# PA230: Reinforcement Learning

Petr Novotný

"Good and evil, reward and punishment, are the only motives to a rational creature; these are the spur and reins whereby all mankind are set on work and guided."

John Locke, Some Thoughts Concerning Education (1693)



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  - mainly binary classification (accepted/not accepted)
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  - can (but do not have to) be done in pairs (pairs can differ across the individual assignments)
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- Exam:
  - oral
  - each attempt counts ? (unlike the Brázdil system)
  - in general, knowledge of anything mentioned on the slides can be required, unless explicitly marked with "nex" (like the Brázdil system)

### Team

• Lecturer: Petr Novotný



• HW team:



Martin Kurečka



Václav Nevyhoštěný



Vít Unčovský

#### Communication

Official discord server:



https://discord.gg/9mxTgYhcdB

- Official communication forum of the course: falls under the university ethical guidelines.
- Use your real name for posting (you can set-up an account under your IS email if necessary).

### Reading

- Compulsory:
  - these slides,
  - material explicitly prescribed by these slides (not much).
- Recommended:
  - Sutton & Barto: Reinforcement Learning: An Introduction (2nd ed.), available at http://incompleteideas.net/book/RLbook2020.pdf
    - henceforth referenced as "S&B"
  - slides by David Silver https://www.davidsilver.uk/teaching/
  - CMU slides https://www.andrew.cmu.edu/course/10-703/
  - more specific literature recommendations will be given for each topic later

What, Why, When, How,

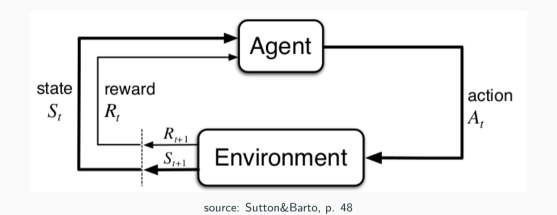
**Reinforcement Learning:** 

& Other Questions

### Types of machine learning

- unsupervised
  - spot "useful" patterns in data
- supervised
  - given labeled data, predict labels on unlabeled data
- reinforcement
  - · agents and decision-making
  - agency = "the ability to take action or to choose what action to take" (Cambridge Dictionary)

### **General RL scheme**



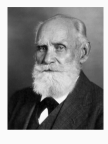
Keywords: sequential, dynamic, subject to uncertainty

### RL: Objective and approach

• Objective: Design a decision policy (= agent behavior) which prescribes to the agent how to act in different situations (states), typically so as to achieve some goal.

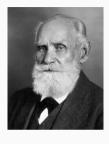
# RL: Objective and approach

- Objective: Design a decision policy (= agent behavior) which prescribes to the agent how to act in different situations (states), typically so as to achieve some goal.
- Approach: Start with (± random) behavior and adapt it based on past experience via the law of effect:
  - actions with good/bad consequences for the agent are more/less likely to be repeated by the agent (within the same context)



(1849-1936) classical conditioning

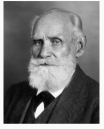
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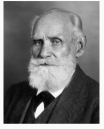
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J.B. Watson (1878-1958) behaviorist manifesto



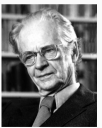
I.P. Pavlov (1849-1936) classical conditioning



E. Thorndike (1874-1949) law of effect



J.B. Watson (1878-1958) behaviorist manifesto



B.F. Skinner (1904-1990) radical behaviorism, reinforcement, rewards

# Learning by trying & XX dilemma

Underlying the RL approach is the idea of learning by trying:

- first, act more or less randomly (exploration)
  - integral part of early human development
- continually adapt behavior according to experience and feedback from the environment (exploitation)
  - strength of feedback pprox strength of behavior adaptation

Balancing exploration and exploitation (XX) is a recurring theme in RL.

# Incomplete history of RL in computer science I

A. Turing

"Learning by trying" machines and software, ad hoc approaches:



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And many more. . . Recommended: S&B: Sec. 1.7.

# Incomplete history of RL in computer science II

#### Mathematical foundations of sequential decision making:



R. Bellman (1920-1984)



R. Howard (b. 1934)

- Formalization via Markov decision processes (MDPs)
- value iteration (attributed to Bellman, 1957)
- policy iteration (attributed to Howard, 1960)

# Incomplete history of RL in computer science III

Since late 1980's: synthesis – learning by trial in MDPs



R. Sutton



A. Barto



...and many more.

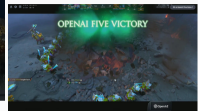


C. Watkins

Q-learning



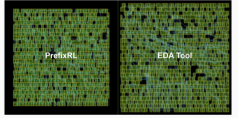


















# Words of caution (and controversy) (nex)





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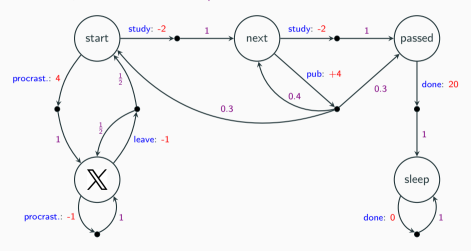


**Mathematical Foundations** 

of Sequential Decision-Making

# **MDP Example**

MDP with actions, rewards and transition probabilities.



#### **Markov Decision Process**

Given a set X, we denote  $\mathcal{D}(X)$  the set of all probability distributions over X.

#### Definition 1

A Markov decision process (MDP) is a tuple (S, A, p, r) where

- $\bullet$   $\mathcal{S}$  is a set of states,
- A is a set of actions,
- $p: \mathcal{S} \times \mathcal{A} \to \mathcal{D}(\mathcal{S})$  is a probabilistic transition function,
- $r: \mathcal{S} \times \mathcal{A} \to \mathbb{R}$  is a reward function.

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The p, r can be partial functions: action a is enabled in state s if both p(s, a) and r(s, a) are defined. We denote by  $\mathcal{A}(s)$  the set of all actions enabled in s.

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  - ullet agent selects action  $a_t \in \mathcal{A}(s_t)$
  - the environment responds with next state  $s_{t+1} \sim p(s_t, a_t)$  and with immediate reward  $r_{t+1} = r(s_t, a_t)$
  - t is incremented and the process repeats in the same fashion forever

Thus, the agent produces a trajectory  $\tau = s_0, a_0, r_1, s_1, a_1, r_2, s_2, a_2, r_3, s_3, \dots$ 

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au is produced randomly (due to p and possibly also agent choices being probabilistic): it is a random variable and so are its components: we define random variables

- $S_t$  = state at time step t
- $A_t$  = action at time step t
- ullet  $R_t = \text{reward received just before entering } S_t$

### **Policies**

#### Definition 2

A history is a finite prefix of a trajectory ending in a state, i.e., an object of type

$$s_0, a_0, r_1, s_1, a_1, r_2, \ldots, a_{t-1}, r_t, s_t \in (\mathcal{S} \cdot \mathcal{A} \cdot \mathbb{R})^* \mathcal{S}.$$

we denote by last(h) the last state of a history h.

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#### Definition 3

A policy is a function  $\pi: (S \cdot A \cdot \mathbb{R})^*S \to \mathcal{D}(A)$  which to each history h assigns a probability distribution over  $\mathcal{A}(last(h))$ .

A policy is by definition an infinite object!

### **MD** policies

#### Definition 4

A policy  $\pi$  is:

- memoryless if  $\pi(h) = \pi(h')$  whenever last(h) = last(h') (we can view memoryless policies as objects of type  $\pi \colon \mathcal{S} \to \mathcal{D}(\mathcal{A})$ );
- deterministic if  $\pi(h)$  always assigns probability 1 to one action, and zero to all others (we can view det. policies of objects of type  $\pi: (S \cdot A \cdot \mathbb{R})^*S \to A$ ).

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#### Definition 5

A policy  $\pi$  is MD (memoryless deterministic) if it is both memoryless and deterministic.

# Dynamics of MDPs (more precise)

Given a distribution  ${\mathcal I}$  of initial states and a policy  $\pi$ 

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- in each time step t, let  $h_t$  be the history produced so far; then:
  - agent selects action  $a_t \in \mathcal{A}(s_t)$  according to  $\pi$ , i.e.  $a_t \sim \pi(h_t)$
  - the environment responds with next state  $s_{t+1} \sim p(s_t, a_t)$  and with immediate reward  $r_{t+1} = r(s_t, a_t)$ , the history is extended by  $a_t, r_t, s_{t+1}$ ,
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In particular, each policy  $\pi$  together with a distribution  $\mathcal{I}$  of initial states induce a probability measure  $\mathbb{P}^{\pi}$  over the trajectories of the MDP.<sup>1</sup>

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#### Exercise 6

In the "study" MDP, consider an MD policy  $\pi$  s.t.  $\pi(start) = study$  and  $\pi(next) = pub$ . Compute the following quantities:

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•  $\mathbb{E}^{\pi}[R_3]$ 

### Memorylessness

In this course, we will almost exclusively focus on memoryless policies. Hence, from now on, policy = memoryless policy. General policies will be referred to as history-dependent policies should the need arise.

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Why memoryless?

Intuition: Markov property of MDPs: next step depends only on the current state and on action performed in the current step. Hence, intuitively there is no need for a policy to remember the past so as to "play well".

The sufficiency of memoryless policies does not extended to more general/complex decision-making settings (not covered in this course), such as:

- partially observable MDPs
- non-stationary environments
- quantile/risk-aware MDPs, etc.

# Returns (payoffs)

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#### Definition 7

Let  $\gamma \in [0,1)$  be a discount factor.

For a trajectory  $\tau=s_0, a_0, r_1, s_1, a_1, r_2, s_2, a_2, r_3, s_3, \ldots$  we define the discounted return (or payoff) of  $\tau$  to be the quantity

$$G(\tau) = r_1 + \gamma \cdot r_2 + \gamma^2 \cdot r^3 + \cdots + \gamma^3 \cdot r^4 = \sum_{i=0}^{\infty} \gamma^i \cdot r_{i+1}.$$

Equivalently

$$G = \sum_{i=0}^{\infty} \gamma^i \cdot R_{i+1}$$

# Returns (variants)

• Finite horizon (FH): additionally, we are given a finite decision horizon  $H \in \mathbb{N} \cup \{\infty\}$ . The return is that counted only up to step H:

$$G^H = \sum_{i=0}^{H-1} \gamma^i \cdot R_{i+1}$$

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For finite H, the discount factor can be 1.  $H = \infty$  corresponds to the original definition.

• Episodic returns: In episodic tasks, there is a distinguished set  $Term \subseteq S$  of terminal states which is guaranteed to be reached with probability 1 under any policy. We denote by T a random variable denoting the first point in time when we hit a terminal state. We count rewards only up to that time:

$$G^T = \sum_{i=1}^{T-1} \gamma^i \cdot R_{i+1}$$

Can be modeled under original definition by "sink" states.

### Types of returns: discussion

- We will typically omit the superscripts since the type of task considered will be known from the context.
- We have  $G^H \to G$  (pointwise) as  $H \to \infty$ . I.e., finite-horizon returns with high enough H approximate the standard (infinite-horizon) case.
- In real world, we typically deal with FH or episodic tasks: we cannot wait infinite time to learn something from a trajectory. However, the infinite-horizon case can be viewed as a neat mathematical abstraction of the FH&episodic tasks, and the classical sequential decision-making theory is most developed for the infinite horizon case.

# Policy and state values

### Definition 8

Let  $\pi$  be a policy and s a state. The value of  $\pi$  in state s is the quantity

$$\mathbf{v}^{\pi}(\mathbf{s}) = \mathbb{E}^{\pi}[G \mid S_0 = \mathbf{s}].$$

#### Exercise 9

Discuss the values of MD policies in our running example.

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#### Exercise 9

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### Definition 10

The (optimal) value of state s is the quantity

$$v^*(s) = \sup_{\pi} v^{\pi}(s).$$

# **Optimality**

#### Definition 11

Let  $\pi$  be a policy and  $\varepsilon > 0$ . We say that  $\pi$  is  $\varepsilon$ -optimal in state s if

$$v^{\pi}(s) \geq v^{*}(s) - \varepsilon$$
.

We say that  $\pi$  is optimal in s is it is 0-optimal in s, i.e. if

$$v^{\pi}(s)=v^*(s).$$

A policy is ( $\varepsilon$ -)optimal if it is ( $\varepsilon$ -)optimal in every state.

### **Existence of optimal policies**

### Theorem 12: (Classical result, not formally proven here)

Let  $\mathcal M$  be a finite MDP (i.e., the state and action sets are finite) with infinite-horizon returns. Then there exists an optimal MD policy. Moreover, an optimal MD policy can be computed in polynomial time.

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Agent control solved? NO! "Only" works if you can actually construct the MDP model of your environment and fit it into a computer. Otherwise, we use reinforcement learning.

**Exact Planning** 

Value & Policy Iteration

with Known Model:

### Goal of this lecture

Algorithms that compute the optimal value vector  $v^*$  and some optimal MD policy  $\pi^*$  given a full knowledge of an MDP  $\mathcal{M}$ .

# Polynomial-time algorithm

MDPs can be solved by linear programming (LP)

maximize 
$$\vec{c} \cdot \vec{x}$$
 subject to  $A \cdot \vec{x} \leq \vec{b}$ 

- LP can be solved in polynomial time by so-called interior-point algorithms.
- However, we typically use other, MDP-specific algorithms: value iteration (VI) and policy iteration (PI). These are not polynomial-time in general, but typically faster on practical instances.
- Moreover, most truly RL algorithms can be seen as approximate generalizations of VI or PI (or both).

### **Example: Policy evaluation**

#### Exercise 13

Consider all four MD policies in our running "pub or study" example that always try to quit when in X state. Compute the values of these policies in the initial state start.

# Policy evaluation equations

#### Theorem 14

For any memoryless policy  $\pi$  and any state s it holds:

$$v^{\pi}(s) = \sum_{a \in \mathcal{A}(s)} \pi(a|s) \cdot \underbrace{\left[r(s,a) + \gamma \cdot \sum_{s' \in \mathcal{S}} p(s'|s,a) \cdot v^{\pi}(s')\right]}_{\stackrel{\text{def}}{=} q^{\pi}(s,a)}.$$

# **Bellman optimality equations**

#### Theorem 15

The following holds for any state s:

$$v^*(s) = \max_{a \in \mathcal{A}(s)} \underbrace{\left[r(s, a) + \gamma \cdot \sum_{s' \in \mathcal{S}} p(s'|s, a) \cdot v^*(s')\right]}_{\stackrel{\text{def}}{=} q^*(s, a)}$$

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• Note: the policy evaluation equations are a special case of the Bellman ones: given a policy  $\pi$ , we can consider an MDP  $\mathcal{M}^{\pi}$  in which there is a single action \* enabled in each state and the probability of transition  $s \stackrel{*}{\to} s'$  equals  $\sum_{a \in \mathcal{A}(s)} \pi(a|s) \cdot p(s'|s,a)$ . Then  $\mathcal{M}^{\pi}$  mimics the behavior of  $\pi$  in  $\mathcal{M}$  and Bellman eq's in  $\mathcal{M}^{\pi}$  = evaluation equations for  $\pi$  in  $\mathcal{M}$ .

### Bellman optimality equations

#### Theorem 15

The following holds for any state s:

$$v^*(s) = \max_{a \in \mathcal{A}(s)} \underbrace{\left[r(s, a) + \gamma \cdot \sum_{s' \in \mathcal{S}} p(s'|s, a) \cdot v^*(s')\right]}_{\stackrel{\text{def}}{=} q^*(s, a)}$$

- Note: the policy evaluation equations are a special case of the Bellman ones: given a policy  $\pi$ , we can consider an MDP  $\mathcal{M}^{\pi}$  in which there is a single action \* enabled in each state and the probability of transition  $s \stackrel{*}{\to} s'$  equals  $\sum_{a \in \mathcal{A}(s)} \pi(a|s) \cdot p(s'|s,a)$ . Then  $\mathcal{M}^{\pi}$  mimics the behavior of  $\pi$  in  $\mathcal{M}$  and Bellman eq's in  $\mathcal{M}^{\pi}$  evaluation equations for  $\pi$  in  $\mathcal{M}$ .
- But these equations are no longer linear! How do we solve them? Is the solution even unique?

### Bellman update operator

The right-hand-side (RHS) of the Bellman equations can be viewed as an operator  $\Phi \colon \mathbb{R}^{\mathcal{S}} \to \mathbb{R}^{\mathcal{S}}$ : for any  $\vec{x} \in \mathbb{R}^{\mathcal{S}}$ ,  $\Phi(\vec{x})$  is a vector such that for any state s:

$$\Phi(\vec{x})(s) \stackrel{\text{def}}{=} \max_{a \in \mathcal{A}(s)} \left[ r(s, a) + \gamma \cdot \sum_{s' \in \mathcal{S}} p(s'|s, a) \cdot \vec{x}(s') \right]$$

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#### Exercise 16

In our running example, compute  $\Phi(\vec{0})$ .

### Bellman update operator

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#### Exercise 16

In our running example, compute  $\Phi(\vec{0})$ .

Theorem 15 says that the optimal value vector  $v^*$  is a fixed point of  $\Phi$ :

$$v^* = \Phi(v^*).$$

#### Mathematical hammers for Bellman

#### Lemma 17: (not proven here)

For any discount factor  $\gamma \in [0,1)$ , the Bellman operator  $\Phi$  is a contraction, i.e. for any pair of vectors  $\vec{x}, \vec{y}$  it holds

$$\|\vec{x} - \vec{y}\|_{\infty} \le \gamma \cdot \|\Phi(\vec{x}) - \Phi(\vec{y})\|_{\infty}.$$

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#### Theorem 18: Banach fixed point theorem (classical calculus, not proven here)

A contraction mapping from a complete metric space (in particular,  $\mathbb{R}^n$ ) to itself has a unique fixed point.

#### **Exact characterization of** $v^*$

#### Corollary 19

The optimal value vector is a unique solution of the Bellman optimality equations.

#### Exact characterization of $v^*$

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In particular, also the policy evaluation equations have a unique solution, equal to  $v^{\pi}$ . Since the policy evaluation equations are linear, their solution can be computed by Gaussian elimination.

But the general Bellman equations are not linear. How can ve solve them?

## Banach fixpoint theorem (full)

#### Theorem 20: Banach fixed point theorem (full version, not proven here)

A contraction mapping  $\Phi$  from a complete metric space (in particular,  $\mathbb{R}^n$ ) to itself has a unique fixed point  $\vec{z}$ .

Moreover,  $\vec{z}$  is the limit of iterative applications of  $\Phi$  on any initial vector. I.e., for any  $\vec{x}_0 \in \mathbb{R}^n$ , the sequence  $\vec{x}_0, \Phi(\vec{x}_0), \Phi(\Phi(\vec{x}_0)), \Phi^{(3)}(\vec{x}_0), \ldots$  converges to  $\vec{z}$ :

$$z = \lim_{i \to \infty} \Phi^{(i)}(\vec{x}_0)$$

# Value iteration (VI; Bellman, 1957)

#### **Algorithm 1:** Value iteration

```
Input: MDP \mathcal{M} = (\mathcal{S}, \mathcal{A}, p, r)
```

**Output:** Approximation  $\tilde{v}$  of  $v^*$ 

 $x \leftarrow \text{any vector from } \mathbb{R}^{|\mathcal{S}|}$ ;

 $next \leftarrow x$ ;

#### repeat

```
foreach s \in \mathcal{S} do
next(s) \leftarrow \max_{a \in \mathcal{A}(s)} [r(s, a) + \gamma \cdot \sum_{s' \in \mathcal{S}} p(s'|s, a) \cdot \vec{x}(s')];
x \leftarrow next
```

until termination condition;

#### Typical term. conditions:

- after a fixed no. of iterations (i.e., use for-loop with a fixed bound)
- after each component of x changes less then some given  $\varepsilon$

// typically  $\vec{0}$ 

#### How to use VI

By the Banach fixpoint theorem (and Lemma 17), the value of variable x VI converges to  $v^*$ . Can we recognize when is x "close enough" to  $\vec{x}$ ?

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By the Banach fixpoint theorem (and Lemma 17), the value of variable x VI converges to  $v^*$ . Can we recognize when is x "close enough" to  $\vec{x}$ ?

In the following couple of theorems, let  $\vec{x}_0, \vec{x}_1, \vec{x}_2, \ldots$  be the sequence of vectors computed by VI, i.e.  $\vec{x}_0$  is arbitrary and  $\vec{x}_{i+1} = \Phi(\vec{x}_i)$  for all  $i \geq 0$ .

#### Theorem 21: Stopping condition (not proven here)

For any  $\varepsilon > 0$ : if

$$\|\vec{x}_{i+1} - \vec{x}_i\|_{\infty} \le \varepsilon \cdot \frac{1 - \gamma}{\gamma},$$

then

$$\|\vec{x}_{i+1} - v^*\|_{\infty} \le \varepsilon$$

## How to use VI (2)

How fast can we get to the point where we are close enough?

#### Theorem 22: Speed of convergence (not proven here)

For all  $i \ge 0$  it holds

$$\|\vec{x}^n - \mathbf{v}^*\|_{\infty} \leq \frac{\gamma^n}{1-\gamma} \cdot \|\vec{x}_1 - \vec{x}_0\|_{\infty}.$$

In particular, if we terminate VI after

$$i = \left\lceil rac{\log(arepsilon) + \log\left(rac{1-\gamma}{\|ec{arkappa_1} - ec{arkappa_0}\|_{\infty}}
ight)}{\log(\gamma)}
ight
ceil$$

steps, then its output  $x_i$  will be an  $\varepsilon$ -approximation of  $v^*$ .

# How to use VI (3)

Can we actually get some optimal values instead of approximations?

## How to use VI (3)

Can we actually get some optimal values instead of approximations? First, note that VI computes optimal finite-horizon values:

Let  $\mathbf{v}^i = \sup_{\pi} \mathbb{E}^{\pi} [\sum_{i=1}^H \gamma^{i-1} \cdot R_i]$ . The supremum is over all (i.e., history dependent) policies, since in the FH problem an optimal policy needs to track the number of elapsed (and thus remaining) steps: memory is needed for that.

### Theorem 23: (Easy but important exercise)

If 
$$\vec{x}_0 = \vec{0}$$
, then  $\vec{x}_H = v^H$  for all  $H \ge 0$ .

## How to use VI (3)

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#### Theorem 23: (Easy but important exercise)

If  $\vec{x}_0 = \vec{0}$ , then  $\vec{x}_H = v^H$  for all  $H \ge 0$ .

Moreover, let  $\pi^H$  be a deterministic history-dependent policy such that for all  $1 \le i \le H$ , whenever there are i steps remaining till the horizon, the policy  $\pi^H$  selects in state s an action a s.t.

$$a = rg \max_{a \in \mathcal{A}(s)} [r(s, a) + \gamma \cdot \sum_{s' \in \mathcal{S}} p(s'|s, a) \cdot \vec{x}_{i-1}(s')]$$

(with ties broken arbitrarily). Then  $\pi^H$  is an optimal H-step policy.

### **Greedy policies**

Can we actually get optimal policy for the inf. horizon problem?

#### Definition 24: $\vec{x}$ -greedy policy (very important)

Let  $\vec{x} \in \mathbb{R}^{S}$  be any vector. A  $\vec{x}$ -greedy policy is an MD policy  $\pi$  such that in any state s:

$$\pi(s) = \argmax_{a \in \mathcal{A}(s)} [r(s, a) + \gamma \cdot \sum_{s' \in \mathcal{S}} p(s'|s, a) \cdot \vec{x}(s')].$$

## How to use VI (4)

#### Theorem 25: Optimal inf.-horizon policy from VI (not proven here)

There is a number N polynomial in size of the MDP and exponential in the binary encoding size of  $\gamma$  such that a policy  $\pi$  that is  $\vec{x}_N$ -greedy is optimal in every state, i.e.  $v^{\pi} = v^*$ .

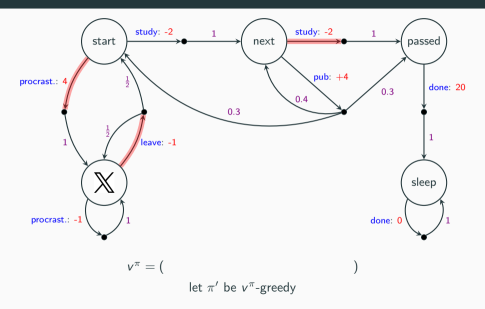
Note that once  $\pi$  is computed,  $v^{\pi}$  can be computed in polynomial time via policy evaluation equations.

Hence, VI can be said to solve MDPs in exponential time (and in polynomial time if the discount factor is assumed to be a fixed constant instead of an input parameter), though the approximate version is typically used in practice.

Note: the fact that some policy  $\pi$  is  $\vec{x}$ -greedy does not mean that  $v^{\pi} \geq \vec{x}!$  Homework: find a counterexample and post it to Discord.

