

Sample-based Distributional Policy Gradient

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Abstract

Distributional reinforcement learning (DRL) is a recent reinforcement learning framework whose success has been supported by various empirical studies. It relies on the idea of replacing the expected return with the return distribution, which captures the intrinsic randomness of the long term rewards. Most of the existing literature on DRL focuses on problems with discrete action space and value based methods. In this work, motivated by applications in control engineering and robotics where the action space is continuous, we propose the sample-based distributional policy gradient (SDPG) algorithm. It models the return distribution using samples via a reparameterization technique widely used in generative modeling. We compare SDPG with the state-of-the-art policy gradient method in DRL, distributed distributional deterministic policy gradients (D4PG). We apply SDPG and D4PG to multiple OpenAI Gym environments and observe that our algorithm shows better sample efficiency as well as higher reward for most tasks¹.

Keywords: Reinforcement learning, optimal control, policy gradients

1. Introduction

As a data-driven counterpart of model-based control, reinforcement learning (RL) has shown potential in solving a variety of complex problems (Mnih et al., 2015; Levine et al., 2016). RL algorithms can be roughly divided into two categories: value function based and policy gradient methods. Value function based algorithms do not explicitly parameterize the policy, but rather obtain the policy from a learned value function. In contrast, policy gradient methods improve a parameterized policy based on the policy gradient theorem (Sutton et al., 2000) and have shown to be more effective in continuous action space control settings. In particular, deep deterministic policy gradient (DDPG) (Lillicrap et al., 2016) utilizes neural networks to parameterize the policy and has been successful in solving continuous control tasks.

Instead of modeling the value function as the expected sum of the discounted rewards, the recently proposed distributional reinforcement learning (DRL) (Bellemare et al., 2017) framework suggests to work with the full distribution of random returns, known as the value or return distribution.

1. <https://github.com/rahulsinghchandraul/SDPG>

Several typical DRL algorithms such as C51 (Bellemare et al., 2017), D4PG (Barth-Maron et al., 2018), QR-DQN (Quantile Regression Deep Q-Network) (Dabney et al., 2018b), and IQN (Implicit Quantile Network) (Dabney et al., 2018a) have shown significant performance improvements over non-distributional counterparts in multiple environments including Atari games and DeepMind Control Suite (Tassa et al., 2018). In DRL, the return distribution is usually represented by a discrete categorical form (Bellemare et al., 2017; Barth-Maron et al., 2018; Qu et al., 2018), quantile function (Dabney et al., 2018b; Zhang et al., 2019; Dabney et al., 2018a), or distribution statistics (Nguyen-Tang et al., 2021). Most of the existing work within DRL framework are value function based and thus are not suitable for tasks with continuous action space. One exception is D4PG (Barth-Maron et al., 2018), an actor-critic type policy gradient algorithm based on DRL. It has demonstrated much better performance (Barth-Maron et al., 2018; Tassa et al., 2018) as compared to its non-distributional counterpart (DDPG). However, it still suffers from various drawbacks such as sample inefficiency and extra burden of parameter tuning, which is largely due to the fact that the return distribution in D4PG is modeled by a discrete categorical distribution or parametrized by Gaussian mixtures.

In this paper, we advocate sample-based representations of return distributions instead of categorical form or quantiles. Our algorithm: sample based distributional policy gradient (SDPG) learns the return distribution by directly generating the return samples via reparameterizing some simple random (e.g. Gaussian) noise samples (Kingma and Welling, 2013). SDPG is an actor-critic type policy gradient based algorithm within DRL framework which employs two neural networks: an actor network to parameterize the policy and a critic network to mimic the target return distribution determined via the distributional Bellman equation based on samples. Since the return distribution is usually 1-dimensional for each state-action pair, we leverage the quantile Huber loss as a surrogate of the Wasserstein distance for comparing return distributions and thereby learning the critic network.

