# CS/Stat 184 Introduction to Reinforcement Learning

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CONTENTS 0.1. OVERVIEW

Welcome to the study of reinforcement learning! This set of lecture notes accompanies the undergraduate course CS/STAT 184 and is intended to be a friendly yet rigorous introduction to this exciting and active subfield of machine learning. Here are some questions you might have before embarking on this journey:

What is reinforcement learning (RL)? Broadly speaking, RL is a subfield of machine learning that studies how an agent can learn to make sequential decisions in an environment.

Why study RL? RL provides a powerful framework for attacking a wide variety of problems, including robotic control, video games and board games, resource management, language modelling, and more. It also provides an interdisciplinary paradigm for studying animal and human behavior. Many of the most stunning results in machine learning, ranging from AlphaGo to ChatGPT, are built on top of RL.

**Is this book for me?** This book assumes familiarity with multivariable calculus, linear algebra, and probability. For Harvard undergraduates, this would be fulfilled by Math 21a, Math 21b, and Stat 110. Stat 111 is strongly recommended but not required. Here is a non-comprehensive list of topics of which this book will assume knowledge:

- Linear Algebra: Vectors, matrices, matrix multiplication, matrix inversion, eigenvalues and eigenvectors, and the Gram-Schmidt process.
- **Multivariable Calculus:** Partial derivatives, gradient, directional derivative, and the chain rule.
- **Probability:** Random variables, probability distributions, expectation, variance, covariance, conditional probability, Bayes' rule, and the law of total probability.

How does reinforcement learning differ from other machine learning paradigms? Here is a list of comparisons:

• Supervised learning. Supervised learning concerns itself with learning a mapping from inputs to outputs (e.g. image classification). Typically the data takes the form of input-output pairs that are assumed to be sampled independently from some generating distribution. In RL, however, the data is generated by the agent interacting with the environment, meaning the observations depend on the agent's behaviour and are not independent from each other. This requires a more general set of tools.

Conversely, supervised learning is a well-studied field that provides many useful tools for RL. For example, it may be useful to use supervised learning to predict how valuable a given state is, or to predict the probability of transitioning to a given state.

# 0.1 Overview

Chapter 1 introduces **Markov Decision Processes**, the dominant mathematical framework for studying RL. We'll discuss **dynamic programming** algorithms for solving MDPs, including

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policy evaluation, policy iteration, and value iteration.

Chapter 2 then discusses **multi-armed bandits**, a simpler problem that is often used as a warm-up to RL.

## 0.2 Notation

We will use the following notation throughout the book. This notation is inspired by Sutton and Barto and AJKS .

- s A state.
- a An action.
- r A reward.
- p A probability.
- $\pi$  A policy.
- V A value function.
- Q An action-value function.
- A An advantage function.
- $\gamma$  A discount factor.
- au A trajectory.
- $\mathcal{S}$  A state space.
- $\mathcal{A}$  An action space.

# 0.3 Challenges of reinforcement learning

**Exploration-exploitation tradeoff.** Should the agent try a new action or stick with the action that it knows is good?

**Prediction.** The agent might want to predict the value of a state or state-action pair.

**Policy computation (control).** In a complex environment, even if the dynamics are known, it can still be challenging to compute the best policy.

## 0.4 Resources

Inspired by the Stat 110 textbook and Stat 111 lecture notes.

This book seeks to provide an intuitive understanding before technical treatment.

Refer to AJ Sutton and Barto, onlin sources.

cite

cite

Add notation

CONTENTS 0.4. RESOURCES

# **C**ontents

# Chapter 1

# **Bandits**

The **multi-armed bandits** (MAB) setting is a simple but powerful setting for studying the basic challenges of RL. In this setting, an agent repeatedly chooses from a fixed set of actions, called **arms**, each of which has an associated reward distribution. The agent's goal is to maximize the total reward it receives over some time period.

States	Actions	Rewards
None	Finite	Stochastic

In particular, we'll spend a lot of time discussing the **Exploration-Exploitation Tradeoff**: should the agent choose new actions to learn more about the environment, or should it choose actions that it already knows to be good?

### **Example 1.0.1: Online advertising**

Let's suppose you, the agent, are an advertising company. You have K different ads that you can show to users; For concreteness, let's suppose there's just a single user. You receive 1 reward if the user clicks the ad, and 0 otherwise. Thus, the unknown reward distribution associated to each ad is a Bernoulli distribution defined by the probability that the user clicks on the ad. Your goal is to maximize the total number of clicks by the user.

examples nical trials, te. etc.

In this chapter, we will introduce the multi-armed bandits setting, and discuss some of the challenges that arise when trying to solve problems in this setting. We will also introduce some of the key concepts that we will use throughout the book, such as regret and exploration-exploitation tradeoffs.

## 1.1 Multi-Armed Bandits

#### Remark 1.1.1: Namesake

The name "multi-armed bandits" comes from slot machines in casinos, which are often called "one-armed bandits" since they have one arm (the lever) and take money from the player.

Let K denote the number of arms. We'll label them  $1, \ldots, K$  and use *superscripts* to indicate the arm index; since we seldom need to raise values to a power, this won't cause much confusion. For simplicity, we'll assume rewards are *bounded* between 0 and 1. Then each arm has an unknown reward distribution  $\nu^k \in \triangle([0,1])$  with mean  $\mu^k = \mathbb{E}_{r \sim \nu^k}[r]$ .

Formally speaking, the agent's interaction with the MAB environment can be described by the following process:

```
# multi-armed bandits
for timestep in range(0, T):
    # Agent chooses an arm
    k = agent.choose_arm()

# Environment generates a reward
    r = env.generate_reward(k)

# Agent observes the reward
    agent.observe_reward(k, r)
```

What's the optimal strategy for the agent, i.e. the one that achieves the highest expected reward? Convince yourself that the agent should try to always pull the arm with the highest expected reward  $\mu^* := \max_{k \in [K]} \mu^k$ .

The goal, then, can be rephrased as to minimize the regret, defined below:

#### **Definition 1.1.1: Regret**

The agent's **regret** after T timesteps is the difference between the total reward it observes and the total reward it *would* have received if it had always pulled the optimal arm:

$$\mathsf{Regret}_T := \sum_{t=0}^{T-1} \mu^{\star} - \mu^{a_t} \tag{1.1}$$

Note that this depends on the *true means* of the pulled arms, *not* the observed rewards.

Often we consider the **expected regret**  $\mathbb{E}[\mathsf{Regret}_T]$ , where the randomness comes from the agent's strategy.

Maybe switch more tradition pseudocode? Or set up so Python "interfaces" near the start?

e Big O on in apIdeally, we'd like to asymptotically achieve **zero regret**, i.e.  $\mathbb{E}[\mathsf{Regret}_T] = o(T)$ .

## 1.1.1 Pure exploration (random guessing)

A trivial strategy is to always choose arms at random (i.e. "pure exploration"). What is the expected regret of this strategy?

$$\begin{split} \mathbb{E}[\mathsf{Regret}_T] &= \sum_{t=0}^{T-1} \mathbb{E}[\mu^\star - \mu^{a_t}] \\ &= T(\mu^\star - \bar{\mu}) > 0 \\ \text{where} \quad \bar{\mu} := \mathbb{E}[\mu^{a_t}] = \frac{1}{K} \sum_{k=1}^K \mu^k \end{split}$$

This scales as  $\Theta(T)$ , i.e. *linear* in the number of timesteps T. There's no learning here: the agent doesn't use any information about the environment to improve its strategy.

## 1.1.2 Pure greedy

How might we improve on pure exploration? Instead, we could try each arm once, and then commit to the one with the highest observed reward. We'll call this the **pure greedy** strategy.

```
# exploration phase
for k in range(K):
    observed_rewards[k] = env.generate_reward(k)
k_hat = argmax(observed_rewards)

# exploitation phase
for t in range(T - K):
    r = env.generate_reward(k_hat)
```

# observed\_rewards is an array of length K

How does the expected regret of this strategy compare to that of pure exploration? We'll do a more general analysis in the following section. Now, for intuition, suppose there's just two arms, with Bernoulli reward distributions given by  $\mu^1 > \mu^2$ .

Let's let  $r^1$  be the random reward from the first arm and  $r^2$  be the random reward from the second. If  $r^1 > r^2$ , then we achieve zero regret. Otherwise, we achieve regret  $T(\mu^1 - \mu^2)$ . Thus, the expected regret is simply:

$$\mathbb{E}[\mathsf{Regret}_T] = \mathbb{P}(r^1 < r^2) \cdot T(\mu^1 - \mu^2) + c$$
  
=  $(1 - \mu^1)\mu^2 \cdot T(\mu^1 - \mu^2) + c$ 

Which is still  $\Theta(T)$ , the same as pure exploration!

Can we do better? For one, we could reduce the variance of the reward estimates by pulling each arm *multiple times*. This is called the **explore-then-commit** strategy.

## 1.1.3 Explore-then-commit

Let's pull each arm  $N_{\text{explore}}$  times, and then commit to the arm with the highest observed average reward. What is the expected regret of this strategy?

```
# avg_reward is an array of length K

# exploration phase
for k in range(K):
    total = 0
    for i in range(N_explore):
        total += env.generate_reward(k)
    avg_reward[k] = total / N_explore
k_hat = argmax(avg_reward)

# exploitation phase
for t in range(T):
    r = env.generate_reward(k_hat)

(Note that the "pure greedy" strategy is just the special case where N_explore = 1.)
```

Let's analyze the expected regret of this strategy by splitting it up into the exploration and exploitation phases.

**Exploration phase.** This phase takes  $N_{\text{explore}}K$  timesteps. Since at each step we incur at most 1 regret, the total regret is at most  $N_{\text{explore}}K$ .

**Exploitation phase.** This will take a bit more effort. We'll ultimately prove that:

- 1. For any total time T,
- 2. We can choose  $N_{\sf explore}$  such that
- 3. With arbitrarily high probability, the regret is sublinear.

Let  $\hat{k}:=\arg\max_{k\in[K]}\hat{\mu}^k$  be the arm we choose to exploit. We know the regret from the exploitation phase is

$$T_{\text{exploit}}(\mu^{\star} - \mu^{\hat{k}})$$
 where  $T_{\text{exploit}} := T - N_{\text{explore}}K$ .

So we'd like to bound  $\mu^\star - \mu^{\hat k} = o(1)$  (as a function of T) in order to achieve sublinear regret. How can we do this?

nk to ap-

Let's use  $\Delta^k = \hat{\mu}^k - \mu^k$  to denote how far the mean estimate for arm k is from the true mean. **Hoeffding's inequality** tells us that, for a given arm k, since the rewards from that arm are i.i.d.,

$$\mathbb{P}\left(|\Delta^k| > \sqrt{\frac{\ln(2/\delta)}{2N_{\text{explore}}}}\right) \le \delta. \tag{1.2}$$

But note that we can't directly apply this to  $\hat{k}$  since  $\hat{k}$  is itself a random variable. Instead, we need to "uniform-ize" this bound across all the arms, i.e. bound the residual across all the arms simultaneously, so that the resulting bound will apply no matter what  $\hat{k}$  "crystallizes" to.

The **union bound** provides a simple way to do this: The probability of error (i.e. the l.h.s. of 1.2) for a *single* arm is at most  $\delta$ , so the probability that *at least one* of the arms is far from the mean is at most  $K\delta$ . Setting  $\delta' := K\delta$  and taking the complement of both sides, we have

$$\mathbb{P}\left(\forall k \in [K], |\Delta^k| \leq \sqrt{\frac{\ln(2K/\delta')}{2N_{\mathsf{explore}}}}\right) \geq 1 - \delta'$$

Then to apply this bound to  $\hat{k}$  in particular, we can apply the useful trick of "adding zero":

$$\begin{split} \mu^{k^\star} - \mu^{\hat{k}} &= \mu^{k^\star} - \mu^{\hat{k}} + (\hat{\mu}^{k^\star} - \hat{\mu}^{k^\star}) + (\hat{\mu}^{\hat{k}} - \hat{\mu}^{\hat{k}}) \\ &= \Delta^{\hat{k}} - \Delta^{k^*} + \underbrace{(\hat{\mu}^{k^\star} - \hat{\mu}^{\hat{k}})}_{\leq 0 \text{ by definition of } \hat{k}} \\ &\leq 2\sqrt{\frac{\ln(2K/\delta')}{2N_{\text{explore}}}} \text{ with probability at least } 1 - \delta' \end{split}$$

Putting this all together, we've shown that, with probability  $1 - \delta'$ ,

$$\mathsf{Regret}_T \leq N_{\mathsf{explore}} K + T_{\mathsf{exploit}} \cdot \sqrt{\frac{2 \ln(2K/\delta')}{N_{\mathsf{explore}}}}.$$

Note that it suffices for  $N_{\rm explore}$  to be on the order of  $\sqrt{T}$  to achieve sublinear regret. In particular, we can find the optimal  $N_{\rm explore}$  by setting the derivative of the r.h.s. to zero:

$$\begin{split} K - T_{\rm exploit} \cdot \frac{1}{2} \sqrt{\frac{2 \ln(2K/\delta')}{N_{\rm explore}^3}} &= 0 \\ N_{\rm explore} &= \left(T_{\rm exploit} \cdot \frac{\sqrt{\ln(2K/\delta')/2}}{K}\right)^{2/3} \end{split}$$

Plugging this into the expression for the regret, we have (still with probability  $1 - \delta'$ )

$$\mathsf{Regret}_T \leq 3T^{2/3} \sqrt[3]{K \ln(2K/\delta')/2}$$

The ETC algorithm is rather "abrupt" in that it switches from exploration to exploitation after a fixed number of timesteps. In practice, it's often better to use a more gradual transition, which brings us to the *epsilon-greedy* algorithm.

