learning is suitable only  $\alpha, \beta \geq 1$ . That makes beta distribution concave and unimodal. This can be ensured by using softplus activation and adding one at the end of actor-network.

# 4.3 Deterministic policy gradients

Deep deterministic policy gradient (DDPG) is one of the methods for exploiting the Actor-Critic framework. Whereas stochastic actor predicts distribution parameters and samples an action DDPG outputs the action directly. Silver [11] has shown that deterministic policy can outperform its stochastic counterparts. A disadvantage of this approach is that it needs an additional policy to ensure action space exploration. Exploration methods are discussed in subsection 4.5.

# 4.4 Wolpetinger policy

Actor-Critic methods and DDPG work well in continuous action spaces, but there are many use cases with large discrete action spaces, such as recommender systems or lidar

planning. Wolpetinger policy is approach how to utilize continuous methods in discrete action space [15]. The whole policy is illustrated in figure 5. An actor doesn't predict action directly, but it predicts so-called proto-action  $\tilde{A}_t$ .

$$\tilde{A}_t = \mu^{\theta}(S_t). \tag{19}$$

Proto action mostly isn't valid action  $\tilde{A}_t \notin \mathcal{A}$ . Thus it is necessary to find the valid action corresponding to the proto action. That is done by computing Euclidean distance to every possible action.

$$\mathcal{A}_{knn} = \underset{a \in \mathcal{A}}{\operatorname{argmin}} |a - \tilde{A}_t|_2. \tag{20}$$

Usually, policy chooses K closest actions to the proto action.  $\mathcal{A}_{knn}$  is the set of closest action to the proto action. The whole set is then assessed by critic and action with highest Q-value is finally picked.

$$A_t = \underset{a \in \mathcal{A}_{knn}}{\operatorname{argmax}} Q^{\omega}(S_t, a) \tag{21}$$

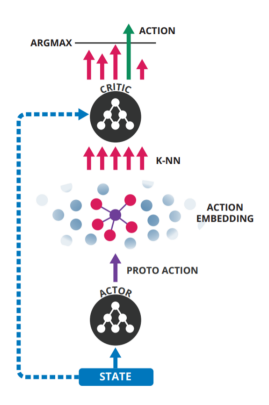


Figure 5: Wolpetinger policy can be described in 3 steps. 1) Actor outputs an action 2) The K nearest valid actions are found 3) Choose the action which is assessed as the best by the critic. Source - [15].

### 4.5 Parameter and action space noise

In large action, space is crucial to emphasize agents exploration. Bad exploration can cause that agent converges prematurely and ends up in a local optimum. DDPG commonly uses stochastic policy to modify actors actions slightly.

$$\hat{A}_t = \mu^{\theta}(S_t) + \mathcal{N}(0, \sigma^2) \tag{22}$$

where  $\mathcal{N}$  is the normal distribution with mean value equal to zero and variance, which is reducing during the training and  $\hat{A}_t$  is perturbed action. Action space noise helps agent to explore the environment.

Another approach is to apply noise directly to actors weights. It can sometimes lead to more consistent exploration and richer behaviors [16].

$$\hat{\theta} = \theta + \mathcal{N}(0, \sigma^2) \tag{23}$$

where  $\hat{\theta}$  is a so-called perturbed actor, which is interacting with an environment. The major issue of parameter space noise is that it is much harder to tune. When we use action space noise, it is easy to estimate its impact on actions (differences between both approaches can be seen in the figure 6). Because of an unpredictable influence of parameter space noise is necessary to use adaptive noise scaling.

$$d = |\hat{A}_t - \mu^{\theta}(S_t)|_2 \tag{24}$$

$$\sigma_{t+1} = \begin{cases} \kappa \sigma_t & \text{if } d \le T \\ \frac{1}{\kappa} \sigma_t & \text{otherwise} \end{cases}$$
 (25)

where  $\kappa$  is scaling factor slightly bigger than one and T is the threshold value, which has to be tuned to a specific environment.

When is necessary to explore action space near to some specific action or include momentum of an environment, it is possible to use Ornstein-Uhlenbeck random process [17].

$$\hat{A}_t = \mu^{\theta}(S_t) + \nu(\rho - \mu^{\theta}(S_t)) + \phi \mathcal{N}(0, 1), \tag{26}$$

where  $\nu, \phi \in [0, 1]$  are constants of the random process and  $\nu$  is mean value around which we want to explore the action space. When  $\nu = 0$  it is basic exploration as in expression (22).