Compared to D4PG, SDPG has the following advantages: i) There is no discretization over the value distributions. The value function network is capable of generating any value distribution, which is not categorical as in D4PG; ii) SDPG does not require the knowledge of the range of the return distribution a priori. In contrast, D4PG requires domain knowledge in terms of bounds on the return distribution; iii) Once the model is trained, the value distribution can be recovered to arbitrary precision by sampling. In contrast, in D4PG, the resolution of the value distribution is fixed once trained. We compare the performance of our algorithm with that of D4PG on multiple OpenAI Gym (Brockman et al., 2016) environments for continuous control tasks. We observe that SDPG exhibits better sample efficiency and performs better than or on-par with D4PG in terms of rewards in almost all the environments.

From a control perspective, our method provides a flexible framework to consider more general optimality criteria other than average control cost/reward in policy gradient methods. For instance, one can easily incorporate risk-measures (Fleming and McEneaney, 1995; Dabney et al., 2018a; Xia, 2016) into the picture by passing the return distribution to a risk utility function.

Related Work: The return distribution was first introduced to study Markov decision processes (Jaquette, 1973; Sobel, 1982). Recently, it was applied to RL problems (Bellemare et al., 2017) which led to DRL. Most of the algorithms proposed under DRL framework are value function based methods that would run into scalability issue for problems with continuous action space. The C51 algorithm (Bellemare et al., 2017), a value based algorithm, represented the return distribution using

a discrete distribution parameterized by 51 number of uniformly spaced atoms in a specified range. Later, the QR-DQN algorithm (Dabney et al., 2018b) proposed to use a discrete set of quantiles to represent the return distribution. QR-DQN was further extended in IQN (Dabney et al., 2018a) to learn the full quantile function. D4PG (Barth-Maron et al., 2018) and Reactor (Gruslys et al., 2017) are the existing policy gradient based methods within DRL framework; D4PG dealt with control problems in continuous action spaces, whereas Reactor was studied in discrete action settings. However, D4PG also utilized the discrete categorical form to represent the return distribution similar to the C51 algorithm (Bellemare et al., 2017), which limits the expressive power of the value distribution network. Distributional policy optimization (DPO) (Tessler et al., 2019) tackled the limitations of policy gradient methods in continuous control that limits the policy representation to parametric distribution classes. To resolve this limitation, DPO used IQN to parameterize the policy, while using a standard mean-value critic to train it. Unlike DPO, Our approach focuses on the expressiveness of the critic, and we discuss the problem of using IQN with comparisons in Section 3.

There have been several works on utilizing samples to represent the return distributions. Generative adversarial networks (GANs) (Goodfellow et al., 2014), one type of sample based generative models, have been employed in value function based approaches: GAN-DQN (Doan et al., 2018), value distribution GAN learning (VDGL) (Freirich et al., 2019), and GAN-DDQN (Hua et al., 2019). GAN-DQN focused on discrete action space and did not show significant improvement over traditional value based methods such as Q-Learning and DQN. VDGL utilized GANs to learn multivariate return distributions. GAN-DDQN combined GAN and IQN to learn the value function for resource allocation in communication systems. DPO parameterizes the policy, while using a standard mean-value critic to train it. An important point to note is that apart from being value based approaches, the existing GAN based DRL methods employ two networks – a generator and a discriminator – for generating return samples by solving a saddle-point problem. In contrast, we utilize quantile Huber loss, as a surrogate of the Wasserstein distance, directly from samples (Deshpande et al., 2018), which results in a single objective rather than saddle-point formulation thereby eliminating the need of a discriminator for learning the return distribution.

2. Background

2.1. Distributional RL

We consider a standard RL problem with underlying model $(\mathcal{X}, \mathcal{A}, R, P, \gamma)$ where \mathcal{X}, \mathcal{A} denote the state and action spaces respectively, $R(x, a)$ is the reward of taking action a at state x , $P(\cdot | x, a)$ is the transition kernel and $0 < \gamma < 1$ is the discount factor. The reward R is assumed to be bounded and can be random in general. The state and action spaces could be either discrete or continuous, though we focus on the more challenging continuous setting. The goal of RL is to find a stationary policy π to maximize the accumulated reward $\mathbb{E} [\sum_{t=0}^{\infty} \gamma^t R(x_t, a_t)]$. Since we focus on continuous-state-action tasks, we restrict the policy π to be deterministic, that is, $a_t = \pi(x_t)$.