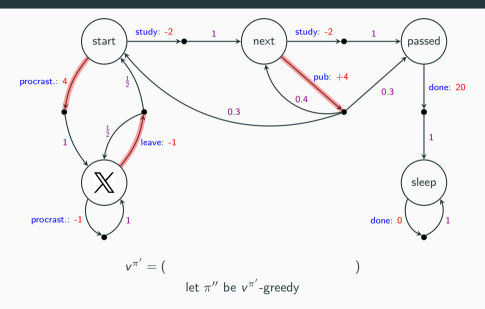
However, for VI it can be shown that if  $\|\vec{x}_{i+1} - \vec{x}_i\|_{\infty} \le \varepsilon \cdot \frac{1-\gamma}{\gamma}$  (stopping condition from Theorem 21), then an  $\vec{x}_{i+1}$ -greedy policy is  $\varepsilon$ -optimal.

# **Policy improvement**



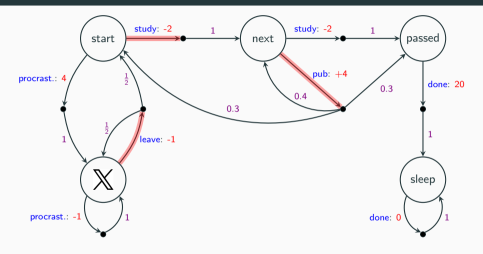
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# **Policy improvement**



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## **Policy improvement**



### Policy improvement theorem

#### Theorem 26: Policy improvement

Let  $\pi$  be a policy. If  $\Phi(v^{\pi}) \geq v^{\pi}$ , then any  $v^{\pi}$ -greedy policy  $\pi_g$  is at least as good as  $\pi$ , i.e.  $\forall s \in \mathcal{S} : v^{\pi_g}(s) \geq v^{\pi}(s)$ .

Moreover, if  $\Phi(v^{\pi})(s) > v^{\pi}(s)$  for some state s, then also  $v^{\pi_g}(s') > v^{\pi}(s')$  for some state s'.

### Returns from a given time step

For the proof of PIT and also many times later, we will need the following notation:

#### Definition 27: Important!

Let  $\tau=s_0,a_0,r_1,s_1,a_1,r_2,s_2,a_2,r_3,s_3,\ldots$  be a trajectory and  $t\in\mathbb{N}$  a time step. We define

$$G_t(\tau) = \sum_{i=t}^{H-1} \gamma^{i-t} \cdot r_{i+1} = r_{t+1} + \gamma r_{t+2} + \gamma^2 r_{t+3} + \cdots,$$

where  $H \in \mathbb{N} \cup \{\infty\}$  or H = T for episodic tasks.

We similarly define, for any policy  $\pi$ :

$$G_t^{\pi} = \mathbb{E}^{\pi}[G_t] = \mathbb{E}^{\pi}[\sum_{i=t}^{H-1} \gamma^{i-t} \cdot R_{i+1}].$$

### Proof of PIT (setup)

We will define a sequence of policies  $\pi_0, \pi_1, \pi_2, \dots$  s.t.:

- $\pi_0 = \pi$
- $\pi_i$  behaves as  $\pi_g$  (i.e., selects the same actions in same states) for the first i steps, then "switches" back to behave as  $\pi$ :

ullet we also define  $\pi_{\infty}=\pi_{oldsymbol{g}}$ 

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We want:  $v^{\pi_{\infty}}(s) \ge v^{\pi}(s)$  for all s.

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We want:  $v^{\pi_{\infty}}(s) \ge v^{\pi}(s)$  for all s.

Not hard to see:  $v^{\pi_i} \to v^{\pi_\infty}$  as  $i \to \infty$  ( $\pi_i$  behaves as  $\pi_\infty$  for longer and longer as i increases + discounting).

It suffices to show:  $v^{\pi_i}(s) \ge v^{\pi}(s)$  for all  $i \in \mathbb{N}$  and all  $s \in \mathcal{S}$ .

$$v^{\pi_i}(s) \geq v^{\pi}(s)$$
 for all  $i \in \mathbb{N}$  and all  $s \in \mathcal{S}$ 

• i = 0: clear

$$v^{\pi_i}(s) \geq v^{\pi}(s)$$
 for all  $i \in \mathbb{N}$  and all  $s \in \mathcal{S}$ 

- i = 0: clear
- i > 0:

$$v^{\pi_i}(s) = \mathbb{E}^{\pi_i}[R_1 + \gamma R_2 + \dots + \gamma^{i-1} R_i + \gamma^i R_{i+1} + \dots \mid S_0 = s]$$

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=  $\mathbb{E}^{\pi_{i}}[R_{1} + \gamma R_{2} + \dots + \gamma^{i-2}R_{i-1} \mid S_{0} = s] + \mathbb{E}^{\pi_{i}}[\gamma^{i-1}R_{i} + \gamma^{i}R_{i+1} \dots \mid S_{0} = s]$ 

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$$\begin{split} v^{\pi_{i}}(s) &= \mathbb{E}^{\pi_{i}}[R_{1} + \gamma R_{2} + \dots + \gamma^{i-1}R_{i} + \gamma^{i}R_{i+1} + \dots \mid S_{0} = s] \\ &= \mathbb{E}^{\pi_{i}}[R_{1} + \gamma R_{2} + \dots + \gamma^{i-2}R_{i-1} \mid S_{0} = s] + \mathbb{E}^{\pi_{i}}[\gamma^{i-1}R_{i} + \gamma^{i}R_{i+1} \dots \mid S_{0} = s] \\ &= \mathbb{E}^{\pi_{i-1}}[R_{1} + \gamma R_{2} + \dots + \gamma^{i-2}R_{i-1} \mid S_{0} = s] + \underbrace{\mathbb{E}^{\pi_{i}}[\gamma^{i-1}R_{i} + \gamma^{i}R_{i+1} \dots \mid S_{0} = s]}_{\text{Suppose we prove } \geq \mathbb{E}^{\pi_{i-1}}[\gamma^{i-1}R_{i} + \gamma^{i}R_{i+1} \dots \mid S_{0} = s]} \end{split}$$

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$$\geq \mathbb{E}^{\pi_{i-1}}[R_1 + \gamma R_2 + \dots + \gamma^{i-2} R_{i-1} + \gamma^{i-1} R_i + \gamma^i R_{i+1} \dots \mid S_0 = s]$$

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## Proof of PIT (induction, behavior at "reset")

We need:  $\mathbb{E}^{\pi_i}[\gamma^{i-1}R_i + \gamma^i R_{i+1} \cdots \mid S_0 = s] \ge \mathbb{E}^{\pi_{i-1}}[\gamma^{i-1}R_i + \gamma^i R_{i+1} \cdots \mid S_0 = s].$ 

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$$\mathbb{E}^{\pi_i}[\gamma^{i-1}R_i + \gamma^i R_{i+1} \cdots \mid S_0 = s] = \gamma^{i-1} \cdot \mathbb{E}^{\pi_i}[R_i + \gamma R_{i+1} \cdots \mid S_0 = s]$$

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$$\mathbb{E}^{\pi_{i}}[\gamma^{i-1}R_{i} + \gamma^{i}R_{i+1} \cdots \mid S_{0} = s] = \gamma^{i-1} \cdot \mathbb{E}^{\pi_{i}}[R_{i} + \gamma R_{i+1} \cdots \mid S_{0} = s]$$

$$= \gamma^{i-1} \cdot \sum_{s' \in S} \mathbb{P}^{\pi_{i}}[S_{i-1} = s' \mid S_{0} = s] \cdot \left(r(s', \pi_{i}(s')) + \gamma \cdot \sum_{s''} p(s'' \mid s', \pi_{i}(s')) \cdot \mathbb{E}^{\pi_{i}}[G_{i} \mid S_{i} = s'']\right)$$

We need:  $\mathbb{E}^{\pi_i}[\gamma^{i-1}R_i + \gamma^i R_{i+1} \cdots \mid S_0 = s] \ge \mathbb{E}^{\pi_{i-1}}[\gamma^{i-1}R_i + \gamma^i R_{i+1} \cdots \mid S_0 = s].$ 

$$\begin{split} & \mathbb{E}^{\pi_{i}}[\gamma^{i-1}R_{i} + \gamma^{i}R_{i+1} \cdots \mid S_{0} = s] = \gamma^{i-1} \cdot \mathbb{E}^{\pi_{i}}[R_{i} + \gamma R_{i+1} \cdots \mid S_{0} = s] \\ & = \gamma^{i-1} \cdot \sum_{s' \in \mathcal{S}} \mathbb{P}^{\pi_{i}}[S_{i-1} = s' \mid S_{0} = s] \cdot \left( r(s', \pi_{i}(s')) + \gamma \cdot \sum_{s''} p(s'' \mid s', \pi_{i}(s')) \cdot \mathbb{E}^{\pi_{i}}[G_{i} \mid S_{i} = s''] \right) \\ & = \gamma^{i-1} \cdot \sum_{s' \in \mathcal{S}} \mathbb{P}^{\pi_{g}}[S_{i-1} = s' \mid S_{0} = s] \cdot \left( r(s', \pi_{g}(s')) + \gamma \cdot \sum_{s''} p(s'' \mid s', \pi_{g}(s')) \cdot \mathbb{E}^{\pi}[G_{i} \mid S_{i} = s''] \right) \end{split}$$

We need: 
$$\mathbb{E}^{\pi_i}[\gamma^{i-1}R_i + \gamma^iR_{i+1}\cdots \mid S_0 = s] \geq \mathbb{E}^{\pi_{i-1}}[\gamma^{i-1}R_i + \gamma^iR_{i+1}\cdots \mid S_0 = s].$$

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$$= \gamma^{i-1} \cdot \sum_{s' \in \mathcal{S}} \mathbb{P}^{\pi_{i}}[S_{i-1} = s' \mid S_{0} = s] \cdot \left(r(s', \pi_{i}(s')) + \gamma \cdot \sum_{s''} p(s'' \mid s', \pi_{i}(s')) \cdot \mathbb{E}^{\pi_{i}}[G_{i} \mid S_{i} = s'']\right)$$

$$= \gamma^{i-1} \cdot \sum_{s' \in \mathcal{S}} \mathbb{P}^{\pi_{g}}[S_{i-1} = s' \mid S_{0} = s] \cdot \left(r(s', \pi_{g}(s')) + \gamma \cdot \sum_{s''} p(s'' \mid s', \pi_{g}(s')) \cdot \mathbb{E}^{\pi}[G_{i} \mid S_{i} = s'']\right)$$

We need: 
$$\mathbb{E}^{\pi_{i}}[\gamma^{i-1}R_{i} + \gamma^{i}R_{i+1} \cdots \mid S_{0} = s] \geq \mathbb{E}^{\pi_{i-1}}[\gamma^{i-1}R_{i} + \gamma^{i}R_{i+1} \cdots \mid S_{0} = s].$$

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$$=\Phi(v^{\pi})(s'), \text{ since } \pi_g \text{ is } v^{\pi}\text{-greedy}$$

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$$\geq v^{\pi}(s')$$
, since  $\Phi(v^{\pi}) \geq v^{\pi}$  by PIT assumption.

 $=\Phi(v^{\pi})(s')$ , since  $\pi_{\sigma}$  is  $v^{\pi}$ -greedy

# Policy iteration (PI; Howard, 1960)

```
Algorithm 2: Policy iteration
Input: MDP \mathcal{M} = (\mathcal{S}, \mathcal{A}, p, r)
Output: Optimal MD policy \pi^* for \mathcal{M}, its value vector v^*
\pi \leftarrow \text{arbitrary MD policy};
v \leftarrow v^{\pi}:
                               // e.g. by solving linear policy evaluation equations
while \Phi(v) \neq v do
    foreach s \in \mathcal{S} do
      \pi(s) \leftarrow \arg\max_{a \in A(s)} [r(s, a) + \gamma \cdot \sum_{s' \in S} p(s'|s, a) \cdot v(s')]
  v \leftarrow v^{\pi}
return \pi, \nu
```

#### Theorem 28

Policy iteration terminates after at most exponentially many iterations. Upon termination, it returns an optimal MD policy.

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• Optimal upon termination:  $v^{\pi}$  is a fixpoint of  $\Phi$  when terminating: optimality follows from Corollary 19.

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- Optimal upon termination:  $v^{\pi}$  is a fixpoint of  $\Phi$  when terminating: optimality follows from Corollary 19.
- Terminates:  $\pi$  always stores an MD policy and there are finitely many of these. We will show that no single MD policy appears in more than one iteration of PI. Consider any iteration and let v, v' be the contents of variable v before and after the iteration. We will show that unless  $\Phi(v) = v$ , it holds v' > v, i.e.  $v' \ge v$  componentwise with strict inequality in some component. Hence,  $v = v^{\pi}$  strictly increases during PI, so no  $\pi$  can appear twice.

 $v' \geq v$ :

### $v' \geq v$ :

We verify assumptions of PIT:  $\Phi(v) \ge v$ . Recall  $v = v^{\pi}$ . For all  $s \in \mathcal{S}$ :

$$\Phi(v)(s) = \max_{a \in \mathcal{A}(s)} [r(s, a) + \gamma \cdot \sum_{s' \in \mathcal{S}} p(s' \mid s, a) \cdot v(s')]$$

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$$\geq r(s, \pi(s)) + \gamma \cdot \sum_{s' \in \mathcal{S}} p(s' \mid s, \pi(s)) \cdot v^{\pi}(s')$$

$$= v^{\pi}(s) = v(s).$$

By PIT,  $v'=v^{\pi'}\geq v^{\pi}=v$  (here  $\pi'$  is the v-greedy policy).

It remains to prove that v' > v or PI terminates. Assume that v' = v. Then for all  $s \in \mathcal{S}$ :

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so PI terminates at this point.

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so PI terminates at this point. Complexity?

### PI&VI: discussion

- We know that MDPs have a linear programming (LP) formulation. PI is basically a variant of a simplex method for this LP, using a special pivoting rule.
- PI typicaly requires less iterations to converge than VI, though each iteration is more expensive (policy eval.)
- Both PI and VI typically work well in practice for MDPs whose explicit transition table fits inside a computer. Which of the two is faster is rather domain-specific.

### PI variants

Can we get rid of the expensive policy evaluation by linear system solving?

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Can we get rid of the expensive policy evaluation by linear system solving?

Yes: we can approximate the value of the current policy  $\pi$  by applying VI on the MDP  $\mathcal{M}^{\pi}$ , for either fixed number of steps or until v does not change much. Often appearing in RL textbooks:

# Policy iteration with approximate evaluation

return v'

```
\pi \leftarrow \text{arbitrary MD policy}; \nu \leftarrow \text{arbitrary vector};
repeat
      v \leftarrow \text{Eval}(\pi, v);
      foreach s \in S do
       \pi(s) \leftarrow \operatorname{arg\,max}_{a \in \mathcal{A}(s)}[r(s, a) + \gamma \cdot \sum_{s' \in \mathcal{S}} p(s'|s, a) \cdot v(s')]
until \pi has not changed;
return \pi, \nu
Function Eval(\pi, v):
      v' \leftarrow v:
      repeat
             foreach s \in S do
                \begin{array}{c} v(s) \leftarrow v'(s); \\ v'(s) \leftarrow r(s, \pi(s)) + \gamma \cdot \sum_{s' \in \mathcal{S}} p(s' \mid s, \pi'(s)) \cdot v(s') \end{array}
      until ||v - v'||_{\infty} \le \varepsilon;
```

### **Convergence of PI variants**

- The algorithm on previous slide still converges to an optimal policy provided that  $\varepsilon$  is small enough.
- If we replaced the " $\pi$  not changed condition" with the original " $\Phi(v) = v$ " condition, the algorithm might not terminate, since the VI is only guaranteed to reach a true fixpoint in the limit. However, v would still converge to  $v^*$  and thus  $\pi$  would eventually become equal to an optimal policy.
- The previous point holds even in the very degenerate case when we do just one iteration of VI per policy evaluation! See next slide.

### **Curiously looking approximate PI**

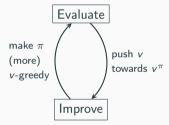
```
v \leftarrow arbitrary vector;
\pi \leftarrow v-greedy MD policy :
repeat
      foreach s \in \mathcal{S} do
       v'(s) \leftarrow r(s, \pi(s)) + \gamma \cdot \sum_{s' \in \mathcal{S}} p(s'|s, \pi(s)) \cdot v(s');
      v \leftarrow v'
      foreach s \in \mathcal{S} do
        \pi(s) \leftarrow \arg\max_{a \in \mathcal{A}(s)} [r(s, a) + \gamma \cdot \sum_{s' \in \mathcal{S}} p(s'|s, a) \cdot v(s')]
until \Phi(v) = v:
return \pi, \nu
```

### **Curiously looking approximate PI**

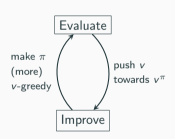
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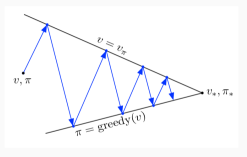
This is just VI in disguise!

### Generalized policy iteration



# **Generalized policy iteration**





Source: Sutton&Barto, p. 87

for Model-Free

**Reinforcement Learning** 

**Tabular Methods** 

### Model-free

#### Model-free

We will still be working with MDPs. But for a bunch of the following lectures, we will not (necessarily) have access to, e.g.:

- a table containing explicit enumeration of all states/actions
- a table containing the description of p or r
- the ability to compute the probability vector  $\delta(s, a)$  or the reward signal r(s, a) given s and a (having this = gray-box model of the MDP)

- know how the states of the MDP look like
  - (e.g. robot state = all possible output values of its sensors)

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  - could be weakened, but simplifies things

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- ullet can, for any  $s\in\mathcal{S}$ , enumerate  $\mathcal{A}(s)$ 
  - could be weakened, but simplifies things
- given  $s \in \mathcal{S}$  and  $a \in \mathcal{A}(s)$ , we can sample the next state  $s' \sim p(s, a)$  and receive the reward r(s, a).

# Sampling from a policy

Given an effective representation of a policy  $\pi$ , we can sample a trajectory  $s_0, a_0, r_1, s_1, a_1, r_2, s_2, a_2, r_3, s_3, \dots$  by performing, for each  $t \in \{0, \dots, T\}$ :

- sample  $a_t \sim \pi(s_t)$
- ullet query the environment for  $s_{t+1} \sim p(s_t, a_t)$  and  $r_{t+1} = r(s_t, a_t)$
- increment t

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- increment t

Tabular = value estimates and policies represented as tables (e.g. Q(s, a) for each state s and action a used in s – explicit representation might only be needed for states/actions actually encountered).

# Basic classification of (tabular) RL algorithms

Three independent axes:

problem	on/off	updates
policy evaluation (value prediction)	on-policy	Monte Carlo
VS.	VS.	
control	off-policy	temporal difference

### Assumptions: successor-dependent rewards & episodic tasks

Since we do no longer have the knowledge of the transition dynamics p, we cannot freely interchange MDPs with rewards functions of type  $\mathcal{S} \times \mathcal{A} \times \mathcal{S} \to \mathbb{R}$  and  $\mathcal{S} \times \mathcal{A} \to \mathbb{R}$  via the equation  $r(s,a) = \sum_{s' \in \mathcal{S}} p(s' \mid s,a) \cdot r(s,a,s')$ .

Hence, to maintain generality (and correspondence to e.g. Gymnasium environments) we will assume reward functions of type  $\mathcal{S} \times \mathcal{A} \times \mathcal{S} \to \mathbb{R}$ .

We will assume episodic returns: each trajectory terminates with probability 1 at some (possibly random) time step T. Termination can be defined e.g. by reaching some terminal state or by running out of some fixed decision horizon (in Gymnasium, this is sometimes called truncation):

$$G = \sum_{i=0}^{T-1} \gamma^i \cdot R_{i+1}.$$

Episode = one high-level iteration of an RL algorithm, corresponding of sampling a single trajectory from some policy.

Monte Carlo Methods

#### MC evaluation

Policy evaluation: given an effective representation of a policy  $\pi$ , estimate  $v^{\pi}$  (or  $q^{\pi}$ ).

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Naive Monte Carlo: Sample from  $\pi$ : if  $\{\tau_1, \tau_2, \dots, \tau_n\}$  are trajectories (episodes) independently sampled under  $\pi$  from the same initial state s, then  $\frac{1}{n} \sum_{i=1}^n G(\tau_i) \to v^{\pi}(s)$  as  $n \to \infty$  due to law of large numbers (LLN).

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But this throws away a lot of valuable information! E.g. what if we want to estimate the whole  $v^{\pi}$ ?

#### First-visit MC

For each s, we estimate  $v^{\pi}(s)$  as an average of sample returns Ret(s) which is formed as follows:

- initially,  $Ret(s) = \emptyset$  for all s
- we then sample trajectories until timeout:
  - for each sampled trajectory  $\tau$  and each state s, we identify the first occurrence of s on  $\tau$ : let this be at timestep t; we add  $G_t(\tau)$  to Ret(s)



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Sub-trajectory starting at the first appearance of s can be seen as a trajectory sampled from  $\pi$  when s is the initial state! (Since we consider memoryless  $\pi$ .)

#### Theorem 29

As  $|Ret(s)| \to \infty$ , the average of Ret(s) converges to  $v^{\pi}(s)$ . Moreover, the average of Ret(s) is an unbiased estimate of  $v^{\pi}(s)$  (as long as  $Ret(s) \neq \emptyset$ ).

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### First-visit MC (pseudocode)

#### First-visit MC prediction, for estimating $V \approx v_{\pi}$

```
Input: a policy \pi to be evaluated
Initialize:
     V(s) \in \mathbb{R}, arbitrarily, for all s \in \mathcal{S}
     Returns(s) \leftarrow \text{an empty list, for all } s \in S
Loop forever (for each episode):
    Generate an episode following \pi: S_0, A_0, R_1, S_1, A_1, R_2, \ldots, S_{T-1}, A_{T-1}, R_T
    G \leftarrow 0
     Loop for each step of episode, t = T - 1, T - 2, \dots 0:
         G \leftarrow \gamma G + R_{t+1}
         Unless S_t appears in S_0, S_1, \ldots, S_{t-1}:
               Append G to Returns(S_t)
               V(S_t) \leftarrow \text{average}(Returns(S_t))
```

Source: Sutton&Barto, p.92

#### **Every-visit MC**

For each s, we estimate  $v^{\pi}(s)$  as an average of sample returns Ret(s) which is formed as follows:

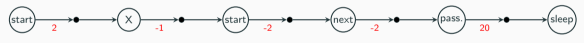
- initially,  $Ret(s) = \emptyset$  for all s
- we then sample trajectories until timeout:
  - for each sampled trajectory  $\tau$  and each state s, and each t such that  $S_t(\tau) = s$  we add  $G_t(\tau)$  to Ret(s)



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  - for each sampled trajectory  $\tau$  and each state s, and each t such that  $S_t(\tau) = s$  we add  $G_t(\tau)$  to Ret(s)



The sample returns added to Ret(s) within the same episode are not independent! Hence, the estimate is biased, though the bias vanishes in the limit:

#### Theorem 30

As  $|Ret(s)| \to \infty$ , the average of Ret(s) converges to  $v^{\pi}(s)$ .

#### MC convergence

Optional reading: More on MC estimate bias, variance, and convergence in:

Singh, S.P. and Sutton, R.S.: Reinforcement Learning with Replacing Eligibility Traces. In *Machine Learning* 22:123–158. Kluwer, 1996.

(Section 3, particularly 3.3 and onwards, you can skip Theorem 4.)

#### **Towards MC control**

Control = computation of "good" policy for a given environment. (Ideally, the policy should get closer to the optimal policy the more episodes we sample.)

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We know (PIT): given a policy  $\pi$  a  $\nu^{\pi}$ -greedy policy is at least as good as  $\pi$ :

$$\pi_g(s) = \argmax_{a \in \mathcal{A}(s)} \left[ \sum_{s' \in \mathcal{S}} p(s' \mid s, a) \cdot \left( r(s, a, s') + \gamma \cdot v^{\pi}(s') \right) \right]$$

Do we have an algo? There is an issue:

### MC control with q-values

Recall:

$$q^{\pi}(s, a) \stackrel{\mathsf{def}}{=} \sum_{s' \in \mathcal{S}} p(s' \mid s, a) \cdot (r(s, a, s') + \gamma \cdot v^{\pi}(s').)$$

Thus, the  $v^{\pi}$ -greedy policy  $\pi_g$  can be defined as:

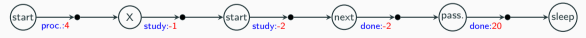
$$\pi_g(s) = \underset{a \in \mathcal{A}(s)}{\operatorname{arg max}} \underbrace{q^{\pi}(s, a)}_{\text{Estimate by MC}}$$

#### MC for q-value estimation

Analogous to value estimation, e.g. first-visit:

For each s, a, we estimate  $q^{\pi}(s, a)$  as an average of sample returns Ret(s, a) which is formed as follows:

- initially,  $Ret(s, a) = \emptyset$  for all s
- we then sample trajectories until timeout:
  - for each sampled trajectory  $\tau$  and each state-action pair (s, a), we identify the first t such that  $S_t(\tau) = s \wedge A_t(\tau) = a$ ; we add  $G_t(\tau)$  to Ret(s)



Similarly for every visit. Convergence guarantees the same as for state values.

