Algorithms for Decision Making
Algorithms for Decision Making

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To our families.
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Preface

This book provides a broad introduction to algorithms for optimal decision making under uncertainty. We cover a wide variety of topics related to decision making, introducing the underlying mathematical problem formulations and the algorithms for solving them. Figures, examples, and exercises are provided to convey the intuition behind the various approaches.

This text is intended for advanced undergraduates and graduate students as well as professionals. The book requires some mathematical maturity and assumes prior exposure to multivariable calculus, linear algebra, and probability concepts. Some review material is provided in the appendix. Disciplines where the book would be especially useful include mathematics, statistics, computer science, aerospace, electrical engineering, and operations research.

Fundamental to this textbook are the algorithms, which are all implemented in the Julia programming language. We have found the language to be ideal for specifying algorithms in human readable form. The priority in the design of the algorithmic implementations was interpretability rather than efficiency. Industrial applications, for example, may benefit from alternative implementations. Permission is granted, free of charge, to use the code snippets associated with this book, subject to the condition that the source of the code is acknowledged. We anticipate that others may want to contribute translations of these algorithms to other programming languages. As translations become available, we will link to them from the book’s webpage.

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Acknowledgments

This textbook has grown from a course on decision making under uncertainty taught at Stanford. We are grateful to the students and teaching assistants who have helped shape the course over the past six years.

The authors wish to thank the many individuals who have provided valuable feedback on early drafts of our manuscript, including Edward Balaban, Jean Betterton, Raunak Bhattacharyya, Kelsey Bing, Maxime Bouton, Simon Chauvin, Shushman Choudhury, Jon Cox, Victoria Dax, Richard Dewey, Dea Dressel, Jayesh Gupta, Arec Jamgochian, Rohan Kapre, Mark Koren, Liam Kruse, Tor Lattimore, Ritchie Lee, Sheng Li, Robert Moss, Francesco Piccoli, Jeffrey Sarnoff, Ransalu Senanayake, Chris Strong, Zach Sunberg, Kemal Ure, Josh Wolff, and Zongzhang Zhang. We would like to also thank Ayan Mukhopadhyay, Kunal Menda, and Sydney Katz for their contributions to the discussion in the introductory chapter. Ross Alexander produced many of the exercises throughout the book. It has been a pleasure working with Elizabeth Swayze from the MIT Press in preparing this manuscript for publication.

The style of this book was inspired by Edward Tufte. Among other stylistic elements, we adopted his wide margins and use of small multiples. The typesetting of this book is based on the Tufte-LaTeX package by Kevin Godby, Bil Kleb, and Bill Wood. The book’s color scheme was adapted from the Monokai theme by Jon Skinner of Sublime Text (https://sublimetext.com) and a palette that better accommodates individuals with color-blindness.¹ For plots, we use the viridis colormap defined by Stéfan van der Walt and Nathaniel Smith.

We have also benefited from the various open source packages on which this textbook depends (see appendix). The typesetting of the code is done with the help of pythontex, which is maintained by Geoffrey Poore. The typeface used

for the algorithms is JuliaMono (https://github.com/cormullion/julia mono). Plotting is handled by pgfplots, which is maintained by Christian Feuersänger.
1 Introduction

Many important problems involve decision making under uncertainty, including aircraft collision avoidance, wildfire management, and disaster response. When designing automated decision-making systems or decision-support systems, it is important to account for the various sources of uncertainty when making or recommending decisions. Accounting for these sources of uncertainty and carefully balancing the multiple objectives of the system can be very challenging. We will discuss these challenges from a computational perspective, aiming to provide the theory behind decision-making models and computational approaches. This chapter introduces the problem of decision making under uncertainty, provides some example applications, and outlines the space of possible computational approaches. The chapter then summarizes how various disciplines have contributed to our understanding of intelligent decision making and highlights areas of potential societal impact. We conclude with an outline of the remainder of the book.

1.1 Decision Making

An agent is something that acts based on observations of its environment. Agents may be physical entities, like humans or robots, or they may be nonphysical entities, such as decision support systems that are implemented entirely in software. As shown in figure 1.1, the interaction between the agent and the world follows an observe-act cycle or loop.

The agent at time $t$ receives an observation of the world, denoted $o_t$. Observations may be made, for example, through a biological sensory process as in humans or by a sensor system like radar in an air traffic control system. Observations are often incomplete or noisy; humans may not see an approaching aircraft or a radar
system might miss a detection through electromagnetic interference. The agent then chooses an action $a_t$ through some decision-making process to be discussed later. This action, such as sounding an alert, may have a nondeterministic effect on the world.

Our focus is on agents that interact intelligently in the world to achieve their objectives over time. Given the past sequence of observations $o_1, \ldots, o_t$ and knowledge about the environment, the agent must choose an action $a_t$ that best achieves its objectives in the presence of various sources of uncertainty,\(^1\) including:

1. **outcome uncertainty**, where the effects of our actions are uncertain,
2. **model uncertainty**, where our model of the problem is uncertain,
3. **state uncertainty**, where the true state of the environment is uncertain, and
4. **interaction uncertainty**, where the behavior of the other agents interacting in the environment is uncertain.

This book is organized around these four sources of uncertainty. Making decisions in the presence of uncertainty is central to the field of *artificial intelligence*\(^2\) as well as many other fields, as outlined in section 1.4. We will discuss a variety of algorithms, or descriptions of computational processes, for making decisions that are robust to uncertainty.

### 1.2 Applications

The decision making framework presented in the previous section can be applied to a wide variety of domains. This section discusses a few conceptual examples with real-world applications. Appendix F outlines additional notional examples that are used throughout this text to demonstrate the algorithms we discuss.

---

\(^1\) We focus on discrete time problems. Continuous time problems are studied in the field of *control theory*. D. E. Kirk, *Optimal Control Theory: An Introduction*. Prentice-Hall, 1970.

1.2.1 Aircraft Collision Avoidance

To help prevent mid-air collisions between aircraft, we want to design a system that can alert pilots to potential threats and direct them how to maneuver. The system communicates with the transponders of other aircraft to identify their positions with some degree of accuracy. Deciding what guidance to provide to the pilots from this information is challenging. There is uncertainty in how quickly the pilots will respond and how strongly they will comply with the guidance. In addition, there is uncertainty in the behavior of the other aircraft in the vicinity. We want our system to alert sufficiently early to provide enough time for the pilots to maneuver the aircraft to avoid collision, but we do not want our system to alert too early and result in many unnecessary maneuvers. Since this system is to be used continuously worldwide, we need the system to provide an exceptional level of safety.

1.2.2 Automated Driving

We want to build an autonomous vehicle that can safely drive in urban environments. The vehicle must rely upon a suite of sensors to perceive its environment to make safe decisions. One type of sensor is lidar, which involves measuring laser reflections off of the environment to determine distances to obstacles. Another type of sensor is a camera, which, through computer vision algorithms, can detect pedestrians and other vehicles. Both of these types of sensors are imperfect and susceptible to noise and occlusions. For example, a parked truck may occlude a pedestrian that may be trying to cross at a crosswalk. Our system must predict the intentions and future paths of other vehicles, pedestrians, and other road users from their observable behavior in order to safely navigate to our destination.

1.2.3 Breast Cancer Screening

Worldwide, breast cancer is the most common cancer in women. Detecting breast cancer early can help save lives, with mammography being the most effective screening tool available. However, mammography carries with it potential risks, including false positives, which can result in unnecessary and invasive diagnostic followup. Research over the years has resulted in various population-based screening schedules based on age in order to balance benefits and risks of the tests. Developing a system that can make personalized recommendations based on age and other factors could improve the effectiveness of breast cancer screening.
on personal risk characteristics and screening history has the potential to result in better outcomes for women.\textsuperscript{5} The success of such a system can be compared to population-wide screening schedules in terms of total expected quality-adjusted life years, the number of mammograms, false-positives, and risk of undetected invasive cancer.

1.2.4 Financial Consumption and Portfolio Allocation

Suppose we want to build a system that recommends to an individual how much of their wealth they should consume that year and how to allocate their investments.\textsuperscript{6} The investment portfolio may include stocks and bonds with different levels of risk and expected return. The evolution of wealth is stochastic due to uncertainty in both earned and investment income, often increasing until near retirement and then steadily decreasing. The enjoyment that comes from the consumption of a unit of wealth in a year typically diminishes with the amount consumed, resulting in a desire to smooth consumption over the lifespan of the individual.

1.2.5 Distributed Wildfire Surveillance

A major challenge in fighting wildfires is situational awareness. The state of the fire evolves over time, influenced by factors such as wind and the distribution of fuel in the environment. Many wildfires span large geographic regions. One concept for monitoring a wildfire is to use a team of drones equipped with sensors to fly above the fire.\textsuperscript{7} The sensing range of individual drones are limited, but the information from the team can be fused to provide a unified snapshot of the situation to inform resource allocation decisions. We would like the team to autonomously determine how to collaborate with each other to provide the best coverage of the fire. Effective monitoring requires deciding how to maneuver to cover areas where new sensor information is likely to be useful; spending time in areas where we are certain the fire is burning or not burning would be uninformative. Identifying important areas to explore requires reasoning about the stochastic evolution of the fire given only imperfect knowledge of the current state of the fire.


\textsuperscript{7} This application was explored by K. D. Julian and M. J. Kochenderfer, “Distributed Wildfire Surveillance with Autonomous Aircraft Using Deep Reinforcement Learning,” \textit{AIAA Journal on Guidance, Control, and Dynamics}, vol. 42, no. 8, pp. 1768–1778, 2019.
1.3 Methods

There are many different methods for designing decision agents. Depending on the application, some may be more appropriate than others. They differ in the responsibilities of the designer and the tasks left to automation. This section briefly overviews a collection of these methods. The book will focus primarily on planning and reinforcement learning, but some of the techniques will involve elements of supervised learning and optimization.

1.3.1 Explicit Programming

The most direct method for designing a decision agent is to anticipate all the different scenarios the agent might find itself in and then explicitly program the agent to do what is desired. The explicit programming approach may work well for simple problems, but it places a large burden on the designer to provide a complete strategy. Various agent programming languages and frameworks have been proposed to make programming agents easier.

1.3.2 Supervised Learning

In some problems, it may be easier to show an agent what to do rather than to write a program for the agent to follow. The designer provides a set of training examples, and an automated learning algorithm must generalize from these examples. This approach is known as supervised learning and has been widely applied to classification problems. This technique is sometimes called behavioral cloning when applied to learning mappings from observations to actions. Behavioral cloning works well when an expert designer actually knows the best course of action for a representative collection of example situations. Although there exists a wide variety of different learning algorithms, they generally cannot perform better than human designers in new situations.

1.3.3 Optimization

Another approach is for the designer to specify the space of possible decision strategies and a performance measure to be maximized. Evaluating the performance of a decision strategy generally involves running a batch of simulations. The optimization algorithm then performs a search in this space for the optimal
strategy. If the space is relatively low dimensional and the performance measure
does not have many local optima, then various local or global search methods may
be appropriate. Although knowledge of a dynamic model is generally assumed in
order to run the simulations, it is not otherwise used to guide the search, which
can be important in complex problems.

1.3.4 Planning

Planning is a form of optimization that uses a model of the problem dynamics
to help guide the search. A broad literature has arisen on planning problems,
much of it focused on deterministic problems. For some problems, it may be
acceptable to approximate the dynamics with a deterministic model. Assuming a
deterministic model allows us to use methods that can more easily scale to high-
dimensional problems. For other problems, accounting for future uncertainty is
absolutely critical. This book focuses entirely on problems in which accounting
for uncertainty is important.

1.3.5 Reinforcement learning

Reinforcement learning relaxes the assumption in planning that a model is known
ahead of time. Instead, the decision-making strategy is learned while the agent
interacts with the world. The designer only has to provide a performance measure;
it is up to a learning algorithm to optimize the behavior of the agent. One of the
interesting complexities that arises in reinforcement learning is that the choice
of action impacts not only the immediate success of the agent in achieving its
objectives but also the agent’s ability to learn about the environment and identify
the characteristics of the problem that it can exploit.

1.4 History

The theory of automating the process of decision making has roots in the dreams
of early philosophers, scientists, mathematicians, and writers. The ancient Greeks
began incorporating automation into myths and stories as early as 800 B.C. The
word automaton was first used in Homer’s Iliad, which contains references to the
notion of automatic machines including mechanical tripods used to serve dinner
guests. In the seventeenth century, philosophers proposed the use of logic rules

8 S. Vasileiadou, D. Kalligeropoulos, and N. Karkanias, “Systems,
Modelling and Control in Ancient Greece: Part 1: Mythical Au-
to automatically settle disagreements. Their ideas created the foundation for mechanized reasoning.

Beginning in the late eighteenth century, inventors began creating automatic machines to perform labor. In particular, a series of innovations in the textile industry led to the development of the automatic loom, which in turn laid the foundation for the first factory robots.\(^9\) In the early nineteenth century, the use of intelligent machines to automate labor began to make its way into science fiction novels. The word *robot* originated in Czech writer Karel Čapek’s play titled *Rossum’s Universal Robots* about machines that could perform work humans would prefer not to do. The play inspired other science fiction writers to incorporate robots into their writing. In the mid-twentieth century, notable writer and professor Isaac Asimov laid out his vision for robotics in his famous *Robot Series*.

A major challenge in practical implementations of automated decision making is accounting for uncertainty. Even at the end of the twentieth century, George Dantzig, most famously known for developing the simplex algorithm, stated in 1991:

> In retrospect it is interesting to note that the original problem that started my research is still outstanding—namely the problem of planning or scheduling dynamically over time, particularly planning dynamically under uncertainty. If such a problem could be successfully solved it could (eventually through better planning) contribute to the well-being and stability of the world.\(^10\)

While decision making under uncertainty still remains an active area of research, over the past few centuries, researchers and engineers have come closer to making the concepts posed by these early dreamers possible. Current state-of-the-art decision making algorithms rely on a convergence of concepts developed in multiple disciplines including economics, psychology, neuroscience, computer science, engineering, mathematics, and operations research. This section highlights some major contributions from these disciplines. The cross-pollination between disciplines has led to many recent advances and will likely continue to support growth in the future.

### 1.4.1 Economics

An understanding of individual decision making is central to economic theory, prompting economists to develop techniques to model human decision making. One such technique is utility theory, which was first introduced in the late

---


eighteenth century. Utility theory provides a means to model and compare the desirability of various outcomes. For example, utility can be used to compare the desirability of various monetary values. In the Theory of Legislation, Jeremy Bentham summarized the nonlinearity in the utility of money:

1st. Each portion of wealth has a corresponding portion of happiness.
2nd. Of two individuals with unequal fortunes, he who has the most wealth has the most happiness.
3rd. The excess in happiness of the richer will not be so great as the excess of his wealth.

By combining the concept of utility with the notion of rational decision making, economists in the mid-twentieth century established a basis for the maximum expected utility principle. This principle is a key concept behind the creation of autonomous decision making agents. Utility theory also gave rise to the development of game theory, which attempts to understand the behavior of multiple agents acting in the presence of one another to maximize their interests.

1.4.2 Psychology

Psychologists also study human decision making, typically from the perspective of human behavior. By studying the reactions of animals to stimuli, psychologists have been developing theories of trial-and-error learning since the nineteenth century. Researchers noticed that animals tended to make decisions based on the satisfaction or discomfort they experienced in previous similar situations. Russian psychologist Ivan Pavlov combined this idea with the concept of reinforcement after observing the salivation patterns of dogs when fed. Psychologists found that a pattern of behavior could be strengthened or weakened using a continuous reinforcement of a particular stimulus. In the mid-twentieth century, mathematician and computer scientist Alan Turing expressed the possibility of allowing machines to learn in the same manner:

The organization of a machine into a universal machine would be most impressive if the arrangements of interference involve very few inputs. The training of a human child depends largely on a system of rewards and punishments, and this suggests that it ought to be possible to carry through the organizing with only two interfering inputs, one for ‘pleasure’ or ‘reward’ (R) and the other for ‘pain’ or ‘punishment’ (P).
The work of psychologists laid the foundation for the field of reinforcement learning, a critical technique used to teach agents to make decisions in uncertain environments.\(^\text{15}\)

### 1.4.3 Neuroscience

While psychologists study human behavior as it happens, neuroscientists focus on the biological processes used to create the behavior. At the end of the nineteenth century, scientists found that the brain is composed of an interconnected network of neurons, which is responsible for its ability to perceive and reason about the world. Artificial intelligence pioneer Nils Nilsson describes the application of these findings to decision making as follows:

> Because it is the brain of an animal that is responsible for converting sensory information into action, it is to be expected that several good ideas can be found in the work of neurophysiologists and neuranatomists who study brains and their fundamental components, neurons.\(^\text{16}\)

In the 1940s, researchers first proposed that neurons could be considered as individual “logic units” capable of performing computational operations when pieced together into a network. This work served as a basis for neural networks, which are used in the field of artificial intelligence to perform a variety of complex tasks.

### 1.4.4 Computer Science

In the mid-twentieth century, computer scientists began formulating the problem of intelligent decision making as a problem of symbolic manipulation through formal logic. The computer program *Logic Theorist*, written in the mid-twentieth century to perform automated reasoning, used this way of thinking to prove mathematical theorems. Herbert Simon, one of its inventors, addressed the symbolic nature of the program by relating it to the human mind:

> We invented a computer program capable of thinking non-numerically, and thereby solved the venerable mind/body problem, explaining how a system composed of matter can have the properties of mind.\(^\text{17}\)


These symbolic systems relied heavily on human expertise. An alternative approach to intelligence, called connectionism, was inspired in part by developments in neuroscience and focuses on the use of artificial neural networks as a substrate for intelligence. With the knowledge that neural networks could be trained for pattern recognition, connectionists attempt to learn intelligent behavior from data or experience rather than the hard-coded knowledge of experts. The connectionist paradigm underpinned the success of AlphaGo, the autonomous program that beat a human professional at the game of Go, as well as much of the development of autonomous vehicles. Algorithms that combine both symbolic and connectionist paradigms remain an active area of research today.

1.4.5 Engineering

The field of engineering has focused on allowing physical systems, such as robots, to make intelligent decisions. World-renowned roboticist Sebastian Thrun describes the components of these systems as follows:

Robotics systems have in common that they are situated in the physical world, perceive their environments through sensors, and manipulate their environment through things that move.18

To design these systems, engineers must address perception, planning, and acting. Physical systems perceive the world by using their sensors to create a representation of the salient features of their environment. The field of state-estimation has focused on using sensor measurements to construct a belief about the state of the world. Planning requires reasoning about the ways to execute the tasks they are designed to perform. The planning process has been enabled by advances in the semiconductor industry spanning many decades.19 Once a plan has been devised, an autonomous agent must act on it in the real world. This task requires both hardware in the form of actuators and algorithms to control the actuators and reject disturbances. The field of control theory has focused on the stabilization of mechanical systems through feedback control.20 Automatic control systems are widely used in industry, from the regulation of temperature in an oven to the navigation of aerospace systems.


1.4.6 Mathematics

In order to make informed decisions in uncertain environments, an agent must be able to quantify its uncertainty. The field of decision making relies heavily on probability theory for this task. In particular, Bayesian statistics plays an important role in this text. In 1763, a paper of Thomas Bayes was published posthumously containing what would later be known as Bayes’ rule. His approach to probabilistic inference fell in and out of favor until the mid-twentieth century, when researchers began to find Bayesian methods useful in a number of settings.\footnote{W. M. Bolstad and J. M. Curran, \textit{Introduction to Bayesian Statistics}. Wiley, 2016.} Mathematician Bernard Koopman found practical use for the theory during World War II.

Every operation involved in search is beset with uncertainties; it can be understood quantitatively only in terms of [...] probability. This may now be regarded as a truism, but it seems to have taken the developments in operational research of the Second World War to drive home its practical implications.\footnote{B. O. Koopman, \textit{Search and Screening: General Principles with Historical Applications}. Pergamon Press, 1980.}

Sampling-based methods (sometimes referred to as Monte Carlo methods) developed in the early twentieth century for large scale calculations as part of the Manhattan Project, made some inference techniques possible that would previously have been intractable. These foundations serve as a basis for Bayesian networks, which increased in popularity later in the twentieth century in the field of artificial intelligence.

1.4.7 Operations Research

Operations research is concerned with finding optimal solutions to decision-making problems such as resource allocation, asset investment, or maintenance scheduling. In the late nineteenth century, researchers began to explore the application of mathematical and scientific analysis to the production of goods and services. The field was accelerated during the Industrial Revolution when companies began to subdivide their management into departments responsible for distinct aspects of overall decisions. During World War II, the optimization of decisions was applied to allocating resources to an army. Once the war came to an end, businesses began to notice that the same operations research concepts previously used to make military decisions could help them optimize their business decisions. This realization led to the development of management science, as described by organizational theorist Harold Koontz:
The abiding belief of this group is that, if management, or organization, or planning, or decision making is a logical process, it can be expressed in terms of mathematical symbols and relationships. The central approach of this school is the model, for it is through these devices that the problem is expressed in its basic relationships and in terms of selected goals or objectives.\(^{23}\)

This desire to be able to better model and understand business decisions sparked the development of a number of algorithms used today such as linear programming, dynamic programming, and queuing theory.\(^{24}\)

1.5 Societal Impact

Algorithmic approaches to decision making have transformed society and will likely continue to an even greater extent in the future. This section briefly highlights a few ways decision making algorithms can contribute to society as well as challenges that remain in ensuring broad benefit.\(^{25}\)

Algorithmic approaches have contributed to environmental sustainability. In the context of energy management, for example, Bayesian optimization has been applied to automated home energy management systems. Algorithms from the field of multi-agent systems are used to predict the operation of smart grids, design markets for trading energy, and predict rooftop solar-power adoption. Algorithms have also been developed to protect biodiversity. For example, neural networks are used to automate wildlife census, game-theoretic approaches are used to combat poaching in forests, and optimization techniques are employed to allocate resources for habitat management.

Decision making algorithms have found success in the field of medicine for decades. Such algorithms have been used for matching residents to hospitals and matching organ donors to patients in need. An early application of Bayesian networks, which we will cover in the first part of this book, was disease diagnosis. Since then, Bayesian networks have been widely used in medicine for diagnosis and prognosis of many diseases such as cervical cancer, breast cancer, and glaucoma. The field of medical image processing has been transformed by deep learning, and recently, algorithmic ideas have played an important role in understanding the spread of disease.

Algorithms have enabled us to understand the growth of urban areas and facilitate their design. Data-driven algorithms have been widely used to improve public infrastructure. For example, stochastic processes have been used to predict


failures in water pipelines, deep learning has improved the management of traffic, Markov decision processes and Monte Carlo methods have been employed to improve emergency response. Ideas from decentralized multi-agent systems have optimized travel routes, and path planning techniques have been used to optimize delivery of goods. A major application of decision making algorithms in transportation has been in the development of autonomous cars and improving the safety of aircraft.

Algorithms for optimizing decisions can amplify the impact of its users, regardless of the nature of their intention. If the objective of the user of these algorithms, for example, is to spread misinformation during a political election, then optimization processes can help facilitate this. However, similar algorithms can be used to monitor and counteract the spread of false information. Sometimes the implementation of these decision making algorithms can lead to downstream consequences that were not intended by their users.  

Although algorithms have the potential to bring significant benefits, there are also challenges associated with their implementation in society. Data-driven algorithms often suffer from inherent biases and blind spots due to the way data is collected. As algorithms become part of our lives, it is important to understand how the risk of bias can be reduced and how the benefits of algorithmic progress can be distributed in a manner that is equitable and fair. Algorithms can also be vulnerable to adversarial manipulation, and it is critical that we design algorithms that are robust to such attacks. It is also important to extend moral and legal frameworks for preventing unintended consequences and assigning responsibility.

1.6 Overview

This book is divided into five parts. The first part addresses the problem of reasoning about uncertainty and objectives in simple decisions at a single point in time. The second part extends decision making to sequential problems, where we must make a sequence of decisions in response to information that we gather along the way about the outcomes of our actions. The third part addresses model uncertainty, where we do not start with a known model and must learn how to act through interaction with the environment. The fourth part addresses state uncertainty, where we do not know the current state of the environment due

to imperfect perceptual information. The final part discusses decision contexts involving multiple agents.

1.6.1 Probabilistic Reasoning

Rational decision making requires reasoning about our uncertainty and objectives. This part of the book begins by discussing how to represent uncertainty as a probability distribution. Real-world problems require reasoning about distributions over many different variables. We will discuss how to construct these models, how to use them to make inferences, and how to learn their parameters and structure from data. We then introduce the foundations of utility theory and show how it forms the basis for rational decision making under uncertainty through the maximum expected utility principle. We then discuss how notions of utility theory can be incorporated into the probabilistic graphical models introduced earlier to form what are called decision networks.

1.6.2 Sequential Problems

Many important problems require that we make a series of decisions. The same principle of maximum expected utility still applies, but optimal decision making in a sequential context requires reasoning about future sequences of actions and observations. This part of the book will discuss sequential decision problems in stochastic environments where the outcomes of our actions are uncertain. We will focus on a general formulation of sequential decision problems under the assumption that the model is known and that the environment is fully observable. We will relax both of these assumptions later. Our discussion will begin with the introduction of the Markov decision process (MDP), the standard mathematical model for sequential decision problems. We will discuss several approaches for finding exact solutions to these types of problems. Because large problems sometimes do not permit exact solutions to be efficiently found, we will discuss a collection of both offline and online approximate solution methods along with a type of method that involves directly searching the space of parameterized decision policies. Finally, we will discuss approaches for validating that our decision strategies will perform as expected when deployed in the real world.
1.6.3 Model Uncertainty

In our discussion of sequential decision problems, we have assumed that the transition and reward models are known. In many problems, however, the dynamics and rewards are not known exactly, and the agent must learn to act through experience. By observing the outcomes of its actions in the form of state transitions and rewards, the agent is to choose actions that maximize its long-term accumulation of rewards. Solving such problems in which there is model uncertainty is the subject of the field of reinforcement learning and the focus of this part of the book. We will discuss several challenges in addressing model uncertainty. First, the agent must carefully balance exploration of the environment with the exploitation of that knowledge gained through experience. Second, rewards may be received long after the important decisions have been made, so credit for later rewards must be assigned to earlier decisions. Third, the agent must generalize from limited experience. We will review the theory and some of the key algorithms for addressing these challenges.

1.6.4 State Uncertainty

In this part, we extend uncertainty to include the state. Instead of observing the state exactly, we receive observations that have only a probabilistic relationship with the state. Such problems can be modeled as a partially observable Markov decision process (POMDP). A common approach to solving POMDPs involves inferring a belief distribution over the underlying state at the current time step and then applying a policy that maps beliefs to actions. This part of the book begins by discussing how to update our belief distribution given a past sequence of observations and actions. It then discusses a variety of exact and approximate methods for solving POMDPs.

1.6.5 Multiagent Systems

Up to this point, there has only been one agent making decisions within the environment. This part expands the previous four parts to multiple agents, discussing the challenges that arise from interaction uncertainty. We begin by discussing simple games, where a group of agents simultaneously each select an action. The result is an individual reward for each agent based on the combined joint action from everyone. The Markov game (MG) represents a generalization of both
simple games to multiple states and the MDP to multiple agents. Consequently, the agents select actions that can stochastically change the state of a shared environment. Algorithms for MGs rely on reinforcement learning due to uncertainty about the policies of the other agents. A *partially observable Markov game (POMG)* introduces state uncertainty, further generalizing MGs and POMDPs, as agents now only receive noisy local observations. The *decentralized partially observable Markov decision process (Dec-POMDP)* focuses the POMG on a collaborative multi-agent team where there is a shared reward among the agents. This part of the book presents these four categories of problems and discusses exact and approximate algorithms that solve them.
PART I:

PROBABILISTIC REASONING

Rational decision making requires reasoning about our uncertainty and objectives. Uncertainty arises from practical and theoretical limitations in our ability to predict future events. For example, predicting exactly how a human operator will respond to advice from a decision support system would require, among other things, a detailed model of how the human brain works. Even the paths of satellites can be difficult to predict. Although Newtonian physics permit highly precise predictions of satellite trajectories, spontaneous failures in the attitude thrusters can result in large deviations from the nominal path, and even small imprecisions can compound over time. To achieve its objectives, a robust decision-making system must account for various sources of uncertainty in the current state of the world and future events. This part of the book begins by discussing how to represent uncertainty using probability distributions. Real-world problems require reasoning about distributions over many different variables. We will discuss how to construct these models, how to use them to make inferences, and how to learn their parameters and structure from data. We then introduce the foundations of utility theory and show how it forms the basis for rational decision making under uncertainty. Utility theory can be incorporated into the probabilistic graphical models introduced earlier to form what are called decision networks. We focus on single-step decisions, reserving discussion of sequential decision problems for the next part of the book.
Computationally accounting for uncertainty requires a formal representation. This chapter discusses how to represent uncertainty. We begin by introducing the notion of degree of belief and show how a set of axioms results in our ability to use probability distributions to quantify our uncertainty.1 We discuss several useful forms of distributions over both discrete and continuous variables. Because many important problems involve probability distributions over a large number of variables, we discuss a way to represent joint distributions efficiently that takes advantage of conditional independence between variables.

2.1 Degrees of Belief and Probability

In problems involving uncertainty, it is essential to be able to compare the plausibility of different statements. We would like to be able to represent, for example, that proposition $A$ is more plausible than proposition $B$. If $A$ represents “my actuator failed”, and $B$ represents “my sensor failed”, then we would write $A \succ B$. Using this basic relation $\succ$, we can define several other relations:

\[
\begin{align*}
A &\prec B \text{ if and only if } B \succ A \\
A &\sim B \text{ if and only if } \text{ neither } A \succ B \text{ nor } B \succ A \\
A &\preceq B \text{ if and only if } A \succ B \text{ or } A \sim B \\
A &\succeq B \text{ if and only if } B \succ A \text{ or } A \sim B
\end{align*}
\]

We want to make certain assumptions about the relationships induced by the operators $\succ$, \sim, and $\prec$. The assumption of universal comparability requires exactly one of the following to hold: $A \succ B$, $A \sim B$, or $A \prec B$. The assumption of transitivity requires that if $A \succeq B$ and $B \succeq C$ then $A \succeq C$. Universal comparability

and transitivity assumptions lead to an ability to represent plausibility by a real-valued function $P$ that has the following two properties:

$$P(A) > P(B) \text{ if and only if } A \succ B \quad (2.5)$$
$$P(A) = P(B) \text{ if and only if } A \sim B \quad (2.6)$$

If we make a set of additional assumptions about the form of $P$, then we can show that $P$ must satisfy the basic axioms of probability (appendix A.2). If we are certain of $A$, then $P(A) = 1$. If we believe $A$ is impossible, then $P(A) = 0$. Uncertainty in the truth of $A$ is represented by values in between the two extrema. Hence, probability masses must lie between 0 and 1 with $0 \leq P(A) \leq 1$.

### 2.2 Probability Distributions

A probability distribution assigns probabilities to different outcomes. There are different ways to represent probability distributions depending on whether they involve discrete or continuous outcomes.

#### 2.2.1 Discrete Probability Distributions

A discrete probability distribution is a distribution over a discrete set of values. We can represent such distributions as a probability mass function, which assigns a probability to every possible assignment of its input variable to a value. For example, suppose we have a variable $X$ that can take on one of $n$ different values: $1, \ldots, n$, or, using colon notation, $1 : n$. A distribution associated with $X$ specifies the $n$ probabilities of the various assignments of values to that variable, in particular $P(X = 1), \ldots, P(X = n)$. Figure 2.1 shows an example of a discrete distribution.

There are constraints on the probability masses associated with discrete distributions. The masses must sum to one:

$$\sum_{i=1}^{n} P(X = i) = 1 \quad (2.7)$$

and $0 \leq P(X = i) \leq 1$ for all $i$. 

---


\(^5\) We will often use this colon notation for compactness. Other texts sometimes use the notation $[1 \ldots n]$ for integer intervals from 1 to $n$. We will also use this colon notation to index into vectors and matrices. For example $x_{1:n}$ represents $x_1, \ldots, x_n$. The colon notation is sometimes used in programming languages, such as Julia and MATLAB.
For notational convenience, we will use lowercase letters and superscripts as shorthand when discussing the assignment of values to variables. For example, \( P(x^3) \) is shorthand for \( P(X = 3) \). If \( X \) is a binary variable, it can take on the value true or false. We will use 0 to represent false and 1 to represent true. For example, we use \( P(x^0) \) to represent the probability \( X \) is false.

The parameters of a distribution govern the probabilities associated with different assignments. For example, if we use \( X \) to represent the outcome of a roll of a six-sided die, then we would have \( P(x^1) = \theta_1, \ldots, P(x^6) = \theta_6 \), with \( \theta_{1:6} \) being the six parameters of the distribution. However, we need only five independent parameters to uniquely specify the distribution over the outcomes of the roll because we know that the distribution must sum to 1.

### 2.2.2 Continuous Probability Distributions

A continuous probability distribution is a distribution over a continuous set of values. Representing a distribution over a continuous variable is a little less straightforward than for a discrete variable. For instance, in many continuous distributions, the probability that a variable takes on a particular value is infinitesimally small. One way to represent a continuous probability distribution is to use a probability density function (see figure 2.2), represented with lowercase letters. If \( p(x) \) is a probability density function over \( X \), then \( p(x) \, dx \) is the probability \( X \) falls within the interval \((x, x + dx)\) as \( dx \to 0 \). Similar to how the probability masses associated with a discrete distribution must sum to 1, a probability density function \( p(x) \) must integrate to 1:

\[
\int_{-\infty}^{\infty} p(x) \, dx = 1
\]  

(2.8)

Another way to represent a continuous distribution is with a cumulative distribution function (see figure 2.3), which specifies the probability mass associated with values below some threshold. If we have a cumulative distribution function \( P \) associated with variable \( X \), then \( P(x) \) represents the probability mass associated with \( X \) taking on a value less than or equal to \( x \). A cumulative distribution function can be defined in terms of a probability density function \( p \) as follows:

\[
cdf_X(x) = P(X \leq x) = \int_{-\infty}^{x} p(x') \, dx'
\]

(2.9)
Related to the cumulative distribution function is the quantile function, also called the inverse cumulative distribution function (see figure 2.4). The value of \( x(\alpha) \) is the value \( x \) such that \( P(X \leq x) = \alpha \). In other words, the quantile function returns the minimum value of \( x \) whose cumulative distribution value exceeds \( \alpha \). Of course, we have \( 0 \leq \alpha \leq 1 \).

There are many different parameterized families of distributions. We outline several in appendix B. A simple distribution family is the uniform distribution \( U(a, b) \), which assigns probability density uniformly between \( a \) and \( b \), and zero elsewhere. Hence, the probability density function is \( p(x) = 1/(b - a) \) for \( x \) in the interval \([a, b]\). We can use \( U(x \mid a, b) \) to represent the density at \( x \). The support of a distribution is the set of values that are assigned non-zero density. In the case of \( U(a, b) \), the support is the interval \([a, b]\). See example 2.1.

The uniform distribution \( U(0, 10) \) assigns equal probability to all values in the range \([0, 10]\) with a probability density function:

\[
U(x \mid 0, 10) = \begin{cases} 1/10 & \text{if } 0 \leq x \leq 10 \\ 0 & \text{otherwise} \end{cases} \tag{2.10}
\]

The probability that a random sample from this distribution is equal to the constant \( \pi \) is essentially zero. However, we can define non-zero probabilities for samples being within some interval, say \([3, 5]\). For example, the probability that a sample lies between 3 and 5 given the distribution plotted below is:

\[
\int_{3}^{5} U(x \mid 0, 10) \, dx = \frac{5 - 3}{10} = \frac{1}{5} \tag{2.11}
\]

The support of this distribution is the interval \([0, 10]\).
Another common distribution for continuous variables is the Gaussian distribution (also called the normal distribution). The Gaussian distribution is parameterized by a mean \( \mu \) and variance \( \sigma^2 \):

\[
p(x) = \mathcal{N}(x \mid \mu, \sigma^2)
\]  
(2.12)

Here, \( \sigma \) is the standard deviation, which is the square root of the variance. The variance is also commonly denoted by \( \nu \). We use \( \mathcal{N}(\mu, \sigma^2) \) to represent a Gaussian distribution with parameters \( \mu \) and \( \sigma^2 \) and \( \mathcal{N}(x \mid \mu, \sigma^2) \) to represent the probability density at \( x \) as given by

\[
\mathcal{N}(x \mid \mu, \sigma^2) = \frac{1}{\sigma} \phi \left( \frac{x - \mu}{\sigma} \right)
\]  
(2.13)

where \( \phi \) is the standard normal density function:

\[
\phi(x) = \frac{1}{\sqrt{2\pi}} \exp \left( -\frac{x^2}{2} \right)
\]  
(2.14)

Appendix B shows plots of Gaussian density functions with different parameters. Although a Gaussian distribution is often convenient because it is defined by only two parameters and makes computation and derivation easy, it has some limitations. It assigns non-zero probability to large positive and negative values, which may not be appropriate for the quantity we are trying to model. For example, we might not want to assign non-zero probabilities for aircraft flying below the ground or at infeasible altitudes. We can use a truncated Gaussian distribution (see figure 2.5) to bound the support of possible values, that is, the range of values assigned non-zero probabilities. The density function is given by

\[
\mathcal{N}(x \mid \mu, \sigma^2, a, b) = \frac{1}{\sigma} \phi \left( \frac{x - \mu}{\sigma} \right) \frac{\Phi \left( \frac{b - \mu}{\sigma} \right) - \Phi \left( \frac{a - \mu}{\sigma} \right)}{\Phi \left( \frac{b - \mu}{\sigma} \right) - \Phi \left( \frac{a - \mu}{\sigma} \right)}
\]  
(2.15)

when \( x \) is within the interval \((a, b)\).

The function \( \Phi \) is the standard normal cumulative distribution function as given by

\[
\Phi(x) = \int_{-\infty}^{x} \phi(x') \, dx'
\]  
(2.16)
The Gaussian distribution is *unimodal*, meaning that there is a point in the distribution at which the density increases on one side and decreases on the other side. There are different ways to represent continuous distributions that are *multimodal*. One way is to use a *mixture model*, which is a mixture of multiple distributions. We mix together a collection of unimodal distributions to obtain a multimodal distribution. A *Gaussian mixture model* is a mixture model that is simply a weighted average of different Gaussian distributions. The parameters of a Gaussian mixture model include the parameters of the Gaussian distribution components $\mu_{1:n}, \sigma^2_{1:n}$ as well as their weights $\rho_{1:n}$. The density is given by

$$p(x \mid \mu_{1:n}, \sigma^2_{1:n}, \rho_{1:n}) = \sum_{i=1}^{n} \rho_i N(x \mid \mu_i, \sigma^2_i)$$  \hspace{1cm} (2.17)

where the weights must sum to 1. Example 2.2 shows a Gaussian mixture model with two components.

We can create a Gaussian mixture model with components $\mu_1 = 5, \sigma_1 = 2$ and $\mu_2 = -5, \sigma_2 = 4$, weighted according to $\rho_1 = 0.6$ and $\rho_2 = 0.4$. Below we plot the density of two components scaled by their weights.

Another approach to representing multimodal continuous distributions is through discretization. For example, we can represent a distribution over a continuous variable as a *piecewise-uniform density*. The density is specified by the bin edges, and a probability mass is associated with each bin. Such a piecewise-uniform distribution is a type of mixture model where the components are uniform distributions.
2.3 Joint Distributions

A joint distribution is a probability distribution over multiple variables. A distribution over a single variable is called a univariate distribution, and a distribution over multiple variables is called a multivariate distribution. If we have a joint distribution over two discrete variables \( X \) and \( Y \), then \( P(x, y) \) denotes the probability that both \( X = x \) and \( Y = y \).

From a joint distribution, we can compute a marginal distribution of a variable or a set of variables by summing out all other variables using what is known as the law of total probability:

\[
P(x) = \sum_y P(x, y)
\]  

(2.18)

This property is used throughout this book.

Real-world decision making often requires reasoning about joint distributions involving many variables. Sometimes there are complex relationships between the variables that are important to represent. We may use different strategies to represent joint distributions depending on whether the variables involve discrete or continuous values.

2.3.1 Discrete Joint Distributions

If the variables are discrete, the joint distribution can be represented by a table like the one shown in table 2.1. That table lists all possible assignments of values to three variables. Each variable can only be 0 or 1, resulting in \( 2^3 = 8 \) possible assignments. As with other discrete distributions, the probabilities in the table must sum to 1. It follows that although there are eight entries in the table, only seven of them are independent. If \( \theta_i \) represents the probability in the \( i \)th row in the table, then we only need the parameters \( \theta_1, \ldots, \theta_7 \) to represent the distribution because we know \( \theta_8 = 1 - (\theta_1 + \ldots + \theta_7) \).

If we have \( n \) binary variables, then we need as many as \( 2^n - 1 \) independent parameters to specify the joint distribution. This exponential growth in the number of parameters makes storing the distribution in memory difficult. In some cases, we can assume that our variables are independent, which means that the realization of one does not affect the probability distribution of the other. If \( X \) and \( Y \) are independent, which is sometimes written \( X \perp Y \), then we know \( P(x, y) = P(x)P(y) \)

<table>
<thead>
<tr>
<th>X</th>
<th>Y</th>
<th>Z</th>
<th>( P(X, Y, Z) )</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0.08</td>
</tr>
<tr>
<td>0</td>
<td>0</td>
<td>1</td>
<td>0.31</td>
</tr>
<tr>
<td>0</td>
<td>1</td>
<td>0</td>
<td>0.09</td>
</tr>
<tr>
<td>0</td>
<td>1</td>
<td>1</td>
<td>0.37</td>
</tr>
<tr>
<td>1</td>
<td>0</td>
<td>0</td>
<td>0.01</td>
</tr>
<tr>
<td>1</td>
<td>0</td>
<td>1</td>
<td>0.05</td>
</tr>
<tr>
<td>1</td>
<td>1</td>
<td>0</td>
<td>0.02</td>
</tr>
<tr>
<td>1</td>
<td>1</td>
<td>1</td>
<td>0.07</td>
</tr>
</tbody>
</table>

8 If our distribution is continuous, then we integrate out the other variables when marginalizing. For example,

\[
p(x) = \int p(x, y) \, dy
\]
for all \( x \) and \( y \). Suppose we have binary variables \( X_1, \ldots, X_n \) that are all independent of each other, resulting in \( P(x_1:n) = \prod_i P(x_i) \). This factorization allows us to represent this joint distribution with only \( n \) independent parameters instead of the \( 2^n - 1 \) required when we cannot assume independence (see table 2.2). Independence can result in an enormous savings in terms of representational complexity, but it is often a poor assumption.

We can represent joint distributions in terms of factors. A factor \( \phi \) over a set of variables is a function from assignments of those variables to the real numbers. In order to represent a probability distribution, the real numbers in the factor must be non-negative. A factor with non-negative values can be normalized such that it represents a probability distribution. Algorithm 2.1 provides an implementation for discrete factors, and example 2.3 demonstrates how they work.

<table>
<thead>
<tr>
<th>( X )</th>
<th>( P(X) )</th>
<th>( Y )</th>
<th>( P(Y) )</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>0.85</td>
<td>0</td>
<td>0.45</td>
</tr>
<tr>
<td>1</td>
<td>0.15</td>
<td>1</td>
<td>0.55</td>
</tr>
<tr>
<td>( Z )</td>
<td>( P(Z) )</td>
<td></td>
<td></td>
</tr>
<tr>
<td>0</td>
<td>0.20</td>
<td></td>
<td></td>
</tr>
<tr>
<td>1</td>
<td>0.80</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Table 2.2. If we know the variables in table 2.1 are independent, we can represent \( P(x, y, z) \) using the product \( P(x)P(y)P(z) \). This representation requires only one parameter for each of the three univariate distributions.

```plaintext
struct Variable
    name::Symbol
    m::Int # number of possible values
end

const Assignment = Dict{Symbol,Int}
const FactorTable = Dict{Assignment,Float64}

struct Factor
    vars::Vector{Variable}
    table::FactorTable
end

function variables(ϕ::Factor)
    [var.name for var in ϕ.vars]
end

function select(a::Assignment, varnames::Vector{Symbol}) =
    Assignment(n=>a[n] for n in varnames)

function assignments(vars::AbstractVector{Variable})
    names = [var.name for var in vars]
    return vec([Assignment(n=>v for (n,v) in zip(names, values))
                       for values in product((1:v.m for v in vars)...)])
end

function normalize!(ϕ::Factor)
    z = sum(p for (a,p) in ϕ.table)
    for (a,p) in ϕ.table
        ϕ.table[a] = p/z
    end
    return ϕ
end
```

Algorithm 2.1. Types and functions relevant to working with factors over a set of discrete variables. A variable is given a name (represented as a symbol) and may take on an integer from 1 to \( m \). An assignment is a mapping from variable names to values represented as integers. A factor is defined by a factor table that assigns values to different assignments involving a set of variables. A factor table is a mapping from assignments to real values. This mapping is represented by a dictionary. Any assignments not contained in the dictionary is given 0 value. Also included in this algorithm block are some utility functions for returning the variable names associated with a factor, selecting a subset of an assignment, enumerating possible assignments, and normalizing factors. As discussed in appendix G.3.3, `product` produces the Cartesian product of a set of collections. It is imported from `Base.Iterators`. 

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We can instantiate the table from table 2.1 using the `Factor` type using the following code:

```plaintext
X = Variable(:x, 2)
Y = Variable(:y, 2)
Z = Variable(:z, 2)
ϕ = Factor([X, Y, Z], FactorTable(
    (x=1, y=1, z=1) => 0.08,
    (x=1, y=2, z=1) => 0.09,
    (x=2, y=1, z=1) => 0.01,
    (x=2, y=2, z=1) => 0.02,

    (x=1, y=1, z=2) => 0.31,
    (x=1, y=2, z=2) => 0.37,
    (x=2, y=1, z=2) => 0.05,
    (x=2, y=2, z=2) => 0.07,
))
```

Another approach to reduce the storage required to represent joint distributions with repeated values is to use a decision tree. A decision tree involving three discrete variables is shown in example 2.4. Although the savings in this example in terms of number of parameters may not be significant, it can become quite substantial when there are many variables and many repeated values.

Example 2.3. Constructing a discrete factor. The construction of the factor table using named tuples takes advantage of the utility functions defined in appendix G.5.

Example 2.4. A decision tree can be a more efficient representation of a joint distribution than a table.
2.3.2 Continuous Joint Distributions

We can also define joint distributions over continuous variables. A rather simple distribution is the *multivariate uniform distribution*, which assigns a constant probability density everywhere there is support. We can use \( \mathcal{U}(\mathbf{a}, \mathbf{b}) \) to represent a uniform distribution over a *box*, which is a Cartesian product of intervals with the \( i \)th interval being \([a_i, b_i]\). This family of uniform distributions is a special type of *multivariate product distribution*, which is a distribution defined in terms of the product of univariate distributions. In this case,

\[
\mathcal{U}(\mathbf{x} \mid \mathbf{a}, \mathbf{b}) = \prod_i \mathcal{U}(x_i \mid a_i, b_i)
\]

We can create a mixture model from a weighted collection of multivariate uniform distributions, just as we can with univariate distributions. If we have a joint distribution over \( n \) variables and \( k \) mixture components, we need to define \( k(2n + 1) - 1 \) independent parameters. For each of the \( k \) components, we need to define the upper and lower bounds for each of the variables in addition to its weight. We can subtract 1 because the weights must sum to 1. Figure 2.6 shows an example that can be represented by five components.

It is also common to represent piecewise constant density functions by discretizing each of the variables independently. The discretization is represented by a set of bin edges for each variable. These bin edges define a grid over the variables. We then associate a constant probability density with each grid cell. The bin edges do not have to be uniformly separated. In some cases, it may be desirable to have increased resolution around certain values. Different variables might have different bin edges associated with them. If there are \( n \) variables and \( m \) bins for each variable, then we need \( m^n - 1 \) independent parameters to define the distribution—in addition to the values that define the bin edges.

In some cases, it may be more memory efficient to represent a continuous joint distribution as a decision tree in a manner similar to what we discussed for discrete joint distributions. The internal nodes compare variables against thresholds and the leaf nodes are density values. Figure 2.7 shows a decision tree that represents the density function in figure 2.6.

Another useful distribution is the *multivariate Gaussian distribution* with the density function

\[
\mathcal{N}(\mathbf{x} \mid \mathbf{\mu}, \Sigma) = \frac{1}{(2\pi)^{n/2} |\Sigma|^{1/2}} \exp \left( -\frac{1}{2} (\mathbf{x} - \mathbf{\mu})^\top \Sigma^{-1} (\mathbf{x} - \mathbf{\mu}) \right)
\]
where \( x \) is in \( \mathbb{R}^n \), \( \mu \) is the mean vector, and \( \Sigma \) is the covariance matrix. The density function above requires that \( \Sigma \) be positive definite (definition is reviewed in appendix A.5). The number of independent parameters is equal to \( n + (n + 1)n/2 \), the number of components in \( \mu \) added to the number of components in the upper triangle of matrix \( \Sigma \). Appendix B shows plots of different multivariate Gaussian density functions. We can also define multivariate Gaussian mixture models. Figure 2.8 shows an example of one with three components.

If we have a multivariate Gaussian with all of the variables independent, then the covariance matrix \( \Sigma \) is diagonal with only \( n \) independent parameters. In fact, we can write the density function as a product of univariate Gaussian densities:

\[
\mathcal{N}(x \mid \mu, \Sigma) = \prod_i \mathcal{N}(x_i \mid \mu_i, \Sigma_{ii})
\]  

(2.21)

### 2.4 Conditional Distributions

The previous section introduced the idea of independence, which can help reduce the number of parameters used to define a joint distribution. However, as was mentioned, independence can be too strong of an assumption. This section will introduce the idea of conditional independence, which can help reduce the number of independent parameters without making assumptions that are as strong. Before discussing conditional independence, we will first introduce the notion of a conditional distribution, which is a distribution over a variable given the value of one or more others.

\[\text{Figure 2.8. Multivariate Gaussian mixture model with three components. The components are mixed together with weights 0.1, 0.5, and 0.4, respectively.}\]
The definition of conditional probability states that
\[
P(x \mid y) = \frac{P(x, y)}{P(y)} \tag{2.22}
\]
where \(P(x \mid y)\) is read as “probability of \(x\) given \(y\).” In some contexts, it is common to refer to \(y\) as evidence.

Since a conditional probability distribution is a probability distribution over one or more variables given some evidence, we know that
\[
\sum_x P(x \mid y) = 1 \tag{2.23}
\]
for a discrete \(X\). If \(X\) is continuous, it integrates to 1.

We can incorporate the definition of conditional probability into equation (2.18) to obtain a slightly different form of the law of total probability:
\[
P(x) = \sum_y P(x \mid y)P(y) \tag{2.24}
\]
for a discrete distribution.

Another useful relationship that follows from the definition of conditional probability is Bayes’ rule:
\[
P(x \mid y) = \frac{P(y \mid x)P(x)}{P(y)} \tag{2.25}
\]
If we have a representation of a conditional distribution \(P(y \mid x)\), we can apply Bayes’s rule to swap the \(y\) and \(x\) to obtain the conditional distribution \(P(x \mid y)\).

We will now discuss a variety of ways to represent conditional probability distributions over discrete and continuous variables.

### 2.4.1 Discrete Conditional Models

A conditional probability distribution over discrete variables can be represented using a table. In fact, we can use the same discrete factor representation that we used in section 2.3.1 for joint distributions. Table 2.3 shows an example of a table representing \(P(X \mid Y, Z)\) with all binary variables. In contrast with a joint table (e.g., table 2.1), the column containing the probabilities need not sum to 1. However, if we sum over the probabilities that are consistent with what we are conditioning on, we must get 1. For example, conditioning on \(y^0\) and \(z^0\) (the evidence), we have
\[
P(x^0 \mid y^0, z^0) + P(x^1 \mid y^0, z^0) = 0.08 + 0.92 = 1 \tag{2.26}
\]
Conditional probability tables can become quite large. If we were to create a table like table 2.3 where all variables can take on $m$ values and we are conditioning on $n$ variables, there would be $m^{n+1}$ rows. However, since the $m$ values of the variable we are not conditioning on must sum to 1, there are only $(m-1)m^n$ independent parameters. There is still an exponential growth with the number of variables on which we condition. When there are many repeated values in the conditional probability table, a decision tree (introduced in section 2.3.1) may be a more efficient representation.

### 2.4.2 Conditional Gaussian Models

A conditional Gaussian model can be used to represent a distribution over a continuous variable given one or more discrete variables. For example, if we have a continuous variable $X$ and a discrete variable $Y$ with values $1 : n$, we could define a conditional Gaussian model as follows:

\[
p(x \mid y) = \begin{cases} 
\mathcal{N}(x \mid \mu_1, \sigma_1^2) & \text{if } y^1 \\
\vdots \\
\mathcal{N}(x \mid \mu_n, \sigma_n^2) & \text{if } y^n 
\end{cases}
\]  

with parameter vector $\theta = [\mu_{1:n}, \sigma_{1:n}]$. All $2n$ of those parameters can be varied independently. If we want to condition on multiple discrete variables, we just need to add more cases and associated parameters.

### 2.4.3 Linear Gaussian Models

The linear Gaussian model of $P(X \mid Y)$ represents the distribution over a continuous variable $X$ as a Gaussian distribution with the mean being a linear function of the value of the continuous variable $Y$. The conditional density function is

\[
p(x \mid y) = \mathcal{N}(x \mid my + b, \sigma^2) 
\]  

with parameters $\theta = [m, b, \sigma]$. The mean is a linear function of $y$ defined by parameters $m$ and $b$. The variance is constant. Figure 2.9 shows an example.
2.4.4 Conditional Linear Gaussian Models

The *conditional linear Gaussian* model combines the ideas of conditional Gaussian and linear Gaussian models to be able to handle conditioning a continuous variable on both discrete and continuous variables. Suppose we want to represent $p(X \mid Y, Z)$, where $X$ and $Y$ are continuous and $Z$ is discrete with values $1 : n$. The conditional density function is then

\[
p(x \mid y, z) = \begin{cases} 
\mathcal{N}(x \mid m_1 y + b_1, \sigma_1^2) & \text{if } z^1 \\
\vdots \\
\mathcal{N}(x \mid m_n y + b_n, \sigma_n^2) & \text{if } z^n
\end{cases}
\]  

Above, the parameter vector $\theta = [m_1 : n, b_1 : n, \sigma_1 : n]$ has $3n$ components.

2.4.5 Sigmoid Models

We can use a *sigmoid*\(^{11}\) model to represent a distribution over a binary variable conditioned on a continuous variable. For example, we may want to represent $P(x^1 \mid y)$, where $x$ is binary and $y$ is continuous. Of course, we could just set a threshold $\theta$ and say $P(x^1 \mid y) = 0$ if $y < \theta$ and $P(x^1 \mid y) = 1$ otherwise. However, in many applications, we may not want to have such a hard threshold that results in assigning zero probability to $x^1$ for certain values of $y$.

Instead of a hard threshold, we could use a *soft threshold* that assigns low probabilities when below a threshold and high probabilities when above a threshold. One way to represent a soft threshold is to use a *logit model*, which produces a sigmoid curve:

\[
P(x^1 \mid y) = \frac{1}{1 + \exp\left(-\frac{2(y-\theta_1)}{\theta_2}\right)}
\]  

The parameter $\theta_1$ governs the location of the threshold, and $\theta_2$ controls the “softness” or spread of the probabilities. Figure 2.10 shows an example plot of $P(x^1 \mid y)$ with a logit model.

2.4.6 Deterministic Variables

Some problems may involve a *deterministic variable* whose value is fixed given evidence. In other words, we assign probability 1 to a value that is a deterministic function of its evidence. Using a conditional probability table to represent

\[^{11}\text{A sigmoid is an ''S''-shaped curve. There are different ways to define such a curve mathematically, but we will focus on the logit model.}\]
a discrete deterministic variable is possible, but it is wasteful. A single variable instantiation will have probability 1 for each parental instantiation, and the remaining entries will be 0. Our implementation can take advantage of this sparsity for a more compact representation. Algorithms in this text using discrete factors treat any assignments missing from the factor table as having value 0, making it so that we only have to store the assignments that have non-zero probability.

