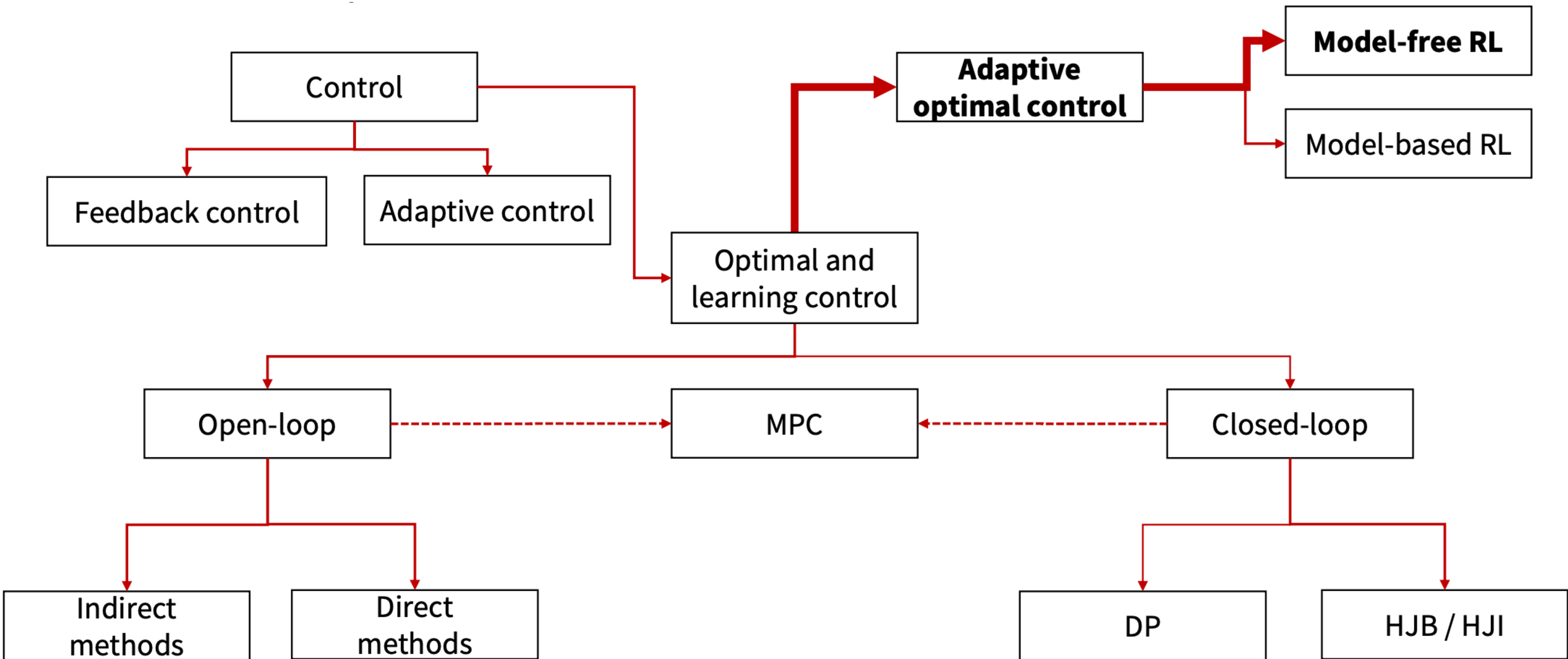


AA203
Optimal and Learning-based Control
Model-free Reinforcement Learning: Value-based Methods

Roadmap



Markov Decision Process

State:

$$x \in \mathcal{X}$$

Action:

$$u \in \mathcal{U}$$

Transition function / Dynamics:

$$T(x_t | x_{t-1}, u_{t-1}) = p(x_t | x_{t-1}, u_{t-1})$$

Typically represented as a tuple

$$\mathcal{M} = (\mathcal{X}, \mathcal{U}, T, R, \gamma)$$

Reward function:

$$r_t = R(x_t, u_t) : \mathcal{X} \times \mathcal{U} \rightarrow \mathbb{R}$$

Discount factor:

$$\gamma \in (0,1)$$

Goal: choose a policy that maximizes cumulative (discounted) reward

$$\pi^* = \underset{\pi}{\operatorname{argmax}} \mathbb{E}_p \left[\sum_{t \geq 0} \gamma^t R(x_t, \pi(x_t)) \right]$$

Review

In previous lectures, we made the distinction between *prediction* (given a policy π , estimate V_π, Q_π) and *control* (learn the optimal policy π^*)

Motivated by Dynamic Programming, we discussed *exact methods* for solving MDPs:

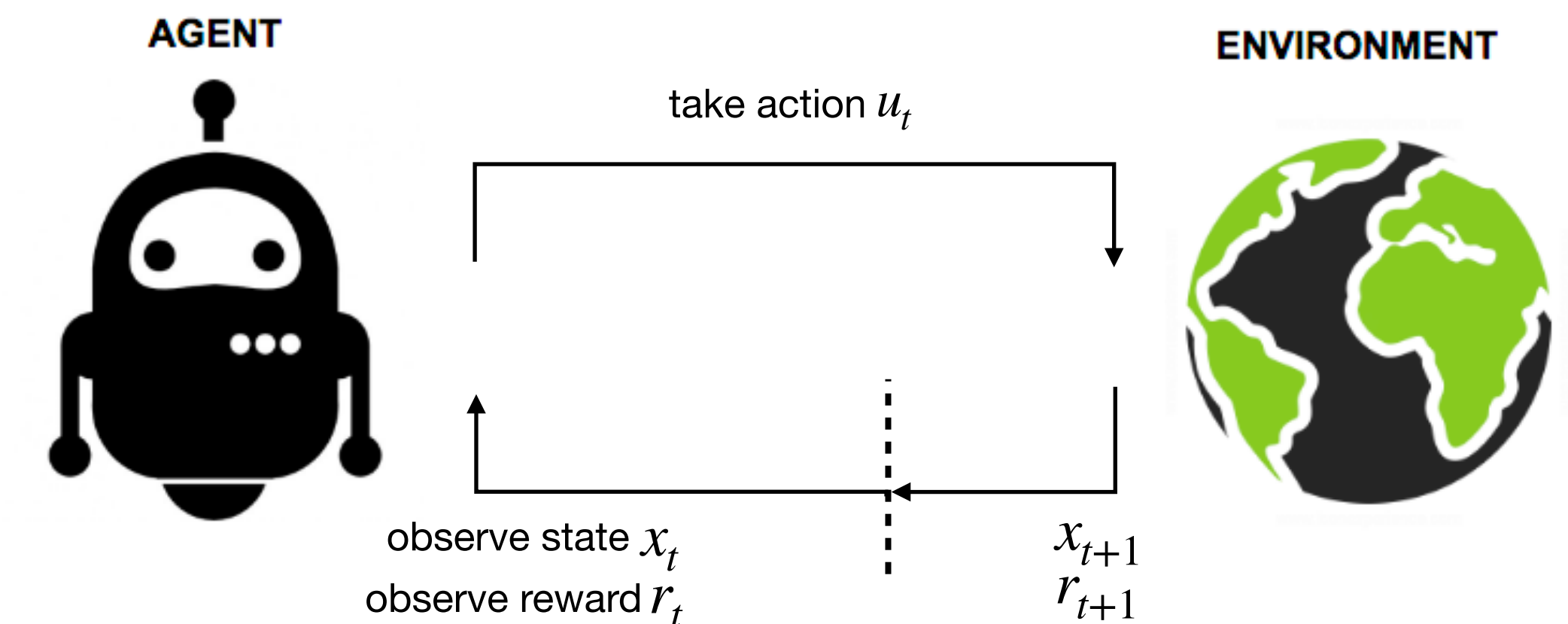
- Policy Iteration
- Value Iteration

Problem	Bellman Equation	Algorithm
Prediction	Bellman Expectation Equation	Iterative Policy Evaluation
Control	Bellman Expectation Equation + Greedy Policy Improvement	Policy Iteration
Control	Bellman Optimality Equation	Value Iteration

Limitation: Update equations (i.e., Bellman equations) require access to dynamics model $T(x_{t+1} | x_t, u_t)$

We saw how to use **sampling and bootstrapping** to approximate the expectations in the update equations:

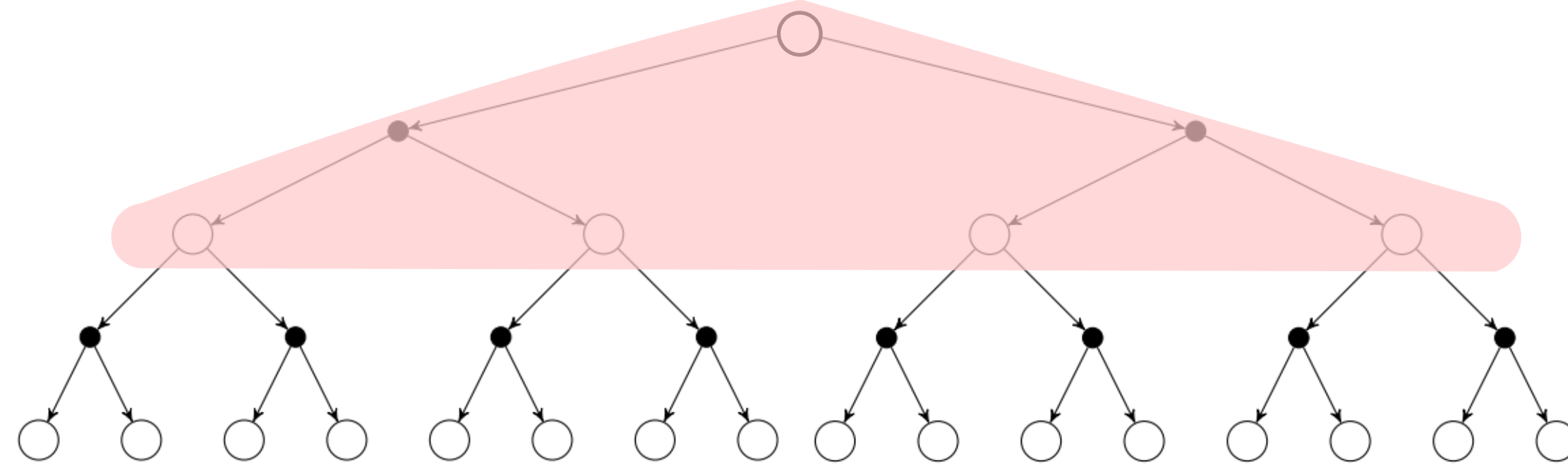
- Monte Carlo (MC) Learning
- Temporal-Difference (TD) Learning



