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Contents

Pı	Preface						
A	bstra	act	iv				
1	Intr	atroduction					
2	Pro	blem description	4				
	2.1	Problem description	4				
		2.1.1 Reinforcement Learning	4				
	2.2	Risk	5				
		2.2.1 Conditional Value-at-Risk (CVaR)	6				
3	The	e algorithm	8				
	3.1	Off-policy Deterministic Actor-Critic	8				
		3.1.1 Preliminaries	8				
		3.1.2 Deterministic policy gradients	9				
		3.1.3 Off-policy Deterministic actor-critic	9				
	3.2	The algorithm: Distributional off-policy deterministic actor critic $$.					
		3.2.1 Distributional Critic	10				
		3.2.2 Actor	11				
	3.3	Other distortion risk measures	13				
	3.4	Truncation	13				
	3.5	Chow	14				
	3.6	Technical Details of the algorithm	16				

4	Bat	ch RL				
	4.1	Details	s of VAE	19		
5	Res	ults				
	5.1	Current results Car				
		5.1.1	Case 1: No velocity penalization	21		
		5.1.2	Case 2: Velocity penalization with probability 1 \dots	22		
		5.1.3	Case 3: Velocity penalization with probability P $\ \ldots \ \ldots$	23		
	5.2	Current results Batch RL HalfCheetah				
Bi	Bibliography					
\mathbf{A}				31		
	A.1	Experiment details				
	A.2	Distributional RL				
		A.2.1	Example showing interest in learning the distribution $\ \ . \ \ . \ \ .$	31		
		A.2.2	Distributional Bellman Operator	33		
		A.2.3	Quantile approximation	35		
		A.2.4	Quantile projection:	36		
		A.2.5	Quantile Regression	36		
		A.2.6	Quantile Regression Temporal Difference Learning	37		

Preface

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Abstract

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Chapter 1

Introduction

In sequential decision-making problems, an agent interacts with an environment by selecting actions and in turn, it observes the state transitions of the system and it receives a reward. The decision-maker or *agent* uses a performance measure to assess how good actions were. The most common optimization criterion is the expectation of the cumulative reward collected by the agent. This leads to the so called *risk-neutral* behavior.

When an agent interacts with the environment, different sources of irreducible randomness are introduced due to modelling errors or inherent stochasticity of the same. Notion of risk in RL is related to the fact that even an optimal policy with respect to the expected return may perform poorly in some cases. Since maximizing the expected return does not necessarily imply the avoidance of rare occurrences of large negative outcomes, we need other criteria to evaluate risk. (García and Fernández, 2015).

Risk-sensitive decision-making provides a promising approach to compute robust and safe policies, essential in safety-critical applications, such as autonomous navigation. However, finding computationally tractable and conceptually meaningful methodologies for such a goal is non-trivial and still a challenge.

We can segment RL algorithms accounting for risk in two main categories: the first transform the optimization criterion to introduce the concept of risk, whereas the second, modifies the exploration process to avoid exploratory actions that can lead to undesirable situations.

We will focus on the first category, which can be divided into 3 subcategories: worst-case criterion, constrained criterion and risk-sensitive criterion.

Worst-case or minimax criterion has been studied by Heger (1994), Coraluppi and Marcus (2000) and Coraluppi and Marcus (1999), in which the objective is to compute a control policy that maximizes the expected return with respect to the worst case scenario. In general, minimax criterion has been found to be too restrictive as it takes into account severe but extremely rare events which may never occur.

Constrained criterion aims to maximize the expected return while keeping other types of expected utilities higher than some given bounds (Altman, 1993). It may be seen as finding the best policy π in the space of considered safe policies. This space may be restricted by different constraints such as ensuring that the expectation of costs (Geibel, 2006) or the variance of return doesn't exceed a given threshold (Tamar et al., 2012). This constraint problems can be converted to equivalent un-

constrained ones by using penalty methods or a Lagrangian approach.

Finally, risk-sensitive criterion uses other utility metrics to be maximized instead of expectation of cumulative rewards. Lot of research has been done using exponential utility functions (Howard and Matheson, 1972; Chung and Sobel, 1987).

Our approach aims to maximize another risk metric, namely the Conditional Value at Risk (CVaR). Non-formally, CVaR of a return distribution at confidence level α can be defined as the expected cumulative reward of outcomes worse than the α -tail of the distribution. CVaR has recently gained a lot of popularity due to its mathematical properties (Artzner et al., 1999), which makes its computation and optimization much easier than for other metrics (Rockafellar and Uryasev, 2000). For example, it has recently been identified by Majumdar and Pavone (2020) as a suitable metric for measuring risk in robotics.

CVaR optimization aims to find the parameters θ that maximizes $\text{CVaR}_{\alpha}(Z)$, where the return distribution Z is parameterized by a controllable parameter θ such that: $Z = f(X; \theta)$.

In the simplest scenarios, where X is not dependant on θ CVaR optimization may be solved using various approaches such as in Rockafellar and Uryasev (2000). However, in RL, the tunable parameter θ also affects the distribution of X and therefore, the standard existing approaches for CVaR optimization are not suitable. Additionally, most of the work with such a goal has been done in the context of MDPs when the model is known by using dynamic programming methods (Chow et al., 2015; Petrik and Subramanian, 2012), and not much research has been done for the RL framework.

Under the RL framework, Morimura et al. (2010) and Morimura, Tetsuro and Sugiyama, Masashi and Kashima, Hisashi and Hachiya, Hirotaka and Tanaka (2010) focused on estimating the density of the returns to handle CVaR risk criteria. However, the resulting distributional-SARSA-with-CVaR algorithm they propose has proved effectiveness only in very simple and discrete MDPs.

Tamar et al. (2015a) proposed a policy gradient (PG) algorithm for CVaR optimization, by deriving a sampling based estimator for the gradient of CVaR and using it to optimize the CVaR via stochastic gradient descent. However, they only considered continuous loss distributions and they used empirical α -quantile to estimate VaR $_{\alpha}$ which is known to be a biased estimator for small samples.

Chow and Ghavamzadeh (2014) also proposed a PG algorithm for mean-CVaR optimization, which has several disadvantages also shared with Tamar et al. (2015b). First, by definition of PG algorithms, they suffer from high variance on the gradient estimates, especially when the trajectories are long. This high variance is even more exacerbated when using very low quantiles α for the CVaR since the averaging on the rewards is effectively only on α N samples. Second, they are both very sample inefficient since only the rewards below the quantile are used to compute the gradient. Third, they are both trajectory-based (not incremental), i.e they only update after observing one or more full trajectories.

Chow and Ghavamzadeh (2014) also proposed both a trajectory-based and incremental actor-critic approaches which help in reducing the variance of PG algorithms. However, they require state augmentation of the original MDP formulation to a state-space $\mathcal{X} \times \mathcal{S}$ where $\mathcal{S} \in \mathbb{R}$ is an additional continuous state that allows to reformulate the Lagrangian objective function as an MDP.

It is also important to be noticed that all the aforementioned algorithms are onpolicy approaches. This is first another source of sample inefficiency because they cannot exploit data from experts or other sources. Additionally they constraint the learning process to happen online while interacting with the environment, which in real-case scenarios, it is paradoxically risky.

The CVaR optimization algorithm that we introduce has 2 main properties that so far, we do not have knowledge any other algorithm for CVaR optimization in the RL setting has; namely, being an off-policy algorithm and using deterministic policies.

Being off-policy means that the algorithm can learn the optimal policy from data collected from another policy. Hence, we can use a more exploratory behavior policy to interact with the environment and collect observations, and then use them to train the optimal target policy. Additionally to the fact of making exploration easier compared to on-policy algorithms, being off-policy allows the possibility to learn from data generated by a conventional non-learning controller or from a human expert, setting mostly called in literature as Batch RL or offline RL. This application is really appealing when working with risky environments, when we do not want to expose the learning agent to its risks at the earlier stages of the learning process.

The second property is the fact that we use a target policy which is deterministic. Using stochastic policies is not natural in many applications and it could increase the variance of the return . When again, working in an stochastic environment and with the goal of finding risk-aware policies, using a policy which is stochastic sounds quite counterintuitive.

In this thesis, we propose a CVaR optimization approach based on a model-free, off-policy deterministic actor-critic algorithm using deep function approximators. Instead of empirically estimating the VaR from the observed rewards or using the CVaR Bellman equation, we build upon recent research in distributional RL (Bellemare et al., 2017; Dabney et al., 2018a,b) to estimate the full *value distribution* (i.e. the distribution of the random return received by a RL agent). A critic learns a parameterization of the value distribution and the actor, is trained towards policies that maximize the CVaR of the returns by sampling from the parameterized value distribution and performing stochastic gradient ascent.

We show algorithm performance in two different settings. First, in the most common off-policy RL setting in which the environment simulator is given and the agent can interact with it to collect new data from time to time. Second, and most powerful, the agent is trained completely offline by using an external dataset and no further interaction with the environment is allowed. For the 2 settings, the algorithm learns risk-sensitive policies in complex high-dimensional, continuous action spaces such as for some modified environments from the Open AI Gym toolkit and from D4RL, a recent suite of tasks and datasets for benchmarking progress in offline RL.

Feinberg (2008)/Taleghan and Dietterich (2018)/nothing

Chapter 2

Problem description

2.1 Problem description

Standard reinforcement learning approaches aim to find policies which maximize the expected cumulative reward. However, this approach neither takes into account variability of the reward around its mean neither sensitivity of the policy to modelling errors.

In this thesis, we change the objective function of the standard RL approach to one that optimizes another metric of the reward distribution which takes into account the risk of the actions taken by the agent. While several metrics have been designed to assess risk, we will focus on a particular risk metric called Conditional Vale at Risk (CVaR).

We, thereby, present a RL algorithm that aims to find policies with optimal CVaR.

2.1.1 Reinforcement Learning

Reinforcement learning is an approach to learn a mapping from states to actions so as to maximize a numerical reward signal. The learner or agent is not told which actions to take but instead must discover which actions yield the most reward by trying them. In most of the cases, actions may affect not only the immediate reward but also the next state and consequently, all subsequent rewards. These two characteristics—trial-and-error search and delayed reward—are the two most important distinguishing features of reinforcement learning (Sutton and Barto, 1998).

Markovian Decision Processes (MDPs)

We formalize the problem of RL by using the framework of Markov decision processes (MDPs) to define the interaction between a learning agent and its environment in terms of states, actions and rewards.