2.5 Bayesian Networks

A Bayesian network can be used to represent a joint probability distribution. The structure of a Bayesian network is defined by a directed acyclic graph consisting of nodes and directed edges. Each node corresponds to a variable. Directed edges connect pairs of nodes, with cycles in the graph being prohibited. The directed edges indicate direct probabilistic relationships. Associated with each node $X_i$ is a conditional distribution $P(X_i \mid \text{pa}(X_i))$, where $\text{pa}(X_i)$ represents the parents of $X_i$ in the graph. Algorithm 2.2 provides an implementation of a Bayesian network data structure. Example 2.5 illustrates the application of Bayesian networks to a satellite-monitoring problem.

```
struct BayesianNetwork
    vars::Vector{Variable}
    factors::Vector{Factor}
    graph::SimpleDiGraph{Int64}
end
```

The chain rule for Bayesian networks specifies how to construct a joint distribution from the local conditional probability distributions. Suppose we have the variables $X_{1:n}$ and want to compute the probability of a particular assignment of all these variables to values $P(x_{1:n})$. The chain rule says

$$P(x_{1:n}) = \prod_{i=1}^{n} P(x_i \mid \text{pa}(x_i))$$

(2.31)

where $\text{pa}(x_i)$ is the particular assignment of the parents of $X_i$ to their values. Algorithm 2.3 provides an implementation for Bayesian networks with conditional probability distributions represented as discrete factors.
Below the caption is a Bayesian network for a satellite-monitoring problem involving five binary variables. Fortunately, battery failure and solar panel failures are both rare, although solar panel failures are somewhat more likely than battery failures. Failures in either can lead to an electrical system failure. There may be causes of electrical system failure other than battery or solar panel failure, such as a problem with the power management unit. An electrical system failure can result in trajectory deviation, which can be observed from the earth by telescope, as well as a communication loss that interrupts the transmission of telemetry and mission data down to various ground stations. Other anomalies not involving the electrical system can result in trajectory deviation and communication loss.

Associated with each of the five variables are five conditional probability distributions. Because $B$ and $S$ do not have any parents, we only need to specify $P(B)$ and $P(S)$. The code below creates a Bayesian network structure with example values for the elements of the associated factor tables. The tuples in the factor tables index into the domains of the variables, which is \{0,1\} for all of the variables. For example, $(e=2,b=1,s=1)$ corresponds to $(e^1,b^0,s^0)$.

```python
B = Variable(':b', 2); S = Variable(':s', 2)
E = Variable(':e', 2)
D = Variable(':d', 2); C = Variable(':c', 2)
vars = [B, S, E, D, C]
factors = [
    Factor([B], FactorTable((b=1,) => 0.99, (b=2,) => 0.01)),
    Factor([S], FactorTable((s=1,) => 0.98, (s=2,) => 0.02)),
    Factor([E,B,S], FactorTable(
        (e=1,b=1,s=1) => 0.90, (e=1,b=1,s=2) => 0.04,
        (e=1,b=2,s=1) => 0.05, (e=1,b=2,s=2) => 0.01,
        (e=2,b=1,s=1) => 0.10, (e=2,b=1,s=2) => 0.96,
        (e=2,b=2,s=1) => 0.95, (e=2,b=2,s=2) => 0.99)),
    Factor([D, E], FactorTable(
        (d=1,e=1) => 0.96, (d=1,e=2) => 0.03,
        (d=2,e=1) => 0.04, (d=2,e=2) => 0.97)),
    Factor([C, E], FactorTable(
        (c=1,e=1) => 0.98, (c=1,e=2) => 0.01,
        (c=2,e=1) => 0.02, (c=2,e=2) => 0.99))
]
graph = SimpleDiGraph(5)
add_edge!(graph, 1, 3); add_edge!(graph, 2, 3)
add_edge!(graph, 3, 4); add_edge!(graph, 3, 5)
bn = BayesianNetwork(vars, factors, graph)
```

Example 2.5. A Bayesian network representing a satellite-monitoring problem. Below is the structure of the network represented as a directed acyclic graph. Associated with each node is a conditional probability distribution.

```
BN

B battery failure
S solar panel failure
E electrical system failure
D trajectory deviation
C communication loss
```

\begin{itemize}
    \item \textbf{$B$}: \text{battery failure}
    \item \textbf{$S$}: \text{solar panel failure}
    \item \textbf{$E$}: \text{electrical system failure}
    \item \textbf{$D$}: \text{trajectory deviation}
    \item \textbf{$C$}: \text{communication loss}
\end{itemize}
2.6 Conditional Independence

The reason that a Bayesian network can represent a joint distribution with fewer independent parameters than would normally be required is due to the conditional independence assumptions encoded in its graphical structure. Conditional independence is a generalization of the notion of independence introduced in section 2.3.1. Variables $X$ and $Y$ are conditionally independent given $Z$ if and only if $P(X, Y \mid Z) = P(X \mid Z)P(Y \mid Z)$. The assertion that $X$ and $Y$ are conditionally independent given $Z$ is written $(X \perp Y \mid Z)$. It is possible to show from this definition that $(X \perp Y \mid Z)$ if and only if $P(X \mid Z) = P(X \mid Y, Z)$. Given $Z$, information about $Y$ provides no additional information about $X$, and vice versa. Example 2.6 provides an example.

We can use a set of rules to determine whether the structure of a Bayesian network implies that two variables must be conditionally independent given a set of other evidence variables. Suppose we want to check whether $(A \perp B \mid C)$ is implied by the network structure, where $C$ is a set of evidence variables. We have

Algorithm 2.3. A function for evaluating the probability of an assignment given a Bayesian network $bn$. For example, if $bn$ is as defined in example 2.5, then $a = (b=1,s=1,e=1,d=2,c=1)$ probability($bn$, Assignment($a$)) returns $0.03422865599999996$. 

The power of Bayesian networks comes from their ability to reduce the number of parameters required to specify a joint probability distribution.

In the satellite example, suppose we want to compute the probability that nothing is wrong, that is, $P(b^0, s^0, e^0, d^0, c^0)$. From the chain rule,

$$P(b^0, s^0, e^0, d^0, c^0) = P(b^0)P(s^0)P(e^0 \mid b^0, s^0)P(d^0 \mid e^0)P(c^0 \mid e^0)$$ \hspace{1cm} (2.32)

If we had fully specified a joint distribution over the five variables $B, S, E, D,$ and $C$, then we would have needed $2^5 - 1 = 31$ independent parameters. The structure assumed in our Bayesian network allows us to specify the joint distribution using only $1 + 1 + 4 + 2 + 2 = 10$ independent parameters. The difference between 10 and 31 does not represent an especially significant savings in the number of parameters, but the savings can become enormous in larger Bayesian networks. The power of Bayesian networks comes from their ability to reduce the number of parameters required to specify a joint probability distribution.

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2021-03-07 23:12:00-08:00, revision 80bf4d6, comments to bugs@algorithmsbook.com
Suppose the presence of satellite trajectory deviation ($D$) is conditionally independent of whether we have a communication loss ($C$) given knowledge of whether we have an electrical system failure ($E$). We would write this ($D \perp C \mid E$). If we know that we have an electrical system failure, then the fact that we observe a loss of communication has no impact on our belief that there is a trajectory deviation. We may have an elevated expectation that there is a trajectory deviation, but that is only because we know that an electrical system failure has occurred.

Example 2.6. Conditional independence in the satellite-tracking problem.

to check all possible undirected paths from $A$ to $B$ for what is called $d$-separation. A path between $A$ and $B$ is $d$-separated by $C$ if any of the following are true:

1. The path contains a chain of nodes, $X \rightarrow Y \rightarrow Z$, such that $Y$ is in $C$.

2. The path contains a fork, $X \leftrightarrow Y \rightarrow Z$, such that $Y$ is in $C$.

3. The path contains an inverted fork (also called a $v$-structure), $X \rightarrow Y \leftrightarrow Z$, such that $Y$ is not in $C$ and no descendant of $Y$ is in $C$. Example 2.7 provides some intuition for this rule.

We say that $A$ and $B$ are $d$-separated by $C$ if all paths between $A$ and $B$ are $d$-separated by $C$. This $d$-separation implies that ($A \perp B \mid C$). Example 2.8 demonstrates this process for checking whether a graph implies a particular conditional independence assumption.

Sometimes the term Markov blanket\footnote{Named after the Russian mathematician Andrey Andreyevich Markov (1856–1922). J. Pearl, *Probabilistic Reasoning in Intelligent Systems: Networks of Plausible Inference*. Morgan Kaufmann, 1988.} of a node $X$ is used to refer to the minimal set of nodes that, if their values were known, makes $X$ conditionally independent of all other nodes. A Markov blanket of a particular node turns out to consist of its parents, its children, and the other parents of its children.

2.7 Summary

- Representing uncertainty as a probability distribution is motivated by a set of axioms related to the comparison of the plausibility of different statements.

- There are many different families of both discrete and continuous probability distributions.

\footnote{An algorithm for efficiently determining $d$-separation is a bit complicated. See algorithm 3.1 in D. Koller and N. Friedman, *Probabilistic Graphical Models: Principles and Techniques*. MIT Press, 2009.}
If we have $X \rightarrow Y \rightarrow Z$ (chain) or $X \leftarrow Y \rightarrow Z$ (fork) with evidence at $Y$, then $X$ and $Z$ are conditionally independent, meaning that $P(X \mid Y, Z) = P(X \mid Y)$. Interestingly, if the directions of the arrows were slightly different with $X \rightarrow Y \leftarrow Z$ (inverted fork), then $X$ and $Z$ may no longer be conditionally independent given $Y$. In other words, it may be the case that $P(B \mid E) \neq P(B \mid S, E)$. To provide some intuition, consider the inverted fork path from battery failure $B$ to solar panel failure $S$ via electrical system failure $E$. Suppose we know that we have an electrical failure. If we know that we do not have a battery failure, then we are more inclined to believe that we have a solar panel failure because it is an alternative cause of our electrical failure. Conversely, if we found out that we do have a battery failure, then our belief that we have a solar panel failure decreases. This effect is referred to as explaining away. Observing a solar panel failure explains away the cause of the electrical system failure.

Suppose we want to determine whether the network shown in the margin implies that $(D \perp B \mid F)$. There are two undirected paths from $D$ to $B$. We need to check both paths for d-separation.

The path $D \leftarrow A \rightarrow C \leftarrow B$ involves a fork $D \leftarrow A \rightarrow C$ followed by an inverted fork $A \rightarrow C \leftarrow B$. There is no evidence at $A$, so there is no d-separation from the fork. Since $F$ is a descendant of $C$, there is not d-separation along the inverted fork. Hence, there is not d-separation along this path.

The second path, $D \rightarrow E \leftarrow C \leftarrow B$, involves an inverted fork $D \rightarrow E \leftarrow C$ and a chain $E \leftarrow C \leftarrow B$. Since $F$ is a descendant of $E$, there is not d-separation along the inverted fork. Because there is no d-separation along the chain part of this path either, there is no d-separation along this path from $D$ to $B$.

In order for $D$ and $B$ to be conditionally independent given $F$, there must be d-separation along all undirected paths from $D$ to $B$. In this case, neither of the two paths have d-separation. Hence, conditional independence is not implied by the network structure.
• Continuous probability distributions can be represented by density functions.
• Probability distribution families can be combined together in mixtures to result in more flexible distributions.
• Joint distributions are distributions over multiple variables.
• Conditional distributions are distributions over one or more variables given values of evidence variables.
• A Bayesian network is defined by a graphical structure and a set of conditional distributions.
• Depending on the structure of the Bayesian network, we can represent joint distributions with fewer parameters due to conditional independence assumptions.

2.8 Exercises

Exercise 2.1. Consider a continuous random variable $X$ that follows the exponential distribution parameterized by $\lambda$ with density $p(x \mid \lambda) = \lambda \exp(-\lambda x)$ with nonnegative support. Compute the cumulative distribution function of $X$.

Solution: We start with the definition of the cumulative distribution function. Since the support of the distribution is lower-bounded by $x = 0$, there is no probability mass in the interval $(-\infty, 0)$, allowing us to adjust the lower bound of the integral to $0$. After computing the integral, we obtain $\text{cdf}_X(x)$:

\[
\text{cdf}_X(x) = \int_{-\infty}^{x} p(x') \, dx' \\
\text{cdf}_X(x) = \int_{0}^{x} \lambda e^{-\lambda x'} \, dx' \\
\text{cdf}_X(x) = -e^{-\lambda x} \big|_{0}^{x} \\
\text{cdf}_X(x) = 1 - e^{-\lambda x}
\]

Exercise 2.2. For the density function in figure 2.6, what are the five components of the mixture? (There are multiple valid solutions.)

Solution: One solution is $\mathcal{U}([-10, -10], [-5, 10]), \mathcal{U}([-5, 0], [0, 10]), \mathcal{U}([-5, -10], [0, 0]), \mathcal{U}([0, -10], [10, 5])$, and $\mathcal{U}([0, 5], [10, 10])$.

Exercise 2.3. Given the following table representation of $P(X, Y, Z)$, generate an equivalent compact decision tree representation.
Solution: We start with the most common probabilities, 0.13, which occurs when $Z = 0$ and $Y = 0$, and 0.05, which occurs when $Z = 0$ and $Y = 1$. We choose to make $Z$ the root of our decision tree and when $Z = 0$ we continue to a $Y$ node. Based on the value of $Y$ we branch to either 0.13 or 0.05. Next, we continue with cases when $Z = 1$. The most common probabilities are 0.02, which occurs when $Z = 1$ and $X = 0$, and 0.12, which occurs when $Z = 1$ and $X = 2$. So, when $Z = 1$, we choose to continue to an $X$ node. Based on the whether $X$ is 0, 1, or 2, we continue to 0.02, a $Y$ node, or 0.12, respectively. Finally, based on the value of $Y$, we branch to either 0.01 or 0.17.

Exercise 2.4. Suppose we want to specify a multivariate Gaussian mixture model with three components defined over four variables. We require that two of the three Gaussian distributions assume independence between the four variables, while the other Gaussian distribution is defined without any independence assumptions. How many independent parameters are required to specify this mixture model?
**Solution:** For a Gaussian distribution over four variables \( n = 4 \) with independence assumptions, we need to specify \( n + n = 2n = 8 \) independent parameters; there are four parameters for the mean vector and four parameters for the covariance matrix (which is equivalent to the mean and variance parameters of four independent univariate Gaussian distributions). For a Gaussian distribution over four variables without independence assumptions, we need to specify \( n + n(n + 1)/2 = 14 \) independent parameters; there are four parameters for the mean vector and ten parameters for the covariance matrix. Additionally, for our three mixture components \( (k = 3) \), we need to specify \( k - 1 = 2 \) independent parameters for the weights. Thus, we need \( 2(8) + 1(14) + 2 = 32 \) independent parameters to specify this mixture distribution.

**Exercise 2.5.** We have three independent variables \( X_{1:3} \) defined by piecewise-constant densities with 4, 7, and 3 bin edges, respectively. How many independent parameters are required to specify their joint distribution?

**Solution:** If we have a piecewise-constant density with \( m \) bins edges, then there are \( m - 1 \) bins and \( m - 2 \) independent parameters. For this problem, there will be \( (4 - 2) + (7 - 2) + (3 - 2) = 8 \) independent parameters.

**Exercise 2.6.** Suppose we have four continuous random variables, \( X_1, X_2, Y_1, \) and \( Y_2 \), and we want to construct a linear Gaussian model of \( X = X_{1:2} \) given \( Y = Y_{1:2} \), i.e. \( p(X \mid Y) \). How many independent parameters are required for this model?

**Solution:** In this case, our mean vector for the Gaussian distribution is two-dimensional and requires four independent parameters for the transformation matrix \( M \) and two independent parameters for the bias vector \( b \). We also require three independent parameters for the covariance matrix \( \Sigma \). In total, we need \( 4 + 2 + 3 = 9 \) independent parameters to specify this model:

\[
p(x \mid y) = \mathcal{N}(x \mid My + b, \Sigma)
\]

**Exercise 2.7.** Given the following Bayesian network where each node can take on one of four values, how many independent parameters are there? What is the percent reduction in the number of independent parameters required when using the following Bayesian network compared to using a full joint probability table?
Solution: The number of independent parameters for each node is equal to \((k-1)k^m\) where \(k\) is the number of values the node can take on and \(m\) is the number of parents that the node has. The variable \(A\) has 3, \(B\) has 12, \(C\) has 48, \(D\) has 3, \(E\) has 12, and \(F\) has 48 and independent parameters. There are 126 total independent parameters for this Bayesian network.

The number of independent parameters required to specify a joint probability table over \(n\) variables that can take on \(k\) values is equal to \(k^n - 1\). Therefore, specifying a joint probability table would require \(4^6 - 1 = 4096 - 1 = 4095\) independent parameters. The percent reduction in the number of independent parameters required is \((4095 - 126)/4095 \approx 96.9\%\).

Exercise 2.8. Given the following Bayesian network, is \(A\) d-separated from \(E\) given \(C\)?

\[
\begin{align*}
A & \quad \rightarrow & \quad B
D & \quad \rightarrow & \quad C
\end{align*}
\]

Solution: There are two paths from \(A\) to \(E\): \(A \rightarrow D \rightarrow E\) and \(A \rightarrow C \rightarrow E\). There is d-separation along the second path, but not the first. Hence, \(A\) is not d-separated from \(E\) given \(C\).

Exercise 2.9. Given the following Bayesian network, determine the Markov blanket of \(B\).

\[
\begin{align*}
D & \quad \rightarrow & \quad B
\end{align*}
\]

Solution: Paths from \(B\) to \(A\) can only be d-separated given \(A\). Paths from \(B\) to \(D\) can only be d-separated given \(D\). Paths from \(B\) to \(E\) and simultaneously \(F\), \(G\), and \(H\) can be efficiently d-separated given \(E\). Paths from \(B\) to \(C\) are naturally d-separated due to a v-structure; however, since \(E\) must be contained in our Markov blanket, paths from \(B\) to \(C\) given \(E\) can only be d-separated given \(C\). So, the Markov blanket of \(B\) is \(\{A, C, D, E\}\).
Exercise 2.10. In a Bayesian network with structure $A \rightarrow B$, is it possible for $A$ to be independent of $B$?

Solution: There is a direct arrow from $A$ to $B$, which indicates that independence is not implied. However, this does not mean that they are not independent. Whether or not $A$ and $B$ are independent depends on the choice of conditional probability tables. We can choose the tables so that there is independence. For example, suppose both variables are binary and $P(a) = 0.5$ is uniform and $P(b \mid a) = 0.5$. Clearly, $P(A)P(B \mid A) = P(A)P(B)$, which means they are independent.
3 Inference

The previous chapter explained how to represent probability distributions. This chapter will show how to use these probabilistic representations for inference. Inference involves determining the distribution over one or more unobserved variables given the values associated with a set of observed variables. This chapter begins by introducing exact inference methods. Because exact inference can be computationally intractable depending on the structure of the network, we will also discuss several algorithms for approximate inference.

3.1 Inference in Bayesian Networks

In inference problems, we want to infer a distribution over query variables given some observed evidence variables. The other nodes are referred to as hidden variables. We often refer to the distribution over the query variables given the evidence as a posterior distribution.

To illustrate the computations involved in inference, recall the Bayesian network from example 2.5, the structure of which is reproduced in figure 3.1. Suppose we have $B$ as a query variable and evidence $D = 1$ and $C = 1$. The inference task is to compute $P(b^1 \mid d^1, c^1)$, which corresponds to computing the probability that we have a battery failure given an observed trajectory deviation and communication loss.

From the definition of conditional probability introduced in equation (2.22), we know

\[
P(b^1 \mid d^1, c^1) = \frac{P(b^1, d^1, c^1)}{P(d^1, c^1)}
\]  

(3.1)
To compute the numerator, we must use a process known as marginalization, where we sum out variables that are not involved, in this case $S$ and $E$:

$$P(b^1, d^1, c^1) = \sum_s \sum_e P(b^1, s, e, d^1, c^1)$$  (3.2)

We know from the chain rule for Bayesian networks introduced in equation (2.31) that

$$P(b^1, s, e, d^1, c^1) = P(b^1)P(s)P(e \mid b^1, s)P(d^1 \mid e)P(c^1 \mid e)$$  (3.3)

All of the components on the right-hand side are specified in the conditional probability distributions associated with the nodes in the Bayesian network. We can compute the denominator in equation (3.1) using the same approach but with an additional summation over the values for $B$.

This process of using the definition of conditional probability, marginalization, and applying the chain rule can be used to perform exact inference in any Bayesian network. We can implement exact inference using factors. Recall that factors represent discrete multivariate distributions. We use the following three operations on factors to achieve this:

- We use the factor product (algorithm 3.1) to combine two factors to produce a larger factor whose scope is the combined scope of the input factors. If we have $\phi(X, Y)$ and $\psi(Y, Z)$, then $\phi \cdot \psi$ will be over $X, Y,$ and $Z$ with $(\phi \cdot \psi)(x, y, z) = \phi(x, y)\psi(y, z)$. The factor product is demonstrated in example 3.1.

- We use factor marginalization (algorithm 3.2) to sum out a particular variable from the entire factor table, removing it from the resulting scope. Example 3.2 illustrates this process.

- We use factor conditioning (algorithm 3.3) with respect to some evidence to remove any rows in the table inconsistent with that evidence. Example 3.3 demonstrates factor conditioning.

These three factor operations are used together in algorithm 3.4 to perform exact inference. It starts by computing the product of all of the factors, conditioning on the evidence, marginalizing out the hidden variables, and normalizing. One potential issue with this approach is the size of the product of all of the factors. The size of the factor product is equal to the product of the number of values each variable can assume. For the satellite example problem, there are only $2^5 = 32$ possible assignments, but many interesting problems would have a factor product that is too large to practically enumerate.
The factor product of two factors produces a new factor over the union of their variables. Below we produce a new factor from two factors that share a variable.

<table>
<thead>
<tr>
<th>$X$</th>
<th>$Y$</th>
<th>$\phi_1(X,Y)$</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>0</td>
<td>0.3</td>
</tr>
<tr>
<td>0</td>
<td>1</td>
<td>0.4</td>
</tr>
<tr>
<td>1</td>
<td>0</td>
<td>0.2</td>
</tr>
<tr>
<td>1</td>
<td>1</td>
<td>0.1</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>$Y$</th>
<th>$Z$</th>
<th>$\phi_2(Y,Z)$</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>0</td>
<td>0.2</td>
</tr>
<tr>
<td>0</td>
<td>1</td>
<td>0.0</td>
</tr>
<tr>
<td>1</td>
<td>0</td>
<td>0.3</td>
</tr>
<tr>
<td>1</td>
<td>1</td>
<td>0.5</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>$X$</th>
<th>$Y$</th>
<th>$Z$</th>
<th>$\phi_3(X,Y,Z)$</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0.06</td>
</tr>
<tr>
<td>0</td>
<td>0</td>
<td>1</td>
<td>0.00</td>
</tr>
<tr>
<td>0</td>
<td>1</td>
<td>0</td>
<td>0.12</td>
</tr>
<tr>
<td>0</td>
<td>1</td>
<td>1</td>
<td>0.20</td>
</tr>
<tr>
<td>1</td>
<td>0</td>
<td>0</td>
<td>0.04</td>
</tr>
<tr>
<td>1</td>
<td>0</td>
<td>1</td>
<td>0.00</td>
</tr>
<tr>
<td>1</td>
<td>1</td>
<td>0</td>
<td>0.03</td>
</tr>
<tr>
<td>1</td>
<td>1</td>
<td>1</td>
<td>0.05</td>
</tr>
</tbody>
</table>
Recall the joint probability distribution $P(X, Y, Z)$ from table 2.1. We can marginalize out $Y$ by summing the probabilities in each row that have matching assignments for $X$ and $Z$. 

<table>
<thead>
<tr>
<th>$X$</th>
<th>$Y$</th>
<th>$Z$</th>
<th>$\phi(X, Y, Z)$</th>
<th>$X$</th>
<th>$Z$</th>
<th>$\phi(X, Z)$</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0.08</td>
<td>0</td>
<td>0</td>
<td>0.17</td>
</tr>
<tr>
<td>0</td>
<td>0</td>
<td>1</td>
<td>0.31</td>
<td>0</td>
<td>1</td>
<td>0.68</td>
</tr>
<tr>
<td>0</td>
<td>1</td>
<td>0</td>
<td>0.09</td>
<td>1</td>
<td>0</td>
<td>0.03</td>
</tr>
<tr>
<td>0</td>
<td>1</td>
<td>1</td>
<td>0.37</td>
<td>1</td>
<td>1</td>
<td>0.12</td>
</tr>
<tr>
<td>1</td>
<td>0</td>
<td>0</td>
<td>0.01</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1</td>
<td>0</td>
<td>1</td>
<td>0.05</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1</td>
<td>1</td>
<td>0</td>
<td>0.02</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1</td>
<td>1</td>
<td>1</td>
<td>0.07</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Example 3.2. A demonstration of factor marginalization.
in_scope(name, φ) = any(name == v.name for v in φ.vars)

function condition(ϕ::Factor, name, value)
    if !in_scope(name, ϕ)
        return ϕ
    end
    table = FactorTable()
    for (a, p) in ϕ.table
        if a[name] == value
            table[delete!(copy(a), name)] = p
        end
    end
    vars = filter(v → v.name != name, ϕ.vars)
    return Factor(vars, table)
end

function condition(ϕ::Factor, evidence)
    for (name, value) in pairs(evidence)
        ϕ = condition(ϕ, name, value)
    end
    return ϕ
end

Algorithm 3.3. Two methods for factor conditioning given some evidence. The first takes a factor ϕ and returns a new factor whose table entries are consistent with the variable named name having value value. The second takes a factor ϕ and applies evidence in the form of a named tuple. The in_scope method returns true if a variable named name is within the scope of the factor ϕ.

Example 3.3. An illustration of setting evidence, in this case for Y, in a factor. The resulting values must be renormalized.
Algorithm 3.4. A naive exact inference algorithm for a discrete Bayesian network $bn$, which takes as input a set of query variable names $query$, and $evidence$ associating values with observed variables. The algorithm computes a joint distribution over the query variables in the form of a factor. We introduce the ExactInference type to allow for $infer$ to be called with different inference methods, as shall be seen in the rest of this chapter.

3.2 Inference in Naive Bayes Models

The previous section presented a general method for performing exact inference in any Bayesian network. This section discusses how this same method can be used to solve classification problems for a special kind of Bayesian network structure known as a naive Bayes model. This structure is shown in figure 3.2. An equivalent but more compact representation is shown in figure 3.3 using a plate, shown as a rounded box. The $i = 1 : n$ in the bottom of the box specifies that the $i$ in the subscript of the variable name is repeated from 1 to $n$.

In the naive Bayes model, the class $C$ is the query variable, and the observed features $O_{1:n}$ are the evidence variables. The naive Bayes model is called naive because it assumes conditional independence between the evidence variables given the class. Using the notation introduced in section 2.6, we can say $(O_i \perp O_j \mid C)$ for all $i \neq j$. Of course, if these conditional independence assumptions do not hold, then we can add the necessary directed edges between the observed features.

We have to specify the prior $P(C)$ and the class-conditional distributions $P(O_i \mid C)$. As done in the previous section, we can apply the chain rule to compute the joint distribution:

$$P(c, o_{1:n}) = P(c) \prod_{i=1}^{n} P(o_i \mid c)$$

(3.4)

Our classification task involves computing the conditional probability $P(c \mid o_{1:n})$. From the definition of conditional probability, we have

$$P(c \mid o_{1:n}) = \frac{P(c, o_{1:n})}{P(o_{1:n})}$$

(3.5)
We can compute the denominator by marginalizing the joint distribution:

\[ P(o_{1:n}) = \sum_c P(c, o_{1:n}) \]  

(3.6)

The denominator in equation (3.5) is not a function of \( C \) and can therefore be treated as a constant. Hence, we can write

\[ P(c \mid o_{1:n}) = \kappa P(c, o_{1:n}) \]  

(3.7)

where \( \kappa \) is a normalization constant such that \( \sum_c P(c \mid o_{1:n}) = 1 \). We often drop \( \kappa \) and write

\[ P(c \mid o_{1:n}) \propto P(c, o_{1:n}) \]  

(3.8)

where the proportional to symbol \( \propto \) is used to represent that the left-hand side is proportional to the right-hand side. Example 3.4 illustrates how inference can be applied to classifying radar tracks.

We can use this method to infer a distribution over classes, but for many applications, we have to commit to a particular class. It is common to classify according to the class with the highest posterior probability, \( \arg \max_c P(c \mid o_{1:n}) \). However, choosing a class is really a decision problem that often should take into account the consequences of misclassification. For example, if we are interested in using our classifier to filter out targets that are not aircraft for the purpose of air traffic control, then we can afford to occasionally let a few birds and other clutter tracks through our filter. However, we would want to avoid filtering out any real aircraft because that could lead to a collision. In this case, we would probably only want to classify a track as a bird if the posterior probability were close to 1. Decision problems will be discussed in chapter 6.

### 3.3 Sum-Product Variable Elimination

A variety of methods can be used to perform efficient inference in more complicated Bayesian networks. One method is known as sum-product variable elimination, which interleaves eliminating hidden variables (summations) with applications of the chain rule (products). It is more efficient to marginalize variables out as early as possible to avoid generating large factors.
Suppose we have a radar track and we want to determine whether it was generated by a bird or an aircraft. We base our inference on airspeed and the amount of heading fluctuation. The prior represents our belief about whether a target is a bird or an aircraft in the absence of any information about the track. Below are example class-conditional distributions for airspeed $v$ as estimated from radar data.

![Graph showing class-conditional distributions for airspeed $v$.]

Suppose from the chain rule we determine:

\[
\begin{align*}
P(\text{bird, slow, little heading fluctuation}) &= 0.03 \\
P(\text{aircraft, slow, little heading fluctuation}) &= 0.01
\end{align*}
\]

Of course, these probabilities do not sum to 1. If we want to determine the probability that a target is a bird given the evidence, then we would make the following calculation:

\[
P(\text{bird} \mid \text{slow, little heading fluctuation}) = \frac{0.03}{0.03 + 0.01} = 0.75
\]
We will illustrate the variable elimination algorithm by computing the distribution \( P(B \mid d^1, c^1) \) for the Bayesian network in figure 3.1. The conditional probability distributions associated with the nodes in the network can be represented by the following factors:

\[
\phi_1(B), \phi_2(S), \phi_3(E, B, S), \phi_4(D, E), \phi_5(C, E)
\] (3.9)

Because \( D \) and \( C \) are observed variables, the last two factors can be replaced with \( \phi_6(E) \) and \( \phi_7(E) \) by setting the evidence \( D = 1 \) and \( C = 1 \).

We then proceed by eliminating the hidden variables in sequence. Different strategies can be used for choosing an ordering, but for this example, we arbitrarily choose the ordering \( E \) and then \( S \). To eliminate \( E \), we take the product of all the factors involving \( E \) and then marginalize out \( E \) to get a new factor:

\[
\phi_8(B, S) = \sum_e \phi_3(e, B, S) \phi_6(e) \phi_7(e)
\] (3.10)

We can now discard \( \phi_3, \phi_6, \) and \( \phi_7 \) because all the information we need from them is contained in \( \phi_8 \).

Next, we eliminate \( S \). Again, we gather all remaining factors that involve \( S \) and marginalize out \( S \) from the product of these factors:

\[
\phi_9(B) = \sum_s \phi_2(s) \phi_8(B, s)
\] (3.11)

We discard \( \phi_2 \) and \( \phi_8 \), and are left with \( \phi_1(B) \) and \( \phi_9(B) \). Finally, we take the product of these two factors and normalize the result to obtain a factor representing \( P(B \mid d^1, c^1) \).

The above procedure is equivalent to computing the following:

\[
P(B \mid d^1, c^1) \propto \phi_1(B) \sum_s \left( \phi_2(s) \sum_e \left( \phi_3(e \mid B, s) \phi_4(d^1 \mid e) \phi_5(c^1 \mid e) \right) \right)
\] (3.12)

This produces the same result as, but is more efficient than, the naive procedure of taking the product of all of the factors and then marginalizing:

\[
P(B \mid d^1, c^1) \propto \sum_s \sum_e \phi_1(B) \phi_2(s) \phi_3(e \mid B, s) \phi_4(d^1 \mid e) \phi_5(c^1 \mid e)
\] (3.13)
The sum-product variable elimination algorithm is implemented in algorithm 3.5. It takes as input a Bayesian network, a set of query variables, a list of observed values, and an ordering of the variables. We first set all observed values. Then, for each variable, we multiply together all factors containing it and then marginalize that variable out. This new factor replaces the consumed factors and we repeat the process for the next variable.

For many networks, variable elimination allows inference to be done in an amount of time that scales linearly with the size of the network, but it has exponential time complexity in the worst case. What influences the amount of computation is the variable elimination order. Choosing the optimal elimination order is NP-hard, meaning that it cannot be done in polynomial time in the worst case (section 3.5). Even if we found the optimal elimination order, variable elimination can still require an exponential number of computations. Variable elimination heuristics generally try to minimize the number of variables involved in the intermediate factors generated by the algorithm.

```
struct VariableElimination
    ordering # array of variable indices
end

function infer(M::VariableElimination, bn, query, evidence)
    Φ = [condition(ϕ, evidence) for ϕ in bn.factors]
    for i in M.ordering
        name = bn.vars[i].name
        if name ∉ query
            inds = findall(ϕ→in_scope(name, ϕ), Φ)
            if !isempty(inds)
                ϕ = prod(Φ[inds])
                deleteat!(Φ, inds)
                ϕ = marginalize(ϕ, name)
                push!(Φ, ϕ)
            end
        end
    end
    return normalize!(prod(Φ))
end
```

3.4 Belief Propagation

An approach to inference known as belief propagation works by propagating “messages” through the network using the sum-product algorithm in order to compute the marginal distributions of the query variables. Belief propagation requires linear time but only provides an exact answer if the network does not have undirected cycles. If the network has undirected cycles, then it can be converted into a tree by combining multiple variables into single nodes by using what is known as the junction tree algorithm. If the number of variables that have to be combined into any one node in the resulting network is small, then inference can be done efficiently. A variation of belief propagation known as loopy belief propagation can provide approximate solutions in networks with undirected cycles. Although this approach does not provide any guarantees and may not converge, it can work well in practice.

3.5 Computational Complexity

We can show that inference in Bayesian networks is NP-hard by using an NP-complete problem called 3SAT. It is easy to construct a Bayesian network from an arbitrary 3SAT problem. For example, consider the following 3SAT formula:

\[
F(x_1, x_2, x_3, x_4) = (x_1 \lor x_2 \lor x_3) \land (\neg x_1 \lor \neg x_2 \lor x_3) \land (x_2 \lor \neg x_3 \lor x_4)
\]

(3.14)

where \(\neg\) represents logical negation (“not”), \(\land\) represents logical conjunction (“and”), and \(\lor\) represents logical disjunction (“or”). The formula consists of a conjunction of clauses, which are disjunctions of what are called literals. A literal is simply a variable or its negation.

Figure 3.4 shows the corresponding Bayesian network representation. The variables are represented by \(X_{1:4}\), and the clauses are represented by \(C_{1:3}\). The distributions over the variables are uniform. The nodes representing clauses have as parents the participating variables. Because this is a 3SAT problem, each clause node has exactly three parents. Each clause node assigns probability 0 to assignments that do not satisfy the clause and probability 1 to all satisfying assignments. The remaining nodes assign probability 1 to true if all their parents

---


5 This formula also appears in example C.3 in appendix C.
are true. The original problem is satisfiable if and only if $P(y^1) > 0$. Hence, inference in Bayesian networks is at least as hard as 3SAT.

![Figure 3.4. Bayesian network representing a 3SAT problem.](image)

The reason we go to the effort of showing that inference in Bayesian networks is NP-hard is so that we know to avoid wasting time looking for an efficient, exact inference algorithm that works on all Bayesian networks. Therefore, research over the past couple decades has focused on approximate inference methods, which are discussed next.

### 3.6 Direct Sampling

Motivated by the fact that exact inference is computationally intractable, many approximation methods have been developed. One of the simplest methods for inference is based on direct sampling, where random samples from the joint distribution are used to arrive at a probability estimate. To illustrate, suppose we want to infer $P(b^1 \mid d^1, c^1)$ from a set of $n$ samples from the joint distribution $P(b, s, e, d, c)$. We use parenthetical superscripts to indicate the index of a sample, where we write $(b^{(i)}, s^{(i)}, e^{(i)}, d^{(i)}, c^{(i)})$ for the $i$th sample. The direct sample estimate is

$$P(b^1 \mid d^1, c^1) \approx \frac{\sum_i (b^{(i)} = 1 \land d^{(i)} = 1 \land c^{(i)} = 1)}{\sum_i (d^{(i)} = 1 \land c^{(i)} = 1)}$$

(3.15)

We use the convention where a logical statement in parentheses is treated numerically as 1 when true and 0 when false. The numerator is the number of samples consistent with $b$, $d$, and $c$ all set to 1, and the denominator is the number of samples consistent with $d$ and $c$ all set to 1.

---

6 Sometimes approaches involving random sampling are referred to as Monte Carlo methods. The name comes from the Monte Carlo Casino in Monaco. An introduction to randomized algorithms and their application to a variety of problem domains is provided by R. Motwani and P. Raghavan, *Randomized Algorithms*. Cambridge University Press, 1995.
Sampling from the joint distribution represented by a Bayesian network is straightforward. The first step involves finding a topological sort of the nodes in the Bayesian network. A topological sort of nodes in a directed acyclic graph is an ordered list such that if there is an edge $A \rightarrow B$, then $A$ comes before $B$ in the list. For example, a topological sort for the network in figure 3.1 is $B, S, E, D, C$. A topological sort always exists, but it may not be unique. Another topological sort for the network is $S, B, E, C, D$.

Once we have a topological sort, we can begin sampling from the conditional probability distributions. Algorithm 3.6 shows how to sample from a Bayesian network given an ordering $X_{1:n}$ that represents a topological sort. We draw a sample from the conditional distribution associated with $X_i$ given the values of the parents that have already been assigned. Because $X_{1:n}$ is a topological sort, we know that all the parents of $X_i$ have already been instantiated, allowing this sampling to be done. Direct sampling is implemented in algorithm 3.7 and is demonstrated in example 3.5.

Algorithm 3.6. A method for sampling an assignment from a Bayesian network $bn$. We also provide a method for sampling an assignment from a factor $\phi$. 

```plaintext
function Base.rand(\phi::Factor)
    tot, p, w = 0.0, rand(), sum(values(\phi.table))
    for (a,v) in \phi.table
        tot += v/w
        if tot >= p
            return a
        end
    end
    return Assignment()
end

function Base.rand(bn::BayesianNetwork)
    a = Assignment()
    for i in topological_sort(bn.graph)
        name, \phi = bn.vars[i].name, bn.factors[i]
        a[name] = rand(condition(\phi, a))[name]
    end
    return a
end
```
Suppose we draw ten random samples from the network in figure 3.1. We are interested in inferring \( P(b_1 \mid d_1, c_1) \). Only two of the ten samples (pointed to in the table) are consistent with the observations \( d_1 \) and \( c_1 \). One sample has \( b = 1 \) and the other sample has \( b = 0 \). From these samples, we infer that \( P(b_1 \mid d_1, c_1) = 0.5 \). Of course, we would want to use more than just two samples to accurately estimate \( P(b_1 \mid d_1, c_1) \).

<table>
<thead>
<tr>
<th>( B )</th>
<th>( S )</th>
<th>( E )</th>
<th>( D )</th>
<th>( C )</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>0</td>
<td>1</td>
<td>1</td>
<td>0</td>
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<td>0</td>
<td>0</td>
<td>1</td>
<td>0</td>
</tr>
</tbody>
</table>

Example 3.5. An example of how direct samples from a Bayesian network can be used for inference.

Algorithm 3.7. The direct sampling inference method, which takes a Bayesian network \( bn \), a list of query variables \( query \), and evidence \( evidence \). The method draws \( m \) samples from the Bayesian network and retains those samples that are consistent with the evidence. A factor over the query variables is returned. This method can fail if no samples that satisfy the evidence are found.
3.7 Likelihood Weighted Sampling

The problem with direct sampling is that we may waste time generating samples that are inconsistent with the observations, especially if the observations are unlikely. An alternative approach is called likelihood weighted sampling, which involves generating weighted samples that are consistent with the observations.

To illustrate, we will again attempt to infer $P(b^1 | d^1, c^1)$. We have a set of $n$ samples, where the $i$th sample is again denoted $(b^{(i)}, s^{(i)}, e^{(i)}, d^{(i)}, c^{(i)})$. The weight of the $i$th sample is $w_i$. The weighted estimate is

$$P(b^1 | d^1, c^1) \approx \frac{\sum_i w_i (b^{(i)} = 1 \land d^{(i)} = 1 \land c^{(i)} = 1)}{\sum_i w_i (d^{(i)} = 1 \land c^{(i)} = 1)} \tag{3.16}$$

$$= \frac{\sum_i w_i (b^{(i)} = 1)}{\sum_i w_i} \tag{3.17}$$

To generate these weighted samples, we begin with a topological sort and sample from the conditional distributions in sequence. The only difference in likelihood weighting is how we handle observed variables. Instead of sampling their values from a conditional distribution, we assign variables to their observed values and adjust the weight of the sample appropriately. The weight of a sample is simply the product of the conditional probabilities at the observed nodes. Likelihood weighted sampling is implemented in algorithm 3.8. Example 3.6 demonstrates inference with likelihood weighted sampling.
struct LikelihoodWeightedSampling
  m # number of samples
end

function infer(M::LikelihoodWeightedSampling, bn, query, evidence)
  table = FactorTable()
  ordering = topological_sort(bn.graph)
  for i in 1:(M.m)
    a, w = Assignment(), 1.0
    for j in ordering
      name, ϕ = bn.vars[j].name, bn.factors[j]
      if haskey(evidence, name)
        a[name] = evidence[name]
        w *= ϕ.table[select(a, variablenames(ϕ))]
      else
        a[name] = rand(condition(ϕ, a))[name]
      end
    end
    b = select(a, query)
    table[b] = get(table, b, 0) + w
  end
  vars = filter(v→v.name ∈ query, bn.vars)
  return normalize!(Factor(vars, table))
end

Algorithm 3.8. The likelihood sampling inference method, which takes a Bayesian network bn, a list of query variables query, and evidence evidence. The method draws m samples from the Bayesian network but sets values from evidence when possible, keeping track of the conditional probability when doing so. These probabilities are used to weight the samples such that the final inference estimate is accurate. A factor over the query variables is returned.
The table below shows five likelihood-weighted samples from the network in figure 3.1. We sample from $P(B)$, $P(S)$, and $P(E \mid B, S)$, as we would with direct sampling. When we come to $D$ and $C$, we assign $D = 1$ and $C = 1$. If the sample has $E = 1$, then the weight is $P(d^1 \mid e^1)P(c^1 \mid e^1)$; otherwise, the weight is $P(d^1 \mid e^0)P(c^1 \mid e^0)$. If we assume

$$P(d^1 \mid e^1)P(c^1 \mid e^1) = 0.95$$
$$P(d^1 \mid e^0)P(c^1 \mid e^0) = 0.01$$

then we may approximate from the samples in the table below

$$P(b^1 \mid d^1, c^1) = \frac{0.95}{0.95 + 0.95 + 0.01 + 0.01 + 0.95} \approx 0.331$$

<table>
<thead>
<tr>
<th></th>
<th>B</th>
<th>S</th>
<th>E</th>
<th>D</th>
<th>C</th>
<th>Weight</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>0</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>$P(d^1 \mid e^1)P(c^1 \mid e^1)$</td>
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<td>0</td>
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<td>$P(d^1 \mid e^1)P(c^1 \mid e^1)$</td>
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<tr>
<td>0</td>
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<td>1</td>
<td>$P(d^1 \mid e^0)P(c^1 \mid e^0)$</td>
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<td>1</td>
<td>$P(d^1 \mid e^0)P(c^1 \mid e^0)$</td>
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<td>0</td>
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<td>1</td>
<td>1</td>
<td>1</td>
<td>$P(d^1 \mid e^1)P(c^1 \mid e^1)$</td>
</tr>
</tbody>
</table>
Although likelihood weighting makes it so that all samples are consistent with the observations, it can still be wasteful. Consider the simple chemical detection Bayesian network shown in figure 3.5, and assume that we detected a chemical of interest. We want to infer \( P(c^1 \mid d^1) \). Because this network is small, we can easily compute this probability exactly by using Bayes’ rule:

\[
P(c^1 \mid d^1) = \frac{P(d^1 \mid c^1)P(c^1)}{P(d^1 \mid c^1)P(c^1) + P(d^1 \mid c^0)P(c^0)} = \frac{0.999 \times 0.001}{0.999 \times 0.001 + 0.001 \times 0.999} = 0.5
\]

If we use likelihood weighting, then 99.9% of the samples will have \( C = 0 \) with a weight of 0.001. Until we get a sample of \( C = 1 \), which has an associated weight of 0.999, our estimate of \( P(c^1 \mid d^1) \) will be 0.

### 3.8 Gibbs Sampling

An alternative approach to inference is to use Gibbs sampling, 8 which is a kind of Markov chain Monte Carlo technique. Gibbs sampling involves drawing samples consistent with the evidence in a way that does not involve weighting. From these samples, we can infer the distribution over the query variables.

Gibbs sampling involves generating a sequence of samples, starting with an initial sample \( x_{1:n}^{(1)} \) generated randomly with the evidence variables set to their observed values. The \( k \)th sample \( x_{1:n}^{(k)} \) depends probabilistically on the previous sample \( x_{1:n}^{(k-1)} \). We modify \( x_{1:n}^{(k-1)} \) in-place to obtain \( x_{1:n}^{(k)} \) as follows. Using any ordering of the unobserved variables, which need not be a topological sort, we assign \( X_i^{(k)} \) a value from the distribution represented by \( P(X_i \mid x_{-i}^{(k)}) \). Here, \( x_{-i}^{(k)} \) represents the values of all other variables except \( X_i \) in sample \( k \). Sampling from \( P(X_i \mid x_{-i}^{(k)}) \) can be done efficiently because we only need to consider the Markov blanket of variable \( X_i \) (section 2.6).

Unlike the other sampling methods discussed so far, the samples produced by this method are not independent. However, it can be proven that, in the limit, samples are drawn exactly from the joint distribution over the unobserved

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2021-03-07 23:12:00-08:00, revision 80bf4d6, comments to bugs@algorithmsbook.com
variables given the observations. Algorithm 3.9 shows how to compute a factor for $P(X_i \mid x_{-i})$. Gibbs sampling is implemented in algorithm 3.10.

```plaintext
function blanket(bn, a, i)
  name = bn.vars[i].name
  val = a[name]
  a = delete!(copy(a), name)
  Φ = filter(ϕ -> in_scope(name, ϕ), bn.factors)
  ϕ = prod(condition(ϕ, a) for ϕ in Φ)
  return normalize!(ϕ)
end
```

Algorithm 3.9. A method for obtaining $P(X_i \mid x_{-i})$ for a Bayesian network $bn$ given a current assignment $a$.

Gibbs sampling can be applied to our running example. We can use our $m$ samples to estimate

$$P(b^1 \mid d^1, c^1) \approx \frac{1}{m} \sum_i (b^{i(i)} = 1)$$

(3.21)

Figure 3.6 compares the convergence of the estimate of $P(c^1 \mid d^1)$ in the chemical detection network using direct, likelihood weighted, and Gibbs sampling. Direct sampling takes the longest to converge. The direct sampling curve has long periods during which the estimate does not change because samples are inconsistent with the observations. Likelihood-weighted sampling converges faster in this example. Spikes occur when a sample is generated with $C = 1$ and then gradually decrease. Gibbs sampling, in this example, quickly converges to the true value of 0.5.

As mentioned earlier, Gibbs sampling, like other Markov chain Monte Carlo methods, produces samples from the desired distribution in the limit. In practice, we have to run Gibbs for some amount of time, called the burn-in period, before converging to a steady state distribution. The samples produced during burn-in are normally discarded. If many samples are to be used from a single Gibbs sampling series, it is common to thin the samples by only keeping every $h$th sample because of potential correlation between samples.
function update_gibbs_sample(a, bn, evidence, ordering)
    for i in ordering
        name = bn.vars[i].name
        if !haskey(evidence, name)
            b = blanket(bn, a, i)
            a[name] = rand(b)[name]
        end
    end
    return a
end

function gibbs_sample(a, bn, evidence, ordering, m)
    for j in 1:m
        a = update_gibbs_sample(a, bn, evidence, ordering)
    end
    return a
end

struct GibbsSampling
    m_samples # number of samples to use
    m_burnin # number of samples to discard during burn-in
    m_skip   # number of samples to skip for thinning
    ordering # array of variable indices
end

function infer(M::GibbsSampling, bn, query, evidence)
    table = FactorTable()
    a = merge(rand(bn), evidence)
    a = gibbs_sample(a, bn, evidence, M.ordering, M.m_burnin)
    for i in 1:(M.m_samples)
        a = gibbs_sample(a, bn, evidence, M.ordering, M.m_skip)
        b = select(a, query)
        table[b] = get(table, b, 0) + 1
    end
    vars = filter(v->v.name ∈ query, bn.vars)
    return normalize!(Factor(vars, table))
end

Algorithm 3.10. Gibbs sampling implemented for a Bayesian network bn with evidence evidence and an ordering ordering. The method iteratively updates the assignment a for m iterations.
3.9 Inference in Gaussian Models

If the joint distribution is Gaussian, we can perform exact inference analytically. Two jointly Gaussian random variables \(a\) and \(b\) can be written

\[
\begin{bmatrix} a \\ b \end{bmatrix} \sim \mathcal{N}\left(\begin{bmatrix} \mu_a \\ \mu_b \end{bmatrix}, \begin{bmatrix} A & C \\ C^\top & B \end{bmatrix}\right)
\]  

(3.22)

The marginal distribution of a multivariate Gaussian is also Gaussian:

\[
a \sim \mathcal{N}(\mu_a, A) \quad b \sim \mathcal{N}(\mu_b, B)
\]  

(3.23)

The conditional distribution of a multivariate Gaussian is also Gaussian with a convenient closed-form solution:

\[
p(a \mid b) = \mathcal{N}\left(a \mid \mu_{a\mid b}, \Sigma_{a\mid b}\right)
\]  

(3.24)

\[
\mu_{a\mid b} = \mu_a + CB^{-1}(b - \mu_b)
\]  

(3.25)

\[
\Sigma_{a\mid b} = A - CB^{-1}C^\top
\]  

(3.26)

Algorithm 3.11 shows how to use these equations to infer a distribution over a set of query variables given evidence. Example 3.7 illustrates how to extract the marginal and conditional distributions from a multivariate Gaussian.
Algorithm 3.11. Inference in a multivariate Gaussian distribution $D$. A vector of integers specifies the query variables in the `query` argument, and a vector of integers specifies the evidence variables in the `evidencevars` argument. The values of the evidence variables are contained in the vector `evidence`. The `Distributions.jl` package defines the `MvNormal` distribution.

Example 3.7. Marginal and conditional distributions for a multivariate Gaussian.

For example, consider

$$
\begin{bmatrix}
x_1 \\
x_2
\end{bmatrix} \sim \mathcal{N}\left(\begin{bmatrix} 0 \\
1
\end{bmatrix}, \begin{bmatrix} 3 & 1 \\
1 & 2
\end{bmatrix}\right)
$$

The marginal distribution for $x_1$ is $\mathcal{N}(0, 3)$, and the marginal distribution for $x_2$ is $\mathcal{N}(1, 2)$.

The conditional distribution for $x_1$ given $x_2 = 2$ is

$$
\mu_{x_1|x_2=2} = 0 + 1 \cdot 2^{-1} \cdot (2 - 1) = 0.5
$$

$$
\Sigma_{x_1|x_2=2} = 3 - 1 \cdot 2^{-1} \cdot 1 = 2.5
$$

$$
x_1 \mid (x_2 = 2) \sim \mathcal{N}(0.5, 2.5)
$$

We can perform this inference calculation using algorithm 3.11 by constructing the joint distribution

$D = \text{MvNormal}([0.0, 1.0], [3.0 1.0; 1.0 2.0])$

and then calling `infer(D, [1], [2], [2.0]).`
3.10 Summary

- Inference involves determining the probability of query variables given some evidence.

- Exact inference can be done by computing the joint distribution over the variables, setting evidence, and marginalizing out any hidden variables.

- Inference can be done efficiently in naive Bayes models, in which a single parent variable affects many conditionally independent children.

- The variable elimination algorithm can make exact inference more efficient by marginalizing out variables in sequence.

- Belief propagation represents another method for inference, in which information is iteratively passed between factors to arrive at a result.

- Inference in a Bayesian network can be shown to be NP-hard through a reduction to the 3SAT problem, motivating the development of approximate inference methods.

- Approximate inference can be done by directly sampling from the joint distribution represented by a Bayesian network, but it may involve discarding many samples that are inconsistent with the evidence.

- Likelihood weighted sampling can reduce computation required for approximate inference by only generating samples that are consistent with the evidence and weighting each sample accordingly.

- Gibbs sampling generates a series of unweighted samples that are consistent with the evidence and can greatly speed approximate inference.

- Exact inference can be done efficiently through matrix operations when the joint distribution is Gaussian.

3.11 Exercises

Exercise 3.1. Suppose Anna is shooting basketball free throws. Before we see her play, we start with an independent uniform prior over the probability she successfully makes a basket per shot. We observe her make three shots with two of them resulting in successful baskets. What is the probability that we assign to her making the next basket?
Solution: We denote the probability of making a basket $\theta$. Since we start with a uniform prior $\text{Beta}(1, 1)$ and observe two baskets and one miss, our posterior is then $\text{Beta}(1 + 2, 1 + 1) = \text{Beta}(3, 2)$. We want to compute the probability of a basket:

$$P(\text{basket}) = \int P(\text{basket} | \theta) \text{Beta}(\theta | 3, 2) \, d\theta = \int \theta \text{Beta}(\theta | 3, 2) \, d\theta$$

The expression above is just the expectation (or mean) of a beta distribution, which gives us $P(\text{basket}) = \frac{3}{5}$.

Exercise 3.2. Given the following Bayesian network and its associated conditional probability distributions, write the equation required to perform exact inference for the query $P(a^1 | d^1)$.

Solution: We first expand the inference expression using the definition of conditional probability.

$$P(a^1 | d^1) = \frac{P(a^1, d^1)}{P(d^1)}$$

We can rewrite the numerator as a marginalization over the hidden variables and we can rewrite the denominator as a marginalization over both the hidden and query variables.

$$P(a^1 | d^1) = \frac{\sum_b \sum_c P(a^1, b, c, d^1)}{\sum_a \sum_b \sum_c P(a, b, c, d^1)}$$

The definition of the joint probability in both the numerator and the denominator can be rewritten using the chain rule for Bayesian networks and the resulting equation can be simplified by removing constants from inside the summations.

$$P(a^1 | d^1) = \frac{\sum_b \sum_c P(a^1) P(b | a^1) P(c | b) P(d^1 | c)}{\sum_a \sum_b \sum_c P(a) P(b | a) P(c | b) P(d^1 | c)} = \frac{P(a^1) \sum_b \sum_c P(b | a^1) P(c | b) P(d^1 | c)}{\sum_a \sum_b \sum_c P(a) P(b | a) P(c | b) P(d^1 | c)} = \frac{P(a^1) \sum_b P(b | a^1) \sum_c P(c | b) P(d^1 | c)}{\sum_a P(a) \sum_b P(b | a) \sum_c P(c | b) P(d^1 | c)}$$

Exercise 3.3. Given the following Bayesian network and its associated conditional probability distributions, write the equation required to perform exact inference for the query $P(c^1, d^1 | a^0, f^1)$.