## 1.1.4 Epsilon-greedy

Instead of doing all of the exploration and then all of the exploitation separately – which additionally requires knowing the time horizon beforehand – we can instead interleave exploration and exploitation by, at each timestep, choosing a random action with some probability. We call this the **epsilon-greedy** algorithm.

```
# epsilon-greedy
# random() samples from the uniform distribution on [0, 1]
for t in range(T):
    if random() < epsilon(t):
        # exploration
        k = random_choice(K)
    else:
        # exploitation
        # calculate averages using element-wise division
        k = argmax(total_reward / num_pulls)
    r = env.generate_reward(k)
    total_reward[k] += r
    num_pulls[k] += 1</pre>
```

Note that  $\epsilon$  can vary over time. In particular we might want to gradually *decrease*  $\epsilon$  as we learn more about the environment over time.

It turns out that setting  $\epsilon_t = \sqrt[3]{K \ln(t)/t}$  also achieves a regret of  $\widetilde{O}(t^{2/3}\sqrt[3]{K})$  (ignoring the logarithmic factors).

In the ETC case, we had to set  $N_{\rm explore}$  based on the total number of timesteps T. But the epsilon-greedy algorithm actually handles the exploration *automatically*: the regret rate holds for any t, and doesn't depend on the final horizon T.

But the way these algorithms explore is rather naive: we've been exploring *uniformly* across all the arms. But what if we could be smarter about it? In particular, what if we could explore more for arms that we're less certain about? This brings us to the **Upper Confidence Bound** (UCB) algorithm.

## 1.1.5 Upper Confidence Bound (UCB)

Insert optime epsilon and a analysis / hi probability b We'll estimate *confidence intervals* for the mean of each arm, and then choose the arm with the highest *upper confidence bound*. This operates on the principle of **the benefit of the doubt** (i.e. optimism in the face of uncertainty): we'll choose the arm that we're most optimistic about.

In particular, we'd like to compute some upper confidence bound  $M^k_t$  for arm k at time t and then choose  $a_t := \arg\max_{k \in [K]} M^k_t$ . But how should we compute  $M^k_t$ ?

In our regret analysis for ETC, we were able to compute this bound using Hoeffding's inequality. Hoeffding's inequality assumes that the number of samples is *fixed*, which was true in ETC. However, in UCB, the number of times we pull each arm depends on the agent's actions, which in turn depend on the random rewards and are therefore stochastic. So we *can't* use Hoeffding's inequality directly.

Instead, we'll apply the same trick we used in the ETC analysis: we'll use the **union bound** to compute a looser upper confidence bound that holds *uniformly* across time and across the different arms. Let's introduce some notation to discuss this.

Let  $N_t^k$  denote the (random) number of times arm k has been pulled within the first t timesteps, and  $\hat{\mu}_t^k$  denote the sample average of those pulls. That is,

$$\begin{split} N_t^k &:= \sum_{\tau=t}^{t-1} \mathbf{1} \{ a_\tau = k \} \\ \hat{\mu}_t^k &:= \frac{1}{N_t^k} \sum_{t=0}^{t-1} \mathbf{1} \{ a_\tau = k \} r_\tau. \end{split}$$

To achieve the "fixed sample size" assumption, we'll need to shift our index from time to number of samples from each arm. In particular, we'll define  $\widetilde{r}_n^k$  to be the nth sample from arm k, and  $\widetilde{\mu}_n^k$  to be the sample average of the first n samples from arm k. Then, for a fixed n, this satisfies the "fixed sample size" assumption, and we can apply Hoeffding's inequality to get a bound on  $\widetilde{\mu}_n^k$ .

So how can we extend our bound on  $\widetilde{\mu}_n^k$  to  $\widehat{\mu}_t^k$ ? Well, we know  $N_t^k \leq t$  (which would be the case if we had pulled arm k every time). So we can apply the same trick as last time, where we uniform-ize across all possible values of  $N_t^k$ . In particular, we let  $\delta':=t\delta$ , giving us

$$\mathbb{P}\left(\forall n \le t, |\widetilde{\mu}_n^k - \mu^k| \le \sqrt{\frac{\ln(2t/\delta')}{2n}}\right) \ge 1 - \delta'$$

Now we can safely set  $n := N_t^k$  to achieve

 $\mathbb{P}\left(|\hat{\mu}_t^k - \mu^k| \le \sqrt{\frac{\ln(2t/\delta')}{2N_t^k}}\right) \ge 1 - \delta'.$ 

ate more on

This bound would then suffice for applying the UCB algorithm! That is, the upper confidence bound for arm k would be

$$M_t^k := \hat{\mu}_t^k + \sqrt{\frac{\ln(2t/\delta')}{2N_t^k}}$$

, where we can choose  $\delta'$  depending on how tight we want the interval to be. A smaller  $\delta'$  would give us a larger yet "more confident" interval, and vice versa.

Intuitively, this prioritizes arms where:

- 1.  $\hat{\mu}_t^k$  is large, i.e. the arm has a high sample average, and we'd choose it for *exploitation*, and
- 2.  $\sqrt{\frac{\ln(2t/\delta')}{2N_t^k}}$  is large, i.e. we're still uncertain about the arm, and we'd choose it for *exploration*.

As desired, this explores in a smarter, *adaptive* way compared to the previous algorithms. Does it achieve lower regret?

First we'll bound the regret incurred at each timestep. Then we'll bound the *total* regret across timesteps.

For the sake of analysis, we'll use a slightly looser bound that applies across the whole time horizon and across all arms. We'll omit the derivation since it's similar to the above (walk through it yourself for practice).

$$\begin{split} \mathbb{P}\left(\forall k \leq K, t < T, |\hat{\mu}_t^k - \mu^k| \leq B_t^k\right) \geq 1 - \delta'' \\ \text{where} \quad B_t^k := \sqrt{\frac{\ln(2TK/\delta'')}{2N_t^k}}. \end{split}$$

Intuitively,  $B_t^k$  denotes the *width* of the CI for arm k at time t. Then, assuming the above uniform bound holds (which occurs with probability  $1 - \delta''$ ), we can bound the regret at each timestep as follows:

$$\begin{split} \mu^{\star} - \mu^{a_t} &\leq \hat{\mu}_t^{k^*} + B_t^{k^*} - \mu^{a_t} & \text{applying UCB to arm } k^{\star} \\ &\leq \hat{\mu}_t^{a_t} + B_t^{a_t} - \mu^{a_t} & \text{since UCB chooses } a_t = \underset{k \in [K]}{\arg\max} \hat{\mu}_t^k + B_t^k \\ &\leq 2B_t^{a_t} & \text{since } \hat{\mu}_t^{a_t} - \mu^{a_t} \leq B_t^{a_t} \text{ by definition of } B_t^{a_t} \end{split}$$

Summing this across timesteps gives

$$\begin{aligned} \mathsf{Regret}_T & \leq \sum_{t=0}^{T-1} 2B_t^{a_t} \\ & = \sqrt{2 \ln(2TK/\delta'')} \sum_{t=0}^{T-1} (N_t^{a_t})^{-1/2} \\ & \sum_{t=0}^{T-1} (N_t^{a_t})^{-1/2} = \sum_{t=0}^{T-1} \sum_{k=1}^{K} \mathbf{1}\{a_t = k\} (N_t^k)^{-1/2} \\ & = \sum_{k=1}^{K} \sum_{n=1}^{N_t^k} n^{-1/2} \\ & \leq K \sum_{n=1}^{T} n^{-1/2} \\ & \leq K \sum_{n=1}^{T} n^{-1/2} \\ & \sum_{t=0}^{T} n^{-1/2} \leq 1 + \int_{1}^{T} x^{-1/2} \, \mathrm{d}x \\ & = 1 + (2\sqrt{x})_1^T \\ & = 2\sqrt{T} - 1 \\ & \leq 2\sqrt{T} \end{aligned}$$

Putting everything together gives

$$\begin{aligned} \mathsf{Regret}_T & \leq 2K\sqrt{2T\ln(2TK/\delta'')} & \text{with probability } 1 - \delta'' \\ & = \widetilde{O}(\sqrt{T}) \end{aligned}$$

In fact, we can do a more sophisticated analysis to show  $\operatorname{Regret}_T = \widetilde{O}(\sqrt{TK})$ .

# 1.2 Thompson sampling

include?

# Chapter 2

# **Markov Decision Processes**

# **Contents**

Earlier, we said that RL is a subfield of machine learning that studies how an agent can learn to make sequential decisions in an environment. How can we *formalize* this task in a way that is both *sufficiently general* yet also tractable enough for *fruitful analysis*? What are the key properties we'd like to capture?

In this chapter, we'll turn to **Markov decision processes** as a simple yet general formalism for solving decision problems. The key idea is that the agent interacts with the environment by taking actions, which in turn affect the environment and the agent's future observations. We'll see that this simple idea leads to a rich set of problems and algorithms.

Let's begin with a motivating example of

#### **Definition 2.0.1: Markov Decision Process**

The key components of a Markov decision process are:

- 1. The **state** that the agent interacts with. We use S to denote the set of possible states, called the **state space**.
- 2. The **agent** and the **actions** that it can take. We use A to denote the set of possible actions, called the **action space**.
- 3. The **reward** signal. In this course we'll take it to be a deterministic function of a state-action pair, i.e.  $r: \mathcal{S} \times \mathcal{A} \to \mathbb{R}$ . In general, though, the reward function can also be stochastic, and it can also accept the *resulting* state as an argument; that is,  $r: \mathcal{S} \times \mathcal{A} \times \mathcal{S} \to \Delta(\mathbb{R})$ .
- 4. The **state transitions** (a.k.a. **dynamics**) that describe what state we **transition to** after taking an action. We'll denote this by  $P: \mathcal{S} \times \mathcal{A} \to \Delta(\mathcal{S})$ .
- 5. A discount factor  $\gamma \in [0,1)$ . We'll see later that this ensures that the *return*, or total reward, is well-defined in infinite-horizon problems.
- 6. Some initial state distribution  $\rho \in \Delta(S)$ .

Combined together, we call these a Markov decision process

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$$M = (\mathcal{S}, \mathcal{A}, P, r, \gamma, \rho).$$

The reason we call it a *Markov* decision process is that the transition function only depends on the "current" state and action. Formally, this implies that the state process satisfies the **Markov property**, that is,

$$\mathbb{P}(s_{t+1} \mid (s_{\tau}, a_{\tau})_{\tau=0}^{t}) = P(s_{t+1} \mid s_{t}, a_{t}).$$

### Example 2.0.1: Examples of MDPs

**Board games and video games** are often MDPs. For example, in chess or Go, the state of the game only depends on the pieces on the board and not on the previous history. Several possible reward functions could be possible, e.g. +1 upon winning the game and 0 otherwise, or to receive reward upon taking the opponent's pieces. The state transitions are based on the opponent's moves.

**Robotic control** can be framed as an MDP task. In this setting, physics provides the state transitions. A possible action might be activating a motor to move forwards. The reward function could be designed based on the task; for example, one could reward the robot for arriving at a desired location.

We'll distinguish between **finite-horizon** MDPs, where the agent eventually enters a **terminal state**, and **infinite-horizon** MDPs, where the agent might keep going on and on.