# Infinite exploration and exploring starts

Issue: MC only estimates  $q^{\pi}(s, a)$  if:

- ullet s guaranteed to be visited with positive probability in each episode
- $\pi(a \mid s) > 0$ .

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#### Definition 31: Infinite exploration

A RL algorithm has infinite exploration (IE) if, during the infinite execution of the algorithm, each state-action pair (s, a) is visited infinitely often with probability 1.

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- $\pi(a \mid s) > 0$ .

#### Definition 31: Infinite exploration

A RL algorithm has infinite exploration (IE) if, during the infinite execution of the algorithm, each state-action pair (s, a) is visited infinitely often with probability 1.

One way of achieving IE is through exploring starts (ES): each episode begins with (typically uniformly) randomly selected  $s_0$  and  $a_0$ . This is achievable when training, e.g., in simulated environments but might be difficult/impossible in real-world environments.

### MC control with exploring starts

#### Monte Carlo ES (Exploring Starts), for estimating $\pi \approx \pi_*$

```
Initialize:
     \pi(s) \in \mathcal{A}(s) (arbitrarily), for all s \in \mathcal{S}
     Q(s, a) \in \mathbb{R} (arbitrarily), for all s \in S, a \in \mathcal{A}(s)
     Returns(s, a) \leftarrow \text{empty list, for all } s \in S, \ a \in \mathcal{A}(s)
Loop forever (for each episode):
     Choose S_0 \in \mathcal{S}, A_0 \in \mathcal{A}(S_0) randomly such that all pairs have probability > 0
     Generate an episode from S_0, A_0, following \pi: S_0, A_0, R_1, \ldots, S_{T-1}, A_{T-1}, R_T
     G \leftarrow 0
     Loop for each step of episode, t = T - 1, T - 2, \dots, 0:
          G \leftarrow \gamma G + R_{t+1}
           Unless the pair S_t, A_t appears in S_0, A_0, S_1, A_1, ..., S_{t-1}, A_{t-1}:
                Append G to Returns(S_t, A_t)
                Q(S_t, A_t) \leftarrow \operatorname{average}(Returns(S_t, A_t))
                \pi(S_t) \leftarrow \operatorname{arg\,max}_{a} Q(S_t, a)
```

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### **IE** through $\varepsilon$ -soft policies

Exploring starts are not always feasible. Alternative: make the sampled policy itself exploratory.

Definition 32:  $\varepsilon$ -soft policy

A policy  $\pi$  is  $\varepsilon$ -soft if for every  $s \in \mathcal{S}$  and every  $a \in \mathcal{A}(s)$  it holds  $\pi(a|s) \geq \frac{\varepsilon}{|\mathcal{A}(s)|}$ .

# IE through $\varepsilon$ -soft policies

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#### Definition 32: $\varepsilon$ -soft policy

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#### Definition 33: $\varepsilon$ -greedy policy

Let  $v \in \mathbb{R}^{\mathcal{S}}$  be a value vector. A policy  $\pi$  is  $v - \varepsilon$ -greedy if for every state  $s \in \mathcal{S}$  there is action  $a^* = \arg\max_{a \in \mathcal{A}(s)} \sum_{s' \in \mathcal{S}} p(s' \mid s, a) \cdot (r(s, a, s') + \gamma \cdot v(s'))$  such that for any action  $a \in \mathcal{A}(s)$  it holds:

$$\pi(a|s) = egin{cases} rac{arepsilon}{\mathcal{A}(s)} & ext{if } a 
eq a^* \ 1 - arepsilon + rac{arepsilon}{\mathcal{A}(s)} & ext{if } a = a^*. \end{cases}$$

Interpretation: with prob.  $\varepsilon$ : play uniformly at random; with prob.  $1-\varepsilon$ : play greedily.

#### $\varepsilon$ -softing

#### **Definition 34**

Let  $\pi$  be a policy. An  $\varepsilon$ -softing of  $\pi$  is a policy  $\pi_{\varepsilon}$  defined as follows: in each state s

- with probability  $\varepsilon$ ,  $\pi_{\varepsilon}$  selects an action uniformly at random;
- with probability  $1 \varepsilon$ ,  $\pi_{\varepsilon}$  selects  $a \sim \pi(s)$ .

I.e., an  $\varepsilon$ -greedy policy can be alternatively defined as  $\varepsilon$ -softing of a greedy policy.

# MC control with $\varepsilon$ -greedy policies

```
Algorithm parameter: small \varepsilon > 0
Initialize:
    \pi \leftarrow \text{an arbitrary } \varepsilon\text{-soft policy}
    Q(s,a) \in \mathbb{R} (arbitrarily), for all s \in S, a \in A(s)
    Returns(s, a) \leftarrow \text{emptv list, for all } s \in S, a \in A(s)
Repeat forever (for each episode):
    Generate an episode following \pi: S_0, A_0, R_1, \ldots, S_{T-1}, A_{T-1}, R_T
    G \leftarrow 0
    Loop for each step of episode, t = T-1, T-2, \ldots, 0:
         G \leftarrow \gamma G + R_{t+1}
         Unless the pair S_t, A_t appears in S_0, A_0, S_1, A_1, ..., S_{t-1}, A_{t-1}:
              Append G to Returns(S_t, A_t)
              Q(S_t, A_t) \leftarrow \text{average}(Returns(S_t, A_t))
              A^* \leftarrow \operatorname{arg\,max}_a Q(S_t, a)
                                                                                     (with ties broken arbitrarily)
              For all a \in \mathcal{A}(S_t):
                       \pi(a|S_t) \leftarrow \begin{cases} 1 - \varepsilon + \varepsilon/|\mathcal{A}(S_t)| & \text{if } a = A^* \\ \varepsilon/|\mathcal{A}(S_t)| & \text{if } a \neq A^* \end{cases}
```

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### Policy iteration for $\varepsilon$ -soft policies

#### Theorem 35

Let  $\pi$  be an  $\varepsilon$ -soft policy and let  $\pi'$  be a  $v^{\pi}$ - $\varepsilon$ -greedy policy. Than  $v^{\pi'} \geq v^{\pi}$  (componentwise). Moreover, the two value vectors are equal if and only if bot  $\pi$  and  $\pi'$  are optimal among all  $\varepsilon$ -soft policies; i.e. if, for every state s:

$$v^{\pi}(s) = \sup_{ar{\pi} \text{ that is } \epsilon\text{-soft}} v^{ar{\pi}}(s).$$

Proof: Required reading: Sutton&Barto, p.101-103.

### Incremental computing of averages

Given a sample  $\{n_1, n_2, \dots, n_{k+1}\}$  and average  $A = avg(\{n_1, n_2, \dots, n_k\})$ , how to compute  $A' = avg(\{n_1, n_2, \dots, n_k, n_{k+1}\})$  without recomputing the average of the whole sample?

$$A' =$$

### Incremental computing of averages

Given a sample  $\{n_1, n_2, \ldots, n_{k+1}\}$  and average  $A = avg(\{n_1, n_2, \ldots, n_k\})$ , how to compute  $A' = avg(\{n_1, n_2, \ldots, n_k, n_{k+1}\})$  without recomputing the average of the whole sample?

$$A' = \frac{k}{k+1} \cdot A + \frac{n_{k+1}}{k+1}.$$

### On-policy vs. off-policy

- On-policy algorithms: track one "policy variable"  $\pi$ ; the policy stored in  $\pi$  is used to interact with the environment (i.e., to sample episodes) and at the same time we learn something about it (e.g. its value vector).
  - Corresponds to the generalized policy iteration scheme.
  - All the MC algos we have seen so far.

## On-policy vs. off-policy

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  - Corresponds to the generalized policy iteration scheme.
  - All the MC algos we have seen so far.
- Off-policy algorithms: track more (typically two) different policy variables:
  - behavior policy: used to sample episodes
  - target policy: which we want to learn about

## Off-policy evaluation

We are given effective representations of:

- a behavior policy  $\beta$ ,
- a target policy  $\pi$ .

The task is to estimate  $v^{\pi}$  by sampling episodes from  $\beta$ . We cannot sample from  $\pi!$  (E.g.  $\pi$  too risky or expensive to sample from.)

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- a behavior policy  $\beta$ ,
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The task is to estimate  $v^{\pi}$  by sampling episodes from  $\beta$ . We cannot sample from  $\pi!$  (E.g.  $\pi$  too risky or expensive to sample from.)

#### Assumptions:

- given (s, a), we can effectively compute  $\pi(a|s)$  and  $\beta(a|s)$  (or at least estimate via sampling)
- coverage:  $\forall s \in \mathcal{S}, a \in \mathcal{A}(s)$ : if  $\pi(a|s) > 0$ , then also  $\beta(a|s) > 0$

## Importance sampling

#### Definition 36: Importance ratio

Let  $\tau=s_0, a_0, r_1, s_1, a_1, r_2, s_2, a_2, r_3, s_3, \ldots$  be a trajectory. The importance-sampling ratio of  $\tau$  is the quantity

$$\rho(\tau) \stackrel{\text{def}}{=} \frac{\mathbb{P}^{\pi}[\tau \mid S_0 = s_0]}{\mathbb{P}^{\beta}[\tau \mid S_0 = s_0]} 
= \frac{\mathbb{P}^{\pi}[A_0 = a_0, S_1 = s_1, A_1 = a_1, \dots, A_{T-1} = a_{T-1}, S_T = s_T \mid S_0 = s_0]}{\mathbb{P}^{\beta}[A_0 = a_0, S_1 = s_1, A_1 = a_1, \dots, A_{T-1} = a_{T-1}, S_T = s_T \mid S_0 = s_0]}.$$

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$$\rho(\tau) \stackrel{\text{def}}{=} \frac{\mathbb{P}^{\pi}[\tau \mid S_0 = s_0]}{\mathbb{P}^{\beta}[\tau \mid S_0 = s_0]} \\
= \frac{\mathbb{P}^{\pi}[A_0 = a_0, S_1 = s_1, A_1 = a_1, \dots, A_{T-1} = a_{T-1}, S_T = s_T \mid S_0 = s_0]}{\mathbb{P}^{\beta}[A_0 = a_0, S_1 = s_1, A_1 = a_1, \dots, A_{T-1} = a_{T-1}, S_T = s_T \mid S_0 = s_0]}.$$

 $\rho(\tau)$  can be computed without the knowledge of MDP transition probabilities!

$$\rho(\tau) = \frac{\pi(a_0 \mid s_0) \cdot p(s_1 \mid s_0, a_0) \cdot \pi(a_1 \mid s_1) \cdot p(s_2 \mid s_1, a_1) \cdots}{\beta(a_0 \mid s_0) \cdot p(s_1 \mid s_0, a_0) \cdot \beta(a_1 \mid s_1) \cdot p(s_2 \mid s_1, a_1) \cdots}$$

## Importance ratio from time t

#### **Definition 37**

Let  $\tau=s_0, a_0, r_1, s_1, a_1, r_2, s_2, a_2, r_3, s_3, \ldots$  be a trajectory. By  $\tau_{i..j}$  we denote the subtrajectory of  $\tau$  starting in time step i and ending in timestep j. By  $\tau_{i..}$  we denote the suffix of  $s_i, a_i, r_{i+1}, s_{i+1}, a_{i+1}, \ldots$ 

## Importance ratio from time t

#### **Definition 37**

Let  $\tau=s_0, a_0, r_1, s_1, a_1, r_2, s_2, a_2, r_3, s_3, \ldots$  be a trajectory. By  $\tau_{i..j}$  we denote the subtrajectory of  $\tau$  starting in time step i and ending in timestep j. By  $\tau_{i..}$  we denote the suffix of  $s_i, a_i, r_{i+1}, s_{i+1}, a_{i+1}, \ldots$ 

#### Definition 38: Importance ratio from time *t*

Let  $\tau = s_0, a_0, r_1, s_1, a_1, r_2, s_2, a_2, r_3, s_3, \dots$  be a trajectory and t a time step. The importance-sampling ratio of  $\tau$  from t is the quantity

$$\rho_{t}(\tau) \stackrel{def}{=} \frac{\mathbb{P}^{\pi}[\tau_{t..} \mid S_{0} = s_{t}]}{\mathbb{P}^{\beta}[\tau_{t..} \mid S_{0} = s_{t}]} \\
= \frac{\mathbb{P}^{\pi}[A_{0} = a_{t}, S_{1} = s_{t+1}, A_{1} = a_{t+1}, \dots, A_{T-1-t} = a_{T-1}, S_{T-t} = s_{T} \mid S_{0} = s_{t}]}{\mathbb{P}^{\beta}[A_{0} = a_{t}, S_{1} = s_{t+1}, A_{1} = a_{t+1}, \dots, A_{T-1-t} = a_{T-1}, S_{T-t} = s_{T} \mid S_{0} = s_{t}]}.$$

#### Theorem 39

For any  $s \in \mathcal{S}$  it holds:

$$\mathbb{E}^{\beta}[\rho\cdot G\mid S_0=s]=v^{\pi}(s).$$

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Proof:

$$\mathbb{E}^{\beta}[\rho \cdot G \mid S_0 = s] = \sum_{\tau} \mathbb{P}^{\beta}[\tau \mid S_0 = s] \cdot \rho(\tau) \cdot G(\tau)$$

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Proof:

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$$= \sum_{\tau} \mathbb{P}^{\beta}[\tau \mid S_0 = s] \cdot \frac{\mathbb{P}^{\pi}[\tau \mid S_0 = s]}{\mathbb{P}^{\beta}[\tau \mid S_0 = s]} \cdot G(\tau)$$

#### Theorem 39

For any  $s \in \mathcal{S}$  it holds:

$$\mathbb{E}^{\beta}[\rho\cdot G\mid S_0=s]=v^{\pi}(s).$$

Proof:

$$\begin{split} \mathbb{E}^{\beta}[\rho \cdot G \mid S_0 = s] &= \sum_{\tau} \mathbb{P}^{\beta}[\tau \mid S_0 = s] \cdot \rho(\tau) \cdot G(\tau) \\ &= \sum_{\tau} \mathbb{P}^{\beta}[\tau \mid S_0 = s] \cdot \frac{\mathbb{P}^{\pi}[\tau \mid S_0 = s]}{\mathbb{P}^{\beta}[\tau \mid S_0 = s]} \cdot G(\tau) \\ &= \sum_{\tau} \mathbb{P}^{\pi}[\tau \mid S_0 = s] \cdot G(\tau) = \mathbb{E}^{\pi}[G \mid S_0 = s] = v^{\pi}(s). \end{split}$$

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Proof:

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Easily integrates into both first-visit and every visit MC: sample from  $\beta$  and store  $\rho_t(\tau) \cdot G_t(\tau)$  in  $Ret(s_t)$ .

## Weighted importance sampling

First-visit variant: for each state s, we keep a set of samples Sam(s). Each sample is a tuple  $(\tau, t)$  – trajectory and time step.

- initially,  $Sam(s) = \emptyset$  for all s
- we then sample trajectories until timeout:
  - for each sampled trajectory au and each state s, and the smallest t such that  $S_t( au) = s$  we add ( au, t) to Sam(s)

Throughout the algorithm, the value of state s is estimated as

$$\textit{WIS}(s) = rac{\displaystyle\sum_{( au,t) \in \textit{Sam}(s)} 
ho_t( au) \cdot \textit{G}_t( au)}{\displaystyle\sum_{( au,t) \in \textit{Sam}(s)} 
ho_t( au)}$$

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ho_t( au) \cdot G_t( au)}{\displaystyle\sum_{( au,t) \in Sam(s)} 
ho_t( au)}$$

#### Exercise 40

Compare ordinary/weighted importance sampling after single sample.

## Weighted importance sampling – correctness

The weighted sampling is clearly a biased estimator. However, the bias vanishes in the limit:

#### Theorem 41

With probability 1: as  $|Sam(s)| \to \infty$ , we have that  $WIS(s) \to v^{\pi}(s)$ .

Proof:

# Ordinary vs. weighted sampling

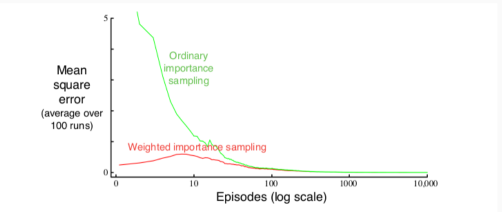


Figure 5.3: Weighted importance sampling produces lower error estimates of the value of a single blackjack state from off-policy episodes.  $\blacksquare$ 

source: Sutton&Barto, p. 106

## Importance sampling: summary

But ordinary and weighted importance sampling can be adapted to every-visit MC.

#### Bias & Convergence:

- First visit:
  - ordinary IS: unbiased, i.e. also converges
  - weighted IS: biased, but converges in the limit
- Every visit:
  - both ordinary and weighted: biased (due to EV), but converges in the limit

## Weighted IS: incremental implementation

Instead of recomputing the weighted average for each new sample, WIS(s) can be updated by keeping keep just two variables:

- V current value of WIS(s), initially arbitrary
- C the sum of importance ratios, initially 0

Upon arrival of new sample  $(\tau', t')$ , we update V, C into new values V', C' by setting:

$$C' = C + \rho_{t'}(\tau')$$

$$V' = V + \frac{\rho_{t'}(\tau')}{C'} \cdot (G_{t'}(\tau') - V).$$

## Off-policy evaluation with weighted IS

#### Off-policy MC prediction (policy evaluation) for estimating $Q \approx q_{\pi}$

```
Input: an arbitrary target policy \pi
Initialize, for all s \in \mathcal{S}, a \in \mathcal{A}(s):
     Q(s,a) \in \mathbb{R} (arbitrarily)
     C(s,a) \leftarrow 0
Loop forever (for each episode):
     b \leftarrow any policy with coverage of \pi
     Generate an episode following b: S_0, A_0, R_1, \ldots, S_{T-1}, A_{T-1}, R_T
     G \leftarrow 0
     W \leftarrow 1
     Loop for each step of episode, t = T - 1, T - 2, \dots, 0, while W \neq 0:
          G \leftarrow \gamma G + R_{t+1}
          C(S_t, A_t) \leftarrow C(S_t, A_t) + W
          Q(S_t, A_t) \leftarrow Q(S_t, A_t) + \frac{W}{C(S_t, A_t)} [G - Q(S_t, A_t)]
          W \leftarrow W \frac{\pi(A_t|S_t)}{h(A_t|S_t)}
```

# Off-policy control with weighted IS

Required reading: Sutton&Barto, Section 5.7.

**Temporal Difference Methods** 

## **TD:** Motivation

Let us first focus on policy evaluation.

MC: zero bias (at least in the limit), but potentially high variance: many samples needed to converge. Also, to update estimates, it must wait till the end of each episode.

TD methods retain the focus on sampling but combine it with bootstrapping.

## Incremental update notation

### Definition 42: Notation for updates

In the context of RL algorithms will denote by  $V^n(s)$  (resp.  $Q^n(s,a)$ ) the algorithm's estimate of  $v^{\pi}(s)$  (resp.  $q^{\pi}(s,a)$ ) after n-th update of this estimate.

## MC vs. TD(0) update

On-policy MC (incremental) update using sampled trajectory

$$\tau = s_0, a_0, r_1, s_1, a_1, r_2, s_2, a_2, r_3, s_3, \ldots$$

$$V^{n+1}(s_t) \leftarrow (1 - \alpha_n)V^n(s_t) + \alpha_n G_t(\tau) = V^n(s_t) + \alpha_n \cdot \left[\underbrace{G_t(\tau) - V^n(s_t)}_{\text{update target}}\right],$$

where  $\alpha_n = n/(n+1)$ .

## MC vs. TD(0) update

On-policy MC (incremental) update using sampled trajectory

$$\tau = s_0, a_0, r_1, s_1, a_1, r_2, s_2, a_2, r_3, s_3, \ldots$$

$$V^{n+1}(s_t) \leftarrow (1 - \alpha_n)V^n(s_t) + \alpha_n G_t(\tau) = V^n(s_t) + \alpha_n \cdot \left[\underbrace{G_t(\tau) - V^n(s_t)}_{\text{update error}}\right],$$

where  $\alpha_n = n/(n+1)$ .

TD(0) update in the same situation, with  $\alpha_n$  "suitably chosen" (possibly constant):

$$V^{n+1}(s_t) \leftarrow V^n(s_t) + \alpha_n \cdot \left[ R_{t+1}(\tau) + \gamma \cdot \underbrace{V^n(S_{t+1}(\tau))}_{\text{bootstrap}} - V^n(s_t) \right]$$

## Policy evaluation with TD(0)

## Tabular TD(0) for estimating $v_{\pi}$

```
Input: the policy \pi to be evaluated Algorithm parameter: step size \alpha \in (0,1] Initialize V(s), for all s \in \mathbb{S}^+, arbitrarily except that V(terminal) = 0 Loop for each episode:

Initialize S
Loop for each step of episode:
A \leftarrow \text{action given by } \pi \text{ for } S
Take action A, observe R, S'
V(S) \leftarrow V(S) + \alpha [R + \gamma V(S') - V(S)]
S \leftarrow S'
until S is terminal
```

source: Sutton&Barto, p. 120

### How can it even work?

Really "just" a very asynchronous, sample-based, and " $\alpha$ -dampened" version of value iteration.

$$\mathbb{E}^{\pi}[G_{t}|S_{t}=s] = \mathbb{E}^{\pi}[R_{t+1} + \gamma \cdot G_{t+1} \mid S_{t}=s] = \mathbb{E}^{\pi}[R_{t+1} \mid S_{t}=s] + \gamma \cdot \underbrace{\mathbb{E}^{\pi}[G_{t+1} \mid S_{t}=s]}_{\nu^{\pi}(S_{t+1})}.$$

In expectation, the TD(0) update is the same as VI update in  $\mathcal{M}^{\pi}$ . Thanks to the contractivity of the Bellman operator, VI possesses an error reduction property: after each update, the error of the estimate decreases. Hence, in expectation, the same is true for the TD(0) update.

Formal proof of correctness in optional reading:

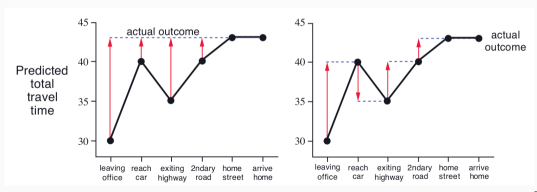
Sutton, R.S.: Learning to Predict by Methods of Temporal Differences. In *Machine Learning* 3:9–44. Kluwer, 1988. (For MDPs with function approximation.)

# Why TD is natural (Sutton&Barto, p. 122-123)

State	Elapsed Time (minutes)	Predicted Time to Go	Predicted Total Time
leaving office, friday at 6	(minutes)	30	30
reach car, raining	5	35	40
exiting highway	20	15	35
2ndary road, behind truck	30	10	40
entering home street	40	3	43
arrive home	43	0	43

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State	Elapsed Time $(minutes)$	Predicted Time to Go	Predicted Total Time
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reach car, raining	5	35	40
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entering home street	40	3	43
arrive home	43	0	43



Left: MC. Right TD(0).

## **On-policy TD control**

#### Recall:

- In control setting, we need to estimate q-values of a policy.
- On-policy: we sample trajectories according to some policy  $\pi$  and then push value estimates towards  $q^{\pi}$ .

To maintain exploration, the policy  $\pi$  will typically be the  $\varepsilon$ -Q-greedy policy for some  $\varepsilon > 0$ , where Q are the current Q values estimates. I.e., throughout the algorithm

$$\pi(a|s) = \begin{cases} 1 - \varepsilon + \frac{\varepsilon}{\mathcal{A}(s)} & \text{if } a = \arg\max_{a' \in \mathcal{A}(s)} Q(s, a') \\ \frac{\varepsilon}{\mathcal{A}(s)} & \text{otherwise.} \end{cases}$$

#### **SARSA**

State-Action-Reward-State-Action. Introduced in Rummery, Niranjan: *On-Line Q-Learning Using Connectionist Systems* (1994).

In each episode, sample a trajectory  $\tau = s_0, a_0, r_1, s_1, a_1, r_2, s_2, a_2, r_3, s_3, \dots, s_T$  according to current policy  $\pi$ ; for each time step  $0 \le t \le T - 1$ , perform the following update:

$$Q^{n+1}(s_t, a_t) = Q^n(s_t, a_t) + \alpha_n \cdot \left[ r_{t+1} + \gamma Q^n(s_{t+1}, a_{t+1}) - Q^n(s_t, a_t) \right]$$

The update can be performed immediately when  $s_{t+1}$  and  $a_{t+1}$  is known (no need to wait for the episode to terminate).