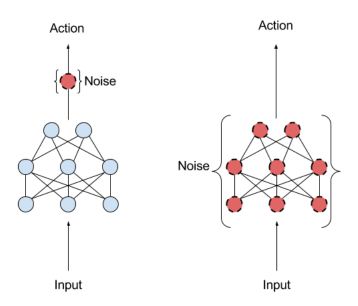


Figure 6: Left - action space noise, Right - parameter space noise. Source - [16].

# 5 Experiment

The experiment aims at using reinforcement learning algorithms for solid-state lidar with steerable rays and a limited number of rays. At first, it was necessary to create an environment where the agent can learn and evaluate [6]. The lidar-gym environment is written in python 3 based on OpenAI gym interface [18]. It uses point clouds from Kitti dataset drives[5]. One episode of learning in environment corresponds to one drive in Kitti dataset. Large point clouds from drives are processed into 3D voxel maps by C++ package [19], which also provides ray tracing engine for the environment. Every voxel map is a 3D array containing real numbers which correspond to the occupancy confidence c of each voxel.

$$c > 0$$
 occupied voxel  
 $c = 0$  unknown occupancy (27)  
 $c < 0$  empty voxel.

The environment also offers visualization of actions using Mayavi [20] and ASCII art. Agents use neural networks as function estimators, which are handled by Tensorflow [21] and Keras [22].

### 5.1 Environments

Lidar-gym implements several environments, which follow the same template with different sizes. Observation space is a local cutout of voxel map, which provides occupancies from sensor's sparse measurements. The sensor is located in the quarter of x-axis and half of y-axis and z-axis of a local cutout. Action space is divided into two parts. First part is dense voxel map reconstructed from observations (sparse measurements). The second part of action space are directions of measuring rays. Each ray has own azimuth and elevation. Environment expects directions in the format of a 2D array of booleans, where true means fired ray. Environments reward is negative logistic loss -L (28). Lidar-gym defines environments, with parameters described in table 1, and visualized in figure 7.

<sup>\*</sup>Using GPU Nvidia 1080Ti.

Name of environment	Large	Small	Toy
Voxel map size [voxels]	$320 \times 320 \times 32$	$160 \times 160 \times 16$	$80 \times 80 \times 8$
Lidar FOV [°]	$120 \times 90$	$120 \times 90$	$120 \times 90$
Densitiy of rays	$160 \times 120$	$120 \times 90$	$40 \times 30$
Lidar range [m]	42	42	42
Number of rays	200	50	15
Voxel size [m]	0.2	0.4	0.8
Episode training time [min]*	120	15	1.5

Table 1: Description of environments

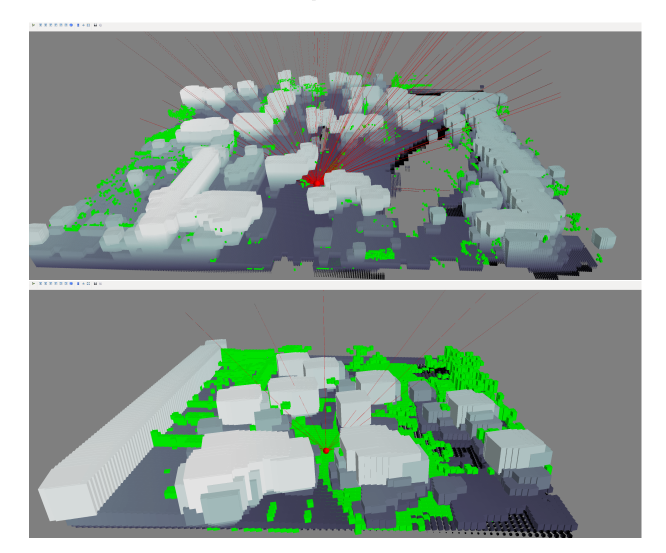


Figure 7: Visualization of two environments: Top - Large, bottom - Toy. Ground truth map is green, reconstructed map is in grey palette, sensor position and rays are red.