MDPs are discrete-time stochastic control processes which provide a mathematical framework for modeling sequential decision making in situations where outcomes are partly random and partly under the control of a decision maker and where 5 2.2. Risk

actions influence not only immediate rewards, but also future ones.

An MDP is defined by a tuple (S, A, R, P, γ) , where S, A are the state and action spaces respectively, R(x, a) is the reward distribution, $P(\cdot|x, a)$ is the transition probability distribution and $\gamma \in [0, 1]$ is the discount factor.

State transitions of an MDP satisfy the Markov property, in which the set of transition probabilities to next states depend only on the current state and action of the system, but are conditionally independent of all previous states and actions. Hence, the state must provide information about all aspects of the past agent-environment interaction that make a difference for the future.

Solving an MDP involves determining a sequence of policies π (mapping states to actions) that maximize an objective function. A commonly considered class of policies in literature are the class of Markovian policies Π_M where at each time-step t the policy π_t is a function that maps state x_t to the probability distribution over the action space \mathcal{A} . In the special case when the policies are time-homogeneous, i.e. $\pi_j = \pi \ \forall j \geq 0$ then the class of policies is known as stationary Markovian $\Pi_{M,S}$. This set of policies, under which actions only depend on current state information and its state-action mapping is time-independent, makes the problem of solving for an optimal policy more tractable and common solution techniques involve dynamic programming algorithms (Bertsekas, 1995) such as Bellman iteration.

When the objective function is given by a risk-neutral expectation of cumulative reward, i.e.:

$$\min_{\pi \in \Pi_H} \mathbb{E}\left[\sum_{t=0}^{\infty} \gamma^t R(x_t, a_t) | x_0, a_t \sim \pi_t(\cdot | h_t)\right]$$
(2.1)

where Π_H represents the set of all history-dependant policies, the Bellman's principle of optimality (Bertsekas, 1995) shows that the optimal policy lies in the class of stationary Markovian policies $\Pi_{M,S}$

When dealing with other types of objective functions that aim towards more risk-sensitive policies, these nice properties doesn't normally hold and require extra mathematical formulations, such as MDP state augmentation (Chow et al., 2015).

2.2 Risk

Standard reinforcement learning agents aim to maximize the expected cumulative reward and hence do not take risk into account. In some scenarios the shape of the reward distribution might be unimportant, since highly different distributions still can have same expectation. However, in real world scenarios, in when catastrophic losses can occur, risk must be taken into account.

We can find 3 types of strategies with respect to risk, namely risk neutral, risk averse and risk seeking

As an example, suppose a participant in a game is told to choose between two doors. One door hides 1000CHF and the other 0CHF. The host also allows the contestant to take 500CHF instead of choosing a door. The two options (choosing between door 1 and door 2, or taking 500CHF) have the same expected value of 500CHF. But it can clearly be seen that the risk among two options is different. Since the expected value is the same, risk neutral contestant is indifferent between these choices. A risk-seeking contestant will maximize its utility from the uncertainty and

hence choose a door, whereas the risk-averse contestant will accept the guaranteed $500\mathrm{CHF}$.

In this thesis we present an algorithm to find risk-sensitive policies that maximize the CVaR of the reward distribution.

2.2.1 Conditional Value-at-Risk (CVaR)

Let Z be a bounded-mean random variable, i.e $\mathbb{E}[|Z|] < \infty$, on a probability space $(\Omega, \mathcal{F}, \mathbb{P})$, with cumulative distribution function $F_Z(z) = \mathbb{P}(Z \leq z)$. We interpret Z as the reward distribution and for convenience we assume that is a continuous random variable meaning that $F_Z(z)$ is everywhere continuous. The value-at-risk (VaR) at confidence level $\alpha \in (0,1)$ is the α -quantile of Z, i.e.:

$$VaR_{\alpha}(Z) = F_Z^{-1}(\alpha) = \inf\{z \mid F_Z(z) \ge \alpha\}$$
(2.2)

The conditional value-at-risk (CVaR) at confidence level $\alpha \in (0,1)$ is defined as the expected reward of outcomes worse than the α -quantile (VaR $_{\alpha}$):

$$CVaR_{\alpha}(Z) = \mathbb{E}[Z|Z \le VaR_{\alpha}] = \frac{1}{\alpha} \int_{0}^{\alpha} F_{Z}^{-1}(\tau)d\tau = \frac{1}{\alpha} \int_{0}^{\alpha} VaR_{\tau}(Z)d\tau \qquad (2.3)$$

The last equality, known as the Acerbi's integral formula for CVaR (Proposition 3.2 in Acerbi and Tasche (2002)), interprets CVaR_{α} as the integral of all the quantiles below the corresponding quantile level α and will become useful later.

While both VaR and CVaR are risk measures, CVaR has superior mathematical properties (monotonicity, translation invariance, positive homogeneity and subadditivity; see Artzner et al. (1999)) which makes CVaR computation and optimization far easier compared to VaR (Rockafellar and Uryasev, 2000). In addition, CVaR takes into account the possibility of tail events where losses exceeds VaR whereas VaR is incapable of distinguishing situations beyond it.

Rockafellar and Uryasev (2000) also showed that CVaR is equivalent to the solution of the following optimization problem:

$$CVaR_{\alpha}(Z) = \max_{\nu} \left\{ \nu + \frac{1}{\alpha} \mathbb{E}_{Z}[[Z - \nu]^{-}] \right\}$$
 (2.4)

where $(x)^{-} = \min(x, 0)$.

In the optimal point it holds that $\nu^* = \operatorname{VaR}_{\alpha}(Z)$.

A useful property of CVaR, is its alternative dual representation Artzner et al. (1999):

$$CVaR_{\alpha}(Z) = \min_{\xi \in U_{CVaR}(\alpha, \mathbb{P})} \mathbb{E}_{\xi}[Z]$$
 (2.5)

where $\mathbb{E}_{\xi}[Z]$ denotes the ξ -weighted expectation of Z, and the risk envelope U_{CVaR} is given by:

$$U_{\text{CVaR}}(\alpha, \mathbb{P}) = \left\{ \xi | \xi(w) \in \left[0, \frac{1}{\alpha}\right], \int_{w \in \Omega} \xi(w) \mathbb{P}(w) dw = 1 \right] \right\}$$
 (2.6)

7 2.2. Risk

Thus, the CVaR of a random variable may be interpreted as the worst case expectation of Z, under a perturbed distribution $\xi \mathbb{P}$.

Our goal in this thesis is to find policies that maximize the CVaR of the return distribution i.e.:

$$\arg \max_{\pi} \text{CVaR}_{\alpha}[Z(x, \pi(x))] \quad \forall x \in \mathcal{X}$$
 (2.7)

To this end, we present an algorithm for CVaR optimization using a distributional variant of the deterministic policy gradient algorithm. The word distributional emphasizes that our approach takes inspiration from the recent advances in distributional RL (Bellemare et al., 2017; Dabney et al., 2018a,b) to estimate the full value distribution (i.e. the distribution of the random return received by a RL agent). Having an estimation of the true value distribution, we compute the CVaR of the current policy via sampling from the estimated value distribution and we move the current policy towards maximizing it via stochastic gradient ascent.

The CVaR dual formulation and the Acerbi's integral formula 2.3 presented above, will be useful to derive our approach to compute the CVaR from parameterized inverse cumulative distributions.

should I repeat all this? Specify that they are NN's actor/critic?

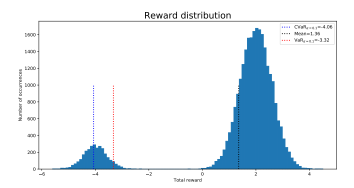


Figure 2.1: Histogram showing VaR, mean and CVaR of a sampled probability distribution. The image shows that the mean doesn't take into account low probability events and the flow of the VaR as a risk metric, since we could move the left-most mode to minus infinity and the VaR will remain the same, whereas the CVaR would change with the shift.

Chapter 3

The algorithm

We introduce an off-policy, model-free algorithm for CVaR optimization using deep function approximators that can learn policies in high-dimensional, continuous action spaces. Our work is based on the deterministic policy gradient algorithm. Specifically, we use an actor-critic approach: the critic uses a distributional variant of RL and it is trained to estimate the whole value distribution, whereas the actor is trained via gradient ascent to maximize the CVaR of this distribution. In the following we briefly describe the standard DPG algorithm 3.1, then we introduce our distributional approach and how we use it to obtain risk-sensitive policies 3.2 and in the last section, we focus more on implementation details of the algorithm, especially explaining more about the neural networks we use and providing a pseudocode 3.6.

decide

Extensive and more theoretical information on the distributional RL approach is addressed in chapter ?? / appendix A

3.1 Off-policy Deterministic Actor-Critic

change title not to repeat with subsection?

3.1.1 Preliminaries

Goal in standard RL is to learn a policy π^* which maximizes the expected return or (discounted) cumulative reward R collected by the agent when acting in an environment E starting from any initial state x. Action-value function for policy π , $Q^{\pi}(x,a)$, is used in many RL algorithms and describes the expected return after taking an action a in state x, and thereafter following policy π :

$$Q^{\pi}(x_t, a_t) = \mathbb{E}_{r_{i \ge t}, x_{i > t} \sim E, a_{i > t} \sim \pi} \left[\sum_{i=t}^{T} \gamma^{(i-t)} r(x_i, a_i) \right]$$
(3.1)

Bellman's equation describes this value Q using the recursive relationship between

the action-value of a state and the action-values of its successor states:

$$Q^{\pi}(x_t, a_t) = \mathbb{E}_{r_t, x_{t+1} \sim E} \left[r(x_t, a_t) \right] + \gamma \mathbb{E}_{a_{t+1} \sim \pi} \left[Q^{\pi}(x_{t+1}, a_{t+1}) \right]$$
 (3.2)

DPG algorithm (Silver et al., 2014) is characterized for using deterministic policies $a_t = \mu_{\theta}(x_t)$.