Solution: We first expand the inference expression using the definition of conditional probability.

$$P(c^1, d^1 | a^0, f^1) = \frac{P(c^1, d^1, a^0, f^1)}{P(a^0, f^1)}$$

We can rewrite the numerator as a marginalization over the hidden variables and we can rewrite the denominator as a marginalization over both the hidden and query variables.

$$P(c^1, d^1 | a^0, f^1) = \frac{\sum_b \sum_c \sum_d P(c^1, d^1, a^0, f^1)}{\sum_a \sum_b \sum_c \sum_d P(a, b, c, d, a^0, f^1)}$$

The definition of the joint probability in both the numerator and the denominator can be rewritten using the chain rule for Bayesian networks and the resulting equation can be simplified by removing constants from inside the summations.

$$P(c^1, d^1 | a^0, f^1) = \frac{\sum_b \sum_c \sum_d P(c^1) P(d^1) P(a^0 | c^1) P(f^1 | c^1, d^1)}{\sum_a \sum_b \sum_c \sum_d P(a) P(b | a) P(c | b) P(d | c) P(a^0 | c^1) P(f^1 | c^1, d^1)} = \frac{P(c^1) \sum_b \sum_c \sum_d P(d^1) P(a^0 | c^1) P(f^1 | c^1, d^1)}{\sum_a \sum_b \sum_c \sum_d P(a) P(b | a) P(c | b) P(d | c) P(a^0 | c^1) P(f^1 | c^1, d^1)} = \frac{P(c^1) \sum_b \sum_c \sum_d P(d^1 | c^1) \sum_f P(f^1 | c^1, d^1)}{\sum_a \sum_b \sum_c \sum_d P(a) \sum_b \sum_c \sum_d P(b | a) \sum_c \sum_d P(c | b) \sum_d P(d | c) P(a^0 | c^1) P(f^1 | c^1, d^1)}$$
We can rewrite the numerator as a marginalization over the hidden variables and we can rewrite the denominator as a marginalization over both the hidden and query variables.

\[
P(c^1, d^1 | a^0, f^1) = \frac{\sum_b \sum_c P(a^0, b, c^1, d^1, e, f^1)}{\sum_b \sum_c \sum_d \sum_e P(a^0, b, c, d, e, f^1)}
\]

The definition of the joint probability in both the numerator and the denominator can be rewritten using the chain rule for Bayesian networks and the resulting equation can be simplified by removing constants from inside the summations. Note, there are multiple possible orderings of the summations in the final equation.

\[
P(c^1, d^1 | a^0, f^1) = \frac{\sum_b \sum_c P(a^0)P(b | a^0, c^1)P(c^1)P(d^1 | a^0)P(e | b, c^1, d^1)P(f^1 | e)}{\sum_b \sum_c \sum_d \sum_e P(a^0)P(b | a^0, c)P(c)P(d | a^0)P(e | b, c, d)P(f^1 | e)}
\]

**Exercise 3.4.** Suppose we are developing an object detection system for an autonomous vehicle driving in a city. Our autonomous vehicle’s perception system reports an object’s size \(S\) (either small, medium, or large) and an object’s speed \(V\) (either slow, moderate, or fast). We want to design a model that will determine the class \(C\) of an object—either a vehicle, pedestrian, or a ball—given observations of the object’s size and speed. Assuming a naive Bayes model with the following class prior and class-conditional distributions, what is the detected class given observations \(S = \text{medium}\) and \(V = \text{slow}\)?

| \(C\)  | \(P(C)\) | \(C\)  | \(S\)    | \(P(S | C)\) | \(C\)  | \(V\)    | \(P(V | C)\) |
|-------|-----------|-------|---------|-------------|-------|---------|-------------|
| vehicle | 0.80     | vehicle | small  | 0.001       | vehicle | slow    | 0.2         |
| pedestrian | 0.19   | vehicle | medium | 0.009       | vehicle | moderate | 0.2         |
| ball     | 0.01     | vehicle | large  | 0.990       | vehicle | fast     | 0.6         |
| pedestrian | 0.200  | pedestrian | small | 0.200       | pedestrian | slow    | 0.5         |
| pedestrian | 0.750  | pedestrian | medium | 0.050       | pedestrian | moderate | 0.4         |
| pedestrian | 0.800  | pedestrian | large  | 0.019       | pedestrian | fast     | 0.1         |
| ball     | 0.199    | ball    | small  | 0.800       | ball     | slow     | 0.4         |
| ball     | 0.001    | ball    | medium |             | ball     | moderate | 0.4         |
| ball     |          | ball    | large  |             | ball     | fast     | 0.2         |

**Solution:** To compute the posterior distribution \(P(c | o_{1:n})\), we use the definition of the joint distribution for a naive Bayes model in equation (3.4):
\[ P(c \mid o_{1:n}) \propto P(c) \prod_{i=1}^{n} P(o_i \mid c) \]

\[ P(\text{vehicle} \mid \text{medium, slow}) \propto P(\text{vehicle}) P(S = \text{medium} \mid \text{vehicle}) P(V = \text{slow} \mid \text{vehicle}) \]
\[ P(\text{vehicle} \mid \text{medium, slow}) \propto (0.80)(0.009)(0.2) = 0.00144 \]

\[ P(\text{pedestrian} \mid \text{medium, slow}) \propto P(\text{pedestrian}) P(S = \text{medium} \mid \text{pedestrian}) P(V = \text{slow} \mid \text{pedestrian}) \]
\[ P(\text{pedestrian} \mid \text{medium, slow}) \propto (0.19)(0.75)(0.5) = 0.07125 \]

\[ P(\text{ball} \mid \text{medium, slow}) \propto P(\text{ball}) P(S = \text{medium} \mid \text{ball}) P(V = \text{slow} \mid \text{ball}) \]
\[ P(\text{ball} \mid \text{medium, slow}) \propto (0.01)(0.199)(0.4) = 0.000796 \]

Since \( P(\text{pedestrian} \mid \text{medium, slow}) \) has the largest probability, the object is classified as a pedestrian.

**Exercise 3.5.** Given the 3SAT formula in equation (3.14) and the Bayesian network structure in figure 3.4, what are the values of \( P(c_1, c_3 \mid x_1, x_0, x_3, x_4) \) and \( P(y_1 \mid d_2, c_3) \)?

**Solution:** We have \( P(c_1, c_3 \mid x_1, x_0, x_3, x_4) = 1 \) because \( x_1, x_0, x_3, x_4 \) makes the third clause true, and \( P(y_1 \mid d_2, c_3) = 0 \) because \( Y = 1 \) requires that both \( D_2 \) and \( C_3 \) be true.

**Exercise 3.6.** Give a topological sort for each of the following directed graphs:

1. \[ \begin{align*}
D & \rightarrow A \\
F & \rightarrow B \\
C & \rightarrow E
\end{align*} \]

2. \[ \begin{align*}
D & \rightarrow A \\
F & \rightarrow B \\
C & \rightarrow E
\end{align*} \]

**Solution:** There are three valid topological sorts for the first directed graph (Bayesian network): \((F, D, A, B, C, E)\), \((D, A, F, B, C, E)\), and \((D, F, A, B, C, E)\). There are no valid topological sorts for the second directed graph since it is cyclic.

**Exercise 3.7.** Suppose we have the following Bayesian network and we are interested in generating an approximation of the inference query \( P(e^1 \mid b^0, d^1) \) using likelihood-weighted sampling. Given the following samples, write the expressions for each of the sample weights. Additionally, write the equation for estimating \( P(e^1 \mid b^0, d^1) \) in terms of the sample weights \( w_i \).
Solution: For likelihood-weighted sampling, the sample weights are the product of the distributions over evidence variables conditioned on the values of their parents. Thus, the general form for our weights is 

\[ P(b^0 | a^i)P(d^1 | b^0, c^i) \]

We then match each of the values for each sample from the joint distribution.

<table>
<thead>
<tr>
<th>A</th>
<th>B</th>
<th>C</th>
<th>D</th>
<th>E</th>
<th>Weight</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>0</td>
<td>0</td>
<td>1</td>
<td>0</td>
<td>(w_1)</td>
</tr>
<tr>
<td>1</td>
<td>0</td>
<td>0</td>
<td>1</td>
<td>0</td>
<td>(w_4)</td>
</tr>
<tr>
<td>0</td>
<td>0</td>
<td>0</td>
<td>1</td>
<td>1</td>
<td>(w_2)</td>
</tr>
<tr>
<td>1</td>
<td>0</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>(w_3)</td>
</tr>
<tr>
<td>0</td>
<td>0</td>
<td>1</td>
<td>1</td>
<td>0</td>
<td>(w_5)</td>
</tr>
<tr>
<td>1</td>
<td>0</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>(w_6)</td>
</tr>
</tbody>
</table>

To estimate \(P(e^1 | b^0, d^1)\), we simply need to sum the weights of samples consistent with the query variable and divide this by the sum all of the weights.

\[
P(e^1 | b^0, d^1) = \frac{\sum_i w_i(e^{(i)} = 1)}{\sum_i w_i} = \frac{w_3 + w_4 + w_6}{w_1 + w_2 + w_3 + w_4 + w_5 + w_6}
\]

Exercise 3.8. Each year, we receive student scores on standardized mathematics \(M\), reading \(R\), and writing \(W\) exams. Using data from prior years, we create the following distribution:

\[
\begin{bmatrix}
M \\
R \\
W
\end{bmatrix} \sim \mathcal{N}
\begin{bmatrix}
81 \\
82 \\
80
\end{bmatrix}
, \quad
\begin{bmatrix}
25 & -9 & -16 \\
-9 & 36 & 16 \\
-16 & 16 & 36
\end{bmatrix}
\]

Compute the parameters of the conditional distribution over a student’s math and reading test scores, given a writing score of 90.
**Solution:** If we let \( \mathbf{a} \) represent the vector of math and reading scores and \( \mathbf{b} \) represent the writing score, the joint and conditional distributions are as follows:

\[
\begin{bmatrix} \mathbf{a} \\ \mathbf{b} \end{bmatrix} \sim \mathcal{N} \left( \begin{bmatrix} \mu_a \\ \mu_b \end{bmatrix}, \begin{bmatrix} A & C \\ C^\top & B \end{bmatrix} \right)
\]

\[p(\mathbf{a} | \mathbf{b}) = \mathcal{N} \left( \begin{bmatrix} \mathbf{a} | b \end{bmatrix}, \Sigma_{a|b} \right)\]

\[
\mu_{a|b} = \mu_a + CB^{-1}(\mathbf{b} - \mu_b)
\]

\[
\Sigma_{a|b} = A - CB^{-1}C^\top
\]

In the example, we have the following definitions:

\[
\begin{align*}
\mu_a &= \begin{bmatrix} 81 \\ 82 \end{bmatrix} \\
\mu_b &= \begin{bmatrix} 80 \end{bmatrix} \\
A &= \begin{bmatrix} 25 & -9 \\ -9 & 36 \end{bmatrix} \\
B &= \begin{bmatrix} 36 \end{bmatrix} \\
C &= \begin{bmatrix} -16 \\ 16 \end{bmatrix}
\end{align*}
\]

Thus, the parameters of our conditional distribution given \( \mathbf{b} = W = 90 \) are:

\[
\begin{align*}
\mu_{M,R|W=90} &= \begin{bmatrix} 81 \\ 82 \end{bmatrix} + \begin{bmatrix} -16 \\ 16 \end{bmatrix} \frac{1}{36}(90 - 80) \\
&\approx \begin{bmatrix} 76.5 \\ 86.4 \end{bmatrix}
\end{align*}
\]

\[
\begin{align*}
\Sigma_{M,R|W=90} &= \begin{bmatrix} 25 & -9 \\ -9 & 36 \end{bmatrix} - \begin{bmatrix} -16 \\ 16 \end{bmatrix} \frac{1}{36} \begin{bmatrix} -16 \\ 16 \end{bmatrix} \\
&\approx \begin{bmatrix} 25 & -9 \\ -9 & 36 \end{bmatrix} - \begin{bmatrix} 7.1 & -7.1 \\ -7.1 & 7.1 \end{bmatrix} \\
&= \begin{bmatrix} 17.9 & -1.9 \\ -1.9 & 28.9 \end{bmatrix}
\end{align*}
\]

Given that the student scores a 90 on the writing test, based on our conditional distribution, we expect the student to earn a 76.5 on the math test with a standard deviation of \( \sqrt{17.9} \) and an 86.4 on the writing test with a standard deviation of \( \sqrt{28.9} \).
4 Parameter Learning

We have assumed so far that the parameters and structure of our probabilistic models were known. This chapter addresses the problem of learning or fitting model parameters from data.¹ We begin by introducing an approach where we identify the parameters of a model that maximize the likelihood of observing the data. After discussing limitations of such an approach, we introduce an alternative Bayesian approach where we start with a probability distribution over the unknown parameters and then update that distribution based on the observed data using the laws of probability. We then discuss probabilistic models that avoid committing to a fixed number of parameters.

4.1 Maximum Likelihood Parameter Learning

In maximum likelihood parameter learning, we attempt to find the parameters of a distribution that maximize the likelihood of observing the data. If \( \theta \) represents the parameters of a distribution, then the maximum likelihood estimate is

\[
\hat{\theta} = \arg \max_{\theta} P(D | \theta) \tag{4.1}
\]

where \( P(D | \theta) \) is the likelihood that our probability model assigns to the data \( D \) occurring when the model parameters are set to \( \theta \).² We often use the “hat” accent to indicate an estimate of a parameter.

There are two challenges associated with maximum likelihood parameter learning. One is to choose an appropriate probability model by which we define \( P(D | \theta) \). We often assume that the samples in our data \( D \) are independently and identically distributed, which means that our samples \( D = o_{1:n} \) are drawn from a

¹ This chapter focuses on learning model parameters from data, which is an important component of the field of machine learning. A broad introduction to the field is provided by K. P. Murphy, Machine Learning: A Probabilistic Perspective. MIT Press, 2012.

² Here, we write \( P(D | \theta) \) as if it is a probability mass associated with a discrete distribution. However, our probability model may be continuous, in which case we are working with densities.
distribution $o_i \sim P(\cdot | \theta)$ with

$$P(D | \theta) = \prod_i P(o_i | \theta) \quad (4.2)$$

Probability models could include, for example, the categorical distributions or Gaussian distributions mentioned in earlier chapters.

The other challenge is performing the maximization in equation (4.3). For many common probability models, we can perform this optimization analytically. Others may be difficult. A common approach is to maximize the log-likelihood, often denoted $\ell(\theta)$. Since the log-transformation is monotonically increasing, maximizing the log-likelihood is equivalent to maximizing the likelihood:

$$\hat{\theta} = \arg \max_\theta \sum_i \log P(o_i | \theta) \quad (4.3)$$

Computing the sum of log-likelihoods is typically much more numerically stable compared to computing the product of many small probability masses or densities. The remainder of this section will demonstrate how to optimize equation (4.3) for different types of distributions.

### 4.1.1 Maximum Likelihood Estimates for Categorical Distributions

Suppose the random variable $C$ represents whether a flight will result in a mid-air collision, and we are interested in estimating the distribution $P(C)$. Because $C$ is either 0 or 1, it is sufficient to estimate the parameter $\theta = P(c^1)$. What we want to do is infer $\theta$ from data $D$. We have a historical database spanning a decade consisting of $n$ flights with $m$ mid-air collisions. Our intuition, of course, tells us that a good estimate for $\theta$ given the data $D$ is $m/n$. The probability of $m$ mid-air collisions out of $n$ flights can be modeled by the *binomial distribution* under the assumption of independence of outcomes between flights:

$$P(D | \theta) = \frac{n!}{m!(n-m)!} \theta^m (1-\theta)^{n-m} \propto \theta^m (1-\theta)^{n-m} \quad (4.4)$$

The maximum likelihood estimate $\hat{\theta}$ is the value for $\theta$ that maximizes the equation above, which is equivalent to maximizing the logarithm of the likelihood:

$$\ell(\theta) \propto \log(\theta^m (1-\theta)^{n-m})$$

$$= m \log \theta + (n-m) \log(1-\theta) \quad (4.5)$$

3 Although it does not matter whether we maximize the natural logarithm (base $e$) or the common logarithm (base 10) in this equation, we will use $\log(x)$ to mean the logarithm of $x$ with base $e$ throughout this book.
We can use the standard technique for finding the maximum of a function by setting the first derivative of $\ell$ to 0 and then solving for $\theta$. The derivative is given by

$$\frac{\partial}{\partial \theta} \ell(\theta) = \frac{m}{\theta} - n - \frac{m}{1 - \theta}$$

(4.7)

We can solve for $\hat{\theta}$ by setting the derivative to 0:

$$\frac{m}{\hat{\theta}} - n - \frac{m}{1 - \hat{\theta}} = 0$$

(4.8)

After a few algebraic steps, we see that, indeed, $\hat{\theta} = m/n$.

Computing the maximum likelihood estimate for a variable $X$ that can assume $k$ values is also straightforward. If $m_{1:k}$ are the observed counts for the $k$ different values, then the maximum likelihood estimate for $P(x^i \mid m_{1:k})$ is given by

$$\hat{\theta}_i = \frac{m_i}{\sum_{j=1}^{k} m_j}$$

(4.9)

### 4.1.2 Maximum Likelihood Estimates for Gaussian Distributions

In a Gaussian distribution, the log-likelihood of the mean $\mu$ and variance $\sigma^2$ is given by

$$\ell(\mu, \sigma^2) \propto -n \log \sigma - \sum_i (o_i - \mu)^2$$

(4.10)

Again, we can set the derivative to 0 with respect to the parameters and solve for the maximum likelihood estimate:

$$\frac{\partial}{\partial \mu} \ell(\mu, \sigma^2) = \frac{\sum (o_i - \hat{\mu})}{\hat{\sigma}^2} = 0$$

(4.11)

$$\frac{\partial}{\partial \sigma} \ell(\mu, \sigma^2) = -\frac{n}{\hat{\sigma}} + \frac{\sum (o_i - \hat{\mu})^2}{\hat{\sigma}^3} = 0$$

(4.12)

After some algebraic manipulation, we get

$$\hat{\mu} = \frac{\sum o_i}{n} \quad \hat{\sigma}^2 = \frac{\sum (o_i - \hat{\mu})^2}{n}$$

(4.13)

Figure 4.1 provides an example of fitting a Gaussian to data.
4.1.3 Maximum Likelihood Estimates for Bayesian Networks

We can apply maximum likelihood parameter learning to Bayesian networks. Here, we will assume that our network is composed of a set of \( n \) discrete variables that we denote \( X_{1:n} \). Our data \( D = \{ o_1, \ldots, o_m \} \) consists of observed samples from those variables. In our network with structure \( G \), \( r_i \) is the number of instantiations of \( X_i \), and \( q_i \) is the number of instantiations of the parents of \( X_i \). If \( X_i \) has no parents, then \( q_i = 1 \). The \( j \)th instantiation of the parents of \( X_i \) is denoted \( \pi_{ij} \).

The factor table for \( X_i \) thus has \( r_i q_i \) entries, resulting in a total of \( \sum_{i=1}^{n} r_i q_i \) parameters in our Bayesian network. Each parameter is written \( \theta_{ijk} \) and determines

\[
P(X_i = k \mid \pi_{ij}) = \theta_{ijk}
\]  

(4.14)

Although there are \( \sum_{i=1}^{n} r_i q_i \) parameters, only \( \sum_{i=1}^{n} (r_i - 1) q_i \) are independent. We use \( \theta \) to represent the set of all the parameters.

We use \( m_{ijk} \) to represent the number of times \( X_i = k \) given parental instantiation \( j \) in the dataset. Algorithm 4.1 provides an implementation of a function for extracting these counts or statistics from a dataset. The likelihood is given in...
4.2 Bayesian Parameter Learning

Bayesian parameter learning addresses some of the drawbacks of maximum likelihood estimation, especially when the amount of data is limited. For example, suppose our aviation safety database was limited to events of the past week, and we found no recorded mid-air collisions. If \( \theta \) is the probability that a flight results in a mid-air collision, then the maximum likelihood estimate would be
Suppose we have a small network $A \rightarrow B \leftarrow C$ and we want to extract the statistics from a data matrix $D$. We can use the following code:

```plaintext
G = SimpleDiGraph(3)
add_edge!(G, 1, 2)
add_edge!(G, 3, 2)
vars = [Variable(:A,2), Variable(:B,2), Variable(:C,2)]
D = [1 2 2 1; 1 2 2 1; 2 2 2 2]
M = statistics(vars, G, D)
```

The output is an array $M$ consisting of these three count matrices, each of size $q_i \times r_i$:

\[
\begin{bmatrix}
2 & 2 \\
0 & 0 \\
2 & 0 \\
0 & 2 \\
\end{bmatrix}
\]

We can compute the maximum likelihood estimate by normalizing the rows in the matrices in $M$:

```plaintext
θ = [mapslices(x->normalize(x,1), Mi, dims=2) for Mi in M]
```

which produces

\[
\begin{bmatrix}
0.5 & 0.5 \\
\text{nan} & \text{nan} \\
1 & 0 \\
0 & 1 \\
\end{bmatrix}
\]

As we can see, the first and second parental instantiations of the second variable $B$ leads to \text{nan} ("not a number") estimates. Because there are no observations of those two parental instantiations in the data, the denominator of equation (4.16) equals zero, making the parameter estimate undefined. Most of the other parameters are not \text{nan}. For example, the parameter $θ_{112} = 0.5$ means that the maximum likelihood estimate of $P(a^2)$ is 0.5.
\( \hat{\theta} = 0 \). Believing that there is zero chance of a mid-air collision is not a reasonable conclusion, unless our prior hypothesis was, for example, that either all flights were safe or all flights resulted in collision.

The Bayesian approach to parameter learning involves estimating \( p(\theta \mid D) \), the posterior distribution over \( \theta \) given our data \( D \). Instead of obtaining a point estimate \( \hat{\theta} \) as in maximum likelihood estimation, we obtain a distribution. This distribution can help us quantify our uncertainty about the true value of \( \theta \). We can convert this distribution into a point estimate by computing the expectation:

\[
\hat{\theta} = \mathbb{E}_{\theta \sim p(\cdot \mid D)}[\theta] = \int \theta p(\theta \mid D) \, d\theta \tag{4.17}
\]

In some cases, however, the expectation may not be an acceptable estimate as illustrated in figure 4.2. An alternative is to use the maximum a posteriori estimate:

\[
\hat{\theta} = \arg \max_{\theta} p(\theta \mid D) \tag{4.18}
\]

This estimate corresponds to a value of \( \theta \) that is assigned the greatest density. This is often referred to as the mode of the distribution. As shown in figure 4.2, the mode may not be unique.

Bayesian parameter learning can be viewed as inference in a Bayesian network with the structure in figure 4.3, which makes the assumption that the observed variables are conditionally independent of each other. As with any Bayesian network, we must specify \( p(\theta) \) and \( P(O_i \mid \theta) \). We often use a uniform prior \( p(\theta) \).

The remainder of this section discusses how to apply Bayesian parameter learning to different models of \( P(O_i \mid \theta) \).

### 4.2.1 Bayesian Learning for Binary Distributions

Suppose we want to learn the parameters of a binary distribution. Here, we will use \( P(o^1 \mid \theta) = \theta \). To infer the distribution over \( \theta \) in the Bayesian network in figure 4.3, we can proceed with the standard method for performing inference...
discussed in the previous chapter. Here, we will assume a uniform prior:

\[
p(\theta \mid o_{1:n}) \propto p(\theta, o_{1:n})
\]

\[
= p(\theta) \prod_{i=1}^{n} P(o_i \mid \theta)
\]

\[
= \prod_{i=1}^{n} P(o_i \mid \theta)
\]

\[
= \prod_{i=1}^{n} \theta^{o_i} (1 - \theta)^{1-o_i}
\]

\[
= \theta^{m} (1 - \theta)^{n-m}
\]

The posterior is proportional to \( \theta^{m} (1 - \theta)^{n-m} \), where \( m \) is the number of times \( O_i = 1 \). To find the normalization constant, we integrate

\[
\int_0^1 \theta^{m} (1 - \theta)^{n-m} \, d\theta = \frac{\Gamma(m+1)\Gamma(n-m+1)}{\Gamma(n+2)}
\]

where \( \Gamma \) is the gamma function. The gamma function is a real-valued generalization of the factorial. If \( n \) is an integer, then \( \Gamma(n) = (n-1)! \). Taking normalization into account, we have

\[
p(\theta \mid o_{1:n}) = \frac{\Gamma(n+2)}{\Gamma(m+1)\Gamma(n-m+1)} \theta^{m} (1 - \theta)^{n-m}
\]

\[
= \text{Beta}(\theta \mid m+1, n-m+1)
\]

The beta distribution \( \text{Beta}(\alpha, \beta) \) is defined by parameters \( \alpha \) and \( \beta \), and curves for this distribution are shown in figure 4.4. The distribution \( \text{Beta}(1,1) \) corresponds to the uniform distribution spanning 0 to 1.

The distribution \( \text{Beta}(\alpha, \beta) \) has mean

\[
\frac{\alpha}{\alpha + \beta}
\]

When \( \alpha \) and \( \beta \) are both greater than 1, the mode is

\[
\frac{\alpha - 1}{\alpha + \beta - 2}
\]
Conveniently, if a beta distribution is used as a prior over a parameter of a binomial distribution, then the posterior is also a beta distribution. In particular, if the prior is given by $\text{Beta}(\alpha, \beta)$ and we make an observation $o_i$, then we get a posterior of $\text{Beta}(\alpha + 1, \beta)$ if $o_i = 1$ and $\text{Beta}(\alpha, \beta + 1)$ if $o_i = 0$. Hence, if we started with a prior given by $\text{Beta}(\alpha, \beta)$ and our data showed that there were $m$ collisions out of $n$ flights, then the posterior would be given by $\text{Beta}(\alpha + m, \beta + n - m)$. The $\alpha$ and $\beta$ parameters in the prior are sometimes called pseudocounts because they are treated similarly to the observed counts of the two outcomes classes in the posterior, although the pseudocounts need not be integers.

Choosing the prior, in principle, should be done without knowledge of the data used to compute the posterior. Uniform priors often work well in practice, although if expert knowledge is available, then it can be encoded into the prior. For example, suppose we had a slightly bent coin and we wanted to estimate $\theta$, the probability that the coin would land heads. Before we collected any data by flipping the coin, we would start with a belief $\theta$ is likely to be around 0.5. Instead of starting with a uniform prior $\text{Beta}(1, 1)$, we might use $\text{Beta}(2, 2)$ (shown in figure 4.4), which gives more weight to values near 0.5. If we were more confident in an estimate near 0.5, then we could reduce the variance of the prior by increasing the pseudocounts. The prior $\text{Beta}(10, 10)$ is much more peaked than $\text{Beta}(2, 2)$. In general, however, the importance of the prior diminishes with the amount of data used to compute the posterior. If we observe $n$ flips and $m$ were heads, then the difference between $\text{Beta}(1 + m, 1 + n - m)$ and $\text{Beta}(10 + m, 10 + n - m)$ is negligible if we do thousands of coin flips.
4.2.2 Bayesian Learning for Categorical Distributions

The Dirichlet distribution\(^4\) is a generalization of the beta distribution and can be used to estimate the parameters of categorical distributions. Suppose \(X\) is a discrete random variable that takes integer values from 1 to \(n\). We define the parameters of the distribution to be \(\theta_{1:n}\), where \(P(x^i) = \theta_i\). Of course, the parameters must sum to 1, and so only the first \(n - 1\) parameters are independent. The Dirichlet distribution can be used to represent both the prior and the posterior distribution and is parameterized by \(\alpha_{1:n}\). The density is given by

\[
\text{Dir}(\theta_{1:n} \mid \alpha_{1:n}) = \frac{\Gamma(\alpha_0)}{\prod_{i=1}^{n} \Gamma(\alpha_i)} \prod_{i=1}^{n} \theta_i^{\alpha_i-1} \tag{4.29}
\]

where \(\alpha_0\) is used to denote the summation of the parameters \(\alpha_{1:n}\).\(^5\) If \(n = 2\), then it is easy to see that equation (4.29) is equivalent to the beta distribution.

It is common to use a uniform prior where all the Dirichlet parameters \(\alpha_{1:n}\) are set to 1. As with the beta distribution, the parameters in the Dirichlet are often referred to as pseudocounts. If the prior over \(\theta_{1:n}\) is given by \(\text{Dir}(\alpha_{1:n})\) and there are \(m_i\) observations of \(X = i\), then the posterior is given by

\[
p(\theta_{1:n} \mid \alpha_{1:n}, m_{1:n}) = \text{Dir}(\theta_{1:n} \mid \alpha_1 + m_1, \ldots, \alpha_n + m_n) \tag{4.30}
\]

The distribution \(\text{Dir}(\alpha_{1:n})\) has a mean vector whose \(i\)th component is

\[
\frac{\alpha_i}{\sum_{j=1}^{n} \alpha_j} \tag{4.31}
\]

When \(\alpha_i > 1\), the \(i\)th component of the mode is

\[
\frac{\alpha_i - 1}{\sum_{j=1}^{n} \alpha_j - n} \tag{4.32}
\]

As we have seen, Bayesian parameter estimation is straightforward for binary and discrete random variables because it involves simply counting the various outcomes in the data. Bayes’ rule can be used to infer the distribution over the parameters for other parametric distributions. Depending on the choice of prior and the form of the parametric distribution, calculating the posterior over the space of parameters also might be done analytically.

\(^4\) This distribution is named after the German mathematician Johann Peter Gustav Lejeune Dirichlet (1805–1859).

\(^5\) See appendix B for plots of Dirichlet distribution densities for different parameters.
4.2.3 Bayesian Learning for Bayesian Networks

We can apply Bayesian parameter learning to discrete Bayesian networks. The prior over the Bayesian network parameters $\theta$ can be factorized:

$$p(\theta \mid G) = \prod_{i=1}^{n} q_i \prod_{j=1}^{p_i} p(\theta_{ij})$$  (4.33)

where $\theta_{ij} = (\theta_{ij1}, \ldots, \theta_{ijr_i})$. The prior $p(\theta_{ij})$, under some weak assumptions, can be shown to follow a Dirichlet distribution $\text{Dir}(\alpha_{ij1}, \ldots, \alpha_{ijr_i})$. Algorithm 4.2 provides an implementation for creating a data structure holding $\alpha_{ijk}$, where all entries are 1, corresponding to a uniform prior.

After observing data in the form of $m_{ijk}$ counts (as introduced in section 4.1.3), the posterior is then:

$$p(\theta_{ij} \mid \alpha_{ij}, m_{ij}) = \text{Dir}(\theta_{ij} \mid \alpha_{ij1} + m_{ij1}, \ldots, \alpha_{ijr_i} + m_{ijr_i})$$  (4.34)

similar to equation (4.30). Example 4.2 demonstrates this process.

Algorithm 4.2. A function for generating a prior $\alpha_{ijk}$ where all entries are 1. The array of matrices that this function returns is of the same form as the statistics generated by algorithm 4.1. To determine the appropriate dimensions, the function takes as input the list of variables $\text{vars}$ and structure $G$.

Example 4.2. Computing the posterior parameters in a Bayesian network.

We can compute the parameters of the posterior associated with a Bayesian network through simple addition of the prior parameters and counts (equation (4.34)). If we use the matrix of counts $M$ obtained in example 4.1, we can add it to the matrices of prior parameters $\alpha = \text{prior(\text{vars}, G)}$ to obtain the set of posterior parameters $M + \alpha$:

$$\begin{bmatrix} 3 & 3 \\ 1 & 1 \\ 1 & 1 \\ 3 & 1 \\ 1 & 3 \end{bmatrix} + \begin{bmatrix} \alpha_{jj1} & \alpha_{jj2} & \alpha_{jj3} \\ \alpha_{jj1} & \alpha_{jj2} & \alpha_{jj3} \\ \alpha_{jj1} & \alpha_{jj2} & \alpha_{jj3} \\ \alpha_{jj1} & \alpha_{jj2} & \alpha_{jj3} \end{bmatrix} = \begin{bmatrix} 4 & 4 \\ 2 & 2 \\ 4 & 4 \\ 4 & 4 \end{bmatrix}$$
4.3 Nonparametric Learning

The previous two sections assumed that the probabilistic model was of a fixed form and that a fixed set of parameters were to be learned from the data. An alternative approach is based on nonparametric methods in which the number of parameters scale with the amount of data. A common nonparametric method is kernel density estimation (algorithm 4.3). Given observations $o_{1:n}$, kernel density estimation represents the density as follows:

$$p(x) = \frac{1}{n} \sum_{i=1}^{n} \phi(x - o_i)$$  \hspace{1cm} (4.35)

where $\phi$ is a kernel function, which integrates to 1. The kernel function is used to assign greater density to values near the observed data points. A kernel function is generally symmetric, meaning that $\phi(x) = \phi(-x)$. A common kernel is the zero-mean Gaussian distribution. When such a kernel is used, the standard deviation is often referred to as the bandwidth, which can be tuned to control the smoothness of the density function. Larger bandwidths generally lead to smoother densities. Bayesian methods can be applied to the selection of the appropriate bandwidth based on the data. The effect of bandwidth choice is shown in figure 4.5.

Algorithm 4.3. The method `gaussian_kernel` returns a zero-mean Gaussian kernel $\phi(x)$ with bandwidth $b$. Kernel density estimation is also implemented for a kernel $\phi$ and list of observations $O$.

4.4 Learning with Missing Data

When learning the parameters of our probabilistic model we may have missing entries in our data.\footnote{Learning with missing data is the subject of a large body of literature. A comprehensive introduction and review is provided by G. Molenberghs, G. Fitzmaurice, M. G. Kenward, A. Tsiatis, and G. Verbeke, eds., Handbook of Missing Data Methodology. CRC Press, 2014.} For example, if we are conducting a survey, some respondents may decide to skip a question. Table 4.1 shows an example of a dataset with missing entries involving three binary variables: $A$, $B$, and $C$. One approach to handling missing data is to discard all the instances that are incomplete, where there are one or more missing entries. Depending on how much of the data is missing, we might have to discard much of our data. In table 4.1, we would have to discard all but one of the rows, which can be wasteful.
Figure 4.5. Kernel density estimation applied to the same dataset using zero-mean Gaussian kernels with different bandwidths. The histogram in blue shows the underlying dataset frequencies and the black lines indicate the probability density from kernel density estimation. Larger bandwidths smooth out the estimate, whereas smaller bandwidths can over-fit to specific samples.
We can learn model parameters from missing data using either a maximum likelihood or a Bayesian approach. If taking a Bayesian maximum a posteriori approach, we want to find the estimate
\[
\hat{\theta} = \arg \max_{\theta} p(\theta \mid D_{\text{obs}}) = \arg \max_{\theta} \sum_{D_{\text{mis}}} p(\theta \mid D_{\text{obs}}, D_{\text{mis}}) P(D_{\text{mis}} \mid D_{\text{obs}})
\]
where \(D_{\text{obs}}\) and \(D_{\text{mis}}\) consist of all of the observed and missing data, respectively. If the data is continuous, then the sum would be replaced by an integral. The marginalization over the missing data can be expensive computationally. The same marginalization also impacts the computational tractability of a Bayesian approach.

This section discusses two general approaches for learning with missing data without having to enumerate over all possible combinations of missing values. The first involves learning the distribution parameters using predicted values of the missing entries. The second involves an iterative approach for improving our parameter estimates.

We will focus on the context where data is missing at random, meaning that the probability that an entry is missing is conditionally independent of its value given the values of the observed variables. An example of a situation that does not adhere to this assumption might include radar data containing measurements of the distance to a target, but the measurement may be missing either due to noise or the target is beyond the sensing range. The fact that an entry is missing is an indication that the value is more likely to be high. Accounting for this form of missingness requires different models and algorithms from what we discuss here.\(^7\)

### 4.4.1 Data Imputation

An alternative to discarding incomplete instances is to impute the values of missing entries. Data imputation is the processes of inferring values for missing entries. One way to view imputation is as an approximation of equation (4.37) where we find
\[
\hat{D}_{\text{mis}} = \arg \max_{D_{\text{mis}}} p(D_{\text{mis}} \mid D_{\text{obs}})
\]
Once we have the imputed missing values, we can then use that data to produce a maximum posteriori estimate

$$
\hat{\theta} = \arg \max_{\theta} p(\theta \mid D_{\text{obs}}) \approx \arg \max_{\theta} p(\theta \mid D_{\text{obs}}, \hat{D}_{\text{mis}}) \quad (4.39)
$$
or, alternatively, we can take a maximum likelihood approach.

Solving equation (4.38) may still be computationally challenging. One simple approach for discrete datasets is to replace missing entries with the most commonly observed value, called the marginal mode. For example, in table 4.1, we might replace all of missing values for $A$ with its marginal mode of 1.

Continuous data often lacks duplicates. However, we can fit a distribution to continuous values and then use the mode of the resulting distribution. For example, we might fit a Gaussian distribution to the data in table 4.2, and then fill in the missing entries with the mean of the observed values associated with each variable. The top-left plot in figure 4.6 illustrates the effect of this approach on two-dimensional data. The red lines show how values with missing first or second components are paired with their imputed counterparts. We can then use the observed and imputed data to arrive at a maximum likelihood estimate of the parameters of a joint Gaussian distribution. As we can see, this method of imputation does not always produce sensible predictions and the learned model is quite poor.

We can often do better if we account for the probabilistic relationships between the observed and unobserved variables. In figure 4.6, there is clearly correlation between the two variables; hence, knowing the value of one variable can help predict the value of the other variable. A common approach to imputation, called nearest-neighbor imputation, is to use the values associated with the instance that is nearest with respect to a distance measure defined on the observed variables. The top-right plot in figure 4.6 uses the Euclidean distance for imputation. This approach tends to lead to better imputations and learned distributions.

An alternative approach is to fit a distribution to the fully observed data and then use that distribution to infer the missing values. We can use the inference algorithms from the previous chapter to perform this inference. For example, if our data is discrete and we can assume a Bayesian network structure, we can use variable elimination or Gibbs sampling to produce a distribution over the missing variables for an instance from the observed variables. From this distribution, we might use the mean or mode to impute the missing values. Alternatively, we can

<table>
<thead>
<tr>
<th>A</th>
<th>B</th>
<th>C</th>
</tr>
</thead>
<tbody>
<tr>
<td>-6.5</td>
<td>0.9</td>
<td>4.2</td>
</tr>
<tr>
<td>?</td>
<td>4.4</td>
<td>9.2</td>
</tr>
<tr>
<td>7.8</td>
<td>?</td>
<td>?</td>
</tr>
<tr>
<td>?</td>
<td>?</td>
<td>?</td>
</tr>
</tbody>
</table>

Table 4.2. Example data with continuous values.

The mode of a distribution is where the probability is at a maximum. For categorical distributions over individual discrete values this is the most frequently occurring value. For continuous distributions it is a local maximum in the probability density function.
Figure 4.6. A demonstration of imputation techniques. Shown are ellipses where the density of the maximum likelihood estimate of the joint distribution is equal to 0.02.
pull a sample from this distribution. If our data is continuous and we can assume that the data is jointly Gaussian, we can use algorithm 3.11 to infer the posterior distribution. The bottom plots in figure 4.6 demonstrate imputation using these posterior mode and posterior sampling approaches.

4.4.2 Expectation-Maximization

The expectation-maximization (EM) category of approaches involves iterative improvement of the distribution parameter estimate \( \hat{\theta} \). We begin with an initial \( \hat{\theta} \), which may be a guess, randomly sampled from a prior distribution over distribution parameters, or estimated using one of the methods discussed in section 4.4.1. At each iteration, we perform a two-step process to update \( \hat{\theta} \).

The first step is called the expectation step (E-step), where we use the current estimate of \( \theta \) to infer completions of the data. For example, if we are modeling our data using a discrete Bayesian network, we can use one of our inference algorithms to infer a distribution over the missing entries for each instance. When extracting the counts, we apply a weighting proportional to the likelihood of the completions as shown in example 4.3. In cases where there are many missing variables, there may be too many possible completions to practically enumerate, making a sampling-based approach attractive. We may also want to use sampling as an approximation method when our variables are continuous.

The second step is called the maximization step (M-step), where we attempt to find a new \( \hat{\theta} \) that maximizes the likelihood of the completed data. If we have a discrete Bayesian network with the weighted counts in the form shown in example 4.3, then we can perform the same maximum likelihood estimate as discussed earlier in this chapter. Alternatively, we can use a maximum a posteriori estimate if we want to incorporate a prior.

This approach is not guaranteed to converge to model parameters that maximize the likelihood of the observed data, but it can work well in practice. To reduce the risk of the algorithm converging to only a local optimum, we can run the algorithm to convergence from many different initial points in the parameter space. We simply choose the resulting parameter estimate in the end that maximizes likelihood.

Expectation-maximization can even be used to impute values for variables that are not observed at all in the data. Such variables are called latent variables. To illustrate, suppose we have a Bayesian network \( Z \rightarrow X \), where \( X \) is continuous...
Suppose we have a binary Bayesian network with $A \rightarrow B$. We start by assuming that $\hat{\theta}$ implies:

\[
P(a^1) = 0.5 \quad P(b^1 \mid a^0) = 0.2 \quad P(b^1 \mid a^1) = 0.6
\]

Using these parameters, we can expand the dataset with missing values (left) to a weighted dataset with all possible individual completions (right).

<table>
<thead>
<tr>
<th>$A$</th>
<th>$B$</th>
<th>weight</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>0</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td>0</td>
<td>?</td>
<td>$P(b^1 \mid a^0) = 0.2$</td>
</tr>
<tr>
<td>?</td>
<td>0</td>
<td>$\alpha P(a^0)P(b^0 \mid a^0) = \alpha 0.4 = 2/3$</td>
</tr>
<tr>
<td>1</td>
<td>0</td>
<td>$\alpha P(a^1)P(b^0 \mid a^1) = \alpha 0.2 = 1/3$</td>
</tr>
</tbody>
</table>

The $\alpha$ in the calculation above is a normalization constant, which enforces that each instance is expanded to instances whose weights sum to one. The count matrices are then:

\[
\begin{bmatrix}
(2 + 2/3) & (1 + 1/3) \\
\end{bmatrix} \quad \begin{bmatrix}
(0.8 + 2/3) & 1.2 \\
1/3 & 1 \\
\end{bmatrix}
\]
and $Z$ is discrete and can take on one of three values. Our model assumes $p(x \mid z)$ is conditional Gaussian. Our dataset contains only values for $X$ but none for $Z$. We start with an initial $\hat{\theta}$ and use it to infer a probability distribution over the values of $Z$ given the value of $X$ for each instance. The distribution over entry completions are then used to update our estimate of the parameters of $P(Z)$ and $P(X \mid Z)$ as illustrated in example 4.4. We iterate to convergence, which is often quite fast. The parameters that we obtain in this example define a Gaussian mixture model, which was introduced in section 2.2.2.

4.5 Summary

- Parameter learning involves inferring the parameters of a probabilistic model from data.

- A maximum likelihood approach to parameter learning involves maximizing a likelihood function, which can be done analytically for some models.

- A Bayesian approach to parameter learning involves inferring a probability distribution over the underlying parameter using Bayes’ rule.

- The beta and Dirichlet distributions are examples of Bayesian priors that are easily updated with evidence.

- In contrast with parametric learning, which assumes a fixed parameterization of a probability model, nonparametric learning uses representations that grow with the amount of data.

- We can approach the problem of learning parameters from missing data using methods such as data imputation or expectation-maximization where we make inferences based on observed values.

4.6 Exercises

Exercise 4.1. Consider a continuous random variable $X$ that follows the Laplace distribution parameterized by $\mu$ and $b$ with density

$$p(x \mid \mu, b) = \frac{1}{2b} \exp\left(-\frac{|x - \mu|}{b}\right)$$
We have a Bayesian network $Z \rightarrow X$, where $Z$ is discrete latent variable with three values and $X$ is continuous with $p(x \mid z)$ modeled as a conditional Gaussian. Hence, we have parameters defining $P(z^1)$, $P(z^2)$, and $P(z^3)$, and a $\mu_i$ and $\sigma_i$ for each of the three Gaussian distributions associated with different values of $Z$. In this example, we use an initial parameter vector $\hat{\theta}$ that specifies $P(z^i) = 1/3$ and $\sigma_i = 1$ for all $i$. We spread out the means with $\mu_1 = -4$, $\mu_2 = 0$, and $\mu_3 = 4$.

Suppose our first instance in our data has $X = 4.2$. We want to infer the distribution over $Z$ for that instance:

$$P(z^i \mid X = 4.2) = \frac{P(z^i) \mathcal{N}(4.2 \mid \mu_i, \sigma_i^2)}{\sum_j P(z^j) \mathcal{N}(4.2 \mid \mu_j, \sigma_j^2)}$$

We compute this distribution for all the instances in our dataset. For the weighted completions, we can obtain a new estimate for $\hat{\theta}$. We estimate $P(z^i)$ by taking the mean across the instances in our dataset. To estimate $\mu_i$ and $\sigma_i$, we use the mean and standard deviation of the values for $X$ over the instances in our dataset, weighted by the probability of $z^i$ associated with the various instances.

We repeat the process until convergence. The plot below shows three iterations. The histogram was generated from the values of $X$. The dark blue function indicates the inferred density. By the third iteration, our parameters of the Gaussian mixture model closely represent the data distribution.

Example 4.4. Expectation maximization applied to learning the parameters of a Gaussian mixture model.
Compute the maximum likelihood estimates of the parameters of a Laplace distribution given a dataset $D$ of $n$ independent observations $x_{1:n}$. Note that $\frac{\partial |u|}{\partial x} = \text{sign}(u)\frac{\partial u}{\partial c}$, where the sign function returns the sign of its argument.

**Solution:** Since the observations are independent, we can write the log-likelihood function as the summation:

$$\ell(\mu, b) = \sum_{i=1}^{n} \log \left( \frac{1}{2b} \exp \left( \frac{-|x_i - \mu|}{b} \right) \right)$$

$$= -\sum_{i=1}^{n} \log 2b - \sum_{i=1}^{n} \frac{|x_i - \mu|}{b}$$

$$= -n \log 2b - \frac{1}{b} \sum_{i=1}^{n} |x_i - \mu|$$

To obtain the maximum likelihood estimates of the true parameters $\mu$ and $b$, we take the partial derivatives of the log-likelihood with respect to each of the parameters, set them to zero, and solve for each parameter. First, we solve for $\hat{\mu}$

$$\frac{\partial}{\partial \mu} \ell(\mu, b) = \frac{1}{b} \sum_{i=1}^{n} \text{sign}(x_i - \mu)$$

$$0 = \frac{1}{b} \sum_{i=1}^{n} \text{sign}(x_i - \hat{\mu})$$

$$0 = \sum_{i=1}^{n} \text{sign}(x_i - \hat{\mu})$$

$$\hat{\mu} = \text{median}(x_{1:n})$$

Now, solving for $\hat{b}$

$$\frac{\partial}{\partial b} \ell(\mu, b) = -\frac{n}{b} + \frac{1}{b^2} \sum_{i=1}^{n} |x_i - \hat{\mu}|$$

$$0 = -\frac{n}{b} + \frac{1}{b^2} \sum_{i=1}^{n} |x_i - \hat{\mu}|$$

$$\frac{n}{\hat{b}} = \frac{1}{b^2} \sum_{i=1}^{n} |x_i - \hat{\mu}|$$

$$\hat{b} = \frac{1}{n} \sum_{i=1}^{n} |x_i - \hat{\mu}|$$

Thus, the maximum likelihood estimates for the parameters of a Laplace distribution are $\hat{\mu}$, the median of the observations, and $\hat{b}$, the mean of absolute deviations from the median.
Exercise 4.2. This question explores the application of maximum likelihood estimation to censored data, where some measurements are only partially known. Suppose we are building electric motors for a quadcopter drone, and we want to produce a model of how long they last until failure. Although there may be more suitable distributions for modeling the reliability of components,\(^\text{10}\) we will use an exponential distribution parameterized by \(\lambda\) with probability density function \(\lambda \exp(-\lambda x)\) and cumulative distribution function \(1 - \exp(\lambda x)\). We fly five drones. Three have motor failures after 132 hours, 42 hours, and 89 hours. We stopped testing the other two after 200 hours without failure; we do not know their failure times, just that they are greater than 200 hours. What is the maximum likelihood estimate for \(\lambda\) given this data?

Solution: This problem has \(n = 3\) fully observed measurements and \(m = 2\) censored measurements. We use \(t_i\) to represent the \(i\)th fully observed measurement and \(t_j\) to represent the \(j\)th censored measurement. The likelihood of a single measurement above \(t_j\) is the complement of the cumulative distribution function, which is simply \(\exp(-\lambda t_j)\). Hence, the likelihood of the data is

\[
\left(\prod_{i=1}^{n} \lambda e^{-\lambda t_i}\right) \left(\prod_{j=1}^{m} e^{-\lambda t_j}\right)
\]

We use our standard approach of maximizing the log-likelihood, which is given by

\[
\ell(\lambda) = \sum_{i=1}^{n} (\log \lambda - \lambda t_i) + \sum_{j=1}^{m} -\lambda t_j
\]

The derivative with respect to \(\lambda\) is

\[
\frac{\partial \ell}{\partial \lambda} = \frac{n}{\lambda} - \sum_{i=1}^{n} t_i - \sum_{j=1}^{m} t_j
\]

Setting this derivative to 0, we can solve for \(\lambda\) to obtain the maximum likelihood estimate:

\[
\lambda = \frac{n}{\sum_{i=1}^{n} t_i + \sum_{j=1}^{m} t_j} = \frac{3}{132 + 42 + 89 + 200 + 200} \approx 0.00452
\]

The mean of the exponential distribution is \(1/\lambda\), making the mean in our problem 221 hours.

Exercise 4.3. We have a Bayesian network where the variables \(X_{1:3}\) can take on values in \{1, 2\} and \(X_4\) can take on values in \{1, 2, 3\}. Given the dataset \(D\) of observations \(o_{1:m}\), generate the maximum likelihood estimates of the associated conditional distribution parameters \(\theta\).
Solution: We can generate count matrices $M_i$ of size $q_i \times r_i$ for each node by iterating through the dataset and storing the counts. We then normalize each row in the count matrices to yield the matrices containing the maximum likelihood estimates of the parameters:

$$
M_1 = \begin{bmatrix} 3 & 1 \\ 0 & 0 \\ 2 & 2 \\ 0 & 2 \\ 0 & 1 \\ 0 & 1 \end{bmatrix} \\
M_2 = \begin{bmatrix} 0.75 & 0.25 \\ \text{NAN} & \text{NAN} \\ 1.0 & 0.0 \\ 0.0 & 1.0 \\ 0.0 & 1.0 \\ 0.0 & 1.0 \end{bmatrix} \\
M_3 = \begin{bmatrix} 6 & 4 \\ 5 & 0 & 2 \\ 1 & 1 & 1 \end{bmatrix} \\
M_4 = \begin{bmatrix} 0.6 & 0.4 \\ 0.71 & 0.0 & 0.29 \\ 0.33 & 0.33 & 0.34 \end{bmatrix}
$$

Exercise 4.4. We have a biased coin, and we want to estimate the Bernoulli parameter $\phi$ that specifies the probability the coin lands on heads. If the first toss lands on heads ($o_1 = 1$),

- What is the maximum likelihood estimate of $\phi$?
- Using a uniform prior, what is the maximum a posteriori (MAP) estimate of $\phi$?
- Using a uniform prior, what is the expectation of our posterior distribution over $\phi$?

Solution: Since our first toss lands on heads, we have $m = 1$ successes and $n = 1$ trials.

- The maximum likelihood estimate of $\phi$ is $m/n = 1$.
- Using a uniform Beta(1, 1) prior, the posterior distribution is $\text{Beta}(1 + m, 1 + n - m) = \text{Beta}(2, 1)$. The maximum a posteriori (MAP) estimate of $\phi$ or mode of the posterior distribution is

$$
\frac{\alpha - 1}{\alpha + \beta - 2} = \frac{2 - 1}{2 + 1 - 2} = 1
$$
• The mean of the posterior distribution is

\[
\frac{\alpha}{\alpha + \beta} = \frac{2}{2 + 1} = \frac{2}{3}
\]

**Exercise 4.5.** Suppose we are given the following dataset with one missing value. What is the value that will be imputed using marginal mode imputation assuming the marginal distribution is a Gaussian? What is the value that will be imputed using nearest-neighbor imputation?

<table>
<thead>
<tr>
<th>X₁</th>
<th>X₂</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.5</td>
<td>1.0</td>
</tr>
<tr>
<td>?</td>
<td>0.3</td>
</tr>
<tr>
<td>−0.6</td>
<td>−0.3</td>
</tr>
<tr>
<td>0.1</td>
<td>0.2</td>
</tr>
</tbody>
</table>

**Solution:** Assuming the marginal distribution over \(X₁\) is a Gaussian, we can compute the marginal mode, which is the mean parameter of the Gaussian distribution

\[
\mu = \frac{1}{n} \sum_{i=1}^{n} x_i = \frac{0.5 - 0.6 + 0.1}{3} = 0
\]

Thus, for marginal mode imputation, the missing value will be set to 0. For nearest-neighbor imputation, the nearest sample to \(X₂ = 0.3\) is the fourth sample, so the missing value will be set to 0.1.

**Exercise 4.6.** Suppose we are given a dataset over two variables \(X₁, X₂\) with several missing values. We assume that \(X₁, X₂\) are jointly Gaussian and use the fully-observed samples to fit the following distribution

\[
\begin{bmatrix} X₁ \\ X₂ \end{bmatrix} \sim \mathcal{N}\left( \begin{bmatrix} 5 \\ 2 \end{bmatrix}, \begin{bmatrix} 4 & 1 \\ 1 & 2 \end{bmatrix} \right)
\]

What is the value that will be imputed for \(X₁\) for the sample \(X₂ = 1.5\) using posterior mode imputation? What distribution do we need to sample from for posterior sample imputation?

**Solution:** Since we assumed \(X₁, X₂\) are jointly Gaussian, the posterior distribution over \(X₁\) given \(X₂\) is also Gaussian and its mode is the mean parameter of the posterior distribution. We can compute the mean of the posterior distribution as follows:

\[
p(x₁ | x₂) = \mathcal{N}\left( x₁ | \mu_{x₁|x₂}, \sigma^2_{x₁|x₂} \right)
\]

\[
\mu_{x₁|x₂=1.5} = 5 + (1)(2)^{-1}(1.5 - 2) = 4.75
\]

Thus, for posterior mode imputation, the missing value will be set to 4.75. For posterior sample imputation, we will sample a value \(X₁ \sim \mathcal{N}(4.75, 3.5)\).
5 Structure Learning

The previous chapters assumed that the structures of our probabilistic models were known. This chapter discusses methods for learning the structure of models from data. We begin by explaining how to compute the probability of a graphical structure given the data. Generally, we want to maximize this probability. Because the space of possible graphical structures is usually too large to enumerate, we discuss ways to search this space efficiently.

5.1 Bayesian Network Scoring

We want to be able to score a network structure $G$ based on how well it models the data. A maximum a posteriori approach to structure learning involves finding a $G$ that maximizes $P(G \mid D)$. We first explain how to compute a Bayesian network score based on $P(G \mid D)$ to measure how well $G$ models the data. We then explain how to go about searching the space of networks for the highest scoring network. Like inference in Bayesian networks, it can be shown that for general graphs and input data, learning the structure of a Bayesian network is NP-hard.

We compute $P(G \mid D)$ using Bayes’ rule and the law of total probability:

$$
P(G \mid D) \propto P(G)P(D \mid G)$$

$$
= P(G) \int P(D \mid \theta, G)p(\theta \mid G)\,d\theta
$$

Integrating with respect to $\theta$ results in

$$
P(G \mid D) = P(G) \prod_{i=1}^{n} \prod_{j=1}^{q_i} \frac{\Gamma(\alpha_{ij0})}{\Gamma(\alpha_{ij0} + m_{ij0})} \prod_{k=1}^{r_i} \frac{\Gamma(\alpha_{ijk} + m_{ijk})}{\Gamma(\alpha_{ijk})}
$$
In the equation above,

\[ \alpha_{ij0} = \sum_{k=1}^{r_i} \alpha_{ijk} \quad m_{ij0} = \sum_{k=1}^{r_i} m_{ijk} \] (5.4)

Finding the G that maximizes equation (5.2) is the same as finding the G that maximizes what is called the Bayesian score:

\[
\log P(G \mid D) = \log P(G) + \sum_{i=1}^{n} \sum_{j=1}^{q_i} \left( \log \left( \frac{\Gamma(\alpha_{ij0})}{\Gamma(\alpha_{ij0} + m_{ij0})} \right) \right) + \sum_{k=1}^{r_i} \log \left( \frac{\Gamma(\alpha_{ijk} + m_{ijk})}{\Gamma(\alpha_{ijk})} \right) \] (5.5)

The Bayesian score is more convenient to compute numerically because it is easier to add the logarithm of small numbers together than to multiply small numbers together. Many software libraries can compute the logarithm of the gamma function directly.