Due to the high time complexity were all experiments conducted in toy environment. RL agents need significantly more training steps than supervised agents. In OpenAI baselines, [23] are RL agents trained for over million timestamps. One drive in Kitti dataset has on average 200 timestamps. All agents were trained and evaluated on different drives from city part of the dataset.

### 5.2 Mapping agent

Mapping agent is based on work of Zimmermann et al. [2]. It uses a convolutional neural network (CNN) for reconstructing dense map from sparse measurements. 3D convolutional layers are used to learn the features and max-pooling layers to avoid overfitting. Whole CNN architecture is described in figure 8.

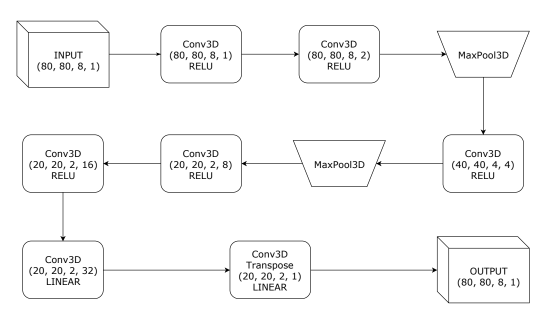


Figure 8: Mapping network architecture

For gradient descent is used logistic loss L between ground truth map Y and predicted dense map  $\hat{Y}$ .

$$L(Y, \hat{Y}) = \sum_{i} w_i \log(1 + \exp(Y_i \hat{Y}_i))$$
(28)

where w are weights which balance importance of occupied and unoccupied voxels. Un-

fortunately, a naive implementation of this loss function is computationally inconvenient and often cause numerical issues as underflow or overflow. To stabilize training following modified loss was used [24].

$$a_i = Y_i \hat{Y}_i$$

$$b_i = \max(0, a_i)$$

$$L = \sum_i w_i (b_i + \log(\exp(-b_i) + \exp(a_i - b_i))).$$
(29)

At first, supervised mapping agent with random ray planning, is trained. Reconstructions of the supervised agent are then used for training RL planning agents and after that is mapping agent retrained with RL agent picking the rays.

### 5.3 Discrete planning agent

Insomuch as the input of the environment for a direction of the rays is 2D binary array, first try is to use a discrete agent. DQN is the most used option for discrete action space, but in this use case, it requires some tweaks. Note that number of possible actions is extremely large. Even in toy environment it is  $\binom{40\times30}{15}\approx10^{34}$  of actions. Thus it is necessary to emphasize on action space exploration. Further arises the problem with  $\epsilon$ -greedy policy, because we are unable to process all possible actions and pick one with the biggest Q-value. To solve this issue is considered one ray as an action and for K rays is TD from (8) now computed as:

$$q(S_t, A_t) = \max_{A_t}^K Q^{\theta}(S_t, A_t)$$

$$\delta_t = R_t + \gamma \overline{q}(S_{t+1}, A_{t+1}) - \overline{q}(S_t, A_t)$$
(30)

where  $\overline{q}$  is average Q value over K actions with maximal Q values. DQN agent implements all available features described in the theoretical part of this thesis as Prioritized experience replay, target network, and double Q learning. Exploration is ensured by action space noise. Parameter values of agent are shown in table ?? and neural network architecture in figure 9. For gradient descent is used Adam optimizer with learning rate  $\lambda$ .

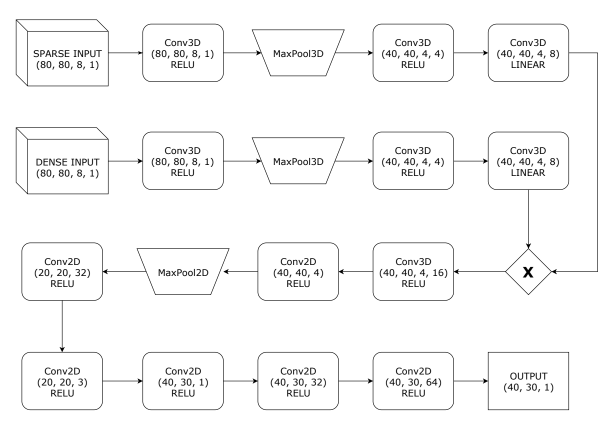


Figure 9: DQN architecture.