The Q-function, denoted by $Q^\pi(x, a)$, describes the expected reward of the agent from taking action $a \in \mathcal{A}$ from state $x \in \mathcal{X}$ at $t = 0$, that is,

$$Q^\pi(x, a) = \mathbb{E} \left[\sum_{t=0}^{\infty} \gamma^t R(x_t, a_t) \right],$$

$$x_t \sim P(\cdot | x_{t-1}, a_{t-1}), a_t = \pi(x_t), x_0 = x, a_0 = a.$$

It satisfies Bellman’s equation (Bellman, 1966)

$$Q^\pi(x, a) = \mathbb{E}R(x, a) + \gamma \mathbb{E}Q^\pi(x', \pi(x') | x, a). \quad (1)$$

Here we have adopted the convention that x' denotes the state succeeding x , i.e., $x' \sim P(\cdot | x, a)$.

In (Bellemare et al., 2017), the authors proposed distributional reinforcement learning (DRL), which relies on a random version of Q-function, defined by

$$Z^\pi(x, a) = \sum_{t=0}^{\infty} \gamma^t R(x_t, a_t), \quad x_t \sim P(\cdot | x_{t-1}, a_{t-1}), a_t = \pi(x_t), x_0 = x, a_0 = a. \quad (2)$$

Clearly, $Q^\pi(x, a) = \mathbb{E}Z^\pi(x, a)$, namely, Q^π is the statistical mean of the random variable Z^π . So in principle, one should be able to recover Q^π using Z^π . The return distribution Z^π contains extra information such as variance that may be used to incorporate risk in the RL framework. The Z function satisfies a modified Bellman’s equation (Bellemare et al., 2017)

$$Z^\pi(x, a) = R(x, a) + \gamma Z^\pi(x', \pi(x') | x, a), \quad (3)$$

where the equation holds in the probability sense.

Multiple approaches have been proposed to model the return distribution $Z^\pi(x, a)$ including distribution quantiles (Dabney et al., 2018b; Zhang et al., 2019) and discrete categorical distribution (Bellemare et al., 2017; Barth-Maron et al., 2018; Qu et al., 2018). In the latter, $Z^\pi(x, a)$ is described a probability vector/histogram with fixed bins. The positions of the bins are chosen a priori that need tuning according to the environment under consideration.

2.2. D4PG

DRL was extended to the policy gradient framework in (Barth-Maron et al., 2018). In the policy gradient framework, the policy π is modeled by a network π_θ directly. For continuous state-action tasks, a widely used method is deterministic policy gradient, which relies on the deterministic policy gradient theorem (Silver et al., 2014). Let $J(\theta)$ be the average return with control strategy π_θ , then

$$\nabla J(\theta) = \mathbb{E} \left[\nabla_\theta \pi_\theta(x) \nabla_a Q^\pi(x, a) |_{a=\pi_\theta(x)} \right]. \quad (4)$$

This theorem is generalized to the DRL setting (Barth-Maron et al., 2018), stated as

$$\nabla J(\theta) = \mathbb{E} \left[\nabla_\theta \pi_\theta(x) \mathbb{E}[\nabla_a Z^\pi(x, a) |_{a=\pi_\theta(x)}] \right], \quad (5)$$

which follows directly from (4). The distributed distributional deterministic policy gradients (D4PG) (Barth-Maron et al., 2018) algorithm is based on this extension (5). It is an actor-critic type algorithm in which the critic learns the return distribution Z^π via a neural network. The return Z is modeled by a categorical discretization or mixture of Gaussians for each (x, a) pair. The actor π_θ is updated via the generalized policy gradient theorem (5) with the expectation being replaced by empirical average.