- For prediction:

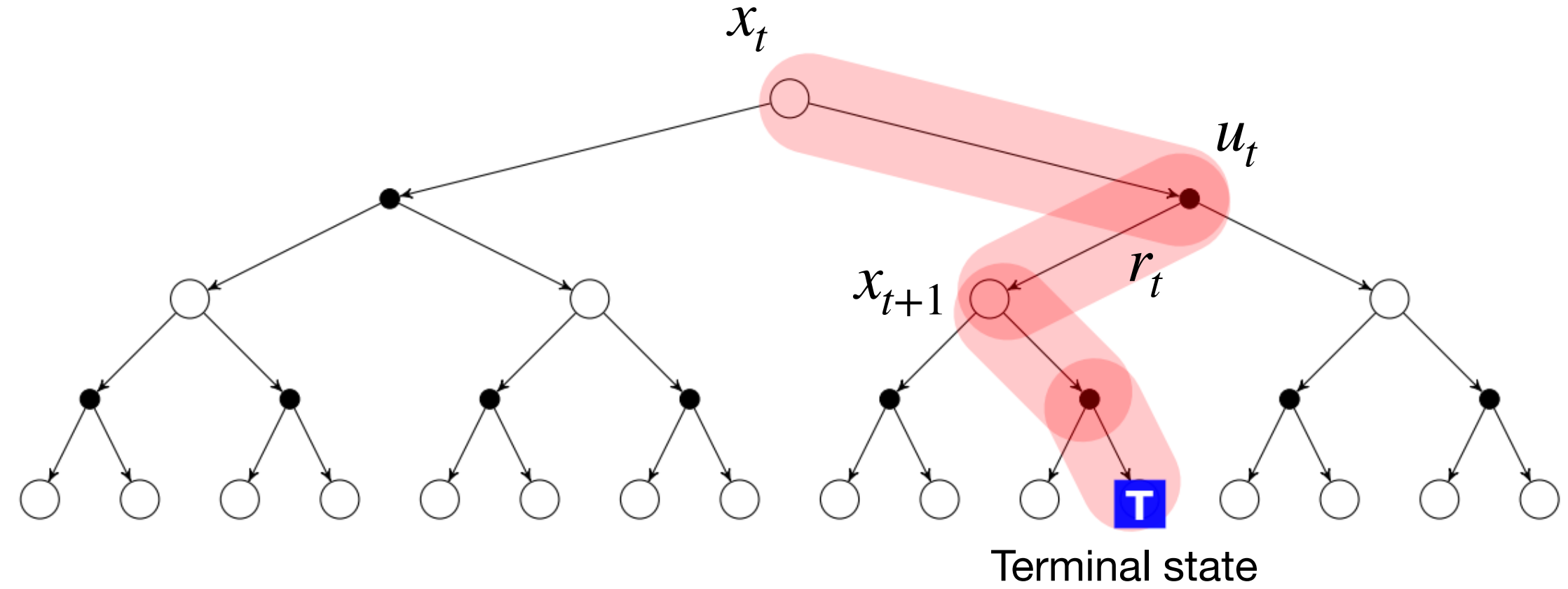
Dynamic Programming backup

$$\hat{V}(x_t) \leftarrow \mathbb{E}_{x_{t+1}} [R_t + \gamma \hat{V}(x_{t+1})]$$



Monte Carlo backup

$$\hat{V}(x_t) \leftarrow \hat{V}(x_t) + \alpha (G_t - \hat{V}(x_t))$$



- **Sampling:** define the update through samples to approximate expectations

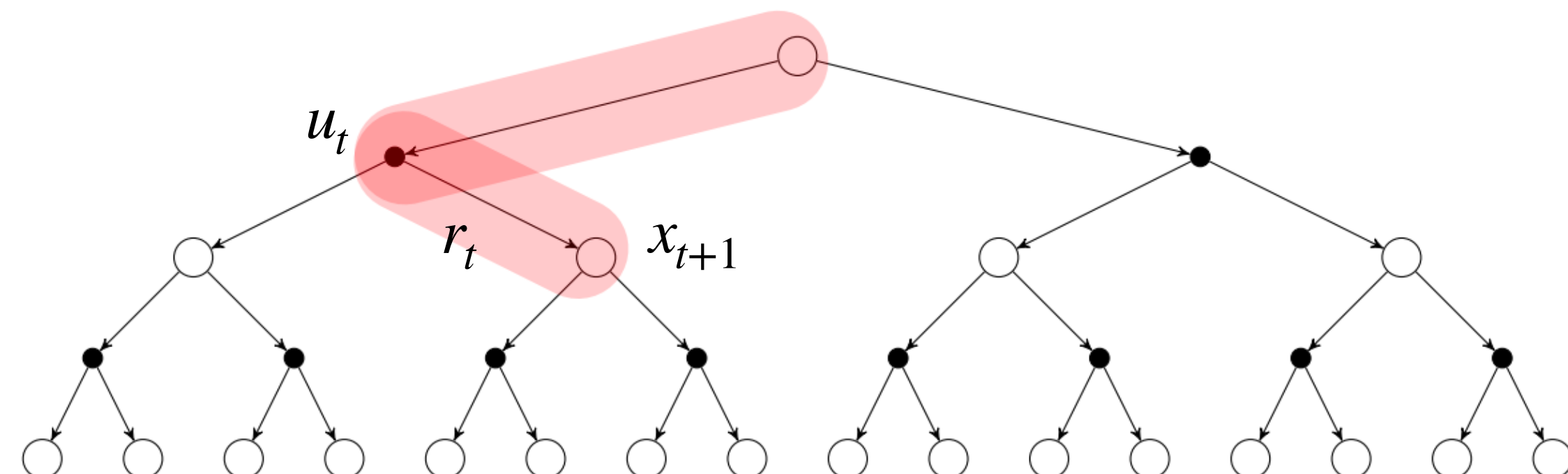
- MC samples
- TD samples
- DP does not sample

- **Bootstrapping:** define the update through an estimate

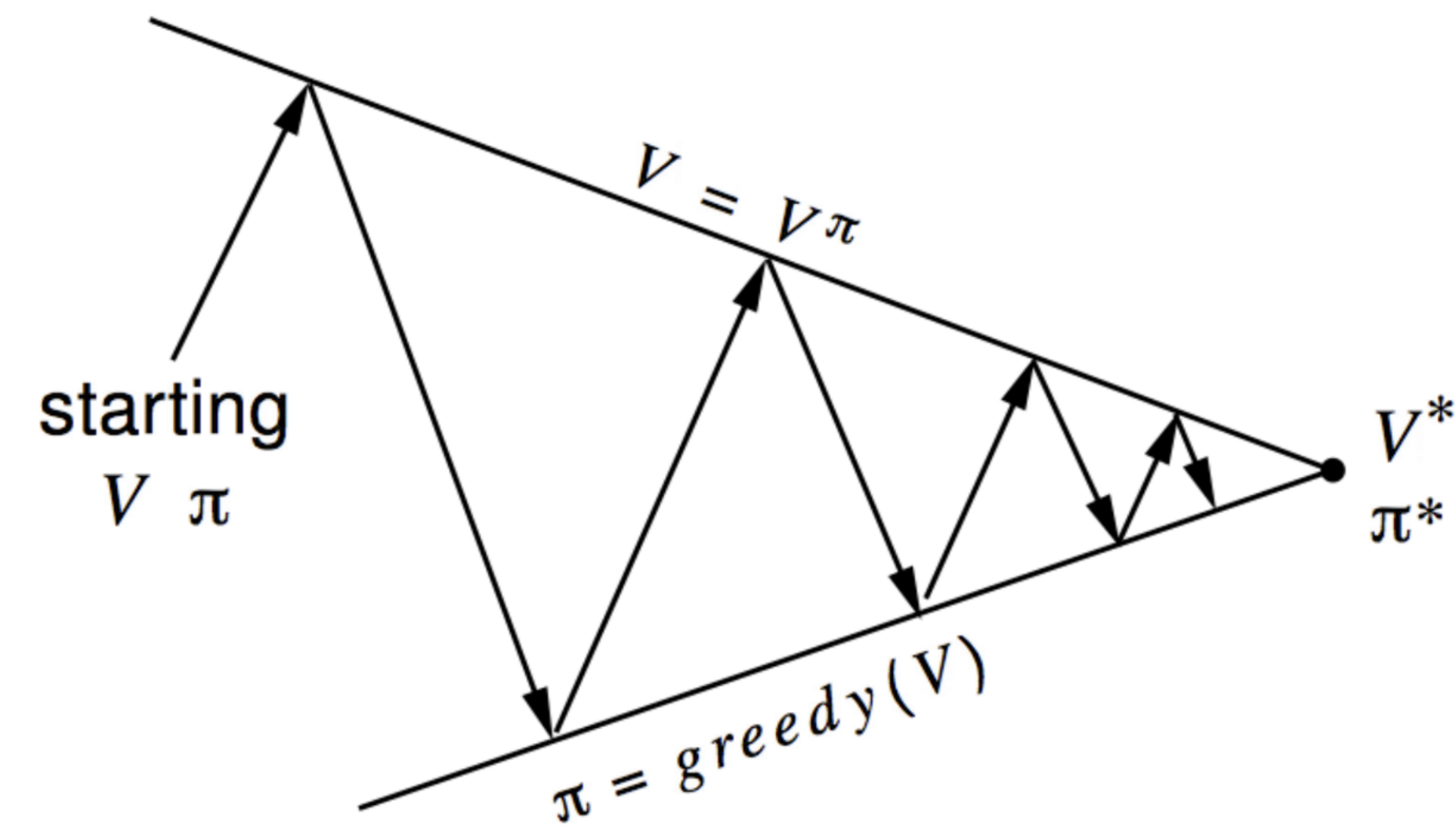
- MC does not bootstrap
- TD bootstraps
- DP bootstraps

