

Meta Reinforcement Learning

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Abstract—Meta reinforcement learning is a research area of reinforcement learning. It applies meta learning approaches on reinforcement learning tasks to generalize the training result of reinforcement learning and adapt it to unseen tasks. Currently, multiple meta reinforcement learning methodologies are being researched, such as optimization-based approach, hyperparameter-based approach and so on. In this report, we first provide some fundamental knowledge about meta reinforcement learning. Then we give an overview of state-of-the-art meta reinforcement learning algorithms. Finally, a couple of algorithms are discussed in detail and their experimental results are also presented.

Disclaimer: Section...by Ruoheng Ma and Section...by Meng Zhang.

I. INTRODUCTION

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II. FUNDAMENTAL KNOWLEDGE

In this section, we provide some fundamental knowledge for ease of understanding the algorithms discussed in the following sections.

A. Formulation of Reinforcement Learning and Meta Learning

In reinforcement learning, a value function

$$G_t =$$

no predefined value function is provided to tell the score of an action. In practice, a proxy of the true value function, known as a *return*, is used for estimation and optimization for the value function.

B. Hyperparameter

C. empty

III. META-GRADIENT REINFORCEMENT LEARNING

One of the algorithm in meta reinforcement learning is the meta-gradient reinforcement learning introduced in [1] by Google DeepMind. The main idea of this algorithm is to update the meta-parameters η to achieve a better learning performance. The details of this algorithm is presented in this section.

In deep reinforcement learning, the environment model is often unknown to the agent. Therefore, the true value function has to be approximated by a function known as *return* with parameter θ . This return function plays an important role in determining the characteristics of the reinforcement learning

algorithm. Usually, the return function is parameterised by a discount factor γ and the bootstrapping parameter λ , which are denoted as meta-parameters η in this paper. Throughout normal training process, these two parameters γ and λ are hand-selected and held fixed. Following are the n -step return[2] depending on fixed γ :

$$g_\eta(\tau_t) = R_{t+1} + \gamma R_{t+2} + \dots + \gamma^{n-1} R_{t+n} + \gamma^n v_\theta(S_{t+n})$$

where $\eta = \{\gamma\}$ and the λ return[3] depending on fixed γ and λ :

$$g_\eta(\tau_t) = R_{t+1} + \gamma(1 - \lambda)v_\theta(S_{t+1}) + \gamma\lambda g_\eta(\tau_{t+1})$$

where $\eta = \{\gamma, \lambda\}$. In addition, the parameter θ is also updated by the following formula, taking fixed η into account:

$$\theta' = \theta + f(\tau, \theta, \eta)$$

where τ is the sample of experience being considered.

In order to achieve better performance, the algorithm introduced in [1] updates η during the training process. Besides an objective function $J(\tau, \theta, \eta)$, whose value the parameter θ is being updated to reduce, a meta-objective defined as $\bar{J}(\tau', \theta', \bar{\eta})$ is also included into this algorithm and the purpose of updating η is to increase its value.

Here is how the algorithm works: First, the algorithm starts with parameter θ and applies the update function on the first sample, resulting in a updated parameter θ' . After that, the performance is measured by $\bar{J}(\tau', \theta', \bar{\eta})$, where τ' is the next sample following τ and $\bar{\eta}$ is a fixed meta-parameter. Then, the gradient of the meta-objective with respect to the meta-parameters η is obtained by applying the chain rule:

$$\frac{\partial \bar{J}(\tau', \theta', \bar{\eta})}{\partial \eta} = \frac{\partial \bar{J}(\tau', \theta', \bar{\eta})}{\partial \theta'} \frac{d\theta'}{d\eta}$$

Finally, $\Delta\eta$ is calculated to maximize the meta-objective:

$$\Delta\eta = -\beta \frac{\partial \bar{J}(\tau', \theta', \bar{\eta})}{\partial \theta'} \frac{\partial f(\tau, \theta, \eta)}{d\eta}$$

where β is the learning rate for updating η .

The validation of meta-gradient algorithm is executed on Arcade Learning Environment with an agent built with the IMPALA framework. The agents are evaluated on 57 different Atari games. The result is shown below:

	η	Human starts		No-op starts	
		$\gamma = 0.99$	$\gamma = 0.995$	$\gamma = 0.99$	$\gamma = 0.995$
IMPALA	$\{\}$	144.4%	211.9%	191.8%	257.1%
Meta-gradient	$\{\lambda\}$	156.6%	214.2%	185.5%	246.5%
		$\bar{\gamma} = 0.99$	$\bar{\gamma} = 0.995$	$\bar{\gamma} = 0.99$	$\bar{\gamma} = 0.995$
Meta-gradient	$\{\gamma\}$	233.2%	267.9%	280.9%	275.5%
Meta-gradient	$\{\gamma, \lambda\}$	221.6%	292.9%	242.6%	287.6%

Fig. 1. Experiment result of meta-gradient algorithm. "Human starts" means episodes are initialized to a state that is randomly sampled from human play, while "No-op starts" means each episode is initialized with a random sequence of no-op actions.