### 5.3.1 Wolpetinger policy

Wolpetinger policy is currently the state-of-the-art method for discrete action spaces because it utilizes actor-critic to continuous methods. It was tested on recommender systems with over a million discrete actions [15]. That is large action space, but significantly smaller than action space of Lidar-gym environment. The obvious problem comes in place during action embedding in formula (20). It is not possible to compute KNN with so many possible actions. When is KNN substituted by picking rays with the highest value, the agent does not converge well. It abandons some possible actions very soon and gets stuck in local optimum.

### 5.4 Continuous planning agent

The discrete action output was substituted by continuous action, which is then mapped into a 2D binary array. This will allow the agent to avoid extremely large discrete action space. Thank this substitution it is possible to exploit actor-critic framework. The output of actor is now 2 by K array where the first row is the elevation and second is the azimuth of each ray. The last layer of the actor-network is tanh function, so its output is an element of [-1,1]. As training method is used DDPG. The neural network architecture is described in the figure 10.

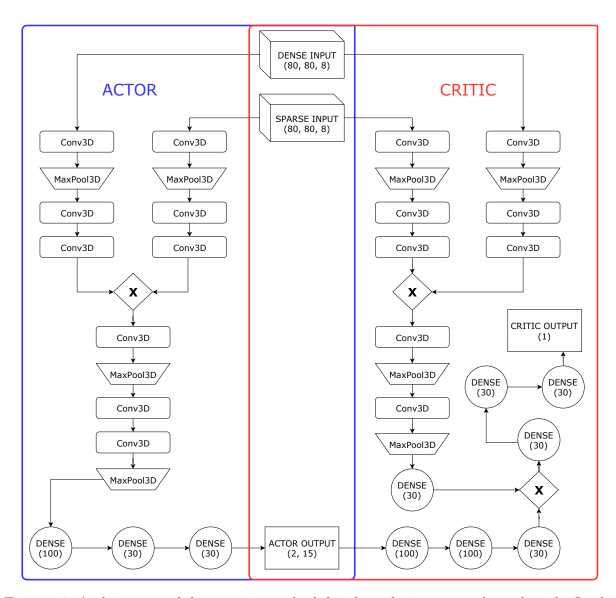


Figure 10: Architecture of the actor is on the left side and critic is on the right side. In the middle can be seen shared inputs and outputs.

To explore action space correctly, it is necessary to apply Ornstein-Uhlenbeck random process. When only Gaussian noise is added, actions tend to converge into the corners very fast as in figure 11. For both actor and critic is used Adam optimizer with learning rates  $\lambda_a$ ,  $\lambda_c$ . To stabilize critic learning is used target network. The continuous agent also uses prioritized experience replay.

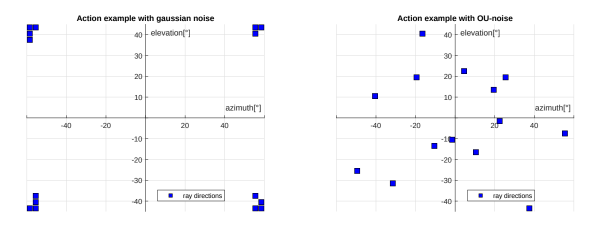


Figure 11: Difference between actions made by agents with different exploration methods. Left figure is an example of action made by an agent with Gaussian noise used for exploration. The action in the right figure is taken by agent trained using Ornstein-Uhlenbeck noise.

# 5.5 Stochastic planning agent

There is an obvious issue with the deterministic actor. For efficient exploring of ground truth map is required to hit as many unique voxels as possible. Thus making several similar actions in a row is not a good strategy. Unfortunately, except first few timestamps of every drive, there is not a big difference between two subsequent observations. It is hard for a neural network to make two different outputs for two similar inputs. The solution to this problem could be a stochastic agent. Stochastic agent outputs parameters of beta distribution and preserves actor-critic framework. The architecture of stochastic agent is similar to DDPG from the previous subsection, only difference is that agent now outputs just two values - distribution parameters. Action is then sampled with distribution probabilities. Stochastic actor output can be seen in figure 12.