Temporal-Difference backup

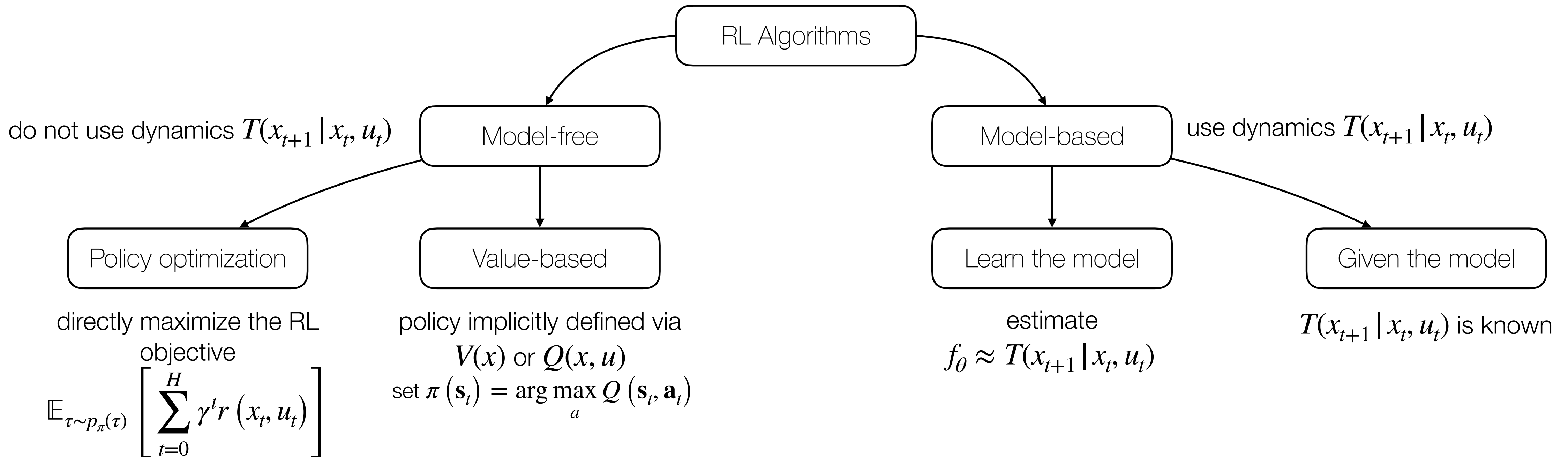
$$\hat{V}(x_t) \leftarrow \hat{V}(x_t) + \alpha (R_t + \gamma \hat{V}(x_{t+1}) - \hat{V}(x_t))$$



- For control:



A taxonomy of RL



Outline

Tabular methods

- On-policy & Off-policy
 - SARSA
 - Q-learning

Value function approximation

Deep (Value-based) RL Methods & Applications

Temporal-Difference Control

- TD learning has several advantages over MC
 - Lower variance
 - Online
 - Incomplete sequences
- Natural idea: use TD instead of MC in our GPI scheme
 - Apply TD to estimate $Q(x, u)$
 - Use ϵ -greedy policy improvement
 - Update every time-step

Updating action-value functions with Sarsa

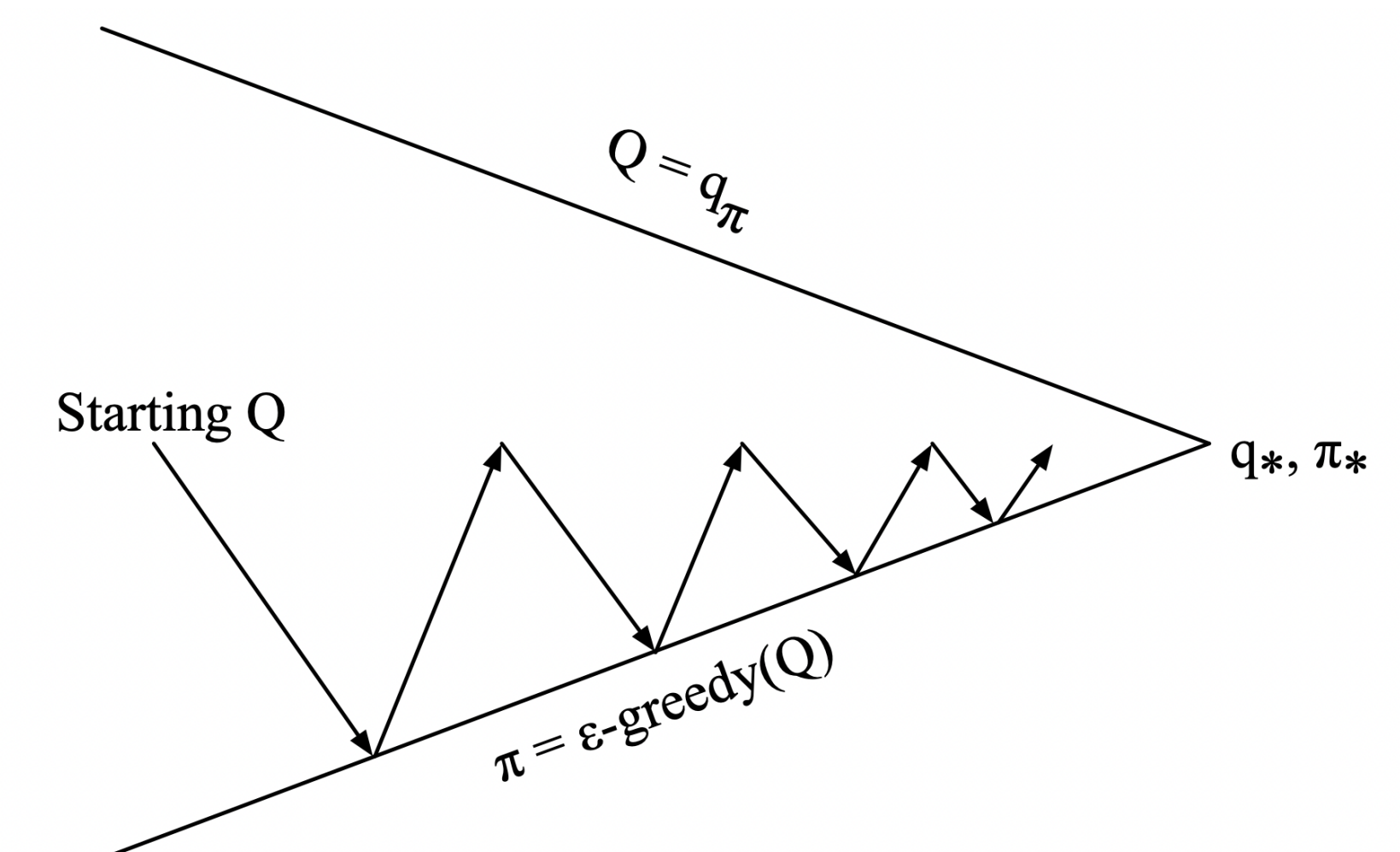
- Uses every element of the quintuple of events, $(x_t, u_t, r_t, x_{t+1}, u_{t+1})$, that make up a transition from one state-action pair to the next through the following update rule

$$Q(x_t, u_t) \leftarrow Q(x_t, u_t) + \alpha \left(r_t + \gamma Q(x_{t+1}, u_{t+1}) - Q(x_t, u_t) \right)$$

Temporal-Difference backup

$$\hat{V}(x_t) \leftarrow \hat{V}(x_t) + \alpha \left(R_t + \gamma \hat{V}(x_{t+1}) - \hat{V}(x_t) \right)$$

- In RL literature, $(x_t, u_t, r_t, x_{t+1}, u_{t+1})$ is often expressed as $(s_t, a_t, r_t, s_{t+1}, a_{t+1})$: hence the name



Every time-step:

Policy evaluation Sarsa, $Q \approx q_\pi$

Policy improvement ϵ -greedy policy improvement

Sarsa algorithm for on-policy control

Initialize $Q(x, u), \forall x \in X, \forall u \in U$, arbitrarily, and $Q(\text{terminal-state}, \cdot) = 0$

Repeat (for each episode):

 Initialize x_t

 Choose u_t from x_t using policy derived from Q (e.g., ϵ -greedy)

 Repeat (for each step of episode):

 Take action u_t , observe r_t, x_{t+1}

 Choose u_{t+1} from u_{t+1} using policy derived from Q (e.g., ϵ -greedy)

$$Q(x_t, u_t) \leftarrow Q(x_t, u_t) + \alpha \left(r_t + \gamma Q(x_{t+1}, u_{t+1}) - Q(x_t, u_t) \right)$$