After the episode ends, make  $\pi$   $\varepsilon$ -Q-greedy.

Conforms to the GVI scheme.

## SARSA pseudocode

#### Sarsa (on-policy TD control) for estimating $Q \approx q_*$

```
Algorithm parameters: step size \alpha \in (0,1], small \varepsilon > 0

Initialize Q(s,a), for all s \in \mathbb{S}^+, a \in \mathcal{A}(s), arbitrarily except that Q(terminal, \cdot) = 0

Loop for each episode:

Initialize S

Choose A from S using policy derived from Q (e.g., \varepsilon-greedy)

Loop for each step of episode:

Take action A, observe R, S'

Choose A' from S' using policy derived from Q (e.g., \varepsilon-greedy)

Q(S,A) \leftarrow Q(S,A) + \alpha \left[R + \gamma Q(S',A') - Q(S,A)\right]

S \leftarrow S'; A \leftarrow A';

until S is terminal
```

source: Sutton&Barto, p. 130

## **GLIE** policies

#### Definition 43: GLIE condition

A RL algorithm is greedy in the limit (GL) if its behavior policy (=target policy in onpolicy algorithms) converges to a 0-greedy policy with increasing number of episodes. A RL algorithm is GLIE if it is GL and IE (infinitely exploring).

Typical ways of ensuring GLIE:

## **GLIE** policies

#### Definition 43: GLIE condition

A RL algorithm is greedy in the limit (GL) if its behavior policy (=target policy in on-policy algorithms) converges to a 0-greedy policy with increasing number of episodes. A RL algorithm is GLIE if it is GL and IE (infinitely exploring).

### Typical ways of ensuring GLIE:

• Dynamically adjust  $\varepsilon$  in  $\varepsilon$ -greedy policy selection" When selecting action in state s, behave  $\varepsilon$ -greedily with  $\varepsilon = \frac{c}{n(s)}$ , where 0 < c < 1 is a constant and n(s) is a number of visits to state s over all the episodes so far.

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- Dynamically adjust  $\varepsilon$  in  $\varepsilon$ -greedy policy selection" When selecting action in state s, behave  $\varepsilon$ -greedily with  $\varepsilon = \frac{c}{n(s)}$ , where 0 < c < 1 is a constant and n(s) is a number of visits to state s over all the episodes so far.
- Use Boltzmann (softmax) exploration:

$$\pi(a \mid s) = \frac{e^{\frac{Q(s,a)}{\eta(s)}}}{\sum_{b \in \mathcal{A}(s)} e^{\frac{Q(s,b)}{\eta(s)}}},$$

where  $\eta$  is a state-dependent and time-varying temperature parameter. We need  $\eta$  to converge to 0 over time, but not too fast (often,  $\eta(s)$  proportional to  $\frac{1}{\log(n(s))}$ ).

## Convergence of SARSA

#### Theorem 44

Consider a GLIE instantiation of SARSA. Moreover, assume that the sequence of learning rates  $(\alpha_n)_{n\in\mathbb{N}}$  satisfies  $\sum_n \alpha_n = \infty$  and  $\sum_n \alpha_n^2 < \infty$ . In this setting, Q converges to  $q^*$  and the behavior policy of SARSA converges to some optimal policy  $\pi^*$ .

For the proof, see optional reading: Singh, Jaakkola, Littman, Szepesvári: Convergence Results for Single-Step On-Policy Reinforcement-Learning Algorithms. In *Machine Learning* 39:287-308. Kluwer, 2000.

Note: learning rate can itself be state/action dependent (omitted for conciseness, constant learning rates preferred in practice).

Surprise: no importance sampling!

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Recall the SARSA update:

$$Q^{n+1}(s_t, a_t) = Q^n(s_t, a_t) + \alpha_n \cdot \left[ r_{t+1} + \gamma Q^n(s_{t+1}, a_{t+1}) - Q^n(s_t, a_t) \right]$$

It pushes Q towards  $q^{\pi}$ , where  $\pi$  is the current policy.

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$$Q^{n+1}(s_t, a_t) = Q^n(s_t, a_t) + \alpha_n \cdot \left[ r_{t+1} + \gamma Q^n(s_{t+1}, a_{t+1}) - Q^n(s_t, a_t) \right]$$

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Idea: push Q directly towards  $q^*$ .

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$$Q^{n+1}(s_t, a_t) = Q^n(s_t, a_t) + \alpha_n \cdot \left[ r_{t+1} + \gamma Q^n(s_{t+1}, a_{t+1}) - Q^n(s_t, a_t) \right]$$

It pushes Q towards  $q^{\pi}$ , where  $\pi$  is the current policy.

Idea: push Q directly towards  $q^*$ .

We could do this e.g. by a VI-like update:

$$Q^{n+1}(s_t, a_t) = Q^n(s_t, a_t) + \alpha_n \cdot \Big[ \max_{a \in \mathcal{A}(s)} \sum_{s' \in \mathcal{S}} p(s' \mid s, a) \big( r(s, a, s') + \gamma V(s') \big) - Q^n(s_t, a_t) \Big].$$

Two problems:

- We do not calculate v-estimates. (Must somehow replace with Q)
- ullet We must get rid of transition probabilities and instead use the sampled  $a_t$  and  $r_{t+1}$ .

### Q-learning update

Solution: push the max towards the bootstrap.

Q-learning (Watkins, 1989): given a sampled trajectory  $s_0$ ,  $a_0$ ,  $r_1$ ,  $s_1$ ,  $a_1$ ,  $r_2$ ,  $s_2$ ,  $a_2$ ,  $r_3$ ,  $s_3$ , . . ., for every t we update:

$$Q^{n+1}(s_t, a_t) = Q^n(s_t, a_t) + \alpha_n \cdot \left[ r_{t+1} + \gamma \cdot \left( \max_{a \in \mathcal{A}(s_{t+1})} Q^n(s_{t+1}, a) \right) - Q^n(s_t, a_t) \right]$$

### Q-learning pseudocode

#### Q-learning (off-policy TD control) for estimating $\pi \approx \pi_*$

Algorithm parameters: step size  $\alpha \in (0,1]$ , small  $\varepsilon > 0$ Initialize Q(s,a), for all  $s \in \mathbb{S}^+$ ,  $a \in \mathcal{A}(s)$ , arbitrarily except that  $Q(terminal,\cdot) = 0$ 

Loop for each episode:

Initialize S

Loop for each step of episode:

Choose A from S using policy derived from Q (e.g.,  $\varepsilon$ -greedy)

Take action A, observe R, S'

$$Q(S, A) \leftarrow Q(S, A) + \alpha [R + \gamma \max_{a} Q(S', a) - Q(S, A)]$$

$$S \leftarrow S'$$

until S is terminal

source: Sutton&Bato, p. 131

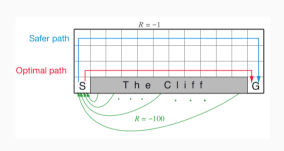
### **Q**-learning convergence

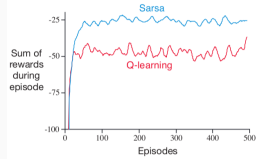
#### Theorem 45

Consider any Q-learning instantiation with infinite exploration. Assume that the sequence of learning rates  $(\alpha_n)_{n\in\mathbb{N}}$  satisfies  $\sum_n \alpha_n = \infty$  and  $\sum_n \alpha_n^2 < \infty$ . In this setting, Q converges to  $q^*$ . Moreover, if the behavior policy is GL, then it converges to an optimal policy  $\pi^*$ .

Proof in optional reading: Watkins, Dayan: Q-Learning. In *Machine Learning* 8:279-292. Kluwer, 1992.

## SARSA vs. Q-learning (SB: p. 132)

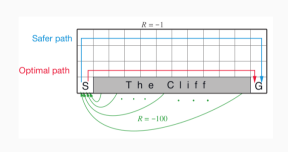


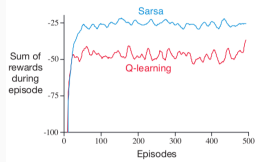


Left: greedy policies learned by SARSA and Q-learning.

Right: in-training performance with a 0.1-greedy behavior policy.

## SARSA vs. Q-learning (SB: p. 132)





Left: greedy policies learned by SARSA and Q-learning.

Right: in-training performance with a 0.1-greedy behavior policy.

(Rough) takeaway: Q-learning more aggressive in finding optimal policy, can lead to risky behavior. Possibly advantageous when environment not too stochastic or if in-training performance has less importance (simulator vs. real world).

## Maximization bias in Q-learning

Q-learning is "risky" not only due to exploration, but also because it is optimistic in the face of uncertainty. TBD

The positive bias only disappears in the limit.

### **Double Q-learning**

Idea: use two independent value estimates  $Q_1$ ,  $Q_2$  and decouple action selection from evaluation in the bootstrap. During each update, we randomly select one of these for update, which is also used to select the maximizing action in bootstrap. The other is used as the bootstrap estimate.

### **Double Q-learning**

Idea: use two independent value estimates  $Q_1$ ,  $Q_2$  and decouple action selection from evaluation in the bootstrap. During each update, we randomly select one of these for update, which is also used to select the maximizing action in bootstrap. The other is used as the bootstrap estimate.

I.e., in each time step t we perform one of these updates, each with probability  $\frac{1}{2}$ : either

$$Q_1(s_t, a_t) = Q_1(s_t, a_t) + \alpha_n \cdot \left[ r_{t+1} + \gamma \cdot \frac{Q_2(s_{t+1}, (\max_{a \in \mathcal{A}(s_{t+1})} Q_1(s_{t+1}, a))) - Q_1(s_t, a_t) \right]$$

or

$$Q_2(s_t, a_t) = Q_2(s_t, a_t) + \alpha_n \cdot \left[ r_{t+1} + \gamma \cdot Q_1(s_{t+1}, (\max_{a \in \mathcal{A}(s_{t+1})} Q_2(s_{t+1}, a))) - Q_2(s_t, a_t) \right].$$

Behavior policy = e.g.  $\varepsilon$ -greedy w.r.t.  $Q_1 + Q_2$ .

### Double Q-learning pseudocode

#### Double Q-learning, for estimating $Q_1 \approx Q_2 \approx q_*$

Algorithm parameters: step size  $\alpha \in (0,1]$ , small  $\varepsilon > 0$ 

Initialize  $Q_1(s,a)$  and  $Q_2(s,a)$ , for all  $s \in S^+, a \in A(s)$ , such that  $Q(terminal, \cdot) = 0$ 

Loop for each episode:

Initialize S

Loop for each step of episode:

Choose A from S using the policy  $\varepsilon$ -greedy in  $Q_1 + Q_2$ 

Take action A, observe R, S'

With 0.5 probability:

$$Q_1(S, A) \leftarrow Q_1(S, A) + \alpha \Big(R + \gamma Q_2(S', \operatorname{argmax}_a Q_1(S', a)) - Q_1(S, A)\Big)$$

else:

$$Q_2(S, A) \leftarrow Q_2(S, A) + \alpha \Big( R + \gamma Q_1 \big( S', \operatorname{argmax}_a Q_2(S', a) \big) - Q_2(S, A) \Big)$$
  
$$S \leftarrow S'$$

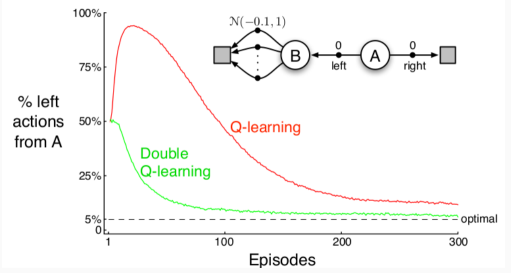
until S is terminal

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## Why Double Q-learning helps

TBD

## Double Q-learning: experiment



source: Sutton&Barto, p. 135

# **Between Monte Carlo** and TD:

*n*-Step and  $\lambda$ -Returns

### MC vs TD(0) update targets

Given a trajectory  $s_0, a_0, r_1, s_1, a_1, r_2, s_2, a_2, r_3, s_3, ...$ :



Update for time step *t* in:

• MC = discounted return from t till the end of trajectory, e.g. for t = 1:

$$r_2 + \gamma r_3 + \gamma^2 r_4 + \dots + \gamma^{T-2} r_T$$

unbiased, but high variance + need the whole trajectory

### MC vs TD(0) update targets

Given a trajectory  $s_0, a_0, r_1, s_1, a_1, r_2, s_2, a_2, r_3, s_3, ...$ :



Update for time step *t* in:

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unbiased, but high variance + need the whole trajectory

• TD(0) = 1-step reward and then (discounted) bootstrap:

$$r_2 + \gamma V(s_2)$$

#### n-step return

Idea: use *n*-step discounted return and then bootstrap

#### Definition 46

Let  $\tau = s_0, a_0, r_1, s_1, a_1, r_2, s_2, a_2, r_3, s_3, \dots$  be a trajectory and  $n \in \mathbb{N} \setminus \{0\}$ .

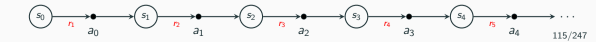
An *n*-step return of  $\tau$  from time step t is the quantity

$$G_{t:t+n}(\tau) = r_{t+1} + \gamma \cdot r_{t+2} + \gamma^2 \cdot r_{t+3} + \cdots + \gamma^{n-1} r_{t+n} + \gamma^n \cdot V(s_{t+n}).$$

We also define a Q-estimate-based version:

$$G_{t:t+n}(\tau) = r_{t+1} + \gamma \cdot r_{t+2} + \gamma^2 \cdot r_{t+3} + \cdots + \gamma^{n-1} r_{t+n} + \gamma^n \cdot Q(s_{t+n}, a_{t+n}).$$

(Which of the two is used will be clear from the context.)



### n-step TD policy evaluation

Similar to TD(0), but using *n*-step return targets:

given a trajectory  $\tau = s_0, a_0, r_1, s_1, a_1, r_2, s_2, a_2, r_3, s_3, \ldots$ , for each  $0 \le t < T$  we perform an update

$$V(s_t) \leftarrow V(s_t) + \alpha [G_{t:t+n}(\tau) - V(s_t)].$$

Note 1: for n = 1 we get exactly TD(0).

### n-step TD policy evaluation

Similar to TD(0), but using *n*-step return targets:

given a trajectory  $\tau = s_0, a_0, r_1, s_1, a_1, r_2, s_2, a_2, r_3, s_3, \ldots$ , for each  $0 \le t < T$  we perform an update

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Note 1: for n = 1 we get exactly TD(0).

Note 2: for n > 1, we cannot update  $V(s_t)$  directly at step t + 1. We need to obtain  $r_{t+1}, \ldots, r_{t+n}, s_{t+n}$  first, i.e. we can perform the update after step t + n.

### n-step TD policy evaluation

Similar to TD(0), but using *n*-step return targets:

given a trajectory  $\tau = s_0, a_0, r_1, s_1, a_1, r_2, s_2, a_2, r_3, s_3, \ldots$ , for each  $0 \le t < T$  we perform an update

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Note 1: for n = 1 we get exactly TD(0).

Note 2: for n > 1, we cannot update  $V(s_t)$  directly at step t + 1. We need to obtain  $r_{t+1}, \ldots, r_{t+n}, s_{t+n}$  first, i.e. we can perform the update after step t + n.

Note 3: if t + n > T, we truncate the sum in  $G_{t:t+n}$  at  $r_T$ , i.e. in such a case  $G_{t:t+n} = G_t$ .

### *n*-step TD policy evaluation: pseudocode

```
n-step TD for estimating V \approx v_{\pi}
Input: a policy \pi
Algorithm parameters: step size \alpha \in (0,1], a positive integer n
Initialize V(s) arbitrarily, for all s \in S
All store and access operations (for S_t and R_t) can take their index mod n+1
Loop for each episode:
   Initialize and store S_0 \neq \text{terminal}
   T \leftarrow \infty
   Loop for t = 0, 1, 2, \dots:
       If t < T, then:
           Take an action according to \pi(\cdot|S_t)
           Observe and store the next reward as R_{t+1} and the next state as S_{t+1}
           If S_{t+1} is terminal, then T \leftarrow t+1
       \tau \leftarrow t - n + 1 (\tau is the time whose state's estimate is being updated)
       If \tau > 0:
           G \leftarrow \sum_{i=\tau+1}^{\min(\tau+n,T)} \gamma^{i-\tau-1} R_i
           If \tau + n < T, then: G \leftarrow G + \gamma^n V(S_{\tau+n})
           V(S_{\tau}) \leftarrow V(S_{\tau}) + \alpha \left[ G - V(S_{\tau}) \right]
   Until \tau = T - 1
```

source: Sutton&Barto, p.144

### *n*-step TD policy evaluation: performance

19-state symmetric random walk:

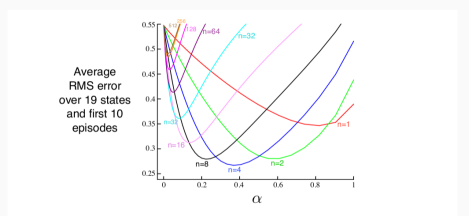


Figure 7.2: Performance of n-step TD methods as a function of  $\alpha$ , for various values of n, on a 19-state random walk task (Example 7.1).

source: Sutton&Barto, p.145

### *n*-step SARSA (on-policy control)

Uses Q-value-bootstrapped *n*-step returns.

For a sampled trajectory  $\tau=s_0, a_0, r_1, s_1, a_1, r_2, s_2, a_2, r_3, s_3, \ldots$  and for all time steps  $0 \le t < T$  we perform an update

$$Q(s_t, a_t) \leftarrow Q(s_t, a_t) + \alpha [G_{t:t+n} - Q(s_t, a_t)].$$

We sample trajectories according to a policy  $\pi$  that is  $\varepsilon$ -greedy w.r.t. current Q-estimates:

$$\pi(a|s) = \begin{cases} 1 - \varepsilon + \frac{\varepsilon}{|\mathcal{A}(s)|} & \text{if } a = \arg\max_{a' \in \mathcal{A}(s)} Q(s, a') \text{ (ties broken in principled way)} \\ \frac{\varepsilon}{|\mathcal{A}(s)|} & \text{otherwise.} \end{cases}$$

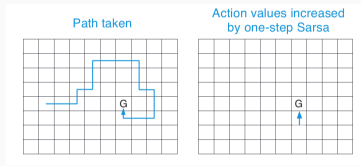
 $\boldsymbol{\pi}$  is redefined in this way after each episode

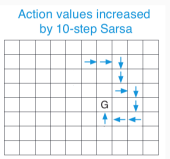
### n-step SARSA (pseudocode)

```
Loop for each episode:
    Initialize and store S_0 \neq \text{terminal}
    Select and store an action A_0 \sim \pi(\cdot|S_0)
   T \leftarrow \infty
   Loop for t = 0, 1, 2, ...:
        If t < T, then:
            Take action A_t
            Observe and store the next reward as R_{t+1} and the next state as S_{t+1}
            If S_{t+1} is terminal, then:
                T \leftarrow t + 1
            else:
                Select and store an action A_{t+1} \sim \pi(\cdot|S_{t+1})
        \tau \leftarrow t - n + 1 (\tau is the time whose estimate is being updated)
        If \tau > 0:
           G \leftarrow \sum_{i=\tau+1}^{\min(\tau+n,T)} \gamma^{i-\tau-1} R_i
            If \tau + n < T, then G \leftarrow G + \gamma^n Q(S_{\tau+n}, A_{\tau+n})
            Q(S_{\tau}, A_{\tau}) \leftarrow Q(S_{\tau}, A_{\tau}) + \alpha \left[ G - Q(S_{\tau}, A_{\tau}) \right]
            If \pi is being learned, then ensure that \pi(\cdot|S_{\tau}) is \varepsilon-greedy wrt Q
    Until \tau = T - 1
```

source: Sutton&Barto, p. 147

### n-step SARSA (speed of signal propagation)





source: Sutton&Barto, p. 147

### n-step Q-learning

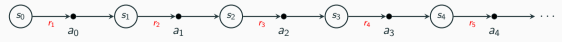
The Q-learning update for a trajectory  $\tau = s_0, a_0, r_1, s_1, a_1, r_2, s_2, a_2, r_3, s_3, \ldots$  and time step t:

$$Q(s_t, a_t) \leftarrow Q(s_t, a_t) + \alpha \cdot [r_{t+1} + \gamma \cdot \max_{a \in \mathcal{A}(s_{t+1})} Q(s_{t+1}, a) - Q(s_t, a_t)].$$

Naive extension to n-step returns

$$Q(s_t, a_t) \leftarrow Q(s_t, a_t) + \alpha \cdot [r_{t+1} + \gamma r_{t+2} + \dots + \gamma^{t+n-1} \cdot r_{t+n} + \gamma^{t+n} \cdot \max_{a \in \mathcal{A}(s_{t+1})} Q(s_{t+n+1}, a) - Q(s_t, a_t)]$$

does not really correspond to Q-learning, since some of the actions  $a_{t+1}, \ldots, a_{t+n}$  might not be Q-greedy (the behavior policy is  $\varepsilon$ -greedy, so some actions might be exploratory). Hence, we are no longer pushing Q towards the Q-value of an optimal policy.



### n-step Q-learning: correct

Idea: apply the Q-learning bootstrap at the first occurrence of a non-Q-greedy action.

I.e., for each episode:

- make  $\pi$  an  $\varepsilon$ -Q-greedy policy
- sample a trajectory  $\tau = s_0, a_0, r_1, s_1, a_1, r_2, s_2, a_2, r_3, s_3, \dots$  from  $\pi$
- for each time step  $0 \le t < T$ :
  - identify the smallest  $n' \in \{t+1, t+2, \dots, t+n\}$  such that  $a_{n'}$  is not a Q-greedy action
  - perform the update

$$Q(s_t, a_t) \leftarrow Q(s_t, a_t) + \alpha \cdot [r_{t+1} + \gamma r_{t+2} + \dots + \gamma^{n'-1} r_{n'} + \gamma^{n'} \cdot \max_{a \in \mathcal{A}(s_{n'})} Q(s_{n'}, a) - Q(s_t, a_t)]$$

(if n' > T, do the standard MC update).



#### $\lambda$ -returns: idea

By varying n, the n-step returns provide a nice tradeoff between bias and variance (and update speed). But the choice of optimal n is mostly a guesswork.

Idea: find a notion of return which combines n-step returns for multiple n's. E.g., a suitable convex combination of individual n-step returns. This leads to the notion of  $\lambda$ -returns.

We will focus only on policy evaluation, though  $\lambda$ -returns can be used also in control.

#### $\lambda$ -returns: definition

Recall:  $G_{t:t+n}$  is the *n*-step return from timestep t.

#### Definition 47: $\lambda$ -return

Let  $\lambda \in [0,1]$ . A  $\lambda$ -return from timestep t is the random variable

$$G_t^{\lambda} = (1 - \lambda) \sum_{n=1}^{\infty} \lambda^{n-1} G_{t:t+n}.$$

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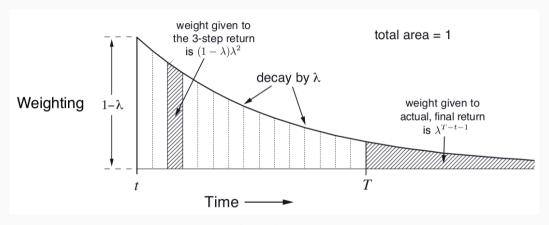
$$G_t^{\lambda} = (1 - \lambda) \sum_{n=1}^{\infty} \lambda^{n-1} G_{t:t+n}.$$

Note that due to truncation at  $t + n \ge T$ , the  $\lambda$ -return can be more explicitly written as

$$G_t^{\lambda} = (1 - \lambda) \sum_{n=1}^{T-t-1} \lambda^{n-1} \cdot G_{t:t+n} + \lambda^{T-t-1} \cdot G_t.$$

$$\underbrace{\begin{pmatrix} s_0 \end{pmatrix}}_{n} \xrightarrow{a_0} \underbrace{\begin{pmatrix} s_1 \end{pmatrix}}_{n} \xrightarrow{s_2} \underbrace{\begin{pmatrix} s_2 \end{pmatrix}}_{n} \xrightarrow{s_3} \underbrace{\begin{pmatrix} s_3 \end{pmatrix}}_{n} \xrightarrow{s_4} \underbrace{\begin{pmatrix} s_4 \end{pmatrix}}_{n}}_{n} \underbrace{\begin{pmatrix} s_4 \end{pmatrix}}_{n} \underbrace{\begin{pmatrix} s_4 \end{pmatrix}}_{n} \xrightarrow{s_4 \end{pmatrix}}_{n} \underbrace{\begin{pmatrix} s_4 \end{pmatrix}}_{n} \underbrace{\begin{pmatrix}$$

### $\lambda$ -return as discounting of n-step returns



source: Sutton&Barto, p. 290

## (Forward-view) $TD(\lambda)$

Like TD(0), but uses  $\lambda$ -returns.