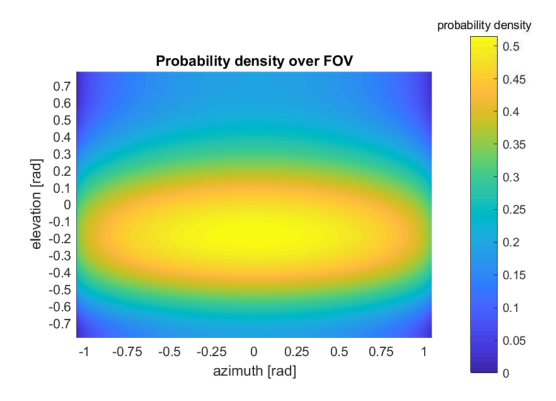


Figure 12: Probability density of picking ray, with specific azimuth and elevation.

# 5.6 Comparison of methods

Several methods were tried. Unfortunately most of the methods are not doing well in Lidar-gym environment. The worst performance had the DQL. That is probably caused by an extremely large set of possible actions. Workaround made in formula (30), did not help agent to converge. Continuous DDPG agent also did not converge successfully. It is hard for DDPG agent to change direction of rays efficiently in subsequent timestamps and it often makes several same actions in a row, that degrades performance a lot. Only stochastic agent was able to outperform a random agent. Summary of agents performances is made in figure 13. It turns out that it is very difficult to get satisfying results. Stochastic agent converge, but it does not use any kind of advanced strategy. Action space of this agent does not even allow him to do any sophisticated action. Therefore were also conducted experiments with extended stochastic agent, which have used beta distribution for each ray separately, but with no significant success. Another advantage of simple stochastic agent is its scalability. This agent can be adjusted to the large environment much easier than any other used agent. Parameters used for these agents are described in table 2.

Parameter	Value
$\gamma$	0.09
$\lambda_{critic}$	0.001
$\lambda_{actor}$	0.0001
au	0.01
Memory size	4096
Batch size	8

Table 2: Parameters used for learning.

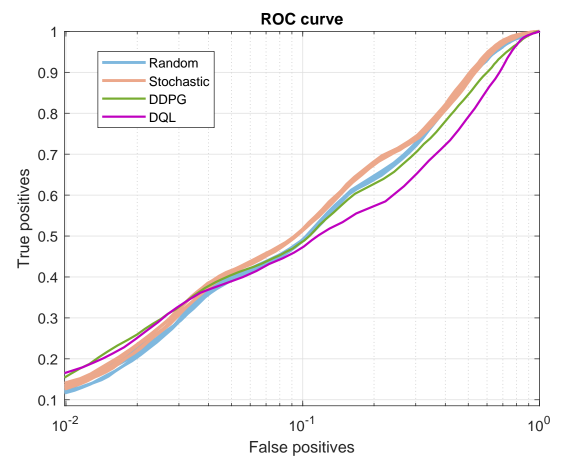


Figure 13: False positives were calculated using ground truth map and true positives using all voxels, which could be possibly hit by sensor. Five different evaluations are displayed for non-deterministic agents.

### 6 Conclusion

At the beginning of the thesis was defined the concept of reinforcement learning. Further, methods as DQL and DDPG were discussed. Also several methods which helps neural networks to converge were described. The experiment focused on training planning agents for choosing rays. All deterministic agents performed poorly in this specific task. Stochastic agent successfully outperformed random planner. Action space size and time-complexity were two major blockers during the training. Any of trained RL agents can not compete with prioritized greedy planner proposed by Zimmermann et al. [2].

### 6.1 Future work

Experiment which would be worth conducting is an agents, which stands between simple and extended stochastic agent. Extended stochastic agent has action space consisting of 60 real numbers (15 rays with azimuth and elevation and for each alpha, beta parameters). That is very likely too much for network architecture used by agent. On the other hand, when only one distribution is outputted for all rays, it does not allow agent to create advanced policies. Solution could be to output for example three different distributions and each use for five rays. Another improvement could be achieved by adjusting neural network architecture. Especially splitting and merging of neural network, can have significant impact on performance.

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# Appendix A CD Content

In Table 3 are listed names of all root directories on CD.

Directory name	Description
thesis	the thesis in pdf format
thesis_sources	latex source codes

Table 3: CD Content

# Appendix B List of abbreviations

In Table 4 are listed abbreviations used in this thesis.

Abbreviation	Meaning
API	application programming interface

Table 4: Lists of abbreviations