We call the total reward the **return**. For finite-horizon MDPs, we can just add up the rewards:

$$G_t := R_t + R_{t+1} + \dots + R_T,$$

where T is the number of time steps and  $R_t := r(S_t, A_t)$ . However, for infinite-horizon problems (i.e.  $T = \infty$ ), in order for this to be well-defined, we need to *discount* future rewards:

$$G_t := R_t + \gamma R_{t+1} + \dots + \gamma^{\tau - t} R_{\tau} + \dots = \sum_{\tau = t}^{\infty} \gamma^{\tau - t} R_{\tau}.$$

Can you see why this ensures that  $G_t$  is finite?

Note that we recover the finite-horizon definition by letting  $\gamma = 1$  and T be finite.

Our key goal in a reinforcement learning task is to maximize expected return.

Why can't we just maximize the current reward at each timestep, i.e. use a greedy strategy? Well, in RL as in real life, often making greedy decisions (e.g. procrastinating) will leave you worse off than if you make some short-term sacrifices for long-term gains.

We call the "video recording" of states, actions, and rewards a trajectory

$$\xi_t = (s_{\tau}, a_{\tau}, r_{\tau})_{\tau=0}^t$$

## 2.1 Policies and value functions

A **policy**  $\pi$  describes the agent's strategy: which actions it takes in a given situation.

Policies can either be **deterministic** (in the same situation, the agent will always take the same action) or **stochastic** (in the same situation, the agent will sample an action from a distribution).

What do I mean by "situation"? In the most general setting, this could include all of the states, actions, and rewards in the trajectory so far.

However, due to the Markov assumption, the state transitions only depend on the current state. Thus a **stationary** policy  $\pi: \mathcal{S} \to \Delta(\mathcal{A})$  — one that only depends on the current state — can do just as well.

Fix a policy  $\pi$ . We'd like a concise way to refer to the expected return when *starting in a given state* and acting according to  $\pi$ . We call this the **value function** of  $\pi$  and denote it by

$$V^{\pi}(s) := \underset{\pi}{\mathbb{E}}[G_0 \mid S_0 = s]$$

We start at time 0 without loss of generality; can you see why we could have chosen to start at any time?

Similarly, we can define the **action-value function** of  $\pi$  (aka the **Q-function**) as the expected return when starting in a given state and taking a given action:

$$Q^{\pi}(s,a) := \mathbb{E}[G_0 \mid S_0 = s, A_0 = a]$$

## 2.1.1 Bellman self-consistency equations

Note that we can break down the return as

$$G_t = R_t + \gamma G_{t+1}$$

the reward from the *current time-step* plus the total reward from from *future time-steps*. It turns out that this simple observation, along with linearity of expectation, gives us a set of equations to solve for the value function analytically!

Let's expand out the definition of the value function to see what I mean. Let's first consider the simple case where  $\pi:\mathcal{S}\to\mathcal{A}$  is deterministic:

$$V^{\pi}(s) := \underset{\pi}{\mathbb{E}}[G_0 \mid S_0 = s]$$

$$= r(s, \pi(s)) + \gamma \underset{s' \sim P(\cdot \mid s, \pi(s))}{\mathbb{E}} \underset{\pi}{\mathbb{E}}[G_1 \mid S_1 = s']$$

$$= r(s, \pi(s)) + \gamma \sum_{s' \in \mathcal{S}} P(s' \mid s, \pi(s)) V^{\pi}(s').$$

This is a set of |S| equations (one per state) in |S| unknowns (the value of each state), which we can solve for  $V^{\pi}$ .

For stochastic policies, we simply average out over the relevant quantities:

$$V^{\pi}(s) := \mathbb{E}[G_0 \mid S_0 = s]$$

$$= \mathbb{E}[R_0 + \gamma G_1 \mid S_0 = s]$$

$$= \mathbb{E}_{a \sim \pi(\cdot \mid s)} \left[ r(s, a) + \gamma \mathbb{E}_{s' \sim P(\cdot \mid s, a)} \mathbb{E}[G_1 \mid S_1 = s'] \right]$$

$$= \sum_{a} \pi(a \mid s) \left[ r(s, a) + \gamma \sum_{s'} P(s' \mid s, a) V^{\pi}(s') \right].$$

These are called the **Bellman self-consistency equations**. They encapsulate that the value function's prediction of the current state must be consistent with its prediction of other states.

Can you write the Bellman self-consistency equations for the action-value function?

# 2.2 Tabular MDPs

When the state and action space are finite and small, we can think of the value function and Q-function as *lookup tables* with each cell corresponding to the value of a state (or state-action pair). We can neatly express quantities as vectors and matrices:

$$r \in \mathbb{R}^{|\mathcal{S}| \times |\mathcal{A}|}, \quad P \in [0, 1]^{(|\mathcal{S}| \times |\mathcal{A}|) \times |\mathcal{S}|}, \quad \rho \in [0, 1]^{|\mathcal{S}|}, \quad \pi \in [0, 1]^{|\mathcal{A}| \times |\mathcal{S}|}, \quad V^{\pi} \in \mathbb{R}^{|\mathcal{S}|}, \quad Q^{\pi} \in \mathbb{R}^{|\mathcal{S}| \times |\mathcal{A}|}.$$

Make sure that these dimensions make sense!

Note that when the policy is deterministic, by definition, the actions can be determined from the state, and so we can chop off the action dimension in most cases:

$$r^{\pi} \in \mathbb{R}^{|\mathcal{S}|}, \quad P^{\pi} \in [0, 1]^{|\mathcal{S}| \times |\mathcal{S}|}, \quad \rho \in [0, 1]^{|\mathcal{S}|}, \quad \pi \in \mathcal{A}^{|\mathcal{S}|}, \quad V^{\pi} \in \mathbb{R}^{|\mathcal{S}|}.$$

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Then, rewriting the system of Bellman equations in this notation gives

$$V^{\pi} = r^{\pi} + \gamma P^{\pi} V^{\pi} \implies V^{\pi} = (I - \gamma P^{\pi})^{-1} r^{\pi}.$$

Note that we've assumed that  $I - \gamma P^{\pi}$  is invertible. Can you see why this is the case?

Recall that a linear operator, i.e. a square matrix, is invertible if and only if its null space is trivial; that is, it doesn't map any nonzero vector to zero. In this case, we can see that  $I - \gamma P^{\pi}$  is invertible because it maps any nonzero vector to a vector with at least one nonzero element.

# 2.3 Optimality

### Theorem 2.3.1: Value Iteration

Initialize:

$$V^0 \sim ||V^0||_{\infty} \in [0, 1/1 - \gamma]$$

Iterate until convergence:

$$V^{t+1} \leftarrow \mathcal{J}(V^t)$$

#### **Analysis**

This algorithm runs in  $O(|\mathcal{S}|^3)$  time since we need to perform a matrix inversion.

#### Theorem 2.3.2: Exact Policy Evaluation

Represent the reward from each state-action pair as a vector

$$R^{\pi} \in \mathbb{R}^{|\mathcal{S}|}$$
  $R_s^{\pi} = r(s, \pi(s))$ 

Also represent the state transitions

$$P^{\pi} \in \mathbb{R}^{|\mathcal{S} \times \mathcal{S}}$$
  $P^{\pi}_{s,s'} = P(s'|s, \pi(s))$ 

That is, row i of  $P^{\pi}$  is a distribution over the *next state* given that the current state is  $s_i$  and we choose an action using policy  $\pi$ .

Using this notation, we can express the Bellman consistency equation as

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$$\begin{pmatrix} \vdots \\ V^{\pi}(s) \\ \vdots \end{pmatrix} = \begin{pmatrix} \vdots \\ r(s, \pi(s)) \\ \vdots \end{pmatrix} + \gamma \begin{pmatrix} \vdots \\ P(s' \mid s, \pi(s)) \\ \vdots \end{pmatrix} \begin{pmatrix} \vdots \\ V^{\pi}(s') \\ \vdots \end{pmatrix}$$

$$V^{\pi} = R^{\pi} + \gamma P^{\pi} V^{\pi}$$

$$(I - \gamma P^{\pi}) V^{\pi} = R^{\pi}$$

$$V^{\pi} = (I - \gamma P^{\pi}) R^{\pi}$$

if  $I - \gamma P^{\pi}$  is invertible, which we can prove is the case.

## Theorem 2.3.3: Iterative Policy Evaluation

How can we calculate the value function  $V^{\pi}$  of a policy  $\pi$ ?

Above, we saw an exact function that runs in  $O(|\mathcal{S}|^2)$ . But say we really need a fast algorithm, and we're okay with having an approximate answer. Can we do better? Yes!

Using the same notation as above, let's initialize  $V^0$  such that the elements are drawn uniformly from  $[0,1/(1-\gamma)]$ .

Then we can iterate the fixed-point equation we found above:

$$V^{t+1} \leftarrow R + \gamma P V^t$$

How can we use this fast approximate algorithm?

### Theorem 2.3.4: Policy Iteration

Remember, for now we're only considering policies that are stationary and deterministic. There's  $|\mathcal{S}|^{|\mathcal{A}|}$  of these, so let's start off by choosing one at random. Let's call this initial policy  $\pi^0$ , using the superscript to indicate the time step.

Now for  $t = 0, 1, \ldots$ , we perform the following:

1. Policy Evaluation: First use the algorithm from earlier to calculate  $V^{\pi^t}(s)$  for all states s. Then use this to calculate the state-action values:

$$Q^{\pi^t}(s, a) = r(s, a) + \gamma \sum_{s'} P(s' \mid s, a) V^{\pi^t}(s')$$

2. *Policy Improvement*: Update the policy so that, at each state, it chooses the action with the highest action-value:

$$\pi^{t+1}(s) = \arg\max_{a} Q^{\pi^t}(s, a)$$

In other words, we're setting it to act greedily with respect to the new Q-function.

What's the computational complexity of this?

## 2.4 Finite Horizon MDPs

Suppose we're only able to act for H timesteps.

Now, instead of discounting, all we care about is the (average) total reward that we get over this time.

$$\mathbb{E}[\sum_{t=0}^{H-1} r(s_t, a_t)]$$

To be more precise, we'll consider policies that depend on the time. We'll denote the policy at timestep h as  $\pi_h : \mathcal{S} \to \mathcal{A}$ . In other words, we're dropping the constraint that policies must be stationary.

This is also called an episodic model.

Note that since our policy is nonstationary, we also need to adjust our value function (and Q-function) to account for this. Instead of considering the total infinite-horizon discounted reward like we did earlier, we'll instead consider the *remaining* reward from a given timestep onwards:

$$V_h^{\pi}(s) = \mathbb{E}\left[\sum_{\tau}^{H-1} r(s_{\tau}, a_{\tau}) \mid s_h = s, a_{\tau} = \pi_h(s_h)\right]$$
$$Q_h^{\pi}(s, a) = \mathbb{E}\left[\sum_{\tau}^{H-1} r(s_{\tau}, a_{\tau}) \mid (s_h, a_h) = (s, a)\right]$$

We can also define our Bellman consistency equations, by splitting up the total reward into the immediate reward (at this time step) and the future reward, represented by our state value function from that next time step:

$$Q_h^{\pi}(s, a) = r(s, a) + \underset{s' \sim P(s, a)}{\mathbb{E}} [V_{h+1}^{\pi}(s')]$$

### Theorem 2.4.1: Computing the optimal policy

We can solve for the optimal policy using dynamic programming.

• Base case. At the end of the episode (time step H-1), we can't take any more actions, so the Q-function is simply the reward that we obtain:

$$Q_{H-1}^{\star}(s,a) = r(s,a)$$

so the best thing to do is just act greedily and get as much reward as we can!

$$\pi_{H-1}^{\star}(s) = \arg \max_{a} Q_{H-1}^{\star}(s, a)$$

Then  $V_{H-1}^{\star}(s)$ , the optimal value of state s at the end of the trajectory, is simply whatever action gives the most reward.

$$V_{H-1}^{\star} = \max_{a} Q_{H-1}^{\star}(s, a)$$

• Recursion. Then, we can work backwards in time, starting from the end, using our consistency equations!

Note that this is exactly just value iteration and policy iteration combined, since our policy is nonstationary, so we can exactly specify its decisions at each time step!