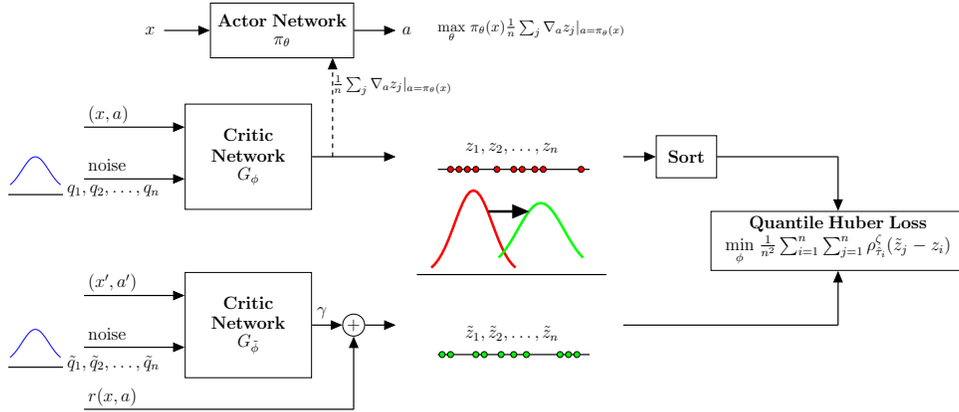


Figure 1: Flow diagram of SDPG.

2.3. Optimal transport and quantile Huber loss

Given two random variable X, Y in the Euclidean space associated with probability distribution μ_X, μ_Y , the Optimal transport (OT) (Villani, 2003) problem seeks the solution to

$$\min_{\nu \in \Pi(\mu_X, \mu_Y)} \int c(x, y) d\nu(x, y) \quad (6)$$

with $\Pi(\mu_X, \mu_Y)$ denoting the set of feasible joint distributions of X and Y . The unit cost function $c(x, y)$ is often taken to be $\|x - y\|^p$, $1 \leq p < \infty$, in which case, (6) defines the Wasserstein distance (Villani, 2003) $W_p(\mu_X, \mu_Y)$ between μ_X and μ_Y .

Computing the Wasserstein distance in general requires solving a linear programming (6), which could be potentially very expensive. One exception is the one-dimensional problem, which has a closed-form solution. When only samples generated by μ_X, μ_Y are available, their Wasserstein distance can be approximated as follows. Let $\{x_1, x_2, \dots, x_n\}, \{y_1, y_2, \dots, y_n\}$ be i.i.d. samples corresponding to μ_X, μ_Y respectively, and $\{\tilde{x}_i\}$ ($\{\tilde{y}_i\}$) be the ascending sorted version of $\{x_i\}$ ($\{y_i\}$), then

$$W_p(\mu_X, \mu_Y)^p \approx \frac{1}{n} \sum_{i=1}^n \|\tilde{x}_i - \tilde{y}_i\|^p. \quad (7)$$

However as noted in (Dabney et al., 2018b), (7) does not give unbiased approximation of the Wasserstein distance. In general,

$$\operatorname{argmin}_{\mu_X} \mathbb{E}[W_p(\mu_X, \hat{\mu}_Y)] \neq \operatorname{argmin}_{\mu_X} W_p(\mu_X, \mu_Y),$$

that is, minimizing the distance to the empirical distribution $\hat{\mu}_Y$ composed of samples from one distribution is not equivalent to minimizing the distance to that distribution μ_Y itself (Dabney et al., 2018b). This hinders the use of stochastic gradient methods to minimize the Wasserstein distance. Utilizing the fact that the return distribution is 1-dimensional, one can leverage the quantile Huber loss (Huber, 1964; Dabney et al., 2018b) as an approximation to the Wasserstein-1 distance. The quantile Huber loss acts as a surrogate of the Wasserstein-1 distance, minimization of which

effectively minimizes the latter. This is given by

$$\frac{1}{n^2} \sum_{i=1}^n \sum_{j=1}^n \rho_{\hat{\tau}_i}^\zeta(\tilde{x}_i - \tilde{y}_j), \quad (8)$$

where $\rho_{\hat{\tau}_i}^\zeta(v) = |\hat{\tau}_i - \delta_{\{v < 0\}}| L_\zeta(v)$,

$$L_\zeta(v) = \begin{cases} 0.5 v^2 & \text{if } |v| < \zeta \\ \zeta(|v| - 0.5 \zeta) & \text{otherwise,} \end{cases}$$

and $\tau_i = \frac{i}{n}, \hat{\tau}_i = \frac{1}{2}(\tau_i + \tau_{i-1}), i = 1, 2, \dots, n$.