IV. EVOLVED POLICY GRADIENT

Another algorithm for meta reinforcement learning is evolved policy gradient (EPG)[4]. Unlike meta gradient algorithm which focuses on learning the return function, EPG aims to learn a surrogate loss function $L_\phi = L_{\phi + \sigma \epsilon_i}$, which is parameterized by parameter ϕ along with a small Gaussian noise $\epsilon_i \sim \mathcal{N}(0, \mathbf{I})$ as perturbed terms.

EPG contains an inner loop and an outer loop during its execution. In inner loop, the algorithm functions as normal policy gradient algorithm that tries to update parameter θ according to the following formula:

$$\theta^* = \arg \min_{\theta} \mathbb{E}_{\tau \sim \mathcal{M}, \pi_{\theta}} [L_{\phi}(\pi_{\theta}, \tau)]$$

where θ^* is the updated parameter θ , \mathcal{M} is a sampled markov decision process, τ is an episode of \mathcal{M} with horizon H . The objective of the inner loop is to minimize the loss function L_{ϕ} . In outer loop, the loss function parameter ϕ is updated as shown below:

$$\phi^* = \arg \max_{\phi} \mathbb{E}_{\mathcal{M} \sim p(\mathcal{M})} \mathbb{E}_{\tau \sim \mathcal{M}, \pi_{\theta^*}} [R_{\tau}]$$

where $p(\mathcal{M})$ is a distribution over MDPs, π_{θ^*} is the agent's policy trained with the loss function L_{ϕ} and $R_{\tau} = \sum_{t=0}^H \gamma^t r_t$ is the discounted episodic return of τ . The goal of the outer loop is to achieve high expected returns in the MDP distribution.

Pseudocode of EPG algorithm is shown below:

```

Algorithm 1: Evolved Policy Gradients (EPG)
1 [Outer Loop] for epoch  $e = 1, \dots, E$  do
2   Sample  $\epsilon_w \sim \mathcal{N}(0, \mathbf{I})$  and calculate the loss parameter  $\phi + \sigma \epsilon_w$  for  $v = 1, \dots, V$ 
3   Each worker  $w = 1, \dots, W$  gets assigned noise vector  $\lceil wV/W \rceil$  as  $\epsilon_w$ 
4   for each worker  $w = 1, \dots, W$  do
5     Sample MDP  $\mathcal{M}_w \sim p(\mathcal{M})$ 
6     Initialize buffer with  $N$  zero tuples
7     Initialize policy parameter  $\theta$  randomly
8     [Inner Loop] for step  $t = 1, \dots, U$  do
9       Sample initial state  $s_t \sim p_0$  if  $\mathcal{M}_w$  needs to be reset
10      Sample action  $a_t \sim \pi_{\theta}(\cdot | s_t)$ 
11      Take action  $a_t$  in  $\mathcal{M}_w$  and receive  $r_t, s_{t+1}$ , and termination flag  $d_t$ 
12      Add tuple  $(s_t, a_t, r_t, d_t)$  to buffer
13      if  $t \bmod M = 0$  then
14        With loss parameter  $\phi + \sigma \epsilon_w$ , calculate losses  $L_i$  for steps  $i = t - M, \dots, t$ 
15        using buffer tuples  $i - N, \dots, i$ 
16        Sample minibatches  $mb$  from last  $M$  steps shuffled, compute  $L_{mb} = \sum_{j \in mb} L_j$ ,
17        and update the policy parameter  $\theta$  and memory parameter (Eq. (5))
18      In  $\mathcal{M}_w$ , using trained policy  $\pi_{\theta}$ , sample several trajectories and compute mean return  $R_w$ 
19    Update the loss parameter  $\phi$  (Eq. (6))
20 Output: Loss  $L_{\phi}$  that trains  $\pi$  from scratch according to inner loop scheme, on MDPs  $\sim p(\mathcal{M})$ 

```

Fig. 2. EPG algorithm.