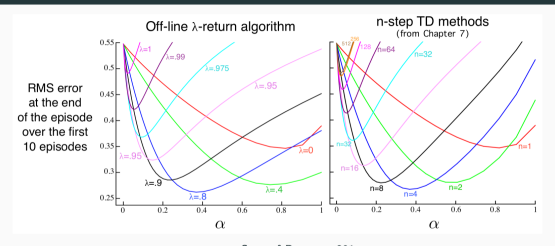
Given a trajectory  $\tau = s_0, a_0, r_1, s_1, a_1, r_2, s_2, a_2, r_3, s_3, \dots$  sampled from the evaluated policy  $\pi$ , we perform, for each time step t an update:

$$V(s_t) \leftarrow V(s_t) + \alpha(G_t^{\lambda}(\tau) - V(s_t)).$$

Note:

- for  $\lambda = 0$ , this is exactly the TD(0) update,
- for  $\lambda = 1$ , this is exactly the MC update,
- $G_t^{\lambda}(\tau)$  depends on the whole suffix of  $\tau_{t..}$ , hence the update can be only performed at the end of the episode. (We will show a workaround later.)

## $\mathsf{TD}(\lambda)$ vs. *n*-step $\mathsf{TD}$ on 19-state random walk



source: Sutton&Barto, p. 291

## Forward vs. backward view $TD(\lambda)$

Backward-view  $\mathsf{TD}(\lambda) = \mathsf{an}$  algorithm performing roughly the same updates as Forward-view  $\mathsf{TD}(\lambda)$  in an online fashion  $(V(s_t)$  can be updated by time t+1).

# Forward vs. backward view $TD(\lambda)$

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# Forward vs. backward view $TD(\lambda)$

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Implemented using eligibility traces: state-wise signals that indicate how much is the current state eligible for an update (sort of state-wise modulation of the learning rate).

We are more keen to update states that:

- appear often along the trajectory (frequency heuristic)
- were visited in the recent past (recency heuristic)

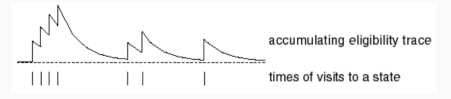
# Accumulating eligibility trace

#### Definition 48: (Accumulating) eligibility trace

For a trajectory  $\tau = s_0, a_0, r_1, s_1, a_1, r_2, s_2, a_2, r_3, s_3, \ldots, \lambda \in [0, 1]$ , and a state  $s \in \mathcal{S}$ , an (accumulating) eligibility trace is a sequence of values  $E_0(s), E_1(s), E_2(s), \ldots$  defined inductively as follows:

$$E_0(s)=0$$
 and for  $t>0$   $E_t(s)=\gamma\cdot\lambda\cdot E_{t-1}(s)+\mathbb{I}(S_t( au)=s),$ 

where  $\mathbb{I}(S_t(\tau) = s)$  is the indicator of the t-th state of  $\tau$  being s, i.e.  $\mathbb{I}(S_t(\tau) = s) = 1$  if  $s_t = s$  and  $\mathbb{I}(S_t(\tau) = s) = 0$  otherwise.



source: slides of D. Silver (Model-free prediction)

#### Backward-view TD( $\lambda$ ): idea

- $E_t(s)$  denotes how much is s eligible for an update after playing t-th action along the run (i.e., action  $a_{t-1}$ ).
- In time step t, all states with non-zero eligibility signal will have their estimates updated in proportion to the learning rate and the strength of the eligibility signal.
- The update target is the standard TD(0) target for time t. I.e., for each timestep t and each state q, we perform the update

$$V(q) \leftarrow V(q) + \alpha \cdot E_t(q) \cdot [r_{t+1} + \gamma \cdot V(s_{t+1}) - V(s_t)].$$

# Backward-view TD( $\lambda$ ): pseudocode

```
Input: policy \pi to evaluate
```

**Output:** Estimate V of  $v^{\pi}$ 

initialize V arbitrarily

#### repeat

```
s \leftarrow sample uniformly (ES) or according to init. distr.
initialize E to be uniformly zero
while s not terminal do
     a \leftarrow \text{sample from } \pi(s)
     s' \leftarrow \text{sample from } p(s, a)
     r \leftarrow r(s, a, s')
     foreach q \in \mathcal{S} (Only q's visited so far) do
         E(q) = \gamma \cdot \lambda \cdot E(q) + \mathbb{I}(q = s)
       V(q) \leftarrow V(q) + \alpha \cdot E(q) \cdot [r + \gamma \cdot V(s') - V(s)]
     s \leftarrow s'
```

#### Forward vs. backward view

If  $\lambda=0$ , then  $E_t(q)=\mathbb{I}(S_t(\tau)=q)$ , i.e. the backward-view update at time point t is

$$V(s_t) \leftarrow V(s_t) + \alpha \cdot [r + \gamma \cdot V(s_{t+1}) - V(s_t)],$$

while for all states other than  $s_t$ , no update is performed. I.e., backward TD(0) is exactly the same thing as forward TD(0).

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while for all states other than  $s_t$ , no update is performed. I.e., backward TD(0) is exactly the same thing as forward TD(0).

For general  $\lambda$  the correspondence is more subtle:

#### Theorem 49: Forward-backward view correspondence

Assume that in the backward view, all the updates along the trajectory are performed offline, i.e. only after the end of the episode, and in a batch, i.e. concurrently, using the pre-episode estimates in right-hand sides.

Then, for any  $\lambda \in (0,1)$ , this offline backward  $\mathsf{TD}(\lambda)$  performs the same updates as forward  $\mathsf{TD}(\lambda)$ .

# Offline backward and forward view (source: D. Silver slides)

Given the batch nature of updates, it suffices to show that the forward update target at time t equals the sum of all updates "triggered" by a visit to state  $s_t$ .

$$G_{t}^{\lambda} - V(S_{t}) = -V(S_{t}) + (1 - \lambda)\lambda^{0} (R_{t+1} + \gamma V(S_{t+1})) + (1 - \lambda)\lambda^{1} (R_{t+1} + \gamma R_{t+2} + \gamma^{2} V(S_{t+2})) + (1 - \lambda)\lambda^{2} (R_{t+1} + \gamma R_{t+2} + \gamma^{2} R_{t+3} + \gamma^{3} V(S_{t+3})) + ...$$

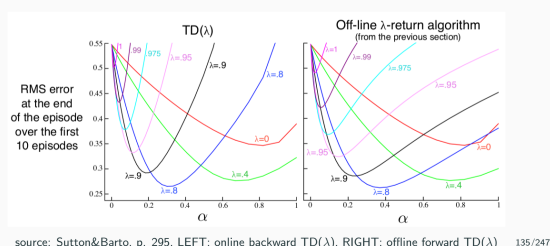
$$= -V(S_{t}) + (\gamma \lambda)^{0} (R_{t+1} + \gamma V(S_{t+1}) - \gamma \lambda V(S_{t+1})) + (\gamma \lambda)^{1} (R_{t+2} + \gamma V(S_{t+2}) - \gamma \lambda V(S_{t+2})) + (\gamma \lambda)^{2} (R_{t+3} + \gamma V(S_{t+3}) - \gamma \lambda V(S_{t+3})) + ...$$

$$= (\gamma \lambda)^{0} (R_{t+1} + \gamma V(S_{t+1}) - V(S_{t})) + (\gamma \lambda)^{1} (R_{t+2} + \gamma V(S_{t+2}) - V(S_{t+1})) + (\gamma \lambda)^{2} (R_{t+3} + \gamma V(S_{t+3}) - V(S_{t+2})) + ...$$

$$= \delta_{t} + \gamma \lambda \delta_{t+1} + (\gamma \lambda)^{2} \delta_{t+2} + ...$$

#### Online vs. offline backward view on 19-state RW

In practice, we want to use the online backward algorithm, which only approximates the forward view. Nevertheless, it performs acceptably:



source: Sutton&Barto, p. 295, LEFT: online backward TD( $\lambda$ ), RIGHT: offline forward TD( $\lambda$ )

• There are other types of eligibility traces (replacing, dutch, ...), yielding different algorithms.

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- $\lambda$ -returns and eligibility traces can be generalized to control setting SARSA( $\lambda$ ), Q( $\lambda$ ); optional reading: Sutton&Barto, Sec. 12.7-12.10.

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- $\lambda$ -returns and eligibility traces can be generalized to control setting SARSA( $\lambda$ ), Q( $\lambda$ ); optional reading: Sutton&Barto, Sec. 12.7-12.10.
- There is a true online backward  $TD(\lambda)$  version. Here, true online=having perfect equivalence with the forward view. However, the equivalence is w.r.t. a more complex notion of  $\lambda$ -return (truncated  $\lambda$ -return) and uses more complex version of eligibility traces than presented here. Outperforms both forward and backward algorithms presented here. Optional reading: van Seijen, Sutton: True Online  $TD(\lambda)$ . In Proceedings of ICML'14.

# First Steps Towards Deep RL:

Value-Based

**On-Policy Methods** 

# Working with huge MDPs

E.g. original Atari games have  $160 \times 192$  resolution with 128 colors: observable state space of size  $2^{7 \cdot 160 \cdot 192} = 2^{215040}$  (though only a fraction reachable and resolution typically scaled down in benchmarks – however, state typically encompass last 3 frames so as to provide some info on movement).

State space can be even continuous (position, velocity,...).

Most states will not be seen - we need the ability to generalize from experience to unseen/rarely seen states.

## **Huge MDP representation**

From now on, states of the MDP will be represented by vectors from  $\mathbb{R}^n$ . The vectorized representation is chosen in a domain-specific manner, e.g.:

- Atari = one component per pixel per frame
- continuous navigation = agent coordinates, velocity, etc.
- small discrete MDPs can be represented by one-hot encoding

For simplicity, we will still assume that the action space is discrete, and reasonably small, though many algorithms can be adapted for continuous actions (acceleration, etc).

#### **Function approximators**

The value functions have types:

$$v^{\pi}, v^* \colon \mathbb{R}^n \to \mathbb{R} \qquad q^{\pi}, q^* \colon \mathbb{R}^n \times \mathcal{A} \to \mathbb{R}.$$

In RL, we need to approximate these functions.

#### Definition 50

A function approximator (FA) for functions of type  $X \to Y$  is a class of functions  $f \subseteq Y^X$  parameterized by a some set of parameter vectors  $\Theta \subseteq \mathbb{R}^n$ .

Each concrete parameter vector  $\theta \in \Theta$  defines a concrete function  $f_{\theta} \in f$ , i.e.  $f = \{f_{\theta} \mid \theta \in \Theta\}$ .

For FA f, we often write  $f_{\theta}(x) = f(x, \theta)$  to stress the fact that the output of  $f_{\theta}$  depends on both the input x and on  $\theta$ . Hence, FA for type  $X \to y$  can be itself seen as a function of type  $X \times \Theta \to Y$ .

#### Function approximators in RL

Our algorithms will use mainly these types of function approximators:

- $V : \mathbb{R}^n \times \Theta \to \mathbb{R}$  to approximate  $v^{\pi}$  or  $v^{\theta}$
- $Q \colon (\mathbb{R}^n imes \mathcal{A}) imes \Theta o \mathbb{R}$  to approximate  $q^\pi$  or  $q^ heta$

The typical task is to find  $\theta \in \Theta$  such that  $V_{\theta} = V(\cdot, \theta)$  is a "good" approximation for  $v^{\pi}$  or  $v^{\theta}$ , and similarly for  $Q_{\theta}$ .

The parametrization  $\Theta$  will depend on the concrete form of function approximator used.

## Forms of function approximators

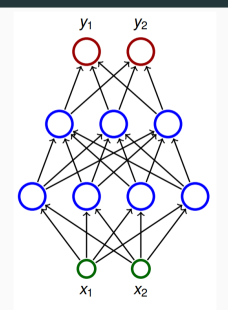
- tabular
  - $\theta =$  the vector containing the contents of the table
- linear

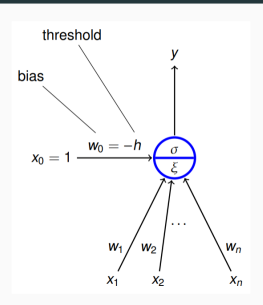
$$ullet$$
  $\Theta = \mathcal{S} = \mathbb{R}^n$  and e.g.  $V_{ heta}(s) = heta^ op \cdot s$ 

- neural nets
  - $\theta = NN$  weights and biases
- decision trees
- . . .

We require the approximators to be differentiable and to admit a training method suitable for non-stationary data.

# Neural nets (source: slides by T. Brázdil)





## Policy evaluation with FAs

Task: given a policy  $\pi$  and FA  $V: \mathbb{R}^n \times \Theta \to \mathbb{R}$ , find  $\theta$  s.t.  $V_{\theta}$  is "close" to  $v^{\pi}$ .

"Closeness" can be expressed using various loss functions. Typically, we want to minimize the mean squared error (MSE):

$$\mathit{MSE}(v^{\pi}, V_{\theta}) = \frac{1}{2} \mathbb{E}_{s \sim \mu} \big[ (v^{\pi}(s) - V_{\theta}(s))^2 \big] = \frac{1}{2} \sum_{s \in \mathcal{S}} \mu(s) \cdot \big[ (v^{\pi}(s) - V_{\theta}(s))^2 \big],$$

where  $\mu$  is some distribution over states expressing how much do we care about errors in particular states.

A local minimum of MSE can be found gradient descent: making successive step in the direction opposite to the gradient of MSE.

#### **Recall:** gradients

#### Definition 51

Given a scalar function  $f(x_1, \ldots, x_n, \theta_1, \ldots, \theta_m) \colon \mathbb{R}^n \times \Theta \to \mathbb{R}$  (where  $\Theta \subseteq \mathbb{R}^m$ ), the gradient of f w.r.t. parameters  $\theta = (\theta_1, \ldots, \theta_m)$  is the vector function

$$\nabla_{\theta} f = (\frac{\partial f}{\partial \theta_1}, \dots, \frac{\partial f}{\partial \theta_m}) \text{ of type } \mathbb{R}^n \times \Theta \to \mathbb{R}^m$$

When f is a function approximator defined by a neural net, the value of the gradient  $\nabla_{\theta} f(x, \theta)$  at a given point  $(x, \theta) = (x_1, \dots, x_n, \theta_1, \dots, \theta_m)$  can be computed by backpropagation (under some usual conditions like smoothness, etc.).

To (locally) minimize  $MSE(v^{\pi}, V_{\theta})$ , it suffices to perform (sufficiently small) steps in the negative direction of the current gradient, i.e., repeatedly perform updates:

$$\theta \leftarrow \theta - \alpha \cdot \nabla_{\theta} \mathit{MSE}(v^{\pi}, V_{\theta}) = \theta - \alpha \cdot \nabla_{\theta} \frac{1}{2} \cdot \mathbb{E}_{s \sim \mu} \big[ \big( v^{\pi}(s) - V_{\theta}(s) \big)^2 \big]$$

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$$= \theta - \frac{\alpha}{2} \cdot \mathbb{E}_{s \sim \mu} \left[ \nabla_{\theta} \left( v^{\pi}(s) - V_{\theta}(s) \right)^{2} \right]$$

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$$= \theta - \frac{\alpha}{2} \cdot \mathbb{E}_{s \sim \mu} \left[ \nabla_{\theta} \left( v^{\pi}(s) - V_{\theta}(s) \right)^{2} \right]$$

$$= \theta + \alpha \cdot \mathbb{E}_{s \sim \mu} \left[ \left( v^{\pi}(s) - V_{\theta}(s) \right) \cdot \nabla_{\theta} V_{\theta}(s) \right]$$

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$$= \theta - \frac{\alpha}{2} \cdot \mathbb{E}_{s \sim \mu} [\nabla_{\theta} (v^{\pi}(s) - V_{\theta}(s))^{2}]$$

$$= \theta + \alpha \cdot \mathbb{E}_{s \sim \mu} [(v^{\pi}(s) - V_{\theta}(s)) \cdot \nabla_{\theta} V_{\theta}(s)]$$

The expected value above is typically impossible to evaluate in practice. Instead we estimate it by samples  $\Rightarrow$  stochastic gradient descent.

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$$= \theta - \frac{\alpha}{2} \cdot \mathbb{E}_{s \sim \mu} [\nabla_{\theta} (v^{\pi}(s) - V_{\theta}(s))^{2}]$$

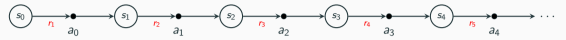
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The expected value above is typically impossible to evaluate in practice. Instead we estimate it by samples  $\Rightarrow$  stochastic gradient descent.

We typically take  $\mu(s)$  representing the overall fraction of time spent in s when behaving according to  $\mu$ . Hence,  $\mathbb{E}_{s\sim\mu}$  can be estimated by sampling a trajectory from  $\mu$  and performing the update for each s on the trajectory in an every-visit fashion.

## Stochastic gradient policy evaluation + MC instantiation

We keep sampling trajectories  $\tau$  from  $\pi$ :



For each timestep t we perform the update of parameters

$$\theta \leftarrow \theta + \alpha \cdot \left[ \left( v^{\pi}(s_t) - V_{\theta}(s_t) \right) \cdot \nabla_{\theta} V_{\theta}(s_t) \right].$$

Problem: in policy evaluation setting, we do not know  $v^{\pi}(s_t)$ . Hence, we estimate it using RL targets.

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Problem: in policy evaluation setting, we do not know  $v^{\pi}(s_t)$ . Hence, we estimate it using RL targets.

The simplest is the Monte Carlo target: estimate  $s_t$  by the discounted return of the sampled trajectory from  $s_t$ , i.e. perform updates of the form

$$\theta \leftarrow \theta + \alpha \cdot \left[ \left( G_t(s_t) - V_{\theta}(s_t) \right) \cdot \nabla_{\theta} V_{\theta}(s_t) \right].$$

# Gradient Monte Carlo policy evaluation: pseudocode

#### **Algorithm 3:** Gradient MC evaluation

**Input:** Policy  $\pi$ , FA  $V: \mathcal{S} \times \Theta \rightarrow \mathbb{R}$ , step size  $\alpha$ 

**Output:** Approximation  $V_{\theta}$  of  $v^{\pi}$ 

initialize  $\theta$  arbitrarily;

#### repeat

until timeout;

# Semi-gradient TD(0)

In the gradient update formula

$$\theta \leftarrow \theta + \alpha \cdot \left[ \left( \mathbf{v}^{\pi}(\mathbf{s}_{t}) - V_{\theta}(\mathbf{s}_{t}) \right) \cdot \nabla_{\theta} V_{\theta}(\mathbf{s}_{t}) \right].$$

we can also estimate  $v^{\pi}(s_t)$  with the TD(0) target:

$$\theta \leftarrow \theta + \alpha \cdot \left[ \left( \mathbf{r_{t+1}} + \gamma \cdot \mathbf{V_{\theta}(s_{t+1})} - V_{\theta}(s_t) \right) \cdot \nabla_{\theta} V_{\theta}(s_t) \right].$$

This yield the semi-gradient  $\mathsf{TD}(0)$  policy evaluation algorithm.

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This yield the semi-gradient TD(0) policy evaluation algorithm.

Why semi-gradient?

## Gradient vs. semi-gradient TD(0)

Recall that our ultimate goal is to minimize

$$\mathit{MSE}(v^\pi, V_\theta) = \frac{1}{2} \mathbb{E}_{s \sim \mu} \big[ (v^\pi(s) - V_\theta(s))^2 \big].$$

The gradient of this loss is

$$abla_{ heta} rac{1}{2} \mathbb{E}_{s \sim \mu} ig[ (v^{\pi}(s) - V_{ heta}(s))^2 ig] = rac{1}{2} \mathbb{E}_{s \sim \mu} ig[ 
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abla_{ heta} (v^\pi(s) - V_ heta(s))^2 ig],$$

Estimation with sample trajectory  $\tau = s_0, a_0, r_1, s_1, a_1, r_2, s_2, a_2, r_3, s_3, \ldots$  and substituting  $v^{\pi}(s)$ 

with the 
$$TD(0)$$
 target would yield

 $\nabla_{\theta} \textit{MSE}(v^{\pi}, V_{\theta}) \approx \frac{1}{2} \nabla_{\theta} (r_{t+1} + \gamma V_{\theta}(s_{t+1}) - V_{\theta}(s_{t}))^{2} = (r_{t+1} + \gamma V_{\theta}(s_{t+1}) - V_{\theta}(s_{t})) \cdot (\gamma \nabla_{\theta} V_{\theta}(s_{t+1}) - \nabla_{\theta} V_{\theta}(s_{t}))$ different update then semi-gradient TD(0)! However, this full gradient:

- is more expensive to compute (2 backpropagations per update);
- $\bullet$  does not really express TD(0) idea (the update target is not fixed).

# Semi-gradient TD(0): pseudocode

#### **Algorithm 4:** Semi-gradient TD(0) evaluation

```
Input: Policy \pi, FA V: \mathcal{S} \times \Theta \to \mathbb{R}, step size \alpha
Output: Approximation V_{\theta} of v^{\pi}
```

initialize  $\theta$  arbitrarily:

#### repeat

```
s \leftarrow \text{initial state};

a \sim \pi(s);

while s not terminal do

s' \sim p(s, a);

r \leftarrow r(s, a, s');

a' \sim \pi(s');

\theta \leftarrow \theta + \alpha \cdot [r + \gamma \cdot V(s', \theta) - V(s, \theta)] \cdot \nabla_{\theta} V(s, \theta);

s \leftarrow s'; a \leftarrow a'
```

until timeout;

## On-policy control with function approximation

Semi-gradient SARSA uses the same idea as TD(0), but with Q-approximator, i.e.

$$Q: \mathbb{R}^n \times \mathcal{A} \times \Theta \to \mathbb{R}$$
.

Behavior policy = e.g.  $\varepsilon$ -greedy with respect to the current Q. For a sampled trajectory  $\tau =$ 



we perform, in each timestep t, an update

$$\theta \leftarrow \theta + \alpha \cdot \left[ \left( r_{t+1} + \gamma \cdot Q_{\theta}(s_{t+1}, a_{t+1}) - Q_{\theta}(s_t, a_t) \right) \cdot \nabla_{\theta} Q_{\theta}(s_t, a_t) \right].$$

## Semi-gradient SARSA: pseudocode

### Algorithm 5: Semi-gradient SARSA

```
Input: FA Q: \mathcal{S} \times \mathcal{A} \times \Theta \rightarrow \mathbb{R}, step size \alpha
```

**Output:** Approximation  $Q_{\theta}$  of  $q^*$ 

initialize  $\theta$  arbitrarily;

#### repeat

```
\begin{split} s &\leftarrow \text{initial state;} \\ \pi &\leftarrow \text{policy } \varepsilon\text{-greedy w.r.t. } Q_{\theta}; \\ a &\sim \pi(s); \\ \text{while } s \text{ not terminal do} \\ & s' \sim p(s,a); \\ & r \leftarrow r(s,a,s'); \\ & a' \sim \pi(s'); \\ & \theta \leftarrow \theta + \alpha \cdot [r + \gamma \cdot Q_{\theta}(s',a') - Q_{\theta}(s,a)] \cdot \nabla_{\theta} Q_{\theta}(s,a); \\ & s \leftarrow s'; \ a \leftarrow a' \end{split}
```

until timeout;

### Representing actions in DNNs

How to represent actions in the (say, DNN) function approximator Q is largely a domain-dependent engineering choice.

If the set of actions  $\mathcal{A} = \{a^1, \dots, a^k\}$  is discrete and reasonably small, we can consider a net which inputs a state (i.e., n input neurons when  $\mathcal{S} = \mathbb{R}^n$ ) and outputs an  $|\mathcal{A}|$ -dimensional vector (i.e., one output neuron per action), so that the output of the i-th neuron on input s is interpreted as  $Q(s, a^i)$ .

I.e., in such a case we consider Q to be function of type  $Q \colon \mathcal{S} \times \Theta \to \mathbb{R}^{|\mathcal{A}|}$ .

### On-policy semi-gradient methods: concluding remarks

The presented algorithms can be instantiated also with other types of returns, such as:

- *n*-step returns
  - *n*-step SARSA update:

$$\theta \leftarrow \theta + \alpha \cdot \left[ \left( \underbrace{r_{t+1} + \gamma \cdot r_{t+2} + \dots + \gamma^{n-1} r_{t+n} + \gamma^n \cdot Q_{\theta}(s_{t+n}, a_{t+n})}_{G_{t:t+n}, \theta} - Q_{\theta}(s_t, a_t) \right) \cdot \nabla_{\theta} Q_{\theta}(s_t, a_t) \right]$$

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- forward-view  $\lambda$ -returns
  - SARSA( $\lambda$ ) update:  $\theta \leftarrow \theta + \alpha \cdot \left[ \left( (1 \lambda) \sum_{n=1}^{\infty} \lambda^{n-1} G_{t:t+n,\theta} Q_{\theta}(s_t, a_t) \right) \cdot \nabla_{\theta} Q_{\theta}(s_t, a_t) \right]$

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The presented algorithms can be instantiated also with other types of returns, such as:

- *n*-step returns
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- backward-view  $\lambda$ -returns (optional reading: Sutton&Barto, sections 12.2 and 12.7)

Off-Policy Control with

Value-Based

**Approximators:** 

**DQNs** and Friends

### Off-policy methods with function approximation

...are tricky to get right, already in the case of policy evaluation. The training can become very unstable.