A variety of different graph priors have been explored in the literature, although a uniform prior is often used in practice, in which case \( \log P(G) \) can be dropped from the computation of the Bayesian score in equation (5.5). Algorithm 5.1 provides an implementation.

```python
function bayesian_score_component(M, α)
    p = sum(loggamma.(α + M))
    p -= sum(loggamma.(α))
    p += sum(loggamma.(sum(α,dims=2)))
    p -= sum(loggamma.(sum(α,dims=2) + sum(M,dims=2)))
    return p
end

function bayesian_score(vars, G, D)
    n = length(vars)
    M = statistics(vars, G, D)
    α = prior(vars, G)
    return sum(bayesian_score_component(M[i], α[i]) for i in 1:n)
end
```

Algorithm 5.1. An algorithm for computing the Bayesian score for a list of variables vars and a graph G given data D. This method uses a uniform prior \( \alpha_{ijk} = 1 \) for all \( i, j, \) and \( k \) as generated by algorithm 4.2. The \text{loggamma} function is provided by \text{SpecialFunctions.jl}. The previous chapter introduced the \text{statistics} and \text{prior} functions. Note that \( \log(\Gamma(\alpha)/\Gamma(\alpha + m)) = \log \Gamma(\alpha) - \log \Gamma(\alpha + m) \) and that \( \log \Gamma(1) = 0. \)

A byproduct of optimizing the structure with respect to the Bayesian score is that we are able to find the right balance in the model complexity given the available data. We do not want a model that misses out on capturing important relationships between variables, but we also do not want a model that has too many parameters to be adequately learned from limited data.
To illustrate how the Bayesian score helps us balance model complexity, consider the network in figure 5.1. The value of $A$ weakly influences the value of $B$, and $C$ is independent of the other variables. We sample from this “true” model to generate data $D$ and then try to learn the model structure. There are 25 possible network structures involving three variables, but we will focus on the scores for the models in figure 5.2.

![Network Diagrams](image)

Figure 5.1. A simple Bayesian network to illustrate how the Bayesian network score helps us balance model complexity.

Figure 5.2. Three Bayesian network structures with varying levels of complexity.

Figure 5.3 shows how the Bayesian scores of the completely connected and unconnected models compare to the true model as the amount of data increases. In the plot, we subtract the score of the true model, so values above 0 indicate that the model provides a better representation than the true model given the available data. The plot shows that the unconnected model does better than the true model when there are fewer than $5 \times 10^3$ samples. The completely connected model never does better than the true model, but it starts to do better than the unconnected model at about $10^4$ samples because there are sufficient data to adequately estimate its seven independent parameters.

### 5.2 Directed Graph Search

In *directed graph search*, we search the space of directed acyclic graphs for one that maximizes the Bayesian score. The space of possible Bayesian network structures grows superexponentially.

With 10 nodes, there are $4.2 \times 10^{18}$ possible directed acyclic graphs. With 20 nodes, there are $2.4 \times 10^{72}$. Except for Bayesian networks with few nodes, we cannot enumerate the space of possible structures to find the highest scoring network. Therefore, we have to rely on a search strategy.

---

Fortunately, search is a general problem, and a wide variety of different generic search algorithms have been studied over the years.

One of the most common search strategies is called K2. The search (algorithm 5.2) runs in polynomial time but does not guarantee finding a globally optimal network structure. It can use any scoring function, but it is often used with the Bayesian score because of its ability to balance the complexity of the model with the amount of data available. K2 begins with a graph with no directed edges and then iterates over the variables according to a provided ordering, greedily adding parents to the nodes in a way that maximally increases the score. It is common for K2 to impose an upper bound on the number of parents for any one node to reduce the required computation. The original K2 algorithm assumed a unit uniform Dirichlet prior with $\alpha_{ijk} = 1$ for all $i$, $j$, and $k$, but any prior can be used in principle.

A general search strategy is local search, which is sometimes called hill climbing. Algorithm 5.3 provides an implementation. We start with an initial graph and then move to the highest scoring neighbor. The neighborhood of a graph consists of the graphs that are only one basic graph operation away, where the basic graph operations include introducing an edge, removing an edge, and reversing an edge. Of course, not all operations are possible from a particular graph, and operations

---

5 The name comes from the fact that it is an evolution of a system called Kutató. The algorithm was introduced by G.F. Cooper and E. Herskovits, “A Bayesian Method for the Induction of Probabilistic Networks from Data,” Machine Learning, vol. 4, no. 9, pp. 309–347. 1992.
Algorithm 5.2. K2 search of the space of directed acyclic graphs using a specified variable ordering. This variable ordering imposes a topological ordering in the resulting graph. The `fit` function takes an ordered list variables `vars` and a dataset `D`. The method starts with an empty graph and iteratively adds the next parent that maximally improves the Bayesian score.

```plaintext
def K2Search
    ordering::Vector{Int} # variable ordering
end

function fit(method::K2Search, vars, D)
    G = SimpleDiGraph(length(vars))
    for (k,i) in enumerate(method.ordering[2:end])
        y = bayesian_score(vars, G, D)
        while true
            y_best, j_best = -Inf, 0
            for j in method.ordering[1:k]
                if !has_edge(G, j, i)
                    add_edge!(G, j, i)
                    y' = bayesian_score(vars, G, D)
                    if y' > y_best
                        y_best, j_best = y', j
                    end
                    rem_edge!(G, j, i)
                end
            end
            if y_best > y
                y = y_best
                add_edge!(G, j_best, i)
            else
                break
            end
        end
    end
    return G
end
```
that introduce cycles into the graph are invalid. The search continues until the current graph scores no lower than any of its neighbors.

An *opportunistic* version of local search is implemented in algorithm 5.3. Rather than generating all graph neighbors at every iteration, this method generates a single random neighbor and accepts it if its Bayesian score is greater than that of the current graph.

```
struct LocalDirectedGraphSearch
    G  # initial graph
    k_max # number of iterations
end

function rand_graph_neighbor(G)
    n = nv(G)
    i = rand(1:n)
    j = mod1(i + rand(2:n)-1, n)
    G' = copy(G)
    has_edge(G, i, j) ? rem_edge!(G', i, j) : add_edge!(G', i, j)
    return G'
end

function fit(method::LocalDirectedGraphSearch, vars, D)
    G = method.G
    y = bayesian_score(vars, G, D)
    for k in 1:method.k_max
        G' = rand_graph_neighbor(G)
        y' = is_cyclic(G') ? -Inf : bayesian_score(vars, G', D)
        if y' > y
            y, G = y', G'
        end
    end
    G
end
```

Algorithm 5.3. Local directed graph search, which starts with an initial directed graph $G$ and opportunistically moves to a random graph neighbor whenever its Bayesian score is greater. It repeats this process for $k_{\text{max}}$ iterations. Random graph neighbors are generated by either adding or removing a single edge. Edge addition can result in a graph with cycles, in which case we assign a score of $-\infty$.

Localsearch can get stuck in *local optima*, preventing it from finding the globally optimal network structure. Various strategies have been proposed for addressing local optima, including the following:

- **Randomized restart.** Once a local optima has been found, simply restart the search at a random point in the search space.

- **Simulated annealing.** Instead of always moving to the neighbor with greatest fitness, the search can visit neighbors with lower fitness according to some randomized exploration strategy. As the search progresses, the randomness in

---

6 The field of optimization is quite vast, and many methods have been developed for addressing local optima. This textbook provides an overview: M. J. Kochenderfer and T. A. Wheeler, *Algorithms for Optimization*. MIT Press, 2019.
the exploration decreases according to some schedule. This approach is called simulated annealing because of its inspiration from annealing in metallurgy.

- **Genetic algorithms.** The procedure begins with an initial random population of points in the search space represented as binary strings. Each bit in a string indicates the presence or absence of an arrow between two nodes. String manipulation thus allows for searching the space of directed graphs. The individuals in the population reproduce at a rate proportional to their score. Individuals selected for reproduction have their strings recombined randomly through genetic crossover. Genetic crossover involves selecting a crossover point on two randomly selected individuals and then swapping the strings after that point. Mutations are also introduced randomly into the population by randomly flipping bits in the strings. The process of evolution continues until a satisfactory point in the search space is found.

- **Memetic algorithms.** This approach is sometimes called *genetic local search* and is simply a combination of genetic algorithms and local search. After genetic recombination, local search is applied to the individuals.

- **Tabu search.** Previous methods can be augmented to maintain a *tabu list* containing recently visited points in the search space. The search algorithm avoids neighbors in the tabu list.

Some search strategies may work better than others on certain datasets, but in general finding the global optima remains NP-hard. Many applications, however, do not require the globally optimal network structure. A locally optimal structure is often acceptable.

### 5.3 Markov Equivalence Classes

As discussed earlier, the structure of a Bayesian network encodes a set of conditional independence assumptions. An important observation to make when trying to learn the structure of a Bayesian network is that two different graphs can encode the same independence assumptions. As a simple example, the two-variable network $A \rightarrow B$ has the same independence assumptions as $A \leftarrow B$. Solely on the basis of the data, we cannot justify the direction of the edge between $A$ and $B$. 
If two networks encode the same conditional independence assumptions, we say they are Markov equivalent. It can be proven that two graphs are Markov equivalent if and only if they have (1) the same edges without regard to direction and (2) the same immoral v-structure. An immoral v-structure is a v-structure $X \rightarrow Y \leftarrow Z$ with $X$ and $Z$ not directly connected, as shown in figure 5.4. A Markov equivalence class is a set containing all the directed acyclic graphs that are Markov equivalent to each other. A method for checking Markov equivalence is given in algorithm 5.4.

In general, two structures belonging to the same Markov equivalence class may be given different scores. However, if the Bayesian score is used with Dirichlet priors such that $\kappa = \sum_i \sum_k x_{ijk}$ is constant for all $i$, then two Markov equivalent structures are assigned the same score. Such priors are called BDe, and a special case is the BDeu prior, which assigns $x_{ijk} = \kappa / (q_i r_i)$. Although the commonly used uniform prior $x_{ijk} = 1$ does not always result in identical scores being assigned to structures in the same equivalence class, they are often fairly close. A scoring function that assigns the same score to all structures in the same class is called score equivalent.

5.4 Partially Directed Graph Search

A Markov equivalence class can be represented as a partially directed graph, sometimes called an essential graph or a directed acyclic graph pattern. A partially directed graph can contain both directed edges and undirected edges. An example of a partially directed graph that encodes a Markov equivalence class is shown in figure 5.5. A directed acyclic graph $G$ is a member of the Markov equivalence class encoded by a partially directed graph $G'$ if and only if $G$ has the same edges as $G'$ without regard to direction and has the same immoral v-structures as $G'$.
Function `are_markov_equivalent(G, H)`

```python
function are_markov_equivalent(G, H)
    if nv(G) != nv(H) || ne(G) != ne(H) ||
        !(all(has_edge(H, e) || has_edge(H, reverse(e)) for e in edges(G))
    return false
end
for c in 1:nv(G)
    parents = inneighbors(G, c)
    for (a, b) in subsets(parents, 2)
        if !(has_edge(G, a, b) && !(has_edge(G, b, a) &&
            !(has_edge(H, a, c) && has_edge(H, b, c)))
        return false
    end
end
return true
end
```

Algorithm 5.4. A method for determining whether the directed acyclic graphs $G$ and $H$ are Markov equivalent. The `subsets` function from `IterTools.jl` returns all subsets of a given set and a specified size.

Figure 5.5. Markov equivalence class and examples of members and a nonmember. The nonmember does not belong to the Markov equivalence class because it introduces an immoral v-structure $A \rightarrow B \leftarrow C$ that is not indicated in the partially directed graph.
Instead of searching the space of directed acyclic graphs, we can search the space of Markov equivalence classes represented by partially directed graphs.\(^9\) Although the space of Markov equivalence classes is, of course, smaller than the space of directed acyclic graphs, it is not significantly smaller; the ratio of directed acyclic graphs to equivalence classes asymptotes to around 3.7 fairly quickly.\(^10\)

A problem with hill climbing in the space of directed acyclic graphs is that the neighborhood may consist of other graphs that are in the same equivalence class with the same score, which can lead to the search becoming stuck in a local optimum. Searching the space of equivalence classes allows us to jump to different directed acyclic graphs outside of the current equivalence class.

Any of the general search strategies presented in section 5.2 can be used. If a form of local search is used, then we need to define the local graph operations that define the neighborhood of the graph. Examples of local graph operations include:

- If an edge between \(X\) and \(Y\) does not exist, add either \(X \rightarrow Y\) or \(X \leftarrow Y\).
- If \(X \rightarrow Y\) or \(X \leftarrow Y\), then remove the edge between \(X\) and \(Y\).
- If \(X \rightarrow Y\), then reverse the direction of the edge to get \(X \leftarrow Y\).
- If \(X \rightarrow Y \rightarrow Z\), then add \(X \rightarrow Y \leftarrow Z\).

To score a partially directed graph, we generate a member of its Markov equivalence class and compute its score.

\section*{5.5 Summary}

- Fitting a Bayesian network to data requires selecting the Bayesian network structure that dictates the conditional dependencies between variables.
- Bayesian approaches to structure learning maximize the Bayesian score, which is related to the probability of the graph structure given a dataset.
- The Bayesian score promotes simpler structures for smaller datasets and supports more complicated structures for larger datasets.
- The number of possible structures is superexponential in the number of variables, and finding a structure that maximizes the Bayesian score is NP-hard.
• Directed graph search algorithms like K2 and local search can be efficient but do not guarantee optimality.

• Methods like partially directed graph search traverse the space of Markov equivalence classes, which may be more efficient than searching the larger space of directed acyclic graphs.

5.6 Exercises

Exercise 5.1. How many neighbors does an edgeless directed acyclic graph with \( m \) nodes have?

Solution: Of the three basic graph operations, we can only add edges. We can add any edge to an edgeless directed acyclic graph and it will remain acyclic. There are \( m(m - 1) = m^2 - m \) node pairs, and therefore that many neighbors.

Exercise 5.2. How many networks are in the neighborhood of the following Bayesian network?

\[
\begin{array}{c}
A \\
\ \ \ \downarrow \\
C \\
\ \ \ \ \ \ \ \ |
\end{array}
\begin{array}{c}
B \\
\ \ \ \ \ \ \ |
\ \ \ \ \ \ \ |
\ \ \ \ \ \ \ |
\ \ \ \ \ \ \ |
C \\
\ \ \ \ \ \ \ |
D
\end{array}
\]

Solution: We can perform the following graph operations:

• add \( A \to D, D \to A, D \to C \)
• remove \( A \to B, A \to C, B \to C, D \to B \)
• flip \( A \to B, B \to C, D \to B \)

Thus, there are 10 Bayesian networks in the neighborhood.

Exercise 5.3. Suppose we start local search with Bayesian network \( G \). What is the fewest number of iterations of local search that could be performed to converge to the optimal Bayesian network \( G^* \)?

\[
G \quad \begin{array}{c}
A \\
\ \ \ \ \ \ \ |
\ \ \ \ \ \ \ |
\ \ \ \ \ \ \ |
\ \ \ \ \ \ \ |
B \\
\ \ \ \ \ \ \ |
C \\
\ \ \ \ \ \ \ |
\ \ \ \ \ \ \ |
\ \ \ \ \ \ \ |
D \\
\ \ \ \ \ \ \ |
E \\
\ \ \ \ \ \ \ |
F
\end{array}
\]

\[
G^* \quad \begin{array}{c}
A \\
\ \ \ \ \ \ \ |
\ \ \ \ \ \ \ |
\ \ \ \ \ \ \ |
\ \ \ \ \ \ \ |
B \\
\ \ \ \ \ \ \ |
C \\
\ \ \ \ \ \ \ |
\ \ \ \ \ \ \ |
\ \ \ \ \ \ \ |
D \\
\ \ \ \ \ \ \ |
E \\
\ \ \ \ \ \ \ |
F
\end{array}
\]
Solution: At each iteration, local search can move from the original network to a network in its neighborhood, which is at most one edge operation from the original network. Since there are three differences between the edges of $G$ and $G^*$, performing local search from $G$ would require a minimum of three iterations to arrive at $G^*$. One potential minimal sequence of local search iterations could be flipping $A \rightarrow B$, removing $B \rightarrow C$, and adding $E \rightarrow D$. We assume the graphs formed with these edge operations yielded the highest Bayesian scores of all graphs in the considered neighborhood.

Exercise 5.4. Draw the partially directed acyclic graph representing the Markov equivalence class of the following Bayesian network. How many graphs are in this Markov equivalence class?

![Diagram](A B C D E F)

Solution: The Markov equivalence class can be represented with the following partially directed acyclic graph:

![Diagram](A B C D E F)

There are six networks in this Markov equivalence class, which are shown below:

![Networks](A B C D E F)

Exercise 5.5. Give an example of a partially directed acyclic graph with four nodes that does not define a (non-empty) Markov equivalence class.

Solution: Consider the following partially directed acyclic graph:
We cannot replace the undirected edge with a directed edge because doing so would introduce a new v-structure.
6 Simple Decisions

This chapter introduces the notion of simple decisions, where we make a single decision under uncertainty.\(^1\) We will study the problem of decision making from the perspective of utility theory, which involves modeling the preferences of an agent as a real-valued function over uncertain outcomes.\(^2\) This chapter begins by discussing how a small set of constraints on rational preferences can lead to the existence of a utility function. This utility function can be inferred from a sequence of preference queries. We then introduce the maximum expected utility principle as a definition of rationality, a central concept in decision theory that will be used as a driving principle for decision making in this book.\(^3\) We show how decision problems can be represented as decision networks and show an algorithm for solving for an optimal decision. The concept of value of information is introduced, which measures the utility gained through observing additional variables. The chapter concludes with a brief discussion of how human decision making is not always consistent with the maximum expected utility principle.

6.1 Constraints on Rational Preferences

We began our discussion on uncertainty in chapter 2 by identifying the need to compare our degree of belief between different statements. This chapter requires the ability to compare the degree of desirability of two different outcomes. We state our preferences using the following operators:

- \( A \succ B \) if we prefer \( A \) over \( B \).
- \( A \sim B \) if we are indifferent between \( A \) and \( B \).
- \( A \succeq B \) if we prefer \( A \) over \( B \) or are indifferent.

---

\(^1\) Simple decisions are simple in contrast to sequential problems, which are the focus of the rest of the book. Simple decisions are not necessarily simple to solve.


\(^3\) A textbook surveying the field of decision theory is provided by M. Peterson, An Introduction to Decision Theory. Cambridge University Press, 2009.
Just as beliefs can be subjective, so can preferences. In addition to comparing events, our preference operators can be used to compare preferences over uncertain outcomes. A lottery is a set of probabilities associated with a set of outcomes. For example, if $S_{1:n}$ is a set of outcomes and $p_{1:n}$ are their associated probabilities, then the lottery involving these outcomes and probabilities is written

$$[S_1 : p_1; \ldots ; S_n : p_n]$$

(6.1)

The existence of a real-valued measure of utility emerges from a set of assumptions about preferences. From this utility function, it is possible to define what it means to make rational decisions under uncertainty. Just as we imposed a set of constraints on beliefs, we will impose some constraints on preferences:

- **Completeness.** Exactly one of the following holds: $A \succ B$, $B \succ A$, or $A \sim B$.
- **Transitivity.** If $A \succeq B$ and $B \succeq C$, then $A \succeq C$.
- **Continuity.** If $A \succeq C \succeq B$, then there exists a probability $p$ such that $[A : p; B : 1 - p] \sim C$.
- **Independence.** If $A \succ B$, then for any $C$ and probability $p$, $[A : p; C : 1 - p] \succeq [B : p; C : 1 - p]$.

These are constraints on rational preferences. They say nothing about the preferences of actual human beings; in fact, there is strong evidence that humans are not always rational (section 6.7). Our objective in this book is to understand rational decision making from a computational perspective so that we can build useful systems. The possible extension of this theory to understanding human decision making is only of secondary interest.

### 6.2 Utility Functions

Just as constraints on the comparison of plausibility of different statements lead to the existence of a real-valued probability measure, constraints on rational preferences lead to the existence of a real-valued utility measure. It follows from our constraints on rational preferences that there exists a real-valued utility function $U$ such that


• \( U(A) > U(B) \) if and only if \( A \succ B \), and

• \( U(A) = U(B) \) if and only if \( A \sim B \).

The utility function is unique up to a positive affine transformation. In other words, for any constants \( m > 0 \) and \( b \), \( U'(S) = mU(S) + b \) if and only if the preferences induced by \( U' \) are the same as \( U \). Utilities are like temperatures: you can compare temperatures using Kelvin, Celsius, or Fahrenheit, all of which are affine transformations of each other.

It follows from the constraints on rational preferences that the utility of a lottery is given by

\[
U([S_1 : p_1; \ldots; S_n : p_n]) = \sum_{i=1}^{n} p_i U(S_i) \tag{6.2}
\]

Example 6.1 applies this equation to compute the utility of outcomes involving a collision avoidance system.

Example 6.1. An example of a lottery involving the outcomes of a collision avoidance system.

Suppose we are building a collision avoidance system. The outcome of an encounter of an aircraft is defined by whether the system alerts (\( A \)) and whether a collision occurs (\( C \)). Because \( A \) and \( C \) are binary, there are four possible outcomes. So long as our preferences are rational, we can write our utility function over the space of possible lotteries in terms of four parameters: \( U(a^0, c^0) \), \( U(a^1, c^0) \), \( U(a^0, c^1) \), and \( U(a^1, c^1) \). For example,

\[
U([a^0, c^0 : 0.5; a^1, c^0 : 0.3; a^0, c^1 : 0.1; a^1, c^1 : 0.1])
\]

is equal to

\[
0.5U(a^0, c^0) + 0.3U(a^1, c^0) + 0.1U(a^0, c^1) + 0.1U(a^1, c^1)
\]

If the utility function is bounded, then we can define a normalized utility function where the best possible outcome is assigned utility 1 and the worst possible outcome is assigned utility 0. The utility of each of the other outcomes is scaled and translated as necessary.
6.3 Utility Elicitation

In building a decision making or decision support system, it is often helpful to infer the utility function from a human or a group of humans. This approach is called utility elicitation or preference elicitation. One way to go about doing this is to fix the utility of the worst outcome $S$ to 0 and the best outcome $\bar{S}$ to 1. So long as the utilities of the outcomes are bounded, we can translate and scale the utilities without altering our preferences. If we want to determine the utility of outcome $S$, then we determine the probability $p$ such that $S \sim [\bar{S} : p; S : 1 - p]$. It then follows that $U(S) = p$. Example 6.2 applies this process to determine the utility function associated with a collision avoidance problem.

Example 6.2. Utility elicitation applied to collision avoidance.

It may be tempting to use monetary values to infer utility functions. For example, if we are building a decision support system for managing wildfires, it may be tempting to define a utility function in terms of the monetary cost incurred by property damage and the monetary cost for deploying fire suppression resources. However, it is well known in economics that the utility of wealth, in general, is not linear. If there were a linear relationship between utility and wealth, then decisions should be made in terms of maximizing expected monetary value. Someone who tries to maximize expected monetary value would have no use for insurance because the expected monetary values of insurance policies are generally negative.

Instead of trying to maximize expected wealth, we generally want to maximize the expected utility of wealth. Of course, different people have different utility functions. Figure 6.1 shows an example utility function. For small amounts of wealth, the curve is roughly linear where $100 is about twice as good as $50. For larger amounts of wealth, the curve tends to flatten out; after all, $1000 is worth

---


less to a billionaire than it is to the average person. The flattening of the curve is
sometimes referred to as diminishing marginal utility.

When discussing monetary utility functions, the three terms below are often
used. To illustrate, assume $A$ represents being given $\$50$ and $B$ represents a $50\%$
chance of winning $\$100$.

- **Risk neutral.** The utility function is linear. There is no preference between $\$50$
  and the $50\%$ chance of winning $\$100$ ($A \sim B$).

- **Risk seeking.** The utility function is convex. There is a preference for the $50\%$
  chance of winning $\$100$ ($A \prec B$).

- **Risk averse.** The utility function is concave. There is a preference for the $\$50$
  ($A \succ B$).

There are several common functional forms for modeling risk aversion of scalar
quantities, such as wealth or the availability of hospital beds. A very simple form
is exponential utility where

$$U(x) = 1 - e^{-\lambda x}$$

(6.3)

with $\lambda > 0$. Although it has a convenient mathematical form, it is generally not
viewed as a plausible model of the utility of wealth. An alternative is the power
utility where

$$U(x) = \frac{x^{1-\lambda} - 1}{1-\lambda}$$

(6.4)

with $\lambda \geq 0$ and $\lambda \neq 1$. The logarithmic utility

$$U(x) = \log x$$

(6.5)

can be viewed as a special case of the power utility where $\lambda \to 1$. Figure 6.2 shows
a plot of the power utility function with the logarithmic utility as a special case.

### 6.4 Maximum Expected Utility Principle

We are interested in the problem of making rational decisions with imperfect
knowledge of the state of the world. Suppose we have a probabilistic model
$P(s' | o, a)$, which represents the probability that the state of the world becomes
$s'$ given that we observe $o$ and take action $a$. We have a utility function $U(s')$ that
encodes our preferences over the space of outcomes. Our expected utility of taking action \( a \) given observation \( o \) is given by

\[
EU(a \mid o) = \sum_{s'} P(s' \mid a, o) U(s')
\]  

(6.6)

The principle of maximum expected utility says that a rational agent should choose the action that maximizes expected utility:

\[
a^* = \arg \max_a EU(a \mid o)
\]  

(6.7)

Because we are interested in building rational agents, equation (6.7) plays a central role in this book. Example 6.3 applies this principle to a simple decision problem.

6.5 Decision Networks

A decision network, sometimes called an influence diagram, is a generalization of a Bayesian network to include action and utility nodes so that we may compactly represent the probability and utility models defining a decision problem. The state, action, and observation spaces in the previous section may be factored, and the structure of a decision network captures the relationships between the various components.

Decision networks are composed of three types of nodes:

- A chance node corresponds to a random variable (indicated by a circle).
- A decision node corresponds to a decision variable (indicated by a square).
- A utility node corresponds to a utility variable (indicated by a diamond), and cannot have children.

There are three kinds of directed edges:

- A conditional edge ends in a chance node and indicates that the uncertainty in that chance node is conditioned on the values of all of its parents.
- An informational edge ends in a decision node and indicates that the decision associated with that node is made with knowledge of the values of its parents. (These edges are often drawn with dashed lines and are sometimes omitted from diagrams for simplicity.)

\[\text{© 2022 Massachusetts Institute of Technology, shared under a a Creative Commons CC-BY-NC-ND license. 2021-03-07 23:12:00-08:00, revision 80bf4d6, comments to bugs@algorithmsbook.com}\]
Suppose we are trying to decide whether to bring an umbrella on our vacation given the weather forecast for our destination. We observe the forecast $o$, which may be either rain or sun. Our action $a$ is either to bring our umbrella or leave our umbrella. The resulting state $s'$ is a combination of whether we brought our umbrella and whether there is sun or rain at our destination. Our probabilistic model is as follows:

| $o$       | $a$       | $s'$        | $P(s' | a, o)$ |
|-----------|-----------|-------------|---------------|
| forecast rain | bring umbrella | rain with umbrella | 0.9           |
| forecast rain | leave umbrella | rain without umbrella | 0.9          |
| forecast rain | bring umbrella | sun with umbrella | 0.1           |
| forecast rain | leave umbrella | sun without umbrella | 0.1          |
| forecast sun  | bring umbrella | rain with umbrella | 0.2           |
| forecast sun  | leave umbrella | rain without umbrella | 0.2          |
| forecast sun  | bring umbrella | sun with umbrella | 0.8           |
| forecast sun  | leave umbrella | sun without umbrella | 0.8          |

As shown in the table, we assume that our forecast is imperfect; rain forecasts are right 90% of the time and sun forecasts are right 80% of the time. In addition, we assume that bringing an umbrella does not affect the weather, though some may question this assumption. The utility function is as follows:

<table>
<thead>
<tr>
<th>$s'$</th>
<th>$U(s')$</th>
</tr>
</thead>
<tbody>
<tr>
<td>rain with umbrella</td>
<td>−0.1</td>
</tr>
<tr>
<td>rain without umbrella</td>
<td>−1</td>
</tr>
<tr>
<td>sun with umbrella</td>
<td>0.9</td>
</tr>
<tr>
<td>sun without umbrella</td>
<td>1</td>
</tr>
</tbody>
</table>

We can compute the expected utility of bringing our umbrella if we forecast rain using equation (6.6):

$$EU(\text{bring umbrella} \mid \text{forecast rain}) = 0.9 \times -0.1 + 0.1 \times 0.9 = 0$$

Likewise, we can compute the expected utility of leaving our umbrella if we forecast rain using equation (6.6):

$$EU(\text{leave umbrella} \mid \text{forecast rain}) = 0.9 \times -1 + 0.1 \times 1 = -0.8$$

Hence, we will want to bring our umbrella.
• A *functional edge* ends in a utility node and indicates that the utility node is
determined by the outcomes of its parents.

Like Bayesian networks, decision networks cannot have cycles. The utility associ-
ated with an action is equal to the sum of the values at all the utility nodes.
Example 6.4 illustrates how a decision network can model the problem of whether
to treat a disease given the results of diagnostic tests.

We have a set of results from diagnostic tests that may indicate the presence of
a particular disease. Given what is known about the tests, we need to decide
whether to apply a treatment. The utility is a function of whether a treatment
is applied and whether the disease is actually present. Conditional edges
connect $D$ to $O_1$, $O_2$, and $O_3$. Informational edges are not shown explicitly in
the figure, but they would connect the observations to $T$. Functional edges
connect $T$ and $D$ to $U$.

<table>
<thead>
<tr>
<th>$T$</th>
<th>$D$</th>
<th>$U(T,D)$</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>0</td>
<td>1</td>
<td>-10</td>
</tr>
<tr>
<td>1</td>
<td>0</td>
<td>-1</td>
</tr>
<tr>
<td>1</td>
<td>1</td>
<td>-1</td>
</tr>
</tbody>
</table>

Solving a simple problem (algorithm 6.1) requires iterating over all possible
decision instantiations to find a decision that maximizes expected utility. For each
instantiation, we evaluate the associated expected utility. We begin by instantiating
the action nodes and observed chance nodes. We can then apply any inference
algorithm to compute the posterior over the inputs to the utility function. The
expected utility is the sum of the values at the utility nodes. Example 6.5 shows
how this process can be applied to our running example.

A variety of methods have been developed to make evaluating decision net-
works more efficient. Early papers by Shachter provide
algorithms for evaluating decision
networks. R.D. Shachter, “Eval-
uating Influence Diagrams,” Op-
erations Research, vol. 34, no. 6,
pp. 871–882, 1986. R.D. Shachter,
“Probabilistic Inference and Influ-
ence Diagrams,” Operations Re-
search, vol. 36, no. 4, pp. 589–604,

---

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Algorithm 6.1. A definition of a simple problem as a decision network. A decision network is defined by a Bayesian network with sets of chance, decision, and utility variables. Utility variables are generally treated as deterministic variables. Because variables in our Bayesian network data structure take on values from $1 : r_i$, the utilities variables are mapped to real values by the `utilities` field. For example, if we have a utility variable $u_1$, the $i$th utility associated with that variable is `utilities[u1][i]`. The `solve` function takes as input the problem, evidence, and an inference method. It returns a tuple indicating the best assignment to the decision variables and its associated expected utility.

<table>
<thead>
<tr>
<th>struct SimpleProblem</th>
</tr>
</thead>
<tbody>
<tr>
<td>bn::BayesianNetwork</td>
</tr>
<tr>
<td>chance_vars::Vector{Variable}</td>
</tr>
<tr>
<td>decision_vars::Vector{Variable}</td>
</tr>
<tr>
<td>utility_vars::Vector{Variable}</td>
</tr>
<tr>
<td>utilities::Dict{Symbol, Vector{Float64}}</td>
</tr>
</tbody>
</table>

function solve(P::SimpleProblem, evidence, M)
    query = [var.name for var in P.utility_vars]
    U(a) = sum(P.utilities[uname][a[uname]] for uname in query)
    best = (a=nothing, u=-Inf)
    for assignment in assignments(P.decision_vars)
        evidence = merge(evidence, assignment)
        φ = infer(M, P.bn, query, evidence)
        u = sum(p*U(a) for (a, p) in φ.table)
        if u > best.u
            best = (a=assignment, u=u)
        end
    end
    return best
end

6.6 Value of Information

We make our decisions based on what we observe. In many applications, it is natural to want to quantify the value of information, which is how much observing additional variables is expected to increase our utility.\footnote{R.A. Howard, “Information Value Theory,” IEEE Transactions on Systems Science and Cybernetics, vol. 2, no. 1, pp. 22–26, 1966.} For example, in the disease treatment application in example 6.5, we assumed we have only observed $o_1^1$. Given the positive result from that one diagnostic test alone, we may decide against treatment. However, it may be beneficial to administer additional diagnostic tests to reduce the risk of not treating a disease that is really present.

In computing the value of information, we will use $EU^*(o)$ to denote the expected utility of an optimal action given observation $o$. The value of information...
We can use equation (6.6) to compute the expected utility of treating a disease for the decision network in example 6.4. For now, we will assume that we have the result from only the first diagnostic test and that it came back positive. If we wanted to make the knowledge of the first diagnostic test explicit in the diagram, then we would draw an informational edge from $O_1$ to $T$, and we would have

$$EU(t^1 | o^1_1) = \sum_{o_3} \sum_{o_2} \sum_d P(d, o_2, o_3 | t^1, o^1_1) U(t^1, d, o^1_1, o_2, o_3)$$

We can use the chain rule for Bayesian networks and the definition of conditional probability to compute $P(d, o_2, o_3 | t^1, o^1_1)$. Because the utility node depends only on whether the disease is present and whether we treat it, we can simplify $U(t^1, d, o^1_1, o_2, o_3)$ to $U(t^1, d)$. Hence,

$$EU(t^1 | o^1_1) = \sum_d P(d | t^1, o^1_1) U(t^1, d)$$

Any of the exact or approximate inference methods introduced in the previous chapter can be used to evaluate $P(d | t^1, o^1_1)$. To decide whether to apply a treatment, we compute $EU(t^1 | o^1_1)$ and $EU(t^0 | o^1_1)$ and make the decision that provides the highest expected utility.
about variable $O'$ given $o$ is

$$VOI(O' \mid o) = \left( \sum_{o'} P(o' \mid o) EU^*(o,o') \right) - EU^*(o)$$  \hspace{1cm} (6.8)$$

In other words, the value of information about a variable is the increase in expected utility if that variable is observed. Algorithm 6.2 provides an implementation.

Algorithm 6.2. A method for decision network evaluation, which takes a simple problem $\mathcal{P}$, a dictionary containing observed chance variables and their values $\text{evidence}$, and an inference strategy $M$. The method returns an assignment of decision values that maximizes the expected utility given the evidence.

The value of information is never negative. The expected utility can only increase if additional observations can lead to different optimal decisions. If observing a new variable $O'$ makes no difference in the choice of action, then $EU^*(o,o') = EU^*(o)$ for all $o'$, in which case equation (6.8) evaluates to 0. For example, if the optimal decision is to treat the disease regardless of the outcome of the diagnostic test, then the value of observing the outcome of the test is 0.

The value of information only captures the increase in expected utility from making an observation. There may be a cost associated with making a particular observation. Some diagnostic tests may be inexpensive, such as a temperature reading; other diagnostic tests are more costly and invasive, such as a lumbar puncture. The value of information obtained by a lumbar puncture may be much greater than a temperature reading, but the costs of the tests should be taken into consideration.

The value-of-information metric is an important and often used metric for choosing what to observe. Sometimes the value-of-information metric is used to determine an appropriate sequence of observations. After each observation, the value of information is determined for the remaining unobserved variables. The unobserved variable with the greatest value of information is then selected for observation. If there are costs associated with making different observations, then
these costs are subtracted from the value of information when determining which variable to observe. The process continues until it is no longer beneficial to observe any more variables. The optimal action is then chosen. This greedy selection of observations is only a heuristic and may not represent the truly optimal sequence of observations. The optimal selection of observations can be determined by using the techniques for sequential decision making introduced in later chapters.

6.7 Irrationality

Decision theory is a normative theory that is prescriptive, not a descriptive theory that is predictive of human behavior. Human judgment and preference often do not follow the rules of rationality outlined in section 6.1. Even human experts may have an inconsistent set of preferences, which can be problematic when designing a decision support system that attempts to maximize expected utility. Example 6.6 shows that certainty often exaggerates losses that are certain relative to losses that are merely probable. This certainty effect works with gains as well. A smaller gain that is certain is often preferred over a much greater gain that is only probable, in a way that the axioms of rationality are necessarily violated.

Example 6.7 demonstrates the framing effect, where people decide on options based on whether they are presented as a loss or as a gain. Many other cognitive biases can lead to deviations from what is prescribed by utility theory. Special care must be given when trying to elicit utility functions from human experts to build decision support systems. Although the recommendations of the decision support system may be rational, they may not exactly reflect human preferences in certain situations.

6.8 Summary

- Rational decision making combines probability and utility theory.
- The existence of a utility function follows from constraints on rational preferences.
- A rational decision is one that maximizes expected utility.
- Decision problems can be modeled using decision networks, which are extensions of Bayesian networks that include actions and utilities.

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2021-03-07 23:12:00-08:00, revision 80bf4d6, comments to bugs@algorithmsbook.com


Tversky and Kahneman studied the preferences of university students who answered questionnaires in a classroom setting. They presented students with questions dealing with the response to an epidemic. The students were to reveal their preference between the following two outcomes:

- \( A \): 100\% chance of losing 75 lives
- \( B \): 80\% chance of losing 100 lives

Most preferred \( B \) over \( A \). From equation (6.2), we know

\[
U(\text{lose 75}) < 0.8U(\text{lose 100}) \tag{6.9}
\]

They were then asked to choose between the following two outcomes:

- \( C \): 10\% chance of losing 75 lives
- \( D \): 8\% chance of losing 100 lives

Most preferred \( C \) over \( D \). Hence, \( 0.1U(\text{lose 75}) > 0.08U(\text{lose 100}) \). We multiply both sides by 10 and get

\[
U(\text{lose 75}) > 0.8U(\text{lose 100}) \tag{6.10}
\]

Of course, equations (6.9) and (6.10) result in a contradiction. We have made no assumption about the actual value of \( U(\text{lose 75}) \) and \( U(\text{lose 100}) \)—we did not even assume that losing 100 lives was worse than losing 75 lives. Because equation (6.2) follows directly from the von Neumann-Morgenstern axioms in section 6.1, there must be a violation of at least one of the axioms—even though many people who select \( B \) and \( C \) seem to find the axioms agreeable.
Tversky and Kahneman demonstrated the *framing effect* using a hypothetical scenario in which an epidemic is expected to kill 600 people. They presented students with the following two outcomes:

- $E$: 200 people will be saved
- $F$: $\frac{1}{3}$ chance that 600 people will be saved and $\frac{2}{3}$ chance that no people will be saved

The majority of students chose $E$ over $F$. They then asked them to choose between:

- $G$: 400 people will die
- $H$: $\frac{1}{3}$ chance that nobody will die and $\frac{2}{3}$ chance that 600 people will die

The majority of students chose $H$ over $G$, even though $E$ is equivalent to $G$ and $F$ is equivalent to $H$. This inconsistency is due to how the question is framed.
6.9. exercises

- Solving a simple decision involves inference in Bayesian networks, and is thus NP-hard.
- The value of information measures the gain in expected utility should a new variable be observed.
- Humans are not always rational.

6.9 Exercises

Exercise 6.1. Suppose we have a utility function $U(s)$ with a finite maximum value $\overline{U}$ and a finite minimum value $\underline{U}$. What is the corresponding normalized utility function $\hat{U}(s)$ that preserves the same preferences?

Solution: A normalized utility function has a maximum value of 1 and a minimum value of 0. Preferences are preserved under affine transforms, so we determine the affine transform of $U(s)$ that matches the unit bounds. This transform is:

$$\hat{U}(s) = \frac{U(s) - \underline{U}}{\overline{U} - \underline{U}} = \frac{1}{\overline{U} - \underline{U}}U(s) - \frac{\underline{U}}{\overline{U} - \underline{U}}$$

Exercise 6.2. If $A \succeq C \succeq B$ and the utilities of each outcome are $U(A) = 450$, $U(B) = -150$, $U(C) = 60$, what is the lottery over $A$ and $B$ that will make us indifferent between the lottery and $C$?

Solution: A lottery over $A$ and $B$ is defined as $[A : p; B : 1 - p]$. In order to satisfy indifference between the lottery and $C$ ($[A : p; B : 1 - p] \sim C$), we must have $U([A : p; B : 1 - p]) = U(C)$. Thus, we must compute $p$ that satisfies the equality

$$U([A : p; B : 1 - p]) = U(C)$$
$$pU(A) + (1 - p)U(B) = U(C)$$

$$p = \frac{U(C) - U(B)}{U(A) - U(B)} = \frac{60 - (-150)}{450 - (-150)} = 0.35$$

This implies that the lottery $[A : 0.35; B : 0.65]$ is equally as desired as $C$.

Exercise 6.3. Suppose for a utility function $U$ over three outcomes $A$, $B$, and $C$ that $U(A) = 5$, $U(B) = 20$, and $U(C) = 0$. We are given a choice between a lottery that gives us a 50% probability of $B$ and a 50% probability of $C$ and a lottery that guarantees $A$. Compute the preferred lottery and show that under the positive affine transformation with $m = 2$ and $b = 30$ that we maintain preference for the same lottery.
Solution: The first lottery is given by \([A : 0.0; B : 0.5; C : 0.5]\) and the second lottery is given by \([A : 1.0; B : 0.0; C : 0.0]\). The original utilities for each lottery are given by
\[
U([A : 0.0; B : 0.5; C : 0.5]) = 0.0U(A) + 0.5U(B) + 0.5U(C) = 10
\]
\[
U([A : 1.0; B : 0.0; C : 0.0]) = 1.0U(A) + 0.0U(B) + 0.0U(C) = 5
\]
Thus, since \(U([A : 0.0; B : 0.5; C : 0.5]) > U([A : 1.0; B : 0.0; C : 0.0])\), we prefer the first lottery. Under the positive affine transformation \(m = 2, b = 30\), our new utilities can be computed as
\[
U'(A) = 40, U'(B) = 70, U'(C) = 30
\]
Exercise 6.4. Prove that the power utility function in equation (6.4) is risk averse for all \(x > 0\) and \(\lambda > 0\) with \(\lambda \neq 1\).

Solution: Risk aversion implies that the utility function is concave, which requires that the second derivative of the utility function is negative. The utility function and its derivatives are computed as follows
\[
U(x) = \frac{x^{1-\lambda} - 1}{1 - \lambda}
\]
\[
\frac{dU}{dx} = \frac{1}{x^\lambda}
\]
\[
\frac{d^2U}{dx^2} = -\lambda x^{\lambda - 1}
\]
For \(x > 0\) and \(\lambda > 0\), \(\lambda \neq 1\), \(x^{\lambda + 1}\) is a positive number raised to a positive exponent, which is guaranteed to be positive. Multiplying this by \(-\lambda\) guarantees that the second derivative is negative. Thus, for all \(x > 0\) and \(\lambda > 0\), \(\lambda \neq 1\), the power utility function is risk averse.

Exercise 6.5. Using the parameters given in example 6.3, compute the expected utility of bringing our umbrella if we forecast sun and the expected utility of leaving our umbrella if we forecast sun. What is the action that maximizes our expected utility given we forecast sun?

Solution:
\[
EU\text{bring umbrella | forecast sun} = 0.2 \times -0.1 + 0.8 \times 0.9 = 0.7
\]
\[
EU\text{leave umbrella | forecast sun} = 0.2 \times -1.0 + 0.8 \times 1.0 = 0.6
\]
The action that maximizes our expected utility if we forecast sun is to bring our umbrella!
**Exercise 6.6.** Suppose we are trying to optimally decide whether or not to feed \((F)\) our new puppy based on the likelihood that the puppy is hungry \((H)\). We can observe whether or not the puppy is whining \((W)\) and whether or not someone else has recently fed the puppy \((R)\). The utilities of each combination of feeding and hunger and the decision network representation are provided below.

<table>
<thead>
<tr>
<th>(F)</th>
<th>(H)</th>
<th>(U(F, H))</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>0</td>
<td>0.0</td>
</tr>
<tr>
<td>0</td>
<td>1</td>
<td>−1.0</td>
</tr>
<tr>
<td>1</td>
<td>0</td>
<td>−0.5</td>
</tr>
<tr>
<td>1</td>
<td>1</td>
<td>−0.1</td>
</tr>
</tbody>
</table>

Given that \(P(h^1 \mid w^1) = 0.8\), if we observe the puppy whining \((w^1)\), what are the expected utilities of not feeding the puppy \((f^0)\) and feeding the puppy \((f^1)\)? What is the optimal action?

**Solution:** We start with the definition of expected utility and recognize that the utility depends only on \(H\) and \(F\):

\[
EU(f^0 \mid w^1) = \sum_h P(h \mid w^1)U(f^0, h)
\]

Now, we can compute the expected utility of feeding the puppy given it is whining and, in a similar fashion as above, the expected utility of not feeding the puppy given it is whining

\[
EU(f^0 \mid w^1) = 0.2 \times 0.0 + 0.8 \times −1.0 = −0.8
\]
\[
EU(f^1 \mid w^1) = 0.2 \times −0.5 + 0.8 \times −0.1 = −0.18
\]

Thus, the optimal action is to feed the puppy \((f^1)\) since this maximizes our expected utility \(EU^*(w^1) = −0.18\).

**Exercise 6.7.** Using the results from exercise 6.6, if \(P(r^1 \mid w^1) = 0.2\), \(P(h^1 \mid w^1, r^0) = 0.9\), and \(P(h^1 \mid w^1, r^1) = 0.3\), what is the value of information of asking someone else if the puppy has recently been fed given that we observe the puppy to be whining \((w^1)\)?

**Solution:** We are interested in computing

\[
VOI(R \mid w^1) = \left(\sum_r P(r \mid w^1)EU^*(w^1, r)\right) - EU^*(w^1)
\]
We start by computing $EU(f \mid w^1, r)$ for all $f$ and $r$. Following a similar derivation as in exercise 6.6, we have

$$EU(f^0 \mid w^1, r^0) = \sum_h P(h \mid w^1, r^0)U(f^0, h)$$

So, for each combination of $F$ and $R$, we have the following expected utilities

$$EU(f^0 \mid w^1, r^0) = \sum_h P(h \mid w^1, r^0)U(f^0, h) = 0.1 \times 0.0 + 0.9 \times -1.0 = -0.9$$

$$EU(f^1 \mid w^1, r^0) = \sum_h P(h \mid w^1, r^0)U(f^1, h) = 0.1 \times -0.5 + 0.9 \times -0.1 = -0.14$$

$$EU(f^0 \mid w^1, r^1) = \sum_h P(h \mid w^1, r^1)U(f^0, h) = 0.7 \times 0.0 + 0.3 \times -1.0 = -0.3$$

$$EU(f^1 \mid w^1, r^1) = \sum_h P(h \mid w^1, r^1)U(f^1, h) = 0.7 \times -0.5 + 0.3 \times -0.1 = -0.38$$

The optimal expected utilities are

$$EU^*(w^1, r^0) = -0.14$$

$$EU^*(w^1, r^1) = -0.3$$

Now, we can compute the value of information

$$VOI(R \mid w^1) = 0.8(-0.14) + 0.2(-0.3) - (-0.18) = 0.008$$
PART II:

SEQUENTIAL PROBLEMS

Up to this point, we have assumed that we make a single decision at one point in time, but many important problems require that we make a series of decisions. The same principle of maximum expected utility still applies, but optimal decision making in a sequential context requires reasoning about future sequences of actions and observations. This part of the book will discuss sequential decision problems in stochastic environments. We will focus on a general formulation of sequential decision problems under the assumption that the model is known and that the environment is fully observable. We will relax both of these assumptions later. Our discussion will begin with the introduction of the Markov decision process, the standard mathematical model for sequential decision problems. We will discuss several approaches for finding exact solutions. Because large problems sometimes do not permit exact solutions to be efficiently found, we will discuss a collection of both offline and online approximate solution methods along with a type of method that involves directly searching the space of parameterized decision policies. Finally, we will discuss approaches for validating that our decision strategies will perform as expected when deployed in the real world.
7 Exact Solution Methods

This chapter introduces a model known as a Markov decision process (MDP) to represent sequential decision problems where the effects of our actions are uncertain. We begin with a description of the model, which specifies both the stochastic dynamics of the system as well as the utility associated with its evolution. Different algorithms can be used to compute the utility associated with a decision strategy and to search for an optimal strategy. Under certain assumptions, we can find exact solutions to Markov decision processes. Later chapters will discuss approximation methods that tend to scale better to larger problems.

7.1 Markov Decision Processes

In an MDP (algorithm 7.1), we choose action $a_t$ at time $t$ based on observing state $s_t$. We then receive a reward $r_t$. The action space $A$ is the set of possible actions, and the state space $S$ is the set of possible states. Some of the algorithms assume these sets are finite, but this is not required in general. The state evolves probabilistically based on the current state and action we take. The assumption that the next state depends only on the current state and action and not on any prior state or action is known as the Markov assumption.

An MDP can be represented using a decision network as shown in figure 7.1. There are information edges (not shown in the figure) from $A_{1:t-1}$ and $S_{1:t}$ to $A_t$. The utility function is decomposed into rewards $R_{1:t}$. We focus on stationary MDPs in which $P(S_{t+1} \mid S_t, A_t)$ and $P(R_t \mid S_t, A_t)$ do not vary with time. Stationary MDPs can be compactly represented by a dynamic decision diagram as shown in figure 7.2. The state transition model $T(s' \mid s, a)$ represents the probability of transitioning from state $s$ to $s'$ after executing action $a$. The reward function $R(s, a)$ represents the expected reward received when executing action $a$ from state $s$. 

\[ \text{Figure 7.1. Markov decision process decision diagram.} \]

\[ \text{Figure 7.2. Stationary Markov decision process decision diagram.} \]
The reward function is a deterministic function of $s$ and $a$ because it represents an expectation, but rewards may be generated stochastically in the environment or even depend upon the resulting next state.\(^2\) Example 7.1 shows how to frame a collision avoidance problem as an MDP.

The problem of aircraft collision avoidance can be formulated as an MDP. The states represent the positions and velocities of our aircraft and the intruder aircraft, and the actions represent whether we climb, descend, or stay level. We receive a large negative reward for colliding with the other aircraft and a small negative reward for climbing or descending.

Given knowledge of the current state, we must decide whether an avoidance maneuver is required. The problem is challenging because the positions of the aircraft evolve probabilistically and we want to make sure that we start our maneuver early enough to avoid collision but late enough so that we avoid unnecessary maneuvering.

Example 7.1. Aircraft collision avoidance framed as an MDP.

Algorithm 7.1. Data structure for an MDP. We will use the TR field later to sample the next state and reward given the current state and action: $s', r = TR(s, a)$.

The rewards in an MDP are treated as components in an additively decomposed utility function. In a finite horizon problem with $n$ decisions, the utility associated with a sequence of rewards $r_{1:n}$ is simply

\[
\sum_{t=1}^{n} r_t \tag{7.1}
\]

The sum of rewards is sometimes called the return.

In an infinite horizon problem in which the number of decisions is unbounded, the sum of rewards can become infinite.\(^3\) There are several ways to define utility in terms of individual rewards in infinite horizon problems. One way is to impose

\(^2\) For example, if the reward depends on the next state as given by $R(s, a, s')$, then the (expected) reward function would be

\[
R(s, a) = \sum_{s'} T(s' \mid s, a) R(s, a, s')
\]

\(^3\) Suppose strategy $A$ results in a reward of 1 per time step and strategy $B$ results in a reward of 100 per time step. Intuitively, a rational agent should prefer strategy $B$ over strategy $A$, but both provide the same infinite expected utility.
a discount factor $\gamma$ between 0 and 1. The utility is given by

$$\sum_{t=1}^{\infty} \gamma^{t-1} r_t \quad (7.2)$$

This value is sometimes called the discounted return. So long as $0 \leq \gamma < 1$ and the rewards are finite, the utility will be finite. The discount factor makes it so that rewards in the present are worth more than rewards in the future, a concept that also appears in economics.

Another way to define utility in infinite horizon problems is to use the average reward given by

$$\lim_{n \to \infty} \frac{1}{n} \sum_{t=1}^{n} r_t \quad (7.3)$$

This average return formulation can be attractive because we do not have to choose a discount factor, but there is often not a practical difference between this formulation and discounted return with a discount factor close to 1. Because the discounted return is often computationally simpler to work with, we will focus on the discounted formulation.

A policy tells us what action to select given the past history of states and actions. The action to select at time $t$, given the history $h_t = (s_1:t, a_1:t-1)$, is written $\pi_t(h_t)$. Because the future states and rewards depend only on the current state and action (as made apparent in the conditional independence assumptions in figure 7.1), we can restrict our attention to policies that depend only on the current state. In addition, we will primarily focus on deterministic policies because, in MDPs, there is guaranteed to exist an optimal policy that is deterministic. Later chapters discuss stochastic policies, where $\pi_t(a_t \mid s_t)$ denotes the probability the policy assigns to taking action $a_t$ in state $s_t$ at time $t$.

In infinite horizon problems with stationary transitions and rewards, we can further restrict our attention to stationary policies, which do not depend on time. We will write the action associated with stationary policy $\pi$ in state $s$ as $\pi(s)$, without the temporal subscript. In finite horizon problems, however, it may be beneficial to select a different action depending on how many time steps are remaining. For example, when playing basketball, it is generally not a good strategy to attempt a half-court shot unless there are only a couple seconds remaining in the game.
The expected utility of executing $\pi$ from state $s$ is denoted $U^\pi(s)$. In the context of MDPs, $U^\pi$ is often referred to as the value function. An optimal policy $\pi^*$ is a policy that maximizes expected utility:

$$\pi^*(s) = \arg \max_\pi U^\pi(s)$$

(7.4)

for all states $s$. Depending on the model, there may be multiple policies that are optimal. The value function associated with an optimal policy $\pi^*$ is called the optimal value function and is denoted $U^*$.

An optimal policy can be found by using a computational technique called dynamic programming,\(^5\) which involves simplifying a complicated problem by breaking it down into simpler sub-problems in a recursive manner. Although we will focus on dynamic programming algorithms for MDPs, dynamic programming is a general technique that can be applied to a wide variety of other problems. For example, dynamic programming can be used in computing a Fibonacci sequence and finding the longest common subsequence between two strings.\(^6\) In general, algorithms that use dynamic programming for solving MDPs are much more efficient than brute force methods.

### 7.2 Policy Evaluation

Before we discuss how to go about computing an optimal policy, we will first discuss policy evaluation, where we compute the value function $U^\pi$. Policy evaluation can be done iteratively. If the policy is executed for a single time step, the utility is $U^\pi_1(s) = R(s, \pi(s))$. Further steps can be obtained by applying the lookahead equation:

$$U^\pi_{k+1}(s) = R(s, \pi(s)) + \gamma \sum_{s'} T(s' | s, \pi(s)) U^\pi_k(s')$$

(7.5)

This equation is implemented in algorithm 7.2. Iterative policy evaluation is implemented in algorithm 7.3. Several iterations are shown in figure 7.3.

The value function $U^\pi$ can be computed to arbitrary precision given sufficient iterations. For an infinite horizon, we have

$$U^\pi(s) = R(s, \pi(s)) + \gamma \sum_{s'} T(s' | s, \pi(s)) U^\pi(s')$$

(7.6)

\(^4\) Doing so is consistent with the maximum expected utility principle introduced in section 6.4.

\(^5\) The term dynamic programming was coined by the American mathematician Richard Ernest Bellman (1920–1984). Dynamic refers to the fact that the problem is time-varying and programming refers to a methodology to find an optimal program or decision strategy. R. Bellman, *Eye of the Hurricane: an Autobiography*. World Scientific, 1984.

Algorithm 7.2. Functions for computing the lookahead state-action value from a state \( s \) given an action \( a \) using an estimate of the value function \( U \) for the MDP \( \mathcal{P} \). The second version handles the case when \( U \) is a vector.