The algorithm works as follows: Assumed that W workers are working in the inner loop. Then, at the beginning of each epoch in the outer loop, V multivariate normal vectors $\epsilon_v \in \mathcal{N}(0, \mathbf{I})$ of the same dimension as the loss function parameter ϕ are generated and assigned to V loss functions $L_w = L_{\phi + \sigma \epsilon_v}$ respectively, with v being the v -th generated perturbed parameters.

Afterwards, the inner loop launches, each worker samples a random MDP from the task distribution $\mathcal{M}_w \sim p(\mathcal{M})$ and trains the policy π_{θ} along with the loss function L_w given from the outer loop and updates the parameter θ as follows:

$$\theta \leftarrow \theta - \delta_{in} \cdot \nabla_{\theta} L_w(\pi_{\theta}, \tau_{t-M}, \dots, t)$$

At the end of the inner-loop training, the return R_w is returned by each worker and is aggregated in the outer loop. Then, the parameter ϕ of the loss function is updated according to the rule shown below:

$$\phi \leftarrow \phi + \delta_{out} \cdot \frac{1}{V} \sum_{v=1}^V F(\phi + \sigma \epsilon_v) \epsilon_v$$

$$\text{where } F(\phi + \sigma \epsilon_v) = \frac{R_{(v-1)*W/V+1} + \dots + R_{v*W/V}}{W/V}.$$

The architecture of the loss function can be seen in the following figure:

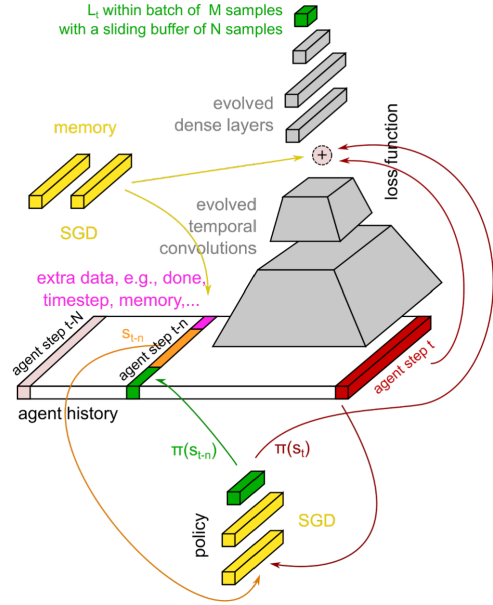


Fig. 3. Architecture of loss function L_{ϕ} .

As depicted in the diagram above, the architecture contains a memory unit, which is actually a single-layer neural network accepting constant input vector and is used to store the loss function value, as well as an experience buffer with limited storage, which stores the agent's N most recent experience steps, in the form of a list of tuples (s_t, a_t, r_t, d_t) at time step t , where d_t is the trajectory termination flag.

In addition, the architecture consists of temporal convolutional layers which generate a context vector $f_{context}$, and dense layers, which output the loss. The dense layer outputs the loss function L_t at time step t by taking a batch of M sequential samples $\{s_i, a_i, d_i, mem, f_{context}, \pi_\theta(\cdot|s_i)\}_{i=t-M}^t$. To generate the context vector, first, the data samples $\{s_i, a_i, d_i, mem, \pi_\theta(\cdot|s_i)\}_{i=t-N}^t$ are stack together to form a matrix, second, this matrix is processed by the temporal convolutional layers which outputs the context vector $f_{context}$.

This algorithm is evaluated under several randomized continuous control MuJoCo environments. For example, the following diagrams show results on RandomHopper, RandomWalker and RandomReacher.

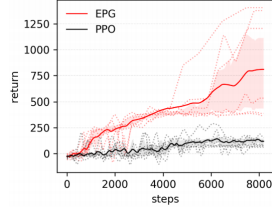


Fig. 4. RandomHopper testtime training over 128 (policy updates) x64 (update frequency) = 8196 timesteps: PPO vs no-reward EPG.

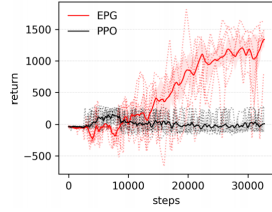


Fig. 5. RandomWalker testtime training over 256 (policy updates) x128 (update frequency) = 32768 timesteps: PPO vs no-reward EPG.

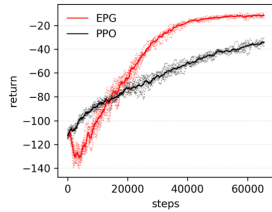


Fig. 6. RandomReacher testtime training over 512 (policy updates) x128 (update frequency) = 65536 timesteps: PG vs no-reward EPG.

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V. MAML

VI. CONCLUSION

The conclusion goes here.

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