Algorithm 1 SDPG

Require: Learning rates α and β , batch size M , sample size n , exploration constant δ ,
Initialize the actor network (π) parameters θ , critic network (G) parameters ϕ randomly
Initialize target networks $(\tilde{\theta}, \tilde{\phi}) \leftarrow (\theta, \phi)$
for the number of environment steps **do**
 Sample M number of transitions $\{(x_t^i, a_t^i, r_t^i, x_{t+1}^i)\}_{i=1}^M$ from the replay pool
 Sample noise $\{q_j^i\}_{j=1}^n \sim \mathbb{P}_q$ and $\{\tilde{q}_j^i\}_{j=1}^n \sim \mathbb{P}_q$, for $i = 1, \dots, M$
 Apply Bellman update to create samples: $\tilde{z}_j^i = r_t^i + \gamma G_{\tilde{\phi}}(\tilde{q}_j^i | (x_{t+1}^i, \pi_{\tilde{\theta}}(x_{t+1}^i)))$ for $j = 1, 2, \dots, n$
 Generate samples $z_j^i = G_\phi(q_j^i | (x_t^i, a_t^i))$ for $j = 1, 2, \dots, n$
 Sort the samples z^i in ascending order
 Update G_ϕ by stochastic gradient descent with learning rate β :
 $\frac{1}{M} \sum_{k=1}^M \frac{1}{n^2} \sum_{i=1}^n \sum_{j=1}^n \rho_{\hat{\tau}_i}^\zeta(z_j^k - z_i^k)$
 Update π_θ by stochastic gradient ascent with learning rate α :
 $\frac{1}{M} \sum_{i=1}^M \pi_\theta(x_t^i) \frac{1}{n} \sum_{j=1}^n \left[\nabla_a z_j^i \right]_{a=\pi_\theta(x_t^i)}$
 Update target networks $(\tilde{\theta}, \tilde{\phi}) \leftarrow (\theta, \phi)$
end for

repeat
 Observe (x_t, a_t, x_{t+1}) and draw reward r_t
 Sample action $a_{t+1} = \pi_\theta(x_{t+1}) + \delta \mathcal{N}(0, 1)$
 Store $(x_t, a_t, r_t, x_{t+1}, a_{t+1})$ in replay pool
until learner finishes

2.4. Reparameterization

Reparameterization is an effective method to model random variables, especially in cases where the goal is to sample from a target distribution instead of modeling them directly using function approximations. Briefly, to model a random variable X with distribution μ_X , the reparameterization trick seeks a neural network to map a simple random variable ϵ (e.g. Gaussian) to the target random variable X , that is, $X = G(\epsilon)$. The hope is that after training, the random variable $G(\epsilon)$ has distribution μ_X . This is extremely useful for sampling purposes because one only needs to sample from the simple distribution ϵ in order to generate samples of X .

3. Algorithm

Our algorithm SDPG consists of two networks: a critic network to learn the return distribution and an actor network to learn the policy. A flow diagram of the SDPG algorithm is shown in Figure 1. We model the return distribution by samples reparameterized via a simple noise distribution \mathbb{P}_q (we use zero-mean Gaussian distribution with unit variance in our implementation). The critic network G_ϕ , parameterized by ϕ , generates the return samples z_1, z_2, \dots, z_n for each state and action pair by transforming the noise samples q_1, q_2, \dots, q_n . These generated samples are compared against the samples $\tilde{z}_1, \tilde{z}_2, \dots, \tilde{z}_n$ generated using the distributional Bellman equation (3) employed in the target critic network $G_{\tilde{\phi}}$. The critic network is learned by minimizing the quantile Huber loss as defined in (8). Thus, the loss function for the critic network is

$$L_{critic}(\phi) = \mathbb{E} \left[\frac{1}{n^2} \sum_{i=1}^n \sum_{j=1}^n \rho_{\hat{\tau}_i}^\zeta(\tilde{z}_j - z_i) \right], \quad (9)$$

where $z_1 \geq z_2 \geq \dots \geq z_n$ are samples after sorting. We emphasize that the sorting is important here to associate each sample with a reasonable $\hat{\tau}$. This is different to (Dabney et al., 2018a) where $\hat{\tau}$ itself is the random seed over $[0, 1]$.