In general, behaving according to a deterministic policy does not ensure adequate exploration and may lead to suboptimal solutions. However, if the policy is deterministic, we can remove the inner expectation with respect to a_{t+1} in 3.2 and then, crucially, the expected cumulative reward in the next-state depends only on the environment and not on the policy distribution used to create the samples.

$$Q^{\pi}(x_t, a_t) = \mathbb{E}_{r_t, x_{t+1} \sim E} \left[r(x_t, a_t) + \gamma Q^{\pi}(x_{t+1}, \pi(x_{t+1})) \right]$$
(3.3)

This means that it is possible to learn the value function Q^{π} off-policy, i.e. using environment interactions which are generated by acting under a different stochastic behavior policy β (where $\beta \neq \pi$) which ensures enough exploration. An advantage of off-policy algorithms is that we can treat the problem of exploration independently from the learning algorithm.

To learn the optimal policy, Q-learning Watkins and Dayan (1992), a commonly used off-policy algorithm, first learns the optimal value function Q^* by iteratively applying the Bellman optimality operator to the current Q estimate:

$$Q(x_t, a_t) \leftarrow \mathbb{E}_{r_t, x_{t+1} \sim E} \left[r(x_t, a_t) + \gamma \max_{a_{t+1}} Q(x_{t+1}, a_{t+1}) \right]$$
(3.4)

which is a contraction mapping proved to converge exponentially to Q^* , and then derives the optimal policy π^* from it via the greedy policy $a^* = \operatorname{argmax} Q^*(x, a)$.

When dealing with continuous actions, it is not possible to apply Q-learning straightforward because finding the greedy policy requires an optimization of a at every timestep, which is too slow to be practical with large action spaces. In this case, policy gradient methods are used in which a parameterized policy is learnt to be able to select actions without consulting the value function.

3.1.2 Deterministic policy gradients

The deterministic policy gradient described by Silver et al. (2014) updates the parameters of the deterministic policy π_{θ} via gradient ascent to maximize an objective function $J(\pi_{\theta})$:

$$J(\pi_{\theta}) = \mathbb{E}_{x \sim \rho^{\pi}} [Z(x, \pi_{\theta}(x))] \tag{3.5}$$

$$\nabla_{\theta} J(\pi_{\theta}) = \mathbb{E}_{x \sim \rho^{\pi}} \left[\nabla_{\theta} \pi_{\theta}(x) \nabla_{a} Q^{\pi}(x, a) |_{a = \pi_{\theta}(x)} \right]$$
(3.6)

where ρ^{π} is the discounted state distribution when acting under policy π .

3.1.3 Off-policy Deterministic actor-critic

When we both learn approximations of Q and policy, the method is called deterministic actor-critic. The actor is the learned policy which is updated with respect to the current value estimate, or critic. Sutton and Barto (1998).

In the off-policy setting, the critic parameterized by θ^Q estimates the action-value function $Q^{\pi}(x, a)$ off-policy from trajectories generated by a behavior policy β (discussed in 3.1.1) using the Bellman equation. It learns by minimizing the loss:

$$\mathcal{L}(\theta^Q) = \mathbb{E}_{x_t \sim \rho^\beta, a_t \sim \beta, r_t \sim E} \left[(Q(x_t, a_t | \theta^Q) - y_t)^2 \right]$$
(3.7)

$$y_t = r(x_t, a_t) + \gamma Q(x_{t+1}, \pi(x_{t+1}) | \theta^Q)$$
(3.8)

The actor, parameterized by θ^{π} , updates its parameters via gradient ascent by using the off-policy deterministic policy gradient Silver et al. (2014):

$$J_{\beta}(\pi|\theta^{\pi}) = \int_{\mathcal{X}} \rho^{\beta}(s) Q^{\pi}(x, \pi(x|\theta^{\pi})) dx$$

$$\nabla_{\theta^{\pi}} J_{\beta}(\pi|\theta^{\pi}) \approx \mathbb{E}_{x \sim \rho^{\beta}} \left[\nabla_{\theta^{\pi}} \pi(x, |\theta^{\pi}) \nabla_{a} Q^{\pi}(x, a)|_{a = \pi(x|\theta^{\pi})} \right]$$
(3.9)

where ρ^{β} is the discounted state distribution when acting under behavior policy β . By propagating the gradient through both policy and Q, the actor learns an approximation to the maximum of the value function under target policy π averaged over the state distribution of the behavior policy β .

A term that depends on $\nabla_{\theta^{\pi}} Q^{\pi}(x, a)$ has been dropped in 3.9, following a justification given by Degris et al. (2012) that argues that this is a good approximation since it can preserve the set of local optima to which gradient ascent converges.

3.2 The algorithm: Distributional off-policy deterministic actor critic

We present the CVaR optimization algorithm which is the main contribution of this thesis. The algorithm is based on the original off-policy deterministic actor critic explained in previous subsection 3.1.3, but introduces a distributional critic, which estimates the whole value distribution instead of only its expected value. With this extra information, the actor can learn to maximize other metrics than the expected value, specifically the CVaR. We proceed to present the components of the algorithm.

3.2.1 Distributional Critic

We use a distributional variant of the standard critic function, which maps from state-action pairs to distributions, inspired by the implicit quantile network (IQN) introduced in Dabney et al. (2018b).

IQN is a deterministic parametric function trained to reparameterize samples from a base distribution, e.g $\tau \in U([0,1])$, to the respective quantile values of a target distribution.

We define Z(x,a) as the random variable representing the return, with cumulative distribution function $F(z) := P(Z \le z)$ and we define $F_Z^{-1}(\tau) := Z(x,a;\tau)$ as its quantile function (or inverse cumulative distribution function) at $\tau \in [0,1]$. Thus, for $\tau \in U([0,1])$, the resulting state-action return distribution sample is $Z(x,a;\tau) \sim Z(x,a)$.

The critic IQN $Z(x, a; \tau | \theta^Z)$ parameterized by θ^Z is hence a parametric function used to represent the quantile function at specific quantile levels.

As in Dabney et al. (2018b),we train the critic IQN using the sampled quantile regression loss (Koenker, 2005) on the pairwise temporal-difference (TD)-errors. For two samples $\tau, \tau' \sim U([0,1])$, and current policy $\pi_{\theta^{\pi}}$, the sampled TD error is:

$$\delta^{\tau,\tau'} = r + \gamma Z(x_{t+1}, \pi(x_{t+1}|\theta^{\pi}); \tau'|\theta^{Z}) - Z(x_{t}, a_{t}; \tau|\theta^{Z})$$
(3.10)

Then, we compute the loss over the quantile samples:

$$\mathcal{L}_{QR}(x_t, a_t, r_t, x_{t+1}) = \frac{1}{N} \sum_{i=1}^{N} \sum_{j=1}^{N'} \rho_{\tau_i}^{\kappa}(\delta^{\tau_i, \tau_j'})$$
(3.11)

where N and N' are the number of iid samples $\tau_i, \tau'_j \sim U([0, 1])$ used to estimate the loss and where ρ_{τ}^{κ} is the quantile Huber loss.

Quantile Huber loss acts as an assymetric squared loss in an interval $[-\kappa, \kappa]$ around zero and reverts to a standard quantile loss outside this interval.

The Huber loss is given by Huber (1964):

$$\mathcal{L}_{\kappa}(u) = \begin{cases} \frac{1}{2}u^2 & \text{if } |u| \leq \kappa \\ \kappa(|u| - \frac{1}{2}\kappa) & \text{otherwise} \end{cases}$$
 (3.12)

Then the quantile Huber loss is the asymmetric variant of the Huber loss:

$$\rho_{\tau}^{\kappa}(u) = \left| \tau - [u < 0] \right| \mathcal{L}_{\kappa}(u), \forall u \in \mathbb{R}$$
(3.13)

where $[\cdot]$ is the Iverson bracket. Quantile loss penalizes overestimation errors (u<0) with weight $1-\tau$ and underestimation errors (u>0)with weight τ . By minimizing 3.11 via stochastic gradient descent with respect to θ^Z we aim to move towards the true quantile function.

3.2.2 Actor

The policy is updated via deterministic policy gradient ascent. We modify equation (3.9), to include the action-value distribution.

$$\nabla_{\theta^{\pi}} J_{\beta}(\pi | \theta^{\pi}) \approx \mathbb{E}_{x \sim \rho^{\beta}} \left[\nabla_{\theta^{\pi}} \pi(x, | \theta^{\pi}) \nabla_{a} Q^{\pi}(x, a) |_{a = \pi(x | \theta^{\pi})} \right]$$
(3.14)

$$= \mathbb{E}_{x \sim \rho^{\beta}} \left[\nabla_{\theta^{\pi}} \pi(x, |\theta^{\pi}) \mathbb{E} \left[\nabla_{a} Z^{\pi}(x, a | \theta^{Z}) \right] \right]_{a = \pi(x | \theta^{\pi})}$$
 (3.15)

Step from (3.14) to (3.15) comes by the fact that

$$Q^{\pi}(x, a) = \mathbb{E}[Z^{\pi}(x, a)] \tag{3.16}$$

With our goal of CVaR optimization in mind:

$$\arg\max_{\pi} \text{CVaR}_{\alpha}[Z(x, \pi(x))] \quad \forall x \in \mathcal{X}$$
 (3.17)

we can make use of the information provided by the Z distribution to optimize other objective functions rather than the expected value.

Second, can the contraction mapping results for a fixed grid of quantiles given by Dabney et al. (2018) be extended to the more general class of approximate quantile functions studied in this work?