#### **Analysis**

Total computation time  $O(H|\mathcal{S}|^2|\mathcal{A}|)$ 

# Chapter 3

# **Linear Quadratic Regulators**

## 3.1 Motivation

Have you ever tried balancing a pen upright on your palm? If not, try it! It's a lot harder than seems. Unlike the cases we studied the previous chapter, the state space and action space aren't finite, or even discrete. Instead, they are continuous and uncountably infinite. In addition, the state transitions governing the system – that is, the laws of physics – are nonlinear and complex.

We'll keep this motivating example in mind throughout the chapter, although reframing it in terms of the following classic *control problem:* 

### **Example 3.1.1: CartPole**

Consider a pole balanced on a cart. The state s consists of just four continuous values:

- 1. The position of the cart;
- 2. The velocity of the cart;
- 3. The angle of the pole;
- 4. The angular velocity of the pole.

We can *control*  $^{a}$  the cart by applying a horizontal force a.

**Goal:** Stabilize the cart around an ideal state and action  $(s^*, a^*)$ .

Beyond this simple scenario, there are many real-world examples that involve continuous control:

• **Robotics.** Autonomous driving; Controlling a drone's position; Automation in warehouses and manufacturing; Humanoid robots with joints.

<sup>&</sup>lt;sup>a</sup>Controls are the continuous analogue to actions in the discrete setting. In control theory, the state and controls are typically denoted as x and u, but we'll stick with the s and a notation to highlight the similarity with the discrete case.

- **Temperature.** Controlling the temperature in a room; Keeping greenhouses warm; Understanding weather patterns.
- Games. Sports; MMORPGs (Massively Multiplayer Online Role-Playing Games); Board games.
- Finance. Stock trading; Portfolio management; Risk management.

How can we teach computers to solve these kinds of problems?

In the last chapter, we developed efficient dynamic programming algorithms (value iteration and policy iteration) for calculating  $V^*$  and  $\pi^*$  in the finite setting. In this chapter, we'll derive similar results in the continuous case by imposing some additional structure on the problem.

Note that we're still assuming that the entire environment is *known* – that is, we understand 'how the world works'. We'll get to the unknown case in the next chapter.

# 3.2 Optimal control

Recall that an MDP is defined by its state space S, action space A, state transitions P, reward function r, and discount factor  $\gamma$  or time horizon T. What are the equivalents in the control setting?

- The state and action spaces are *continuous* rather than finite. That is,  $\mathcal{S} = \mathbb{R}^{n_s}$  and  $\mathcal{A} = \mathbb{R}^{n_a}$ , where  $n_s$  and  $n_a$  are the number of coordinates to specify a single state or action respectively.
- We call the state transitions the **dynamics** of the system. In the most general case, these might change across timesteps and also include some stochastic **noise**  $w_t$ . We denote these dynamics as the function  $f_t$ , such that  $s_{t+1} = f_t(s_t, a_t, w_t)$ . Of course, we can simplify to cases where the dynamics are deterministic/noise-free (no  $w_t$  term) or are stationary/time-homogeneous (the same function f across timesteps).
- Instead of a reward function, it's more intuitive to consider a **cost function**  $c_t : \mathcal{S} \times \mathcal{A} \to \mathbb{R}$  that describes *how far away* we are from our **goal state-action pair**  $(s^*, a^*)$ . An important special case is when the cost is time-homogeneous; that is, it remains the same function c at each timestep.
- We seek to minimize the *undiscounted* cost with a *time horizon* T. Note that we end an episode at  $s_T$  there is no  $a_T$ , and so we denote the cost for the final state as  $c_T(s_T)$ .

With all of these components, we can now formulate the **optimal control problem:** find a time-dependent policy to minimize the expected undiscounted cost over T timesteps.

### **Definition 3.2.1: Optimal control problem**

$$\begin{aligned} \min_{\pi_0,\dots,\pi_{T-1}:\mathcal{S}\to\mathcal{A}} \quad & \mathbb{E} \left[ \left( \sum_{t=0}^{T-1} c_t(s_t, a_t) \right) + c_T(s_T) \right] \\ \text{where} \quad & s_{t+1} = f_t(s_t, a_t, w_t), \\ & a_t = \pi_t(s_t) \\ & s_0 \sim \mu_0 \\ & w_t \sim \text{noise} \end{aligned} \tag{3.1}$$

#### 3.2.1 Discretization

How does this relate to the finite horizon case? If  $s_t$  and  $a_t$  were discrete, then we'd be able to work backwards using the DP algorithms we saw before. As a matter of fact, let's consider what happens if we discretize the problem. For intuition, suppose  $n_s = n_a = 1$  (that is, states and actions are real numbers). To make  $\mathcal S$  and  $\mathcal A$  discrete, let's choose some small positive  $\epsilon$ , and simply round states and actions to the nearest multiple of  $\epsilon$ . For example, if  $\epsilon = 0.01$ , then we're just rounding s and s to two decimal spaces. If both these state and action spaces can be bounded, then the resulting sets are actually finite, so now we can use our previous tools for MDPs.

But is this actually a feasible solution? Even if our  $\mathcal S$  and  $\mathcal A$  are finite, the existing algorithms might take unfeasibly long to complete. Suppose our state and action spaces are bounded by some constants  $\max_{s\in\mathcal S}\|s\|\leq B_s$  and  $\max_{a\in\mathcal A}\|a\|\leq B_a$ . Then using our rounding method, we must divide each dimension into intervals of length  $\varepsilon$ , resulting in  $(B_s/\varepsilon)^{n_s}$  and  $(B_a/\varepsilon)^{n_a}$  total points. To get a sense of how quickly this grows, let's consider  $\varepsilon=0.01, n_s=n_a=10$ . Then the number of elements in our transition matrix is  $|\mathcal S|^2|\mathcal A|=(100^{10})^2(100^{10})=10^{60}!$  Try finding a computer that'll fit that in memory! (For reference, 32 GB of memory can store  $10^9$  32-bit floating point numbers.)

So as we've seen, discretizing the problem isn't a feasible solution as soon as our action and state spaces are even moderately high-dimensional. How can we do better?

Note that by discretizing the state and action spaces, we implicitly assumed that rounding each state or action vector by some tiny amount  $\varepsilon$  wouldn't change the behavior much; namely, that the functions involved were relatively *continuous*. Can we use this continuous structure in other ways? This brings us to the topic of **Linear Quadratic Regulators**, a widely used and studied tool in control theory.

<sup>&</sup>lt;sup>1</sup>Formally, we can consider an  $\epsilon$ -net over the original continuous space. Let V be some normed space. A subset  $V_{\epsilon} \subseteq V$  is called an  $\epsilon$ -net if for all  $v \in V$ , there exists a  $v_{\epsilon} \in V_{\epsilon}$  such that  $\|v - v_{\epsilon}\| \le \epsilon$ . The rounding example given is technically a 0.005-net.

# 3.3 The Linear Quadratic Regulator Problem

The optimal control problem stated above seems very difficult to solve. The cost function might not be convex, making optimization difficult, and the state transitions might be very complex, making it difficult to satisfy the constraints. Is there a relevant simplification that we can analyze?

We'll show that a natural structure to impose is *linear dynamics* and a *quadratic cost function* (in both arguments). This is called the **linear quadratic regulator** (LQR) model, and is a popular tool in control theory. In fact, some people even design systems to be linear in order to use results from LQR!

Why are these assumptions useful? As we'll see later in the chapter, it lets us *locally approximate* nonlinear dynamics and cost functions using their *Taylor approximations* (up to first and second order respectively). We'll also find that even for more complex setups, we can generalize the algorithms for LQR to get surprisingly good solutions.

### Definition 3.3.1: The linear quadratic regulator

Linear, time-homogeneous dynamics:

$$s_{t+1} = f(s_t, a_t, w_t) = As_t + Ba_t + w_t$$

Quadratic, time-homogeneous cost function: <sup>a</sup>

$$c(s_t, a_t) = \begin{cases} s_t^\top Q s_t + a_t^\top R a_t & t < T \\ s_T^\top Q s_T & t = T \end{cases}$$

We want c to be a convex function (easy to optimize) in both  $s_t$  and  $a_t$ , so we'll set Q and R to both be symmetric.  $^b$ 

Intuitively, the cost function punishes states and actions that are far away from the origin (i.e. both the state and action are zero vectors). More generally, we'll want to replace the origin with a *goal* state and action  $(s^*, a^*)$ . This can easily be done by replacing  $s_t$  with  $(s_t - s^*)$  and  $a_t$  with  $(a_t - a^*)$  in the expression above.

#### **Isotropic Gaussian noise:**

$$w_t \sim \mathcal{N}(0, \sigma^2 I)$$

Putting everything together, the optimization problem we want to solve is:

$$\min_{\substack{\pi_0, \dots, \pi_{T-1}: \mathcal{S} \to \mathcal{A}}} \quad \mathbb{E}\left[\left(\sum_{t=0}^{T-1} s_t^\top Q s_t + a_t^\top R a_t\right) + s_T^\top Q s_T\right]$$
where 
$$s_{t+1} = A s_t + B a_t + w_t$$

$$a_t = \pi_t(s_t)$$

$$w_t \sim \mathcal{N}(0, \sigma^2 I)$$

$$s_0 \sim \mu_0.$$

<sup>a</sup>For some intuition into this expression, consider the simple case where  $a_t$  and  $s_t$  are scalars (and so are Q and R), so  $c(s_t, a_t) = Qs_t^2 + Ra_t^2$ . If this notation is unfamiliar to you, we recommend this tutorial on quadratic forms from Khan Academy!

<sup>b</sup>Note that it suffices for them to be positive definite, but symmetry makes some later expressions much nicer.

So how do we go about analyzing this system? A good first step might be to introduce *value functions*, analogous to those in the previous chapter, to reason about the behavior of the system over the time horizon.

### **Definition 3.3.2: Value functions for LQR**

Given a policy  $\pi = (\pi_0, \dots, \pi_{t-1})$ , we can define the value function  $V_t^{\pi} : \mathcal{S} \to \mathbb{R}$  as

$$V_t^{\pi}(s) = \mathbb{E}\left[\left(\sum_{i=t}^{T-1} c(s_i, a_i)\right) + c(s_T)\right]$$

$$= \mathbb{E}\left[\left(\sum_{i=t}^{T-1} s_i^{\top} Q s_i + a_i^{\top} R a_i\right) + s_T^{\top} Q s_T\right]$$
conditional on  $s_t = s$ 

$$a_i = \pi_i(s_i) \quad \forall i > t.$$

The expression inside the expectation is called the  $\mathbf{cost}$ - $\mathbf{to}$ - $\mathbf{go}$ , since it's just the total cost starting from timestep t.

The Q function additionally conditions on the first action we take:

$$Q_t^{\pi}(s, a) = \mathbb{E}\left[\left(\sum_{i=t}^{T-1} c(s_i, a_i)\right) + c(s_T)\right]$$

$$= \mathbb{E}\left[\left(\sum_{i=t}^{T-1} s_i^{\top} Q s_i + a_i^{\top} R a_i\right) + s_T^{\top} Q s_T\right]$$
conditional on  $(s_t, a_t) = (s, a)$ 

$$a_i = \pi_i(s_i) \quad \forall i > t$$

As in the previous chapter, these will be instrumental in constructing optimal policy  $\pi$  via dynamic programming.

# 3.4 Optimality and the Ricatti Equation

In this section, we'll derive the optimal policy in the LQR setting. We'll do this through induction, which directly translates into a recursive dynamic programming algorithm for actually calculating the optimal policy. Along the way, we'll prove that the optimal value function is *quadratic*, and

that the optimal policy is a linear function of the state.

### Definition 3.4.1: Optimal value functions for LQR

The **optimal value function** is the one that, at any time, in any state, achieves *minimum* cost across all policies:

$$V_t^{\star}(s) = \min_{\pi_t, \dots, \pi_{T-1}: \mathcal{S} \to \mathcal{A}} V_t^{\pi}(s)$$

$$= \min_{\pi_t, \dots, \pi_{T-1}} \mathbb{E} \left[ \left( \sum_{i=t}^{T-1} s_t^{\top} Q s_t + a_t^{\top} R a_t \right) + s_T^{\top} Q s_T \right]$$
conditional on  $a_i = \pi_i(s_i) \quad \forall i \ge t$ 

$$s_t = s$$

### Theorem 3.4.1: Optimal value function in LQR is quadratic and convex

$$V_t^{\star}(s) = s^{\top} P_t s + p_t$$

for some time-dependent  $P_t \in \mathbb{R}^{n_s \times n_s}$  and  $p_t \in \mathbb{R}^{n_s}$  where  $P_t$  is symmetric. Note that there is no linear term.