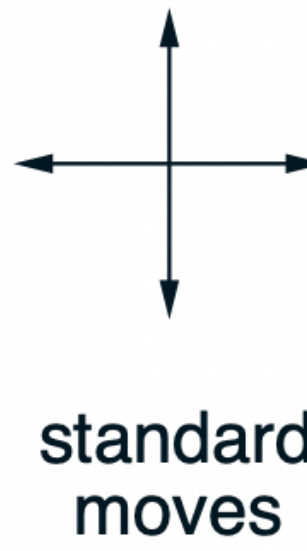
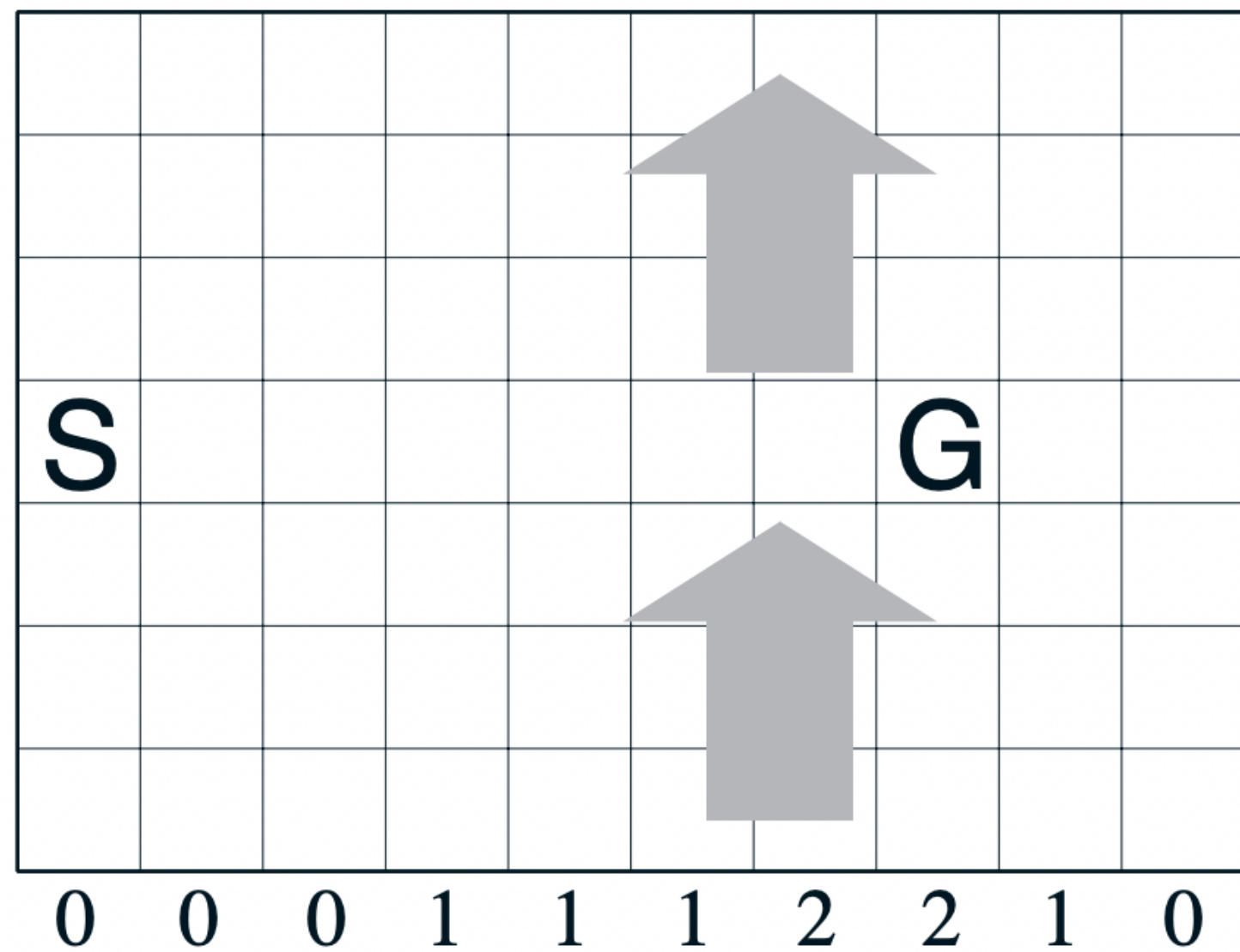
$x_t \leftarrow x_{t+1}; u_t \leftarrow u_{t+1}$

 until x_t is terminal

On-policy: evaluate or improve the policy that is used to make decisions

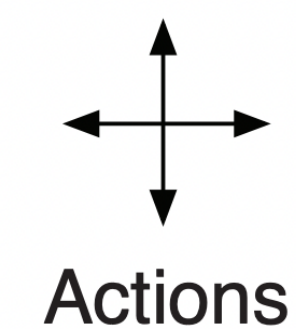
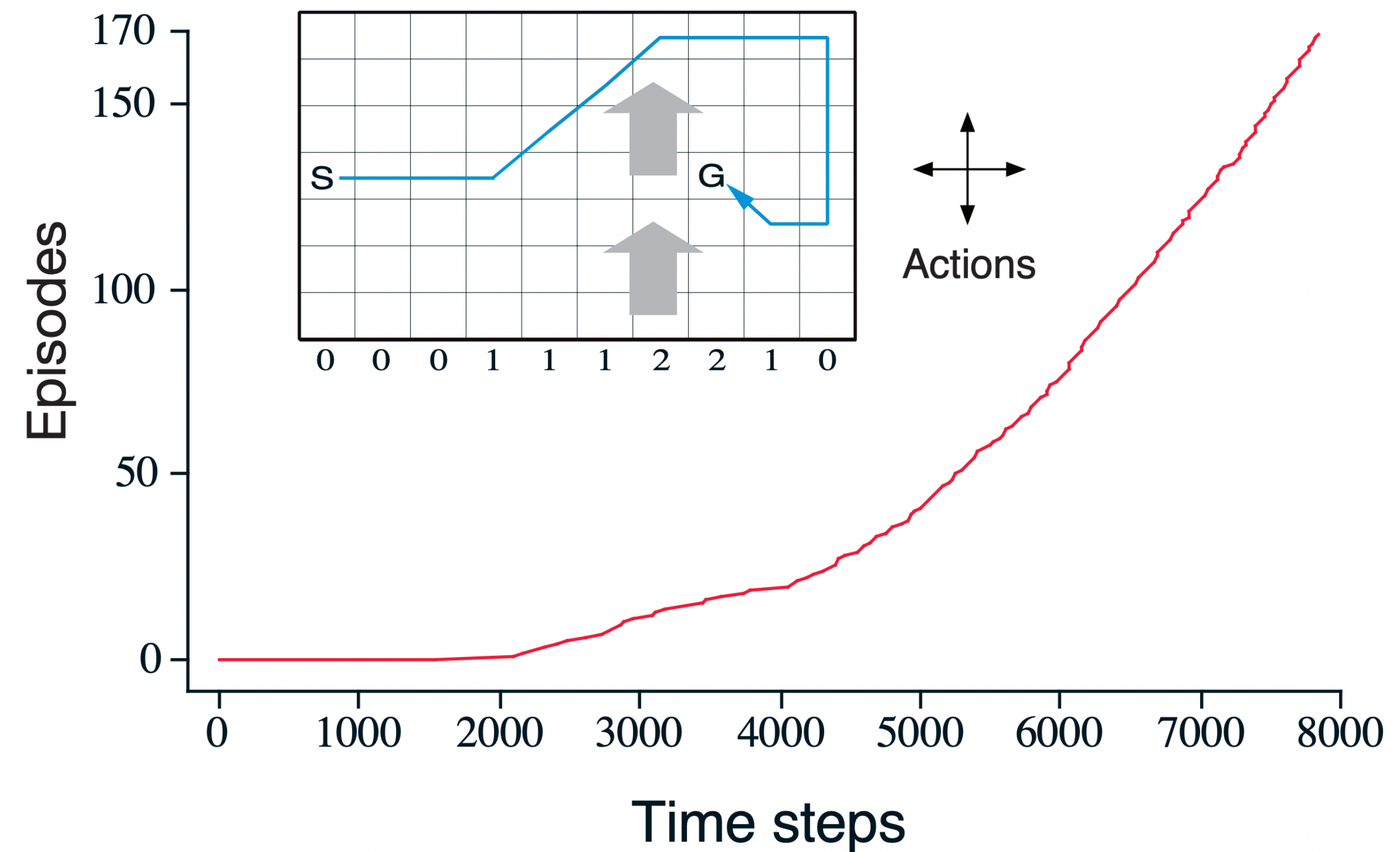
Off-policy: evaluate or improve a policy different from that used to generate the data

Windy Gridworld example



- Reward -1 until goal is reached
- $\epsilon = 0.1$
- $\alpha = 0.5$
- $\gamma = 1$

Windy Gridworld example



Question:

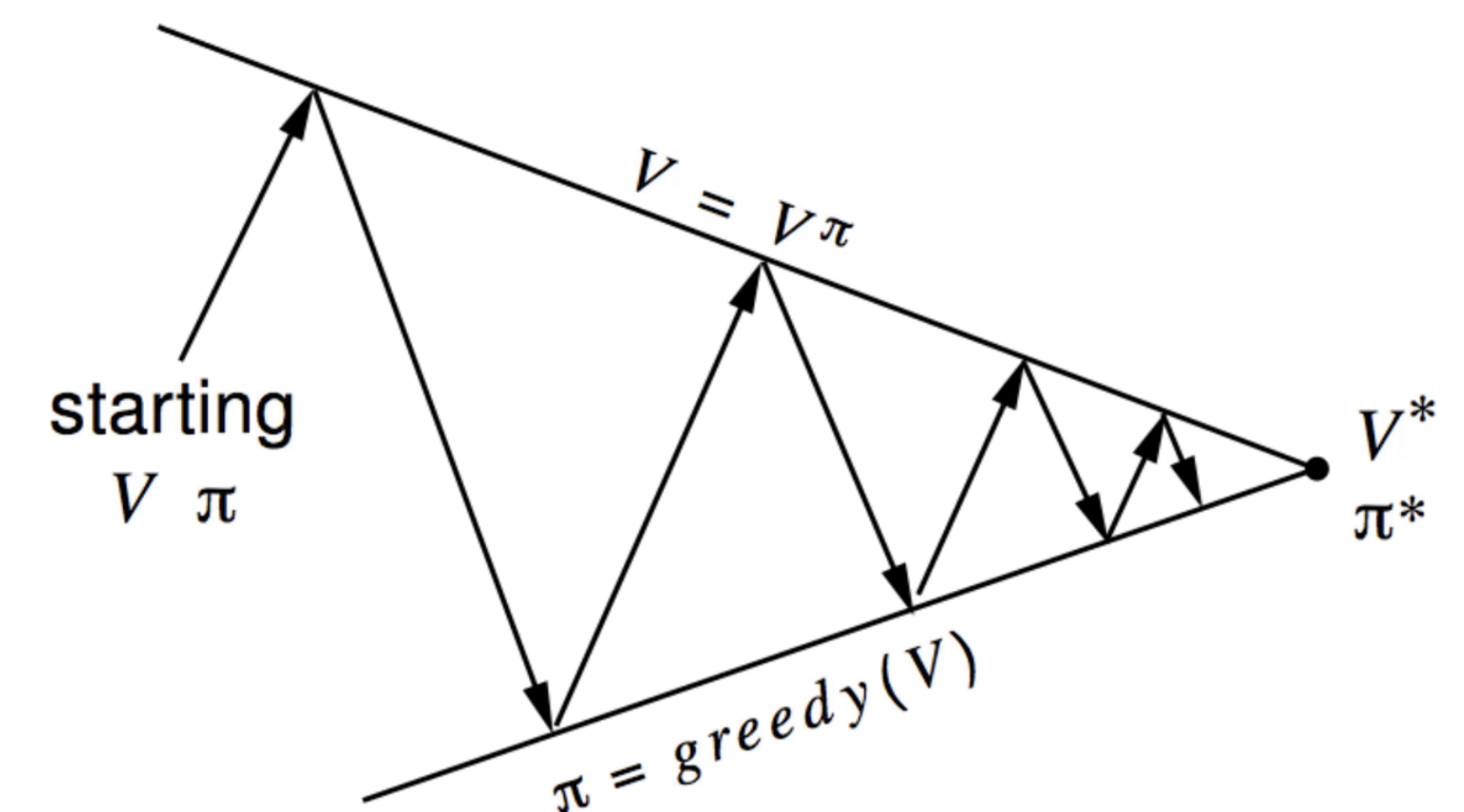
Would MC methods easily apply to this problem? And why?