For on-policy (semi)-gradient methods, one can typically prove convergence to correct/optimal values at least in the case of linear function approximation (though not in the more general case of NN approximators).

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For on-policy (semi)-gradient methods, one can typically prove convergence to correct/optimal values at least in the case of linear function approximation (though not in the more general case of NN approximators).

Off-policy semi-gradient methods, such as:

- TD with importance sampling (not covered here), or
- Q-learning with function approximators (will be covered a bit later),

can diverge already with linear function approximators.

### Divergence examples (high-level)

- Baird's counterexample: semi-gradient TD with importance sampling can diverge in presence of linear FAs
- Moreover, the divergence is not due to the instability of (semi)-gradient descent. Tsitsiklis and Van Roy's counterexample shows divergence even in the case where each update completely replaces the current  $\theta$  with the optimal  $\theta^*$  which minimizes the MSE between  $V_{\theta}$  and the TD(0) update target. The problem lies in the off-policy distribution of updates.
- Counterexamples explained in optional reading: Sutton&Barto, Sec. 11.2.

### **Deadly triad**

Identified by Sutton&Barto: risk of training instability and divergence steeply rises when combining:

- function approximation,
- bootstrapping, and
- off-policy training.

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Identified by Sutton&Barto: risk of training instability and divergence steeply rises when combining:

- function approximation,
- bootstrapping, and
- off-policy training.

But often we want to do just that. :)

Practical solution: Happily do the deadly triad, but use insights from supervised learning to develop additional techniques that help stabilize the training.

### Deep Q-Networks (DQN)

2013 arXiv tech. report, there is also follow-up 2015 Nature paper

### Playing Atari with Deep Reinforcement Learning

Volodymyr Mnih Koray Kavukcuoglu David Silver Alex Graves Ioannis Antonoglou

Daan Wierstra Martin Riedmiller

DeepMind Technologies

 $\{ \texttt{vlad}, \texttt{koray}, \texttt{david}, \texttt{alex}. \texttt{graves}, \texttt{ioannis}, \texttt{daan}, \texttt{martin}. \texttt{riedmiller} \} \texttt{ @ deepmind}. \texttt{com}$ 

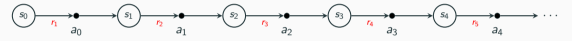
#### **Abstract**

We present the first deep learning model to successfully learn control policies directly from high-dimensional sensory input using reinforcement learning. The model is a convolutional neural network, trained with a variant of Q-learning, whose input is raw pixels and whose output is a value function estimating future rewards. We apply our method to seven Atari 2600 games from the Arcade Learning Environment, with no adjustment of the architecture or learning algorithm. We find that it outperforms all previous approaches on six of the games and surpasses a human expert on three of them.

### **Q-learning with function approximators**

Same semi-gradient idea as in TD(0), SARSA: adjust  $\theta$  to bring  $Q_{\theta}(s, a)$  closer to the fixed Q-learning update target.

I.e., for a sampled trajectory



and its timestep t, the update is

$$\theta \leftarrow \theta + \alpha \cdot \left[ \left( r_{t+1} + \gamma \cdot \max_{a \in \mathcal{A}(s_{t+1})} Q_{\theta}(s_{t+1}, a) - Q_{\theta}(s_t, a_t) \right) \cdot \nabla_{\theta} Q_{\theta}(s_t, a_t) \right].$$

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But performing the updates based solely on the current step would be susceptible to instability due to the presence of the deadly triad.

### **Deep Q-learning challenges**

From Mnih et al. Playing Atari with Deep Reinforcement Learning:

However reinforcement learning presents several challenges from a deep learning perspective. Firstly, most successful deep learning applications to date have required large amounts of hand-labelled training data. RL algorithms, on the other hand, must be able to learn from a scalar reward signal that is frequently sparse, noisy and delayed. The delay between actions and resulting rewards, which can be thousands of timesteps long, seems particularly daunting when compared to the direct association between inputs and targets found in supervised learning. Another issue is that most deep learning algorithms assume the data samples to be independent, while in reinforcement learning one typically encounters sequences of highly correlated states. Furthermore, in RL the data distribution changes as the algorithm learns new behaviours, which can be problematic for deep learning methods that assume a fixed underlying distribution.

### **Experience replay**

Originated in the work of Long-Ji Lin, e.g.: *Reinforcement Learning for Robots Using Neural Networks*, dissertation, 1993.

#### Definition 52: Experience

An experience is a 4-tuple  $(s, a, r, s') \in \mathcal{S} \times \mathcal{A} \times \mathcal{S} \times \mathbb{R}$  interpreted as ("state", "action played in it", "reward obtained", "next state observed").

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- DQN does not perform update based only on the current step. Instead, for each sampled trajectory  $\tau = s_0, a_0, r_1, s_1, a_1, r_2, s_2, a_2, r_3, s_3, \dots$  and each timestep t it:
  - first stores the one-step experience  $(s_t, a_t, r_{t+1}, s_{t+1})$  in a data structure  $\mathcal{B}$  called replay buffer;
  - then, sample a random minibatch of experiences  $B \subseteq \mathcal{B}$  of a given minibatch size Bsize
  - perform a minibatch-gradient-descent update w.r.t. B: compute the gradient of the Q-learning loss for each  $e \in B$  and then update  $\theta$  in the direction of an average gradient over the whole B.

### Minibatch update

Fix a minibatch B.

For each  $e = (s, a, r, s') \in B$  we compute the gradient of the Q-learning loss  $\nabla_{\theta} \mathcal{L}(\theta, e)$  at point e:

$$\nabla_{\theta} \mathcal{L}(\theta, e) = \nabla_{\theta} \frac{1}{2} \left( \underbrace{r + \gamma \cdot \max_{b \in \mathcal{A}(s')} Q_{\theta}(s', b)}_{\text{fixed target}} - Q_{\theta}(s, a) \right)^{2}$$

$$= \left[ \left( \underbrace{r + \gamma \cdot \max_{b \in \mathcal{A}(s')} Q_{\theta}(s', b)}_{=0 \text{ if s terminal}} - Q_{\theta}(s, a) \right) \cdot \nabla_{\theta} Q_{\theta}(s, a) \right]$$

We then perform an update in the direction of average gradient:

$$\theta \leftarrow \theta + \alpha \cdot \frac{1}{|B|} \sum_{e \in B} \nabla_{\theta} \mathcal{L}(\theta, e).$$

### **Experience replay rationale**

- helps decorrelate the DNN training data
- helps to prevent catastrophic forgetting
- improves data efficiency via experience re-use

Why good match for deep Q-learning? Experience replay is by design off-policy since we train on old data, which were sampled from different policy than the current one.

### Replay buffer implementation

The replay buffer  $\mathcal{B}$  is typically not unbounded, but has a fixed capacity  $\mathcal{B}$ size. Replacement is eventually needed. If  $\mathcal{B}$  is full, the oldest experience if removed ( $\mathcal{B} = \text{queue}$ ).

How to sample the minibatches?

- Original DQN: uniformly from B.
- Alternative: prioritized experience replay: each experience is assigned a priority (several heuristics exist). An experience is sampled into a minibatch with probability proportional to its priority.

### DQN: 2013 pseudocode

#### Algorithm 6: DQN with replay buffer

```
Input: Black-box MDP \mathcal{M} = (\mathcal{S}, \mathcal{A}, p, r), approximator Q; hyperparam's \varepsilon, \mathcal{B}size, Bsize, \dots
```

**Output:** Approximation  $Q_{\theta}$  of  $q^*$ 

initialize  $\theta$  arbitrarily; initialize empty replay buffer  $\mathcal B$  of capacity  $\mathcal B$ size;

#### repeat

```
\begin{split} s \leftarrow & \text{initial state;} \\ \textbf{while } s \text{ not terminal do} \\ & \pi \leftarrow & \text{policy } \varepsilon\text{-greedy w.r.t. } Q_{\theta}; \\ & a \sim \pi(s); \\ & s' \sim p(s,a); \\ & r \leftarrow r(s,a,s'); \\ & \text{store } (s,a,r,s') \text{ in } \mathcal{B}; \\ & \text{sample a minibatch } \mathcal{B} \text{ of size } \textit{Bsize from replay buffer } \mathcal{B}; \\ & \text{perform the minibatch update } \theta \leftarrow \theta + \alpha \cdot \frac{1}{\textit{Bsize}} \sum_{e \in \mathcal{B}} \nabla_{\theta} \mathcal{L}(\theta,e) \text{ (see this slide)}; \\ & s \leftarrow s'; \end{split}
```

until timeout;

### DQN: 2013 results

	B. Rider	Breakout	Enduro	Pong	Q*bert	Seaquest	S. Invaders
Random	354	1.2	0	-20.4	157	110	179
Sarsa [3]	996	5.2	129	-19	614	665	271
Contingency [4]	1743	6	159	-17	960	723	268
DQN	4092	168	470	20	1952	1705	581
Human	7456	31	368	-3	18900	28010	3690
HNeat Best [8]	3616	52	106	19	1800	920	1720
HNeat Pixel [8]	1332	4	91	-16	1325	800	1145
DQN Best	5184	225	661	21	4500	1740	1075

Table 1: The upper table compares average total reward for various learning methods by running an  $\epsilon$ -greedy policy with  $\epsilon=0.05$  for a fixed number of steps. The lower table reports results of the single best performing episode for HNeat and DQN. HNeat produces deterministic policies that always get the same score while DQN used an  $\epsilon$ -greedy policy with  $\epsilon=0.05$ .

Mnih et al. (2013, arXiv)

### Target networks: another stabilizing factor in DQNs

Introduced in the reviewed version of DQN paper:

Mnih et al.: Human-level control through deep reinforcement learning. *Nature*, 518 (2015).

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Performing the (minibatch) Q-update looks like supervised learning:

$$\text{change} \quad \theta \quad \text{ so that } \quad Q_{\theta}(s,a) \quad \text{ gets closer to the fixed target} \quad r + \gamma \cdot \max_{b \in \mathcal{A}(s')} Q_{\theta}(s',b),$$

where (s, a, r, s') is the processed experience.

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Performing the (minibatch) Q-update looks like supervised learning, but:

change 
$$\theta$$
 so that  $Q_{\theta}(s,a)$  gets closer to the fixed target  $r + \gamma \cdot \max_{b \in \mathcal{A}(s')} \underbrace{Q_{\theta}(s',b)}_{b \in \mathcal{A}(s')}$ , the "label" changes with each update! where  $(s,a,r,s')$  is the processed experience.

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### Target networks: idea

To stabilize learning, we use two networks: the main network, and the target network. They have the same architecture (denote it by Q), but their weights may differ during the execution of the algorithm.

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- $\bullet$   $\theta$  weights of main network
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### Usage:

• The target network is used only to compute TD targets when computing losses:

$$\nabla_{\theta} \mathcal{L}(\theta, e) = \left(r + \gamma \cdot \max_{b \in \mathcal{A}(s')} Q(s', b, \hat{\theta}) - Q(s, a, \theta)\right) \cdot \nabla_{\theta} Q(s, a, \theta)$$

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$$abla_{ heta} \mathcal{L}( heta, e) = \left(r + \gamma \cdot \max_{b \in A(s')} Q(s', b, \hat{\theta}) - Q(s, a, \theta)\right) \cdot \nabla_{\theta} Q(s, a, \theta)$$

• At the start, and also in periodic intervals (but not after each update!) the two networks are synchronized by performing  $\hat{\theta} \leftarrow \theta$ . Other than this,  $\hat{\theta}$  stays fixed, the gradient steps are only used to update  $\theta$  (i.e., the main network).

### DQN: 2015 pseudocode

#### Algorithm 7: DQN with replay buffer and target network

```
Input: Black-box MDP \mathcal{M} = (\mathcal{S}, \mathcal{A}, p, r), approximator Q; hyperparam's \varepsilon, \mathcal{B}size, \mathcal{B}size, \mathcal{C},...
Output: Approximation Q_{\theta} of a^*
initialize \theta arbitrarily: \hat{\theta} \leftarrow \theta; counter \leftarrow C:
initialize empty replay buffer \mathcal{B} of capacity \mathcal{B}size:
repeat
     s \leftarrow \text{initial state}:
     while s not terminal do
           if counter = 0 then \hat{\theta} \leftarrow \theta; counter \leftarrow C else counter \leftarrow counter - 1;
           \pi \leftarrow \text{policy } \varepsilon\text{-greedy w.r.t. } Q_{\theta}:
           a \sim \pi(s):
           s' \sim p(s, a):
           r \leftarrow r(s, a, s'):
           store (s, a, r, s') in \mathcal{B};
```

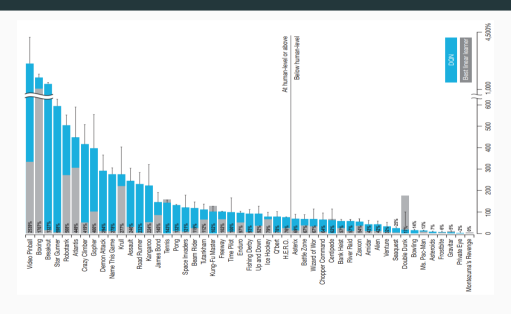
perform the minibatch update  $\theta \leftarrow \theta + \alpha \cdot \frac{1}{B \text{size}} \sum_{e \in B} \nabla_{\theta} \mathcal{L}(\theta, e)$ , where  $\nabla_{\theta} \mathcal{L}(\theta, e) = (r + \gamma \cdot \max_{b \in \mathcal{A}(s')} Q(s', b, \hat{\theta}) - Q(s, a, \theta)) \cdot \nabla_{\theta} Q(s, a, \theta)$ ;

sample a minibatch B of size Bsize from replay buffer B;

 $s \leftarrow s'$ :

until timeout;

### DQN: 2015 results



## Engineering behind DQN for Atari: partial observability

True state = current state (program counter + variable values) of the game program.

We do not see this – only the frames rendered on screen.

Solving partially observable environments requires (per some POMDP theory) making decisions based on the whole history of observations. This is computationally demanding (recurrent NNs...).

## Engineering behind DQN for Atari: partial observability

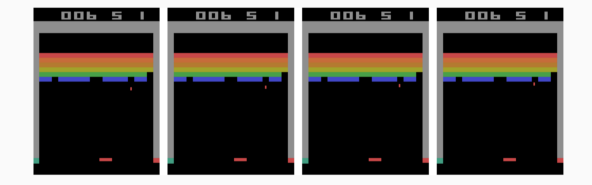
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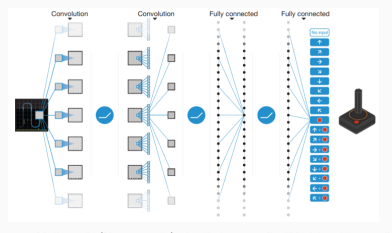
DQN for Atari solves this by feeding the last 4 observed frames into the NN. This is typically enough to deduce the dynamics of the current play.

### **DQN:** dynamics from limited frame history



### Engineering behind DQN for Atari: the network

Inputs four 84x84px images, then 3 convolutional layers, then two fully connected layers. all with ReLU activations. Outputs Q-estimate for each action.



source: Mnih et al. (Nature, 2015), details in appendix "Model architecture"

### DQN for Atari: dirty engineering tricks

- Preprocessing: 210x160 RGB color images are converted to grayscale and resized to 84x84 resolution.
- Frame skipping: the agent only observes and acts in every K-th frame, for the frames in between, the last selected action is repeated without providing the frame to the agent. (In the paper, K = 4.)
- Reward clipping: all positive one-step Atari rewards are clipped to +1, all negative ones are clipped to -1 (Atari gives integer rewards).
- TD error clipping: for each update, the Q-learning error  $r + \gamma \cdot \max b \in \mathcal{A}(s')Q(s',b) Q(s,a)$  is clipped to [-1,1].

# DQN for Atari: selected hyperparameters (nex)

minibatch size Bsize	32
replay buffer size <i>Bsize</i>	1,000,000
target network update frequency $C$	10,000
discount factor $\gamma$	0.99
update frequency (steps between two minibatch updates)	4
learning rate	0.00025
initial $arepsilon$	1
final $arepsilon$ (linear decay)	0.1
final decay frame	1,000,000
random policy played for init.	50,000 frames
max. do-nothing actions at episode start	30

### **RAINBOW** of heuristics

Multitude of heuristics for the improvement of DQN were developed over time. Some of them make sense also in the context of other deep RL algorithms.

The RAINBOW agent combines six such heuristics to further improve the DQN performance on Atari games.

Rainbow: Combining Improvements in Deep Reinforcement Learning				
Matteo Hessel	Joseph Modayil	Hado van Hasselt	Tom Schaul	Georg Ostrovski
DeepMind	DeepMind	DeepMind	DeepMind	DeepMind
Will Dabney	Dan Horgan	Bilal Piot	Mohammad Azar	David Silver
DeepMind	DeepMind	DeepMind	DeepMind	DeepMind

(In proceedings of AAAI 2018.)

### Rainbow heuristics

- dueling networks architecture
- double DQN
- prioritized experience replay
- n-step rewards
- distributional learning
- noisy networks

### **Action advantage**

Idea: imagine that for some state s, the Q-values of all actions are high. Then s should be in some sense valuable in itself.

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### Definition 53: Advantage

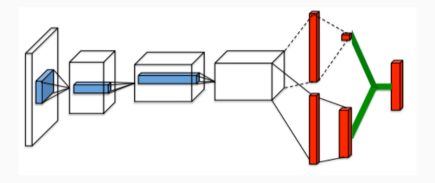
Let  $\pi$  be a policy. An advantage function  $adv^{\pi} \colon \mathcal{S} \times \mathcal{A} \to \mathbb{R}$  is defined as

$$adv^{\pi}(s,a)=q^{\pi}(s,a)-v^{\pi}(s).$$

Dueling architecture splits certain layers of the neural network into two "streams", one estimating (something like)  $v^{\pi}(s)$  and one estimating (something like)  $adv^{\pi}(s,a)$ . The final layer combines these estimates to produce an estimate of  $q^{\pi}(s,a)$ .

## **Dueling architecture**

Wang et al.: *Dueling Network Architectures for Deep Reinforcement Learning*. In proceedings of ICML'16.



# Dueling architecture: theory and training

We have 
$$Q_{\theta,\alpha,\beta}(s,a) = aggregate(V_{\theta,\alpha}(s), A_{\theta,\beta}(s,a))$$
, where

- ullet heta convolutional (or other feature extraction) layer parameters
- ullet  $\alpha$  value channel parameters
- $\bullet \;\; \beta$  advantage channel parameters

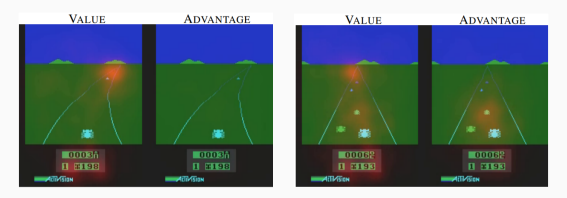
# Dueling architecture: theory and training

We have  $Q_{\theta,\alpha,\beta}(s,a) = aggregate(V_{\theta,\alpha}(s),A_{\theta,\beta}(s,a))$ , where

- ullet heta convolutional (or other feature extraction) layer parameters
- ullet lpha value channel parameters
- ullet eta advantage channel parameters

The whole network Q is trained to estimate  $q^{\pi}$  (where  $\pi$  is the target policy) using any deep RL algorithm (e.g. DQN, in which case  $\pi$  is the optimal policy). There is nothing new from RL perspective here, all the novelty is inside the network. The factorization into state value and advantage is supposed to help the network "focus" on features that are important to recognize valuable states and features that help us rank actions.

## **Dueling architecture: Atari example**



From Wang et al.: *Dueling Network Architectures for Deep Reinforcement Learning*. In proceedings of ICML'16.

- The whole net is trained end-to-end to predict  $q^{\pi}$ .
- How do we ensure the value/advantage channels are trained to predict state values/advantages?

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- By a suitable choice of aggregator:
  - $Q_{\theta,\alpha,\beta}(s,a) = V_{\theta,\alpha}(s) + A_{\theta,\beta}(s,a)$  does not work: e.g.  $V_{\theta,\alpha}$  could converge to constant 0 and  $A_{\theta,\beta}$  to  $q^{\pi}$ .

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$$Q_{ heta,lpha,eta}(s,a) = V_{ heta,lpha}(s) + A_{ heta,eta}(s,a) - \max_{b\in\mathcal{A}(s)}A_{ heta,eta}(s,b),$$

the training then indeed pushes  $V_{\theta,\alpha}$  to  $v^{\pi}$  and  $A_{\theta,\beta}$  to  $adv^{\pi}+c$  where c is some constant.

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the training then indeed pushes  $V_{\theta,\alpha}$  to  $v^{\pi}$  and  $A_{\theta,\beta}$  to  $adv^{\pi}+c$  where c is some constant. Issues: not differentiable, update sensitive to the value of maximizing action changes.

## **Aggregation in Rainbow**

The point: aggregating layer should anchor the sum of the channels to same baseline value derived non-trivially from the advantages (if all advantages shift up/down, so should the baseline). Rainbow uses mean advantage baseline:

$$Q_{ heta,lpha,eta}(s,a) = V_{ heta,lpha}(s) + A_{ heta,eta}(s,a) - rac{1}{|\mathcal{A}(s)|} \sum_{b \in \mathcal{A}(s)} A_{ heta,eta}(s,b)$$

pushes the value channel to predict  $\frac{1}{|\mathcal{A}(s)|} \sum_{a \in \mathcal{A}(s)} q^{\pi}(s, a)$ .

### Double DQN

van Hasselt, Guez, Silver: *Deep Reinforcement Learning with Double Q-learning*. In proceedings of AAAI 2016.

Similar idea to tabular Double Q-learning (use different estimates for selecting maximizing action in bootstrap and for evaluating the bootstrap), but instead of independently updated networks uses main and target networks. I.e., for experience (s, a, r, s'), the update is:

$$\theta \leftarrow \theta + \alpha \cdot [r + \gamma \cdot Q_{\hat{\theta}}(s', \arg \max_{b \in \mathcal{A}(s')} Q_{\theta}(s', b)) - Q_{\theta}(s, a)] \nabla_{\theta} Q_{\theta}(s, a),$$

where  $\hat{\theta}$  is the parameter vector of the target network.

## Prioritized experience replay (Schaul et al., ICLR'16)

Each experience e=(s,a,r,s') in the replay buffer is assigned a priority according to its TD-error

$$p_{e} = |r + \gamma \cdot \max_{b \in \mathcal{A}(s')} Q_{\hat{\theta}}(s', b) - Q_{\theta}(s, a)| + \varepsilon$$

 $(\varepsilon > 0$  ensures all priorities are positive).

The probability of sampling an experience e from the buffer is set to  $\frac{p_e^{\alpha}}{\sum_{e' \in \mathcal{B}} p_{e'}^{\alpha}}$ , where  $\alpha > 0$  is a hyperparameter controlling the degree of prioritization.

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The probability of sampling an experience e from the buffer is set to  $\frac{p_e^{\alpha}}{\sum_{e' \in \mathcal{B}} p_{e'}^{\alpha}}$ , where  $\alpha > 0$  is a hyperparameter controlling the degree of prioritization.

Prioritization induces bias: the sampled experiences no longer follow the same distribution as sampled trajectories. We can correct this by using importance sampling during updates:

$$\theta \leftarrow \theta + \alpha \cdot \left(\frac{1}{|\mathcal{B}|} \cdot \frac{1}{p_e^{\alpha}}\right)^{\beta} \cdot \left[r + \gamma \cdot Q_{\hat{\theta}}(s', \operatorname{arg\,max}_{b \in \mathcal{A}(s')} Q_{\theta}(s', b)) - Q_{\theta}(s, a)\right] \cdot \nabla_{\theta} Q_{\theta}(s, a),$$

where  $\beta > 0$  determines the degree of IS correction (annealed to 1 during training).

### n-step returns

Self-explanatory, use n-step return with Q-learning bootstrap when computing TD target.

How to combine with replay buffer? Each experience stores a single step.

### n-step returns

Self-explanatory, use *n*-step return with Q-learning bootstrap when computing TD target.

How to combine with replay buffer? Each experience stores a single step.

### Solutions:

- Store experiences in  $\mathcal{B}$  sequentially, with each sampled experience, retrieve also the next n-1 ones (up to episode termination). Requires careful implementation.
- Naive: each element of  $\mathcal{B}$  consists of n consecutive experiences (space inefficient).