The actor network π_θ , parameterized by θ , outputs the action $\pi_\theta(x)$ given a state x . The actor network receives feedback from the critic network G_ϕ in terms of the gradients of the return distribution with respect to the actions determined by the policy. This feedback is used to update the actor network by applying a distributional form of the policy gradient theorem given by (5). Therefore, the gradient of the actor network loss function is

$$\nabla_\theta L_{actor}(\theta) = \mathbb{E} \left[\nabla_\theta \pi_\theta(x) \frac{1}{n} \sum_{j=1}^n [\nabla_a z_j] \Big|_{a=\pi_\theta(x)} \right]. \quad (10)$$

The SDPG algorithm is summarized in Algorithm 1. The network parameters of actor and critic networks are updated alternatively in stochastic gradient ascent/descent fashion.

In contrast to the categorical parameterization of the return distribution considered in D4PG, we use samples to represent the return distribution. Since the return distribution is required to be differentiable with respect to the network parameters in order to be learned, we utilize reparameterization trick discussed in Section 2.4 to model the return distribution via random noise input. This allows us to learn a continuous distribution via samples as opposed to a discrete-valued categorical distribution in D4PG. Moreover, D4PG requires a projection step in every iteration during training in order to make the target distribution resulting from the distributional Bellman equation coincide with the support of categorical parameterized distribution being learned, which introduces extra approximation errors (Rowland et al., 2019); SDPG eliminates the need of such a projection step during training. Furthermore, the range of the discretized grid required in D4PG must be tuned according to the reward values for each environment; SDPG does not require such tuning. Another advantage of SDPG is that one can recover the return distribution to arbitrary precision by sampling from the trained critic network. However, the resolution of the return distribution is fixed in D4PG and the critic network has to be trained again from the scratch if one wants to change the resolution.

Other than sample based reparameterization, the implicit quantile network (IQN) (Dabney et al., 2018a), which learns the reward distribution in terms of the inverse cumulative distribution function

(CDF) for each state-action pair, can also be extended to policy gradient method for continuous action space. The resulting algorithm is similar to SDPG and is named IQN policy gradient (IQN-PG); details are skipped due to page limit. One drawback of the IQN-PG is that the CDFs learned during training are not monotone and thus are not valid CDFs. In our experiments, IQN-PG has comparable performance to SDPG.

In principle, one can adopt a conditional GAN (Gauthier, 2014) method to learn the return distribution; this however requires an extra discriminator network. We believe this is an overkill for DRL framework as the return distributions are one dimensional. Our strategy using quantile Huber loss, as a surrogate of the Wasserstein distance, to learn return distributions directly is a better choice.

4. Experimental Results

We compare the performance of SDPG with D4PG and IQN-PG algorithms on a range of challenging continuous control tasks from the OpenAI Gym environments. Noise was injected to increase the randomness of the dynamics.

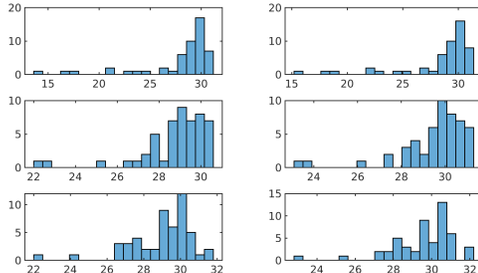


Figure 2: Histograms of the 51 samples from random state-action pairs after training on BipedalWalker-v2: (left) histograms of samples by the critic network and (right) corresponding histograms with the distributional Bellman equation.

For both actor and critic networks, we use a two layer feedforward neural network with hidden layer sizes of 400 and 300, respectively, and rectified linear units (ReLU) between each hidden layer. We used batch normalization on all the layers. Moreover, the output of the actor network is passed through a hyperbolic tangent (Tanh) activation unit.