Algorithm 7.3. Iterative policy evaluation, which iteratively computes the value function for a policy \( \pi \) for MDP \( \mathcal{P} \) with discrete state and action spaces using \( k_{\text{max}} \) iterations.

Figure 7.3. Iterative policy evaluation used to evaluate an east-moving policy on the hex world problem (appendix F.1). The arrows indicate the direction recommended by the policy (i.e., always move east), and the colors indicate the values associated with the states. The values change with each iteration.
at convergence. Convergence can be proven using the fact that the update in equation (7.5) is a contraction mapping (reviewed in appendix A.15). 7

Policy evaluation can be done without iteration by solving a system of equations in matrix form:

\[
U^\pi = R^\pi + \gamma T^\pi U^\pi
\]  

(7.7)

where \(U^\pi\) and \(R^\pi\) are the utility and reward functions represented in vector form with \(|S|\) components. The \(|S| \times |S|\) matrix \(T^\pi\) contains state transition probabilities where \(T^\pi_{ij}\) is the probability of transitioning from the \(i\)th state to the \(j\)th state.

The value function is obtained as follows:

\[
U^\pi - \gamma T^\pi U^\pi = R^\pi
\]

(7.8)

\[
(I - \gamma T^\pi)U^\pi = R^\pi
\]

(7.9)

\[
U^\pi = (I - \gamma T^\pi)^{-1} R^\pi
\]

(7.10)

This method is implemented in algorithm 7.4. Solving for \(U^\pi\) in this way requires \(O(|S|^3)\) time. The method is used to evaluate a policy in figure 7.4.

---

7 See exercise 7.12.

---

Algorithm 7.4. Exact policy evaluation, which computes the value function for a policy \(\pi\) for an MDP \(\mathcal{P}\) with discrete state and action spaces.

Figure 7.4. Exact policy evaluation used to evaluate an east-moving policy for the hex world problem. The exact solution contains lower values than what was contained in the first few steps of iterative policy evaluation in figure 7.3. If we ran iterative policy evaluation for more iterations, it would converge to the same value function.

---

### 7.3 Value Function Policies

The previous section showed how to compute a value function associated with a policy. This section shows how to extract a policy from a value function, which we later use when generating optimal policies. Given a value function \(U\), which...
may or may not correspond to the optimal value function, we can construct a policy $\pi$ that maximizes the lookahead equation introduced in equation (7.5):

$$\pi(s) = \arg\max_a \left( R(s,a) + \gamma \sum_{s'} T(s' | s,a) U(s') \right)$$

(7.11)

We refer to this policy as a greedy policy with respect to $U$. If $U$ is the optimal value function, then the extracted policy is optimal. Algorithm 7.5 implements this idea.

An alternative way to represent a policy is to use the action value function, sometimes called the Q-function. The action value function represents the expected return when starting in state $s$, taking action $a$, and then continuing with the greedy policy with respect to $Q$:

$$Q(s,a) = R(s,a) + \gamma \sum_{s'} T(s' | s,a) U(s')$$

(7.12)

From this action value function, we can obtain the value function,

$$U(s) = \max_a Q(s,a)$$

(7.13)

as well as the policy,

$$\pi(s) = \arg\max_a Q(s,a)$$

(7.14)

Storing $Q$ explicitly for discrete problems requires $O(|S| \times |A|)$ storage instead of $O(|S|)$ storage for $U$, but we do not have to use $R$ and $T$ to extract the policy.

Policies can also be represented using the advantage function, which quantifies the advantage of taking an action in comparison to the greedy action. It is defined in terms of the difference between $Q$ and $U$:

$$A(s,a) = Q(s,a) - U(s)$$

(7.15)

Greedy actions have zero advantage, and non-greedy actions have negative advantage. Some algorithms that we will discuss later in the book use $U$ representations, but others will use $Q$ or $A$. 

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Algorithm 7.5. A value function policy extracted from a value function $U$ for an MDP $\mathcal{P}$. The `greedy` function will be used in other algorithms. This two-argument version of `findmax` is defined in appendix G.5.

7.4 Policy Iteration

Policy iteration (algorithm 7.6) is one way to compute an optimal policy. It involves iterating between policy evaluation (section 7.2) and policy improvement through lookahead (algorithm 7.5). Policy iteration is guaranteed to converge given any initial policy. It converges because there are finitely many policies and every iteration improves the policy if it is possible to be improved. Figure 7.5 demonstrates policy iteration on the hex world problem.
Policy iteration tends to be expensive because we must evaluate the policy in each iteration. A variation of policy iteration is called modified policy iteration, which approximates the value function using iterative policy evaluation instead of exact policy evaluation. A parameter in the modified policy iteration algorithm is the number of policy evaluation iterations to perform between steps of policy improvement. If this parameter is 1, then this approach is identical to value iteration.

### 7.5 Value Iteration

Value iteration is an alternative to policy iteration that is often used because of its simplicity. Unlike policy improvement, value iteration updates the value function directly. It begins with any bounded value function $U$, meaning that $|U(s)| < \infty$ for all $s$. One common initialization is $U(s) = 0$ for all $s$.

The value function can be improved by applying the Bellman equation:

$$U_{k+1}(s) = \max_a \left( R(s, a) + \gamma \sum_{s'} T(s' | s, a) U_k(s') \right)$$  \hspace{1cm} (7.16)

This backup procedure is implemented in algorithm 7.7.

```python
function backup(\mathcal{P}::MDP, U, s)
    return maximum(lookahead(\mathcal{P}, U, s, a) for a in \mathcal{P}.A)
end
```

Repeated application of this update is guaranteed to converge on an optimal policy. Like iterative policy evaluation, we can use the fact that the update is a

---


contraction mapping to prove convergence.\(^{10}\) This optimal policy is guaranteed to satisfy:

\[
U^*(s) = \max_a \left( R(s, a) + \gamma \sum_{s'} T(s' | s, a) U^*(s') \right)
\]  

(7.17)

Further applications of Bellman’s equation once this equality holds do not change the policy. An optimal policy can be extracted from \(U^*\) using equation (7.11). Value iteration is implemented in algorithm 7.8 and is applied to the hex world problem in figure 7.6.

The implementation in algorithm 7.8 stops after a fixed number of iterations, but it is also common to terminate the iterations early based on the maximum change in value called the Bellman residual, \(\|U_{k+1} - U_k\|_\infty\). If the Bellman residual drops below some threshold \(\delta\), then the iterations terminate. A Bellman residual of \(\delta\) guarantees that the optimal value function obtained by value iteration is within \(\epsilon = \delta \gamma / (1 - \gamma)\) of \(U^*\).\(^{11}\) Discount factors closer to 1 significantly inflate this error, leading to slower convergence. If we heavily discount future reward (\(\gamma\) closer to 0), then we do not need to iterate as much into the future. This effect is demonstrated in example 7.2.

Knowing the maximum deviation of the estimated value function from the optimal value function, \(\|U_k - U^*\|_\infty < \epsilon\), allows us to bound the maximum deviation of reward obtained under the extracted policy \(\pi\) from an optimal policy \(\pi^*\), the policy loss \(\|U^\pi - U^*\|_\infty\). The policy loss is bounded by \(2\epsilon\).\(^{12}\)

```plaintext
struct ValueIteration
    k_max # maximum number of iterations
end

function solve(M::ValueIteration, P::MDP)
    U = [0.0 for s in P.S]
    for k = 1:M.k_max
        U = [backup(P, U, s) for s in P.S]
    end
    return ValueFunctionPolicy(P, U)
end
```

\(^{10}\) See exercise 7.13.

\(^{11}\) See exercise 7.8.

\(^{12}\) To see why, see exercise 7.9.
Figure 7.6. Value iteration on the hex world problem to obtain an optimal policy. Each hex is colored according to the value function and arrows indicate the policy that is greedy with respect to that value function.
Consider a simple variation of the hex world problem consisting of a straight line of tiles with a single consuming tile at the end producing 10 reward. The discount factor directly affects the rate at which reward from the consuming tile propagates down the line to the other tiles, and thus how quickly value iteration converges.

$\gamma = 0.9$

$\gamma = 0.5$

Example 7.2. An example showing the effect of the discount factor on convergence of value iteration. In each case value iteration was run until the Bellman residual was less than 1.
7.6 Asynchronous Value Iteration

Value iteration tends to be computationally intensive, as every entry in the value function $U_k$ is updated in each iteration to obtain $U_{k+1}$. In asynchronous value iteration, only a subset of the states are updated in each iteration. Asynchronous value iteration is still guaranteed to converge on the optimal value function provided that each state is updated infinitely often.

One common asynchronous value iteration method, Gauss-Seidel value iteration (algorithm 7.9), sweeps through an ordering of the states and applies the Bellman update in place:

$$U(s) \leftarrow \max_a \left( R(s,a) + \gamma \sum_{s'} T(s' \mid s,a) U(s') \right)$$ (7.18)

The computational savings lies in not having to construct a second value function in memory in each iteration. Gauss-Seidel value iteration can converge more quickly than standard value iteration depending on the ordering chosen. In some problems, the state contains a time index that increments deterministically forward in time. If we apply Gauss-Seidel value iteration starting at the last time index and work our way backwards, this process is sometimes called backwards induction value iteration. An example of the impact of the state ordering is given in example 7.3.

```plaintext
struct GaussSeidelValueIteration
    k_max # maximum number of iterations
end

function solve(M::GaussSeidelValueIteration, P::MDP)
    U = [0.0 for s in S]
    for k = 1:M.k_max
        for (s, i) in enumerate(P.S)
            U[i] = backup(P, U, s)
        end
    end
    return ValueFunctionPolicy(P, U)
end
```

Algorithm 7.9. Asynchronous value iteration, which updates states in a different manner to value iteration, often saving computation time. The method terminates after $k_{\text{max}}$ iterations.

---

13 A poor ordering in Gauss-Seidel value iteration cannot cause the algorithm to be slower than standard value iteration.
Consider the linear variation of the hex world problem from example 7.2. We can solve the same problem using asynchronous value iteration. The ordering of the states directly affects the rate at which reward from the consuming tile propagates down the line to the other tiles, and thus how quickly the method converges.

Example 7.3. An example showing the effect of the state ordering on convergence of asynchronous value iteration. In this case, evaluating right to left allows for convergence in far fewer iterations.
7.7 Linear Program Formulation

The problem of finding an optimal policy can be formulated as a linear program, which is an optimization problem with a linear objective function and a set of linear equality or inequality constraints. Once a problem is represented as a linear program, we can use one of many different linear programming solvers.\footnote{For an overview of linear programming, see R. Vanderbei, *Linear Programming, Foundations and Extensions*, 4th ed. Springer, 2014.}

To show how we can convert the Bellman equation into a linear program, we begin by replacing the equality in the Bellman equation with a set of inequality constraints while minimizing $U(s)$ at each state $s$:\footnote{Intuitively, we want to push the value $U(s)$ at all states $s$ down to convert the inequality constraints into equality constraints. Hence, we minimize the sum of all utilities.}

$$\begin{align*}
\text{minimize} & \sum_s U(s) \\
\text{subject to} & U(s) \geq \max_a \left( R(s, a) + \gamma \sum_{s'} T(s' \mid s, a) U(s') \right) \quad \text{for all } s
\end{align*} \quad (7.19)$$

The variables in the optimization are the utilities at each state. Once we know those utilities, we can extract an optimal policy using equation (7.11).

The maximization in the inequality constraints can be replaced by a set of linear constraints, making it a linear program:

$$\begin{align*}
\text{minimize} & \sum_s U(s) \\
\text{subject to} & U(s) \geq R(s, a) + \gamma \sum_{s'} T(s' \mid s, a) U(s') \quad \text{for all } s \text{ and } a
\end{align*} \quad (7.20)$$

In the linear program above, the number of variables is equal to the number of states and the number of constraints is equal to the number of states times the number of actions. Because linear programs can be solved in polynomial time,\footnote{This was proven by L.G. Khachiyan, “Polynomial Algorithms in Linear Programming,” *USSR Computational Mathematics and Mathematical Physics*, vol. 20, no. 1, pp. 53–72, 1980. Modern algorithms tend to be more efficient in practice.} MDPs can be solved in polynomial time. Although a linear programming approach provides this asymptotic complexity guarantee, it is often more efficient in practice to simply use value iteration. Algorithm 7.10 provides an implementation.

7.8 Linear Systems with Quadratic Reward

So far, we have assumed discrete state and action spaces. This section relaxes this assumption, allowing for continuous, vector-valued states and actions. The Bellman equation for discrete problems can be modified as follows:\footnote{This section assumes the problem is undiscounted and finite horizon, but these equations can be easily generalized.}

$$\begin{align*}
\text{minimize} & \sum_s U(s) \\
\text{subject to} & U(s) \geq R(s, a) + \gamma \sum_{s'} T(s' \mid s, a) U(s') \quad \text{for all } s \text{ and } a
\end{align*} \quad (7.20)$$
Algorithm 7.10. Linear program formulation. For convenience in specifying the linear program, we define a function for converting an MDP into its tensor form, where the states and actions consist of integer indices, the reward function is a matrix, and the transition function is a three-dimensional tensor. It uses the JuMP.jl package for mathematical programming. The optimizer is set to use GLPK.jl, but others can be used instead. We also define the default solve behavior for MDPs to use this formulation.

\[ U_{h+1}(s) = \max_a \left( R(s, a) + \int T(s' \mid s, a) U_h(s') \, ds' \right) \]  

(7.21)

where \( s \) and \( a \) in equation (7.16) are replaced with their vector equivalents, the summation is replaced with an integral, and \( T \) provides a probability density rather than a probability mass. Computing equation (7.21) is not straightforward for an arbitrary continuous transition distribution and reward function.

In some cases, exact solution methods do exist for MDPs with continuous state and action spaces.\(^\text{18}\) In particular, if a problem has linear dynamics and has quadratic reward, then the optimal policy can be efficiently found in closed form. Such a system is known in control theory as a linear quadratic regulator (LQR) and has been well studied.\(^\text{19}\)

A problem has linear dynamics if the transition function has the form:

\[ T(s' \mid s, a) = T_s s + T_a a + w \]  

(7.22)

where \( T_s \) and \( T_a \) are matrices that determine the mean of the next state \( s' \) given \( s \) and \( a \), and \( w \) is a random disturbance drawn from a zero mean, finite variance distribution that does not depend on \( s \) and \( a \). One common choice is the multivariate Gaussian.

\(^\text{18}\) For a detailed overview, see Chapter 4 of Volume I of D. Bertsekas, Dynamic Programming and Optimal Control. Athena Scientific, 2007.

A reward function is quadratic if it can be written in the form:

\[ R(s, a) = s^\top R_s s + a^\top R_a a \]  

(7.23)

where \( R_s \) and \( R_a \) are matrices that determine how state and action component combinations contribute reward. We additionally require that \( R_s \) be negative semidefinite and \( R_a \) be negative definite. Such a reward function penalizes states and actions that deviate from 0.

Problems with linear dynamics and quadratic reward are common in control theory where one often seeks to regulate a process such that it does not deviate far from a desired value. The quadratic cost assigns much higher cost to states far from the origin than those near it. The optimal policy for problem with linear dynamics and quadratic reward has an analytic, closed-form solution. Many MDPs can be approximated with linear quadratic MDPs and solved, often yielding reasonable policies for the original problem.

Substituting the transition and reward functions into equation (7.21) produces:

\[ U_{h+1}(s) = \max_a \left( s^\top R_s s + a^\top R_a a + \int p(w) U_h(T_s s + T_a a + w) \, dw \right) \]  

(7.24)

where \( p(w) \) is the probability density of the random, zero-mean disturbance \( w \).

The optimal one-step value function is:

\[ U_1(s) = \max_a \left( s^\top R_s s + a^\top R_a a \right) = s^\top R_s s \]  

(7.25)

for which the optimal action is \( a = 0 \).

We will show through induction that \( U_h(s) \) has a quadratic form \( s^\top V_h s + q_h \) with symmetric matrices \( V_h \). For the one-step value function, \( V_1 = R_s \) and \( q_1 = 0 \). Substituting this quadratic form into equation (7.24) yields:

\[ U_{h+1}(s) = s^\top R_s s + \max_a \left( a^\top R_a a + \int p(w) \left( (T_s s + T_a a + w)^\top V_h (T_s s + T_a a + w) + q_h \right) \, dw \right) \]  

(7.26)

This can be simplified by expanding and using the fact that \( \int p(w) \, dw = 1 \) and \( \int w p(w) \, dw = 0 \):

\[ U_{h+1}(s) = s^\top R_s s + s^\top T_s^\top V_h T_s s \]

\[ + \max_a \left( a^\top R_a a + 2s^\top T_s^\top V_h T_a a + a^\top T_a^\top V_h T_a a \right) \]

(7.27)

\[ + \int p(w) \left( w^\top V_h w \right) \, dw + q_h \]
We can obtain the optimal action by differentiating with respect to \( a \) and setting it to 0:\(^{21}\)

\[
0 = \left( R_a + R_a^\top \right) a + 2T_a^\top V_hT_s s + \left( T_a^\top V_h T_a + \left( T_a^\top V_h T_a \right)^\top \right) a
\]

\[
= 2R_a a + 2T_a^\top V_h T_s s + 2T_a^\top V_h T_a a
\]

Solving for the optimal action yields:\(^{22}\)

\[
a = - \left( R_a + T_a^\top V_h T_a \right)^{-1} T_a^\top V_h T_s s
\]

Substituting the optimal action into \( U_{h+1}(s) \) yields the quadratic form that we were seeking \( U_{h+1}(s) = s^\top V_{h+1}s + q_{h+1} \), with\(^{23}\)

\[
V_{h+1} = R_s + T_s^\top V_h T_s - \left( T_a^\top V_h T_s \right)^\top \left( R_a + T_a^\top V_h T_a \right)^{-1} \left( T_a^\top V_h T_s \right)
\]

and

\[
q_{h+1} = \sum_{i=1}^{h} E_w [w^\top V_i w]
\]

If \( w \sim \mathcal{N}(0, \Sigma) \), then

\[
q_{h+1} = \sum_{i=1}^{h} \text{Tr}(\Sigma V_i)
\]

We can compute \( V_h \) and \( q_h \) up to any horizon \( h \) starting from \( V_1 = R_s \) and \( q_1 = 0 \) and iterating using the equations above. The optimal action for an \( h \)-step policy comes directly from equation (7.29):

\[
\pi_h(s) = - \left( T_a^\top V_{h-1} T_a + R_a \right)^{-1} T_a^\top V_{h-1} T_s s
\]

Note that the optimal action is independent of the zero-mean disturbance distribution. The variance of the disturbance, however, does affect the expected utility. Algorithm 7.11 provides an implementation. Example 7.4 demonstrates this process on a simple problem with linear Gaussian dynamics.

### 7.9 Summary

- Discrete Markov decision processes with bounded rewards can be solved exactly through dynamic programming.
Algorithm 7.11. A method that computes an optimal policy for an \( h_{\text{max}} \)-step horizon MDP with stochastic linear dynamics parameterized by matrices \( T_s \) and \( T_a \) and quadratic reward parameterized by matrices \( R_s \) and \( R_a \). The method returns a vector of policies where entry \( h \) produces the optimal first action in an \( h \)-step policy.

7.10 Exercises

Exercise 7.1. Show that for an infinite sequence of constant rewards \( (r_t = r \text{ for all } t) \) the infinite horizon discounted return converges to \( r/(1 - \gamma) \).
Consider a continuous MDP where the state is composed of a scalar position and velocity $s = [x, v]$. Actions are scalar accelerations $a$ that are each executed over a time step $\Delta t = 1$. Find an optimal 5-step policy from $s_0 = [-10, 0]$ given a quadratic reward:

$$R(s, a) = -x^2 - v^2 - 0.5a^2$$

such that the system tends towards rest at $s = 0$.

The transition dynamics are:

$$\begin{bmatrix} x' \\ v' \end{bmatrix} = \begin{bmatrix} x + v\Delta t + \frac{1}{2}a\Delta t^2 + w_1 \\ v + a\Delta t + w_2 \end{bmatrix} = \begin{bmatrix} 1 & \Delta t \\ 0 & 1 \end{bmatrix} \begin{bmatrix} x \\ v \end{bmatrix} + \begin{bmatrix} 0.5\Delta t^2 \\ \Delta t \end{bmatrix} [a] + w$$

where $w$ is drawn from a zero-mean multivariate Gaussian distribution with covariance $0.1I$.

The reward matrices are $R_s = -I$ and $R_a = -[0.5]$.

The resulting optimal policies are:

$$\pi_1(s) = \begin{bmatrix} 0 \\ 0 \end{bmatrix} s$$
$$\pi_2(s) = \begin{bmatrix} -0.286 \\ -0.857 \end{bmatrix} s$$
$$\pi_3(s) = \begin{bmatrix} -0.462 \\ -1.077 \end{bmatrix} s$$
$$\pi_4(s) = \begin{bmatrix} -0.499 \\ -1.118 \end{bmatrix} s$$
$$\pi_5(s) = \begin{bmatrix} -0.504 \\ -1.124 \end{bmatrix} s$$
Solution: We can prove that the infinite sequence of discounted constant rewards converges to \( r/(1 - \gamma) \) in the following steps

\[
\sum_{t=1}^{\infty} \gamma^{t-1}r_t = r + \gamma r + \gamma^2 r + \cdots
\]

\[
= r + \gamma \sum_{t=1}^{\infty} \gamma^{t-1}r_t
\]

We can move the summation to the left side and factor out \((1 - \gamma)\):

\[
(1 - \gamma) \sum_{t=1}^{\infty} \gamma^{t-1}r_t = r
\]

\[
\sum_{t=1}^{\infty} \gamma^{t-1}r_t = \frac{r}{1 - \gamma}
\]

Exercise 7.2. Suppose we have a Markov decision process consisting of five states \( s_1 \) to \( s_5 \) and two actions, to stay \((a_S)\) and continue \((a_C)\). We have the following

\[
T(s_i \mid s_i, a_S) = 1 \text{ for } i \in \{1, 2, 3, 4\}
\]

\[
T(s_{i+1} \mid s_i, a_C) = 1 \text{ for } i \in \{1, 2, 3, 4\}
\]

\[
T(s_5 \mid s_5, a) = 1 \text{ for all actions } a
\]

\[
R(s_i, a) = 0 \text{ for } i \in \{1, 2, 3, 5\} \text{ and for all actions } a
\]

\[
R(s_4, a_S) = 0
\]

\[
R(s_4, a_C) = 10
\]

What is the discount factor \( \gamma \) if the optimal value \( U^*(s_1) = 1 \)?

Solution: The optimal value of \( U^*(s_1) \) is associated with following the optimal policy \( \pi^* \) starting from \( s_1 \). Given the transition model, the optimal policy from \( s_1 \) is to continue until reaching \( s_5 \), which is a terminal state where we can no longer transition to another state or accumulate additional reward. Thus, the optimal value of \( s_1 \) can be computed as

\[
U^*(s_1) = \sum_{t=1}^{\infty} \gamma^{t-1}r_t
\]

\[
U^*(s_1) = R(s_1, a_C) + \gamma R(s_2, a_C) + \gamma^2 R(s_3, a_C) + \gamma^3 R(s_4, a_C) + \gamma^4 R(s_5, a_C) + \cdots
\]

\[
U^*(s_1) = 0 + \gamma^1 \times 0 + \gamma^2 \times 0 + \gamma^3 \times 10 + \gamma^4 \times 0 + 0
\]

\[
1 = 10\gamma^3
\]

Thus, the discount factor is \( \gamma = 0.1^{1/3} \approx 0.464 \)

Exercise 7.3. What is the time complexity of performing \( k \) steps of iterative policy evaluation?
Solution: Iterative policy evaluation requires computing the lookahead equation:

\[ U_{k+1}^\pi(s) = R(s, \pi(s)) + \gamma \sum_{s'} T(s' \mid s, \pi(s)) U_k^\pi(s') \]

Updating the value at a single state requires summing over all \(|S|\) states. For a single iteration over all states, we must do this operation \(|S|\) times. Thus, the time complexity of \(k\) steps of iterative policy evaluation is \(O(k|S|^2)\).

Exercise 7.4. Suppose we have an MDP with six states \(s_{1:6}\) and four actions \(a_{1:4}\). Using the following tabular form of the action value function \(Q(s, a)\), compute \(U(s), \pi(s),\) and \(A(s, a)\).

<table>
<thead>
<tr>
<th>(Q(s, a))</th>
<th>(a_1)</th>
<th>(a_2)</th>
<th>(a_3)</th>
<th>(a_4)</th>
</tr>
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<tr>
<td>(s_1)</td>
<td>0.41</td>
<td>0.46</td>
<td>0.37</td>
<td>0.37</td>
</tr>
<tr>
<td>(s_2)</td>
<td>0.50</td>
<td>0.55</td>
<td>0.46</td>
<td>0.37</td>
</tr>
<tr>
<td>(s_3)</td>
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<td>0.50</td>
<td>0.38</td>
<td>0.44</td>
</tr>
<tr>
<td>(s_4)</td>
<td>0.41</td>
<td>0.50</td>
<td>0.33</td>
<td>0.41</td>
</tr>
<tr>
<td>(s_5)</td>
<td>0.50</td>
<td>0.60</td>
<td>0.41</td>
<td>0.39</td>
</tr>
<tr>
<td>(s_6)</td>
<td>0.71</td>
<td>0.70</td>
<td>0.61</td>
<td>0.59</td>
</tr>
</tbody>
</table>

Solution: We can compute \(U(s), \pi(s),\) and \(A(s, a)\) using the following equations

\[ U(s) = \max_a Q(s, a) \quad \pi(s) = \arg \max_a Q(s, a) \quad A(s, a) = Q(s, a) - U(s) \]

<table>
<thead>
<tr>
<th>(s)</th>
<th>(U(s))</th>
<th>(\pi(s))</th>
<th>(A(s, a_1))</th>
<th>(A(s, a_2))</th>
<th>(A(s, a_3))</th>
<th>(A(s, a_4))</th>
</tr>
</thead>
<tbody>
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<td>0.46</td>
<td>(a_2)</td>
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<td>0.00</td>
<td>-0.09</td>
<td>-0.09</td>
</tr>
<tr>
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<td>(a_2)</td>
<td>-0.05</td>
<td>0.00</td>
<td>-0.09</td>
<td>-0.18</td>
</tr>
<tr>
<td>(s_3)</td>
<td>0.60</td>
<td>(a_1)</td>
<td>0.00</td>
<td>-0.10</td>
<td>-0.22</td>
<td>-0.16</td>
</tr>
<tr>
<td>(s_4)</td>
<td>0.50</td>
<td>(a_2)</td>
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<td>0.00</td>
<td>-0.17</td>
<td>-0.09</td>
</tr>
<tr>
<td>(s_5)</td>
<td>0.60</td>
<td>(a_2)</td>
<td>-0.10</td>
<td>0.00</td>
<td>-0.19</td>
<td>-0.21</td>
</tr>
<tr>
<td>(s_6)</td>
<td>0.71</td>
<td>(a_1)</td>
<td>0.00</td>
<td>-0.01</td>
<td>-0.10</td>
<td>-0.12</td>
</tr>
</tbody>
</table>

Exercise 7.5. Suppose we have a three-tile, straight-line hex world (appendix F.1) where the rightmost tile is an absorbing state. When we take any action in the rightmost state, we get a reward of 10 and we are transported to a fourth terminal state where we no longer receive any reward. Use a discount factor of \(\gamma = 0.9\). Perform a single step of policy iteration where the initial policy \(\pi\) has us move east in the first tile, northeast in the second tile, and southwest in the third tile. For the policy evaluation step, write out the transition matrix \(T^\pi\) and the reward vector \(R^\pi\) and then solve the infinite horizon value function \(U^\pi\) directly using matrix inversion. For the policy improvement step, compute the updated policy \(\pi'\) by maximizing the lookahead equation.

Solution: For the policy evaluation step, we use equation (7.10), repeated below:

\[ U^\pi = (I - \gamma T^\pi)^{-1} R^\pi \]
Forming the transition matrix $T^\pi$ and reward vector $R^\pi$ with an additional state for the terminal state, we can solve for the infinite horizon value function $U^\pi$

$$U^\pi = \begin{pmatrix} 1 & 0 & 0 & 0 \\ 0 & 1 & 0 & 0 \\ 0 & 0 & 1 & 0 \\ 0 & 0 & 0 & 1 \end{pmatrix} - (0.9) \begin{pmatrix} 0.3 & 0.7 & 0 & 0 \\ 0 & 0.85 & 0.15 & 0 \\ 0 & 0 & 1 & 0 \\ 0 & 0 & 0 & 1 \end{pmatrix}^{-1} \begin{pmatrix} -0.3 \\ -0.85 \\ 10 \\ 0 \end{pmatrix} \approx \begin{pmatrix} 1.425 \\ 2.128 \end{pmatrix}$$

For the policy improvement step, we apply equation (7.11) using the updated value function. The actions in the $\arg\max$ term correspond to $a_E, a_{NE}, a_{NW}, a_W, a_{SW},$ and $a_{SE}$:

$$\pi(s_1) = \arg \max \{1.425, 0.527, 0.283, 0.283, 0.283, 0.527\} = a_E$$
$$\pi(s_2) = \arg \max \{6.575, 2.128, 0.970, 1.172, 0.970, 2.128\} = a_E$$
$$\pi(s_3) = \arg \max \{10, 10, 10, 10, 10, 10\} = a_*, \text{ all actions are equally desirable}$$

**Exercise 7.6.** Perform two steps of value iteration to the problem in exercise 7.5, starting with an initial value function $U_0(s) = 0$ for all $s$.

**Solution:** We need to use the Bellman equation (equation (7.16)) to perform iterative backups of the value function. The actions in the $\max$ term correspond to $a_E, a_{NE}, a_{NW}, a_W, a_{SW},$ and $a_{SE}$. For our first iteration, the value function is zero for all states, so we only need to consider the reward component:

$$U_1(s_1) = \max(-0.3, -0.85, -1, -1, -1, -0.85) = -0.3$$
$$U_1(s_2) = \max(-0.3, -0.85, -0.85, -0.3, -0.85, -0.85) = -0.3$$
$$U_1(s_3) = \max(10, 10, 10, 10, 10, 10) = 10$$

For the second iteration

$$U_2(s_1) = \max(-0.57, -1.12, -1.27, -1.27, -1.27, -1.12) = -0.57$$
$$U_2(s_2) = \max(5.919, 0.271, -1.12, -0.57, -1.12, 0.271) = 5.919$$
$$U_2(s_3) = \max(10, 10, 10, 10, 10, 10) = 10$$

**Exercise 7.7.** Apply one sweep of asynchronous value iteration to the problem in exercise 7.5, starting with an initial value function $U_0(s) = 0$ for all $s$. Update the states from right to left.
Solution: We use the Bellman equation (equation (7.16)) to iteratively update the value function over each state following our ordering. The actions in the max term correspond to $a_E, a_{NE}, a_{NW}, a_W, a_{SW},$ and $a_{SE}$:

\[
U(s_3) = \max(10, 10, 10, 10, 10) = 10 \\
U(s_2) = \max(6, 0.5, 0.85, -0.3, -0.85, 0.5) = 6 \\
U(s_1) = \max(3.48, -0.04, -1, -1, -1, -0.04) = 3.48
\]

Exercise 7.8. Prove that a Bellman residual of $\delta$ guarantees that the value function obtained by value iteration is within $\delta \gamma / (1 - \gamma)$ of $U^*(s)$ at every state $s$.

Solution: For a given $U_k$, suppose we know that $\|U_k - U_{k-1}\|_\infty < \delta$. We can bound the improvement in the next iteration:

\[
U_{k+1}(s) - U_k(s) = \max_a \left( R(s,a) + \gamma \sum_{s'} T(s' | s,a) U_k(s') \right) \\
- \max_a \left( R(s,a) + \gamma \sum_{s'} T(s' | s,a) U_{k-1}(s') \right) \\
< \max_a \left( R(s,a) + \gamma \sum_{s'} T(s' | s,a) U_k(s') \right) \\
- \max_a \left( R(s,a) + \gamma \sum_{s'} T(s' | s,a) (U_k(s') - \delta) \right) \\
= \delta \gamma
\]

Similarly,

\[
U_{k+1}(s) - U_k(s) > \max_a \left( R(s,a) + \gamma \sum_{s'} T(s' | s,a) U_k(s') \right) \\
- \max_a \left( R(s,a) + \gamma \sum_{s'} T(s' | s,a) (U_k(s') + \delta) \right) \\
= -\delta \gamma
\]

The accumulated improvement after infinite iterations is thus bounded by:

\[
\|U^*(s) - U_k(s)\|_\infty < \sum_{i=1}^{\infty} \delta \gamma^i = \frac{\delta \gamma}{1 - \gamma}
\]

A Bellman residual of $\delta$ thus guarantees that the optimal value function obtained by value iteration is within $\delta \gamma / (1 - \gamma)$ of $U^*$.
Exercise 7.9. Prove that if \( \| U - U^* \|_\infty < \epsilon \) for an estimate of the value function \( U \), then the policy \( \pi \) extracted from \( U \) using equation (7.11) has a policy loss bound of \( \| U^\pi - U^* \|_\infty < 2\epsilon \).

**Solution:** We are given \( \| U - U^* \|_\infty < \epsilon \). The worst-case deviation between the policy extracted from \( U \) and an optimal policy occurs when the policy from \( U \) has value just under \( U + \epsilon \) and the true optimal policy has value just above \( U - \epsilon \). In this case, \( U^\pi - U^* < U + \epsilon - (U - \epsilon) = 2\epsilon \).

Exercise 7.10. Show how a LQR problem with a quadratic reward function \( R(s, a) = s^T R_s s + a^T R_a a \) can be reformulated so that the reward function includes linear terms in \( s \) and \( a \).

**Solution:** We can introduce an additional state dimension that is always equal to 1, yielding a new system with linear dynamics:

\[
\begin{bmatrix}
s' \\
1
\end{bmatrix} = \begin{bmatrix}
T_s & 0 \\
0 & 1
\end{bmatrix} \begin{bmatrix}
s \\
1
\end{bmatrix} + T_a a
\]

The reward function of the augmented system can now have linear state reward terms:

\[
\begin{bmatrix}
s \\
1
\end{bmatrix}^T R_{\text{augmented}} \begin{bmatrix}
s \\
1
\end{bmatrix} = s^T R_s s + 2r_{s,\text{linear}} s + r_{s,\text{scalar}}
\]

Similarly, we can include an additional action dimension that is always 1 to obtain linear action reward terms.

Exercise 7.11. Why does the optimal policy obtained in example 7.4 produce actions with greater magnitude when the horizon is greater?

**Solution:** The problem in example 7.4 has quadratic reward that penalizes deviations from the origin. The longer the horizon, the greater the negative reward is that can be accumulated, making it more worthwhile to reach the origin sooner.

Exercise 7.12. Prove that iterative policy evaluation converges to the solution of equation (7.6).

**Solution:** Consider iterative policy evaluation applied to a policy \( \pi \) as given in equation (7.5):

\[
U^\pi_{k+1}(s) = R(s, \pi(s)) + \gamma \sum_{s'} T(s' \mid s, a) U^\pi_k(s')
\]
Let us define an operator \( B_\pi \) and rewrite the above as \( U_{k+1}^\pi = B_\pi U_k^\pi \). We can show that \( B_\pi \) is a contraction mapping:

\[
B_\pi U^\pi(s) = R(s, \pi(s)) + \gamma \sum_{s'} T(s' \mid s, a) U^\pi(s')
\]

\[
= R(s, \pi(s)) + \gamma \sum_{s'} T(s' \mid s, a) (U^\pi(s') - \hat{U}^\pi(s') + \hat{U}^\pi(s'))
\]

\[
= B_\pi \hat{U}^\pi(s) + \gamma \sum_{s'} T(s' \mid s, a) (U^\pi(s') - \hat{U}^\pi(s'))
\]

\[
\leq B_\pi \hat{U}^\pi(s) + \gamma \|U^\pi - \hat{U}\|_\infty
\]

Hence, \( \|B_\pi U^\pi - B_\pi \hat{U}^\pi\|_\infty \leq \alpha \|U^\pi - \hat{U}\|_\infty \) for \( \alpha = \gamma \), implying that \( B_\pi \) is a contraction mapping. As discussed in appendix A.15, \( \lim_{t \to \infty} B_\pi^t U^\pi \) converges to a unique fixed point \( U^\pi \) for which \( U^\pi = B_\pi U^\pi \).

**Exercise 7.13.** Prove that value iteration converges to a unique solution.

**Solution:** The value iteration update (equation (7.16)) is:

\[
U^{k+1}(s) = \max_a \left( R(s, a) + \gamma \sum_{s'} T(s' \mid s, a) U_k(s') \right)
\]

We will denote the Bellman operator as \( B \) and rewrite an application of Bellman’s equation as \( U_{k+1} = BU_k \). As with the previous problem, if \( B \) is a contraction mapping, then repeated application of \( B \) to \( U \) will converge to a unique fixed point.

We can show that \( B \) is a contraction mapping:

\[
BU(s) = \max_a \left( R(s, a) + \gamma \sum_{s'} T(s' \mid s, a) U(s') \right)
\]

\[
= \max_a \left( R(s, a) + \gamma \sum_{s'} T(s' \mid s, a) (U(s') - \hat{U}(s') + \hat{U}(s')) \right)
\]

\[
\leq B\hat{U}(s) + \gamma \max_a \sum_{s'} T(s' \mid s, a) (U(s') - \hat{U}(s'))
\]

\[
\leq B\hat{U}(s) + \alpha \|U - \hat{U}\|_\infty
\]

for \( \alpha = \gamma \max_s \max_a \sum_{s'} T(s' \mid s, a) \), with \( 0 \leq \alpha < 1 \). Hence, \( \|BU - B\hat{U}\|_\infty \leq \alpha \|U - \hat{U}\|_\infty \), which implies \( B \) is a contraction mapping.

**Exercise 7.14.** Show that the point to which value iteration converges corresponds to the optimal value function.

**Solution:** Let \( U \) be the value function produced by value iteration. We want to show that \( U = U^* \). At convergence, we have \( BU = U \). Let \( U_0 \) be a value function that maps all states to 0. For any policy \( \pi \), it follows from the definition of \( B_\pi \) that \( B_\pi U_0 \leq BU_0 \). Similarly, \( B_\pi^t U_0 \leq B^t U_0 \). Because \( B_\pi^t U_0 \to U^* \) and \( B^t U_0 \to U \) as \( t \to \infty \), it follows that \( U^* \leq U \), which can only be the case if \( U = U^* \).
Exercise 7.15. Suppose we have a linear Gaussian problem with quadratic reward with disturbance $\mathbf{w} \sim \mathcal{N}(0, \Sigma)$. Show that the scalar term in the utility function has the form:

$$q_{h+1} = \sum_{i=1}^{h} \mathbb{E}_{\mathbf{w}} \left[ \mathbf{w}^\top \mathbf{V}_i \mathbf{w} \right] = \sum_{i=1}^{h} \text{Tr}(\Sigma \mathbf{V}_i)$$

You may want to use the trace trick:

$$\mathbf{x}^\top \mathbf{A} \mathbf{x} = \text{Tr} \left( \mathbf{x}^\top \mathbf{A} \mathbf{x} \right) = \text{Tr} \left( \mathbf{A} \mathbf{x} (\mathbf{x}^\top) \right)$$

Solution: The above equation is true if $\mathbb{E}_{\mathbf{w}} \left[ \mathbf{w}^\top \mathbf{V}_i \mathbf{w} \right] = \text{Tr}(\Sigma \mathbf{V}_i)$. Our derivation is:

$$\mathbb{E}_{\mathbf{w} \sim \mathcal{N}(0, \Sigma)} \left[ \mathbf{w}^\top \mathbf{V}_i \mathbf{w} \right] = \mathbb{E}_{\mathbf{w} \sim \mathcal{N}(0, \Sigma)} \left[ \text{Tr} \left( \mathbf{w}^\top \mathbf{V}_i \mathbf{w} \right) \right]$$

$$= \mathbb{E}_{\mathbf{w} \sim \mathcal{N}(0, \Sigma)} \left[ \text{Tr} \left( \mathbf{V}_i \mathbf{w} \mathbf{w}^\top \right) \right]$$

$$= \text{Tr} \left( \mathbb{E}_{\mathbf{w} \sim \mathcal{N}(0, \Sigma)} \left[ \mathbf{V}_i \mathbf{w} \mathbf{w}^\top \right] \right)$$

$$= \text{Tr} \mathbf{V}_i \mathbb{E}_{\mathbf{w} \sim \mathcal{N}(0, \Sigma)} \left[ \mathbf{w} \mathbf{w}^\top \right]$$

$$= \text{Tr} \mathbf{V}_i \Sigma$$

$$= \text{Tr}(\Sigma \mathbf{V}_i)$$

Exercise 7.16. What is the role of the scalar term $q$ in the LQR optimal value function, as given in equation (7.31):

$$q_{h+1} = \sum_{i=1}^{h} \mathbb{E}_{\mathbf{w}} \left[ \mathbf{w}^\top \mathbf{V}_i \mathbf{w} \right]$$

Solution: A matrix $\mathbf{M}$ is positive definite if for all non-zero $\mathbf{x}$, that $\mathbf{x}^\top \mathbf{M} \mathbf{x} > 0$. In equation (7.31), every $\mathbf{V}_i$ is negative semi-definite, so $\mathbf{w}^\top \mathbf{V}_i \mathbf{w} \leq 0$ for all $\mathbf{w}$. Thus, these $q$ terms are guaranteed to be non-positive. This should be expected, as in LQR problems it is impossible to obtain positive reward, and we seek instead to minimize cost.

The $q$ scalars are offsets in the quadratic optimal value function:

$$U(\mathbf{s}) = \mathbf{s}^\top \mathbf{V} \mathbf{s} + q$$

Each $q$ represents the baseline reward around which the $\mathbf{s}^\top \mathbf{V} \mathbf{s}$ term fluctuates. We know that $\mathbf{V}$ is negative definite, so $\mathbf{s}^\top \mathbf{V} \mathbf{s} \leq 0$, and $q$ thus represents the expected reward that one could obtain if one were at the origin, $\mathbf{s} = 0$. 

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8 Approximate Value Functions

Up to this point, we have assumed that the value function can be represented as a table. Tables are only useful representations for small discrete problems. Problems with larger state spaces may require an infeasible amount of memory, and the exact methods discussed in the previous chapter may require an infeasible amount of computation. For such problems, we often have to resort to approximate dynamic programming, where the solution may not be exact.¹ One way to approximate solutions is to use value function approximation, which is the subject of this chapter. We will discuss different approaches to approximating the value function and how to incorporate dynamic programming to derive approximately optimal policies.

8.1 Parametric Representations

We will use \( U_\theta(s) \) to denote our parametric representation of the value function, where \( \theta \) is the vector of parameters. There are many different ways to represent \( U_\theta(s) \), several of which will be mentioned later in this chapter. Assuming we have such an approximation, we can extract an action according to:

\[
\pi(s) = \arg \max_a \left( R(s, a) + \gamma \sum_{s'} T(s' \mid s, a) U_\theta(s') \right)
\] (8.1)

Value function approximations are often used in problems with continuous state spaces, in which case the summation above may be replaced with an integral. The integral can be approximated using transition model samples.

An alternative to the computation in equation (8.1) is to approximate the action value function \( Q(s, a) \). If we use \( Q_\theta(s, a) \) to represent our parametric

¹ A deeper treatment of this field is provided by W. B. Powell, *Approximate Dynamic Programming: Solving the Curses of Dimensionality*, 2nd ed. Wiley, 2011.
approximation, we can obtain an action according to:

$$\pi(s) = \arg \max_a Q_{\theta}(s, a)$$  \hfill (8.2)

This chapter discusses how we can apply dynamic programming at a finite set of states $S = s_1:n$ to arrive at a parametric approximation of the value function over the full state space. Different schemes can be used to generate this set. If the state space is relatively low-dimensional, we can define a grid. Another approach is to use random sampling from the state space. However, some states are more likely to be encountered than others and are therefore more important in constructing the value function. We can bias the sampling towards more important states by running simulations with some policy, perhaps initially random, from a plausible set of initial states.

An iterative approach can be used to enhance our approximation of the value function at the states in $S$. We alternate between improving our value estimates at $S$ through dynamic programming and refitting our approximation at those states. Algorithm 8.1 provides an implementation where the dynamic programming step consists of Bellman backups as done in value iteration (section 7.5). A similar algorithm can be created for action value approximations $Q_{\theta}$.

Algorithm 8.1. Approximate value iteration for an MDP with parameterized value function approximation $U_{\theta}$. We perform backups (defined in algorithm 7.7) at the states in $S$ to obtain a vector of utilities $U$. We then call $\text{fit!}(U_{\theta}, S, U)$, which modifies the parametric representation $U_{\theta}$ to better match the value of the states in $S$ to the utilities in $U$. Different parametric approximations have different implementations for $\text{fit!}$.

```plaintext
struct ApproximateValueIteration
    U_{\theta} # initial parameterized value function that supports fit!
    S # set of discrete states for performing backups
    k_max # maximum number of iterations
end

function solve(M::ApproximateValueIteration, P::MDP)
    U_{\theta}, S, k_max = M.U_{\theta}, M.S, M.k_max
    for k in 1:k_max
        U = [backup(P, U_{\theta}, s) for s in S]
        fit!(U_{\theta}, S, U)
    end
    return ValueFunctionPolicy(P, U_{\theta})
end
```

All of the parametric representations discussed in this chapter can be used with algorithm 8.1. To be used with that algorithm, a representation needs to support the evaluation of $U_{\theta}$ and the fitting of $U_{\theta}$ to estimates of the utilities at the points in $S$.  

---

We can group the parametric representations into two categories. The first category includes local approximation methods where $\theta$ corresponds to the values at the states in $S$. To evaluate $U_\theta(s)$ at an arbitrary state $s$, we take a weighted sum of the values stored in $S$. The second category includes global approximation methods where $\theta$ is not directly related to the values at the states in $S$. In fact, $\theta$ may have far fewer or even far more components than there are states in $S$.

Both local approximation as well as many global approximations can be viewed as a linear function approximation $U_\theta(s) = \theta^\top \beta(s)$, where methods differ in how they define the vector function $\beta$. In local approximation methods, $\beta(s)$ determines how to weight the utilities of the states in $S$ to approximate the utility at state $s$. In many global approximation methods, $\beta(s)$ is viewed as a set of basis functions that are combined together in a linear fashion to obtain an approximation for an arbitrary $s$.

We can also approximate the action value function using a linear function, $Q_\theta(s,a) = \theta^\top \beta(s,a)$. In the context of local approximations, we can provide approximations over continuous action spaces by choosing a finite set of actions $A \subset A$. Our parameter vector $\theta$ would then consist of $|S| \times |A|$ components, each corresponding to a state-action value. Our function $\beta(s,a)$ would return a vector with the same number of components that specifies how to weight together our finite set of state-action values to obtain an estimate of the utility associated with state $s$ and action $a$.

### 8.2 Nearest Neighbor

A simple approach to local approximation is to use the value of the state in $S$ that is the nearest neighbor of $s$. In order to use this approach, we need a distance metric (see appendix A.3). We use $d(s,s')$ to denote the distance between two states $s$ and $s'$. The approximate value function is then $U_\theta(s) = \theta_i$, where $i = \arg\min_{j \in 1:m} d(s_j,s)$. Figure 8.1 shows an example of a value function represented using the nearest neighbor scheme.

We can generalize this approach to average together the values of the $k$-nearest neighbors. This approach still results in piecewise constant value functions, but different values for $k$ can result in better approximations. Figure 8.1 shows examples of value functions approximated with different values for $k$. Algorithm 8.2 provides an implementation.

![Nearest Neighbor](image-url)
mutable struct NearestNeighborValueFunction
    k  # number of neighbors
    d  # distance function d(s, s')
    S  # set of discrete states
    θ  # vector of values at states in S
end

function (Uθ::NearestNeighborValueFunction)(s)
    dists = [Uθ.d(s, s′) for s′ in Uθ.S]
    ind = sortperm(dists)[1:Uθ.k]
    return mean(Uθ.θ[i] for i in ind)
end

function fit!(Uθ::NearestNeighborValueFunction, S, U)
    Uθ.θ = U
    return Uθ
end

Algorithm 8.2. The \( k \)-nearest neighbors method, which approximates the value of a state \( s \) based on the \( k \) closest states in \( S \) as determined by a distance function \( d \). The vector \( θ \) contains the values of the states in \( S \). Greater efficiency can be achieved by using specialized data structures, such as \( kd \)-trees, implemented in NearestNeighbors.jl.

8.3 Kernel Smoothing

Another local approximation method is kernel smoothing, where the utilities of the states in \( S \) are smoothed over the entire state space. This method requires defining a kernel function \( k(s, s') \) that relates pairs of states \( s \) and \( s' \). We generally want \( k(s, s') \) to be higher for states that are closer together because those values tell us how to weight together the utilities associated with the states in \( S \). This method results in the linear approximation:

\[
U_θ(s) = \sum_{i=1}^{m} \theta_i \beta_i(s) = \theta^T \beta(s) \quad (8.3)
\]

where

\[
\beta_i(s) = \frac{k(s, s_i)}{\sum_{j=1}^{m} k(s, s_j)} \quad (8.4)
\]

Algorithm 8.3 provides an implementation.

There are many different ways we can define a kernel function. We can define our kernel to simply be the inverse of the distance between states:

\[
k(s, s') = \max(d(s, s'), \epsilon)^{-1} \quad (8.5)
\]
where $\epsilon$ is a small positive constant to avoid dividing by zero when $s = s'$. Figure 8.2 shows value approximations using several different distance functions. As we can see, kernel smoothing can result in smooth value function approximations, in contrast with $k$-nearest neighbors. Figure 8.3 applies this kernel to a discrete hex world problem and shows the outcome of a few iterations of approximate value iteration (algorithm 8.1). Figure 8.4 shows a value function and policy learned for the mountain car problem (appendix F.4) with a continuous state space.

Another common kernel is the Gaussian kernel:

$$k(s, s') = \exp\left(-\frac{d(s, s')^2}{2\sigma^2}\right)$$  \hspace{1cm} (8.6)

where $\sigma$ controls the degree of smoothing.

```plaintext
mutable struct LocallyWeightedValueFunction
    k # kernel function k(s, s')
    S # set of discrete states
    θ # vector of values at states in S
end

function (Uθ::LocallyWeightedValueFunction)(s)
    w = normalize([Uθ.k(s, s') for s' in Uθ.S], 1)
    return Uθ.θ .• w
end

function fit!(Uθ::LocallyWeightedValueFunction, S, U)
    Uθ.θ = U
    return Uθ
end
```

Algorithm 8.3. Locally weighted value function approximation defined by a kernel function $k$ and a vector of utilities $\theta$ at states in $S$. 

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2021-03-07 23:12:00-08:00, revision 80bf4d6, comments to bugs@algorithmsbook.com
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Figure 8.3. Local approximation value iteration used to iteratively improve an approximate value function on the hex world problem. The five outlined states are used to approximate the value function. The value of the remaining states are approximated using the distance function $\|s - s'\|_2^2$. The resulting policy is reasonable but nevertheless sub-optimal.

Figure 8.4. A utility function and policy obtained by learning the action values for a finite set of states (white) in the mountain car problem using the distance function $\|s - s'\|_2 + 0.1$. 
8.4 Linear Interpolation

Linear interpolation is another common approach to local approximation. The one-dimensional case is straightforward, where the approximated value for a state \( s \) between two states \( s_1 \) and \( s_2 \) is:

\[
U_\theta(s) = \alpha \theta_1 + (1 - \alpha) \theta_2
\]

with \( \alpha = (s_2 - s) / (s_2 - s_1) \). This case is shown in both figure 8.5 and figure 8.6.

Linear interpolation can be extended to a multidimensional grid. In the two-dimensional case, called bilinear interpolation, we interpolate between four vertices. Bilinear interpolation is done through single-dimensional linear interpolation, once in each axis, requiring the utility of four states at the grid vertices. This interpolation is shown in figure 8.7.

Given four vertices with coordinates \( s_1 = [x_1, y_1] \), \( s_2 = [x_1, y_2] \), \( s_3 = [x_2, y_1] \), and \( s_4 = [x_2, y_2] \), and a sample state \( s = [x, y] \), the interpolated value is:

\[
U_\theta(s) = \begin{cases}
\theta_{12} = 1D \text{ interpolation between } \theta_1 \text{ and } \theta_2 \text{ along vertical axis.} \\
\theta_{24} = 1D \text{ interpolation between } \theta_2 \text{ and } \theta_4 \text{ along horizontal axis.} \\
\theta_{13} = 1D \text{ interpolation between } \theta_1 \text{ and } \theta_3 \text{ along horizontal axis.} \\
\theta_{34} = 1D \text{ interpolation between } \theta_3 \text{ and } \theta_4 \text{ along vertical axis.}
\end{cases}
\]

\[U_\theta(s) = \begin{cases}
\text{1D interpolation between } \theta_{12} \text{ and } \theta_{34} \text{ along horizontal axis.} \\
\text{1D interpolation between } \theta_{13} \text{ and } \theta_{24} \text{ along vertical axis.}
\end{cases}\]
\[ U_\theta(s) = \alpha \theta_{12} + (1 - \alpha) \theta_{34} \]  
   \[ = \frac{x_2 - x}{x_2 - x_1} \theta_{12} + \frac{x - x_1}{x_2 - x_1} \theta_{34} \]  
   \[ = \frac{x_2 - x}{x_2 - x_1} (\alpha \theta_1 + (1 - \alpha) \theta_2) + \frac{x - x_1}{x_2 - x_1} (\alpha \theta_3 + (1 - \alpha) \theta_4) \]  
   \[ = \frac{x_2 - x}{x_2 - x_1} \left( \frac{y_2 - y}{y_2 - y_1} \theta_1 + \frac{y - y_1}{y_2 - y_1} \theta_2 \right) + \frac{x - x_1}{x_2 - x_1} \left( \frac{y_2 - y}{y_2 - y_1} \theta_3 + \frac{y - y_1}{y_2 - y_1} \theta_4 \right) \]  
   \[ = \frac{(x_2 - x)(y_2 - y)}{(x_2 - x_1)(y_2 - y_1)} \theta_1 + \frac{(x_2 - x)(y - y_1)}{(x_2 - x_1)(y_2 - y_1)} \theta_2 + \frac{(x - x_1)(y_2 - y)}{(x_2 - x_1)(y_2 - y_1)} \theta_3 + \frac{(x - x_1)(y - y_1)}{(x_2 - x_1)(y_2 - y_1)} \theta_4 \] (8.12)

The resulting interpolation weighs each vertex according to area of its opposing quadrant, as shown in figure 8.8.

**Multilinear interpolation** in \( d \) dimensions is similarly achieved by linearly interpolating along each axis, requiring \( 2^d \) vertices. Here too, the utility of each vertex is weighted according to the volume of the opposing hyperrectangle. Multilinear interpolation is implemented in algorithm 8.4. Figure 8.9 demonstrates this approach on a two-dimensional state space.

### 8.5 Simplex Interpolation

Multilinear interpolation can be inefficient in high dimensions. Rather than weighting the contributions of \( 2^d \) points, **simplex interpolation** considers only \( d + 1 \) points in the neighborhood of a given state to produce a continuous surface that matches the known sample points.

We start with a multidimensional grid and divide each cell into \( d! \) **simplexes**, which are multidimensional generalizations of triangles defined by the convex hull of \( d + 1 \) vertices. This process is known as **Coxeter-Freudenthal-Kuhn triangulation**, and it ensures that any two simplexes that share a face will produce equivalent values across the face, thus producing continuity when interpolating, as shown in figure 8.10.