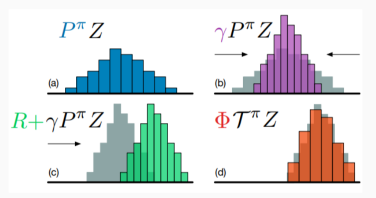
Given consecutive experiences

$$(s_t, a_t, r_{t+1}, s_{t+1}), (s_{t+1}, a_{t+1}, r_{t+2}, s_{t+2}), \ldots, ((s_{t+n-1}, a_{t+n-1}, r_{t+n}, s_{t+n})), \text{ perform update } s_t, s_{t+1}, s_{t+$$

$$\theta \leftarrow \theta + \alpha \cdot \left[ r_{t+1} + \gamma r_{t+2} + \dots + \gamma^{n-1} r_{t+n} + \gamma^n \max_{b \in \mathcal{A}(s_{t+n})} Q_{\hat{\theta}}(s_{t+n}, b) - Q_{\theta}(s_t, a_t) \right] \nabla_{\theta} Q_{\theta}(s_t, a_t).$$

## **Distributional learning**

Very rough idea: instead of expected returns, predict (discretized) distribution of returns.



Source: Bellemare, Dabney, Munos: *A Distributional Perspective on Reinforcement Learning*. In proceedings of ICML'17.

We still optimize the expected value of the distribution, but the NN processes richer information (neurological inspiration).

### Noisy nets

### Fortunato et al.: Noisy Networks for Exploration . In proceedings of ICRL'17.

An alternative way of achieving exploration (without  $\varepsilon$ -greedy policies). Replaces linear layers  $y = W \cdot x + b$  with noisy layers of the form

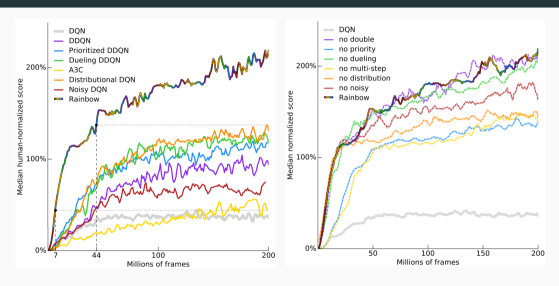
$$y = (\mu_w + \sigma_w \odot \varepsilon_w) \cdot x + \mu_b + \sigma_b \odot \varepsilon_b,$$

where matrices  $\mu_w, \sigma_w$  and vectors  $\mu_b, \sigma_b$  are learnable, matrix  $\varepsilon_w$  and vector  $\varepsilon_b$  consist of random noise, and  $\odot$  represents component-wise multiplication.

The loss function of the DQN training is then encapsulated in expectation over the noise.

Interesting point: the net can learn to adjust  $\sigma$ 's and thus the degree of exploration over time.

## Rainbow: evaluation and ablations (Hessel et al., 2017)



**Policy Gradient Methods** 

# Value-based vs. policy-based methods

So far: focus on approximating  $q^*$  via some parameterized estimate  $Q_\theta$ , policy the defined by  $Q_\theta$ , e.g.  $(\varepsilon$ -)-greedy...

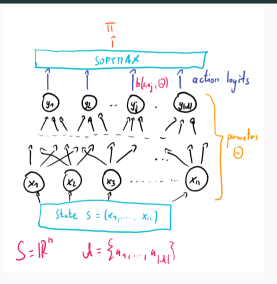
## Value-based vs. policy-based methods

So far: focus on approximating  $q^*$  via some parameterized estimate  $Q_\theta$ , policy the defined by  $Q_\theta$ , e.g. ( $\varepsilon$ -)-greedy...

In policy gradient methods we work directly with some parameterized representation of a policy  $\pi_{\theta}$ , and update  $\theta$  so as to improve some performance characteristic of  $\pi_{\theta}$  (e.g., expected return).

In particular, the policy  $\pi$  can be represented by a function approximator  $\pi_{\theta} \colon \mathcal{S} \times \Theta \to \mathcal{D}(\mathcal{A})$ .

## Standard NN + softmax representation



$$\pi_{ heta}(a|s) = rac{e^{h(s,a, heta)}}{\sum_{b \in \mathcal{A}} e^{h(s,b, heta)}},$$

where  $h(s, a, \theta)$  is the logit ("preference") of action a

## General policy gradient scheme

We want to find  $\theta$  that maximizes some performance measure (or objective function)  $J(\theta)$  of  $\pi_{\theta}$ . The obvious choice for J is the expected return:

$$J(\theta) = v^{\pi_{\theta}} = \mathbb{E}^{\pi_{\theta}}[G],$$

thought some algorithms use a different surrogate objective function.

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$$J(\theta) = v^{\pi_{\theta}} = \mathbb{E}^{\pi_{\theta}}[G],$$

thought some algorithms use a different surrogate objective function.

The optimization problem

$$\max_{\theta} J(\theta)$$

can be (locally) solved using gradient ascent: repeatedly perform updates

$$\theta \leftarrow \theta + \alpha \cdot \nabla_{\theta} J(\theta)$$
.

It is thus necessary to compute or approximate  $\nabla_{\theta} J(\theta)$ : this is the scope of various policy gradient theorems.

# Gradient of expected return: possible version

$$abla_{ heta} J( heta) = 
abla_{ heta} \mathbb{E}^{\pi_{ heta}}[G] =$$

# Gradient of expected return (cont'd)

## Vanilla MC policy gradient

### Algorithm 8: Vanilla MC policy gradient

**Input:** Black-box MDP  $\mathcal{M} = (\mathcal{S}, \mathcal{A}, p, r)$ , policy parametrization  $\pi_{\theta}$ , learning rate  $\alpha$ 

**Output:** Approximation  $\pi_{\theta}$  of  $\pi^*$ 

initialize  $\theta$  arbitrarily;

### repeat

```
s_0 \sim initial distribution; generate episode \tau = s_0, a_0, r_1, s_1, a_1, r_2, s_2, a_2, r_3, s_3, \dots, r_T, s_T using \pi_{\theta}; \theta \leftarrow \theta + \alpha \cdot G(\tau) \cdot \sum_{t=0}^{i} \nabla_{\theta} \log \pi_{\theta}(a_j|s_j)
```

until timeout;

## **Score function form**

# Step-wise gradient

$$\nabla_{\theta} J(\theta) = \nabla_{\theta} \mathbb{E}^{\pi_{\theta}}[G] = \nabla_{\theta} \mathbb{E}^{\pi_{\theta}}[\sum_{t=0}^{\infty} \gamma^{t} R_{t+1}]$$

# Step-wise gradient (cont'd)

### REINFORCE: better MC policy gradient

### Algorithm 9: REINFORCE (Williams, 1992)

**Input:** Black-box MDP  $\mathcal{M} = (\mathcal{S}, \mathcal{A}, p, r)$ , policy parametrization  $\pi_{\theta}$ , learning rate  $\alpha$ 

**Output:** Approximation  $\pi_{\theta}$  of  $\pi^*$ 

initialize  $\theta$  arbitrarily;

### repeat

### until timeout;

# Baseline in policy gradient

### Theorem 54: Baseline theorem

Let  $b_{\theta}(s) \colon \mathcal{S} \times \theta \to \mathbb{R}$  be any function. Then for any t:

$$\mathbb{E}^{\pi_{ heta}}ig[b_{ heta}(s_t)\cdot
abla_{ heta}log\pi_{ heta}(a_t|s_t)ig]=0.$$

As a consequence

$$egin{aligned} 
abla_{ heta} J( heta) &= \mathbb{E}^{\pi_{ heta}} ig[ \sum_{t=0}^{T} \gamma^{t} \cdot G_{t} \cdot 
abla_{ heta} \log \pi_{ heta}(a_{t}|s_{t}) ig] = \ &= \mathbb{E}^{\pi_{ heta}} ig[ \sum_{t=0}^{T} \gamma^{t} \cdot ig( G_{t} - b_{ heta}(s_{t}) ig) \cdot 
abla_{ heta} \log \pi_{ heta}(a_{t}|s_{t}) ig] \end{aligned}$$

The gradient estimates using baseline have the same expectations as the standard REINFORCE estimate, but might have a lower variance if baseline selected correctly.

### **State-value baseline**

Good choice is

$$b(s):=v^{\pi_{\theta}}(s),$$

reducing the estimate variance by correcting the return for a bias caused by being in a certain state.

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Problem: we do not know  $v^{\pi_{\theta}}$ .

#### State-value baseline

Good choice is

$$b(s):=v^{\pi_{\theta}}(s),$$

reducing the estimate variance by correcting the return for a bias caused by being in a certain state.

Problem: we do not know  $v^{\pi_{\theta}}$ .

Solution: Learn  $v^{\pi_{\theta}}$  online using a separate function approximator V, e.g. via gradient (every-visit) Monte Carlo.

#### REINFORCE with a state-value baseline

#### **Algorithm 10:** REINFORCE with baseline

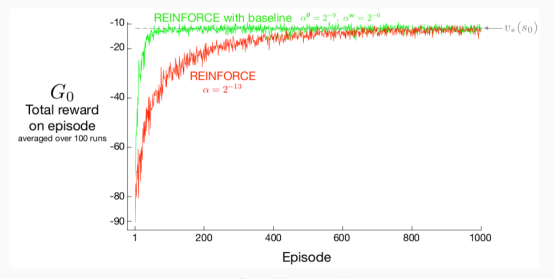
**Input:** Black-box MDP  $\mathcal{M}=(\mathcal{S},\mathcal{A},p,r)$ , policy parametrization  $\pi_{\theta}$ , value parametrization  $V_{\eta}$ , learning rates  $\alpha_{\pi}$ ,  $\alpha_{V}$  for the two approximators

**Output:** Approximation  $\pi_{\theta}$  of  $\pi^*$  initialize  $\theta$  and  $\eta$  arbitrarily;

#### repeat

until timeout;

## REINFORCE: baseline effect experiment



source: Sutton&Barto, p. 330

Comparing returns to some state-dependent baseline is reminiscent of what happened in the dueling network architecture.

Comparing returns to some state-dependent baseline is reminiscent of what happened in the dueling network architecture.

In particular, one can derive another form of the policy gradient, showing that

$$abla_{ heta} J( heta) \propto \mathbb{E}_{s \sim \pi, a \sim \pi_{ heta}}[q^{\pi_{ heta}}(s, a) \cdot 
abla_{ heta} \log \pi(a|s)].$$

Comparing returns to some state-dependent baseline is reminiscent of what happened in the dueling network architecture.

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abla_{ heta} \log \pi(a|s)].$$

Inputting a state-value baseline yields

$$abla_{ heta} J( heta) \propto \mathbb{E}_{s \sim \pi, a \sim \pi_{ heta}} [\underbrace{(q^{\pi_{ heta}}(s, a) - v^{\pi}(s))}_{ ext{ad}v^{\pi}(s, a)} \cdot 
abla_{ heta} \log \pi(a|s)].$$

Comparing returns to some state-dependent baseline is reminiscent of what happened in the dueling network architecture.

In particular, one can derive another form of the policy gradient, showing that

$$\nabla_{\theta} J(\theta) \propto \mathbb{E}_{s \sim \pi, a \sim \pi_{\theta}}[q^{\pi_{\theta}}(s, a) \cdot \nabla_{\theta} \log \pi(a|s)].$$

Inputting a state-value baseline yields

$$abla_{ heta} J( heta) \propto \mathbb{E}_{s \sim \pi, a \sim \pi_{ heta}} [\underbrace{(q^{\pi_{ heta}}(s, a) - v^{\pi}(s))}_{ ext{ad}v^{\pi}(s, a)} \cdot 
abla_{ heta} \log \pi(a|s)].$$

Hence, policy-gradient-type algorithms often formulate the update of policy parameters in the form

$$\theta \leftarrow \theta + \alpha \cdot A_{\theta,n}^t(s_t, a_t) \cdot \nabla_{\theta} \log \pi_{\theta}(a_t|s_t),$$

where  $A_{\theta,\eta}^t$  is some approximator called advantage estimate. E.g. in PG with baseline,  $A_{(\theta,\eta)}^t = \gamma^t \cdot (G_t - V_{\eta}(s_t))$ .

### **Proof of Baseline theorem**

# Actor-critic: Policy gradient with bootstrapping

Recall the REINFORCE-with-baseline update:

$$\theta \leftarrow \theta + \alpha \cdot \gamma^t \cdot (G_t - V_{\eta}(s_t)) \cdot \nabla_{\theta} \log \pi_{\theta}(a_t|s_t)$$

 $G_t$  is a possible source of variance: let's remove it (at the cost of introducing bias) via bootstrapping! E.g. TD(0):

$$\theta \leftarrow \theta + \alpha \cdot \gamma^t \cdot (r_t + \gamma \cdot V_{\eta}(s_{t+1}) - V_{\eta}(s_t)) \cdot \nabla_{\theta} \log \pi_{\theta}(a_t | s_t).$$

Here, the value network  $V_{\eta}$  both estimates the baseline value of the current state, and (via boostrap) the quality of the played action: we call it a critic, while the policy network  $\pi_{\theta}$  is called an actor.

Note: we can use the same bootstrap to update critic parameters.

# Basic Actor-Critic (AC) algorithm: pseudocode

#### Algorithm 11: One-step AC

**Input:** Black-box MDP  $\mathcal{M} = (\mathcal{S}, \mathcal{A}, p, r)$ , policy parametrization  $\pi_{\theta}$ , value parametrization  $V_{\eta}$ , learning rates  $\alpha_{\pi}$ ,  $\alpha_{V}$  for the two approximators

**Output:** Approximation  $\pi_{\theta}$  of  $\pi^*$ 

initialize  $\theta$  and  $\eta$  arbitrarily;

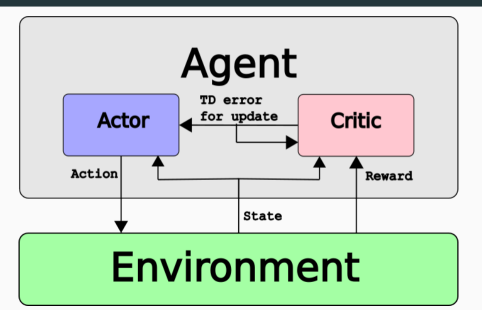
#### repeat

until timeout:

```
\begin{split} s &\sim \text{initial distribution;} \\ D &\leftarrow 1; \\ \textbf{while } s \text{ is not terminal do} \\ & | \quad a \sim \pi_{\theta}(s); \\ s' &\sim p(s,a); \\ \delta &\leftarrow r(s,a,s') + \gamma \cdot V_{\eta}(s') - V_{\eta}(s); \\ \eta &\leftarrow \eta + \alpha_{V} \cdot \delta \cdot \nabla_{\eta} V_{\eta}(s_{t}); \\ \theta &\leftarrow \theta + \alpha_{\pi} \cdot D \cdot \delta \cdot \nabla_{\theta} \log \pi_{\theta}(a_{t}|s_{t}); \\ D &\leftarrow \gamma \cdot D; \\ s &\leftarrow s' \end{split}
```

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# **AC:** picture



#### Comments on AC

• The algorithm presented on previous slide is just a basic variant: actor-critic framework covers a wide range of algorithms: soft Actor-Critic (SAC), A2C, A3C, deep deterministic policy gradient (DDPG), twin-delayed DDPG (TD3), PPO (next time),...

- Entropy regularization: add to the objective function a term representing the entropy of the policy: we prefer "more" randomized policies to encourage exploration (used e.g. in SAC).
  - $J_{ENTR}(\theta) = \mathbb{E}^{\pi}[G] + \beta \cdot \mathbb{E}_{s \sim \pi_{\theta}}[H(\pi(s))]$

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- Using parallel agents whose gradients are averaged for each update (A2C).

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- Off-policy training by using replay buffer (e.g. in SAC, DDPG, TD3).
- Using parallel agents whose gradients are averaged for each update (A2C).
- Using *n*-step (e.g. A2C) or  $\lambda$ -returns in bootstrap (PPO).
- Using different  $J(\theta)$  than just expected return (SAC, TRPO, PPO).

Note that the above algorithms typically differ from "vanilla" AC in more aspects then presented above. We shall see soon on the case of PPO.

**Taming Unstable Gradients with** 

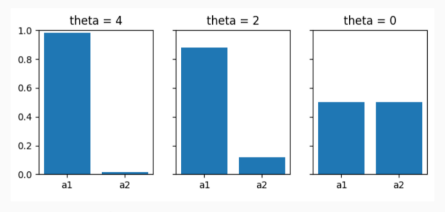
Trust Regions: TRPO, PPO

### Limitations of policy gradient methods

- Basic policy gradient methods are prone to large variance of gradient estimates.
- These can be mitigated to some degree, e.g. by using n-step or  $\lambda$ -returns in advantage estimation.
- Even then, the methods can yield large updates which can destabilize the training. We are never sure the parameter updates will actually lead to an improvement of a policy:

# Sensitivity of policy to parameters

For two actions and 
$$\pi(a|s) = \begin{cases} \frac{1}{1-e^{-\theta}} & a = a_1 \\ 1 - \frac{1}{1-e^{-\theta}} & a = a_2, \end{cases}$$
 we get



Source: slides by Emma Brunskill, Lecture 6, https://web.stanford.edu/class/cs234/modules.html

# Direct policy improvement via surrogate objectives

- We will present algorithms that use different performance metric so as increase the chance of policy improvement on update.
- Moreover, we will design the performance metric so that its gradient can be easily computed by an automated differentiation tool without the need to derive the formulas manually.

### Overall structure of presented algorithms

- The algorithms will generate a sequence of policies  $\pi_1$ ,  $\pi_2$ ,  $\pi_3$  such that each policy will be (likely) an improvement over the previous one.
- Each step from  $\pi_i$  to  $\pi_{i+1}$  entails finding a policy  $\pi_{i+1}$  that optimizes some performance metric  $\mathcal{L}_{\pi_i}$  that is dependent on the previous policy  $\pi_i$ ! (Cf. standard policy gradient: the performance metric J evaluated only the current policy).
- I.e. a run of such an algorithm entails solving (using gradient-based methods) multiple optimization problems: one per each policy update.

- We will now focus on the single improvement step: we will denote by
  - $\pi = \pi_{\theta}$  the previous policy  $(\pi_i)$
  - $\pi' = \pi_{\theta'}$  the new policy  $(\pi_{i+1})$  we seek.

 $\theta$  is treated as a constant,  $\theta'$  as variables!

# Roles of advantages in policy improvement

#### Theorem 55

Let  $\theta, \theta'$  be two parameter vectors and  $\pi = \pi_{\theta}, \pi' = \pi_{\theta'}$ .

Then

$$J(\theta') - J(\theta) = \underbrace{\mathbb{E}_{\tau \sim \pi'} \Big[ \sum_{t=0}^{\infty} \gamma^t \cdot \mathsf{adv}^{\pi}(s_t, a_t) \Big]}_{\stackrel{\mathsf{def}}{=} \mathcal{L}_{\pi}(\pi')}.$$

## Another loss surrogate

To ensure that update from  $\theta$  to  $\theta'$  is an improvement, we want to maximize

$$\mathcal{L}_{\pi}(\pi') = \mathbb{E}_{ au \sim \pi'} \Big[ \sum_{t=0}^{\infty} \gamma^t \cdot \mathsf{adv}^{\pi}(s_t, a_t) \Big]$$

But we cannot sample from  $\pi'$ , neither can we easily compute the gradient of the loss by automated differentiation.

Trick: the loss function  $\mathcal{L}_{\pi}$  behaves similarly to the following loss function  $\widetilde{\mathcal{L}}_{\pi}$  for all points  $\pi'$  that are "close enough" to  $\pi$ :

$$\widetilde{\mathcal{L}}_{\pi}(\pi') = \mathbb{E}_{ au \sim \pi} \Big[ \sum_{t=0}^{\infty} \gamma^t \cdot \mathsf{adv}^{\pi}(s_t, a_t) \cdot \frac{\pi'(a_t|s_t)}{\pi(a_t|s_t)} \Big]$$

.

# Closeness of $\mathcal{L}_{\pi}$ and $\widetilde{\mathcal{L}}_{\pi}$

$$\mathcal{L}_{\pi}(\pi) = \mathbb{E}_{\tau \sim \pi'} \Big[ \sum_{t=0}^{\infty} \gamma^{t} \cdot \mathsf{adv}^{\pi}(\mathsf{s}_{t}, \mathsf{a}_{t}) \Big] \qquad \qquad \widetilde{\mathcal{L}}_{\pi} = \mathbb{E}_{\tau \sim \pi} \Big[ \sum_{t=0}^{\infty} \gamma^{t} \cdot \mathsf{adv}^{\pi}(\mathsf{s}_{t}, \mathsf{a}_{t}) \cdot \frac{\pi'(\mathsf{a}_{t}|\mathsf{s}_{t})}{\pi(\mathsf{a}_{t}|\mathsf{s}_{t})} \Big]$$

"Behaves similarly" can be formalized as follows:

#### Theorem 56

It holds  $\mathcal{L}_{\pi}(\pi) = \widetilde{\mathcal{L}}_{\pi}(\pi)$ .

Moreover the gradients  $\nabla_{\theta'}\mathcal{L}_{\pi}$  and  $\nabla_{\theta'}\widehat{\mathcal{L}}_{\pi}$  are equal at point  $\pi$ .

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Proof: the first part is trivial. The second part is technical and requires converting the expectation into expectation-over-states form, see

Optional reading: Schulman, Levine, Abbeel, Jordan, Moritz: *Trust Region Policy Optimization*. In Proceedings of ICML'15.

## **Trust Region Methods**

Hence, the constrained optimization problems

maximize 
$$\mathcal{L}_{\pi}(\pi')$$
 subject to  $\pi'$  close to  $\pi$  maximize  $\widetilde{\mathcal{L}}_{\pi}(\pi')$  subject to  $\pi'$  close to  $\pi$ 

have approximately the same optimal solutions. We want to solve the first, and will proceed by solving the second.

The set of  $\pi'$  that are "close enough" to  $\pi$  is called a trust region. What "close enough" means differs among algorithms. Exact bounds on the error of the approximation in terms of KL divergence between  $\pi$  and  $\pi'$  was given in

Optional reading: Achiam, Held, Tamar, Abbeel: *Constrained Policy Optimization*. In Proceedings of ICML'17.

# Optimizing the loss by sampling and automated differentiation

$$\widetilde{\mathcal{L}}_{\pi} = \mathbb{E}_{\tau \sim \pi} \Big[ \sum_{t=0}^{\infty} \gamma^{t} \cdot \mathsf{adv}^{\pi}(s_{t}, a_{t}) \cdot \frac{\pi'(a_{t}|s_{t})}{\pi(a_{t}|s_{t})} \Big]$$

We replace the true advantage  $adv^{\pi}$  with an advantage estimator  $A_{\eta}$  (neural net, more on that later).

Instead of optimizing the true loss  $\widetilde{\mathcal{L}}_{\pi}$ , we optimize a sample loss  $\widehat{\mathcal{L}}_{\pi}$ : for a trajectory  $\tau = s_0, a_0, r_1, s_1, a_1, r_2, s_2, a_2, r_3, s_3, \ldots$ :

$$\widetilde{\mathcal{L}}_{\pi}(\pi') pprox \widehat{\mathcal{L}}_{\pi}(\pi') = \sum_{t=0}^{\infty} \gamma^{t} \cdot A_{\eta}( au, t) \cdot rac{\pi'(a_{t}|s_{t})}{\pi(a_{t}|s_{t})}$$

(More trajectories can be sampled, in which case we optimize the average sample loss over the trajectories.) We then let an automated gradient-based optimizer find  $\pi' = \pi_{\theta'}$  maximizing  $\widehat{\mathcal{L}}_{\pi}(\pi')$ . Note that the only term in  $\widehat{\mathcal{L}}_{\pi}$  that depends on the optimized parameters  $\theta'$  are the likelihood ratios! Hence, the gradient can be computed easily.

### Trust region policy optimization (TRPO)

Schulman, Levine, Abbeel, Jordan, Moritz: *Trust Region Policy Optimization*. In Proceedings of ICML'15.

To perform update from  $\pi$  to  $\pi'$ , theoretical TRPO:

- ullet samples a trajectory au (or a batch of trajectories) from  $\pi$
- uses the trajectory to:
  - ullet update the advantage estimator  $A_\eta$  (i.e., update its parameters  $\eta$ )
  - ullet construct the loss  $\widehat{\mathcal{L}}_{\pi}$
- solves the optimization problem

maximize 
$$\widehat{\mathcal{L}}_{\pi}(\pi') - \beta \cdot D_{\mathit{KL}}(\pi, \pi')$$

( $D_{KL}$  - KL divergence,  $\beta$  - suitable constant) via a black-box gradient-based optimizer.

This update loop is performed until timeout. Practical TRPO makes several changes to individual steps of the above scheme.