### Theorem 3.4.2: Optimal policy in LQR is linear

$$\pi_t^{\star}(s) = -K_t s$$

for some  $K_t \in \mathbb{R}^{k \times d}$ . (The negative is due to convention.)

We'll derive both of these theorems simultaneously via an *inductive* proof. We'll start from the final timestep T as our base case. Then, we'll show that if the theorems hold at time t+1, they must hold for time t. This is called the *inductive hypothesis*, and by proving it, the theorems will 'ripple down' to all earlier timesteps, and we'll have shown that these theorems are always true. As an additional bonus, as mentioned above, our proof will naturally produce a DP algorithm that allows us to calculate the optimal value function and policy.

Base case:  $V_T^{\star}(s) = s^{\top} P_T s + p_T$ . At the final timestep, there are no possible actions to take, and so  $V_T^{\star}(s) = c(s) = s^{\top} Q s$ . Thus  $P_T = Q$  and  $p_T$  is the zero vector.

**Inductive hypothesis:** We seek to show that the inductive step holds for both theorems: If  $V_{t+1}^{\star}(s)$  is quadratic and convex, then  $V_t^{\star}(s)$  must also be quadratic and convex, and  $\pi_t^{\star}(s)$  must be linear. We'll break this down into the following steps:

- **Step 1.** Show that  $Q_t^{\star}(s, a)$  is quadratic and convex (in both s and a).
- **Step 2.** Derive the optimal policy  $\pi_t^\star(s) = \arg\min_a Q_t^\star(s,a)$  and show that it's linear.
- **Step 3.** Show that  $V_t^{\star}(s)$  is quadratic and convex.

This is essentially the same proof that we wrote in the finite-horizon MDP setting, except

now the state and action are continuous instead of finite.

We first assume our theorems are true at time t+1. That is,

$$V_{t+1}^{\star}(s) = s^{\top} P_{t+1} s + p_{t+1}$$
 for all states  $s \in \mathcal{S}$ .

**Step 1.** We'll start off by demonstrating that  $Q_t^{\star}(s)$  is quadratic and convex. Recall that the definition of  $Q_t^{\star}: \mathcal{S} \times \mathcal{A} \to \mathbb{R}$  is

$$Q_t^{\star}(s, a) = c(s, a) + \underset{s' \sim f(s, a, w_{t+1})}{\mathbb{E}} V_{t+1}^{\star}(s').$$

We know  $c(s,a) := s^\top Q s + a^\top R a$ . Let's consider the average value over the next timestep. The only randomness in the dynamics comes from the noise  $w_{t+1}$ , so we can write out this expected value as:

$$\begin{split} & \underset{s' \sim f(s, a, w_{t+1})}{\mathbb{E}} V_{t+1}^{\star}(s') \\ &= \underset{w_{t+1} \sim \mathcal{N}(0, \sigma^2 I)}{\mathbb{E}} V_{t+1}^{\star}(As + Ba + w_{t+1}) & \text{definition of } f \\ &= \underset{w_{t+1}}{\mathbb{E}} [(As + Ba + w_{t+1})^{\top} P_{t+1}(As + Ba + w_{t+1}) + p_{t+1}]. & \text{inductive hypothesis} \end{split}$$

Summing and combining like terms, we get

$$Q_{t}^{\star}(s, a) = s^{\top}Qs + a^{\top}Ra + \underset{w_{t+1}}{\mathbb{E}} [(As + Ba + w_{t+1})^{\top}P_{t+1}(As + Ba + w_{t+1}) + p_{t+1}]$$

$$= s^{\top}(Q + A^{\top}P_{t+1}A)s + a^{\top}(R + B^{\top}P_{t+1}B)a + 2s^{\top}A^{\top}P_{t+1}Ba$$

$$+ \underset{w_{t+1}}{\mathbb{E}} [w_{t+1}^{\top}P_{t+1}w_{t+1}] + p_{t+1}.$$

Note that the terms that are linear in  $w_t$  have mean zero and vanish. Now consider the remaining expectation over the noise. By expanding out the product and using linearity of expectation, we can write this out as

$$\mathbb{E}_{w_{t+1}}[w_{t+1}^{\top} P_{t+1} w_{t+1}] = \sum_{i=1}^{d} \sum_{j=1}^{d} (P_{t+1})_{ij} \mathbb{E}_{w_{t+1}}[(w_{t+1})_i (w_{t+1})_j].$$

When dealing with these *quadratic forms*, it's often helpful to consider the terms on the diagonal (i = j) separately from those off the diagonal. On the diagonal, the expectation becomes

$$(P_{t+1})_{ii} \mathbb{E}(w_{t+1})_i^2 = (P_{t+1})_{ii} \operatorname{Var}((w_{t+1})_i) = \sigma^2(P_{t+1})_{ii}.$$

Off the diagonal, since the elements of  $w_{t+1}$  are independent, the expectation factors, and since each element has mean zero, the term disappears:  $(P_{t+1})_{ij} \mathbb{E}(w_{t+1})_i \mathbb{E}(w_{t+1})_j = 0$ . Thus, the only terms left are the ones on the diagonal, so the sum of these can be expressed as the trace of  $\sigma^2 P_{t+1}$ :

$$\mathbb{E}_{w_{t+1}} w_{t+1}^{\top} P_{t+1} w_{t+1} = \text{Tr}(\sigma^2 P_{t+1}).$$

Substituting this back into the expression for  $Q_t^{\star}$ , we have:

$$Q_t^{\star}(s,a) = s^{\top}(Q + A^{\top}P_{t+1}A)s + a^{\top}(R + B^{\top}P_{t+1}B)a + 2s^{\top}A^{\top}P_{t+1}Ba + \operatorname{Tr}(\sigma^2P_{t+1}) + p_{t+1}.$$
(3.2)

As we hoped, this expression is quadratic in s and a. Furthermore, we'd like to show that it's also convex w.r.t. a in order to make the optimization much easier. This is fairly straightforward:

## Theorem 3.4.3: $Q_t^{\star}$ is convex

Consider the part of Equation 3.2 that is quadratic in a, namely  $a^{\top}(R+B^{\top}P_{t+1}B)a$ . Then  $Q_t^{\star}$  is convex w.r.t. a if  $R+B^{\top}P_{t+1}B$  is positive definite (PD).

It suffices to show that it is symmetric, which we do as follows. Recall that in our definition of LQR, we assumed that R is symmetric (see Definition 3.3.1). Also note that since  $P_{t+1}$  is symmetric (by the inductive hypothesis), so too must be  $B^\top P_{t+1} B$ . (If this isn't clear, try proving it as an exercise!) Since the sum of two symmetric matrices is also symmetric, we have that  $R + B^\top P_{t+1} B$  is symmetric, and so  $Q_t^\star$  is convex w.r.t. a. A similar proof shows that  $Q + A^\top P_{t+1} A$  is symmetric, and so  $Q_t^\star$  is also convex in s.

**Step 2.** Now let's move on to the next part of the next part of proving the inductive hypothesis: showing that  $\pi_t^{\star}(s) = \arg\min_a Q_t^{\star}(s,a)$  is linear. Since  $Q_t^{\star}$  is convex, finding its minimum over a is easy: we can just take the gradient w.r.t. a and set it to zero. First, we calculate the gradient:

$$\nabla_a Q_t^{\star}(s, a) = \nabla_a [a^{\top} (R + B^{\top} P_{t+1} B) a + 2s^{\top} A^{\top} P_{t+1} B a]$$
  
=  $2(R + B^{\top} P_{t+1} B) a + (2s^{\top} A^{\top} P_{t+1} B)^{\top}$ 

Setting this to zero, we get

$$0 = (R + B^{\top} P_{t+1} B) a + B^{\top} P_{t+1} A s$$

$$\pi_t^{\star}(s) := a = -(R + B^{\top} P_{t+1} B)^{-1} B^{\top} P_{t+1} A s$$

$$= -K_t s,$$
(3.3)

where  $K_t = (R + B^{\top} P_{t+1} B)^{-1} B^{\top} P_{t+1} A$ .

Note that this optimal policy has an interesting property: in addition to being independent of the starting distribution  $\mu_0$  (which also happened for our finite-horizon MDP solution), it's also fully deterministic and isn't affected by noise! (Compare this with the discrete MDP case, where calculating our optimal policy required taking an expectation over the state transitions.)

**Step 3.** To complete our inductive proof, we must show that the inductive hypothesis is true at time t; that is, we must prove that  $V_t^{\star}(s)$  is quadratic. Using the identity  $V_t^{\star}(s) = Q_t^{\star}(s, \pi^{\star}(s))$ , we have:

$$V_t^{\star}(s) = Q_t^{\star}(s, \pi^{\star}(s))$$

$$= s^{\top} (Q + A^{\top} P_{t+1} A) s + (-K_t s)^{\top} (R + B^{\top} P_{t+1} B) (-K_t s) + 2s^{\top} A^{\top} P_{t+1} B (-K_t s)$$

$$+ \operatorname{Tr}(\sigma^2 P_{t+1}) + p_{t+1}$$

Note that w.r.t. s, this is the sum of a quadratic term and a constant, which is exactly what we were aiming for!

To conclude our proof, let's concretely specify the values of  $P_t$  and  $p_t$ . The constant term is clearly  $p_t = \text{Tr}(\sigma^2 P_{t+1}) + p_{t+1}$ . We can simplify the quadratic term by substituting in  $K_t$ . Notice that when we do this, the  $(R + B^\top P_{t+1} B)$  term in the expression is cancelled out by its inverse, and the remaining terms combine to give what is known as the *Riccati equation*:

### Theorem 3.4.4: Riccati equation

$$P_{t} = Q + A^{\top} P_{t+1} A - A^{\top} P_{t+1} B (R + B^{\top} P_{t+1} B)^{-1} B^{\top} P_{t+1} A.$$

There are several nice things to note about the Riccati equation:

- 1. It's defined **recursively.** Given  $P_T$ , the dynamics defined by A and B, and the state coefficients Q, we can recursively calculate  $P_t$  across all timesteps.
- 2.  $P_t$  often appears in calculations surrounding optimality, such as  $V_t^{\star}, Q_t^{\star}$ , and  $\pi_t^{\star}$ .
- 3. Together with A, B, and the action coefficients R, it fully defines the optimal policy.

Now we've shown that  $V_t^{\star}(s) = s^{\top} P_t s + p_t$ , which is quadratic and convex, and this concludes our proof.  $\blacksquare$ 

In summary, we just demonstrated that:

- The optimal value function  $V_t^*$  is convex at all t.
- $\bullet$  The optimal  $Q\text{-function }Q_t^\star$  is convex (in both arguments) at all t.
- The optimal policy  $\pi_t^{\star}$  is linear at all t.
- All of these quantities can be calculated using a symmetric matrix  $P_t$  for each timestep, which can be defined recursively using the Riccati equation.

Before we move on to some extensions of LQR, let's consider how the state at time t behaves when we act according to this optimal policy.

## 3.4.1 Expected state at time t

Suppose you're about to go to bed, and the thermostat in your room is controlled by an optimal policy under LQR. A very reasonable question to ask would be: what's the expected state (i.e. temperature) of the room at a given time t? Certainly you don't want to freeze or boil overnight!

To answer this question, let's first express the state at time t in a cleaner way in terms of the

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history. Note that having linear dynamics makes it easy to expand terms backwards in time:

$$s_{t} = As_{t-1} + Ba_{t-1} + w_{t-1}$$

$$= A(As_{t-2} + Ba_{t-2} + w_{t-2}) + Ba_{t-1} + w_{t-1}$$

$$= \cdots$$

$$= A^{t}s_{0} + \sum_{i=0}^{t-1} A^{i}(Ba_{t-i-1} + w_{t-i-1}).$$

Let's consider the average state at this time, given all the past states and actions. Since we assume that  $\mathbb{E} w_t = 0$  (this is the zero vector in d dimensions), when we take an expectation, the  $w_t$  term vanishes due to linearity, and so we're left with

$$\mathbb{E}[s_t \mid s_{0:(t-1)}, a_{0:(t-1)}] = A^t s_0 + \sum_{i=0}^{t-1} A^i B a_{t-i-1}.$$

If we choose actions according to our optimal policy, this becomes

$$\mathbb{E}[s_t \mid s_0, a_t = -K_t s_t] = \left(\prod_{i=0}^{t-1} (A - BK_i)\right) s_0.$$

This introdces the quantity  $A-BK_i$ , which shows up frequently in control theory. For example, one important question is: will  $s_t$  remain bounded, or will it go to infinity as time goes on? To answer this, let's imagine that these  $K_i$ s are equal (call this matrix K). Then the expression above becomes  $(A-BK)^t s_0$ . Now consider the maximum eigenvalue  $\lambda_{\max}$  of A-BK. If  $|\lambda_{\max}|>1$ , then there's some nonzero initial state  $\bar{s}_0$ , the corresponding eigenvector, for which

$$\lim_{t \to \infty} (A - BK)^t \bar{s}_0 = \lambda_{\max}^t \bar{s}_0 = \infty.$$

By then, your room is *definitely* on fire! Otherwise, if  $|\lambda_{\max}| < 1$ , then it's impossible for your original state to explode as dramatically (assuming it's properly normalized).