Off-policy learning

- Evaluate target policy $\pi(u | x)$ to compute $V_\pi(x)$ or $Q_\pi(x, u)$ while following behavior policy $\mu(u | x)$, i.e.,
$$\{x_1, u_1, r_1, \dots, x_T\} \sim \mu, \text{ "the data we observe is obtained under policy } \mu\text{"}$$

Why is this important?

- Learn from observing humans or other agents
- Re-use experience generated from old policies $\pi_1, \pi_2, \dots, \pi_{t-1}$
- Learn about **optimal** policy while following **exploratory** policy



Off-policy learning of action-values

- We consider off-policy learning of action-values $Q(x, u)$
- As in Sarsa, we use the behavior policy μ to obtain $(x_t, u_t, r_t, x_{t+1}, \textcolor{red}{u'_{t+1}})$, but we consider an alternative successor action $\textcolor{red}{u'_{t+1}} \sim \pi(u'_{t+1} | x_{t+1})$
- And update $Q(x, u)$ towards value of alternative action

$$Q(x_t, u_t) \leftarrow Q(x_t, u_t) + \alpha \left(\textcolor{red}{r_t + \gamma Q(x_{t+1}, u'_{t+1}) - Q(x_t, u_t)} \right)$$

Q-learning

Specifically, in Q-learning

- The target policy π is chosen as the **greedy** policy w.r.t. $Q(x, u)$

$$\pi(x_{t+1}) = \operatorname{argmax}_{u'_{t+1}} Q(x_{t+1}, u'_{t+1})$$

- The behavior policy μ is chosen as the **ϵ -greedy** policy w.r.t. $Q(x, u)$

Which leads to the following Q-learning target and update:

$$\begin{aligned} & r_{t+1} + \gamma Q(x_{t+1}, u'_{t+1}) \\ &= r_{t+1} + \gamma Q\left(x_{t+1}, \operatorname{argmax}_{u'_{t+1}} Q(x_{t+1}, u'_{t+1})\right) \end{aligned}$$

$$= r_{t+1} + \gamma \max_{u'_{t+1}} Q(x_{t+1}, u'_{t+1})$$

$$Q(x_t, u_t) \leftarrow Q(x_t, u_t) + \alpha \left(r_t + \gamma \max_{u'_{t+1}} Q(x_{t+1}, u'_{t+1}) - Q(x_t, u_t) \right)$$

Q-learning algorithm for off-policy control

Initialize $Q(x, u), \forall x \in X, \forall u \in U$, arbitrarily, and $Q(\text{terminal-state}, \cdot) = 0$

Repeat (for each episode):

 Initialize x_t

 Repeat (for each step of episode):

 Choose u_t from x_t using policy derived from Q (e.g., ϵ -greedy)

 Take action u_t , observe r_t, x_{t+1}

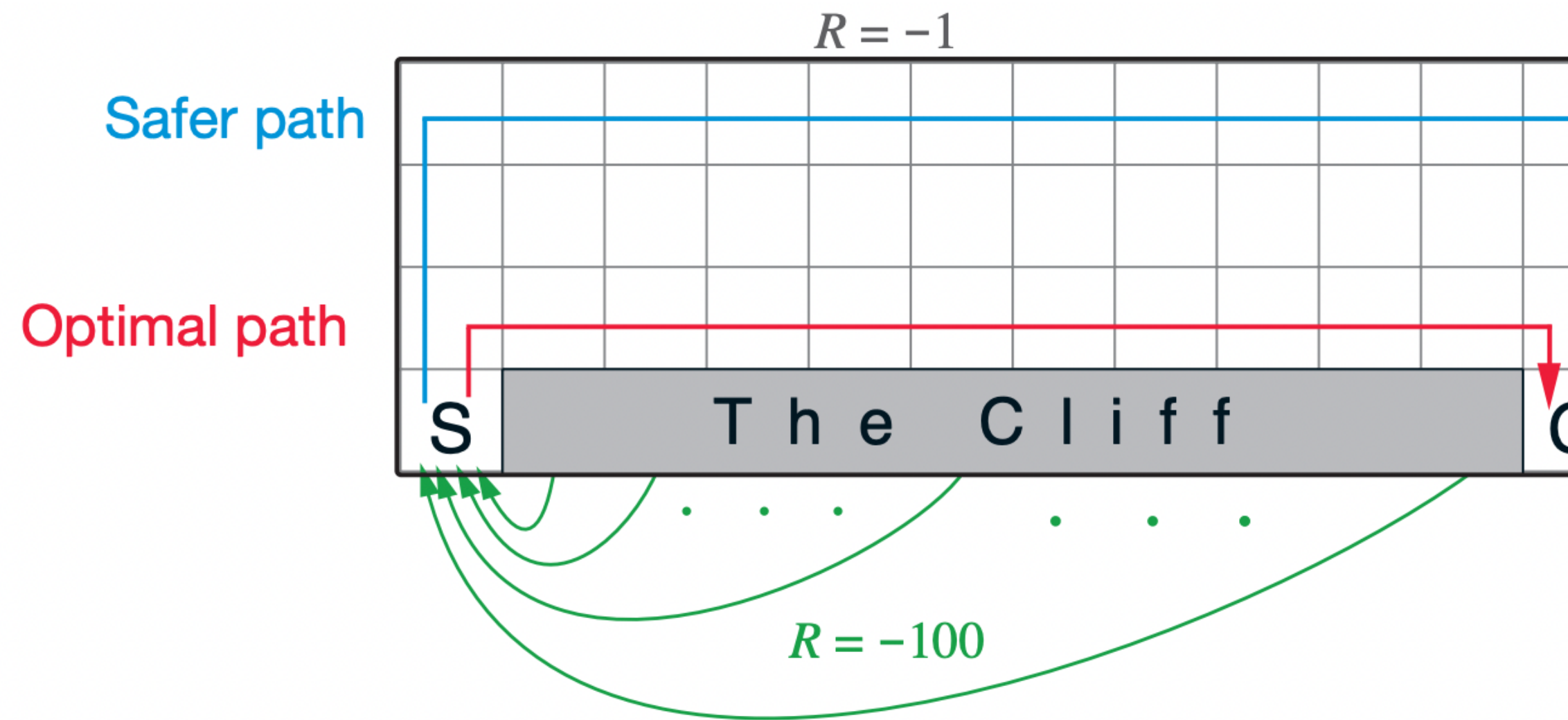
$$Q(x_t, u_t) \leftarrow Q(x_t, u_t) + \alpha \left(r_t + \gamma \max_{u'_{t+1}} Q(x_{t+1}, u'_{t+1}) - Q(x_t, u_t) \right)$$