To approach this, we use as a performance objective the distorted expectation of Z(x,a) under the distortion risk measure $\phi:[0,1]\to[0,1]$, with identity corresponding to risk-neutrality, i.e.:

$$Q_{\phi}(x,a) = \mathbb{E}_{\tau \sim U([0,1])}[Z_{\phi(\tau)}(x,a)]$$
(3.18)

which is actually equivalent to the expected value of $F_{Z(x,a)}^{-1}$ weighted by ϕ , i.e.:

$$Q_{\phi}(x,a) = \int_{0}^{1} F_{Z}^{-1}(\tau)d\phi(\tau)$$
 (3.19)

When we use as mapping $\phi(\tau) = \alpha \tau$, 3.19 corresponds to the CVaR of Z(x,a) as already presented in 2.3, and as a reminder:

$$CVaR_{\alpha}(Z) = \frac{1}{\alpha} \int_{0}^{\alpha} F_{Z}^{-1}(\tau) d\tau$$
 (3.20)

Again, 3.20 is the Acerbi's integral formula for CVaR, which states that CVaR at confidence level α can be interpreted as the integral of all the quantiles below the corresponding α . We can hence approximate (3.20) via sampling, by taking K samples of $\tau \sim U[0, \alpha]$:

$$CVaR_{\alpha}(Z) \approx \frac{1}{\alpha} \frac{1}{K} \sum_{i=1}^{K} Z(x, a; \tau_i) \qquad \tau_i \sim U[0, \alpha] \quad \forall i \in [1, K]$$
 (3.21)

Therefore, we arrive to the formula for the deterministic risk-sensitive policy gradient, for the parameters θ^{π} of the actor:

call it like that?

$$J_{\beta}^{CVAR}(\pi|\theta^{\pi}) = \int_{\mathcal{X}} \rho^{\beta}(s) \text{CVaR}_{\alpha}(Z^{\pi}(x, \pi(x|\theta^{\pi})) dx$$

$$\nabla_{\theta^{\pi}} J_{\beta}(\pi|\theta^{\pi}) \approx \mathbb{E}_{x \sim \rho^{\beta}} \left[\nabla_{\theta^{\pi}} \pi(x, |\theta^{\pi}) \nabla_{a} \left[\frac{1}{\alpha} \frac{1}{K} \sum_{i=1}^{K} Z(x, a; \tau_{i}) |\theta^{Z}) \right] |_{a = \pi(x|\theta^{\pi})} \right] \quad (3.22)$$
where $\tau_{i} \in U([0, \alpha]) \ \forall i \in [1, K]$

The algorithm can be summed up with:

- 1. Update distributional critic IQN via off-policy quantile-regression TD-learning: $\delta^{\tau,\tau'} = r + \gamma Z(x_{t+1}, \pi(x_{t+1}|\theta^{\pi}); \tau'|\theta^{Z}) Z(x_{t}, a_{t}; \tau|\theta^{Z})$
- 2. Update actor via deterministic risk-sensitive policy gradient ascent:

$$\nabla_{\theta^{\pi}} J_{\beta}(\pi | \theta^{\pi}) \approx \mathbb{E}_{x \sim \rho^{\beta}} \left[\nabla_{\theta^{\pi}} \pi(x, | \theta^{\pi}) \nabla_{a} \left[\frac{1}{\alpha} \frac{1}{K} \sum_{i=1}^{K} Z(x, a; \tau_{i}) | \theta^{Z} \right) \right] |_{a = \pi(x | \theta^{\pi})} \right]$$

Having the two main steps above, we again remark the capability of the algorithm to be implemented off-policy. We act in the environment using a more-exploratory behavior policy β and we learn a target policy π (where $\beta \neq \pi$). In contrast to stochastic off-policy actor-critic algorithms (Degris et al., 2012), we can avoid importance sampling both in the actor and the critic. This is due to the deterministic essence of the policies, which removes the integral over actions in the policy gradient and the expected value over actions in the Bellman equation for the critic.

3.3 Other distortion risk measures

In this section we present other interesting distortion risk measures ϕ that can be used to find policies optimizing other metrics rather than the CVaR. As discussed, evaluating under different distortion risk measures is equivalent to changing the distribution used to sample the quantile levels τ . The fact of being able to switch the objective function just by changing the sampling distribution for τ gives to the algorithm a lot of versatility. Algorithms that directly derive policies for CVaR optimization (Chow and Ghavamzadeh, 2014; Tamar et al., 2015a) don't show such property.

For the CVaR we have the distribution:

$$CVaR(\alpha, \tau) = \alpha \tau \quad \tau \sim U([0, 1])$$
(3.23)

A distortion risk measure proposed by Wang (2000) can easily switch between risk-averse (for $\eta < 0$) and risk-sensitive (for $\eta > 0$) distortions:

$$Wang(\eta, \tau) = \Phi(\Phi^{-1}(\tau) + \eta) \quad \tau \sim U([0, 1])$$
(3.24)

where Φ is the cumulative distribution of the standard Normal distribution.

While both Wang and CVaR heavily shift the distribution mass towards the tails of the distribution, CVaR entirely ignores all values corresponding to $\tau > \alpha$, whereas Wang gives to these non-zero, but vanishingly small probability (Dabney et al., 2018b).

Another, distortion risk measure would be a simple power formula for risk-averse (for $\eta < 0$) or risk-seeking(for $\eta > 0$) policies:

$$\operatorname{Pow}(\eta, \tau) = \begin{cases} \tau^{\frac{1}{1+|\eta|}} & \text{if } \eta \ge 0\\ 1 - (1-\tau)^{\frac{1}{1+|\eta|}} & \text{otherwise} \end{cases} \quad \tau \sim U([0, 1])$$
 (3.25)

3.4 Truncation

When changing the sampling distribution for τ to $U([0, \alpha])$ to compute the CVaR of the distribution, we make use of both the dual formulation 2.5 and Acerbi's integral formulas for CVaR.3.20.

A second approach would be to use the Rockafellar and Uryasev (2000) optimization problem for CVaR presented in 2.4 and as a reminder:

$$CVaR_{\alpha}(Z) = \max_{\nu} \left\{ \nu + \frac{1}{\alpha} \mathbb{E}_{Z}[[Z - \nu]^{-}] \right\}$$
(3.26)

which in the optimal point it holds that $\nu^* = \operatorname{VaR}_{\alpha}(Z)$.

Given the fact that our critic network estimates the $\operatorname{VaR}_{\alpha}(Z)$ of the return distribution, we can compute the CVaR using 3.26 by sampling uniformly from the whole quantile distribution, substracting the current estimated VaR for the confidence level α and truncating the result, that is to say, we can compute CVaR via sampling by:

$$\text{CVaR}_{\alpha}(Z) \approx \nu + \frac{1}{\alpha} \frac{1}{K} \sum_{i=1}^{K} \left[Z(x, a; \tau_i | \theta^Z) - \nu \right]^-$$
(3.27)

where $\tau_i \sim U([0,1])$ and $\nu = Z(x,a;\alpha|\theta^Z)$.

A disadvantage of such approach is sample-inefficiency, since due to truncation, only approximately αK samples will be used to compute the policy gradient.

This is a shared disadvantage with the algorithm presented in Chow and Ghavamzadeh (2014). For the sake of comparison we present it in the next section.

3.5 Chow

The way they formulated the problem considers costs and not rewards. Hence they want to minimize (and not maximize) the cost and reduce the right-tail $(1-\alpha)$ of the cost distribution

Chow and Ghavamzadeh (2014) present several policy gradient algorithms to solve the optimization problem:

$$\min_{\alpha} V^{\theta}(x^{0})$$
 subject to $\text{CVaR}_{\alpha}(D^{\theta}(x^{0})) \leq \beta$ (3.28)

for a given confidence level $\alpha \in (0,1)$ and loss tolerance $\beta \in \mathbb{R}$: where

$$V^{\theta}(x^0) = \mathbb{E}[D^{\theta}(x)] \tag{3.29}$$

$$D^{\theta}(x) = \sum_{k=0}^{\infty} \gamma^k C(x_k, a_k) | x_0 = x, a \sim \mu_{\theta}$$
 (3.30)

$$CVaR_{\alpha}(Z) = \min_{\nu \in \mathbb{R}} H_{\alpha}(Z, \nu) \equiv \min_{\nu \in \mathbb{R}} \{ \nu + \frac{1}{1 - \alpha} \mathbb{E}[(Z - \nu)^{+}] \}$$
 (3.31)

By Theorem 16 in Rockafellar and Uryasev (2002) and using the Lagrangian relaxation procedure (Bertsekas, 1976) 3.31 can be converted to the following unconstrained problem:

$$\max_{\lambda \ge 0} \min_{\theta, \nu} \mathcal{L}(\theta, \lambda, \nu) \equiv V^{\theta}(x^{0}) + \lambda (H_{\alpha}(D^{\theta}(x^{0}), \nu) - \beta)$$
(3.32)

where λ is the Lagrangian multiplier.

The goal of the authors is to find the saddle point of $\mathcal{L}(\theta, \lambda, \nu)$, i.e. $(\theta^*, \lambda^*, \nu^*)$ which is achieved by descending in (θ, ν) and ascending in α using the gradients of $\mathcal{L}(\theta, \lambda, \nu)$ with respect to the corresponding parameters.

For the sake of comparison with our algorithm, we will focus on the two actor-critic algorithms they propose, and ignore the policy gradient algorithm they propose which has high variance.

To address the problem, they approximate some quantities in the gradient estimates by linear combinations of basis functions adn update the parameters (linear coefficients) incrementally (after each state-action transition).

Their approach is to change the original MDP $\mathcal{M}=(\mathcal{X},\mathcal{A},C,P,P_0)$ to the augmented MDP $\hat{\mathcal{M}}=(\hat{\mathcal{X}},\hat{\mathcal{A}},\hat{C},\hat{P},\hat{P}_0)$ where $\hat{\mathcal{X}}=\mathcal{X}\times\mathbb{R},\;\hat{\mathcal{A}}=\mathcal{A},\;\hat{P}_0(x,s)=P_0(x)\mathbb{1}\{s_0=s\}$ and

$$\hat{\mathcal{C}}(x,s,a) = \begin{cases} \frac{\lambda(-s)^+}{1-\alpha} & \text{if } x = x_T\\ C(x,a) & \text{otherwise} \end{cases}$$
 (3.33)

$$\hat{P}(x', s'|x, s, a) = \begin{cases} P(x'|x, a) & \text{if } s' = \frac{s - C(x, a)}{\gamma} \\ 0 & \text{otherwise} \end{cases}$$
(3.34)

3.5. Chow

where S is the finite space of the augmented state s, s_0 is the initial state of the augmented MDP, x_T is the target state of the original MDP \mathcal{M} and s_T is the spart of the state when a policy μ reaches a target state x_T , i.e. $s_T = \frac{1}{\gamma^T}(\nu - \sum_{k=0}^{T-1} \gamma^k C(x_k, a_k))$, such as $s_0 = \nu$.

The augmented state s keeps track of the cumulative CVaR constraint cost and allows to reformulate the CVaR Lagrangian problem as an MDP. In the augmented MDP, the value function of policy μ_{θ} in state (x^0, ν) describes the whole CVaR Lagrangian cost:

$$V^{\theta}(x^{0}, \nu) = \mathbb{E}[D^{\theta}(x^{0})] + \frac{\lambda}{1 - \alpha} \mathbb{E}[(D^{\theta}(x - \nu)^{+})]$$
 (3.35)

and it holds that:

$$\nabla_{\theta} V^{\theta}(x^{0}, \nu) = \nabla_{\theta} \mathcal{L}(\theta, \lambda, \nu) \tag{3.36}$$

 $V^{\theta}(x^0, \nu)$ is called the critic and represented using linear approximation: $V^{\theta}(x^0, s) \approx v^T \phi(x, s)$.