---

Algorithm 8.4. A method for conducting multilinear interpolation to estimate the value of state vector \( s \) for known state values \( \theta \) over a grid defined by a lower-left vertex \( o \) and vector of widths \( \delta \). Vertices of the grid can all be written \( o + \delta \cdot i \) for some non-negative integral vector \( i \). The package Interpolations.jl also provides multilinear and other interpolation methods.

```plaintext
mutable struct MultilinearValueFunction
    o # position of lower-left corner
    δ # vector of widths
    θ # vector of values at states in S
end

function (Uθ :: MultilinearValueFunction)(s)
    o, δ, θ = Uθ.o, Uθ.δ, Uθ.θ
    Δ = (s - o)/δ
    # Multidimensional index of lower-left cell
    i = min.(floor.(Int, Δ) .+ 1, size(θ) .- 1)
    vertex_index = similar(i)
    d = length(s)
    u = 0.0
    for vertex in 0:2^d-1
        weight = 1.0
        for j in 1:d
            # Check whether jth bit is set
            if vertex & (1 << (j-1)) > 0
                vertex_index[j] = i[j] + 1
                weight *= Δ[j] - i[j] + 1
            else
                vertex_index[j] = i[j]
                weight *= i[j] - Δ[j]
            end
        end
        u += θ[vertex_index...] * weight
    end
    return u
end

function fit!(Uθ :: MultilinearValueFunction, S, U)
    Uθ.θ = U
    return Uθ
end
```
To illustrate, suppose we have translated and scaled the cell containing a state such that the lowest vertex is 0 and the diagonally opposite vertex is 1. There is a simplex for each permutation of \(1 : d\). The simplex given by permutation \(p\) is the set of points \(x\) satisfying:

\[
0 \leq x_{p_1} \leq x_{p_2} \leq \cdots \leq x_{p_d} \leq 1
\]

(8.13)

Figure 8.11 shows the simplexes obtained for the unit cube.

Simplex interpolation first translates and scales a state vector \(s\) to the unit hypercube of its corresponding cell to obtain \(s'\). It then sorts the entries in \(s'\) to determine which simplex contains \(s'\). The utility at \(s'\) can then be expressed by a unique linear combination of the vertices of that simplex.

Example 8.1 provides an example of simplex interpolation. The process is implemented in algorithm 8.5.

Consider a three-dimensional simplex given by the permutation \(p = [3, 1, 2]\) such that points within the simplex satisfy \(0 \leq x_3 \leq x_1 \leq x_2 \leq 1\). This simplex has vertices \((0, 0, 0)\), \((0, 1, 0)\), \((1, 1, 0)\), and \((1, 1, 1)\).

Any point \(s\) belonging to the simplex can thus be expressed by a weighting of the vertices:

\[
\begin{bmatrix}
  s_1 \\
  s_2 \\
  s_3
\end{bmatrix} = w_1 \begin{bmatrix} 0 \\ 0 \\ 0 \end{bmatrix} + w_2 \begin{bmatrix} 0 \\ 1 \\ 0 \end{bmatrix} + w_3 \begin{bmatrix} 1 \\ 1 \\ 0 \end{bmatrix} + w_4 \begin{bmatrix} 1 \\ 1 \\ 1 \end{bmatrix}
\]

We can determine the values of the last three weights in succession:

\[
w_4 = s_3 \quad w_3 = s_1 - w_4 \quad w_2 = s_2 - w_3 - w_4
\]

We obtain \(w_1\) by enforcing that the weights sum to one.

If \(s = [0.3, 0.7, 0.2]\), then the weights are:

\[
w_4 = 0.2 \quad w_3 = 0.1 \quad w_2 = 0.4 \quad w_1 = 0.3
\]
mutable struct SimplexValueFunction
    o # position of lower-left corner
    δ # vector of widths
    θ # vector of values at states in S
end

function (Uθ::SimplexValueFunction)(s)
    Δ = (s - Uθ.o)./Uθ.δ
    # Multidimensional index of upper-right cell
    i = min.(floor.(Int, Δ) .+ 1, size(Uθ.θ) .- 1) .+ 1
    u = 0.0
    s′ = (s - (Uθ.o + Uθ.δ.*(i.-2)))./Uθ.δ
    p = sortperm(s′) # increasing order
    w_tot = 0.0
    for j in p
        w = s′[j] - w_tot
        u += w*Uθ.θ[i...]
        i[j] -= 1
        w_tot += w
    end
    u += (1 - w_tot)*Uθ.θ[i...]
    return u
end

function fit!(Uθ::SimplexValueFunction, S, U)
    Uθ.θ = U
    return Uθ
end

Algorithm 8.5. A method for conducting simplex interpolation to estimate the value of state vector $s$ for known state values $θ$ over a grid defined by a lower-left vertex $o$ and vector of widths $δ$. Vertices of the grid can all be written $o + δ .* i$ for some non-negative integral vector $i$. Simplex interpolation is also implemented in the general GridInterpolations.jl package.
8.6 Linear Regression

A simple global approximation approach is *linear regression*, where \( U_\theta(s) \) is a linear combination of *basis functions*.\(^4\) These basis functions are generally a nonlinear function of the state \( s \) and are combined together into a vector function \( \beta(s) \) or \( \beta(s,a) \), resulting in the approximations:

\[
U_\theta(s) = \theta^\top \beta(s) \quad Q_\theta(s,a) = \theta^\top \beta(s,a)
\]  

(8.14)

Although our approximation is linear with respect to the basis functions, the resulting approximation may be nonlinear with respect to the underlying state variables. Figure 8.12 illustrates this concept. Example 8.2 provides an example of global linear value approximation using polynomial basis functions for the continuous mountain car problem, resulting in a nonlinear value function approximation with respect to the state variables.

[Image of Figure 8.12]

Adding more basis functions generally improves the ability to match the target utilities at the states in \( S \), but too many basis functions can lead to poor approximations at other states. Principled methods exist for choosing an appropriate set of basis functions for our regression model.\(^5\)

Fitting linear models involves determining the vector \( \theta \) that minimizes the squared error of the predictions at the states in \( S = s_{1:m} \). If the utilities associated with those states are denoted \( u_{1:m} \), then we want to find the \( \theta \) that minimizes

\[
\sum_{i=1}^{m} (\hat{U}_\theta(s_i) - u_i)^2 = \sum_{i=1}^{m} (\theta^\top \beta(s_i) - u_i)^2
\]  

(8.15)

\(^4\) Also commonly referred to as *features*.

We can approximate the value function for the mountain car problem using a linear approximation. The problem has a continuous state space with two dimensions consisting of position $x$ and speed $v$. Here are the basis functions up to degree six:

$$
\beta(s) = [1, x, v, x^2, xv, v^2, x^3, x^2v, xv^2, v^3, x^4, x^3v, x^2v^2, xv^3, v^4, x^5, x^4v, x^3v^2, x^2v^3, xv^4, v^5, x^6, x^5v, x^4v^2, x^3v^3, x^2v^4, xv^5, v^6]
$$

Below is a plot of an approximate value function fit to state-value pairs from an expert policy:

Example 8.2. An example of using a linear approximation to the mountain car value function. The choice of basis functions makes a big difference. The optimal value function for the mountain car is nonlinear, with a spiral shape and discontinuities. Even sixth-degree polynomials do not produce a perfect fit.
The optimal $\theta$ can be computed through some simple matrix operations. We first construct a matrix $X$ where each of the $m$ rows $X_i =$ contains $\beta(s_i)$. It can be shown that the value of $\theta$ that minimizes the squared error is:

$$\theta = \left(X^TX\right)^{-1}X^T u_{1:m} = X^+ u_{1:m} \tag{8.16}$$

where $X^+$ is the Moore-Penrose pseudoinverse of matrix $X$. The pseudoinverse is often implemented by first computing the singular value decomposition, $X = U \Sigma U^*$. We then have

$$X^+ = U \Sigma^+ U^* \tag{8.17}$$

The pseudoinverse of the diagonal matrix $\Sigma$ is obtained by taking the reciprocal of each nonzero element of the diagonal and then transposing the result.

Figure 8.13 shows how the utilities of states in $S$ are fit with several different basis function families. Different choices of basis functions result in different errors.

Algorithm 8.6 provides an implementation for evaluating and fitting linear regression models of the value function. Example 8.3 demonstrates this approach on the mountain car problem.

### 8.7 Neural Network Regression

Neural network regression relieves us of having to construct an appropriate set of basis functions as required in linear regression. Instead, a neural network is used to represent our value function. For a review of neural networks, see appendix D. The input to the neural network would be the state variables, and the output
### 8.8 Summary

- For large or continuous problems, we can attempt to find approximate policies represented by parameterized models of the value function.

- The approaches taken in this chapter involve iteratively applying steps of dynamic programming at a finite set of states and refining our parametric approximation.

- Local approximation techniques approximate the value function based on the values of nearby states with known values.

- A variety of local approximation techniques include nearest neighbor, kernel smoothing, linear interpolation, and simplex interpolation.
We can apply linear regression to learn a value function for the mountain car problem. The optimal value function has the form of a spiral, which can be difficult to approximate with polynomial basis functions (see example 8.2). We use Fourier basis functions whose components have the form:

\[
\begin{align*}
    b_0(x) &= \frac{1}{2} \\
    b_{s,i}(x) &= \sin(2\pi i x / T) \quad \text{for } i = 1, 2, \ldots \\
    b_{c,i}(x) &= \cos(2\pi i x / T) \quad \text{for } i = 1, 2, \ldots
\end{align*}
\]

where \(T\) is the width of the component’s domain. The multidimensional Fourier basis functions are all combinations of the one-dimensional components across the state-space axes. Below we use a 8th order approximation, so \(i\) ranges up to 8. The expert policy is to accelerate in the direction of motion.

Example 8.3. Linear regression using Fourier bases used to approximate the value function for the mountain car problem (appendix F.4). Value functions (top row) and resulting policies (bottom row) are shown. The globally approximated value function is a poor fit despite using 8th order Fourier basis functions. The resulting approximate policy is not a close approximation to the expert policy. The small timestep in the mountain car problem causes even small changes in the value function landscape to affect the policy. Optimal utility functions often have complex geometries that can be difficult to capture with global basis functions.
• Global approximation techniques include linear regression and neural network regression.

• Nonlinear utility functions can be obtained when using linear regression when combined with an appropriate selection of nonlinear basis functions.

• Neural network regression relieves us of having to specify basis functions, but fitting them is more complex and generally requires us to use gradient descent to tune our parametric approximation of the value function.

8.9 Exercises

Exercise 8.1. A tabular representation is a special case of linear approximate value functions. Show how, for any discrete problem, a tabular representation can be framed as a linear approximate value function.

Solution: Consider a discrete MDP with $m$ states $s_{1:m}$ and $n$ actions $a_{1:n}$. A tabular representation associates a value with each state or state-action pair. We can recover the same behavior using a linear approximate value function. We associate an indicator function with each state or state-action pair, whose value is $1$ when the input is the given state or state-action pair and $0$ otherwise:

$$
\beta_i(s) = (s = s_i) = \begin{cases} 
1 & \text{if } s = s_i \\
0 & \text{otherwise}
\end{cases}
$$
or

$$
\beta_{ij}(s, a) = ((s, a) = (s_i, a_j)) = \begin{cases} 
1 & \text{if } (s, a) = (s_i, a_j) \\
0 & \text{otherwise}
\end{cases}
$$

Exercise 8.2. Suppose we have a problem with continuous state and action spaces and that we would like to construct both a linear approximation and a global approximation of the action value function $Q(s, a) = \theta^T \beta(s, a)$. For global approximation, we choose the basis functions

$$
\beta(s, a) = [1, s, a, s^2, sa, a^2]
$$

Given a set of one-hundred states $S = s_{1:100}$ and a set of five actions $A = a_{1:5}$, how many parameters are in $\theta$ for a local approximation method? How many parameters are in $\theta$ for the specified global approximation method?

Solution: In local approximation methods, the state-action values are the parameters. We will have $|S| \times |A| = 100 \times 5 = 500$ parameters in $\theta$. In global approximation methods, the coefficients of the basis functions are the parameters. Since there are six components in $\beta(s, a)$, we will have six parameters in $\theta$. 
Exercise 8.3. We are given the following states \( s_1 = (4, 5), s_2 = (2, 6), \) and \( s_3 = (-1, -1), \) and their corresponding values, \( U(s_1) = 2, U(s_2) = 10, \) and \( U(s_3) = 30. \) Compute the value at state \( s = (1, 2) \) using 2-nearest neighbor local approximation with an \( L_1 \) distance metric, with an \( L_2 \) distance metric, and with an \( L_\infty \) distance metric.

Solution: We tabulate the distances from \( s \) to the points \( s' \in S \) below:

<table>
<thead>
<tr>
<th>( s' \in S )</th>
<th>( L_1 )</th>
<th>( L_2 )</th>
<th>( L_\infty )</th>
</tr>
</thead>
<tbody>
<tr>
<td>( s_1 = (4, 5) )</td>
<td>6</td>
<td>( \sqrt{18} )</td>
<td>3</td>
</tr>
<tr>
<td>( s_2 = (2, 6) )</td>
<td>5</td>
<td>( \sqrt{17} )</td>
<td>4</td>
</tr>
<tr>
<td>( s_3 = (-1, -1) )</td>
<td>5</td>
<td>( \sqrt{13} )</td>
<td>3</td>
</tr>
</tbody>
</table>

Using the \( L_1 \) norm, we estimate \( U(s) = (10 + 30) / 2 = 20. \) Using the \( L_2 \) norm, we estimate \( U(s) = (10 + 30) / 2 = 20. \) Using the \( L_\infty \) norm, we estimate \( U(s) = (2 + 30) / 2 = 16. \)

Exercise 8.4. We would like to estimate the value at a state \( s \) given the values at a set of two states \( S = \{s_1, s_2\}. \) If we want to use local approximation value iteration, which of the following weighting functions are valid? If they are invalid, how could the weighting functions be modified to make them valid?

- \( \beta(s) = [1, 1] \)
- \( \beta(s) = [1 - \lambda, \lambda] \) where \( \lambda \in [0, 1] \)
- \( \beta(s) = \left[ e^{(s-s_1)^2}, e^{(s-s_2)^2} \right] \)

Solution: The first set of weighting functions is not valid, as it violates the constraint \( \sum \beta_i(s) = 1. \) We can modify the weighting functions by normalizing them by their sum

\[
\beta(s) = \left[ \frac{1}{1+1}, \frac{1}{1+1} \right] = \left[ \frac{1}{2}, \frac{1}{2} \right]
\]

The second set of weighting functions is valid. The third set of weighting functions is not valid, as it violates the constraint \( \sum \beta_i(s) = 1. \) We can modify the weighting functions by normalizing them by their sum

\[
\beta(s) = \left[ \frac{e^{(s-s_1)^2}}{e^{(s-s_1)^2} + e^{(s-s_2)^2}}, \frac{e^{(s-s_2)^2}}{e^{(s-s_1)^2} + e^{(s-s_2)^2}} \right]
\]

Exercise 8.5. Prove that bilinear interpolation is invariant under (nonzero) linear grid scaling.
Solution: It is straightforward to show that the interpolated value is invariant to a linear scaling on one or both axes, e.g. \( \tilde{U}_\theta(\tilde{s}) = U_\theta(s) \). We show this by substituting all \( x \)- and \( y \)-values by their scaled versions \( \tilde{x} = \beta x \) and \( \tilde{y} = \gamma y \), and showing that the grid scalings cancel out:

\[
\tilde{U}_\theta(\tilde{s}) = \frac{(\tilde{x}_2 - \tilde{x}_1)(\tilde{y}_2 - \tilde{y}_1)}{(\tilde{x}_2 - \tilde{x}_1)(\tilde{y}_2 - \tilde{y}_1)} \theta_1 + \frac{(\tilde{x}_2 - \tilde{x}_1)(\tilde{y} - \tilde{y}_1)}{(\tilde{x}_2 - \tilde{x}_1)(\tilde{y}_2 - \tilde{y}_1)} \theta_2 + \frac{(\tilde{x} - \tilde{x}_1)(\tilde{y}_2 - \tilde{y}_1)}{(\tilde{x}_2 - \tilde{x}_1)(\tilde{y}_2 - \tilde{y}_1)} \theta_3 + \frac{(\tilde{x} - \tilde{x}_1)(\tilde{y} - \tilde{y}_1)}{(\tilde{x}_2 - \tilde{x}_1)(\tilde{y}_2 - \tilde{y}_1)} \theta_4
\]

\[
U_\theta(s) = \frac{\beta(x_2 - x_1)\gamma(y_2 - y_1)}{\beta(x_2 - x_1)\gamma(y_2 - y_1)} \theta_1 + \frac{\beta(x_2 - x_1)\gamma(y - y_1)}{\beta(x_2 - x_1)\gamma(y_2 - y_1)} \theta_2 + \frac{\beta(x - x_1)\gamma(y_2 - y_1)}{\beta(x_2 - x_1)\gamma(y_2 - y_1)} \theta_3 + \frac{\beta(x - x_1)\gamma(y - y_1)}{\beta(x_2 - x_1)\gamma(y_2 - y_1)} \theta_4
\]

\[
\tilde{U}_\theta(\tilde{s}) = U_\theta(s)
\]

Exercise 8.6. Given the following four states \( s_1 = [0, 5] \), \( s_2 = [0, 25] \), \( s_3 = [1, 5] \), and \( s_4 = [1, 25] \), and a sample state \( s = [0.7, 10] \), generate the interpolant equation \( U_\theta(s) \) for arbitrary \( \theta \).

Solution: The general form for bilinear interpolation is given in equation (8.12) and reproduced below. To generate the interpolant, we substitute our values into the equation and simplify

\[
U_\theta(s) = \frac{(x_2 - x)(y_2 - y)}{(x_2 - x_1)(y_2 - y_1)} \theta_1 + \frac{(x_2 - x)(y - y_1)}{(x_2 - x_1)(y_2 - y_1)} \theta_2 + \frac{(x - x_1)(y_2 - y)}{(x_2 - x_1)(y_2 - y_1)} \theta_3 + \frac{(x - x_1)(y - y_1)}{(x_2 - x_1)(y_2 - y_1)} \theta_4
\]

\[
U_\theta(s) = \frac{(1 - 0.7)(25 - 10)}{(1 - 0)(25 - 5)} \theta_1 + \frac{(1 - 0.7)(10 - 5)}{(1 - 0)(25 - 5)} \theta_2 + \frac{(0.7 - 0)(25 - 10)}{(1 - 0)(25 - 5)} \theta_3 + \frac{(0.7 - 0)(10 - 5)}{(1 - 0)(25 - 5)} \theta_4
\]

\[
U_\theta(s) = \frac{9}{40} \theta_1 + \frac{3}{40} \theta_2 + \frac{21}{40} \theta_3 + \frac{7}{40} \theta_4
\]

Exercise 8.7. Following example 8.1, what are the simplex interpolant weights for a state \( s = [0.4, 0.95, 0.6] \)?

Solution: For the given state \( s \), we have \( 0 \leq x_1 \leq x_3 \leq x_2 \leq 1 \), and so our permutation vector is \( p = [1, 3, 2] \). The vertices of our simplex can be generated by starting from \((0, 0, 0)\) and changing each 0 to a 1 in reverse order of the permutation vector. Thus, the vertices of the simplex are \((0, 0, 0), (0, 1, 0), (0, 1, 1), \) and \((1, 1, 1)\).

Any point \( s \) belonging to the simplex can thus be expressed by a weighting of the vertices

\[
\begin{bmatrix}
  s_1 \\
  s_2 \\
  s_3
\end{bmatrix} = w_1 \begin{bmatrix} 0 \\
  0 \\
  0
\end{bmatrix} + w_2 \begin{bmatrix} 0 \\
  1 \\
  0
\end{bmatrix} + w_3 \begin{bmatrix} 0 \\
  0 \\
  1
\end{bmatrix} + w_4 \begin{bmatrix} 1 \\
  1 \\
  1
\end{bmatrix}
\]
We can determine the values of the weights in reverse order, finally solving for \( w_1 \) by applying the constraint that the weights must sum to one. We can then compute the weights for \( s = [0.4, 0.95, 0.6] \)

\[
\begin{align*}
    w_4 &= s_1 \\
    w_3 &= s_3 - w_4 \\
    w_2 &= s_2 - w_3 - w_4 \\
    w_1 &= 1 - w_2 - w_3 - w_4 \\
    w_4 &= 0.4 \\
    w_3 &= 0.2 \\
    w_2 &= 0.35 \\
    w_1 &= 0.05
\end{align*}
\]
9 Online Planning

The solution methods we have discussed so far compute policies offline, before any actions are executed in the real problem. Even offline approximation methods can be intractable in many high-dimensional problems. This chapter discusses online planning methods that find actions based on reasoning about states reachable from the current state. The reachable state space is often orders of magnitude smaller than the full state space, which can significantly reduce storage and computational requirements compared to offline methods. We will discuss a variety of algorithms that aim to make online planning efficient, including pruning the state space, sampling, and planning more deeply along trajectories that appear more promising.

9.1 Receding Horizon Planning

In receding horizon planning, we plan from the current state to some maximum fixed horizon or depth $d$. We then execute the action from our current state, transition to the next state, and replan. The online planning methods discussed in this chapter follow this receding horizon planning scheme. They differ in how they explore different courses of action.

A challenge in applying receding horizon planning is determining the appropriate depth. Deeper planning generally requires more computation. For some problems, a shallow depth can be quite effective; the fact that we replan at each step can compensate for our lack of longer-term modeling. In other problems, greater planning depths may be necessary so that our planner can be driven towards goals or away from unsafe states as illustrated in example 9.1.
Suppose we want to apply receding horizon planning to aircraft collision avoidance. The objective is to provide descend or climb advisories when necessary to avoid collision. A collision occurs when our altitude relative to the intruder $h$ is within $\pm 50$ m and the time to potential collision $t_{\text{col}} = 0$. We want to plan deeply enough so that we can provide an advisory sufficiently early to avoid collision with a high degree of confidence. The plots below show the actions that would be taken by a receding horizon planner with different depths.

If the depth is $d = 10$, we only provide advisories within 10 s of collision. Due to the limitations of the vehicle dynamics and the uncertainty of the behavior of the other aircraft, providing advisories this late compromises safety. With $d = 20$, we can do better, but there are cases where we would want to alert a little earlier to further reduce collision risk. There is no motivation to plan deeper than $d = 40$ because we do not need to advise any maneuvers that far in advance of potential collision.
9.2 Lookahead with Rollouts

The previous chapter involved extracting policies that are greedy with respect to an approximate value function $U$ through the use of one-step lookahead. A simple online strategy involves acting greedily with respect to values estimated through simulation to depth $d$. In order to run a simulation, we need a policy to simulate. Of course, we do not know the optimal policy, but we can use what is called a rollout policy instead. Rollout policies are typically stochastic, with actions drawn from a distribution $a \sim \pi(s)$. To produce these rollout simulations, we use a generative model $s' \sim T(s,a)$ to generate successor states $s'$ from the distribution $T(s' \mid s,a)$. This generative model can be implemented through draws from a random number generator, which can be easier to implement in practice compared to explicitly representing the distribution $T(s' \mid s,a)$.

Algorithm 9.1 combines one-step lookahead with values estimated through rollout. This approach often results in better behavior than that of the original rollout policy, but optimality is not guaranteed. It can be viewed as an approximate form of policy improvement used in the policy iteration algorithm (section 7.4). A simple variation of this algorithm is to use multiple rollouts to arrive at a better estimate of the expected discounted return. If we run $m$ simulations to estimate the action values, the time complexity is $O(m \times |A| \times d)$.

9.3 Forward Search

Forward search determines the best action to take from an initial state $s$ by expanding all possible transitions up to depth $d$. These expansions form a search tree. Such search trees have a worst-case branching factor of $|S| \times |A|$, yielding a computational complexity of $O((|S| \times |A|)^d)$. Figure 9.1 shows a search tree applied to a problem with three states and two actions.

Algorithm 9.2 calls itself recursively to the specified depth. Once reaching the specified depth, it uses an estimate of the utility provided by the function $U$. If we simply want to plan to the specified horizon, we set $U(s) = 0$. If our problem requires planning beyond the depth we can afford to compute online, we can use an estimate of the value function obtained offline using, for example, one of the value function approximations described in the previous chapter. Combining online and offline approaches in this way is sometimes referred to as hybrid planning.

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2021-03-07 23:12:00-08:00, revision 80bf4d6, comments to bugs@algorithmsbook.com
Algorithm 9.1. A function that runs a rollout of policy $\pi$ in problem $\mathcal{P}$ from state $s$ to depth $d$. It returns the total discounted reward. This function can be used with the `greedy` function (introduced in algorithm 7.5) to generate an action that is likely to be an improvement over the original rollout policy. We will use this algorithm later for problems other than MDPs, requiring us to only have to modify `randstep` appropriately.

Figure 9.1. A forward search tree for a problem with three states and two actions.
struct ForwardSearch
    ℙ # problem
    d # depth
    U # value function at depth d
end

function forward_search(ℙ, s, d, U)
    if d ≤ 0
        return (a=nothing, u=U(s))
    end
    best = (a=nothing, u=-Inf)
    U′(s) = forward_search(ℙ, s, d-1, U).u
    for a in ℙ.𝒜
        u = lookahead(ℙ, U′, s, a)
        if u > best.u
            best = (a=a, u=u)
        end
    end
    return best
end

(π:ForwardSearch)(s) = forward_search(π.ℙ, s, π.d, π.U).a

Algorithm 9.2. The forward search algorithm for finding an approximately optimal action online for a problem ℙ from a current state s. The search is performed to depth d, at which point the terminal value is estimated with an approximate value function U. The returned named tuple consists of the best action a and its finite-horizon expected value u. The problem type is not constrained to be an MDP; section 22.2 uses this same algorithm in the context of partially observable problems with a different implementation for lookahead.

Figure 9.2. Forward search applied to the hex world problem with four different maximum depths. The search can visit a node multiple times. The actions and colors for visited states were chosen according to the shallowest, highest-value node in the search tree for that state. The initial state has an additional black border.
9.4 Branch and Bound

Branch and bound (algorithm 9.3) attempts to avoid the exponential computational complexity of forward search. It prunes branches by reasoning about bounds on the value function. The algorithm requires knowing a lower bound on the value function \( U(s) \) and an upper bound on the action value function \( Q(s, a) \). The lower bound is used to evaluate the states at the maximum depth. This lower bound is propagated upwards through the tree through Bellman updates. If we find that the upper bound of an action at a state is lower than the lower bound of a previously explored action from that state, then we need not explore that action, allowing us to prune the associated subtree from consideration.

```
struct BranchAndBound
  ℙ   # problem
  d   # depth
  Ulo # lower bound on value function at depth d
  Qhi # upper bound on action value function
end

function branch_and_bound(ℙ, s, d, Ulo, Qhi)
  if d <= 0
    return (a=nothing, u=Ulo(s))
  end
  U'(s) = branch_and_bound(ℙ, s, d-1, Ulo, Qhi).u
  best = (a=nothing, u=-Inf)
  for a in sort(ℙ.𝒜, by=a→Qhi(s,a), rev=true)
    if Qhi(s, a) < best.u
      return best # safe to prune
    end
    u = lookahead(ℙ, U', s, a)
    if u > best.u
      best = (a=a, u=u)
    end
  end
  return best
end

(π::BranchAndBound)(s) = branch_and_bound(π.ℙ, s, π.d, π.Ulo, π.Qhi).a
```

Algorithm 9.3. The branch and bound algorithm for finding of an approximately optimal action online for a discrete MDP ℙ from a current state s. The search is performed to depth d with value function lower bound Ulo and action value function upper bound Qhi. The returned named tuple consists of the best action a and its finite-horizon expected value u. This algorithm is also used for POMDPs.

Branch and bound will give the same result as forward search, but it can be more efficient depending on how many branches are pruned. The worst-case complexity of branch and bound is still the same as forward search. To facilitate
pruning, actions are traversed in descending order by upper bound. Tighter bounds will generally result in more pruning as shown in example 9.2.

Consider applying branch and bound to the mountain car problem. We can use the value function of a heuristic policy for the lower bound $U(s)$, such as a heuristic policy that always accelerates in the direction of motion. For our upper bound $Q([x, v], a)$, we can use the return expected when accelerating toward the goal with no hill. Branch and bound visits about a third as many states as forward search.

Example 9.2. An example of branch and bound applied to the mountain car problem (appendix F.4). Branch and bound can achieve a significant speedup over forward search.

### 9.5 Sparse Sampling

A method known as *sparse sampling*\(^3\) (algorithm 9.4) attempts to reduce the branching factor of forward search and branch and bound. Instead of branching on all possible next states, we only consider a limited number of samples of the next state. Although the sampling of the next state results in an approximation, this method can work well in practice and can significantly reduce computation. If we draw $m$ samples of the next state for each action node in the search tree, the computational complexity is $O\left((m \times |A|)^d\right)$, which is still exponential in the depth but no longer depends on the size of the state space. Figure 9.3 shows an example.

### 9.6 Monte Carlo Tree Search

*Monte Carlo tree search* (algorithm 9.5) avoids the exponential complexity in the horizon by running $m$ simulations from the current state.\(^4\) During these simulations, the algorithm updates estimates of the action value function $Q(s, a)$ and a record of the number of times a particular state-action pair has been selected, $N(s, a)$. After running these $m$ simulations from our current state $s$, we simply choose the action that maximizes our estimate of $Q(s, a)$.

A simulation (algorithm 9.6) begins by traversing the explored state space, consisting of the states for which we have estimates of $Q$ and $N$. We follow some exploration strategy to choose actions from the various states. A common approach is to select the action that maximizes the $UCB1$ exploration heuristic:\(^5\)


Algorithm 9.4. The sparse sampling algorithm for finding of an approximately optimal action online for a discrete problem $\mathcal{P}$ from a current state $s$ to depth $d$ with $m$ samples per action. The returned named tuple consists of the best action $a$ and its finite-horizon expected value $u$.

Figure 9.3. Sparse sampling with $m = 10$ applied to the hex world problem. Tiles are colored according to their value if they were visited. The bordered tile is the initial state. Compare to forward search in figure 9.2.
Algorithm 9.5. The Monte Carlo tree search policy for finding of an approximately optimal action from a current state $s$. This two-argument version of $\text{argmax}$ is defined in appendix G.5.

$$Q(s, a) + c \sqrt{\frac{\log N(s)}{N(s, a)}}$$

(9.1)

where $N(s) = \sum_a N(s, a)$ is the total visit count to $s$ and $c$ is an exploration parameter that scales the value of unexplored actions. The second term corresponds to an exploration bonus. If $N(s, a) = 0$, the bonus is defined to be infinity. With $N(s, a)$ in the denominator, the exploration bonus is higher for actions that have not been tried as frequently. Algorithm 9.7 implements this exploration strategy. We will discuss many other exploration strategies later in chapter 15.

As we take actions specified by algorithm 9.7, we step into new states sampled from the generative model $T(s, a)$, similar to the sparse sampling method. We increment the visit count $N(s, a)$ and update $Q(s, a)$ to maintain the mean value.

At some point, we will either reach the maximum depth or a state that we have not yet explored. If we reach an unexplored state $s$, we initialize $N(s, a)$ and $Q(s, a)$ to zero for each action $a$. We may modify algorithm 9.6 to initialize these counts and value estimates to some other values based on prior expert knowledge of the problem. After initializing $N$ and $Q$, we then arrive at a value estimate through rollout of some policy using the process outlined in section 9.2.

Examples 9.3 to 9.7 work through an illustration of Monte Carlo tree search applied to the 2048 problem. Figure 9.4 shows a search tree generated by running Monte Carlo tree search on 2048. Example 9.8 discusses the impact of using different strategies for estimating values.
Consider using Monte Carlo tree search to play 2048 (appendix F.2) with a maximum depth $d = 10$, an exploration parameter $c = 100$, and a uniform random rollout policy. Our first simulation expands the starting state. The count and value are initialized for each action from the initial state:

![2048 Starting State](image)

**Algorithm 9.6.** A simulation in Monte Carlo tree search starting from state $s$ to depth $d$.

**Algorithm 9.7.** An exploration policy used in Monte Carlo tree search when determining which nodes to traverse through the search tree. The policy is determined by a dictionary of state-action visitation counts $N$ and values $Q$, as well as an exploration parameter $c$. When $N[(s,a)] = 0$, the policy returns infinity.

**Example 9.3.** An example of solving 2048 with Monte Carlo tree search.
The second simulation begins by selecting the best action from the initial state according to our exploration strategy in equation (9.1). Because all states have the same value, we arbitrarily choose the first action, left. We then sample a new successor state and expand it, initializing the associated counts and value estimates. A rollout is run from the successor state and its value is used to update the value of left.

Example 9.4. A (continued) example of solving 2048 with Monte Carlo tree search.
The third simulation begins by selecting the second action, down, because it has infinite value due to the exploration bonus given for unexplored actions. The first action has finite value:

\[
Q(s_0, \text{left}) + c\sqrt{\frac{\log N(s_0)}{N(s_0, \text{left})}} = 48 + 100\sqrt{\frac{\log 2}{1}} \approx 117.315
\]

We take the down action and sample a new successor state, which is expanded. A rollout is run from the successor state and its value is used to update the value of down.

Example 9.5. A (continued) example of solving 2048 with Monte Carlo tree search.
The next two simulations select right and up, respectively. This results in:

Example 9.6. A (continued) example of solving 2048 with Monte Carlo tree search.
Example 9.7. A (continued) example of solving 2048 with Monte Carlo tree search.

There are variations of this basic Monte Carlo tree search algorithm to better handle large action and state spaces. Instead of expanding all of the actions, we can use progressive widening. The number of actions considered from state $s$ is limited to $\theta_1 N(s)^{\theta_2}$, where $\theta_1$ and $\theta_2$ are hyper-parameters. Similarly, we can limit the number of states that result from taking action $a$ from state $s$ in the same way, using what is called double progressive widening. If the number of states that have been simulated from state $s$ after action $a$ is below $\theta_3 N(s,a)^{\theta_4}$, then we sample a new state; otherwise, we sample one of the previously sampled states with probability proportional to the number of times it has been visited. This strategy can be used to handle large as well as continuous action and state spaces.6

9.7 Heuristic Search

Heuristic search (algorithm 9.8) uses $m$ simulations of a greedy policy with respect to a value function $U$ from the current state $s$.7 The value function $U$ is initialized to an upper bound of the value function $\bar{U}$, which is referred to as a heuristic. As we run these simulations, we update our estimate of $U$ through lookahead. After running these simulations, we simply select the greedy action from $s$ with respect

---


Figure 9.4. A Monte Carlo tree search tree on 2048 after 100 simulations. In general, Monte Carlo tree search for MDPs produces a search graph because there can be multiple ways to reach the same state. The colors in the tree indicate the estimated values at the nodes, with high values in blue and low values in red. The tree is shallow with a fairly high branching factor because 2048 has many reachable states for each action.
Rollouts are not the only means by which we can estimate utilities in Monte Carlo tree search. Custom evaluation functions can often be crafted for specific problems to help guide the algorithm. For example, we can encourage Monte Carlo tree search to order its tiles in 2048 using evaluation functions that return the weighted sum across tile values:


heuristic A weights


heuristic B weights


The plot below compares Monte Carlo tree search on 2048 using rollouts with a uniform random policy, rollouts with a one-step lookahead policy, the two evaluation functions above, and using the current board score.

Example 9.8. The performance of Monte Carlo tree search varies with the number of simulations and as the board evaluation method is changed. Heuristic board evaluations tend to be efficient and can more effectively guide search when run counts are low. Lookahead rollout evaluations take about 18 times longer than heuristic evaluations.
9.8 Labeled Heuristic Search

Labeled heuristic search (algorithm 9.9) is a variation of heuristic search that runs simulations with value updates while labeling states based on whether their value is solved.\(^8\) We say that a state \(s\) is solved if their utility residual falls below some threshold \(\delta > 0:\)

\[
|U_{k+1}(s) - U_k(s)| < \delta
\]  

(9.2)

We run simulations with value updates until the current state is solved. In contrast with the heuristic search in the previous section that runs a fixed number of iterations, this labeling process focuses computational effort in the most important areas of the state space.

Simulations in labeled heuristic search (algorithm 9.10) begin by running to a maximum depth of $d$ by following a policy that is greedy with respect to our estimated value function $U$, similar to the heuristic search in the previous section. We may stop a simulation before a depth of $d$ if we reach a state that has been labeled as solved in a prior simulation.
### Algorithm 9.10. Simulations are run from the current state to a maximum depth \( d \). We stop a simulation at depth \( d \) or if we encounter a state that is in the set \( \text{solved} \). After a simulation, we call \( \text{label!} \) on the states we visited in reverse order.

```plaintext
function simulate!(π::LabeledHeuristicSearch, U, solved, s)
    visited = []
    for d in 1:π.d
        if s ∈ solved
            break
        end
        push!(visited, s)
        a, u = greedy(π.\( \mathcal{P} \), U, s)
        U[s] = u
        s = rand(π.\( \mathcal{P} \).T(s, a))
    end
    while !isempty(visited)
        if label!(π, U, solved, pop!(visited))
            break
        end
    end
end
```

After each simulation, we iterate over the states we visited during that simulation in reverse order, performing a labeling routine on each state and stopping if a state is found that is not solved. The labeling routine (algorithm 9.11) searches the states in the greedy envelope of \( s \), which is defined to be the states reachable from \( s \) under a greedy policy with respect to \( U \). The state \( s \) is considered not solved if there is a state in the greedy envelope of \( s \) whose utility residual is greater than the threshold \( \delta \). If no such state is found, then \( s \) is marked as solved—as well as all states in the greedy envelope of \( s \) because they must have converged as well. If a state with a sufficiently large utility residual is found, then the utilities all of states traversed during the search of the greedy envelope are updated.

Figure 9.6 shows several different greedy envelopes. Figure 9.7 shows the states traversed in a single iteration of labeled heuristic search. Figure 9.8 shows the progression of heuristic search on the hex world problem.

### 9.9 Open-Loop Planning

The online methods discussed in this chapter as well as the offline methods discussed in the previous chapters are examples of closed-loop planning, which involves accounting for future state information in the planning process.\(^{10}\) Often open-loop planning can provide a satisfactory approximation of an optimal closed-loop plan while greatly enhancing computational efficiency by avoiding having

\(^{10}\) The loop in this context refers to the observe-act loop introduced in section 1.1.
Algorithm 9.11. The label! function will attempt to find a state in the greedy envelope of s whose utility residual exceeds some threshold \( \delta \). The function expand computes the greedy envelope of s and determines whether any of those states have utility residuals above the threshold. If a state has a residual that exceeds some threshold, then we update utilities of the states in the envelope. Otherwise, we add that envelope to the set of solved states.

function expand(\( \pi \)::LabeledHeuristicSearch, \( U \), solved, s)
    \( \mathcal{P}, \delta = \pi.\mathcal{P}, \pi.\delta \)
    \( \mathcal{S}, \mathcal{A}, T = \mathcal{P}.\mathcal{S}, \mathcal{P}.\mathcal{A}, \mathcal{P}.T \)
    found, toexpand, envelope = false, Set(s), []
    while !isempty(toexpand)
        s = pop!(toexpand)
        push!(envelope, s)
        a, u = greedy(\( \mathcal{P} \), \( U \), s)
        if abs(\( U[s] - u \)) > \( \delta \)
            found = true
        else
            for s′ in \( \mathcal{S} \)
                if T(s, a, s′) > 0 && s′ \( \notin \) (solved \( \cup \) envelope)
                    push!(toexpand, s′)
                end
            end
        end
    end
    return (found, envelope)
end

function label!(\( \pi \)::LabeledHeuristicSearch, \( U \), solved, s)
    if s \( \in \) solved
        return false
    end
    found, envelope = expand(\( \pi \), \( U \), solved, s)
    if found
        for s in reverse(envelope)
            if U[s] = greedy(\( \mathcal{P} \), \( U \), s).u
                end
            else
                union!(solved, envelope)
            end
        end
    return found
end
end

Figure 9.6. The greedy envelope for \( \delta = 1 \) for several different states visualized for a value function on the hex world problem. The value function was obtained by running basic heuristic search for 10 iterations from the first darkly bordered state to a maximum depth of 8. We find that the size of the greedy envelope can vary widely depending on the state.
Figure 9.7. A single iteration of labeled heuristic search conducts an exploratory run (arrows) followed by labeling (hexagonal border). Only two states are labeled in this iteration: the hidden terminal state and the state with a hexagonal border. Both the exploratory run and the labeling step update the value function.

Figure 9.8. A progression of heuristic search on the hex world problem using $\delta = 1$ and a heuristic $U(s) = 10$. The solved states in each iteration are covered in an opaque gray wash. The set of solved states grows from the terminal reward state back toward the initial state with the dark border.
to reason about the acquisition of future information. Sometimes this open-loop planning approach is referred to as model predictive control.\textsuperscript{11} As with receding horizon control, model predictive control solves the open-loop problem, executes the action from our current state, transition to the next state, and then replans.

Open-loop plans can be represented as a sequence of actions up to some depth $d$. The planning process reduces to an optimization problem

$$\max_{a_{1:d}} U(a_{1:d})$$

where $U(a_{1:d})$ is the expected return when executing the sequence of actions $a_{1:d}$. Depending on the application, this optimization problem may be convex or lends itself to a convex approximation, meaning that it can be solved quickly using a variety of different algorithms.\textsuperscript{12} Later in this section, we will discuss a few different formulations that can be used to transform equation (9.3) into a convex problem.

Open-loop planning can often allow us to devise effective decision strategies in high-dimensional spaces where closed-loop planning is computationally infeasible. Open-loop planning gains this efficiency by not accounting for future information. Example 9.9 provides a simple example of where open-loop planning can result in poor decisions, even when we account for stochasticity.

### 9.9.1 Deterministic Model Predictive Control

A common approximation to make $U(a_{1:d})$ amenable to optimization is to assume deterministic dynamics:

$$\max_{a_{1:d},s_{2:d}} \sum_{t=1}^{d} \gamma^t R(s_t, a_t)$$

subject to $s_{t+1} = T(s_t, a_t), \ t \in 1 : d - 1$

where $s_1$ is the current state and $T(s, a)$ is a deterministic transition function that returns the state that results from taking action $a$ from state $s$. A common strategy for producing a suitable deterministic transition function from a stochastic transition function is to use the most likely transition. If the dynamics in equation (9.4) are linear and the reward function is convex, then the problem is convex.
Consider a problem with nine states, as shown in the margin, with two decision steps starting from the initial state $s_1$. In our decisions, we must decide between going up (blue arrows) and going down (green arrows). The effects of these actions are deterministic, except that if we go up from $s_1$, then we end up in state $s_2$ half the time and in state $s_3$ half the time. We receive a reward of 30 in states $s_5$ and $s_7$ and a reward of 20 in states $s_8$ and $s_9$, as indicated in the figure.

There are exactly four open-loop plans: (up, up), (up, down), (down, up), and (down, down). In this simple example, it is easy to compute their expected utilities:

- $U(\text{up, up}) = 0.5 \times 30 + 0.5 \times 0 = 15$
- $U(\text{up, down}) = 0.5 \times 0 + 0.5 \times 30 = 15$
- $U(\text{down, up}) = 20$
- $U(\text{down, down}) = 20$

According to the set of open-loop plans, it is best to choose down from $s_1$ because our expected reward is 20 instead of 15.

Closed-loop planning, in contrast, takes into account the fact that we can base our next decision on the observed outcome of our first action. If we choose to go up from $s_1$, then we can choose to go down or up depending on whether we end up in $s_2$ or $s_3$, thereby guaranteeing a reward of 30.
Example 9.10 provides an example involving navigating to a goal state while avoiding an obstacle and minimizing acceleration effort. Both the state space and action space are continuous, and we can find a solution in well under a second. Replanning after every step can help compensate for stochasticity or unexpected events. For example, if the obstacle moves, we can readjust our plan as illustrated in figure 9.9.


In this problem, our state $s$ represents our two-dimensional position concatenated with our two-dimensional velocity vector, with $s$ initially set to $[0, 0, 0, 0]$. Our action $a$ is an acceleration vector, where each component must be between $\pm 1$. At each step, we use our action to update our velocity, and we use our velocity to update our position. Our objective is to reach a goal state of $s_{\text{goal}} = [10, 10, 0, 0]$. We plan to $d = 10$ steps with no discounting. With each step, we accumulate a cost of $\|a_t\|^2$ to minimize acceleration effort. At the last step, we want to be as close to the goal state as possible with a penalty of $100\|s_d - s_{\text{goal}}\|^2$. We also have to ensure that we avoid a circular obstacle with radius 2 centered at $[3, 4]$. We can formulate this problem as follows and extract the first action from the plan.

```julia
model = Model(Ipopt.Optimizer)
d = 10
current_state = zeros(4)
goal = [10,10,0,0]
obstacle = [3,4]
@variables model begin
    s[1:4, 1:d]
    -1 ≤ a[1:2, 1:d] ≤ 1
end
# velocity update
@constraint(model, [i=2:d, j=1:2], s[2+j, i] == s[2+j, i-1] + a[j, i-1])
# position update
@constraint(model, [i=2:d, j=1:2], s[j, i] == s[j, i-1] + s[2+j, i-1])
# initial condition
@constraint(model, s[:, 1] .== current_state)
# obstacle
@constraint(model, [i=1:d], sum((s[1:2, i] - obstacle).^2) ≥ 4)
@objective(model, Min, 100*sum((s[:, d] - goal).^2) + sum(a.^2))
optimize!(model)
action = value.(a[:, 1])
```

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2021-03-07 23:12:00-08:00, revision 80bf4d6, comments to bugs@algorithmsbook.com
Figure 9.9. Model predictive control applied to the problem in example 9.10 with the addition of a moving obstacle. The sequence progresses left-to-right, top-to-bottom. Initially, we have a plan that passes to the right of the obstacle, but in the third cell, we see that we must change our mind and pass to the left. We have to maneuver around a little to adjust our velocity vector appropriately with minimal effort. Of course, we could have created a better path (in terms of our utility function) if our planning process knew that the obstacle was moving in a particular direction.
9.9.2 Robust Model Predictive Control

We can change the problem formulation to provide robustness to outcome uncertainty. There are many different robust model predictive control formulations, but one involves choosing the best open-loop plan given the worst-case state transitions. This formulation defines $T(s, a)$ to be an uncertainty set consisting of all possible states that can result from taking action $a$ in state $s$. In other words, the uncertainty set is the support of the distribution $T(\cdot \mid s, a)$. Optimizing with respect to worse-case state transitions requires transforming the optimization problem in equation (9.4) into a minimax or min-max problem:

$$\begin{align*}
\text{maximize} & \quad a_{1:d} \\
\text{minimize} & \quad i, s_{2:d} \\
\text{subject to} & \quad s_{t+1} \in T(s_t, a_t), \quad t \in 1 : d - 1
\end{align*}$$ (9.5)

Unfortunately, this formulation can result in extremely conservative behavior. If we adapt example 9.10 to model the uncertainty in the motion of the obstacle, the accumulation of uncertainty may become quite large even when planning with a relatively short horizon. One way to reduce the accumulation of uncertainty is to restrict the uncertainty set output by $T(s, a)$ to contain only, say, 95% of the probability mass. Another issue with this approach is that the minimax optimization problem is often not convex and difficult to solve.

9.9.3 Multi-Forecast Model Predictive Control

One way to address the computational challenge within the minimax problem in equation (9.5) is to use $m$ forecast scenarios, each of which follows its own deterministic transition function. There are different formulations of this kind of multi-forecast model predictive control, which is a type of hindsight optimization approach. One common approach is to have the deterministic transition functions depend on the step $k$, $T_i(s, a, k)$, which is the same as augmenting the state space to include depth. Example 9.11 demonstrates how this might be done for a linear Gaussian model.

We try to find the best sequence of actions against the worst sampled scenario:

$$\begin{align*}
\text{maximize} & \quad a_{1:d} \\
\text{minimize} & \quad i, s_{2:d} \\
\text{subject to} & \quad s_{k+1} = T_i(s_k, a_k), \quad k \in 1 : d - 1
\end{align*}$$ (9.6)

---


15 It is called hindsight optimization because it represents a solution that is optimizing using knowledge about action outcomes than can only be known in hindsight. E.K.P. Chong, R.L. Givan, and H.S. Chang, “A Framework for Simulation-Based Network Control via Hindsight Optimization,” in IEEE Conference on Decision and Control (CDC), 2000.
Suppose we have a problem with linear Gaussian dynamics:

\[ T(s' \mid s, a) = \mathcal{N}(Ts + Ta, \Sigma) \]

The problem in figure 9.9 is linear with no uncertainty, but if we allow the obstacle to move according to a Gaussian distribution at each step, then the dynamics become linear Gaussian. We can approximate the dynamics using a set of \( m \) forecast scenarios, each consisting of \( d \) steps. We can pull \( m \times d \) samples \( \epsilon_{ik} \sim \mathcal{N}(0, \Sigma) \) and define the deterministic transition functions:

\[ T_i(s, a, k) = Ts + Ta + \epsilon_{ik} \]

This problem can be much easier to solve than the original robust problem.

We can also use a multi-forecast approach to optimize the average case.\(^{16}\) The formulation is similar to equation (9.6), except that we replace the minimization with an expectation and allow different action sequences to be taken for different scenarios with the constraint that the first action must agree:

\[
\begin{align*}
\text{maximize} & \quad 1 \sum_{i=1}^{m} \sum_{k=1}^{d} \gamma^k R(s_k, a_k) \\
\text{subject to} & \quad s_{k+1} = T_i(s_k, a_k, k), \quad k \in 1:d-1, i \in 1:m \\
& \quad a_1^{(i)} = a_1^{(j)}, \quad i \in 1:m, j \in 1:m
\end{align*}
\]  

(9.7)

This formulation can result in robust behavior without being overly conservative while still maintaining computational tractability. Both formulations in equations (9.6) and (9.7) can be made more robust by increasing the number of forecast scenarios \( m \) at the expense of additional computation.

9.10 Summary

- Online methods plan from the current state, focusing computation on states that are reachable.

Example 9.11. Modeling linear Gaussian transition dynamics in multi-forecast model predictive control.

\(^{16}\) This approach was applied to optimizing power flow policies by N. Moehle, E. Busseti, S. Boyd, and M. Wytock, “Dynamic Energy Management,” in Large Scale Optimization in Supply Chains and Smart Manufacturing, Springer, 2019, pp. 69–126.
• Receding horizon planning involves planning to a certain horizon and then replanning with each step.

• Lookahead with rollouts involves acting greedily with respect to values estimated using simulations of a rollout policy; it is computationally efficient compared to other algorithms, but there are no guarantees on performance.

• Forward search considers all state-action transitions up to a certain depth, resulting in computational complexity that grows exponentially in both the number of states and the number of actions.

• Branch and bound uses upper and lower bound functions to prune portions of the search tree that will not lead to a better outcome in expectation.

• Sparse sampling avoids the exponential complexity in the number of states by limiting the number of sampled transitions from every search node.

• Monte Carlo tree search guides search to promising areas of the search space by taking actions that balance exploration with exploitation.

• Heuristic search runs simulations of a policy that is greedy with respect to a value function that is updated along the way using lookahead.

• Labeled heuristic search reduces computation by not reevaluating states whose values have converged.

• Open-loop planning aims to find the best possible sequence of actions and can be computationally efficient if the optimization problem is convex.

9.11 Exercises

Exercise 9.1. Why does branch and bound have the same worst-case computational complexity as forward search?

Solution: In the worst case, branch and bound will never prune, resulting in a traversal of the same search tree as forward search with the same complexity.

Exercise 9.2. Given two admissible heuristics $h_1$ and $h_2$, how can we use both of them in heuristic search?

Solution: Create a new heuristic $h(s) = \min\{h_1(s), h_2(s)\}$ and use it instead. This strictly improves a heuristic search algorithm. Moreover, this new heuristic is guaranteed to be admissible. Both $h_1(s) \geq U^*(s)$ and $h_2(s) \geq U^*(s)$ imply that $h(s) \geq U^*(s)$. 
Exercise 9.3. Given two inadmissible heuristics \( h_1 \) and \( h_2 \), describe a way we can use both of them in heuristic search.

Solution: Would could define a new heuristic \( h_3(s) = \max(h_1(s), h_2(s)) \) to get a potentially admissible or a “less-inadmissible” heuristic. It may be slower to converge, but it may be more likely to not miss out on a better solution.

Exercise 9.4. Suppose we have a discrete MDP with state space \( S \) and action space \( A \) and we want to perform forward search to depth \( d \). Due to computational constraints and the requirement that we must simulate to depth \( d \), we decide to generate new, smaller state and action spaces by re-discretizing the original state and action spaces on a coarser scale with \( |S'| < |S| \) and \( |A'| < |A| \). In terms of the original state and action spaces, what would the size of the new state and action spaces need to be in order to make the computational complexity of forward search approximately depth-invariant with respect to the size of our original state and action spaces, i.e. \( O(|S||A|) \)?

Solution: We need

\[
|S'| = |S|^{\frac{1}{d}} \quad \text{and} \quad |A'| = |A|^{\frac{1}{d}}
\]

This results in the following complexity:

\[
O(|S'|^d|A'|^d) = O \left( \left( |S|^{\frac{1}{d}} \right)^d \left( |A|^{\frac{1}{d}} \right)^d \right) = O(|S||A|)
\]

Exercise 9.5. Building upon the previous exercise, suppose now that we want to keep all of the original actions in our action space and only re-discretize the state space. What would the size of the new state space need to be to make the computational complexity of forward search approximately depth-invariant with respect to the size of our original state and action spaces?

Solution: The computational complexity of forward search is given by \( O \left( (|S||A|)^d \right) \), which can also be written as \( O \left( |S|^d|A|^d \right) \). Thus, in order for our coarser state space to lead to forward search that is approximately depth-invariant with respect to the size of our original state and action spaces, we need

\[
|S'| = \left( \frac{|S|}{|A|^{d-1}} \right)^{\frac{1}{d}}
\]

This gives us:

\[
O \left( |S'|^d|A'|^d \right) = O \left( \left( \frac{|S|}{|A|^{d-1}} \right)^{\frac{1}{d}} |A|^d \right) = O \left( |S| \frac{|A|^d}{|A|^{d-1}} \right) = O(|S||A|)
\]

Exercise 9.6. Will changing the ordering of the action space cause forward search to take different actions? Will changing the ordering of the action space cause branch and bound to take different actions? Can the ordering of the action space affect how many states are visited by branch and bound?
Solution: Forward search enumerates over all possible future actions. It may return different actions if there are ties in their expected utilities. Branch and bound maintains the same optimality guarantee over the horizon as forward search by sorting by upper bound. The ordering of the action space can affect branch and bound’s visitation rate when the upper bound produces the same expected value for two or more actions. Below we show this effect on the modified mountain car problem from example 9.2. The plot compares the number of states visited in forward search to that of branch and bound for different action orderings to depth 6. Branch and bound consistently visits far fewer states than forward search, but action ordering can still affect state visitation.

Exercise 9.7. Is sparse sampling with \( m = |S| \) equivalent to forward search?

Solution: No. While the computational complexities are identical at \( O\left(|S|^d |A|^d \right) \), forward search will branch on all states in the state space, while sparse sampling will branch on \( |S| \) randomly sampled states.

Exercise 9.8. Given an MDP with \( |S| = 10, |A| = 3 \), and a uniform transition distribution \( T(s' | s, a) = 1/|S| \) for all \( s \) and \( a \), what is probability that sparse sampling with \( m = |S| \) samples and depth \( d = 1 \) yields the exact same search tree produced by forward search with depth \( d = 1 \)?

Solution: For both forward search and sparse sampling, we branch on all actions from the current state node. For forward search, at each of these action nodes, we branch on all states, while for sparse sampling, we will branch on \( m = |S| \) sampled states. If these sampled states are exactly equal to the state space, that action branch is equivalent to the branch produced in forward search. Thus, for a single action branch we have:

\[
\begin{align*}
\text{the probability the first state is unique} & \quad \frac{10}{10} \\
\text{the probability the second state is unique (not equal to the first state)} & \quad \frac{9}{10} \\
\text{the probability the third state is unique (not equal to the first or second state)} & \quad \frac{8}{10} \\
\vdots & \quad \vdots
\end{align*}
\]
Since each of these sampled states is independent, this leads to the probability of all unique states in the state space being selected with probability

\[
\frac{10 \times 9 \times 8 \times \cdots}{10 \times 10 \times 10 \times \cdots} = \frac{10!}{10^{10}} \approx 0.000363
\]

Since each of the sampled states across different action branches is independent, the probability that all three action branches sample the unique states in the state space is

\[
\left( \frac{10!}{10^{10}} \right)^3 \approx (0.000363)^3 \approx 4.78 \times 10^{-11}
\]

**Exercise 9.9.** Given the following tables of \(Q(s,a)\) and \(N(s,a)\), use the upper confidence bound in equation (9.1) to compute the optimal action for each state with an exploration parameter of \(c_1 = 10\) and again for \(c_2 = 20\).