 until x_t is terminal

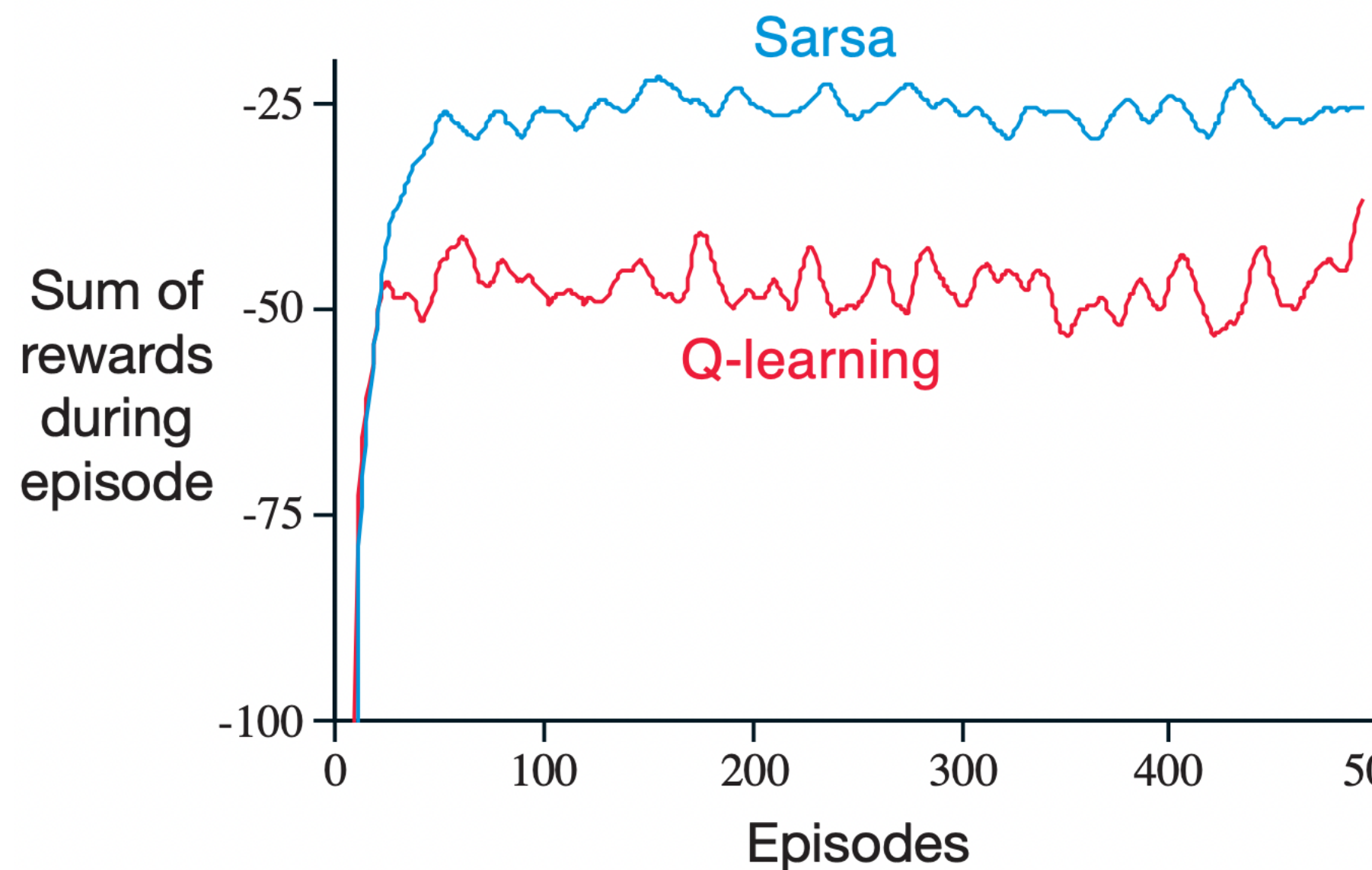
Theorem

*Q-learning control converges to the optimal action-value function,
 $Q(s, a) \rightarrow q_*(s, a)$*

Differences between Sarsa and Q-learning



- Reward -1 until goal is reached, -100 if on “The Cliff”
- $\epsilon = 0.1$
- $\alpha = 0.5$
- $\gamma = 1$



- Sarsa converges to the **optimal ϵ -greedy policy**
- Q-learning converges to the **optimal policy π^*** / value function Q^*

Outline

Tabular methods

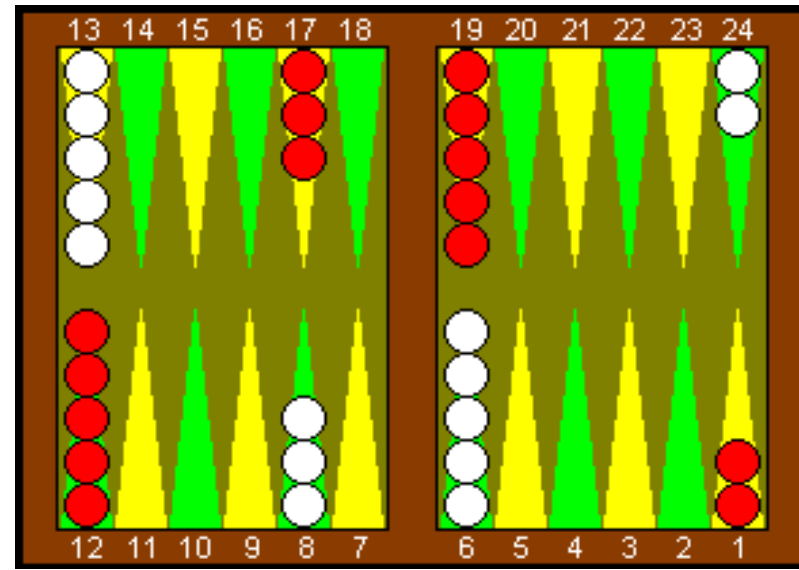
- On-policy & Off-policy
 - SARSA
 - Q-learning

Value function approximation

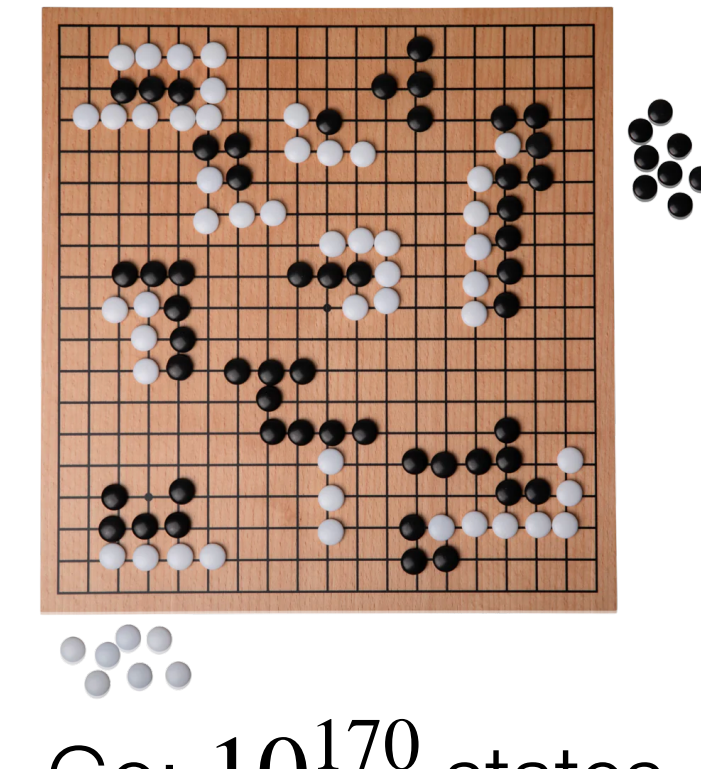
Deep (Value-based) RL Methods & Applications

Solving large-scale problems with RL

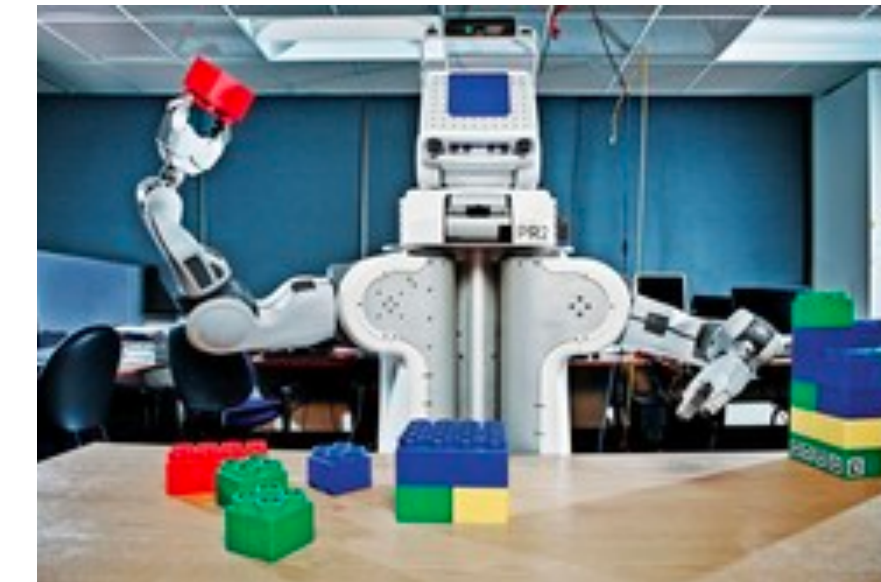
- Reinforcement learning can be used to solve *large* problems, e.g.,



Backgammon: 10^{20} states



Go: 10^{170} states



All those problems where we have a continuous state space

How can we scale the methods for model-free RL we developed over the last lectures?

Value function approximation

- So far we used **lookup tables** to represent value functions:
 - One entry for every state x in $V(x)$
 - One entry for every state-action pair (x, u) in $Q(x, u)$
- In large MDPs, lookup table might be prohibitive. For two main reasons:
 - Memory: too many actions/states to store
 - Sparsity/Curse of dimensionality: learning the value of each state/action pair individually might take too long

Solution:

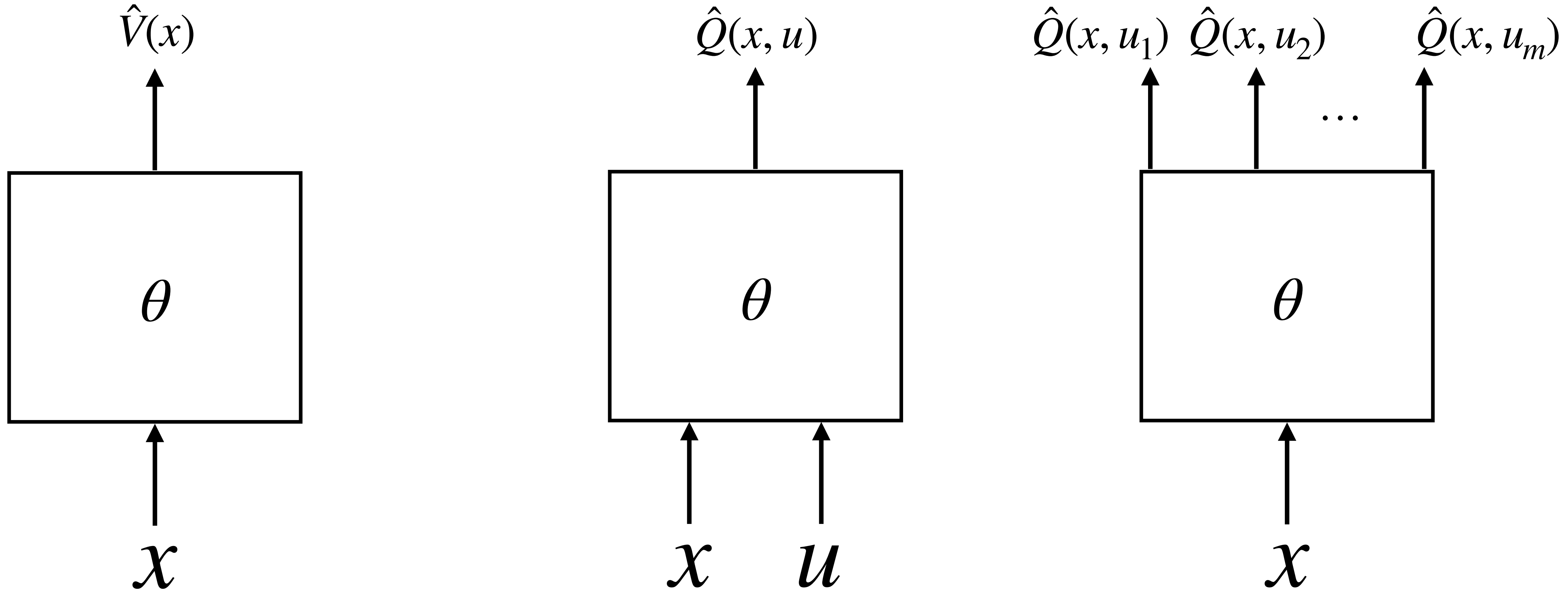
- Estimate the value function through function approximation, i.e., define a parametric function with parameters θ

$$\hat{Q}_\theta(x, u) \approx Q(x, u)$$

$$\hat{V}_\theta(x) \approx V(x)$$

- Represent the value function compactly (depends only on parameters θ)
- Generalize across states (avoid having to visit the entire state-action space by generalizing from seen to unseen states)

Different types of value function approximations



There are many possible function approximators

- Linear regression, Neural network, Random forest, Nearest neighbor, etc.