At every time-step k an action is sampled from a parameterized policy $a_k \sim \mu(\cdot|x_k,s_k;\theta_k)$, we observe cost $\hat{C}(x_k,s_k,a_k)$ and next state $(x_{k+1},s_{k+1})=\hat{P}(\cdot|x_k,s_k,a_k)$ where $s_{k+1}=\frac{s_k-C(x_k,a_k)}{\gamma}$. The critic parameters v are then updated via TD-learning:

$$v_{k+1} = v_k + \varsigma_1 \delta_k \phi(x_k, s_k) \tag{3.37}$$

where
$$\delta_k = \hat{C}(x_k, s_k, a_k) + \gamma v_k^T \phi(x_{k+1}, s_{k+1}) - v_k^T \phi(x_k, s_k)$$
 (3.38)

Rest of parameters θ, λ, ν are updated by computing their gradients (or gradient estimates) with respect to $\mathcal{L}(\theta, \lambda, \nu)$.

$$\nabla_{\theta} \mathcal{L}(\theta_k, \lambda_k, \nu_k) \equiv \frac{1}{1 - \gamma} \nabla_{\theta} \log \mu(a_k | x_k, s_k | \theta) \delta_k$$
 (3.39)

With the remaining gradients we can already see some negative points: They use the unbiased estimate of $\nabla_{\lambda}\mathcal{L}(\theta,\lambda,\nu) = \nu_k - \beta + \frac{1}{(1-\gamma)(1-\alpha)}\mathbb{1}(x_k = x_T)(-s_T)^+$ which is fixed to $\nu_k - \beta$ all along a system trajectory and only changes at the end, which affects the incremental nature of the actor-critic algorithm. Additionally, it will only change λ in the case that $-s_T > 0$

 $\partial_{\nu}\mathcal{L}(\theta,\lambda,\nu)$ estimate can only be applied at the end of a system trajectory, when the terminal state x_T is reached, which prevents from having a fully-incremental algorithm.

$$\partial_{\nu} \mathcal{L}(\theta_k, \lambda_k, \nu_k) \ni \lambda_k - \lambda_k \frac{\mathbb{1}\{s_T \le 0\}}{1 - \alpha}$$
 (3.40)

This is the estimator they use in the semi trajectory-based actor-critic algorithm and it only updates the value in case $s_T \leq 0$. As an alternative, to estimate the subgradient $g(v) \in \partial_{\nu} \mathcal{L}(\theta, \lambda, \nu)$ incrementally they propose a simultaneous perturbation stochastic approximation (SPSA) approximation, which aims to estimate the subgradient $g(\nu)$ using two values of g at $\nu^- = \nu - \Delta$ and $\nu^+ = \nu + \Delta$ where Δ is a positive perturbation. In this case, and using the fact that the critic uses a linear approximation for the value function $V^{\theta}(x^0, s) \approx v^T \phi(x, s)$:

$$\partial_{\nu} \mathcal{L}(\theta_k, \lambda_k, \nu_k) \stackrel{\text{SPSA}}{\approx} \lambda_k + v^T [\phi(x^0, \nu^+) - \phi(x^0, \nu^-)] / 2\Delta$$
 (3.41)

When using more complex function approximators for the critic, computing 3.41 won't be that trivial, hence algorithm is also limited to simple MDPs that don't require complex V^{θ} .

Despite the final computational simplicity of the algorithm, we showed that it relies on lots of approximations and implementing it incrementally it is not straightforward.

It also uses an stochastic policy which as we already discussed is not natural in many appplications and it can increase the variance of the return. Furthermore, when working in an stochastic environment and trying to reduce the risk, using a policy which brings in additional stochasticity sounds a bit counterintuitive. Finally, is an on-policy algorithm which needs to act in the environment using same policy as the one being learnt. Hence, despite some modifications (for example introducing importance sampling), the algorithm cannot never be used in off-line settings.

3.6 Technical Details of the algorithm

The algorithm uses neural networks as non-linear function approximators for both the actor and the critic. To make effective use of large neural networks, we use insights from the DeepDPG algorithm Lillicrap et al. (2016) and in its turn from Deep Q Network (DQN) (Mnih et al., 2015). Before DQN, RL was believed to be unstable or diverge when nonlinear function approximators such as neural networks were used. The instability has several causes. First, correlation in the sequence of observations occurring when samples ar generated from exploring sequentially in an environment (hence i.i.d assumption is violated). Second, non-stationary data distribution since small updates in Q network may significantly change the policy and hereby, the data distribution. And third, correlation between the action-values and the target values during temporal difference backups since same network is being used. These 3 issues are solved by adding 2 innovations: 1) experience replay buffer: the network is trained by sampling batches of random past observations from a buffer, hereby removing correlations in the observation sequence and smoothing over changes in the data distribution. 2) use target Q networks that are only periodically updated, to update the Q network during temporal difference backups to reduce correlations with the target.

Observations (x_t, a_t, r_t, x_{t+1}) were sampled from the environment when acting under behavior policy β and stored in a fixed-sized (we used size of 10^6). When full, oldest samples were discarded. Since the algorithm is off-policy, the buffer can be large, allowing the algorithm to benefit from learning across a set of uncorrelated transitions. Exploration was addressed by constructing an exploratory behavior policy β by adding noise η to the actor policy.

$$\beta(x_t) = \pi(x_t | \theta^{\pi}) + \eta \tag{3.42}$$

where η is sampled from a noise process. In our case, as in Lillicrap et al. (2016),we used an Ornstein-Uhlenbeck process (Uhlenbeck and Ornstein, 1930) with $\theta=0.15$ and $\sigma=0.3$ with exponential decay. Ornstein-Uhlenbeck process models the velocity of a Brownian particle with friction, which results in temporally correlated values centered around the mean $(\mu=0.)$

explain more?

At every training step, a minibatch of observations is sampled from the buffer. To estimate the target value for critic training, we use two target networks: critic target network and actor target network $(\hat{Z}(x,a;\tau|\theta^{\hat{Z}}))$ and $\hat{\pi}(x|\theta^{\hat{\pi}})$ which are initialized as their homologues, but constrained to slowly track the learnt networks, so that to

rename?

stabilize learning. With this goal, weights of target networks are updated "softly", via: $\theta' \leftarrow (1 - \tau)\theta' + \tau\theta$ where $\tau \in [0, 1], \tau \ll 1$.

We additionally, use two insights from TD3 Fujimoto et al. (2018). First, we update the policy and target networks less frequently than the critic network. As Fujimoto et al. (2018) recommends we do one policy and target networks updates for every two critic network updates.

Second, we use target policy smoothing to address a particular failure mode that can happen in DeepDPG. To prevent policy to exploit actions for which the critic overestimated its value, target policy smoothing adds noise to target actions to smooth out Q over similar actions. Specifically target action used for the TD backup:

$$a_{t+1} = \hat{\pi}(x_{t+1}|\theta^{\hat{\pi}} + \text{clip}(\epsilon, -c, c) \quad \epsilon \sim \mathcal{N}(0, \sigma)$$
(3.43)

where we experimentally set c = 0.5 and $\sigma = 0.2$ and $\mathcal{N}(0, \sigma)$ is a Gaussian distribution with mean=0 and standard deviation σ .

To compute the quantile regression loss, we used N'=N=32 quantile levels to sample from target and critic networks. This parameter must be kept relatively big but further increasing it doesn't seem to improve performance. To compute the empirical CVaR from the estimated value distribution $Z(x,a;\tau)|\theta^Z\rangle$, we use K=8 quantile levels.

Training starts after a warm-up of T=25000 environment interactions. During this time, actor networks where not used but random policies where used instead, to enhance exploration. Adam optimizer wsa used for learning the neural networks parameters with a learning rate of 10^-4 and 10^-3 for the actor and the critic respectively. For additional information on the networks used, please see Appendix A

Algorithm 1: How to write algorithms

```
Result: Write here the result
initialization;
while While condition do

instructions;
if condition then

instructions1;
instructions2;
else
instructions3;
end
end
```

From a practical viewpoint, using stochastic policies requires integrating over both state and action spaces to compute the policy gradient, whereas the deterministic case only needs to integrate over the state space. Hence, stochastic policy gradients may require much more samples, especially if the action space has many dimensions.

use replay buffer, target networks as DQN to deal with problems of Q-learning with functions approximators.

Since distorted expectations can be expressed as weighted average over the quantiles Dhaene et al. (2012), we can use a specific sampling base distribution $\beta: [0,1] \to [0,1]$ to sample the quantile levels $\tau \in [0,1]$) from our critic network $Z(x,a;\tau)$.

Chapter 4

Batch RL

- Watch videos Levine
- Papers bear and bcq

We decide to test the capabilities of our algorithm in a *fully* off-policy setting, also called *batch RL setting* or *offline RL*. In this setting, the agent can only learn from a fixed dataset without further interaction with the environment.

Add motivation in doing so: envs where collection of data is expensive, unsafe for robotics or autonomous vehicles

The 'off-policy' algorithm we presented in previous sections falls in the category of off-policy "growing batch learning" in which data is collected by using near-on-policy policies such as ϵ -greedy and stored in a replay buffer. After used for training, the data is replaced with *fresher* data obtained from interaction of the agent with the environment using an updated policy. As a result, the dataset used tends to be heavily correlated to the current policy.

Issues with Batch RL

Most of off-policy algorithms fail to learn in the off-line setting. This is due to a fundamental problem of off-policy RL, called extrapolation error (Fujimoto et al., 2019) or bootstrapping error (Kumar et al., 2019). This error is introduced due to a mismatch between the dataset distribution and the state-action visitation distribution induced by the current target policy. At every train step the Q estimate is updated in the direction to reduce the Bellman error, i.e. the mean squared error between the current value estimate and the expected Q value under the current target policy at the next state. The Q function estimator, however, is valid only when evaluated on actions sampled from the behavior policy, which in the batch-RL case is the distribution of the dataset. Using unfamiliar (unlikely or not contained in the dataset) action (also called out of distribution (OOD) actions in (Kumar et al., 2019)) for the next-state, results on a new Q value estimate which is affected by this extrapolation error, resulting in pathological values that incur large absolute error from the optimal desired Q-value.