# **Proximal Policy Optimization (PPO)**

## Proximal Policy Optimization Algorithms

John Schulman, Filip Wolski, Prafulla Dhariwal, Alec Radford, Oleg Klimov OpenAI {joschu, filip, prafulla, alec, oleg}@openai.com

#### Abstract

We propose a new family of policy gradient methods for reinforcement learning, which alternate between sampling data through interaction with the environment, and optimizing a "surrogate" objective function using stochastic gradient ascent. Whereas standard policy gradient methods perform one gradient update per data sample, we propose a novel objective function that enables multiple epochs of minibatch updates. The new methods, which we call proximal policy optimization (PPO), have some of the benefits of trust region policy optimization (TRPO), but they are much simpler to implement, more general, and have better sample complexity (empirically). Our experiments test PPO on a collection of benchmark tasks, including simulated robotic locomotion and Atari game playing, and we show that PPO outperforms other online policy gradient methods, and overall strikes a favorable balance between sample complexity, simplicity, and wall-time.

#### PPO vs TRPO

PPO follows the same general scheme as TRPO with the following tweaks:

- Explicitly specifies how the advantage should be estimated: generalized advantage estimation (essentialy, truncated offline  $\lambda$ -returns, see later).
- Tweaks the loss function a bit uses different way of ensuring that  $\pi'$  is in the proximity of  $\pi$ .

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PPO comes in two variants depending on how it tweaks the loss function:

• Adaptive KL divergence penalty coefficient: like TRPO, but the coefficient  $\beta$  changes adaptively. Empirically does not perform as well as the second method:

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- Adaptive KL divergence penalty coefficient: like TRPO, but the coefficient  $\beta$  changes adaptively. Empirically does not perform as well as the second method:
- Clipped likelihood ratios.

#### Definition 57: Clipping

The function *CLIP* is defined as follows: 
$$CLIP(x,a,b) = \begin{cases} x & \text{if } a \leq x \leq b \\ a & \text{if } x < a \\ b & \text{if } b < x \end{cases}$$

# Basic PPO empirical loss

For a trajectory  $\tau = s_0, a_0, r_1, s_1, a_1, r_2, s_2, a_2, r_3, s_3, \dots$  denote

$$r_t(\pi,\pi') = \frac{\pi'(a_t|s_t)}{\pi(a_t|s_t)}$$

The main component of PPO loss/performance metric is:

$$\mathcal{L}_{\pi}^{\textit{CLIP}}(\pi') = \sum_{t=0}^{\infty} \min \left( A_{\eta}( au, t) \cdot r_t(\pi, \pi') \,, \, A_{\eta}( au, t) \cdot \textit{CLIP}ig( r_t(\pi, \pi'), 1 - arepsilon, 1 + arepsilon ig) 
ight)$$

(Note that no  $\gamma^t$  is included in the formula: discounting to some degree implicitly encompassed in the advantage estimation.)

# Constraining improvements but not damages

$$\mathcal{L}_{\pi}^{\textit{CLIP}}(\pi') = \sum_{t=0}^{\infty} \min \left( A_{\eta}( au, t) \cdot r_t(\pi, \pi') \,, \, A_{\eta}( au, t) \cdot \textit{CLIP}ig( r_t(\pi, \pi'), 1 - arepsilon, 1 + arepsilon ig) 
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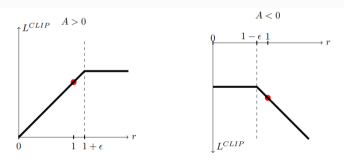


Figure 1: Plots showing one term (i.e., a single timestep) of the surrogate function  $L^{CLIP}$  as a function of the probability ratio r, for positive advantages (left) and negative advantages (right). The red circle on each plot shows the starting point for the optimization, i.e., r=1. Note that  $L^{CLIP}$  sums many of these terms.

### Advantage estimation in PPO

PPO uses generalized advantage estimation = an actor-critic framework using  $\lambda$ -return to estimate the q-value.

Formally, the advantage estimator is built on a value network  $V_{\eta} \colon \mathcal{S} \times \Theta \to \mathbb{R}$ .

Let  $\tau = s_0, a_0, r_1, s_1, a_1, r_2, s_2, a_2, r_3, s_3, \dots$  be a trajectory. An *n*-step advantage from time step t is the quantity

$$A_{\eta}^{t:t+n}(\tau) = r_{t+1} + \gamma r_{t+2} + \cdots + \gamma^{n-1} r_{t+n} + \gamma^n V_{\eta}(s_{t+n}) - V_{\eta}(s_t).$$

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For a parameter  $\lambda \in [0,1]$ , the generalized advantage estimate at time t is

$$GAE_{\eta}^{(\lambda,t)}(\tau) = \sum_{n=1}^{T-t} \lambda^n \cdot A_{\eta}^{t:t+n}(\tau).$$

For a trajectory  $\tau$ , PPO puts  $A_{\eta}(\tau, t) = GAE_{\eta}^{\lambda, t}(\tau)$ .

### **PPO** loss extension

Apart from  $\mathcal{L}_{\pi}^{\textit{CLIP}}\text{, PPO loss consist of two additional terms:}$ 

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1. The value network  $V_{\eta}$  and policy network  $\pi_{\theta}$  might share parameters (e.g. the same feature extraction layers). In this case,  $\eta$  and  $\theta$  should be trained together: for a sampled trajectory  $\tau=s_0, a_0, r_1, s_1, a_1, r_2, s_2, a_2, r_3, s_3, \ldots$ , the PPO loss will incorporate the empirical value loss

$$\mathcal{L}^V(\eta') = \sum_{t=0}^T (V_{\eta'}(s_t) - \underbrace{G_t( au)}_{\mathsf{target}})^2$$

(instead of sample return, the training target might be e.g. TD(0), or  $A_{\eta}(\tau, t) + V_{\eta}(s_t)$ : note that  $\eta$  in the target (current value of the parameter) is a constant).

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2. Original PPO also used entropy regularization, adding sample entropy loss:

$$\mathcal{L}^{entropy}(\pi') = -rac{1}{T}\sum_{t=0}^{T-1}\sum_{a}\pi'(a|s_t)\cdot\log\pi'(a|s_t)$$

#### **Total PPO loss**

Given a sampled trajectory  $\tau = s_0, a_0, r_1, s_1, a_1, r_2, s_2, a_2, r_3, s_3, \ldots$ , the current reference policy  $\pi$ , and hyperparameters  $\beta_1, \beta_2 \in \mathbb{R}^+$ , the canonical PPO loss is:

$$\begin{split} \mathcal{L}_{\tau}^{PPO}(\theta', \eta') &= \mathcal{L}_{\pi}^{\textit{CLIP}}(\pi') - \beta_{1} \cdot \mathcal{L}^{\textit{V}}(\eta') + \beta_{2} \cdot \mathcal{L}^{\textit{entropy}}(\pi') \\ &= \sum_{t=0}^{T} \min \left( A_{\eta}(\tau, t) \cdot r_{t}(\pi, \pi') \,,\, A_{\eta}(\tau, t) \cdot \textit{CLIP}\Big(r_{t}(\pi, \pi'), 1 - \varepsilon, 1 + \varepsilon\Big) \right) \\ &- \beta_{1} \cdot \sum_{t=0}^{T} (V_{\eta'}(s_{t}) - \underbrace{G_{t}(\tau)}_{\textit{target}})^{2} + \beta_{2} \cdot -\frac{1}{T} \sum_{t=0}^{T-1} \sum_{a} \pi'(a|s_{t}) \cdot \log \pi'(a|s_{t}). \end{split}$$

Important note: the advantage estimates  $A_{\eta}$  in  $\mathcal{L}^{CLIP}$  are evaluated before the loss is constructed and are treated as constants inside  $\mathcal{L}^{CLIP}$ ! The value network parameters are only considered variable inside the value loss.

Usually, a batch of  $\tau$  is sampled and average loss  $\frac{1}{|\mathsf{batch}|} \cdot \sum_{\tau \in \mathsf{batch}} \mathcal{L}^{PPO}_{\tau}(\theta', \eta')$  is optimized using gradient-based optimizer (eg., ADAM) to yield a policy update.

# PPO high-level pseudocode

#### Algorithm 12: PPO

**Input:** policy network  $\pi_{\theta}$ , value network  $V_{\eta}$ , hyperparameters  $\lambda$ , |B|,  $\beta_1$ ,  $\beta_2$ ,... initialize  $\theta$ ,  $\eta$ ;

## repeat

```
sample a batch B of trajectories from policy \pi = \pi_{\theta}; for each \tau \in B do  \begin{bmatrix} \text{for each } 0 \leq t \leq T - 1 \text{ do} \\ a_t(\tau) \leftarrow GAE_{\eta}^{\lambda,t}(\tau); \\ \text{define } \mathcal{L}_{\tau}^{CLIP}(\pi') = \sum_{t=0}^{T} \min \left( a_t(\tau) \cdot r_t(\pi, \pi') \,, \, a_t(\tau) \cdot CLIP(r_t(\pi, \pi'), 1 - \varepsilon, 1 + \varepsilon) \right); \\ \text{define } \mathcal{L}_{\tau}^{PPO}(\theta', \eta') = \mathcal{L}_{\pi, \tau}^{CLIP}(\pi') - \beta_1 \cdot \mathcal{L}^{V}(\eta') + \beta_2 \cdot \mathcal{L}^{entropy}(\pi') \\ \text{define } \mathcal{L}^{PPO}(\theta', \eta') = \frac{1}{|B|} \cdot \sum_{\tau \in B} \mathcal{L}^{PPO}_{\tau}(\theta', \eta') \end{bmatrix}
```

#### until timeout;

use ADAM or other optimizer to find  $\theta', \eta'$  approximately maximizing  $\mathcal{L}^{PPO}(\theta', \eta')$ ;  $\theta \leftarrow \theta'$ :  $\eta \leftarrow \eta'$ 

**Exploration vs. Exploitation:** 

A Systematic Approach

When you see a good move, look for a better one.

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Emanuel Lasker (1868-1941), 2nd World Chess Champion

• In RL, the algorithms often need to balance exploration of the MDP state space with exploitation: focusing on behavior that worked well in the past.

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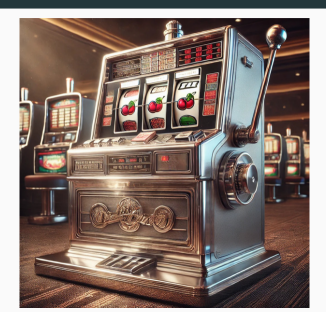
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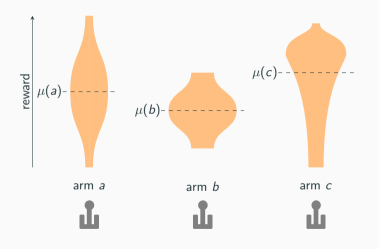
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- EvE dilemma also appears in domains that are not typically modeled as MDPs: recommender systems, disease treatment plans, or games.
- The EvE dilemma is systematically studied using the formalism of multi-armed bandits (MABs).

# One-armed bandit



# Multi-armed bandit (MAB)



#### A MAB is given by:

- a finite set of arms
   A;
- for each arm a a reward distribution  $D_a$  with mean  $\mu(a)$
- the  $D_a$  and  $\mu(a)$  are unknown to the player!

For simplicity, we will assume that rewards are from the interval [0,1].

- Interaction proceeds in discrete time steps 1, 2, 3, ..., T, where T is a termination time which might or might not be known to the player.
- ullet In each time step t, we choose some arm  $a_t \in A$  to pull and receive reward  $r_t \sim D_{a_t}$ .

= like a one-state MDP with stochastic rewards, but we do not known the reward distributions  $D_i$  in advance.

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Clearly, the expected reward is maximized by pulling, in each step t, the arm  $a^*$  with maximal mean reward  $\mu^*$  (we assume all arms have different mean rewards):

$$\mu^* = \max_{a \in A} \mu(a)$$

$$a^* = \arg\max_{a \in A} \mu(a)$$

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However, since we do not know  $D_a$ 's and  $\mu(a)$ 's, we cannot a priori determine which arm is optimal! We need to learn something about the reward distributions by exploring individual arms, while trying to maximize the accumulated returns.

#### **Policies in MABs**

#### Definition 58: MAB policy

A policy in a multi-armed bandit problem is a function  $\pi$  which to each history  $a_1, r_1, a_2, r_2, \ldots, a_t, r_t$  of actions and resulting rewards assigns a distribution over arms.

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The policies we will work with typically based their decisions on the following statistics of the history: for each arm a we keep:

- $N_t(a)$  the number of times the arm a was pulled by time t
- $R_t(a) = \sum_{1 \le i \le t} r_t \cdot \mathbb{I}(a_i = a)$  the total reward accumulated by pulling arm a up to time t
- ullet  $\hat{\mu}_t(a) = rac{R_t(a)}{N_t(a)}$  the empirical mean return of arm a

Node that if  $N_t(a) \to \infty$  as  $t \to \infty$ , then  $\hat{\mu}_t(a) \to \mu(a)$ .

## Example: $\varepsilon$ -greedy policy

Given  $\varepsilon \in [0,1]$ , an  $\varepsilon$ -greedy policy selects arm  $a_{t+1}$ , for any  $t \geq 0$ , as follows:

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Intuitively, this is sub-optimal policy: even for large t, when all  $\hat{\mu}_t(b)$  should be relatively good approximations of true mean rewards, the policy still selects sub-optimal arms at a constant rate. How to formalize this issue?

## Regret

The regret of a policy at time T is the difference between the expected return of the policy and the return we would get by always pulling the optimal arm. Formally

#### Definition 59: Regret

The regret of a policy  $\pi$  at time T is the quantity

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Another form of writing the regret:

$$Regret_{\pi}(T) = T \cdot \mu^* - \sum_{a \in A} \mu(a) \cdot \mathbb{E}^{\pi}[N_T(a)].$$

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Hence,

$$\begin{split} \textit{Regret}_{\varepsilon\text{-greedy}}(T) &= T \cdot \mu^* - \sum_{a \in A} \mu(a) \cdot \mathbb{E}^{\varepsilon\text{-greedy}}[N_T(a)] \\ &= \sum_{a \in A} \mathbb{E}^{\varepsilon\text{-greedy}}[N_T(a)] \cdot \mu^* - \sum_{a \in A} \mu(a) \cdot \mathbb{E}^{\varepsilon\text{-greedy}}[N_T(a)] \\ &= \sum_{a \in A} \mathbb{E}^{\varepsilon\text{-greedy}}[N_T(a)] \cdot (\mu^* - \mu(a)) \geq \mathbb{E}^{\varepsilon\text{-greedy}}[N_T(a)] \cdot \Delta_b \geq T \cdot \varepsilon \cdot \Delta_b. \end{split}$$

## Logarithmic regret

We will now demonstrate a simple policy achieving logarithmic regret.

The policy  $\pi$  we will construct requires the advance knowledge of the termination time T and of the the gap between the optimal and second-best arm:

$$\Delta_{\mathsf{min}} = \min_{b 
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Policy  $\pi$  proceeds in two phases:

- Phase 1 (exploration): in the first  $T_1 = \lceil \frac{\log(T) \cdot |A| \cdot 4}{\Delta_{\min}^2} \rceil$  steps it deterministically cycles through all arms. I.e., if  $A = \{a^1, a^2, \dots, a^k\}$ , then in step i it plays arm  $a^j$  where j = i (mod k) + 1.
- Phase 2 (exploitation) at timestep  $T_1+1$ ,  $\pi$  identifies action a with maximal empirical mean  $\hat{\mu}_{T_1}(a)$  and keeps playing this action for the remaining  $T_2=T-T_1$  steps.

# Upper-bounding the regret

The total regret of  $\pi$  can be decomposed into regrets accumulated in the two phases:

$$\begin{aligned} \textit{Regret}_{\pi}(\textit{T}) &= \textit{T} \cdot \mu^* - \sum_{t=1}^{T} \mathbb{E}^{\pi}[\textit{r}_t] \\ &= \underbrace{\textit{T}_1 \cdot \mu^* - \sum_{t=1}^{T_1} \mathbb{E}^{\pi}[\textit{r}_t]}_{\textit{R}_1, \text{ exploration regret}} \quad + \quad \underbrace{\textit{T}_2 \cdot \mu^* - \sum_{t=T_1+1}^{T} \mathbb{E}^{\pi}[\textit{r}_t]}_{\textit{R}_2, \text{ exploitation regret}} \end{aligned}$$

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Clearly  $R_1 \leq 1 \cdot T_1 \in \mathcal{O}(\log(T))$ .

 $R_2$  depends on whether  $\pi$  correctly classifies the optimal arm at timestep  $T_1+1$ .

- If yes, then  $R_2 = 0$ .
- If no, then  $R_2 \leq T_2 \leq T \in \mathcal{O}(T)$ .

# Bounding the probability of misclassification

We have

$$R_2 \leq T \cdot \mathbb{P}^{\pi}[Mis],$$

where Mis is the event that  $a^*$  does not have the maximal empirical mean after  $\mathcal{T}_1$  steps.

By union bound

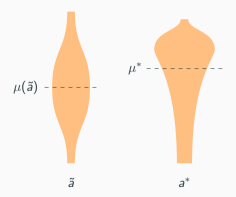
$$\mathbb{P}^{\pi}[\mathit{Mis}] \leq \sum_{b \neq a^*} \mathbb{P}^{\pi}[\hat{\mu}_{\mathcal{T}_1}(b) \geq \hat{\mu}_{\mathcal{T}_1}(a^*)].$$

Clearly, the second-best action, call it  $\tilde{a}$ , has the highest likelihood of achieving higher empirical mean than  $a^*$ . I.e.,

$$\mathbb{P}^{\pi}[\textit{Mis}] \leq (|A|-1) \cdot \mathbb{P}^{\pi}[\hat{\mu}_{\mathcal{T}_1}(\tilde{\textit{a}}) \geq \hat{\mu}_{\mathcal{T}_1}(\textit{a}^*)].$$

It thus suffices to bound the probability of  $\tilde{a}$  overtaking  $a^*$  in empirical mean.

## Probability of empirical deviation



For  $\tilde{a}$  to overtake  $a^*$ , at least one of the arms must have empirical mean after  $T_1$  steps at least  $\frac{\Delta_{\min}}{2}$ -away from their true mean.

Hence, it suffices to bound the probability that an empirical mean deviates too much from the true mean.

## Hoeffding's inequality for large deviations

#### Theorem 60: Hoeffding's inequality

Let D be a distribution taking values in [0,1]. Let  $\mu$  be the mean of D and let  $\hat{\mu}_n = \sum_{i=1}^n x_i$ , where each  $x_i$  is an independent sample from D. Then, for any  $n \in \mathbb{N}^+$  and any  $\delta > 0$ :

$$\mathbb{P}\big[|\hat{\mu}_n - \mu| \ge \delta\big] \le 2e^{-2n\delta^2}$$

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Recall  $T_1 \approx \frac{\log(T) \cdot |A| \cdot 4}{\Delta_{\min}^2}$ . Then, e.g. for  $a^*$ :

$$\mathbb{P}^{\pi}\big[|\mu_{T_1}(a^*) - \mu^*| \geq \frac{\Delta_{\min}}{2}\big] \leq 2e^{-2\frac{T_1}{|A|}\frac{\Delta_{\min}^2}{4}} \approx 2e^{-2\log(T)} = \frac{2}{T^2}.$$

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Hence,  $\mathbb{P}^{\pi}[\mathit{Mis}] \leq C \cdot \frac{1}{T^2}$  for some constant C.

It follows that the exploitation regret  $R_2$  is  $\leq T \cdot \mathbb{P}^{\pi}[\mathit{Mis}] = C' \cdot \frac{1}{T}$  for some constant C'.

## Final bound on regret of $\pi$

The total regret of  $\pi$  is:

$$Regret_{\pi}(T) = \underbrace{R_1(T)}_{\in \mathcal{O}(\log(T))} + \underbrace{R_2(T)}_{\in \mathcal{O}(1)} \in \mathcal{O}(\log(T)).$$

The logarithmic regret is the best possible:

Theorem 61: Lai and Robbins ("Asymptotically efficient adaptive allocation rules", 1985)

Any policy has regret in  $\Omega(\log(T))$ .

The disadvantage of  $\pi$  is that it needs to know both T and  $\Delta_{\min}$  in advance.

Knowledge of T can be discarded by using  $\varepsilon$ -greedy policies with adaptive  $\varepsilon$ : if  $\varepsilon_t = \min\{1, \frac{|A|}{\Delta^2 \cdot \cdot t}\}$ , then the regret is logarithmic (see David Silver's slides).

But there is actually a policy yielding logarithmic regret without the advance knowledge of either T or  $\Delta_{min}$ .

## Optimism in the face of uncertainty

The idea is that <u>under-explored</u> arms should be explored more: they could be better than we currently think (and if not, exploring them should disprove that).

# Optimism in the face of uncertainty

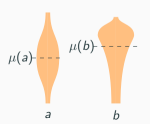
The idea is that under-explored arms should be explored more: they could be better than we currently think (and if not, exploring them should disprove that).

We will seek optimistic estimates of  $\mu(a)$  in the form of upper confidence bounds: we seek an empirical quantity  $U_t(a)$  such that with high probability

$$\mu(a) \leq \underbrace{\hat{\mu}_t(a) + U_t(a)}_{UCB_t(a)}.$$

The policy will always pick action maximizing  $UCB_t$ .

Moreover,  $U_t(a)$  should be as tight as possible given available information. In particular, it should hold that if  $N_t(a) \leq N_t(b)$ , then  $U_t(a) \leq U_t(b)$ .



## One-sided Hoeffding's inequality

#### Theorem 62: Hoeffding's inequality (one-sided)

Let D be a distribution taking values in [0,1]. Let  $\mu$  be the mean of D and let  $\hat{\mu}_n = \sum_{i=1}^n x_i$ , where each  $x_i$  is an independent sample from D. Then, for any  $n \in \mathbb{N}^+$  and any  $\delta > 0$ :

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Let 1-p be some required confidence level. We want to find "tight"  $U_t(a)$  such that  $\mu^* \geq \hat{\mu}_t(a) + U_t(a)$  with probability at most p, i.e.

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$$\mathbb{P}\big[\mu^* - \hat{\mu}_t(\mathsf{a}) \geq U_t(\mathsf{a})\big] \leq \mathsf{p}.$$

By (one-sided) Hoeffding:

$$\mathbb{P}\big[\mu^* - \hat{\mu}_t(a) \geq U_t(a)\big] \leq e^{-2N_t(a)U_t^2(a)}.$$

### **Deriving UCB formula**

From the previous we have:

$$\mathbb{P}\big[\mu^* - \hat{\mu}_t(a) \geq U_t(a)\big] \leq e^{-2N_t(a)U_t^2(a)}.$$

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Performing substitution  $p = e^{-2N_t(a)U_t^2(a)}$  we derive the expression for  $U_t(a)$ :

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$$U_t(a) = \sqrt{\frac{log(1/p)}{2N_t(a)}}$$

For this  $U_t(a)$ , it holds  $\mathbb{P}[\mu^* - \mu_t(a) \geq U_t(a)] \leq p$ , i.e.

$$\mathbb{P}[\mu^* \leq \hat{\mu}_t(a) + U_t(a)] \geq 1 - p.$$

We want to increase the confidence over time, i.e. p should be a function of t with  $p \to 0$  as  $t \to \infty$ . The standard approach is to put  $p = \frac{1}{t^c}$  for a suitable constant c. Then

$$U_t(a) = \sqrt{\frac{\log(1/p)}{2N_t(a)}} = \sqrt{\frac{c\log(t)}{2N_t(a)}} = \sqrt{\frac{c}{2}}\sqrt{\frac{\log t}{N_t(a)}} = C \cdot \sqrt{\frac{\log t}{N_t(a)}}$$

### **Summary of UCB policy**

In each step t, the UCB policy selects the arm with the highest upper confidence bound, i.e. arm a such that

$$a = \operatorname*{arg\,max}_{a \in A} UCB_t(a) = \operatorname*{arg\,max}_{a \in A} \left( \hat{\mu}_t(a) + C \cdot \sqrt{\frac{\log t}{N_t(a)}} \right),$$

where C is a hyperparameter known as exploration constant. It can be shown that for [0,1]-valued rewards, choosing  $C=\sqrt{2}$  suffices to achieve logarithmic regret

#### Theorem 63

When the reward distributions are over the interval [0,1], then for  $C=\sqrt{2}$  it holds

$$Regret_{UCB}(T) \in \mathcal{O}(\log T).$$

Proof in optional reading: Auer, Cesa-Bianchi, Fischer: Finite-time Analysis of the Multiarmed Bandit Problem. In *Machine Learning* (47), 2002.

• UCB-like algorithms with logarithmic regret were developed also for more general cases (e.g. distributions with unbounded support, where typically some shape of the distribution or a known bound on its variance is known).

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• Most prominently, this idea is applied in the context of Monte-Carlo tree search algorithms.