<table>
<thead>
<tr>
<th>(Q(s,a_1))</th>
<th>(Q(s,a_2))</th>
<th>(N(s,a_1))</th>
<th>(N(s,a_2))</th>
</tr>
</thead>
<tbody>
<tr>
<td>(s_1)</td>
<td>10</td>
<td>-5</td>
<td>(s_1)</td>
</tr>
<tr>
<td>(s_2)</td>
<td>12</td>
<td>10</td>
<td>(s_2)</td>
</tr>
</tbody>
</table>

**Solution:** For the first exploration parameter \(c_1 = 10\), we tabulate the upper confidence bound of each state-action pair and select the action maximizing the bound for each state:

<table>
<thead>
<tr>
<th>(UCB(s,a_1))</th>
<th>(UCB(s,a_2))</th>
<th>(\text{arg max}_a UCB(s,a))</th>
</tr>
</thead>
<tbody>
<tr>
<td>(s_1) 10 + 10√(log 31/27) (\approx 13.566)</td>
<td>-5 + 10√(log 31/4) (\approx 4.266)</td>
<td>(a_1)</td>
</tr>
<tr>
<td>(s_2) 12 + 10√(log 50/32) (\approx 15.496)</td>
<td>10 + 10√(log 50/18) (\approx 14.662)</td>
<td>(a_1)</td>
</tr>
</tbody>
</table>

And for \(c_2 = 20\), we have:

<table>
<thead>
<tr>
<th>(UCB(s,a_1))</th>
<th>(UCB(s,a_2))</th>
<th>(\text{arg max}_a UCB(s,a))</th>
</tr>
</thead>
<tbody>
<tr>
<td>(s_1) 10 + 20√(log 31/27) (\approx 17.133)</td>
<td>-5 + 20√(log 31/4) (\approx 13.531)</td>
<td>(a_1)</td>
</tr>
<tr>
<td>(s_2) 12 + 20√(log 50/32) (\approx 18.993)</td>
<td>10 + 20√(log 50/18) (\approx 19.324)</td>
<td>(a_2)</td>
</tr>
</tbody>
</table>
10 Policy Search

Policy search involves searching the space of policies without directly computing a value function. The policy space is often lower-dimensional than the state space and can often be searched more efficiently. Policy optimization optimizes the parameters in a parameterized policy to maximize utility. This parameterized policy can take many different forms, such as neural networks, decision trees, and computer programs. This chapter begins by discussing a way to estimate the value of a policy given an initial state distribution. We will then discuss search methods that do not use estimates of the gradient of the policy, saving gradient methods for the next chapter. Although local search can be quite effective in practice, we will discuss a few alternative optimization approaches that can avoid local optima.¹

10.1 Approximate Policy Evaluation

As introduced in section 7.2, we can compute the expected discounted return when following a policy \( \pi \) from a state \( s \). This expected discounted return \( U^\pi(s) \) can be computed iteratively (algorithm 7.3) or through matrix operations (algorithm 7.4), when the state space is discrete and relatively small. We can use these results to compute the expected discounted return of \( \pi \):

\[
U(\pi) = \sum_s b(s) U^\pi(s)
\]

assuming an initial state distribution \( b(s) \).

We will use this definition of \( U(\pi) \) throughout this chapter. However, we often cannot compute \( U(\pi) \) exactly when the state space is large or continuous. Instead, we can approximate \( U(\pi) \) by sampling trajectories, consisting of states, actions,

¹There are many other optimization approaches, as discussed by M.J. Kochenderfer and T.A. Wheeler, Algorithms for Optimization. MIT Press, 2019.
and rewards when following \( \pi \). The definition of \( U(\pi) \) can be rewritten
\[
U(\pi) = \mathbb{E}_\tau [R(\tau)] = \int p_\pi(\tau) R(\tau) \, d\tau
\]  
(10.2)
where \( p_\pi(\tau) \) is the probability density associated with trajectory \( \tau \) when following policy \( \pi \) starting from initial state distribution \( b \). The trajectory reward \( R(\tau) \) is the discounted return associated with \( \tau \). Figure 10.1 illustrates the computation of \( U(\pi) \) in terms of trajectories sampled from an initial state distribution.

Monte Carlo policy evaluation (algorithm 10.1) involves approximating equation (10.2) with \( m \) trajectory rollouts of \( \pi \):
\[
U(\pi) \approx \frac{1}{m} \sum_{i=1}^{m} R(\tau^{(i)})
\]  
(10.3)
where \( \tau^{(i)} \) is the \( i \)th trajectory sample.

Monte Carlo policy evaluation is stochastic. Multiple evaluations of equation (10.1) with the same policy can give different estimates. Increasing the number of rollouts decreases the variance of the evaluation, as demonstrated in figure 10.2.

We will use \( \pi_\theta \) to denote a policy parameterized by \( \theta \). For convenience, we will use \( U(\theta) \) as shorthand for \( U(\pi_\theta) \) in cases where it is not ambiguous. The parameter \( \theta \) may be a vector or some other more complex representation. For example, we may want to represent our policy using a neural network with a particular structure. We would use \( \theta \) to represent the weights in the network. Many optimization algorithms assume that \( \theta \) is a vector with a fixed number of components.
Other optimization algorithms allow for more flexible representations, including representations like decision trees or computational expressions.\(^2\)

### 10.2 Local Search

A common approach to optimization is local search, where we begin with an initial parameterization and incrementally move from neighbor to neighbor in the search space until convergence. We discussed this type of approach in chapter 5 in the context of optimizing Bayesian network structures with respect to the Bayesian score. Here, we are optimizing policies parameterized by \(\theta\). We are trying to find a \(\theta\) that maximizes \(U(\theta)\).

There are many different local search algorithms, but this section will focus on the Hooke-Jeeves method (algorithm 10.2).\(^3\) This algorithm assumes that our policy is parameterized by an \(n\)-dimensional vector \(\theta\). The algorithm takes a step of size \(\pm \alpha\) in each of the coordinate directions from the current \(\theta\). These \(2n\) points correspond to the neighborhood of \(\theta\). If no improvements to the policy are found, then the step size \(\alpha\) is decreased by some factor. If an improvement is found, it moves to the best point. The process continues until \(\alpha\) drops below some threshold \(\epsilon > 0\). Figure 10.3 illustrates this process.

### 10.3 Genetic Algorithms

A potential issue with local search algorithms like the Hooke-Jeeves method is that the optimization can get stuck in a local optimum. There are a wide variety of approaches that involve maintaining a population consisting of samples of points in the parameter space, evaluating them in parallel with respect to our objective, and then recombining them in some way to drive the population towards a global optimum. A genetic algorithm\(^4\) is one such approach, which derives inspiration from biological evolution. It is a general optimization method, but it has been successful in the context of optimizing policies. For example, this approach has been used to optimize policies for Atari video games, where the policy parameters correspond to weights in a neural network.\(^5\)

A simple version of this approach (algorithm 10.3) begins with a population of \(m\) random parameterizations, \(\theta^{(1)}, \ldots, \theta^{(m)}\). We compute \(U(\theta^{(i)})\) for each sample \(i\) in the population. Since these evaluations potentially involve many rollout

---

\(^2\) We will not be discussing them here, but some are implemented in ExprOptimization.jl.


Algorithm 10.2. Policy search using the Hooke-Jeeves method, which returns a $\theta$ that has been optimized with respect to $U$. The policy $\pi$ takes as input a parameter $\theta$ and state $s$. This implementation starts with an initial $\theta$. The step size $\alpha$ is reduced by a factor of $c$ if no neighbor improves the objective. Iterations are run until the step size is less than $\epsilon$. 

```plaintext
struct HookeJeevesPolicySearch
    \theta # initial parameterization
    \alpha # step size
    c # step size reduction factor
    \epsilon # termination step size
end

function optimize(M::HookeJeevesPolicySearch, \pi, U)
    \theta, \theta', \alpha, c, \epsilon = copy(M.\theta), similar(M.\theta), M.\alpha, M.c, M.\epsilon
    u, n = U(\pi, \theta), length(\theta)
    while \alpha > \epsilon
        copyto!(\theta', \theta)
        best = (i=0, sgn=0, u=u)
        for i in 1:n
            for sgn in (-1,1)
                \theta'[i] = \theta[i] + sgn*\alpha
                u' = U(\pi, \theta')
                if u' > best.u
                    best = (i=i, sgn=sgn, u=u')
                end
            end
            \theta'[i] = \theta[i]
        end
        if best.i != 0
            \theta[best.i] += best.sgn*\alpha
            u = best.u
        else
            \alpha *= c
        end
    end
    return \theta
end
```
Suppose we want to optimize a policy for the simple regulator problem described in appendix F.5. We define a stochastic policy \( \pi \) parameterized by \( \theta \) such that the action is generated according to

\[
a \sim \mathcal{N}(\theta_1 s, (|\theta_2| + 10^{-5})^2)
\]  

(10.4)

The following code defines the parameterized stochastic policy \( \pi \), evaluation function \( U \), and method \( M \). It then calls \( \text{optimize}(M, \pi, U) \), which returns an optimized value for \( \theta \). In this case, we use the Hooke-Jeeves method, but the other methods discussed in this chapter can be passed in as \( M \) instead.

```python
function \( \pi(\theta, s) \)
    return rand(Normal(\theta[1]*s, abs(\theta[2]) + 0.00001))
end

b, d, n_rollouts = Normal(0.3, 0.1), 10, 3
U = MonteCarloPolicyEvaluation(\( \mathcal{P} \), b, d, n_rollouts)
\( \theta, \alpha, c, \epsilon = [0.0, 1.0], 0.75, 0.75, 0.01 \)
M = HookeJeevesPolicySearch(\( \theta, \alpha, c, \epsilon \))
\( \theta = \text{optimize}(M, \pi, U) \)
```

Example 10.1. An example of using a policy optimization algorithm to optimize the parameters of a stochastic policy.

Figure 10.3. The Hooke-Jeeves method applied to optimizing a policy in the simple regulator problem discussed in example 10.1. The evaluations at each iteration are shown as white points. Iterations proceed left to right and top to bottom, and the background is colored according to the expected utility.
simulations and are therefore computationally expensive, they are often run in parallel. These evaluations help us identify the *elite samples*, which are the top $m_{elite}$ samples according to $U$.

The population at the next iteration is generated by producing $m - 1$ new parameterizations by repeatedly selecting a random elite sample $\theta$ and perturbing it with isotropic Gaussian noise, $\theta + \sigma \epsilon$, where $\epsilon \sim \mathcal{N}(0, I)$. The best parameterization, unperturbed, is included as the $m$th sample. Because the evaluations involve stochastic rollouts, a variation of this algorithm could involve running additional rollouts to help identify which of the elite samples is truly the best. Figure 10.4 shows several iterations, or *generations*, of this approach on an example problem.

```
struct GeneticPolicySearch
    θs      # initial population
    σ       # initial standard deviation
    m_elite # number of elite samples
    k_max   # number of iterations
end

function optimize(M::GeneticPolicySearch, π, U)
    θs, σ = M.θs, M.σ
    n, m = length(first(θs)), length(θs)
    for k in 1:M.k_max
        us = [U(π, θ) for θ in θs]
        sp = sortperm(us, rev=true)
        θ_best = θs[sp[1]]
        rand_elite() = θs[sp[rand(1:M.m_elite)]]
        θs = [rand_elite() + σ.*randn(n) for i in 1:(m-1)]
        push!(θs, θ_best)
    end
    return last(θs)
end
```

Algorithm 10.3. A genetic policy search method for iteratively updating a population of policy parameterizations $\theta$s, which takes a policy evaluation function $U$, a policy $\pi(\theta, s)$, a perturbation standard deviation $\sigma$, an elite sample count $m_{elite}$, and an iteration count $k_{max}$. The best $m_{elite}$ samples from each iteration are used to generate the samples for the subsequent iteration.

Figure 10.4. Genetic policy search with $\sigma = 0.25$ applied to the simple regulator problem using 25 samples per iteration. The five elite samples in each generation are shown in red, with the best sample drawn with a larger dot.
10.4 Cross Entropy Method

The cross entropy method (algorithm 10.4) involves updating a search distribution over the parameterized space of policies at each iteration. We parameterize this search distribution \(p(\theta | \psi)\) with \(\psi\). This distribution can belong to any family, but a Gaussian distribution is a common choice, where \(\psi\) represents the mean and covariance of the distribution. The objective is to find a \(\psi^*\) that maximizes the expectation of \(U(\theta)\) when \(\theta\) is drawn from the search distribution:

\[
\psi^* = \arg \max_{\psi} \mathbb{E}_{\theta \sim p(\cdot | \psi)} [U(\theta)] = \arg \max_{\psi} \int_{\theta} U(\theta) p(\theta | \psi) d\theta
\] (10.5)

Directly maximizing equation (10.5) is typically computationally infeasible. The approach taken in the cross entropy method is to start with an initial \(\psi\), typically chosen so that the distribution is spread over the relevant parameter space. At each iteration, we draw \(m\) samples from the associated distribution and then update \(\psi\) to fit the elite samples. For the fit, we typically use the maximum likelihood estimate (section 4.1). We stop after some fixed number of iterations or until the search distribution becomes highly focused on an optimum. Figure 10.5 demonstrates the algorithm on a simple problem.

Algorithm 10.4. Cross entropy policy search, which iteratively improves a search distribution initially set to \(p\). This algorithm takes as input a parameterized policy \(\pi(\theta, s)\) and a policy evaluation function \(U\). In each iteration, \(m\) samples are drawn and the top \(m_{\text{elite}}\) are used to refit the distribution. The algorithm terminates after \(k_{\text{max}}\) iterations. The distribution \(p\) can be defined using the Distributions.jl package. For example, we might define \(\mu = [0.0, 1.0]\) \(\Sigma = [1.0, 0.0; 0.0, 1.0]\) \(p = \text{MvNormal} (\mu, \Sigma)\).
10.5 Evolution Strategies

Evolution strategies\(^9\) update a search distribution parameterized by a vector \(\boldsymbol{\psi}\) at each iteration. However, instead of fitting the distribution to a set of elite samples, it updates the distribution by taking a step in the direction of the gradient.\(^10\) The gradient of the objective in equation (10.5) can be computed as follows:\(^11\)

\[
\nabla_{\boldsymbol{\psi}} \mathbb{E}_{\theta \sim p(\cdot | \boldsymbol{\psi})} [U(\theta)] = \mathbb{E}_{\theta \sim p(\cdot | \boldsymbol{\psi})} [U(\theta) \nabla_{\theta} \log p(\theta | \boldsymbol{\psi})] = \nabla_{\boldsymbol{\psi}} \int U(\theta) \log p(\theta | \boldsymbol{\psi}) \, d\theta. \tag{10.10}
\]

The introduction of the logarithm above comes from what is called log derivative trick, which observes that \(\nabla_{\boldsymbol{\psi}} \log p(\theta | \boldsymbol{\psi}) = \nabla_{\boldsymbol{\psi}} p(\theta | \boldsymbol{\psi}) / p(\theta | \boldsymbol{\psi})\). This computation requires knowing \(\nabla_{\boldsymbol{\psi}} \log p(\theta | \boldsymbol{\psi})\), but we can often compute this analytically as discussed in example 10.2.

The search gradient can be estimated from \(m\) samples: \(\theta^{(1)}, \ldots, \theta^{(m)} \sim p(\cdot | \boldsymbol{\psi})\):

\[
\nabla_{\boldsymbol{\psi}} \mathbb{E}_{\theta \sim p(\cdot | \boldsymbol{\psi})} [U(\theta)] \approx \frac{1}{m} \sum_{i=1}^{m} U(\theta^{(i)}) \nabla_{\boldsymbol{\psi}} \log p(\theta^{(i)} | \boldsymbol{\psi}) \tag{10.11}
\]

This estimate depends on the evaluated expected utility, which itself can vary widely. We can make our gradient estimate more resilient with rank shaping, which replaces the utility values with weights based on the relative performance.
of each sample to the other samples in its iteration. The $m$ samples are sorted in descending order of expected utility. Weight $w^{(i)}$ is assigned to sample $i$ according to some weighting scheme with $w^{(1)} \geq \cdots \geq w^{(m)}$. The search gradient becomes:

$$\nabla_\psi \mathbb{E}_{\theta \sim p(\cdot|\psi)} [U(\theta)] \approx \sum_{i=1}^{m} w^{(i)} \nabla_\psi \log p(\theta^{(i)} | \psi)$$  \hspace{1cm} (10.12)

A common weighting scheme is:

$$w^{(i)} = \frac{\max(0, \log \left( \frac{m}{2} + 1 - \log(i) \right))}{\sum_{j=1}^{m} \max(0, \log \left( \frac{m}{2} + 1 - \log(j) \right))} - \frac{1}{m}$$  \hspace{1cm} (10.13)

These weights, shown in figure 10.6, favor better samples and give most samples a small negative weight. Rank-shaping reduces the influence of outliers.

![Figure 10.6. Several weightings constructed using equation (10.13).](image)

Algorithm 10.5 provides an implementation of the evolution strategies method. Figure 10.7 shows an example search progression.

10.6 Isotropic Evolutionary Strategies

The previous section introduced evolutionary strategies that can work with general search distributions. This section will make the assumption that the search distribution is a spherical or isotropic Gaussian, where the covariance matrix of the form $\sigma^2 I$.

Under this assumption, the expected utility of the distribution introduced in equation (10.5) simplifies to:

$$A^\top A = \Sigma$$


In general, if $A^\top A = \Sigma$, then $\theta = \mu + A^\top \epsilon$ transforms $\epsilon \sim \mathcal{N}(0, I)$ into a sample $\theta \sim \mathcal{N}(\mu, \Sigma)$.

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Algorithm 10.5. An evolution strategies method for updating a search distribution $D(\psi)$ over policy parameterizations for policy $\pi(\theta, s)$. This implementation also takes an initial search distribution parameterization $\psi$, the log search likelihood gradient $\nabla \log p(\psi, \theta)$, a policy evaluation function $U$, and an iteration count $k_{\text{max}}$. In each iteration, $m$ parameterization samples are drawn and are used to estimate the search gradient. This gradient is then applied with a step factor $\alpha$. We can use Distributions.jl to define $D(\psi)$. For example, if we want to define $D$ to construct a Gaussian with a given mean $\psi$ and fixed covariance $\Sigma$, we can use $D(\psi) = \text{MvNormal}(\psi, \Sigma)$.

Figure 10.7. Evolution strategies, algorithm 10.5, applied to the simple regulator problem using a multivariate Gaussian search distribution. Samples are shown in white, along with their search gradient contributions $w \nabla \log p$. 
The multivariate normal distribution \( \mathcal{N}(\mu, \Sigma) \) with mean \( \mu \) and covariance \( \Sigma \) is a common distribution family. The likelihood in \( d \) dimensions has the form

\[
p(x \mid \mu, \Sigma) = (2\pi)^{-d/2} |\Sigma|^{-1/2} \exp \left( -\frac{1}{2} (x - \mu)^\top \Sigma^{-1} (x - \mu) \right)
\]

where \( |\Sigma| \) is the determinant of \( \Sigma \). The log likelihood is

\[
\log p(x \mid \mu, \Sigma) = -\frac{d}{2} \log(2\pi) - \frac{1}{2} \log |\Sigma| - \frac{1}{2} (x - \mu)^\top \Sigma^{-1} (x - \mu)
\]

The parameters can be updated using their log likelihood gradients:

\[
\nabla_\mu \log p(x \mid \mu, \Sigma) = \Sigma^{-1} (x - \mu)
\]

\[
\nabla_\Sigma \log p(x \mid \mu, \Sigma) = \frac{1}{2} \Sigma^{-1} (x - \mu)(x - \mu)^\top \Sigma^{-1} - \frac{1}{2} \Sigma^{-1}
\]

The term \( \nabla_\Sigma \) contains the partial derivative of each entry of \( \Sigma \) with respect to the log likelihood.

Directly updating \( \Sigma \) may not result in a positive definite matrix, as is required for covariance matrices. One solution is to represent \( \Sigma \) as a product \( A^\top A \), which guarantees that \( \Sigma \) remains positive semidefinite, and then update \( A \) instead. Replacing \( \Sigma \) by \( A^\top A \) and taking the gradient with respect to \( A \) yields:

\[
\nabla_{(A)} \log p(x \mid \mu, A) = A \left[ \nabla_\Sigma \log p(x \mid \mu, \Sigma) + \nabla_\Sigma \log p(x \mid \mu, \Sigma)^\top \right]
\]
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\[
\mathbb{E}_{\theta \sim \mathcal{N}(\psi, \sigma^2 I)} [U(\theta)] = \mathbb{E}_{\epsilon \sim \mathcal{N}(0, I)} [U(\psi + \sigma \epsilon)]
\] (10.14)

The search gradient reduces to:

\[
\nabla_{\psi} \mathbb{E}_{\theta \sim \mathcal{N}(\psi, \sigma^2 I)} [U(\theta)] = \mathbb{E}_{\theta \sim \mathcal{N}(\psi, \sigma^2 I)} \left[ U(\theta) \nabla_{\psi} \log p(\theta | \psi, \sigma^2 I) \right]
\] (10.15)

\[
= \mathbb{E}_{\epsilon \sim \mathcal{N}(0, I)} \left[ U(\psi + \sigma \epsilon) \frac{1}{\sigma^2} \right]
\] (10.16)

\[
= \frac{1}{\sigma} \mathbb{E}_{\epsilon \sim \mathcal{N}(0, I)} \left[ U(\psi + \sigma \epsilon) \epsilon \right]
\] (10.17)

Algorithm 10.6 provides an implementation of this strategy. This implementation incorporates mirrored sampling.\(^{15}\) We sample \(m/2\) values from the search distribution and then generate the other \(m/2\) samples by mirroring them about the mean. Mirrored samples reduce the variance of the gradient estimate.\(^{16}\) The benefit of this technique is shown in figure 10.8.

```plaintext
struct IsotropicEvolutionStrategies
    # initial mean
    psi # initial standard devidation
    sigma # number of samples
    m # step factor
    k_max # number of iterations
end

function optimize_dist(M::IsotropicEvolutionStrategies, pi, U)
    psi, sigma, m, k_max = M.psi, M.sigma, M.m, M.k_max
    n = length(psi)
    ws = evolution_strategy_weights(2*div(m,2))
    for k in 1:k_max
        es = [randn(n) for i in 1:div(m,2)]
        append!(es, -es) # weight mirroring
        us = [U(pi, psi + sigma*epsilon) for epsilon in es]
        sp = sortperm(us, rev=true)
        V = sum(w.*es[i] for (w,i) in zip(ws,sp)) / sigma
        psi += alpha*V
    end
    return MvNormal(psi, sigma)
end
```


\(^{16}\) This technique was implemented by T. Salimans, J. Ho, X. Chen, S. Sidor, and I. Sutskever, “Evolution Strategies as a Scalable Alternative to Reinforcement Learning,” 2017. arXiv: 1703.03864v2. They included other techniques as well, including weight decay.
10.7 Summary

- Monte Carlo policy evaluation involves computing expected utility associated with a policy using a large number of rollouts from states sampled from an initial state distribution.

- Local search methods such as the Hooke-Jeeves method improve a policy based on small, local changes.

- Genetic algorithms maintain a population of points in the parameter space, recombining them in different ways in attempt to drive the population towards a global optimum.

- The cross entropy method iteratively improves a search distribution over policy parameters by re-fitting the distribution to elite samples at each iteration.

- Evolutionary strategies attempt to improve the search distribution using gradient information from samples from that distribution.

- Isotropic evolutionary strategies make the assumption that the search distribution is an isotropic Gaussian.

10.8 Exercises

Exercise 10.1. In Monte Carlo policy evaluation, how is the variance of the utility estimate affected by the number of samples?
Solution: The variance of Monte Carlo policy evaluation is the variance of the mean of $m$ samples. These samples are assumed independent, and so the variance of the mean is the variance of a single rollout evaluation divided by the sample size:

$$\frac{1}{m} \sum_{i=1}^{m} (u^{(i)} - \mu)^2$$

where $u^{(i)}$ is the $i$th rollout utility and $\mu$ is their sample mean. The sample variance therefore decreases with $1/m$.

Exercise 10.2. What effect does varying the number of samples $m$ and the number of elite samples $m_{\text{elite}}$ have on cross entropy policy search?

Solution: The computational cost per iteration scales linearly with the number of samples. More samples will better cover the search space, resulting in a better chance of identifying better elite samples to improve the policy. The number of elite samples also has an effect. Making all samples elite provides no feedback to the improvement process. Having too few elite samples can lead to premature convergence to a suboptimal solution.

Exercise 10.3. Consider using evolution strategies with a univariate Gaussian distribution, $\theta \sim \mathcal{N}(\mu, \nu)$. What is the search gradient with respect to the variance $\nu$? What issue arises as the variance becomes small?

Solution: The search gradient is the gradient of the log-likelihood:

$$\frac{\partial}{\partial \nu} \log p(x \mid \mu, \nu) = \frac{\partial}{\partial \nu} \log \frac{1}{\sqrt{2\pi\nu}} \exp \left( -\frac{(x - \mu)^2}{2\nu} \right)$$

$$= \frac{\partial}{\partial \nu} \left( -\frac{1}{2} \log(2\pi) - \frac{1}{2} \log(\nu) - \frac{(x - \mu)^2}{2\nu} \right)$$

$$= -\frac{1}{2\nu} + \frac{(x - \mu)^2}{2\nu^2}$$

We find that the gradient goes to infinity as the variance approaches zero. This is a problem because the variance should be small when the search distribution converges. Very large gradients can cause simple ascent methods to overshoot optima.
Exercise 10.4. Equation (10.14) defines the objective in terms of a search distribution \( \theta \sim \mathcal{N}(\psi, \Sigma) \). What advantage does this objective have over directly optimizing \( \theta \) using the expected utility objective in equation (10.1)?

Solution: The added Gaussian noise around the policy parameters can smooth discontinuities in the original objective, which can lead more reliable optimization.

Exercise 10.5. Which of the methods in this chapter are best suited to reflect the fact that multiple types of policies could perform well in a given problem?

Solution: The Hooke-Jeeves method improves a single policy parameterization, so it cannot retain multiple policies. Both the cross entropy method and evolution strategies use search distributions. In order to successfully represent multiple types of policies, a multi-modal distribution would have to be used. One common multi-modal distribution is a mixture of Gaussians. A mixture of Gaussians cannot be fit analytically, but can be reliably fit using expectation maximization as demonstrated in example 4.4. Genetic algorithms can retain multiple policies if the population size is sufficiently large.

Exercise 10.6. Suppose we have a parameterized policy \( \pi_\theta \) that we would like to optimize using the Hooke-Jeeves method. If we initialize our parameter \( \theta = 0 \) and the utility function is \( U(\theta) = -3\theta^2 + 4\theta + 1 \), what is the largest step size \( \alpha \) that would still guarantee policy improvement in the first iteration of the Hooke-Jeeves method?

Solution: The Hooke-Jeeves method evaluates the objective function at the center point \( \pm \alpha \) along each coordinate direction. In order to guarantee improvement in the first iteration of Hooke-Jeeves search, any of the objective function values at the new points must improve the objective function value. For our policy optimization problem, this means we are searching for the largest step size \( \alpha \) such that either \( U(\theta + \alpha) \) or \( U(\theta - \alpha) \) is greater than \( U(\theta) \).

Since the underlying utility function is parabolic and concave, the largest step size that would still lead to improvement is slightly less than the width of the parabola at the current point. Thus, we compute the point on the parabola opposite the current point, \( \theta' \) at which \( U(\theta') = U(\theta) \):

\[
U(\theta) = -3\theta^2 + 4\theta + 1 = -3(0)^2 + 4(0) + 1 = 1
\]

\[
U(\theta) = U(\theta')
\]

\[
1 = -3\theta'^2 + 4\theta' + 1
\]

\[
0 = -3\theta'^2 + 4\theta' + 0
\]

\[
\theta' = \frac{-4 \pm \sqrt{4^2 - 4(-3)(1)}}{2(-3)} = \frac{-4 \pm 4}{-6} = \frac{2 \pm 2}{3} = \left\{0, \frac{4}{3}\right\}
\]

The point on the parabola opposite the current point is thus \( \theta' = \frac{4}{3} \). The distance between \( \theta \) and \( \theta' \) is \( \frac{4}{3} - 0 = \frac{4}{3} \). Thus, the maximal step size we can take and still guarantee improvement in the first iteration is \( \alpha = \frac{4}{3} - \epsilon \) (where \( \epsilon \) is a small number).
Exercise 10.7. Suppose we have a policy parameterized by a single parameter $\theta$. We take an evolution strategies approach with a search distribution that follows a Bernoulli distribution $p(\theta \mid \psi) = \psi^\theta (1 - \psi)^{1-\theta}$. Compute the log-likelihood gradient $\nabla_{\psi} \log p(\theta \mid \psi)$.

Solution: The log-likelihood gradient can be computed as follows:

$$p(\theta \mid \psi) = \psi^\theta (1 - \psi)^{1-\theta}$$
$$\log p(\theta \mid \psi) = \theta \log \psi + (1 - \theta) \log(1 - \psi)$$

$$\nabla_{\psi} \log p(\theta \mid \psi) = \frac{\theta}{\psi} - \frac{1 - \theta}{1 - \psi}$$

Exercise 10.8. Compute the sample weights for search gradient estimation with rank shaping given $m = 3$ samples.

Solution: We first compute the numerator of the first term from equation (10.13), for all $i$

$$i = 1 \quad \max \left(0, \log \left(\frac{3}{2} + 1\right) - \log 1\right) = \log \frac{5}{2}$$
$$i = 2 \quad \max \left(0, \log \left(\frac{3}{2} + 1\right) - \log 2\right) = \log \frac{5}{4}$$
$$i = 3 \quad \max \left(0, \log \left(\frac{3}{2} + 1\right) - \log 3\right) = 0$$

Now, we compute the weights:

$$w^{(1)} = \frac{\log \frac{5}{2}}{\log \frac{5}{2} + \log \frac{5}{4} + 0} - \frac{1}{3} = 0.47$$
$$w^{(2)} = \frac{\log \frac{5}{4}}{\log \frac{5}{2} + \log \frac{5}{4} + 0} - \frac{1}{3} = -0.14$$
$$w^{(3)} = \frac{0}{\log \frac{5}{2} + \log \frac{5}{4} + 0} - \frac{1}{3} = -0.33$$
11 Policy Gradient Estimation

The previous chapter discussed several ways to go about directly optimizing the parameters of a policy to maximize expected utility. In many applications, it is often useful to use the gradient of the utility with respect to the policy parameters to guide the optimization process. This chapter discusses several approaches to estimating this gradient from trajectory rollouts.¹ A major challenge with this approach is the variance of the estimate due to the stochastic nature of the trajectories arising from both the environment and our exploration of it. The next chapter will discuss how to use these algorithms for gradient estimation for the purpose of policy optimization.

11.1 Finite Difference

Finite difference methods estimate the gradient of a function from small changes in its evaluation. Recall that the derivative of a univariate function \( f \) is

\[
\frac{df}{dx}(x) = \lim_{\delta \to 0} \frac{f(x + \delta) - f(x)}{\delta}
\]  

(11.1)

The derivative at \( x \) can be approximated by a sufficiently small step \( \delta > 0 \):

\[
\frac{df}{dx}(x) \approx \frac{f(x + \delta) - f(x)}{\delta}
\]  

(11.2)

This approximation is illustrated in figure 11.1.

The gradient of a multivariate function \( f \) with an input of length \( n \) is:

\[
\nabla f(x) = \left[ \frac{\partial f}{\partial x_1}(x), \ldots, \frac{\partial f}{\partial x_n}(x) \right]
\]  

(11.3)

Finite differences can be applied to each dimension to estimate the gradient.

¹ An additional resource on this topic is provided by M. C. Fu, “Gradient Estimation,” in *Simulation*, S.G. Henderson and B.L. Nelson, eds., Elsevier, 2006, pp. 575–616.
In the context of policy optimization, we want to estimate the gradient of the utility expected from following a policy parameterized by $\theta$:

$$\nabla U(\theta) = \left[ \frac{\partial U}{\partial \theta_1}(\theta), \ldots, \frac{\partial U}{\partial \theta_n}(\theta) \right]$$

(11.4)

$$\approx \left[ \frac{U(\theta + \delta e^{(1)}) - U(\theta)}{\delta}, \ldots, \frac{U(\theta + \delta e^{(n)}) - U(\theta)}{\delta} \right]$$

(11.5)

where $e^{(i)}$ is the $i$th standard basis vector consisting of zeros except for the $i$th component that is set to 1.

As discussed in section 10.1, we need to simulate policy rollouts to estimate $U(\theta)$. We can use algorithm 11.1 to generate trajectories. From these trajectories, we can compute their return and estimate the utility associated with the policy. Algorithm 11.2 implements the gradient estimate in equation (11.5) by simulating $m$ rollouts for each component and averaging the returns.

```plaintext
function simulate(𝒫::MDP, s, π, d)
    τ = []
    for i = 1:d
        a = π(s)
        s′, r = 𝒫.TR(s, a)
        push!(τ, (s, a, r))
        s = s′
    end
    return τ
end
```

Algorithm 11.1. A method for generating a trajectory associated with problem $\mathcal{P}$ starting in state $s$ and executing policy $\pi$ to depth $d$. It creates a vector $\tau$ containing state-action-reward tuples.

A major challenge in arriving at accurate estimates of the policy gradient is the fact that the variance of the trajectory rewards can be quite high. One approach to reduce the resulting variance in the gradient estimate is to have each rollout share the same random generator seeds.\(^2\) This approach can be helpful, for example, in cases where one rollout happens to hit a low-probability transition early on. Other rollouts will have the same tendency due to the shared random generator, and their rewards will tend to be biased in the same way.

Policy representations have a significant effect on the policy gradient. Example 11.1 demonstrates the sensitivity of the policy gradient to the policy parameterization. Finite differences for policy optimization can perform poorly when the parameters differ in scale.

\(^2\) This random seed sharing is used in the PEGASUS algorithm. A.Y. Ng and M. Jordan, “A Policy Search Method for Large MDPs and POMDPs,” in Conference on Uncertainty in Artificial Intelligence (UAI), 2000.
Consider a single-state, single-step MDP with a one-dimensional continuous action space and a reward function $R(s, a) = a$. In this case, larger actions produce higher rewards.

Suppose we have a stochastic policy $\pi_\theta$ that samples its action according to a uniform distribution between $\theta_1$ and $\theta_2$ for $\theta_2 > \theta_1$. The expected value is:

$$U(\theta) = \mathbb{E}[a] = \int_{\theta_1}^{\theta_2} a \frac{1}{\theta_2 - \theta_1} \, da = \frac{\theta_1 + \theta_2}{2}$$

The policy gradient is:

$$\nabla U(\theta) = [1/2, 1/2]$$

The policy could be reparameterized to draw actions from a uniform distribution between $\theta_1$ and $100\theta_2$, for $100\theta_2 > \theta_1$. Now the expected reward is $(\theta_1 + 100\theta_2)/2$ and the policy gradient is $[1/2, 50]$.

Both policies are identical from the agent’s perspective, but the two have wildly different gradients. Finding a suitable perturbation scalar for the second policy is much more difficult, as the parameters vary widely in scale.
11.2 Regression Gradient

Instead of estimating the gradient at $\theta$ by taking a fixed step along each coordinate axis as done in the previous section, we can use linear regression\(^3\) to estimate the gradient from the results of random perturbations from $\theta$. These perturbations are stored in a matrix:

$$\Delta \Theta = \begin{bmatrix} (\Delta \theta^{(1)})^\top \\ \vdots \\ (\Delta \theta^{(m)})^\top \end{bmatrix}$$

(11.6)

More policy parameter perturbations will tend to produce better gradient estimates.\(^4\)

For each of these perturbations, we perform a rollout and estimate the change in utility:\(^5\)

$$\Delta U = \left[ U(\theta + \Delta \theta^{(1)}) - U(\theta), \ldots, U(\theta + \Delta \theta^{(m)}) - U(\theta) \right]$$

(11.7)

The policy gradient estimate using linear regression is then:\(^6\)

$$\nabla U(\theta) \approx \Delta \Theta^+ \Delta U$$

(11.8)

Algorithm 11.3 provides an implementation of this approach where the perturbations are drawn uniformly from a hypersphere with radius $\delta$. Example 11.2 demonstrates this approach on a simple function.

### Algorithm 11.3

A method for estimating a policy gradient using finite differences for an MDP $\mathcal{P}$, a stochastic parameterized policy $\pi(\theta, s)$, and policy parameterization vector $\theta$. Policy variation vectors are generated by normalizing normally-distributed samples and scaling by a perturbation scalar $\delta$. A total of $m$ parameter perturbations are generated, and each is evaluated in a rollout from an initial state drawn from $b$ to depth $d$ and compared to the original policy parameterization.

```
struct RegressionGradient
    P # problem
    b # initial state distribution
    d # depth
    m # number of samples
    δ # step size
end

function gradient(M::RegressionGradient, π, θ)
    P, b, d, m, δ, γ = M.P, M.b, M.d, M.m, M.δ, M.γ
    ΔΘ = [δ .* normalize(randn(length(θ)), 2) for i = 1:m]
    R(τ) = sum(r * γ^k for (k, (s, a, r)) in enumerate(τ))
    U(θ) = R(simulate(P, rand(b), s→π(θ,s), d))
    ΔU = [U(θ + Δθ) - U(θ) for Δθ in ΔΘ]
    return pinv(reduce(hcat, ΔΘ))' * ΔU
end
```
We would like to apply the regression gradient to estimate the gradient of a simple one-dimensional function \( f(x) = x^2 \) evaluated at \( x_0 = 2 \) from \( m = 20 \) samples. To imitate the stochasticity inherent in policy evaluation, we add noise to the function evaluations. We generate a set of disturbances \( \Delta X \) sampled from \( \mathcal{N}(0, \delta^2) \) and evaluate \( f(x_0 + \Delta x) - f(x_0) \) for each disturbance \( \Delta x \) in \( \Delta X \). We can then estimate the one-dimensional gradient (or derivative) \( \Delta X^+ \Delta F \) with this code:

\[
\begin{align*}
f(x) &= x^2 + 1e-2 \times \text{randn}() \\
m &= 20 \\
\delta &= 1e-2 \\
\Delta X &= [\delta \times \text{randn()} \text{ for } i = 1:m] \\
x0 &= 2.0 \\
\Delta F &= [f(x0 + \Delta x) - f(x0) \text{ for } \Delta x \text{ in } \Delta X] \\
\text{pinv}(\Delta X) \times \Delta F
\end{align*}
\]

The samples and linear regression are shown below. The slope of the regression line is close to the exact solution of 4.

Example 11.2. Demonstration of the regression gradient method on a one-dimensional function.
11.3 Likelihood Ratio

The likelihood ratio approach to gradient estimation uses an analytical form of \( \nabla \pi_\theta \) to improve our estimate of \( \nabla U(\theta) \). Recall from equation (10.2) that

\[
U(\theta) = \int p_\theta(\tau) R(\tau) \, d\tau \tag{11.9}
\]

Hence,

\[
\nabla U(\theta) = \nabla_\theta \int p_\theta(\tau) R(\tau) \, d\tau \tag{11.10}
\]

\[
= \int \nabla_\theta p_\theta(\tau) R(\tau) \, d\tau \tag{11.11}
\]

\[
= \int p_\theta(\tau) \frac{\nabla_\theta p_\theta(\tau)}{p_\theta(\tau)} R(\tau) \, d\tau \tag{11.12}
\]

\[
= \mathbb{E}_\tau \left[ \frac{\nabla_\theta p_\theta(\tau)}{p_\theta(\tau)} R(\tau) \right] \tag{11.13}
\]

Applying the log derivative trick, we have

\[
\nabla U(\theta) = \mathbb{E}_\tau [\nabla_\theta \log p_\theta(\tau) R(\tau)] \tag{11.14}
\]

We can estimate the expectation above through rolling out trajectories. For each trajectory sample \( \tau^{(i)} \), we need to compute the product \( \nabla_\theta \log p_\theta(\tau^{(i)}) R(\tau^{(i)}) \). Recall that \( R(\tau^{(i)}) \) is the return associated with trajectory \( \tau^{(i)} \). If we have a stochastic policy, the gradient \( \nabla_\theta \log p_\theta(\tau^{(i)}) \) is

\[
\nabla_\theta \log p_\theta(\tau) = \sum_{k=1}^{d} \nabla_\theta \log \pi_\theta(a^{(k)} \mid s^{(k)}) \tag{11.15}
\]

because \( p_\theta(\tau) \) has the form:

\[
p_\theta(\tau) = p(s^{(1)}) \prod_{k=1}^{d} T(s^{(k+1)} \mid s^{(k)}, a^{(k)}) \pi_\theta(a^{(k)} \mid s^{(k)}) \tag{11.16}
\]

where \( s^{(k)} \) and \( a^{(k)} \) are the \( k \)th state and action, respectively, in trajectory \( \tau \). Algorithm 11.4 provides an implementation where \( m \) trajectories are sampled to arrive at a gradient estimate. Example 11.3 illustrates the process.
If we have a deterministic policy, the gradient requires computing:

\[
\nabla_\theta \log p_\theta(\tau) = \nabla_\theta \log \left[ p(s^{(1)}) \prod_{k=1}^{d} T(s^{(k+1)} \mid s^{(k)}, \pi_\theta(s^{(k)})) \right]
\]

(11.17)

\[
= \sum_{k=1}^{d} \nabla_\theta \pi_\theta(s^{(k)}) \frac{\partial}{\partial a^{(k)}} \log T(s^{(k+1)} \mid s^{(k)}, a^{(k)})
\]

(11.18)

The equation above requires knowing transition likelihood, which is in contrast with equation (11.15) for stochastic policies.

```
struct LikelihoodRatioGradient
  𝒫 # problem
  b # initial state distribution
  d # depth
  m # number of samples
  ∇logπ # gradient of log likelihood
end

function gradient(M::LikelihoodRatioGradient, π, θ)
  𝒫, b, d, m, ∇logπ, γ = M.𝒫, M.b, M.d, M.m, M.∇logπ, M.γ
  πθ(s) = π(θ, s)
  R(τ) = sum(τ*γ^k-1) for (k, (s,a,r)) in enumerate(τ)
  ∇U(τ) = sum(∇logπ(θ, a, s) for (s,a) in τ)*R(τ)
  return mean(∇U(simulate(𝒫, rand(b), πθ, d)) for i in 1:m)
end
```

Algorithm 11.4. A method for estimating a policy gradient of a policy π(s) for an MDP 𝒫 with initial state distribution b using the likelihood ratio trick. The gradient with respect to the parameterization vector θ is estimated from m rollouts to depth d using the log policy gradients ∇logπ.

11.4 Reward-to-Go

The likelihood ratio policy gradient method is unbiased but has high variance. Example 11.4 reviews bias and variance. The variance generally increases significantly with rollout depth due to the correlation between actions, states, and rewards across time steps. The reward-to-go approach attempts to reduce the variance in the estimate.

To derive this approach, we begin by expanding equation (11.14):

\[
\nabla U(θ) = \mathbb{E}_\tau \left[ \left( \sum_{k=1}^{d} \nabla_\theta \log \pi_\theta(a^{(k)} \mid s^{(k)}) \right) \left( \sum_{k=1}^{d} r^{(k)} \gamma^{k-1} \right) \right]
\]

(11.19)

Let \( f^{(k)} \) replace \( \nabla_\theta \log \pi_\theta(a^{(k)} \mid s^{(k)}) \) for convenience. We then expand:
Consider the single-step, single-state problem from example 11.1. Suppose we have a stochastic policy \( \pi_\theta \) that samples its action according to a Gaussian distribution \( \mathcal{N}(\theta_1, \theta_2^2) \) where \( \theta_2^2 \) is the variance.

The log policy likelihood is:

\[
\log \pi_\theta (a | s) = \log \left( \frac{1}{\sqrt{2\pi\theta_2^2}} \exp \left( -\frac{(a - \theta_1)^2}{2\theta_2^2} \right) \right) = -\frac{(a - \theta_1)^2}{2\theta_2^2} - \frac{1}{2} \log (2\pi\theta_2^2)
\]

The gradient of the log policy likelihood is:

\[
\frac{\partial}{\partial \theta_1} \log \pi_\theta (a | s) = \frac{a - \theta_1}{\theta_2^2}
\]
\[
\frac{\partial}{\partial \theta_2} \log \pi_\theta (a | s) = \frac{(a - \theta_1)^2 - \theta_2^2}{\theta_2^3}
\]

Suppose we run three rollouts with \( \theta = [0, 1] \), taking actions \( \{0.5, -1, 0.7\} \) and receiving the same rewards \( (R(s, a) = a) \). The estimated policy gradient is:

\[
\nabla U(\theta) \approx \frac{1}{m} \sum_{i=1}^{m} \nabla \theta \log p_\theta (\tau^{(i)}) R(\tau^{(i)})
\]
\[
= \frac{1}{3} \left( \begin{bmatrix} 0.5 & -1.0 & 0.7 \end{bmatrix} \begin{bmatrix} 0.5 \end{bmatrix} + \begin{bmatrix} -0.75 \end{bmatrix} \begin{bmatrix} -1.0 \end{bmatrix} + \begin{bmatrix} 0.0 \end{bmatrix} \begin{bmatrix} 0.7 \end{bmatrix} \right)
\]
\[
= [0.58, -0.244]
\]
When estimating a quantity of interest from a collection of simulations, we generally want to use a scheme that has both low bias and low variance. In this chapter, we want to estimate $\nabla U(\theta)$. Generally, with more simulation samples, we can arrive at a better estimate. Some methods can lead to bias, where—even with infinitely many samples—it does not lead to an accurate estimate. Sometimes methods with non-zero bias may still be attractive if they have low variance, meaning they require fewer samples to converge.

Below are plots of the estimates from four notional methods for estimating $\nabla U(\theta)$. The true value is 17.5, as indicated by the red line. For all four methods, we ran 100 simulations 100 times. The variance decreases as the number of samples increases. The blue regions indicate the 5% to 95% and 25% to 75% empirical quantiles of the estimates.
\[ \nabla U(\theta) = \mathbb{E}_\tau \left[ \left( \sum_{k=1}^{d} f^{(k)} \right) \left( \sum_{k=1}^{d} r^{(k)} \gamma^{k-1} \right) \right] \]
\[ = \mathbb{E}_\tau \left[ \left( f^{(1)} + f^{(2)} + f^{(3)} + \ldots + f^{(d)} \right) \left( r^{(1)} + r^{(2)} \gamma + r^{(3)} \gamma^2 + \ldots + r^{(d)} \gamma^{d-1} \right) \right] \]

The first reward, \( r^{(1)} \), is only affected by the first action. Thus, its contribution to the policy gradient should not depend on subsequent timesteps. We can remove other such causality-violating terms:\(^{10}\)

\[ \nabla U(\theta) = \mathbb{E}_\tau \left[ \begin{array}{c} f^{(1)} r^{(1)} + f^{(1)} r^{(2)} \gamma + f^{(1)} r^{(3)} \gamma^2 + \ldots + f^{(1)} r^{(d)} \gamma^{d-1} \\ f^{(2)} r^{(1)} + f^{(2)} r^{(2)} \gamma + f^{(2)} r^{(3)} \gamma^2 + \ldots + f^{(2)} r^{(d)} \gamma^{d-1} \\ f^{(3)} r^{(1)} + f^{(3)} r^{(2)} \gamma + f^{(3)} r^{(3)} \gamma^2 + \ldots + f^{(3)} r^{(d)} \gamma^{d-1} \\ \vdots \\ f^{(d)} r^{(1)} + f^{(d)} r^{(2)} \gamma + f^{(d)} r^{(3)} \gamma^2 + \ldots + f^{(d)} r^{(d)} \gamma^{d-1} \end{array} \right] \]

Algorithm 11.5 provides an implementation.

Notice that the reward-to-go for a state-action pair \((s, a)\) under a policy parameterized by \(\theta\) is really an approximation of the state-action value from that state, \(Q_\theta(s, a)\). The action value function, if known, can be used to obtain the policy gradient:

\[ \nabla U(\theta) = \mathbb{E}_\tau \left[ \sum_{k=1}^{d} \nabla \log \pi_\theta(a^{(k)} | s^{(k)}) \left( \sum_{\ell=k}^{d} r^{(\ell)} \gamma^{\ell-1} \right) \right] \]

\[ = \mathbb{E}_\tau \left[ \sum_{k=1}^{d} \nabla \log \pi_\theta(a^{(k)} | s^{(k)}) \gamma^{k-1} \left( \sum_{\ell=k}^{d} r^{(\ell)} \gamma^{\ell-k} \right) \right] \]

\[ = \mathbb{E}_\tau \left[ \sum_{k=1}^{d} \nabla \log \pi_\theta(a^{(k)} | s^{(k)}) \gamma^{k-1} r^{(k)}_{\text{to-go}} \right] \]

\[ = \mathbb{E}_\tau \left[ \sum_{k=1}^{d} \nabla \log \pi_\theta(a^{(k)} | s^{(k)}) \gamma^{k-1} Q_\theta(s^{(k)}, a^{(k)}) \right] \]
11.5 Baseline Subtraction

We can further build upon the approach presented in the previous section by subtracting a baseline value from the reward-to-go\(^{11}\) to reduce the variance of the gradient estimate. This subtraction does not bias the gradient.

We now subtract a baseline \( r_{\text{base}}(s^{(k)}) \):

\[
\nabla U(\theta) = E_{\tau} \left[ \sum_{k=1}^{d} \nabla_\theta \log \pi_\theta(a^{(k)} | s^{(k)}) \gamma^{k-1} \left( r^{(k)}_{\text{to-go}} - r_{\text{base}}(s^{(k)}) \right) \right] \tag{11.28}
\]

To show that baseline subtraction does not bias the gradient, we first expand:

\[
\nabla U(\theta) = E_{\tau} \left[ \sum_{k=1}^{d} \nabla_\theta \log \pi_\theta(a^{(k)} | s^{(k)}) \gamma^{k-1} r^{(k)}_{\text{to-go}} - \sum_{k=1}^{d} \nabla_\theta \log \pi_\theta(a^{(k)} | s^{(k)}) \gamma^{k-1} r_{\text{base}}(s^{(k)}) \right] \tag{11.29}
\]

The linearity of expectation states that \( E[a + b] = E[a] + E[b] \), so it is sufficient to prove that equation (11.29) is equivalent to equation (11.26), if for each step \( k \), the expected associated baseline term is 0:

\[
E_{\tau} \left[ \nabla_\theta \log \pi_\theta(a^{(k)} | s^{(k)}) \gamma^{k-1} r_{\text{base}}(s^{(k)}) \right] = 0 \tag{11.30}
\]

We begin by converting the expectation into nested expectations, as illustrated in figure 11.2:

\[
E_{\tau} \left[ \nabla_\theta \log \pi_\theta(a^{(k)} | s^{(k)}) \gamma^{k-1} r_{\text{base}}(s^{(k)}) \right] = E_{\tau_{k+1:d}} \left[ E_{\tau_{k+1}} \left[ \nabla_\theta \log \pi_\theta(a^{(k)} | s^{(k)}) \gamma^{k-1} r_{\text{base}}(s^{(k)}) \right] \right] \tag{11.31}
\]

\(^{11}\) We could also subtract a baseline from a state-action value.
Therefore, subtracting a term down the equations in our derivation, we define

\[ E_\tau[f(\tau)] = E_{\tau_1:k}[E_{\tau_{k+1:d}}[f(\tau)]] \]

We continue with our derivation, using the same log derivative trick that was used in section 11.3:

\[
E_{\tau_1:k}\left[ E_{\tau_{k+1:d}}[\nabla_\theta \log \pi_\theta(a^{(k)} | s^{(k)}) \gamma^{k-1}r_{\text{base}}(s^{(k)})] \right] \\
= E_{\tau_1:k}\left[ \gamma^{k-1}r_{\text{base}}(s^{(k)}) E_{\tau_{k+1:d}}[\nabla_\theta \log \pi_\theta(a^{(k)} | s^{(k)})] \right] \\
= E_{\tau_1:k}\left[ \gamma^{k-1}r_{\text{base}}(s^{(k)}) E_{a^{(k)}}[\nabla_\theta \log \pi_\theta(a^{(k)} | s^{(k)})] \right] \\
= E_{\tau_1:k}\left[ \gamma^{k-1}r_{\text{base}}(s^{(k)}) \int \nabla_\theta \log \pi_\theta(a^{(k)} | s^{(k)}) \pi_\theta(a^{(k)} | s^{(k)}) da^{(k)} \right] \\
= E_{\tau_1:k}\left[ \gamma^{k-1}r_{\text{base}}(s^{(k)}) \int \nabla_\theta \pi_\theta(a^{(k)} | s^{(k)}) \pi_\theta(a^{(k)} | s^{(k)}) da^{(k)} \right] \\
= E_{\tau_1:k}\left[ \gamma^{k-1}r_{\text{base}}(s^{(k)}) \nabla_\theta \int \pi_\theta(a^{(k)} | s^{(k)}) da^{(k)} \right] \\
= E_{\tau_1:k}\left[ \gamma^{k-1}r_{\text{base}}(s^{(k)}) \nabla_\theta 1 \right] \\
= E_{\tau_1:k}\left[ \gamma^{k-1}r_{\text{base}}(s^{(k)}) 0 \right]
\]

Therefore, subtracting a term \( r_{\text{base}}(s^{(k)}) \) does not bias the estimate. This derivation assumed continuous state and action spaces. The same result applies to discrete spaces.

We can choose a different \( r_{\text{base}}(s) \) for every component of the gradient, and will select them to minimize the variance. For simplicity, we will drop the dependence on \( s \) and treat each baseline component as constant.\(^{12}\) For compactness in writing down the equations in our derivation, we define

\[
\ell_i(a, s, k) = \gamma^{k-1} \frac{\partial}{\partial \theta_i} \log \pi_\theta(a | s)
\]

\(^{12}\) Some methods approximate a state-dependent baseline using \( r_{\text{base}}(s^{(k)}) = \Phi(s^{(k)})^T \mathbf{w} \). Selecting appropriate baseline functions tends to be difficult. J. Peters and S. Schaal, “Reinforcement Learning of Motor Skills with Policy Gradients,” *Neural Networks*, vol. 21, no. 4, pp. 682–697, 2008.
The variance of the \(i\)th component of our gradient estimate in equation (11.28) is

\[
\mathbb{E}_{a,s,r_{\text{to-go}},k} \left[ (\ell_i(a,s,k)(r_{\text{to-go}} - r_{\text{base},i}))^2 \right] - \mathbb{E}_{a,s,r_{\text{to-go}},k} \left[ \ell_i(a,s,k)(r_{\text{to-go}} - r_{\text{base},i}) \right]^2
\]

where the expectation is over the \((a,s,r_{\text{to-go}})\) tuples in our trajectory samples, and \(k\) is each tuple’s depth.

We have just shown that the second term is zero. Hence, we can focus on choosing \(r_{\text{base},i}\) to minimize the first term by taking the derivative with respect to the baseline and setting it to zero:

\[
\frac{\partial}{\partial r_{\text{base},i}} \mathbb{E}_{a,s,r_{\text{to-go}},k} \left[ (\ell_i(a,s,k)(r_{\text{to-go}} - r_{\text{base},i}))^2 \right] = \frac{\partial}{\partial r_{\text{base},i}} \left( \mathbb{E}_{a,s,r_{\text{to-go}},k} \left[ \ell_i(a,s,k)^2 r_{\text{to-go}} \right] - 2 \mathbb{E}_{a,s,r_{\text{to-go}},k} \left[ \ell_i(a,s,k)^2 r_{\text{to-go}} r_{\text{base},i} \right] + r_{\text{base},i} \mathbb{E}_{a,s,k} \left[ \ell_i(a,s,k)^2 \right] \right)
\]

\[
= -2 \mathbb{E}_{a,s,r_{\text{to-go}},k} \left[ \ell_i(a,s,k)^2 r_{\text{to-go}} \right] + 2 r_{\text{base},i} \mathbb{E}_{a,s} \left[ \ell_i(a,s,k)^2 \right] = 0
\]

Solving for \(r_{\text{base},i}\) yields the baseline component that minimizes the variance:

\[
r_{\text{base},i} = \frac{\mathbb{E}_{a,s,r_{\text{to-go}},k} \left[ \ell_i(a,s,k)^2 r_{\text{to-go}} \right]}{\mathbb{E}_{a,s,k} \left[ \ell_i(a,s,k)^2 \right]}
\]

It is common to use likelihood ratio policy gradient estimation with this baseline subtraction (algorithm 11.6).\(^\text{13}\) Figure 11.3 compares the methods we discussed.