It is good to notice, that for an on-policy settings, extrapolation error is generally something positive, since it leads to a beneficial exploration. In this case, if the value function is overestimating the value at a (state-action) pair, the current policy will

19 4.1. Details of VAE

lead the agent to that pair, collect the data at that point and hence, the value estimate will be corrected afterwards. In the off-policy setting, the correction step is not possible due to the inability of collecting new data.

Our approach To overcome this issue, we inspire ourselves on the approach presented in Fujimoto et al. (2019), where a generative model G_w is trained to generate actions with high similarity to the dataset. For the generative model we use a conditional variational auto-encoder (VAE) Kingma and Welling (2014) which generates action samples as a reasonable approximation to $\underset{a}{\operatorname{argmax}} P_{\mathcal{B}}^G(a|s)$, where $P_{\mathcal{B}}^G(a|s)$ is the conditioned marginal likelihood.

4.1 Details of VAE

A variational autoencoder aims to maximize the marginal log-likelihood $\log p(X) = \sum_{i=1}^{N} \log p(x_i)$, where X is the dataset with iid samples $\{x_1, x_2, ... x_N\}$. It is assumed that data is generated by some random process, involving an unobserved continuous random variable **z**. The process consists of two steps: (1) a value z_i is generated from some prior distribution p(z) and (2) a value x_i is generated from some conditional distribution p(x|z). Given that the true probabilities are unknown, a recognition model $q(z|x;\phi)$ is introduced as an approximation to the intractable true posterior $p(z|x;\theta)$.

The recognition model $q(z|x;\phi)$ is called an *encoder*, since given a datapoint x it produces a *random latent vector* z. $p(x|z;\theta)$ is called a *decoder*, since given the random latent vector z it reconstructs the original sample x.

Since computing the desired marginal $p(X;\theta)$ is intractable, VAE algorithm optimizes a lower bound instead:

$$\log p(X;\theta) \ge \mathcal{L}(\theta,\phi;X) = \mathbb{E}_{q(z|X;\phi)}[\log p(X|z;\theta)] - D_{KL}(q(z|X;\phi) \mid\mid p(z;\theta)) \tag{4.1}$$

For our implementation, the prior $p(z;\theta)$ is chosen to be a multivariate normal distribution $\mathcal{N}(0,Id)$, hence it lacks parameters.

For the probabilistic encoders and decoders we use neural networks. For the encoder $q(z|x_i;\phi)$ we used a neural network with Gaussian output, specifically a multivariate Gaussian with a diagonal covariance structure $\mathcal{N}(z|\mu(X),\sigma^2(X)Id)$, where μ and σ are the outputs of the neural network, i.e. nonlinear functions of datapoint $x_i := (state_i, action_i)$ and ϕ . To sample from the posterior $z_i \sim q(z|x_i;\phi)$ we use the reparameterization trick: $z_i = g(x_i, \epsilon; \phi) = \mu_i + \sigma_i \odot \epsilon$ where $\epsilon \sim \mathcal{N}(0, Id)$ and \odot is the element-wise product. For the decoder $p(x|z;\theta)$ we used another neural network with deterministic output, i.e. nonlinear function of datapoint $\hat{x}_i := (state_i, z_i)$ and θ .

The VAE is trained to maximize reconstruction loss and a KL-divergence term according to the distribution of the latent vector:

When it comes to training the VAE, both recognition model parameters ϕ and the generative model parameters θ are learnt jointly to maximize the variational lower bound $\mathcal{L}(\theta, \phi; X)$ via gradient ascent which includes the expected reconstruction

rewrite

error loss and the KL-divergence term according to the distribution of the latent vectors. When both prior and posterior are Gaussian, KL-divergence can be computed analytically:

$$-D_{KL}(q(z|X;\phi) \mid\mid p(z;\theta)) = \frac{1}{2} \sum_{j=1}^{J} (1 + \log((\sigma_j)^2) - (\mu_j)^2 - (\sigma_j)^2)$$
(4.2)

where J is the dimensionality of z, and μ_j , σ_j represent the jth element of these vectors. The expected reconstruction error $\mathbb{E}_{q(z|X;\phi)}[\log p(X|z;\theta)]$ requires estimation by sampling, and we will use the mean-squared error between the $action_i$ from the dataset and the reconstructed action.

Finally, when acting during evaluation or deployment, random values of z will be sampled from the multivariate normal and passed through the decoder to produce actions.

add:

- New actor network
- VAE(state) + perturbation_level × DeterministicActorNN(vae_action, state)
- Train DeterministicActorNN as in the algorithm presented previously for off-policy RL.
- Remove randn sampling of z in deployment? It works :)

Chapter 5

Results

5.1 Current results Car

Problem: A car with fully-observable 2D-state: [position, velocity] needs to move from initial position $x_0 = 0$ m and initial velocity $v_0 = 0$ m ts⁻¹ to goal position $x_F = 2.0$ m. The action taken at every time-step ts, with a discretization of $t_d = 0.1$, determines the car acceleration. The control input a is constrained to range between [-1.0,1.0]m ts⁻². Per every time-step passed before it reaches the goal, the car receives a penalization reward $R_{\rm ts} = -10$ If the car reaches the goal position, it receives a reward $R_F = +270$ and the episode ends. Otherwise, after $T_F = 400$ ts the episode ends (with no extra penalization).

5.1.1 Case 1: No velocity penalization

Using both DDPG and CVAR-DDPG algorithms, the car arrives at the goal position. Both with a maximum acceleration kept throughout the whole episode.

For this setup we have:

$$x = x_0 + v_0 \frac{\text{ts}}{10} + 0.5a \left(\frac{\text{ts}}{10}\right)^2$$

In the optimal case, the car keeps an acceleration of $1 \,\mathrm{m\,ts^{-2}}$ for the whole episode, and hence reaches $x_F = 2 \mathrm{m}$ with 20 time-steps. Hence the final cumulative reward $G_T = (20+1)R_{\mathrm{ts}} + R_F = 60$.

Starting from $x_0 = 0$ m, the car reaches a velocity of 1m ts^{-1} after 10 time-steps, at $x_{\tau=10} = 0.5$. Keeping velocity 1m ts^{-1} through the rest of the episode, it reaches the goal position after 14 time-steps. Hence the final cumulative reward $G_T = (10 + 14 + 1)R_{\tau} + R_F = 74$. The reward values were chosen in order to make sure that, for this Case 2 setting, driving with a velocity higher than 1m ts^{-1} never induces higher cumulative rewards.

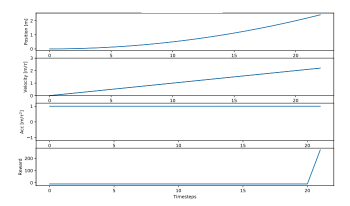


Figure 5.1: Car trajectory using DDPG and algorithm without velocity penalization. (Same behavior for CVAR-DDPG algorithm).

5.1.2 Case 2: Velocity penalization with probability 1

The experiment is carried out to ensure the two algorithms manage to learn the new reward function when there is no uncertainty. In this setup, when the car velocity exceeds $1 \mathrm{m} \, \mathrm{ts}^{-1}$, it receives a penalization of $R_v = -20$. We expect both algorithms to perform similarly since there is no reward uncertainty. As expected, both DDPG and CVAR-DDPG algorithms learn to accelerate with maximum value till a velocity of $1 \mathrm{m} \, \mathrm{ts}^{-1}$ is reached, and then they keep the velocity constant until the goal is reached.

Starting from $x_0 = 0$ m, the car reaches a velocity of 1m ts^{-1} after 10 time-steps, at $x_{\text{ts}=10} = 0.5$. Keeping velocity 1m ts^{-1} through the rest of the episode, it reaches the goal position after 14 time-steps. Hence the final cumulative reward $G_T = (10 + 14 + 1)R_{\text{ts}} + R_F = 20$. The reward values were chosen in order to make sure that, for this Case 2 setting, driving with a velocity higher than 1m ts^{-1} never induces higher cumulative rewards.

For this Case 2 setting, the trained models were saved using Early stopping with maximal reward in episode evaluation as a metric and with a patience of 100 episodes. The quantiles used for learning the actor for the CVAR-DDPG algorithm were sampled uniformly $\sim U[0,1]$

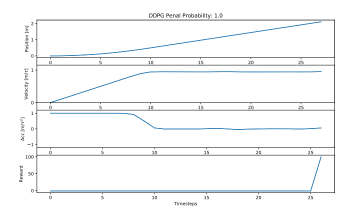


Figure 5.2: Car trajectory using DDPG algorithm and velocity penalization with probability 1

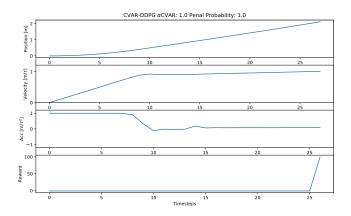


Figure 5.3: Car trajectory using CVAR-DDPG algorithm and velocity penalization with probability 1. (α -CVAR = 1)

5.1.3 Case 3: Velocity penalization with probability P

The experiment is carried out to show the risk-sensitiveness property of the CVAR-DDPG algorithm.

The models saved were the ones that obtained a maximum CVAR (with a window of 10 episodes) of the cumulative rewards during evaluation The quantiles used for learning the actor for the CVAR-DDPG algorithm were sampled uniformly $\backsim U[0,\alpha]$ where $\alpha=0.1$

The models saved were the ones that obtained a maximum CVAR (with a window of 10 episodes) of the cumulative rewards during evaluation

The quantiles used for learning the actor for the CVAR-DDPG algorithm were sampled uniformly $\backsim U[0,\alpha]$ where $\alpha=0.2$.

For P = 0.2 the CVAR-DDPG algorithm learns to saturate the velocity, even though

the probability of a penalization is low, whereas the DDPG algorithm doesn't, and keeps a linear increase of the velocity during the whole episode. The CVAR algorithm reaches its maximum CVAR of 64.0 at episode 220, whereas the DDPG reaches its maximum CVAR value of 49.0 at episode 1435.