We've now formulated an optimal solution for the typical, time-homogeneous case of LQR, and considered the expected state under the optimal policy. However, this simple case is insufficient for more complex tasks. In the following sections, we'll consider some motivating examples, and extensions of LQR where some assumptions are relaxed.

# 3.5 Extensions

In this section, we'll consider settings where some of the assumptions we made above are relaxed. Specifically, we'll consider:

- 1. **Time-dependency,** where the dynamics and cost function might change depending on the timestep.
- 2. General quadratic cost, where we allow for linear terms and a constant term.
- 3. Tracking a goal trajectory rather than aiming for a single goal state-action pair.

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## 3.5.1 Time-dependent dynamics and cost function

So far, we've considered the *time-homogeneous* case, where the dynamics and cost function stay the same at every timestep. However, this might not always be the case. For example, if we want to preserve the temperature in a greenhouse, the outside forces are going to change depending on the time of day. As another example, in many sports or video games, the rules and scoring system might change during overtime. To address these sorts of problems, we can loosen the time-homogeneous restriction, and consider the case where the dynamics and cost function are *time-dependent*. Our analysis remains almost identical; in fact, we can simply add a time index to the matrices A and B that determine the dynamics and the matrices A and B that determine the dynamics and verify this claim!)

The modified problem is now defined as follows:

$$\arg\min_{\pi_0,\dots,\pi_{T-1}:S\to\mathcal{A}} \quad \mathbb{E}\left[\left(\sum_{t=0}^{T-1}(s_t^\top Q_t s_t) + a_t^\top R_t a_t\right) + s_T^\top Q_T s_T\right]$$
where  $s_{t+1} = f_t(s_t, a_t, w_t) = A_t s_t + B_t a_t + w_t$ 

$$s_0 \sim \mu_0$$

$$a_t = \pi_t(s_t)$$

$$w_t \sim \mathcal{N}(0, \sigma^2 I).$$

The derivation of the optimal value functions and the optimal policy remains almost exactly the same, and we can modify the Riccati equation accordingly:

#### Theorem 3.5.1: Time-dependent Riccati Equation

$$P_t = Q_t + A_t^{\top} P_{t+1} A_t - A_t^{\top} P_{t+1} B_t (R_t + B_t^{\top} P_{t+1} B_t)^{-1} B_t^{\top} P_{t+1} A_t.$$

Note that this is just the time-homogeneous Riccati equation (Theorem 3.4.4), but with the time index added to each of the relevant matrices.

Additionally, by allowing the dynamics to vary across time, we gain the ability to *locally approximate* nonlinear dynamics at each timestep. We'll discuss this later in the chapter.

# 3.5.2 More general quadratic cost functions

Our original cost function had only second-order terms w.r.t. the state and action. We can also consider more general quadratic cost functions that also have first-order terms and a constant term. Combining this with time-dependent dynamics results in the following expression, where we introduce a new matrix  $M_t$  for the cross term, linear coefficients  $q_t$  and  $r_t$  for the state and action respectively, and a constant term  $c_t$ :

$$c_t(s_t, a_t) = (s_t^{\top} Q_t s_t + s_t^{\top} M_t a_t + a_t^{\top} R_t a_t) + (s_t^{\top} q_t + a_t^{\top} r_t) + c_t.$$
(3.4)

Similarly, we can also include a constant term  $v_t \in \mathbb{R}^{n_s}$  in the dynamics (note that this is *fixed* at each timestep, unlike the noise  $w_t$ ):

$$s_{t+1} = f_t(s_t, a_t, w_t) = A_t s_t + B_t a_t + v_t + w_t.$$

The derivation of the optimal solution in this case will be left as a homework exercise.

### 3.5.3 Tracking a predefined trajectory

So far, we've been trying to get the robot to stay as close as possible to the origin, or more generally a goal state-action pair  $(s^\star, a^\star)$ . However, consider applying LQR to autonomous driving. Now, we want the desired state to change over time, instead of remaining in one location. Otherwise, it wouldn't be a very useful vehicle! In these cases, we want the robot to follow a predefined *trajectory* of states and actions  $(s_t^\star, a_t^\star)_{t=0}^{T-1}$ . To do this, we'll modify the cost function accordingly:

$$c_t(s_t, a_t) = (s_t - s_t^{\star})^{\top} Q(s_t - s_t^{\star}) + (a_t - a_t^{\star})^{\top} R(a_t - a_t^{\star}).$$

Note that this punishes states and actions that are far from the intended trajectory. By expanding out these multiplications, we can see that this is actually a special case of the more general quadratic cost function we discussed above (Equation 3.4):

$$M_t = 0, q_t = -2Qs_t^*, r_t = -2Ra_t^*, c_t = (s_t^*)^\top Q(s_t^*) + (a_t^*)^\top R(a_t^*).$$

## 3.6 The infinite-horizon setting

Another assumption we've made is that the task has a *finite horizon* T. How about tasks that might matter indefinitely, where we want to minimize the expected cost over *all future timesteps?* Consider, for example, controlling the long-term value of a portfolio, or managing acidity levels in a lake.

In the previous chapter, we dealt with such **infinite-horizon** cases by *discounting* future rewards. This time, we'll take a different approach by considering the limit of a *finite*-horizon task as  $T \to \infty$ . As it turns out, our derivation is exactly analogous to value iteration from the previous chapter! However, here the structure is nice enough that we don't need a discount factor  $\gamma$  to deal with limits analytically. (Note that we must normalize by 1/T to keep the total cost bounded.)

In the discounted case, analogously to taking the limit as  $T\to\infty$ , we consider the limit as the discount factor  $\gamma$  approaches 1. This is because as  $\gamma\to 1$ , time discounting becomes less and less important – just like the horizon T vanishing into the distance – and we're left with the undiscounted case. (Note that for the total reward to remain bounded, we must normalize the sum by  $1-\gamma$ .) Let's consider value iteration in this setting, which uses the Bellman operator  $\mathcal J$  to update V:

$$V_{t+1}(s) = \lim_{\gamma \to 1} (\mathcal{J}V_t)(s) = \lim_{\gamma \to 1} \max_{a} \left( r(s, a) + \gamma \mathop{\mathbb{E}}_{s' \sim P(s, a)} V_t(s') \right).$$

This is exactly analogous to the *Riccati equation* (3.4.4)! Instead of thinking of  $P_{t+1}$  as defining the value function for the *next timestep*, though, we think of it as the *next version* of the value function in an iterative algorithm. Then both of these algorithms are doing the same thing: iteratively refining the value function by acting greedily w.r.t. the current iteration.

By going through the same derivation in section 3.4, we'll see that  $P_t$  (defined recursively by the Riccati equation) converges to a fixed value P. For an intuitive perspective on why this happens, let's suppose you have an upcoming project deadline. When it's still a few months away, you might not pay much attention to it, and behave as you normally do. But as the deadline gets closer and closer, suddenly the horizon becomes more and more relevant, and you'll spend more time thinking about it. The infinite-horizon case is just where the deadline is infinitely far away, so you can just behave "like normal" all the time!

	Control	Finite MDP
State and action spaces	Continuous	Finite
Optimization problem	Minimize finite-horizon undiscounted cost	Maximize infinite-horizon discounted reward <i>or</i> finite-horizon discounted reward
Approaching the infinite- horizon, undiscounted setting	$\lim_{T \to \infty} \frac{1}{T} \mathbb{E} \left( \sum_{t=0}^{T-1} c(s_t, a_t) \right)$	$\lim_{\gamma \to 1} (1 - \gamma) \mathbb{E} \sum_{t=0}^{\infty} \gamma^t r(s_t, a_t)$
Iterative algorithm for optimal value	Riccati equations $P \leftarrow Q + A^\top PA - A^\top PB(R + B^\top PB)^{-1}B^\top PA$	Value iteration $V(s) \leftarrow \max_{a} \left[ r(s, a) + \gamma \mathbb{E}_{s' \sim P(s, a)} V(s') \right]$

Figure 3.1: A comparison between the continuous control and finite MDP settings.

## 3.7 Approximating nonlinear dynamics

As its name might suggest, LQR works best when the dynamics are linear and the cost function is quadratic. In these settings, we've shown a way to analytically derive the optimal policy. However, let's return to the CartPole example from the start of the chapter (Example 3.1.1). The dynamics (physics) aren't linear, and we might also want to specify a cost function that's more complex than just quadratic.

Concretely, let's consider a *noise-free* problem since, as we saw, the noise doesn't affect the optimal policy. Let's assume the dynamics and cost function are stationary, and ignore the terminal state for simplicity:

#### **Definition 3.7.1: Nonlinear control problem**

$$\min_{\pi_0, \dots, \pi_{T-1}: \mathcal{S} \to \mathcal{A}} \quad \mathbb{E} \left[ \sum_{t=0}^{T-1} c(s_t, a_t) \right]$$
 where  $s_{t+1} = f(s_t, a_t)$  
$$a_t = \pi_t(s_t)$$
 
$$s_0 \sim \mu_0$$
 
$$c(s, a) = d(s, s^\star) + d(a, a^\star).$$

Here, d denotes some general measure of distance to the goal state and action  $(s^*, a^*)$ .

This is now only slightly simplified from the general optimal control problem (see 3.2.1). Here, we don't know an analytical form for the dynamics f or the cost function c, but we assume that we're able to query/sample/simulate them to get their values at a given state and action. How can we adapt LQR to this more general nonlinear case?

### 3.7.1 Local linearization

As we briefly mentioned in the introduction, part of the reason we designed the LQR problem the way we did was because we can take any *locally continuous* function, and approximate it using a Taylor expansion of low-order polynomials. By taking a linear approximation of f and a quadratic approximation of f, we're back to the regime of LQR with these derived matrices in terms of their gradients and Hessians accordingly.

**Assumptions.** This approach assumes that f is differentiable and that c is twice-differentiable, both around  $(s^\star, a^\star)$ . Additionally, since a Taylor expansion generally gets less and less accurate the further you stray from the point of expansion, this means that we also assume that all states are close to the optimal state  $s^\star$ , and that we can stay close using actions that are close to  $a^\star$ . If this seems like a strong set of restrictions, it is! But we'll save this discussion for the next section.

If you're unfamiliar with Taylor expansions, we recommend taking a calculus course; we'll use them here without much further introduction. Linearizing the dynamics around  $(s^*, a^*)$  gives:

$$f(s,a) \approx f(s^*, a^*) + \nabla_s f(s^*, a^*)(s - s^*) + \nabla_a f(s^*, a^*)(a - a^*)$$
$$(\nabla_s f(s,a))_{ij} = \frac{df_i(s,a)}{ds_j}, \quad i, j \le n_s \qquad (\nabla_a f(s,a))_{ij} = \frac{df_i(s,a)}{da_j}, \quad i \le n_s, j \le n_a$$

and quadratizing the cost function around  $(s^*, a^*)$  gives:

$$\begin{split} c(s,a) &\approx c(s^\star,a^\star) & \text{constant} \\ &+ \nabla_s c(s^\star,a^\star)(s-s^\star) + \nabla_a c(s^\star,a^\star)(a-a^\star) & \text{linear} \\ &+ \frac{1}{2}(s-s^\star)^\top \nabla_{ss} c(s^\star,a^\star)(s-s^\star) \\ &+ \frac{1}{2}(a-a^\star)^\top \nabla_{aa} c(s^\star,a^\star)(a-a^\star) & \text{quadratic} \\ &+ (s-s^\star)^\top \nabla_{sa} c(s^\star,a^\star)(a-a^\star) \end{split}$$

where the gradients and Hessians are defined as

$$(\nabla_s c(s,a))_i = \frac{dc(s,a)}{ds_i}, \quad i \le n_s$$

$$(\nabla_a c(s,a))_i = \frac{dc(s,a)}{da_i}, \quad i \le n_a$$

$$(\nabla_s c(s,a))_{ij} = \frac{d^2 c(s,a)}{ds_i ds_j}, \quad i,j \le n_s$$

$$(\nabla_{aa} c(s,a))_{ij} = \frac{d^2 c(s,a)}{da_i da_j}, \quad i,j \le n_a$$

$$(\nabla_{aa} c(s,a))_{ij} = \frac{d^2 c(s,a)}{da_i da_j}, \quad i,j \le n_a$$

We note that this cost can be expressed in the general quadratic form seen in Equation 3.4. We leave it as an exercise to derive the corresponding matrices and vectors Q, R, M, q, r, c. To calculate these gradients and Hessians in practice, we use a method known as **finite differencing** for numerically computing derivatives. Namely, we can simply use the definition of derivative, and see how the function changes as we add or subtract a tiny  $\delta$  to the input.