In all experiments we use learning rates of $\alpha = \beta = 1 \times 10^{-4}$, batch size $M = 256$, exploration constant $\delta = 0.3$, and $\zeta = 1$. We use a replay table of size $R = 1 \times 10^6$ for all the domains except $R = 0.2 \times 10^6$ for Pendulum and LunarLander. Across all the tasks, for D4PG we use 51 atoms to represent the categorical distribution and similarly, for SDPG we use $n = 51$ number of samples to represent return distributions. Moreover, we run each task for a maximum of 1000 steps per episode. Note that SDPG is a centralized algorithm at this moment for a single agent. Thus, the distributed feature in D4PG is deactivated for fair comparison. One can easily establish a distributed version of SDPG. Since SDPG requires sorting operation during training, SDPG takes a little more time per episode (roughly $\times 1.3$) than D4PG.

4.1. Training and Evaluation

First we demonstrate the ability of the critic network to learn the return distribution utilizing the quantile Huber loss (8) based on the distributional Bellman equation. Figure 2 shows the histograms of the samples generated by the learned critic network and the corresponding histograms of samples

SDPG

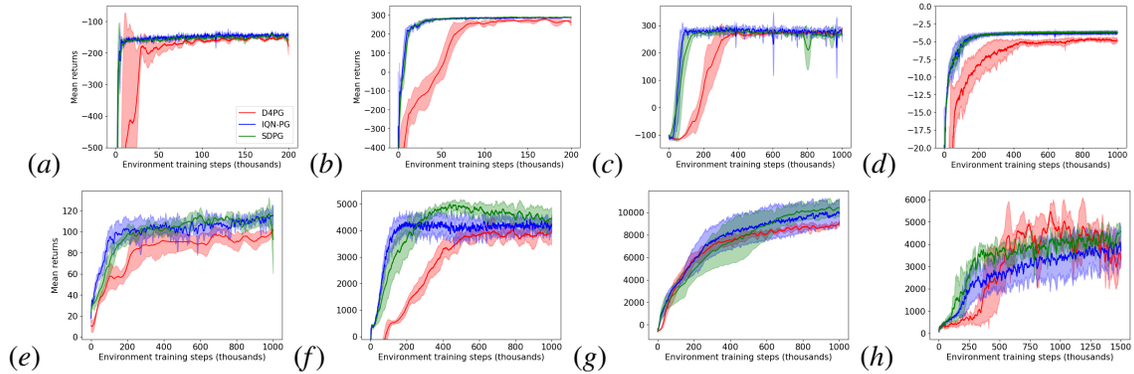


Figure 3: Mean returns on different environments: (a) Pendulum-v0, (b) LunarLander-v2, (c) BipedalWalker-v2, (d) Reacher-v2, (e) Swimmer-v2, (f) Ant-v2, (g) HalfCheetah-v2, and (h) Humanoid-v2. The shaded region represents standard deviation of the average returns over 5 seeds. The curves are smoothed uniformly for visual clarity. Note that in the original D4PG paper, the environments are from DeepMind Control Suite (Tassa et al., 2018) with the rewards bounded in the range $[0, 1]$ for all the domains.

Table 1: Comparison of average maximal returns \pm one standard deviation over 5 different trials.

Domain	Simulator	Train Steps	Reward		
			D4PG	IQN-PG	SDPG
Pendulum	Classic control	0.2×10^6	-144.63 ± 6.72	-128.81 ± 3.35	-142.46 ± 5.21
LunarLander	Box2D	0.2×10^6	283.22 ± 1.63	290.68 ± 1.72	291.71 ± 0.90
BipedalWalker	Box2D	1.0×10^6	289.66 ± 3.26	302.55 ± 6.64	301.91 ± 4.38
Reacher	MuJoCo	1.0×10^6	-4.40 ± 0.28	-3.41 ± 0.15	-3.62 ± 0.20
Swimmer	MuJoCo	1.0×10^6	115.52 ± 4.94	126.90 ± 5.91	128.38 ± 6.20
Ant	MuJoCo	1.0×10^6	5058.84 ± 93.57	4991.20 ± 288.57	5762.83 ± 194.67
HalfCheetah	MuJoCo	1.0×10^6	9565.06 ± 209.77	10081.30 ± 1362.21	10607 ± 845.04
Humanoid	MuJoCo	1.5×10^6	6064.14 ± 192.84	4755.66 ± 623.44	5093.62 ± 402.25

generated based on the distributional Bellman equation on BipedalWalker-v2 domain. Clearly, the histograms match almost perfectly which demonstrates that the critic network in SDPG successfully learns the target return distribution determined via the distributional Bellman equation.