Approximating value fn. by (stochastic) gradient descent

Goal: find the parameter vector θ that minimizes the mean-squared error between the estimated value $\hat{V}_\theta(x)$ and the true value $V_\pi(x)$

$$J(\theta) = \mathbb{E}_\pi \left[\left(V_\pi(x) - \hat{V}_\theta(x) \right)^2 \right]$$

Gradient descent converges to a local minimum

$$\begin{aligned} \Delta\theta &= -\frac{1}{2}\alpha \nabla_\theta J(\theta) \\ &= \alpha \mathbb{E}_\pi \left[\left(V_\pi(x) - \hat{V}_\theta(x) \right) \nabla_\theta \hat{V}_\theta(x) \right] \end{aligned}$$

Stochastic GD samples the gradient

$$\Delta\theta = \alpha \left(V_\pi(x) - \hat{V}_\theta(x) \right) \nabla_\theta \hat{V}_\theta(x)$$

Approximating value fn. by (stochastic) gradient descent

In the previous slide, we assumed to know the true value function $V_\pi \rightarrow$ in RL there is no supervisor, only reward

In practice, we use a *target* for V_π

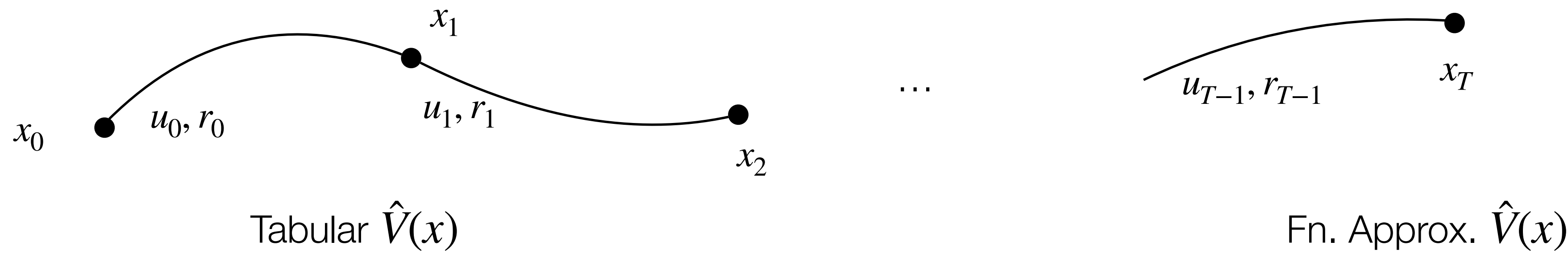
- Monte-Carlo: the target is the return

$$\Delta\theta = \alpha \left(\textcolor{red}{G_t} - \hat{V}_\theta(x_t) \right) \nabla_\theta \hat{V}_\theta(x_t)$$

- Temporal-Difference: the target is the TD target

$$\Delta\theta = \alpha \left(\textcolor{red}{r_t + \gamma \hat{V}_\theta(x_{t+1})} - \hat{V}_\theta(x_t) \right) \nabla_\theta \hat{V}_\theta(x_t)$$

Intuition



MC

1) Compute return $G_t = r_t + \gamma r_{t+1} + \dots, \forall t$

2) Update estimate

$$\hat{V}(x) = \left[\hat{V}(x_t) + \alpha(G_t - \hat{V}(x_t)) \right]$$

1) Collect dataset $\mathcal{D} = \{(x_t, G_t)\}$

2) Update θ

$$\theta = \theta + \alpha \left(G_t - \hat{V}_\theta(x_t) \right) \nabla_\theta \hat{V}_\theta(x_t)$$

TD

1) Compute target $r_t + \gamma \hat{V}(x_{t+1}), \forall t$

2) Update estimate

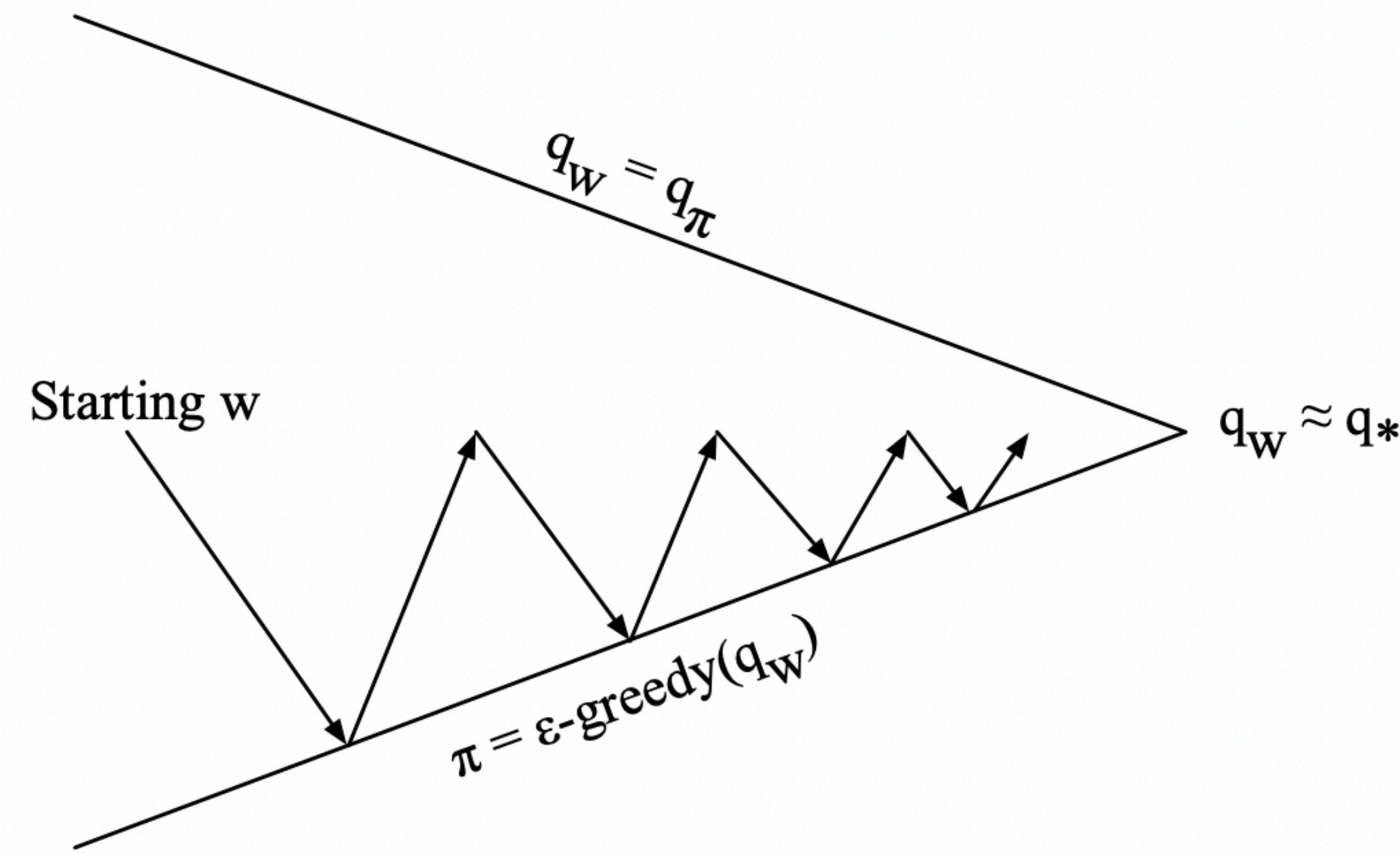
$$\hat{V}(x) = \left[\hat{V}(x_t) + \alpha(r_t + \gamma \hat{V}(x_{t+1}) - \hat{V}(x_t)) \right]$$

1) Collect dataset $\mathcal{D} = \{(x_t, r_t + \gamma \hat{V}_\theta(x_t))\}$

2) Update estimate

$$\theta = \theta + \alpha \left(r_t + \gamma \hat{V}_\theta(x_t) - \hat{V}_\theta(x_t) \right) \nabla_\theta \hat{V}_\theta(x_t)$$

Control with function approximation



Policy evaluation **Approximate** policy evaluation, $\hat{q}(\cdot, \cdot, \mathbf{w}) \approx q_\pi$
Policy improvement ϵ -greedy policy improvement

Action-value function approximation

Exactly the same intuitions apply when we try to approximate the action value function:

- Minimize the mean-squared error between the estimated value $\hat{Q}_\theta(x, u)$ and the true value $Q_\pi(x, u)$

$$J(\theta) = \mathbb{E}_\pi \left[Q_\pi(x, u) - \hat{Q}_\theta(x, u) \right]$$

- Use stochastic gradient descent to find a local minimum

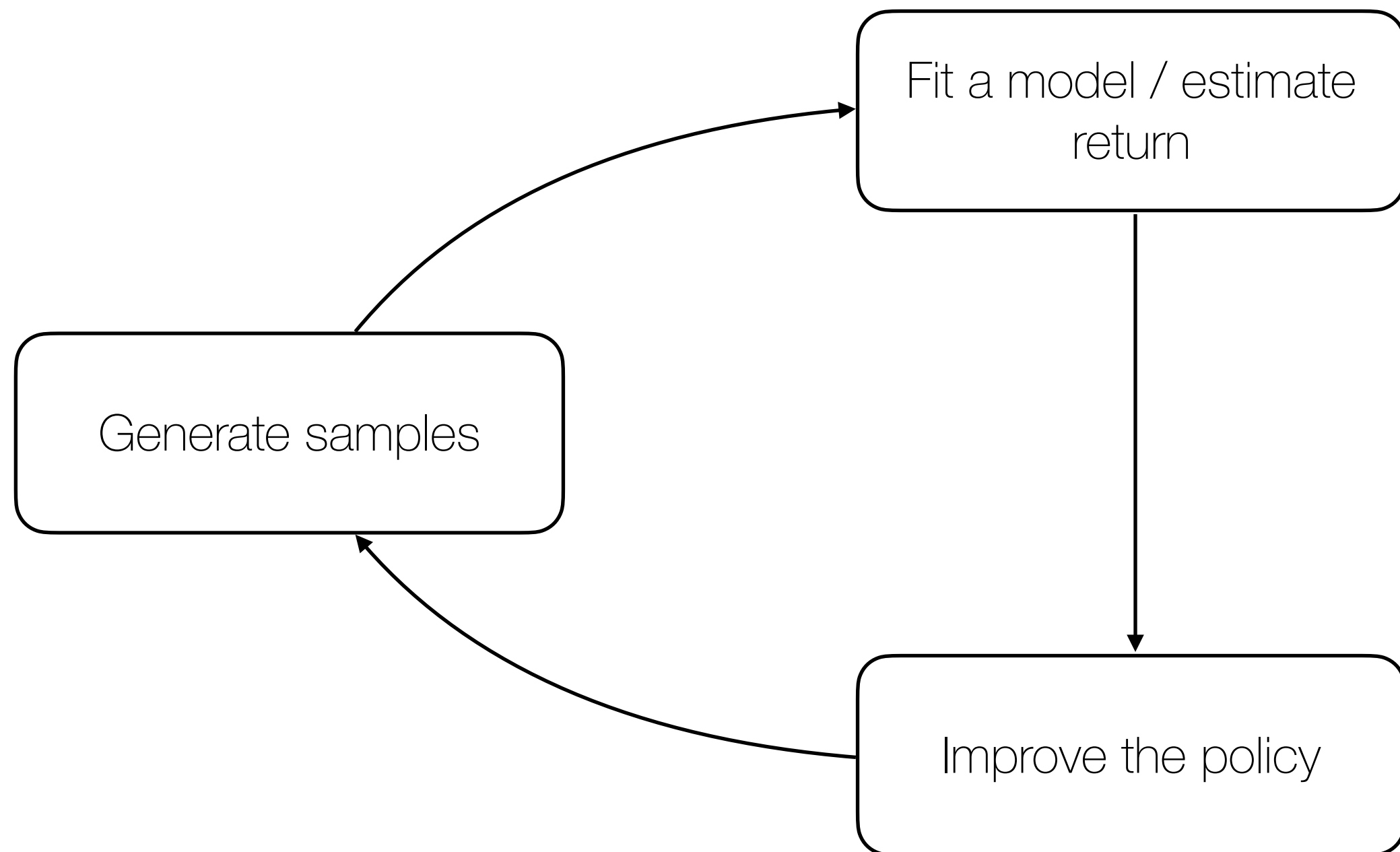
$$\Delta\theta = \alpha \left(Q_\pi(x, u) - \hat{Q}_\theta(x, u) \right) \nabla_\theta \hat{Q}_\theta(x, u)$$

Fitted Q-Iteration: update θ via stochastic gradient descent on TD target

$$\Delta\theta = \alpha \left(r_t + \gamma \max_{u'_{t+1}} Q_\theta(x_{t+1}, u'_{t+1}) - \hat{Q}_\theta(x_t, u_t) \right) \nabla_\theta \hat{Q}_\theta(x_t, u_t)$$

The skeleton of fitted Q-learning

Run the policy and observe
 (x_t, u_t, r_t, x_{t+1})



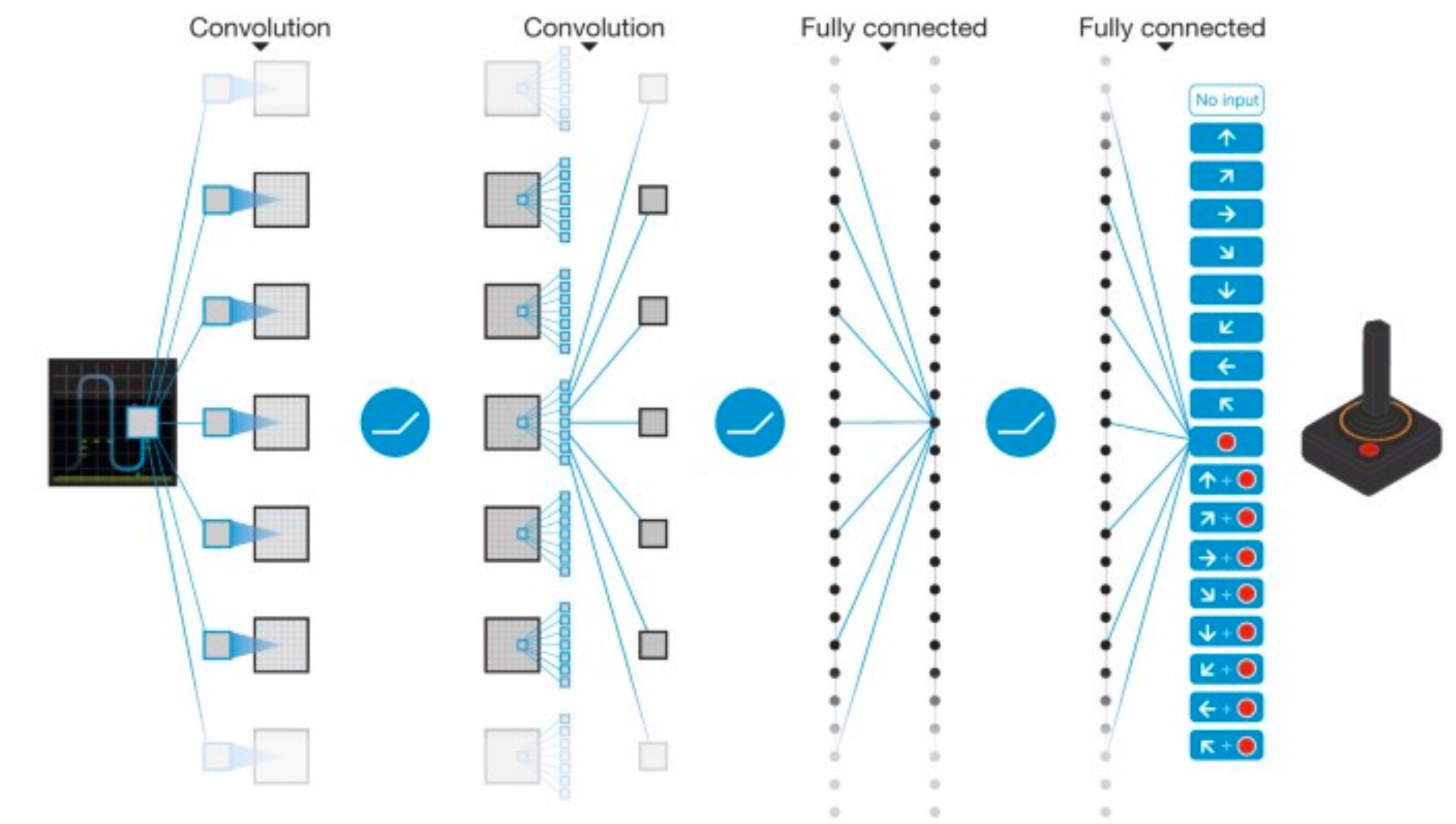
Set target
 $y_t \leftarrow r_t + \gamma \max_u Q_\theta(x_{t+1}, u)$

Update θ to minimize
 $J(\theta) = \mathbb{E}_\pi [y_t - Q_\theta(x_t, u_t)]$

set $\pi(x_t) = \arg \max_a Q_\theta(x_t, u_t)$

Deep Q-Networks (DQN)

One of the most popular Deep RL algorithms and arguably one of the first successes of RL with neural networks



(1) Use **deep neural nets** to represent Q_θ in Q-learning

(2) Uses **experience replay** and **fixed Q-targets**

Deep Q-Networks (DQN)

(2) Uses **experience replay** and **fixed Q-targets**

- These two ideas turned out to be very important to stabilize training. Specifically, these concepts attempt to solve two issues:
 - i) Samples within a trajectory are highly correlated → makes supervised learning unstable
 - ii) The target $r_t + \gamma \max_{u'_{t+1}} Q_\theta(x_{t+1}, u'_{t+1})$ is a moving target (i.e., as we update θ , the target of our regression also changes)

Intuitively:

- Take action u_t according to ϵ -greedy policy
- Store transition (x_t, u_t, r_t, x_{t+1}) in replay memory \mathcal{D}
- Sample batch of transitions $\{(x_t, u_t, r_t, x_{t+1})_i\}_{i=1}^B$ from \mathcal{D} (**Experience replay decorrelates data**)
- Compute Q-learning targets w.r.t. old, fixed parameters ϕ
- Optimize MSE between Q-network prediction and Q-learning targets (**Fixed targets stabilize the objective**)

$$J(\theta) = \mathbb{E}_{(x_t, u_t, r_t, x_{t+1}) \sim \mathcal{D}} \left[r_t + \gamma \max_u Q_\phi(x_{t+1}, u) - \hat{Q}_\theta(x_t, u_t) \right]$$

Next time

- Model-free RL: policy optimization methods