However, simply taking the second-order approximation of the cost function is insufficient; we also need to ensure that it is locally convex, that is, the Hessians  $\nabla_{ss}c(s^{\star},a^{\star})$  and  $\nabla_{aa}c(s^{\star},a^{\star})$  are positive definite.

**Local convexification.** Recall that an equivalent definition of positive definite matrices is that all of their eigenvalues are positive. One way to naively force  $\nabla_{ss}c(s^*,a^*)$  and  $\nabla_{aa}c(s^*,a^*)$  to be positive definite is to simply remove any negative eigenvalues. Note that both of these Hessians are symmetric, and so they can be decomposed in terms of their eigenbasis. Thus by removing their eigenvalues and adding a small 'lower bound' to the eigenvalues (so that the surface will have some minimum amount of curvature), we obtain

$$\nabla_{ss} c(s^*, a^*) = \sum_{i=1}^{n_s} \sigma_i u_i u_i^{\top}$$

$$\nabla_{ss} \widetilde{c}(s^*, a^*) = \sum_{\substack{i=1 \ \sigma_i > 0}}^{n_s} \sigma_i u_i u_i^{\top} + \lambda I.$$

We can use a similar approach to convexify  $\nabla_{aa}\widetilde{c}(s^\star,a^\star)$ . Now that we have a convex quadratic approximation to the cost function, and a linear approximation to the state transitions, we can simply apply the time-homogenous LQR methods we derived before.

But what happens when our assumptions break down, namely when we enter states far away from  $s^*$  and want to use actions far from  $a^*$ ? As we mentioned above, our Taylor approximation will typically become less accurate. To address this, we'll need to do a Taylor approximation around different points at each time step.

### 3.7.2 Iterative LQR

**Iterative LQR** is a way to resolve the issues with local linearization. The key idea is to linearize around different points at each timestep, making creating a time-dependent approximation of the dynamics. We'll break it into a few steps:

- **Step 1.** Form an LQR around the current candidate trajectory  $(\bar{s}_t^i, \bar{a}_t^i)_{t=0}^{T-1}$  using local approximation.
- **Step 2.** Apply the solution to time-dependent LQR from subsection 3.5.1 to obtain an optimal policy  $\pi^i$ .
- **Step 3.** Generate a new trajectory  $(\widetilde{a}_t)_{t=0}^{T-1}$  using  $\pi^i$ .
- **Step 4.** Compute a better candidate trajectory  $\mathbf{a}^{i+1}$  by interpolating between  $\mathbf{a}^i$  and  $\widetilde{\mathbf{a}}$ .

Now the question becomes: How do we choose the best *waypoints*  $(\bar{s}_t, \bar{a}_t)$  to get the best approximation?

We'll once again use an iterative approach, similarly to value iteration, where at each step we update the waypoints *greedily* w.r.t. the current iteration. Let's use a superscript to denote the iteration of the algorithm. We can start off by initializing some (bad) sequence of waypoints  $\bar{a}_0^0,\ldots,\bar{a}_{T-1}^0$  by some approximate method such as local linearization. Applying these actions at their respective timesteps gives a sample trajectory

$$\bar{s}_0, \bar{a}_0, \bar{s}_1, \bar{a}_1, \dots, \bar{s}_{T-1}, \bar{a}_{T-1}$$
 where  $\bar{s}_0^0 = \bar{s}_0 = \underset{s_0 \sim \mu_0}{\mathbb{E}}[s_0], \quad \bar{s}_{t+1}^0 = f(\bar{s}_t^0, \bar{a}_t^0).$ 

Now, at each timestep t, we linearize f and quadratize (and convexify) c around the point generated in the sample trajectory, using the same Taylor expansion techniques we saw in the previous section:

$$\begin{split} f_t(s,a) &\approx f(\bar{s}_t^i, \bar{a}_t^i) + \nabla_s f(\bar{s}_t^i, \bar{a}_t^i)(s - \bar{s}_t^i) + \nabla_a f(\bar{s}_t^i, \bar{a}_t^i)(a - \bar{a}_t^i) \\ c_t(s,a) &\approx c(\bar{s}_t^i, \bar{a}_t^i) + \left[s - \bar{s}_t^i \quad a - \bar{a}_t^i\right] \begin{bmatrix} \nabla_s c(\bar{s}_t^i, \bar{a}_t^i) \\ \nabla_a c(\bar{s}_t^i, \bar{a}_t^i) \end{bmatrix} \\ &+ \frac{1}{2} \left[s - \bar{s}_t^i \quad a - \bar{a}_t^i\right] \begin{bmatrix} \nabla_{ss} c(\bar{s}_t^i, \bar{a}_t^i) & \nabla_{sa} c(\bar{s}_t^i, \bar{a}_t^i) \\ \nabla_{as} c(\bar{s}_t^i, \bar{a}_t^i) & \nabla_{aa} c(\bar{s}_t^i, \bar{a}_t^i) \end{bmatrix} \begin{bmatrix} s - \bar{s}_t^i \\ a - \bar{a}_t^i \end{bmatrix} \end{split}$$

Now let's use the time-dependent LQR solution to compute an optimal policy  $\pi_0^i, \ldots, \pi_{T-1}^i$ . We can then generate a new sample trajectory by taking actions according to this optimal policy:

$$\bar{s}_0^{i+1} = \bar{s}_0, \qquad \tilde{a}_t = \pi_t^i(\bar{s}_t^{i+1}), \qquad \bar{s}_{t+1}^{i+1} = f(\bar{s}_t^{i+1}, \tilde{a}_t).$$

Note that the states are drawn by sampling from the *true* dynamics, and also that we've denoted these actions as  $\tilde{a}_t$  and aren't directly using them for the next iteration  $a_t^{i+1}$ . Rather, we want

to interpolate between them and the actions from the previous iteration  $\bar{a}_0^i, \ldots, \bar{a}_{T-1}^i$ . This is so that the cost will *increase monotonically*, since if the new policy turns out to actually be worse, we can stay closer to the previous trajectory. (Can you think of an intuitive example where this might happen?) Formally, we want to find  $\alpha \in [0,1]$  to generate the next iteration of actions:

$$\begin{aligned} \min_{\alpha \in [0,1]} \quad & \sum_{t=0}^{T-1} c(s_t, \bar{a}_t^{i+1}) \\ \text{where} \quad & s_{t+1} = f(s_t, \bar{a}_t^{i+1}) \\ & \bar{a}_t^{i+1} = \alpha \bar{a}_t^i + (1-\alpha) \bar{a}_t \\ & s_0 = \bar{s}_0. \end{aligned}$$

Note that this is only optimizing over the closed interval [0,1], so by the Extreme Value Theorem it's guaranteed to have a global maximum.

The final output of this algorithm is a policy  $\pi^{n_{\text{steps}}}$  derived after  $n_{\text{steps}}$  of the algorithm. Though the proof is somewhat complex, one can show that for many nonlinear control problems, this solution converges to a locally optimal solution (in the policy space).

## 3.8 Programming and Implementation

Not sure how much to include here yet. WIP. Walk through a basic Python solution to OpenAl Gym and CartPole? (Or save this for homework?)

## 3.9 Exercises

- 1. Consider a cleaning robot with one wheel on each side of its body. Your pet has made a mess nearby, and you want to steer the robot to go clean it up. Let's represent the state of the robot as a 3-dimensional vector containing its (x,y) coordinates and its angle  $\theta$ , relative to some global reference frame. We can control the robot using its linear velocity v (change in x, y) and angular velocity w (change in  $\theta$ ). For simplicity, we'll assume the robot is perfectly ideal and there's no noise in the system.
  - (a) Formally describe the true of the system as a function  $f: \mathcal{S} \times \mathcal{A} \to \mathcal{S}$ . Assume the system works in timesteps of dt = 1s and that all distances are measured in meters.
  - (b) Linearize the dynamics that you derived using a first-order Taylor approximation. Is this approximation stationary or time-dependent?
  - (c) Suppose the mess is at location  $(x^\star,y^\star)=(2,2)$ . We want to reach that state using a small amount of energy. Write down the quadratic cost function using  $Q=\mathrm{diag}(1,1,0.5)$ , expressing that we care about the final position more than the state, and R=0.2I to penalize large action steps.

Now that our LQR is set up, let's find the optimal policy.

CHAPTER 3. LQR 3.9. EXERCISES

(a) Let the horizon be T=50 timesteps. Write down the recursive update formula for  $P_t$  (Hint: use the Riccati equation).

(b) Write the closed form solution of the optimal policy at time T-1. Verify that this is a linear function of the state.

2. (Todo.)

# Chapter 4

# **Policy Gradients**

### 4.1 Motivation

The scope of our problem has been gradually expanding.

- 1. In the first chapter, we considered *bandits* with a finite number of arms, where the only stochasticity involved was their rewards.
- 2. In the second chapter, we considered *MDPs* more generally, involving a finite number of states and actions, where the state transitions are Markovian.
- 3. In the third chapter, we considered *continuous* state and action spaces and developed the *Linear Quadratic Regulator*. We then showed how to use it to find *locally optimal solutions* to problems with nonlinear dynamics and non-quadratic cost functions.

Now, we'll continue to investigate the case of finding optimal policies in large MDPs using the self-explanatory approach of *policy optimization*. This is a general term encompassing many specific algorithms we've already seen:

- Policy iteration for finite MDPs,
- *Iterative LQR* for locally optimal policies in continuous control.

Here we'll see some general algorithms that allow us to optimize policies for general kinds of problems. These algorithms have been used in many groundbreaking applications, including AlphaGo, OpenAl Five. These methods also bring us into the domain where we can use *deep learning* to approximate complex, nonlinear functions.

## 4.2 (Stochastic) Policy Gradient Ascent

Let's suppose our policy can be *parameterized* by some parameters  $\theta$ . For example, these might be a preferences over state-action pairs, or in a high-dimensional case, the weights and biases of a deep neural network. We'll talk more about possible parameterizations in section 4.5

Remember that in reinforcement learning, the goal is to *maximize reward*. Specifically, we seek the parameters that maximize the expected total reward, which we can express concisely using the value function we defined earlier:

$$J(\theta) := \underset{s_0 \sim \mu_0}{\mathbb{E}} V^{\pi_{\theta}}(s_0) = \mathbb{E} \sum_{t=0}^{T-1} r_t$$

$$\text{where} \quad s_0 \sim \mu_0$$

$$s_{t+1} \sim P(s_t, a_t),$$

$$a_h = \pi_{\theta}(s_h)$$

$$r_h = r(s_h, a_h).$$

$$(4.1)$$

We call a sequence of states, actions, and rewards a **trajectory**  $\tau = (s_i, a_i, r_i)_{i=0}^{T-1}$ , and the total time-discounted reward is also often called the **return**  $R(\tau)$  of a trajectory. Note that the above is the *undiscounted*, *finite-horizon case*, which we'll continue to use throughout the chapter, but analogous results hold for the *discounted*, *infinite-horizon case*.

Note that when the state transitions are Markov (i.e.  $s_t$  only depends on  $s_{t-1}, a_{t-1}$ ) and the policy is stationary (i.e.  $a_t \sim \pi_{\theta}(s_t)$ ), we can write out the *likelihood of a trajectory* under the policy  $\pi_{\theta}$ :

$$\rho_{\theta}(\tau) = \mu(s_0)\pi_{\theta}(a_0|s_0) 
\times P(s_1|s_0, a_0)\pi_{\theta}(a_1|s_1) 
\times \cdots 
\times P(s_{H-1}|s_{H-2}, a_{H-2})\pi_{\theta}(a_{H-1}|s_{H-1}).$$
(4.2)

This lets us rewrite  $J(\theta) = \mathbb{E}_{\tau \sim \rho_{\theta}} R(\tau)$ .