To compare the three algorithms, we train five different instances of each algorithm with random seeds and each performing one evaluation rollout every 5000 environment steps. Figure 3 shows the comparison of mean returns in different environments. It is evident from the figures that SDPG exhibits significantly better sample efficiency than D4PG on almost all the environments. The performance of SDPG is close to IQN-PG when sample efficiency is concerned. Moreover in terms of average returns, SDPG as well as IQN-PG perform better than D4PG on all the domains except Humanoid-v2. This maybe due to insufficient training steps. The performance of SDPG and IQN-PG for Humanoid-v2 keeps increasing during the entire training process and this trend is expected to continue.

We evaluate the performance of the algorithms based on two criteria: average returns and sample efficiency. Table 1 lists the maximal mean returns (the average of the maximal returns over different trials) along with the standard deviation. The average returns are evaluated every 5000 training

Table 2: Comparison of sample efficiency in terms of number of episodes needed to reach a certain threshold. The episodes reported here are the smallest m for which the mean episode reward over m^{th} and $(m + 10)^{th}$ episodes crosses a certain return threshold. The thresholds are chosen from (Gu et al., 2017).

Environment	Threshold	Episodes		
		D4PG	IQN-PG	SDPG
Pendulum	-150	1605	1142	481
LunarLander	200	2994	1244	1039
BipedalWalker	250	4659	2996	3039
Reacher	-7	36623	24837	11859
Swimmer	90	3492	9158	3867
Ant	3500	5567	4168	4437
HalfCheetah	4700	3217	17585	7692
Humanoid	2500	57142	65915	51191

steps over 100 episodes. We observe that the returns for SDPG and IQN-PG are significantly better than D4PG for all the environments except Humanoid. To compare the sample efficiency of the three algorithms, we list the number of episodes needed to reach certain return threshold in Table 2. The episode numbers reported in the table are averaged over 5 different trials and for each trial the episode number is the number of episodes required before the reward crosses a certain threshold. It is evident that SDPG and IQN-PG require significantly smaller number of episodes than D4PG on many environments.

Lastly, we study the effect of varying number of samples representing the return distribution while training. Figure 4 (a) depicts the training curves with different samples on Ant-v2 domain. For a fixed number of samples, the algorithm is trained for five different seeds: the solid lines represent the mean returns over five trials and the shaded region represent the corresponding standard deviation. Initially, increasing the number of samples improves the performance in terms of efficiency as well as returns, but the trend does not continue for large sample size. Furthermore, Figure 4(b) compares the performance by increasing the number of atoms in D4PG on LunarLander domain. It can be observed that increasing the number of atoms in D4PG does not improve the performance.

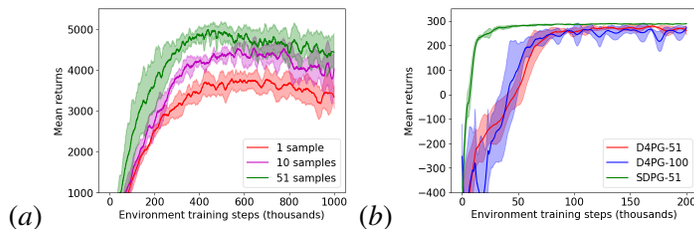


Figure 4: Effect of number of samples: (a) SDPG on Ant-v2 and (b) D4PG on LunarLander-v2.

5. Conclusion

In this paper, driven by applications in continuous action space, we proposed sample-based distributional policy gradient (SDPG) algorithm for learning the policy. This algorithm is a combination of an actor-critic type of policy gradient method and DRL. Our algorithm showed better sample efficiency than D4PG in most environments and performed better than D4PG in terms of average returns.

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