Important issue: Although CVAR-DDPG finds a risk-sensitive trajectory at episode 220, it doesn't converge there and keeps oscillating and even moves towards a risk-neutral behavior later on. The graph in figure 5.7, shows the evolution of the sampled mean of the tail of the sampled cumulative value distribution (CDF). (i.e. we compute via IQN the quantile values from the tail value distribution (VD) and take the mean). The value it converges to coincides with the maximum value of the CVAR we achieved ,but then the actor doesn't seem to behave accordingly.

$$CVaR_{\alpha}(Z) = \frac{1}{\alpha} \int_{0}^{\alpha} F_{Z}^{-1}(\tau) d\tau = \frac{1}{\alpha} \int_{0}^{\alpha} IQN(\tau) d\tau \approx \frac{1}{\alpha} \frac{1}{K} \sum_{i=0}^{K} IQN(\tau_{i})$$
 (5.1)

where $\tau_i \sim U[0, \alpha]$, and IQN is the output of the IQN network for given τ , representing the value of the return for the given quantile.

(Values of the sampled CVAR showed in 5.7 are not divided by α neither K)

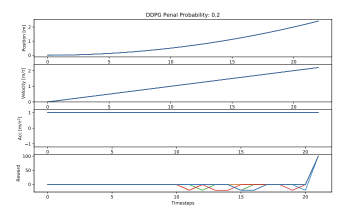


Figure 5.4: Car trajectory using DDPG algorithm and velocity penalization with probability P=0.2

A binomial distribution can be observed.

5.2 Current results Batch RL HalfCheetah

We use one of the D4RL datasets. Specifically halfcheetah-medium-v0, which uses 1M samples from a policy trained to approximately 1/3 the performance of the expert.

We introduce stochasticity in the original cost function in a way that makes the environment stochastic enough to have a meaningful assessment of risk in terms of tail performance. A reward of -100 is given wp 0.05, if the velocity of the cheetah

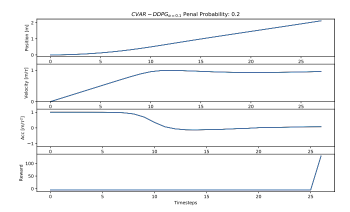


Figure 5.5: Car trajectory using CVAR-DDPG algorithm and velocity penalization with probability 0.2 and (α -CVAR = 0.2)

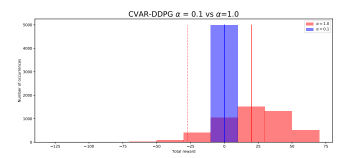


Figure 5.6: Comparison of cumulative rewards achieved with CVAR-DDPG algorithms with $\alpha = 0.2$ and $\alpha = 1$ when the probability of velocity penalization = 0.2. Algorithm with $\alpha = 1$ achieves a higher expected value ($\mu = 20.11$) but has a lower CVAR (CVaR $_{\alpha=0.1}$ =-27.12 compared to the algorithm with $\alpha=0.1$, which has $\mu=0$ and CVaR $_{\alpha=0.1}$ =0.0 5000 episodes were ran after training each algorithm.

is greater than 4. We train using the distributional critic and a policy that consists of a variational autoencoder to sample from the dataset distribution and then a second perturbing network that shifts the action towards maximizing the sampled CVaR. The perturbation is up to 0.5 (paper originally 0.05).

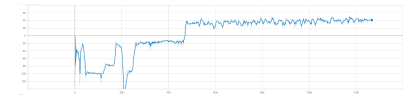


Figure 5.7: Evolution of the sampled mean of the tail of the Cumulative Value Distribution during training epochs

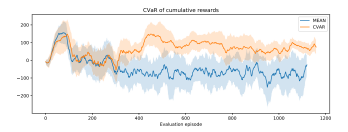


Figure 5.8: Evolution during training of CVaR ($\alpha=0.1$) of the cumulative rewards over 5 evaluation episodes

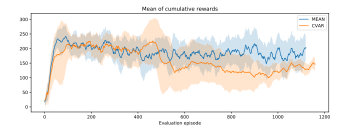


Figure 5.9: Evolution during training of mean of the cumulative rewards over 5 evaluation episodes

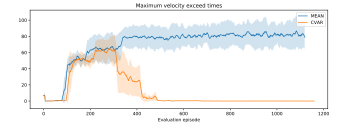


Figure 5.10: Evolution during training of mean of times of maximum velocity exceed over 5 evaluation episodes

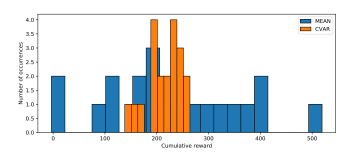


Figure 5.11: Histogram of cumulative rewards during 200 time steps using the trained final policies

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Appendix A

A.1 Experiment details

Extra info about the network architectures?

A.2 Distributional RL

demonstrated the importance of learning the value distribution, i.e., the distribution of the random return received by a RL agent. This differs from the common RL approach which is focused on learning the expected value of this return.

cite papers

Goal in RL is to teach an agent to act so that it maximizes its expected utility, Q Sutton and Barto (1998) Bellman's equation describes this value Q in terms of the expected reward and expected outcome of the random transition $(x, a) \to (X', A')$, showing the particular recursive relationship between the value of a state and the values of its successor states:

$$Q(x,a) = \mathbb{E}[R(x,a)] + \gamma \mathbb{E}[Q(X',A')] \tag{A.1}$$

Distributional RL aims to go beyond the notion of *value* and training to study instead the random return Z.

A.2.1 Example showing interest in learning the distribution

Imagine the example in which we are playing a board game and we roll 2 dices. If we get a 3, we fall in prison and need to pay 2000CHF (i.e. reward of -2000CHF), whereas otherwise we we collect a salary of 200CHF (i.e. reward of +200CHF). If we consider the common reinforcement learning approach and we compute the expected immediate ($\gamma = 1$) reward:

remove and maybe the motivation is done for the doors example in chapter 2

$$\mathbb{E}[R(x)] = \frac{1}{36}(-2000 \,\text{CHF}) + \frac{35}{36}(200 \,\text{CHF}) = 138.88 \,\text{CHF} \tag{A.2}$$

Hence, the expected immediate return is +138.88CHF. However, in any case we will get a return of +138.88CHF. Instead:

Appendix A. 32

$$R(x) = \begin{cases} -2000 \text{ CHF}, & \text{w.p.} \frac{1}{36} \\ 200 \text{ CHF}, & \text{w.p.} \frac{35}{36} \end{cases}$$

We define the random return $Z^{\pi}(x, a)$ as the random variable that represents the sum of discounted rewards obtained by starting from position x taking action a and thereupon following policy π .

add new section

This variable captures intrinsic randomness from:

- 1. Immediate rewards
- 2. Stochastic dynamics
- 3. Possibly an stochastic policy

Having defined $Z^{\pi}(x,a)$, we can clearly see that:

$$Q^{\pi}(x,a) = \mathbb{E}[Z^{\pi}(x,a)] \tag{A.3}$$

Z is also described by a recursive equation, but of a distributional nature:

$$Z^{\pi}(x,a) \stackrel{D}{=} R(x,a) + \gamma Z(x',a') \tag{A.4}$$

where $x' \backsim p(\cdot|x, a)$ and $a' \backsim \pi(\cdot|x')$

where $\stackrel{D}{=}$ denotes that the RV on both sides of the equation share the same probability distribution. The distributional Bellman equation defined in (A.4), states that the distribution of Z is characterized by the interaction of 3 RV's: the random variable reward R, the next state-action (X',A') and its random return Z(X',A'). From here on, we will view Z^{π} as a mapping from state-action pairs to distributions over returns, and we call this distribution the value distribution.

A.2.2 Distributional Bellman Operator

In the policy evaluation setting Sutton and Barto (1998), one aims to find the value function $V\pi$ associated with a given fixed policy π . In the distributional case, we aim to find $Z\pi$. Bellemare et al. (2017) defined the Distributional Bellman operator T^{π} . We view the reward function as a random vector $R \in \mathbb{Z}$ and define the transition operator $P^{\pi}: \mathbb{Z} \to \mathbb{Z}$

$$P^{\pi}Z(x,a) \stackrel{D}{=} Z(X',A') \tag{A.5}$$

$$X' \backsim P(\cdot|x, a) \text{ and } A' \backsim \pi(\cdot|X')$$
 (A.6)

where we use capital letters to emphasize the random nature of the next state-action pair (X',A') Then, the Distributional Bellman operator T^{π} is defined as:

$$T^{\pi}Z(x,a) \stackrel{D}{=} R(x,a) + \gamma P^{\pi}Z(x,a)$$
 (A.7)

Bellemare et al. (2017) showed that (A.7) is a contraction mapping in Wasserstein metric whose unique fixed point is the random return Z^{π} .

Appendix A. 34

Wasserstein metric:

The p-Wasserstein metric W_p , for $p \in [1, \infty]$, also known as the Earth Mover's Distance when p = 1 is an integral probability metric between distributions. The p-Wasserstein distance is characterized as the L^p metric on inverse cumulative distribution functions (CDF). Tht is, the p-Wasserstein metric between distributions U and Y is given by:

$$W_p(U,Y) = \left(\int_0^1 |F_Y^{-1}(w) - F_U^{-1}(w)|^p dw \right)^{\frac{1}{p}}$$
 (A.8)

where for a random variable Y, the inverse CDF F_V^{-1} of Y is defined by:

$$F_Y^{-1}(w) := \inf\{y \in \mathbb{R} \mid w \le F_Y(w)\}$$
(A.9)

where $F_Y(w) = Pr(y \leq Y)$.

Add Figure 2.1 in Dabney et al. (2018a) Unlike the Kullback-Leibler divergence, the Wasserstein metric is a true probability metric and considers both the probability of and the distance between various outcome events, which makes it well-suited to domains where an underlying similarity in outcome is more important than exactly matching likelihoods.

Contraction in \hat{d}_p :

Let \mathcal{Z} be the space of action-value distributions:

$$\mathcal{Z} = \{ Z \mid \mathcal{X} \times \mathcal{A} \to \wp(\mathbb{R})$$
 (A.10)

$$\mathbb{E}[|Z(x,a)|^p < \infty, \forall (x,a), p > 1]\}$$
(A.11)

check first line the \wp

Then, for two action-value distribution $Z_1, Z_2 \in \mathcal{Z}$, the maximal form of the Wasserstein metric is defined by:

$$\hat{d}_p(Z_1, Z_2) := \sup_{x, a} W_p(Z_1(x, a), Z_2(x, a))$$
(A.12)

Bellemare et al. (2017) showed that \hat{d}_p is a metric over value distributions and furthermore, the distributional Bellman operator T^{π} is a contraction in \hat{d}_p . Consider the process $Z_{k+1} := T^{\pi} Z_k$, starting with some $Z_0 \in \mathcal{Z}$.