Now how do we optimize for this function? One very general optimization technique is gradient ascent. Namely, the **gradient** of a function at a given point answers: At this point, which direction should we move to increase the function the most? By repeatedly moving in this direction, we can keep moving up on the graph of this function. Expressing this iteratively, we have:

$$\theta_{t+1} = \theta_t + \eta \nabla_{\theta} J(\pi_{\theta}) \Big|_{\theta = \theta_t},$$

Where  $\eta$  is a *hyperparameter* that says how big of a step to take each time.

In order to apply this technique, we need to be able to evaluate the gradient  $\nabla_{\theta}J(\pi_{\theta})$ . How can we do this?

In practice, it's often impractical to evaluate the gradient directly. For example, in supervised learning,  $J(\theta)$  might be the sum of squared prediction errors across an entire **training dataset**. However, if our dataset is very large, we might not be able to fit it into our computer's memory!

Instead, we can *estimate* a gradient step using some estimator  $\widetilde{\nabla} J(\theta)$ . This is called **stochastic gradient descent** (SGD). Ideally, we want this estimator to be **unbiased**, that is, on average, it matches a single true gradient step:

$$\mathbb{E}[\widetilde{\nabla}J(\theta)] = \nabla J(\theta).$$

If J is defined in terms of some training dataset, we might randomly choose a *minibatch* of samples and use them to estimate the prediction error across the *whole* dataset. (This approach is known as **minibatch SGD**.)

Notice that our parameters will stop changing once  $\nabla J(\theta)=0$ . This implies that our current parameters are 'locally optimal' in some sense; it's impossible to increase the function by moving in any direction. If J is convex, then the only point where this happens is at the *global optimum*. Otherwise, if J is nonconvex, the best we can hope for is a *local optimum*.

We can actually show that in a finite number of steps, SGD will find a  $\theta$  that is "close" to a local optimum. More formally, suppose we run SGD for T steps, using an unbiased gradient estimator. Let the step size  $\eta_t$  scale as  $O(1/\sqrt{t})$ . Then if J is bounded and  $\beta$ -smooth, and the norm of the gradient estimator has a finite variance, then after T steps:

$$\|\nabla_{\theta} J(\theta)\|^2 \le O\left(M\beta\sigma^2/T\right).$$

In another perspective, the local "landscape" of J around  $\theta$  becomes flatter and flatter the longer we run SGD.

## 4.3 REINFORCE and Importance Sampling

Note that the objective function above,  $J(\theta) = \mathbb{E}_{\tau \sim \rho_{\theta}} R(\tau)$ , is very difficult to compute! It requires playing out every possible trajectory, which is clearly infeasible for slightly complex state and action spaces. Can we rewrite this in a form that's more convenient to implement? Specifically, suppose there is some distribution, given by a likelihood  $\rho(\tau)$ , that's easy to sample from (e.g. a database of existing trajectories). We can then rewrite the objective function as follows (all gradients are being taken w.r.t.  $\theta$ ):

$$\begin{split} \nabla J(\theta) &= \nabla \mathop{\mathbb{E}}_{\tau \sim \rho_{\theta}} R(\tau) \\ &= \nabla \mathop{\mathbb{E}}_{\tau \sim \rho} \frac{\rho_{\theta}(\tau)}{\rho(\tau)} R(\tau) & \text{likelihood ratio trick} \\ &= \mathop{\mathbb{E}}_{\tau \sim \rho} \frac{\nabla \rho_{\theta}(\tau)}{\rho(\tau)} R(\tau) & \text{switching gradient and expectation} \end{split}$$

Note that setting  $\rho = \rho_{\theta}$  gives us an alternative form of J that's easier to implement. (Notice the swapped order of  $\nabla$  and  $\mathbb{E}!$ )

$$\nabla J(\theta) = \underset{\tau \sim \rho_{\theta}}{\mathbb{E}} [\nabla \log \rho_{\theta}(\tau) \cdot R(\tau)].$$

Consider expanding out  $\rho_{\theta}$ . Note that taking its  $\log$  turns it into a sum of  $\log$  terms, of which only the  $\pi_{\theta}(a_t|s_t)$  terms depend on  $\theta$ , so we can simplify even further to obtain

$$\nabla J(\theta) = \mathbb{E}_{\tau \sim \rho_{\theta}} \left[ \sum_{t=0}^{T-1} \nabla_{\theta} \log \pi_{\theta}(a_t | s_t) R(\tau) \right]$$

In fact, we can perform one more simplification. Intuitively, the action at step t does not affect the reward at previous timesteps. You can also show rigorously that this is the case, and that we only need to consider the present and future rewards to calculate the policy gradient:

$$\nabla J(\theta) = \underset{\tau \sim \rho_{\theta}}{\mathbb{E}} \left[ \sum_{t=0}^{T-1} \nabla_{\theta} \log \pi_{\theta}(a_t|s_t) \sum_{t'=t}^{T-1} r(s_{t'}, a_{t'}) \right]$$

$$= \underset{\tau \sim \rho_{\theta}}{\mathbb{E}} \left[ \sum_{t=0}^{T-1} \nabla_{\theta} \log \pi_{\theta}(a_t|s_t) Q^{\pi_{\theta}}(s_t, a_t) \right]$$

$$(4.3)$$

Note that in the discounted case, the  $Q^{\pi_{\theta}}$  term must becomes  $\lambda^t Q^{\pi_{\theta}}$ . (Make sure this makes sense!) **Exercise:** Prove that this is equivalent to the previous definitions. Also show that this works in the undiscounted case and for infinite horizon.

This expression allows us to estimate the gradient by sampling a few sample trajectories from  $\pi_{\theta}$ , calculating the likelihoods of the chosen actions, and substituting these into the expression above.

For some intuition into how this method works, recall that we update our parameters according to

$$\theta_{t+1} = \theta_t + \nabla J(\theta_t)$$
  
=  $\theta_t + \underset{\tau \sim \rho_{\theta_t}}{\mathbb{E}} \nabla \log \rho_{\theta_t}(\tau) \cdot R(\tau).$ 

Consider the "good" trajectories where  $R(\tau)$  is large. Then  $\theta$  gets updated so that these trajectories become more likely. To see why, recall that  $\rho_{\theta}(\tau)$  is the likelihood of the trajectory  $\tau$  under the policy  $\pi_{\theta}$ , so evaluating the gradient points in the direction that makes  $\tau$  more likely.

This is an example of **importance sampling:** updating a distribution to put more density on "more important" samples (in this case trajectories).

## 4.4 Baselines and advantages

A central idea from supervised learning is the bias-variance tradeoff. So far, our method is *unbiased*, meaning that its average is the true policy gradient. Can we find ways to reduce the variance of our estimator as well?

We can instead subtract a **baseline function**  $b_t : \mathcal{S} \to \mathbb{R}$  at each timestep t. This modifies the policy gradient as follows:

$$\nabla J(\theta) = \mathbb{E}_{\tau \sim \rho_{\theta}} \left[ \sum_{t=0}^{H-1} \nabla \log \pi_{\theta}(a_t | s_t) \left( \left( \sum_{t'=t}^{H-1} r_t \right) - b_t(s_t) \right) \right].$$

For example, we might want  $b_t$  to estimate the average reward-to-go at a given timestep:  $b_t^\theta = \mathbb{E}_{\tau \sim \rho_\theta} \, R_t(\tau)$ . This way, the random variable  $R_t(\tau) - b_t^\theta$  is centered around zero, making certain algorithms more stable.

As a better baseline, we could instead choose the *value function*. Note that the random variable  $Q_t^{\pi}(s,a) - V_t^{\pi}(s)$ , where the randomness is taken over the actions, is also centered around zero. (Recall  $V_t^{\pi}(s) = \mathbb{E}_{a \sim \pi} Q_t^{\pi}(s,a)$ .) In fact, this quantity has a particular name: the **advantage function**. In a sense, it measures how much better this action does than the average for that policy. We can now alternatively and concisely express the policy gradient as follows. Note that the advantage function effectively replaces the Q-function from Equation 4.3:

$$\nabla J(\theta) = \mathbb{E}_{\tau \sim \rho_{\theta}} \left[ \sum_{t=0}^{T-1} \nabla \log \pi_{\theta}(a_t|s_t) A_t^{\pi_{\theta}}(s_t, a_t) \right]$$

Additionally, note that for an optimal policy  $\pi^*$ , the advantage of a given state-action pair is always nonpositive. (Why?)

## 4.5 Policy parameterizations

What are some different ways we could parameterize our policy?

If both the state and action spaces are finite, perhaps we could simply learn a preference value  $\theta_{s,a}$  for each state-action pair. Then to turn this into a valid distribution, we exponentiate each of them, and divide by the total:

$$\pi_{\theta}^{\mathsf{softmax}}(a|s) = \frac{\exp(\theta_{s,a})}{\sum_{s,a'} \exp(\theta_{s,a'})}.$$

However, this doesn't preserve any structure in the states or actions. While this is flexible, it is also prone to overfitting.

#### 4.5.1 Linear in features

Instead, what if we map each state-action pair into some **feature space**  $\phi(s, a) \in \mathbb{R}^p$ ? Then, to map a feature vector to a probability, we take a linear combination  $\theta \in \mathbb{R}^p$  of the features and take a softmax:

$$\pi_{\theta}^{\text{linear in features}}(a|s) = \frac{\exp(\theta^{\top}\phi(s,a))}{\sum_{a'} \exp(\theta^{\top}\phi(s,a'))}.$$

Another interpretation is that  $\theta$  represents the feature vector of the "ideal" state-action pair, as state-action pairs whose features align closely with  $\theta$  are given higher probability.

The score for this parameterization is also quite elegant:

$$\nabla \log \pi_{\theta}(a|s) = \nabla \left( \theta^{\top} \phi(s, a) - \log \left( \sum_{a'} \exp(\theta^{\top} \phi(s, a')) \right) \right)$$
$$= \phi(s, a) - \underset{a' \sim \pi_{\theta}(s)}{\mathbb{E}} \phi(s, a')$$

Plugging this into our policy gradient expression, we get

$$\nabla J(\theta) = \underset{\tau \sim \rho_{\theta}}{\mathbb{E}} \left[ \sum_{t=0}^{T-1} \nabla \log \pi_{\theta}(a_{t}|s_{t}) A_{t}^{\pi_{\theta}} \right]$$

$$= \underset{\tau \sim \rho_{\theta}}{\mathbb{E}} \left[ \sum_{t=0}^{T-1} \left( \phi(s_{t}, a_{t}) - \underset{a' \sim \pi(s_{t})}{\mathbb{E}} \phi(s_{t}, a') \right) A_{t}^{\pi_{\theta}}(s_{t}, a_{t}) \right]$$

$$= \underset{\tau \sim \rho_{\theta}}{\mathbb{E}} \left[ \sum_{t=0}^{T-1} \phi(s_{t}, a_{t}) A_{t}^{\pi_{\theta}}(s_{t}, a_{t}) \right]$$

Why can we drop the  $\mathbb{E}\,\phi(s_t,a')$  term? By linearity of expectation, consider the dropped term at a single timestep:  $\mathbb{E}_{\tau\sim\rho_{\theta}}\left[\left(\mathbb{E}_{a'\sim\pi(s_t)}\,\phi(s,a')\right)A_t^{\pi_{\theta}}(s_t,a_t)\right]$ . By Adam's Law, we can wrap the advantage term in a conditional expectation on the state  $s_t$ . Then we already know that  $\mathbb{E}_{a\sim\pi(s)}\,A_t^{\pi}(s,a)=0$ , and so this entire term vanishes.

### 4.5.2 Neural policies

More generally, we could map states and actions to unnormalized scores via some parameterized function  $f_{\theta}: \mathcal{S} \times \mathcal{A} \to \mathbb{R}$ , such as a neural network, and choose actions according to a softmax:

$$\pi_{\theta}^{\text{general}}(a|s) = \frac{\exp(f_{\theta}(s, a))}{\sum_{a'} \exp(f_{\theta}(s, a'))}.$$

The score can then be written as

$$\nabla \log \pi_{\theta}(a|s) = \nabla f_{\theta}(s, a) - \underset{a \sim \pi_{\theta}(s)}{\mathbb{E}} \nabla f_{\theta}(s, a')$$