 $T^{\pi}Z: \mathcal{Z} \to \mathcal{Z}$ is a γ -contraction in the Wasserstein metric \hat{d}_p , which implies that not only the first moment (expectation) converges exponentially to Q^{π} , but also in all moments.

Lemma 1: (Lemma 3 in Bellemare et al. (2017)) T^{π} is a γ -contraction: for any two $Z_1, Z_2 \in \mathcal{Z}$,

$$\hat{d}_p(T^{\pi}Z_1, T^{\pi}Z_2) \le \gamma \hat{d}_p(Z_1, Z_2) \tag{A.13}$$

Using Banach's fixed point theorem, it is proven that T^{π} has a unique fixed point, which by inspection must be Z^{π} .

Hence the \hat{d}_p metric is shown to be useful metric for studying behavior of distributional RL algorithms, and to showed their convergence to a fixed point. Moreover,

shows than en effective way to learn a value distribution is to attempt minimize the Wasserstein distance between a distribution Z and its distributional Bellman update $T^{\pi}Z$, analogously to the way that TD-learning attempts to iteratively minimize the L^2 distance between Q and TQ.

We have so fare considered a policy evaluation setting, i.e. trying to learn a value distribution for a fixed policy π , and we studied the behavior of its associated distributional operator T^{π} . In the control setting, i.e., when we try to find a policy π^* that maximizes a value, or its distributional analogous, i.e. that induces an optimal value distribution. However, while all optimal policies attain the same value Q^* , in general there are many optimal value distributions.

The distributional analogue of the Bellman optimality operator converges, in a weak sense, to the set of optimal value distributions, but this operator is *not a contraction* in any metric between distributions.

Let Π^* be the set of optimal policies.

Definition 1: An optimal value distribution is the value distribution of an optimal policy. The set of optimal value distributions is

$$\mathcal{Z}^* \coloneqq \left\{ Z^{\pi^*} \middle| \pi^* \in \Pi^* \right\}$$

Not all value distributions with expectation Q^* are optimal, but they must match the full distribution of the return under some optimal policy. **Definition 2:** A greedy policy π for Z $in\mathcal{Z}$ maximizes the expectation of Z. The set of greedy policies for Z is:

$$\mathcal{G}_{\mathcal{Z}} \coloneqq \left\{\pi \mid \sum_{a} \pi(a|x) \mathbb{E}(Z(x, a) = \max_{a' \in \mathcal{A}} Q(x', a')\right\}$$

We will call a distributional Bellman optimality operator any operator \mathcal{T} which implements a greedy selection rule, i.e.:

$$\mathcal{T}Z = \{\mathcal{T}^{\pi}Z \text{ for some } \pi \in \mathcal{G}_{\mathcal{Z}}\}$$

As in the policy evaluation setting, we are interested in the behavior of the iterates $Z_{k+1} := \mathcal{T}Z_k, Z_0 \in \mathcal{Z}$. Lemma 4 in Bellemare et al. (2017) shows that $\mathbb{E}Z_k$ behaves as expected: **Lemma 4:** Let $Z_1, Z_2 \in \mathcal{Z}$. Then:

$$\|\mathbb{E}\mathcal{T}Z_1 - \mathbb{E}\mathcal{T}Z_2\|_{\infty} \le \gamma \|\mathbb{E}Z_1 - \mathbb{E}Z_2\|_{\infty}$$

and in particular $\mathbb{E}Z_1 \to Q^*$ exponentially quickly. However, Z_k is not assured to converge to a fixed point. Specifically, they provide a number of negative results concerning \mathcal{T} :

Proposition 1: The operator \mathcal{T} is not a contraction.

Proposition 2: Not all optimality operators have a fixed point $Z^* = \mathcal{T}Z^*$ **Proposition 3:** That \mathcal{T} has a fixed point $Z^* = \mathcal{T}Z^*$ is insufficient to guarantee the convergence of $\{Z_k\}$ to Z^*

Another result, shows that we cannot in general minimize the Wasserstein metric, viewed as a loss, using stochastic gradient descent methods. This limitation, is crucial in a practical context, when the value distribution needs to be approximated.

A.2.3 Quantile approximation

Dabney et al. (2018a) used the theory of quantile regression Koenker (2005), to design an algorithm applicable in a stochastic approximation setting. Quantile

Appendix A. 36

regression is used to estimate the quantile function at precisely chosen points. Then the Bellman update is applied onto this parameterized quantile distribution. This combined operator is proven to be a contraction and the estimated quantile function is shown to converge to the true value distribution when minimized using stochastic approximation.

A.2.4 Quantile projection:

Our current aim is to estimate quantiles of the target distribution, i.e. the values of the return that divide the value distribution in equally sized parts. We will call it a quantile distribution, and we will let $\mathcal{Z}_{\mathcal{Q}}$ be the space of quantile distributions. We denote the cumulative probabilities associated with such a distribution by $\tau_1, \tau_2...\tau_N$, so that $\tau_i = \frac{i}{N}$ for i = 1, ...N.

by $\tau_1, \tau_2...\tau_N$, so that $\tau_i = \frac{i}{N}$ for i = 1, ...N. Formally, let $\theta : \mathcal{X} \times \mathcal{A} \to \mathbb{R}^N$ be some parametric model. A quantile distribution $Z_\theta \in \mathcal{Z}_Q$ maps each state-action pair (x,a) to a uniform probability distribution supported on $\{\theta_i(x,a)\}$. Hence we can approximate it by a uniform mixture of N Diracs:

$$Z_{\theta}(x,a) := \frac{1}{N} \sum_{i=1}^{N} \delta_{\theta_i(x,a)}$$
(A.14)

with each θ_i assigned a fixed quantile. We aim to learn the support of these Diracs, i.e. learn $\theta_i \forall i, a, x$. We will do it by quantifying the projection of an arbitrary value distribution $Z \in \mathcal{Z}$ onto $\mathcal{Z}_{\mathcal{Q}}$, that is:

$$\prod_{W_1} Z := \underset{Z_{\theta} \in \mathcal{Z}_{\mathcal{O}}}{\operatorname{arg\,min}} W_1(Z, Z_{\theta}) \tag{A.15}$$

This projection \prod_{W_1} is the quantile projection.

We can quantify the projection between a distribution with bounded first moment Y and U, a uniform distribution over N Diracs as in (A.14) with support $\{\theta_1, ... \theta_N\}$ by:

$$W_1(Y, U) = \sum_{i=1}^{N} \int_{\tau_{i-1}}^{\tau_i} |F_Y^{-1}(w) - \theta_i| dw$$
 (A.16)

Lemma 2 in Dabney et al. (2018a) establishes that the values $\{\theta_1, ..., \theta_N\}$ for the returns that minimize $W_1(Y, U)$ are given by $\theta_i = F_Y^{-1}(\hat{\tau}_i)$, where $\hat{\tau}_i = \frac{\tau_{i-1} + \tau_i}{2}$.

A.2.5 Quantile Regression

Quantile regression is a method for approximating quantile functions of a distribution at specific points, i.e. its inverse cumulative distribution function. The quantile regression loss, for quantile $\tau \in [0,1]$, is an asymmetric convex lox function that penalizes underestimation errors with weight τ and overestimation errors with weight $1-\tau$.

For a distribution Z, and given quantile τ , the value of the quantile function $F_Z^{-1}(\tau)$

may be characterized as the minimizer of the quantile regression loss:

$$\mathcal{L}_{QR}^{\tau}(\theta) = \mathbb{E}_{\hat{Z} \sim Z}[\rho_{\tau}(\hat{Z} - \theta)]$$

$$\rho_{\tau}(u) = u(\tau - \delta_{u < 0}), \forall u \in \mathbb{R}$$
(A.17)

Given that the minimizer of the quantile regression loss for τ is $F_Z^{-1}(\tau)$, and using Lemma 2 in Dabney et al. (2018a), which claims that the values of $\{\theta_1,...\theta_N\}$ that minimize $W_1(Z,Z_\theta)$ are given by $\theta_i=F_Y^{-1}(\hat{\tau}_i)$; we can claim that the values of $\{\theta_1,...\theta_N\}$ are the minimizers of the following objective:

$$\sum_{i=1}^{N} \mathbb{E}_{\hat{Z} \sim Z}[\rho_{\hat{\tau}_i}(\hat{Z} - \theta_i)] \tag{A.18}$$

This loss gives unbiased sample gradients and hence, we can find the minimizing $\{\theta_1,...\theta_N\}$ by stochastic gradient descent.

add huberloss

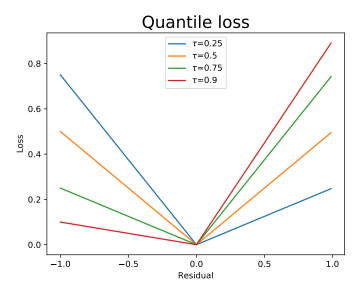


Figure A.1: Quantile loss for different quantile values

Proposition 2 in Dabney et al. (2018a) states that the combined quantile projection \prod_{W_1} with the Bellman update \mathcal{T}^{π} has a unique fixed point \hat{Z}^{π} , and the repeated application of this operator, or its stochastic approximation, converges to \hat{Z}^{π} .

A.2.6 Quantile Regression Temporal Difference Learning

Temporal difference learning updates the estimated value function with a single unbiased sample following policy π . Quantile regression allows to improve the estimate of the quantile function for some target distribution Y(x), by observing samples $y \sim Y(x)$ and minimizing equation (A.17). Using the quantile regression loss, we can obtain an approximation with minimal 1-Wasserstein distance from the original. We can combine this with the distributional Bellman operator to give a

Appendix A. 38

target distribution for quantile regression, creating the quantile regression temporal difference learning algorithm: $\frac{1}{2}$

$$u = r + \gamma z' - \theta_i(x) \tag{A.19}$$

$$\theta_i(x) \leftarrow \theta_i(x) + \alpha(\hat{\tau}_i - \delta_{u<0})$$
 (A.20)

$$a \sim \pi(\cdot|x), r \sim R(x, a), x' \sim P(\cdot|x, a), z' \sim Z_{\theta}(x')$$
(A.21)

where Z_{θ} is a quantile distribution as in (A.14) and $\theta_i(x)$ is the estimated value of $F_{Z^{\pi}(x)}^{-1}(\hat{\tau}_i)$ in state x.