Qualitatively, when considering the gradient contribution of state-action pairs, what we really care about is the relative value of one action over another. If all actions in a particular state produce the same high value, there is no real signal in the gradient, and baseline subtraction can zero that out. We want to identify the actions that produce a higher value than others, irrespective of what the mean value is across actions.

An alternative to the action value is the advantage, \(A(s,a) = Q(s,a) - U(s)\). Using the state value function in baseline subtraction produces the advantage. The policy gradient using the advantage is unbiased and typically has much lower
struct BaselineSubtractionGradient
    P # problem
    b # initial state distribution
    d # depth
    m # number of samples
    \nabla \log \pi # gradient of log likelihood
end

function gradient(M::BaselineSubtractionGradient, \pi, \theta)
    \P, b, d, m, \nabla \log \pi, \gamma = M.P, M.b, M.d, M.m, M.nabla, M.P.gamma
    \pi\theta(s) = \pi(\theta, s)
    \ell(a, s, k) = \nabla \log \pi(\theta, a, s) \gamma^{k-1}
    R(\tau, k) = \sum(\tau[^{\gamma}(j-1) \text{ for } (j, (s, a, r)) \text{ in enumerate}(\tau[k:end]))
    \text{num}er(\tau) = \sum(\ell(a, s, k) \cdot 2 \cdot R(\tau, k) \text{ for } (k, (s, a, r)) \text{ in enumerate}(\tau))
    \text{denom}(\tau) = \sum(\ell(a, s, k) \cdot 2 \text{ for } (k, (s, a)) \text{ in enumerate}(\tau))
    \text{base}(\tau) = \text{num}er(\tau) / \text{denom}(\tau)
    \text{trajs} = [\text{simulate}(\P, \text{rand}(b), \pi\theta, d) \text{ for } i \text{ in } 1:m]
    r\text{base} = \text{mean}(\text{base}(\tau) \text{ for } \tau \text{ in } \text{trajs})
    \nabla U(\tau) = \sum(\ell(a, s, k) \cdot (R(\tau, k) - r\text{base}) \text{ for } (k, (s, a, r)) \text{ in enumerate}(\tau))
    \text{return } \text{mean}(\nabla U(\tau) \text{ for } \tau \text{ in } \text{trajs})
end

Algorithm 11.6. Likelihood ratio gradient estimation with reward-to-go and baseline subtraction for an MDP \P, policy \pi, and initial state distribution b. The gradient with respect to the parameterization vector \theta is estimated from m rollouts to depth d using the log policy gradients \nabla \log \pi.

Figure 11.3. Several policy gradient methods used to optimize policies for the simple regulator problem from the same initial parameterization. Each gradient evaluation ran 6 rollouts to depth 10. The magnitude of the gradient was limited to 1 and step updates were applied with a step size 0.2. The optimal policy parameterization is shown in black.
variance. The gradient computation has this form:

\[
\nabla U(\theta) = \mathbb{E}_\tau \left[ \sum_{k=1}^{d} \nabla \log \pi_\theta (a^{(k)} \mid s^{(k)}) \gamma^{k-1} A_\theta \left(s^{(k)}, a^{(k)}\right) \right]
\]  

(11.44)

As with the state and action value functions, the advantage function is typically unknown. Other methods, covered in chapter 13, are needed to approximate it.

11.6 Summary

- A gradient can be estimated using finite-differences.
- Linear regression can also be used to provide more robust estimates of the policy gradient.
- The policy gradient theorem can be used to derive a form of the policy gradient that separates the gradient of the action likelihood, which can typically be computed.
- The variance of the policy gradient can be significantly reduced using the reward-to-go and baseline subtraction.

11.7 Exercises

Exercise 11.1. If we estimate the expected discounted return of a given parameterized policy \( \pi_\theta \) defined by a \( n \)-dimensional vector of parameters \( \theta \) using \( m \) rollouts, how many total rollouts do we need to perform to compute the policy gradient using a finite difference approach?

Solution: In order to estimate the policy gradient using a finite difference approach, we need to estimate the utility of the policy given the current parameter vector \( U(\theta) \) as well as given all \( n \) variations of the current parameter vector \( U(\theta + \delta e^{(i)}) \) for \( i = 1 : n \). Since we estimate each of these using \( m \) rollouts, we need to perform a total of \( m(n + 1) \) rollouts.

Exercise 11.2. Consider policy gradients of the form:

\[
\nabla_\theta R(\pi) = \mathbb{E}_\tau \left[ \sum_{k=1}^{d} \gamma^{k-1} y \nabla \log \pi_\theta (a^{(k)} \mid s^{(k)}) \right]
\]

Which of the following values of \( y \) result in a valid policy gradient?

(a) \( \gamma^{1-k} \sum_{\ell=1}^{\infty} r^{(\ell)} \gamma^{\ell-1} \)
(b) \( \sum_{\ell=k}^{\infty} r(\ell) \gamma^{\ell-k} \)

(c) \( \left( \sum_{\ell=k}^{\infty} r(\ell) \gamma^{\ell-k} \right) - r_{base}(s(k)) \)

(d) \( U(s(k)) \)

(e) \( Q(s(k), a(k)) \)

(f) \( A(s(k), a(k)) \)

(g) \( r^{(k)} + \gamma U(s^{(k+1)}) - U(s^{(k)}) \)

Solution:

(a) \( \sum_{\ell=1}^{\infty} r^{(\ell)} \) results in the total discounted reward, as

\[
\gamma^{k-1} \sum_{\ell=1}^{\infty} r(\ell) \gamma^{\ell-1} = \sum_{\ell=1}^{\infty} r(\ell) \gamma^{\ell-1}
\]

and produces a valid policy gradient as given in equation (11.19).

(b) \( \sum_{\ell=k}^{\infty} r(\ell) \gamma^{\ell-k} \) is the reward-to-go, and produces a valid policy gradient as given in equation (11.26).

(c) \( \left( \sum_{\ell=k}^{\infty} r(\ell) \right) - b(s^{(k)}) \) is the baseline subtracted reward-to-go, and produces a valid policy gradient as given in equation (11.28).

(d) \( U(s^{(k)}) \) is the state value function, and does not produce a valid policy gradient.

(e) \( Q(s^{(k)}, a^{(k)}) \) is the state-action value function, and produces a valid policy gradient as given in equation (11.27).

(f) \( A(s^{(k)}, a^{(k)}) \) is the advantage function, and produces a valid policy gradient as given in equation (11.44).

(g) \( r^{(k)} + \gamma U(s^{(k+1)}) - U(s^{(k)}) \) is the temporal difference residual, and produces a valid policy gradient because it is an unbiased approximation to the advantage function.

Exercise 11.3. Show that \( \mathbb{E}_{\tau \sim \pi} [f(\tau)] = \mathbb{E}_{\tau_{1:k} \sim \pi} [\mathbb{E}_{\tau_{k+1} \sim \pi} [f(\tau)]] \) for some step \( k \).

Solution: The nested expectations can be proven by writing the expectation in integral form and then converting back:
\[ E_{\tau \sim \pi} [f(\tau)] = \]
\[ = \int p(\tau) f(\tau) \, d\tau \]
\[ = \int \left( p(s^{(1)}) \prod_{k=1}^{d} p(s^{(k+1)} | s^{(k)}, a^{(k)}) \pi(a^{(k)} | s^{(k)}) \right) f(\tau) \, d\tau \]
\[ = \int \int \int \ldots \int \left( p(s^{(1)}) \prod_{k=1}^{d} p(s^{(k+1)} | s^{(k)}, a^{(k)}) \pi(a^{(k)} | s^{(k)}) \right) f(\tau) \, ds^{(d)} \ldots da^{(2)} \, ds^{(2)} \, da^{(1)} \, ds^{(1)} \]
\[ = E_{\tau_{1:k} \sim \pi} \left[ \int \int \int \ldots \int \left( \prod_{q=k}^{d} p(s^{(q+1)} | s^{(q)}, a^{(q)}) \pi(a^{(q)} | s^{(q)}) \right) f(\tau) \, ds^{(d)} \ldots da^{(k+1)} \, ds^{(k+1)} \, da^{(k)} \, ds^{(k)} \right] \]
\[ = E_{\tau_{1:k} \sim \pi} [E_{\tau_{k:d} \sim \pi} [f(\tau)]] \]

**Exercise 11.4.** Our implementation of the regression gradient (algorithm 11.3) fits a linear mapping from perturbations to the difference in returns, \( U(\theta + \Delta \theta^{(i)}) - U(\theta) \). We evaluate \( U(\theta + \Delta \theta^{(i)}) \) and \( U(\theta) \) for each of the \( m \) perturbations, thus re-evaluating \( U(\theta) \) a total of \( m \) times. How might we reallocate the samples in a more effective manner?

**Solution:** One approach is to evaluate \( U(\theta) \) once and use the same value for each perturbation, thereby conducting only \( m + 1 \) evaluations. Having an accurate estimate of \( U(\theta) \) is particularly important for an accurate regression gradient estimate. An alternative is to still compute \( U(\theta) \) once, but use \( m \) rollouts, thus preserving the total number of rollouts per iteration. This approach uses the same amount of computation as algorithm 11.3, but it can produce a more reliable gradient estimate.
We can use estimates of the policy gradient to drive the search of the parameter space towards an optimal policy. The previous chapter outlined methods for estimating this gradient. This chapter explains how to use these estimates to guide the optimization. We begin with gradient ascent, which simply takes steps in the direction of the gradient at each iteration. A major challenge is in determining the step size. Large steps can lead to faster progress to the optimum, but it can overshoot. The natural policy gradient modifies the direction of the gradient to better handle variable levels of sensitivity across parameter components. We conclude with the trust region method, which starts in exactly the same way as the natural gradient method to obtain a candidate policy. It then searches along the line segment in policy space connecting the original policy to this candidate to find a better policy.

### 12.1 Gradient Ascent Update

We can use gradient ascent (reviewed in appendix A.11) to find a policy parameterized by $\theta$ that optimizes expected utility $U(\theta)$. Gradient ascent is a type of iterated ascent method, which involves taking steps in the parameter space at each iteration in an attempt to improve the quality of the associated policy. All of the methods discussed in this chapter are iterated ascent methods, but they differ in how they take steps. The gradient ascent method discussed in this section takes steps in the direction of $\nabla U(\theta)$, which may be estimated using one of the methods discussed in the previous chapter. The update of $\theta$ is:

$$\theta \leftarrow \theta + \alpha \nabla U(\theta)$$  \hspace{1cm} (12.1)
where $\alpha > 0$ is the step factor. The step length is equal to some step factor $\alpha > 0$ times the magnitude of the gradient.

Algorithm 12.1 provides an implementation of a function that takes this step. This function can be called for either a fixed number of iterations or until $\theta$ or $U(\theta)$ converges. This algorithm, as well as the other algorithms discussed in this chapter, are not guaranteed to converge to the optimal policy. However, there are techniques to encourage convergence to a \textit{locally optimal} policy, which is one where taking an infinitesimally small step in parameter space cannot result in a better policy. One approach is to decay the step factor with each step.\footnote{This approach as well as many others are covered in detail by M. J. Kochenderfer and T. A. Wheeler, \textit{Algorithms for Optimization}. MIT Press, 2019.}

Algorithm 12.1. The gradient ascent method for policy optimization. It takes a step from a point $\theta$ in the direction of the gradient $\nabla U$ with step factor $\alpha$. We can use one of the methods in the previous chapter to compute $\nabla U$.

Very large gradients tend to overshoot the optimum and may occur due to a variety of reasons. Rewards for some problems, such as for the 2048 problem, can vary by orders of magnitude. One approach for keeping the gradients manageable is to use \textit{gradient scaling}, which limits the magnitude of a gradient estimate before using it to the update the policy parameterization. Gradients are commonly limited to having an $L_2$-norm of 1. Another approach is \textit{gradient clipping}, which clamps each element of the gradient when using it to the update the policy. Clipping commonly limits the entries to lie between $\pm 1$. Both techniques are implemented in algorithm 12.2.

Algorithm 12.2. Methods for gradient scaling and clipping. Gradient scaling limits the magnitude of the provided gradient vector $\nabla$ to $L_2_{\text{max}}$. Gradient clipping provides elementwise clamping of the provided gradient vector $\nabla$ to between $a$ and $b$.
12.2 Restricted Gradient Update

The remaining algorithms in this chapter attempt to optimize an approximation of the objective function \( U(\theta) \) subject to a constraint that the policy parameters at the next step \( \theta' \) are not too far away from \( \theta \) at the current step. The constraint takes the form \( g(\theta, \theta') \leq \epsilon \), where \( \epsilon > 0 \) is a free parameter in the algorithm. The methods differ in their approximation of \( U(\theta) \) and the form of \( g \). This section describes a simple restricted step method.

We use the first order Taylor approximation (appendix A.12) obtained from our gradient estimate at \( \theta \) to approximate \( U \):

\[
U(\theta') \approx U(\theta) + \nabla U(\theta)^\top (\theta' - \theta) \tag{12.2}
\]

For the constraint, we use

\[
g(\theta, \theta') = \frac{1}{2} (\theta' - \theta)^\top I (\theta' - \theta) = \frac{1}{2} \| \theta' - \theta \|_2^2 \tag{12.3}
\]

We can view this constraint as limiting the step length to be no more than \( \sqrt{2\epsilon} \)—or, in other words, the feasible region in our optimization is a ball of radius \( \sqrt{2\epsilon} \) centered at \( \theta \).

The optimization problem is then

\[
\begin{align*}
\text{maximize} & \quad U(\theta) + \nabla U(\theta)^\top (\theta' - \theta) \\
\text{subject to} & \quad \frac{1}{2} (\theta' - \theta)^\top I (\theta' - \theta) \leq \epsilon 
\end{align*} \tag{12.4}
\]
In the objective, we can drop $U(\theta)$ since it does not depend on $\theta'$. In addition, we can change the inequality to an equality in the constraint because the linear objective forces the optimal solution to be on the boundary of the feasible region. These changes result in an equivalent optimization problem:

$$\text{maximize} \quad \nabla U(\theta)^\top (\theta' - \theta)$$

subject to $$\frac{1}{2} (\theta' - \theta)^\top I (\theta' - \theta) = \epsilon$$

This optimization problem can be solved analytically:

$$\theta' = \theta + u \sqrt{\frac{2\epsilon}{u^\top u}} = \theta + \sqrt{2\epsilon} \frac{u}{||u||}$$

where the unnormalized search direction $u$ is simply $\nabla U(\theta)$. Of course, we do not know $\nabla U(\theta)$ exactly, but we can use any of the methods described in the previous chapter to estimate it. Algorithm 12.3 provides an implementation.

**Algorithm 12.3.** The update function for the restricted gradient method at $\theta$ for a problem $\mathcal{P}$ with initial state distribution $b$. The gradient is estimated from an initial state distribution $b$ to depth $d$ with $m$ simulations of parameterized policy $\pi(\theta, s)$ with log policy gradient $\nabla \log \pi$. 

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12.3 Natural Gradient Update

The natural gradient method\(^2\) is a variation of the restricted step method in the previous section to better handle situations when some components of the parameter space are more sensitive than others. Sensitivity in this context refers to how much the utility of a policy varies with respect to small changes in one of the parameters. The sensitivity in gradient methods is largely determined by the choice of scaling of the policy parameters. The natural policy gradient method makes the search direction \( \mathbf{u} \) invariant to parameter scaling. Figure 12.2 illustrates the differences between the true gradient and the natural gradient.

![Figure 12.2. A comparison of the true gradient and the natural gradient on the simple regulator problem (see appendix F.5). The true gradient generally points strongly in the negative \( \theta_2 \) direction, whereas the natural gradient generally points towards the optimum (black dot) at \([-1, 0]\). A similar figure is presented in J. Peters and S. Schaal, “Reinforcement Learning of Motor Skills with Policy Gradients,” Neural Networks, vol. 21, no. 4, pp. 682–697, 2008.](image)

The natural policy gradient method uses the same first-order approximation of the objective as the previous section. The constraint, however, is different. The intuition is that we want to restrict changes in \( \theta \) that result in large changes in the distribution over trajectories. A way to measure how much a distribution changes is to use the Kullback-Leibler divergence (appendix A.10). We could impose the constraint

\[
g(\theta, \theta') = D_{KL}(p(\cdot | \theta) \mid \mid p(\cdot | \theta')) \leq \epsilon
\]  

(12.7)

but instead we will use a second-order Taylor approximation:

\[
g(\theta, \theta') = \frac{1}{2}(\theta' - \theta)^\top F_\theta (\theta' - \theta) \leq \epsilon
\]  

(12.8)

where

\[ F_\theta = \int p(\tau \mid \theta) \nabla \log p(\tau \mid \theta) \nabla \log p(\tau \mid \theta)^\top \, d\tau \]  \hspace{1cm} (12.9)

\[ = \mathbb{E}_\tau \left[ \nabla \log p(\tau \mid \theta) \nabla \log p(\tau \mid \theta)^\top \right] \hspace{1cm} (12.10) \]

The resulting optimization problem is

\[
\begin{align*}
\text{maximize} & \quad \nabla U(\theta)^\top (\theta' - \theta) \\
\text{subject to} & \quad \frac{1}{2} (\theta' - \theta)^\top F_\theta (\theta' - \theta) = \epsilon \hspace{1cm} (12.11)
\end{align*}
\]

which looks identical to equation (12.5) except that, instead of the identity matrix \( I \), we have \( F_\theta \). This difference results in an ellipsoid feasible set. Figure 12.3 shows an example in two dimensions.

This optimization problem can be solved analytically and has the same form as the update in the previous section:

\[
\theta' = \theta + u \sqrt{\frac{2\epsilon}{\nabla U(\theta)^\top u}} \hspace{1cm} (12.12)
\]

except that we now have

\[
u = F_\theta^{-1} \nabla U(\theta) \hspace{1cm} (12.13)\]

We can use sampled trajectories to estimate \( F_\theta \) and \( \nabla U(\theta) \). Algorithm 12.4 provides an implementation.

### 12.4 Trust Region Update

This section discusses a method for searching within the trust region defined by the elliptical feasible region from the previous section. This category of approach is referred to as trust region policy optimization (TRPO).\(^4\) It works by computing the next evaluation point \( \theta' \) that would be taken by the natural policy gradient and then conducting a line search along the line segment connecting \( \theta \) to \( \theta' \). A key property of this line search phase is that evaluations of the approximate objective and constraint do not require any additional rollout simulations.

During the line search phase, we no longer use a first order approximation. Instead, we use an approximation derived from an equality involving the advantage function\(^5\)

\[
U(\theta') = U(\theta) + \mathbb{E}_{\tau \sim \pi_{\theta'}} \left[ \sum_{k=1}^{d} A_{\theta}(s^{(k)}, a^{(k)}) \right] \hspace{1cm} (12.14)
\]

\(^3\) This computation can be done using conjugate gradient descent, which reduces computation when the dimension of \( \theta \) is large. S.M. Kakade, “A Natural Policy Gradient,” in Advances in Neural Information Processing Systems (NIPS), 2001.


Algorithm 12.4. The update function for the natural policy gradient given policy $\pi(\theta, s)$ for an MDP $\mathcal{P}$ with initial state distribution $b$. The natural gradient with respect to the parameter vector $\theta$ is estimated from $m$ rollouts to depth $d$ using the log policy gradients $\nabla \log \pi$. The `natural-update` helper method conducts an update according to equation (12.12) given an objective gradient $\nabla f(\tau)$ and a Fisher matrix $F(\tau)$ for a list of trajectories.
Another way to write this is to use $b_{\gamma, \theta}$, which is the discounted visitation distribution of state $s$ under policy $\pi_\theta$, where

$$b_{\gamma, \theta}(s) \propto P(s^{(1)} = s) + \gamma P(s^{(2)} = s) + \gamma^2 P(s^{(3)} = s) + \cdots \quad (12.15)$$

Using the discounted visitation distribution, the objective becomes

$$U(\theta') = U(\theta) + \mathbf{E}_{s \sim b_{\gamma, \theta'}} \left[ \mathbf{E}_{a \sim \pi_{\theta'}(\cdot | s)} \left[ A_\theta(s, a) \right] \right] \quad (12.16)$$

We would like to pull our samples from our policy parameterized by $\theta$ instead of $\theta'$ so that we do not have to run more simulations during the line search. The samples associated with the inner expectation can be replaced with samples from our original policy so long as we appropriately weight the advantage:

$$U(\theta') = U(\theta) + \mathbf{E}_{s \sim b_{\gamma, \theta}} \left[ \mathbf{E}_{a \sim \pi_\theta(\cdot | s)} \left[ \frac{\pi_{\theta'}(a | s)}{\pi_\theta(a | s)} A_\theta(s, a) \right] \right] \quad (12.17)$$

The next step involves replacing the state distribution with $b_{\gamma, \theta}$. The quality of the approximation degrades as $\theta'$ gets further from $\theta$, but it is hypothesized that it is acceptable within the trust region. Since $U(\theta)$ does not depend on $\theta'$, we can drop it from the objective. We can also drop the state value function from the advantage function, leaving us with the action value function. What remains is referred to as the surrogate objective:

$$f(\theta, \theta') = \mathbf{E}_{s \sim b_{\gamma, \theta}} \left[ \mathbf{E}_{a \sim \pi_\theta(\cdot | s)} \left[ \frac{\pi_{\theta'}(a | s)}{\pi_\theta(a | s)} Q_\theta(s, a) \right] \right] \quad (12.18)$$

This equation can be estimated from the same set of trajectories that was used to estimate the natural gradient update. The $Q_\theta(s, a)$ can be estimated from the reward-to-go in the sampled trajectories.

The surrogate constraint in the line search is given by

$$g(\theta, \theta') = \mathbf{E}_{s \sim b_{\gamma, \theta}} \left[ D_{KL}(\pi_{\theta'}(\cdot | s) || \pi_{\theta}(\cdot | s)) \right] \leq \epsilon \quad (12.19)$$

Line search involves iteratively evaluating our surrogate objective $f$ and surrogate constraint $g$ for different points in the policy space. We begin with the $\theta'$ obtained from the same process as the natural gradient update. We then iteratively apply

$$\theta' \leftarrow \theta + \alpha(\theta' - \theta) \quad (12.20)$$
until we have an improvement in our objective with \( f(\theta, \theta') > f(\theta, \theta) \) and our constraint is met with \( g(\theta, \theta') \leq \epsilon \). The step factor \( 0 < \alpha < 1 \) shrinks the distance between \( \theta \) and \( \theta' \) at each iteration, with \( \alpha \) typically set to 0.5.

Algorithm 12.5 provides an implementation of this approach. Figure 12.4 illustrates the relationship between the feasible regions associated with the natural gradient and the line search. Figure 12.5 demonstrates the approach on a regulator problem, and example 12.1 shows an update for a simple problem.

### 12.5 Clamped Surrogate Objective

We can avoid detrimental policy updates from overly optimistic estimates of the trust region surrogate objective by clamping. The surrogate objective from equation (12.18), after exchanging the action value advantage, is

\[
E_{s \sim b_{\gamma, \theta}} \left[ E_{a \sim \pi_{\theta}(\cdot | s)} \left[ \frac{\pi_{\theta'}(a | s)}{\pi_{\theta}(a | s)} A_{\theta}(s, a) \right] \right] \tag{12.21}
\]

The probability ratio \( \frac{\pi_{\theta'}(a | s)}{\pi_{\theta}(a | s)} \) can be overly optimistic. A pessimistic lower bound on the objective can significantly improve performance:

\[
E_{s \sim b_{\gamma, \theta}} \left[ E_{a \sim \pi_{\theta}(\cdot | s)} \left[ \min \left( \frac{\pi_{\theta'}(a | s)}{\pi_{\theta}(a | s)} A_{\theta}(s, a), \operatorname{clamp} \left( \frac{\pi_{\theta'}(a | s)}{\pi_{\theta}(a | s)}, 1 - \epsilon, 1 + \epsilon \right) \right) \right] \right] \tag{12.22}
\]

where \( \epsilon \) is a small positive value and \( \operatorname{clamp}(x, a, b) \) forces \( x \) to be between \( a \) and \( b \). By definition, \( \operatorname{clamp}(x, a, b) = \min\{\max\{x, a\}, b\} \).

Clamping the probability ratio alone does not produce a lower bound; we must also take the minimum of the clamped and original objectives. The lower bound is shown in figure 12.6 together with the original and clamped objectives. The end result of the lower bound is that the change in probability ratio is ignored when it would cause the objective to significantly improve. Using the lower bound thus prevents large, often detrimental, updates in these situations, and removes the need for the trust region surrogate constraint equation (12.19). Without the constraint, we can also eliminate line search and use standard gradient ascent methods.

The gradient of the unclamped objective equation (12.21) with action values is

\[
\nabla_{\theta'} f(\theta, \theta') = E_{s \sim b_{\gamma, \theta}} \left[ E_{a \sim \pi_{\theta}(\cdot | s)} \left[ \frac{\nabla_{\theta'} \pi_{\theta'}(a | s)}{\pi_{\theta}(a | s)} Q_{\theta}(s, a) \right] \right] \tag{12.23}
\]

Algorithm 12.5. The update procedure for trust region policy optimization, which augments the natural gradient with a line search. It generates $m$ trajectories using policy $\pi$ in problem $\mathcal{P}$ with initial state distribution $b$ and depth $d$. To obtain the starting point of the line search, we need the gradient of the log-probability of the policy generating a particular action from the current state, which we denote $\nabla \log \pi$. For the surrogate objective, we need the probability function $p$ that gives the probability our policy generates a particular action from the current state. For the surrogate constraint, we need the divergence between the action distributions generated by $\pi_\theta$ and $\pi_{\theta'}$. At each step of the line search, we shrink the distance between the considered point $\theta'$ and $\theta$ while maintaining the search direction.
12.5. CLAMPED SURROGATE OBJECTIVE

\[
\frac{1}{2} (\theta' - \theta)^\top F_\theta (\theta' - \theta) = \epsilon
\]

\[
g(\theta, \theta') = \epsilon
\]

\[
\nabla U_{\pi_\theta} (\theta)
\]

\[
\mathbf{F}_\theta^{-1} \nabla U(\theta)
\]

\[
\theta'\]

Figure 12.4. Trust region policy optimization searches within the elliptical constraint generated by a second-order approximation of the Kullback-Leibler divergence. After computing the natural policy gradient ascent direction, a line search is conducted to ensure that the updated policy improves the policy reward and adheres to the divergence constraint. The line search starts from the estimated maximum step size and reduces the step size along the ascent direction until a satisfactory point is found.

\[
\theta^* - 1 - 0.5 0 0.5 1
\]

\[
\theta_1
\]

\[
\phi_i
\]

\[
\theta_i
\]

\[
\text{Expected Reward}
\]

\[
0 2 4 6 8
\]

\[
\text{iteration}
\]

Figure 12.5. Trust region policy optimization applied to the simple regulator problem with rollouts to depth 10 with \(\epsilon = 1\) and \(c = 2\). The optimal policy parameterization is shown in black.
Consider applying TRPO to the Gaussian policy $\mathcal{N}(\theta_1, \theta_2^2)$ from example 11.3 to the single-state MDP from example 11.1. Recall that the gradient of the log policy likelihood is:

\[
\frac{\partial}{\partial \theta_1} \log \pi_{\theta}(a \mid s) = \frac{a - \theta_1}{\theta_2^2}, \\
\frac{\partial}{\partial \theta_2} \log \pi_{\theta}(a \mid s) = \frac{(a - \theta_1)^2 - \theta_2^2}{\theta_2^2}.
\]

Suppose we run two rollouts with $\theta = [0, 1]$ (This problem only has one state):

\[
\tau_1 = \{(a = r = -0.532), (a = r = 0.597), (a = r = 1.947)\} \\
\tau_2 = \{(a = r = -0.263), (a = r = -2.212), (a = r = 2.364)\}
\]

The estimated Fisher information matrix is:

\[
F_\theta = \frac{1}{2} \left( \nabla \log p(\tau^{(1)})^\top \nabla \log p(\tau^{(1)}) + \nabla \log p(\tau^{(2)})^\top \nabla \log p(\tau^{(2)}) \right) \\
= \frac{1}{2} \begin{bmatrix} 4.048 & 2.878 \\ 2.878 & 2.046 \end{bmatrix} + \begin{bmatrix} 0.012 & -0.838 \\ -0.838 & 29.529 \end{bmatrix} = \begin{bmatrix} 2.030 & 1.020 \\ 1.019 & 118.115 \end{bmatrix}
\]

The objective function gradient is $[2.030, 1.020]$. The resulting descent direction $u$ is $[1, 0]$. Setting $\epsilon = 0.1$, we compute our updated parameterization vector and obtain $\theta' = [0.314, 1]$.

The surrogate objective function value at $\theta$ is 1.485. Line search begins at $\theta'$, where the surrogate objective function value is 2.110 and the constraint yields 0.049. This satisfies our constraint (as 0.049 < $\epsilon$), so we return the new parameterization.
where \( Q_\theta(s, a) \) can be estimated from reward-to-go. The gradient of the clamped objective equation (12.22) is the same, except there is no contribution from experience tuples for which the objective is actively clamped. That is, if either the reward-to-go is positive and the probability ratio is less than \( 1 + \epsilon \), or if the reward-to-go is negative and the probability ratio is greater than \( 1 - \epsilon \), the gradient contribution is zero.

Like trust region policy optimization, the gradient can be computed for a parameterization \( \theta' \) from experience generated from \( \theta \). Hence, several gradient updates can be run in a row using the same set of sampled trajectories. Algorithm 12.6 provides an implementation.

The clamped surrogate objective is compared to several other surrogate objectives in figure 12.7. The figure includes a line plot for the effective objective for trust region policy optimization,

\[
\mathbb{E}_{s \sim b_{r, \theta}, a \sim \pi_{\theta}(\cdot | s)} \left[ \frac{\pi_{\theta'}(a | s)}{\pi_{\theta}(a | s)} A_\theta(s, a) - \beta D_{KL}(\pi_{\theta}(\cdot | s) \parallel \pi_{\theta'}(\cdot | s)) \right] = (12.24)
\]

which is the trust region policy objective where the constraint is implemented as a penalty for some coefficient \( \beta \). Trust region policy optimization typically uses a hard constraint rather than a penalty because it is difficult to choose a \( \beta \) that performs well within a single problem, let alone across problems.
struct ClampedSurrogateUpdate
    ℋ # problem
    ℬ # initial state distribution
    d # depth
    m # number of trajectories
    π # policy
    p # policy likelihood
    ∇π # policy likelihood gradient
    ɛ # divergence bound
    α # step size
    k_max # number of iterations per update
end

function clamped_gradient(M::ClampedSurrogateUpdate, θ, θ′, τs)
    d, p, ∇π, ɛ, γ = M.d, M.p, M.∇π, M.ɛ, M.ℋ.γ
    R(τ, j) = sum(r*γ^(k-1) for (k,(s,a,r)) in zip(j:d, τ[j:end]))
    ∇f(a,s,r_togo) = begin
        P = p(θ, a,s)
        w = p(θ′,a,s) / P
        if (r_togo > 0 && w > 1+ɛ) || (r_togo < 0 && w < 1-ɛ)
            return zeros(length(θ))
        end
        return ∇π(θ′, a, s) * r_togo / P
    end
    ∇f(τ) = mean(∇f(a,s,R(τ,k)) for (k,(s,a,r)) in enumerate(τ))
    return mean(∇f(τ) for τ in τs)
end

function update(M::ClampedSurrogateUpdate, θ)
    ℋ, ℬ, d, m, π, α, k_max= M.ℋ, M.ℬ, M.d, M.m, M.π, M.α, M.k_max
    πθ(s) = π(θ, s)
    τs = [simulate(ℋ, rand(ℬ), πθ, d) for i in 1:m]
    θ’ = copy(θ)
    for k in 1:k_max
        θ’ += α*clamped_gradient(M, θ, θ’, τs)
    end
    return θ’
end

Algorithm 12.6. An implementation of clamped surrogate policy optimization, which returns a new policy parameterization for policy π(s) of an MDP ℋ with initial state distribution ℬ. This implementation samples m trajectories to depth d, and then uses them to estimate the policy gradient in k_max subsequent updates. The policy gradient using the clamped objective is constructed using the policy gradients ∇p with clamping parameter ɛ.
12.6 Summary

- The gradient ascent algorithm can use the gradient estimates obtained from the methods discussed in the previous chapter to iteratively improve our policy.

- Gradient ascent can be made more robust by scaling, clipping, or forcing the size of the improvement steps to be uniform.

- The natural gradient approach uses a first order approximation of the objective function with a constraint on the divergence between the trajectory distribution at each step, approximated using an estimate of the Fisher information matrix.

- Trust region policy optimization involves augmenting the natural gradient method with a line search to further improve the policy without additional trajectory simulations.

- We can use a pessimistic lower bound of the trust region policy optimization objective to obtain a clamped surrogate objective that performs similarly without the need for line search.

12.7 Exercises

Exercise 12.1. Trust region policy optimization starts its line search from a new parameterization given by a natural policy gradient update. However, TRPO conducts the line search
using a different objective than the natural policy gradient. Show that the gradient of the surrogate objective equation (12.18) used in TRPO is actually the same as the reward-to-go policy gradient equation (11.26).

**Solution:**

The gradient of TRPO’s surrogate objective is:

\[
\nabla_\theta U_{TRPO} = \mathbb{E}_{s \sim b, a \sim \pi_\theta (\cdot | s)} \left[ \nabla_\theta \pi_\theta (a | s) \right] \frac{\nabla_\theta \pi_\theta (a | s)}{\pi_\theta (a | s)} Q_\theta (s, a)
\]

When conducting the initial natural policy gradient update, the search direction is evaluated at \( \theta' = \theta \). Furthermore, the action value is approximated with the reward-to-go:

\[
\nabla_\theta U_{TRPO} = \mathbb{E}_{s \sim b, a \sim \pi_\theta (\cdot | s)} \left[ \nabla_\theta \log \pi_\theta (a | s) \right] r_{\text{to-go}}
\]

Recall that the derivative of \( \log f(x) \) is \( f'(x)/f(x) \). It thus follows that:

\[
\nabla_\theta U_{TRPO} = \mathbb{E}_{s \sim b, a \sim \pi_\theta (\cdot | s)} \left[ \nabla_\theta \log \pi_\theta (a | s) \right] r_{\text{to-go}}
\]

which has the same form as the reward-to-go policy gradient equation (11.26).

**Exercise 12.2.** Perform the calculations of example 12.1. First, compute the inverse of the Fisher information matrix \( F_\theta^{-1} \), compute \( u \), and compute the updated parameters \( \theta' \).

**Solution:** We start by computing the inverse of the Fisher information matrix:

\[
F_\theta^{-1} \approx \frac{1}{0.341(118.115) - 0.332(0.332)} \begin{bmatrix} 118.115 & -0.332 \\ -0.332 & 0.341 \end{bmatrix} \approx \begin{bmatrix} 2.941 & -0.008 \\ -0.008 & 0.008 \end{bmatrix}
\]

Now, we update \( u \) as follows,

\[
u = F_\theta^{-1} \nabla U(\theta) \approx \begin{bmatrix} 2.941 & -0.008 \\ -0.008 & 0.008 \end{bmatrix} \begin{bmatrix} 0.341 \\ 0.332 \end{bmatrix} \approx \begin{bmatrix} 1 \\ 0 \end{bmatrix}
\]

Finally, we estimate the updated parameters \( \theta' \):

\[
\theta' = \theta + \frac{2\epsilon}{\nabla U(\theta)^T u}
\]

\[
\approx \begin{bmatrix} 0 \\ 1 \end{bmatrix} + \frac{1}{\sqrt{\begin{bmatrix} 0.341 & 0.332 \\ 0.332 & 0.341 \end{bmatrix} \begin{bmatrix} 1 \\ 0 \end{bmatrix}}}
\]

\[
\approx \begin{bmatrix} 0 \\ 1 \end{bmatrix} + \sqrt{\begin{bmatrix} 0.2 \\ 0.341 \end{bmatrix} \begin{bmatrix} 0.766 \\ 1 \end{bmatrix}}
\]

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2021-03-07 23:12:00-08:00, revision 80bf4d6, comments to bugs@algorithmsbook.com
Exercise 12.3. Suppose we have the following parameterized policies $\pi_\theta$ and $\pi_{\theta'}$ given in the table below. Given that we sample the following five states, $s_1, s_2, s_1, s_1, s_2$, approximate $\mathbb{E}_s [D_{KL} (\pi_\theta (\cdot | s) \mid\mid \pi_{\theta'} (\cdot | s))]$ using the definition

$$D_{KL} (P \mid\mid Q) = \sum_x P(x) \log \frac{P(x)}{Q(x)}$$

<table>
<thead>
<tr>
<th></th>
<th>$a_1$</th>
<th>$a_2$</th>
<th>$a_3$</th>
<th>$a_4$</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\pi_\theta (a</td>
<td>s_1)$</td>
<td>0.1</td>
<td>0.2</td>
<td>0.3</td>
</tr>
<tr>
<td>$\pi_{\theta'} (a</td>
<td>s_1)$</td>
<td>0.4</td>
<td>0.3</td>
<td>0.2</td>
</tr>
<tr>
<td>$\pi_\theta (a</td>
<td>s_2)$</td>
<td>0.1</td>
<td>0.1</td>
<td>0.6</td>
</tr>
<tr>
<td>$\pi_{\theta'} (a</td>
<td>s_2)$</td>
<td>0.1</td>
<td>0.1</td>
<td>0.5</td>
</tr>
</tbody>
</table>

Solution: First, we compute the KL divergence for a state sample $s_1$:

$$D_{KL} (\pi_\theta (\cdot | s_1) \mid\mid \pi_{\theta'} (\cdot | s_1)) = 0.1 \log \left( \frac{0.1}{0.4} \right) + 0.2 \log \left( \frac{0.2}{0.3} \right) + 0.3 \log \left( \frac{0.3}{0.3} \right) + 0.4 \log \left( \frac{0.4}{0.1} \right) \approx 0.456$$

Now, we compute the KL divergence for a state sample $s_2$:

$$D_{KL} (\pi_\theta (\cdot | s_2) \mid\mid \pi_{\theta'} (\cdot | s_2)) = 0.1 \log \left( \frac{0.1}{0.1} \right) + 0.1 \log \left( \frac{0.1}{0.1} \right) + 0.6 \log \left( \frac{0.6}{0.5} \right) + 0.2 \log \left( \frac{0.2}{0.3} \right) \approx 0.0283$$

Finally, we compute the approximation of the expectation, which is the average KL divergence of the parameterized policies over the $n$ state samples

$$\mathbb{E}_s [D_{KL} (\pi_\theta (\cdot | s) \mid\mid \pi_{\theta'} (\cdot | s))] \approx \frac{1}{n} \sum_{i=1}^{n} D_{KL} \left( \pi_\theta (\cdot | s^{(i)}) \mid\mid \pi_{\theta'} (\cdot | s^{(i)}) \right)$$

$$\approx \frac{1}{5} \left( 0.456 + 0.0283 + 0.456 + 0.456 + 0.0283 \right)$$

$$\approx 0.285$$
13 Actor-Critic Methods

The previous chapter discussed ways to improve a parameterized policy through gradient information estimated from rollouts. This chapter introduces actor-critic methods, which uses an estimate of a value function to help direct the optimization. The actor, in this context, is the policy, and the critic is the value function. Both are trained in parallel. We will discuss several methods that differ in whether they approximate the value function, advantage function, or action value function. Most focus on stochastic policies, but we will also discuss one method that supports deterministic policies that output continuous actions. Finally, we will discuss a way to incorporate an online method for generating more informative trajectories for training the actor and critic.

13.1 Actor-Critic

In actor-critic methods, we have an actor represented by a policy \( \pi_\theta \) parameterized by \( \theta \) with the help of a critic that provides an estimate of the value function \( U_\phi(s) \), \( Q_\phi(s,a) \), or \( A_\phi(s,a) \) parameterized by \( \phi \). We will start this chapter with a simple actor-critic approach where the optimization of \( \pi_\theta \) is done through gradient ascent with the gradient of our objective being the same as in equation (11.44):

\[
\nabla U(\theta) = \mathbb{E}_\tau \left[ \sum_{k=1}^{d} \nabla_{\theta} \log \pi_\theta(a^{(k)} \mid s^{(k)}) \gamma^{k-1} A_\theta(s^{(k)}, a^{(k)}) \right] \tag{13.1}
\]

The advantage when following a policy parameterized by \( \theta \) can be estimated using a set of observed transitions from \( s \) to \( s' \) with reward \( r \):

\[
A_\theta(s, a) = \mathbb{E}_{r,s'} [r + \gamma U_\pi(s') - U_\pi(s)] \tag{13.2}
\]
The $r + \gamma U^\pi_\theta(s') - U^\pi_\theta(s)$ inside the expectation is referred to as the \textit{temporal difference residual}.

The critic allows us to estimate the true value function $U^\pi_\theta$ when following $\pi_\theta$, resulting in the following gradient for the actor:

$$\nabla U(\theta) \approx \mathbb{E}_\tau \left[ \sum_{k=1}^d \nabla_\theta \log \pi_\theta(a^{(k)} \mid s^{(k)}) \gamma^{k-1} \left( r^{(k)} + \gamma U_\phi(s^{(k+1)}) - U_\phi(s^{(k)}) \right) \right]$$ (13.3)

This expectation can be estimated through rollout trajectories as done in chapter 11.

The critic is also updated through gradient optimization. We want to find a $\phi$ that minimizes our loss function:

$$\ell(\phi) = \frac{1}{2} \mathbb{E}_s \left[ (U_\phi(s) - U^\pi_\theta(s))^2 \right]$$ (13.4)

To minimize this objective, we can take steps in the opposite direction of the gradient:

$$\nabla \ell(\phi) = \mathbb{E}_s \left[ (U_\phi(s) - U^\pi_\theta(s)) \nabla_\phi U_\phi(s) \right]$$ (13.5)

Of course, we do not know $U^\pi_\theta$ exactly, but it can be estimated using the reward-to-go along rollout trajectories, resulting in

$$\nabla \ell(\phi) = \mathbb{E}_\tau \left[ \sum_{k=1}^d \left( U_\phi(s^{(k)}) - r^{(k)}_{\text{to-go}} \right) \nabla_\phi U_\phi(s^{(k)}) \right]$$ (13.6)

where $r^{(k)}_{\text{to-go}}$ is the reward-to-go at step $k$ in a particular trajectory $\tau$.

Algorithm 13.1 shows how to estimate $\nabla U(\theta)$ and $\nabla \ell(\phi)$ from rollouts. With each iteration, we step $\theta$ in the direction of $\nabla U(\theta)$ to maximize utility, and we step $\phi$ in the opposite direction of $\nabla \ell(\phi)$ to minimize our loss. This approach can become unstable due to the dependency between the estimation of $\theta$ and $\phi$, but this approach has worked well for a variety of problems. Some implementations update the policy more frequently than the value function to improve stability.

### 13.2 Generalized Advantage Estimation

Generalized advantage estimation (algorithm 13.2) is an actor-critic method that uses a more general version of the advantage estimate shown in equation (13.2) that allows us to balance between bias and variance.\(^1\) Approximation with the temporal difference residual has low variance but introduces bias due to a potentially

### 13.2. Generalized Advantage Estimation

An inaccurate $U_\phi$ used to approximate $U^{\pi_\theta}$. An alternative is to replace $r + \gamma U^{\pi_\theta}(s')$ with the sequence of rollout rewards $r_1, \ldots, r_d$:

$$A_\theta(s, a) = \mathbb{E}_{r_1, \ldots, r_d} \left[ r_1 + \gamma r_2 + \gamma^2 r_3 + \cdots + \gamma^{d-1} r_d - U^{\pi_\theta}(s) \right]$$

$$= \mathbb{E}_{r_1, \ldots, r_d} \left[ -U^{\pi_\theta}(s) + \sum_{\ell=1}^{d} \gamma^{\ell-1} r_\ell \right]$$

We can obtain an unbiased estimate of the expectation above through rollout trajectories as done in the policy gradient estimation methods (chapter 11). However, the estimate is high-variance, meaning we need many samples to arrive at an accurate estimate.

The approach taken by generalized advantage estimation is to balance between these two extremes of using temporal difference residuals and full rollouts. We define $\hat{A}^{(k)}$ to be the advantage estimate obtained from $k$ steps of a rollout and
An alternative way to write \( \hat{A}^{(k)} \) is in terms of an expectation over temporal difference residuals. We can define

\[
\delta_t = r_t + \gamma U(s_{t+1}) - U(s_t)
\]

where \( s_t, r_t, \) and \( s_{t+1} \) are the state, reward, and subsequent state along a sampled trajectory and \( U \) is our value function estimate. Then,

\[
\hat{A}^{(k)}(s, a) = \mathbb{E} \left[ \sum_{\ell=1}^{k} \gamma^{\ell-1} \delta_{\ell} \right]
\]

Instead of committing to a particular value for \( k \), generalized advantage estimation introduces a parameter \( \lambda \in [0, 1] \) that provides an exponentially weighted average of \( \hat{A}^{(k)} \) for \( k \) ranging from 1 to \( d \):

\[
\hat{A}_{\text{GAE}}(s, a) |_{d=1} = \hat{A}^{(1)}
\]

\[
\hat{A}_{\text{GAE}}(s, a) |_{d=2} = (1 - \lambda) \hat{A}^{(1)} + \lambda \hat{A}^{(2)}
\]

\[
\hat{A}_{\text{GAE}}(s, a) |_{d=3} = (1 - \lambda) \hat{A}^{(1)} + \lambda \left( (1 - \lambda) \hat{A}^{(2)} + \lambda \hat{A}^{(3)} \right)
\]

\[
= (1 - \lambda) \hat{A}^{(1)} + \lambda (1 - \lambda) \hat{A}^{(2)} + \lambda^2 \hat{A}^{(3)}
\]

\[
\vdots
\]

\[
\hat{A}_{\text{GAE}}(s, a) = (1 - \lambda) \left( \hat{A}^{(1)} + \lambda \hat{A}^{(2)} + \lambda^2 \hat{A}^{(3)} + \cdots + \lambda^{d-2} \hat{A}^{(d-1)} \right) + \lambda^{d-1} \hat{A}^{(d)}
\]

For an infinite horizon, the generalized advantage estimate simplifies to
\[ A_{\text{GAE}}(s,a) = (1 - \lambda) \left( A^{(1)} + \lambda A^{(2)} + \lambda^2 A^{(3)} + \cdots \right) \]
\[ = (1 - \lambda) \left( \delta_1 (1 + \lambda + \lambda^2 + \cdots) + \gamma \delta_2 (\lambda + \lambda^2 + \cdots) + \gamma^2 \delta_3 (\lambda^2 + \cdots) + \cdots \right) \]  
\[ = (1 - \lambda) \left( \delta_1 \frac{1}{1 - \lambda} + \gamma \delta_2 \frac{\lambda}{1 - \lambda} + \gamma^2 \delta_3 \frac{\lambda^2}{1 - \lambda} + \cdots \right) \]
\[ = E \left[ \sum_{k=1}^{\infty} (\gamma \lambda)^{k-1} \delta_k \right] \]

We can tune the parameter \( \lambda \) to balance between bias and variance. If \( \lambda = 0 \), then we have the high-bias, low-variance estimate for the temporal difference residual from the previous section. If \( \lambda = 1 \), we have the unbiased full rollout with increased variance. Figure 13.1 demonstrates the algorithm with different values for \( \lambda \).

Algorithm 13.2. Generalized advantage estimation for computing both a policy gradient and a value function gradient for an MDP \( \mathcal{P} \) with initial state distribution \( b \). The policy is parameterized by \( \theta \) and has a log-gradient \( \nabla \log \pi \). The value function \( U \) is parameterized by \( \phi \) and has a gradient \( \nabla U \). This method runs \( m \) rollouts to depth \( d \). The generalized advantage is computed with exponential weighting \( \lambda \) using equation (13.21) with a finite horizon. The implementation here is a simplified version of what was presented in the original paper, which included aspects of trust regions when taking steps.
13.3 Deterministic Policy Gradient

The deterministic policy gradient approach involves optimizing a deterministic policy $\pi_\theta(s)$ that produces continuous actions with the help of a critic in the form of a parameterized action value function $Q_\phi(s, a)$. As with the actor-critic methods we discussed so far, we define a loss function with respect to the parameterization $\phi$:

$$\ell(\phi) = \frac{1}{2} \mathbb{E}_{s,a,r,s'} \left[ (r + \gamma Q_\phi(s', \pi_\theta(s')) - Q_\phi(s, a))^2 \right]$$  \hspace{1cm} (13.22)

where the expectation is over the experience tuples generated by rollouts of $\pi_\theta$. This loss function attempts to minimize the residual of $Q_\phi$, similar to how the actor-critic method in the first section tried to minimize the residual of $U_\phi$.

Similar to the other methods, we update $\phi$ by taking a step in the opposite direction of the gradient:

$$\nabla \ell(\phi) = \mathbb{E}_{s,a,r,s'} \left[ (r + \gamma Q_\phi(s', \pi_\theta(s')) - Q_\phi(s, a)) \left( \gamma \nabla_\phi Q_\phi(s', \pi_\theta(s')) - \nabla_\phi Q_\phi(s, a) \right) \right]$$  \hspace{1cm} (13.23)

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2021-03-07 23:12:00-08:00, revision 80bf4d6, comments to bugs@algorithmsbook.com
We thus need a differentiable parameterized action value function from which we can compute \( \nabla_{\phi} Q_\phi (s, a) \), such as a neural network.

For the actor, we want to find a \( \theta \) that maximizes
\[
U(\theta) = \mathbb{E}_{s \sim b_{\pi_\theta}} [Q_\phi (s, \pi_\theta(s))] \tag{13.24}
\]
where the expectation is over the states from the discounted visitation frequency when following \( \pi_\theta \). Again, we can use gradient ascent to optimize \( \theta \) with the gradient given by
\[
\nabla U(\theta) = \mathbb{E}_s [\nabla_{\theta} Q_\phi (s, \pi_\theta(s))] \tag{13.25}
\]
\[
= \mathbb{E}_s [\nabla_{\theta} \pi_\theta(s) \nabla_a Q_\phi (s, a)|_{a=\pi_\theta(s)}] \tag{13.26}
\]
Here, \( \nabla_{\theta} \pi_\theta(s) \) is a Jacobian matrix whose \( i \)th column is the gradient with respect to the \( i \)th action dimension of the policy under the parameterization \( \theta \). An example for this term is given in example 13.1. The gradient \( \nabla_a Q_\phi (s, a)|_{a=\pi_\theta(s)} \) is a vector that indicates how much our estimated action value changes as we perturb the action given by our policy at state \( s \). In addition to the Jacobian, we also need to supply this gradient in order to use this method.

Consider the following deterministic policy for a two-dimensional action space and a one-dimensional state space:
\[
\pi_\theta(s) = \begin{bmatrix}
\theta_1 + \theta_2 s + \theta_3 s^2 \\
\theta_1 + \sin(\theta_4 s) + \cos(\theta_5 s)
\end{bmatrix}
\]

The matrix \( \nabla_{\theta} \pi_\theta(s) \) then has the form:
\[
\nabla_{\theta} \pi_\theta(s) = \begin{bmatrix}
\nabla_{\theta} \pi_\theta(s)|_{a_1} \\
\nabla_{\theta} \pi_\theta(s)|_{a_2}
\end{bmatrix} = \begin{bmatrix}
1 & 1 \\
s & 0 \\
s^2 & 0 \\
0 & \cos(\theta_4 s) s \\
0 & -\sin(\theta_5 s)
\end{bmatrix}
\]

Example 13.1. An example of the Jacobian in the deterministic policy gradient.

As with the other actor-critic methods, we perform gradient descent on \( \ell(\phi) \) and gradient ascent on \( U(\theta) \). For this approach to work in practice, a few additional techniques are needed. One is to generate experiences from a stochastic
policy to allow for better exploration. It is often adequate to simply add zero-mean Gaussian noise to actions generated by our deterministic policy \( \pi_\theta \) as done in algorithm 13.3. To encourage stability when learning \( \theta \) and \( \phi \), we can use experience replay.\(^4\)

An example of this method and the effect of \( \sigma \) on performance is given in example 13.2.

```
struct DeterministicPolicyGradient
    \( \mathcal{P} \) # problem
    b # initial state distribution
    d # depth
    m # number of samples
    \( \nabla \pi \) # gradient of deterministic policy \( \pi(\theta, s) \)
    Q # parameterized value function \( Q(\phi, s, a) \)
    \( \nabla Q \phi \) # gradient of value function with respect to \( \phi \)
    \( \nabla Q a \) # gradient of value function with respect to \( a \)
    \( \sigma \) # policy noise
end

function gradient(M::DeterministicPolicyGradient, \( \pi, \theta, \phi \))
\( \mathcal{P}, b, d, m, \nabla \pi = M.\mathcal{P}, M.b, M.d, M.m, M.\nabla \pi \)
\( Q, \nabla Q \phi, \nabla Q a, \sigma, \gamma = M.Q, M.\nabla Q \phi, M.\nabla Q a, M.\sigma, M.\gamma \)
\( \pi.r\text{and}(s) = \pi(\theta, s) + \sigma.*\text{randn}(\cdot)*I \)
\( \nabla U\theta(\tau) = \text{sum}(\nabla \pi(\theta, s)\nabla Q a(\phi, s, \pi(\theta, s))\gamma^j-1 \text{ for } (j, (s, a, r)) \)
    in enumerate(\( \tau \))
\( \nabla U\phi(\tau, j) = \text{begin} 
    s, a, r = \tau[j]
    s' = \tau[j+1][1]
    a' = \pi(\theta, s')
    \delta = r + \gamma*Q(\phi, s', a') - Q(\phi, s, a)
    \text{return } \delta*(\gamma*\nabla Q a(\phi, s', a') - \nabla Q a(\phi, s, a)) 
\text{end} \)
\( \nabla U\phi(\tau) = \text{sum}(\nabla U\phi(\tau, j) \text{ for } j \text{ in } 1:\text{length}(\tau)-1) \)
\( \text{trajs } = [\text{simulate}(\mathcal{P}, \text{rand}(b), \pi.\text{rand}, d) \text{ for } i \text{ in } 1:m] \)
\( \text{return } \text{mean}(\nabla U\theta(\tau) \text{ for } \tau \text{ in trajs}), \text{mean}(\nabla U\phi(\tau) \text{ for } \tau \text{ in trajs}) \)
end
```

13.4 Actor-Critic with Monte Carlo Tree Search

We can extend concepts from online planning (chapter 22) to the actor-critic setting in which we improve a parameterized policy \( \pi_\theta(a \mid s) \) and a parameterized value function \( U_\phi(s) \).\(^5\) This section discusses the application of Monte Carlo tree search (section 9.6) to learning a stochastic policy with a discrete action space. We


\(^5\) Deterministic policy gradient used \( Q_\phi \), but this approach uses \( U_\phi \) like the other actor-critic methods discussed in this chapter.
Consider applying the deterministic policy gradient algorithm to the simple regulator problem. Suppose we use a simple parameterized deterministic policy \( \pi_{\theta}(s) = \theta_1 \) and the parameterized state-action value function:

\[
Q_{\phi}(s, a) = \phi_1 + \phi_2 s + \phi_3 s^2 + \phi_4 (s + a)^2
\]

Below we plot a progression of the deterministic policy gradient algorithm starting with \( \theta = [0] \) and \( \phi = [0, 1, 0, -1] \) for different values of \( \sigma \). Each iteration was run with 5 rollouts to depth 10 with \( \gamma = 0.9 \).

For this simple problem, the policy quickly converges to optimality nearly irrespective of \( \sigma \). However, if \( \sigma \) is either too small or too large, the value function takes longer to improve. In the case of very small values of \( \sigma \), our policy conducts insufficient exploration from which to effectively learn the value function. For larger values of \( \sigma \) we explore more, but also tend to make poor move choices more frequently.
use our parameterized policy and value function to guide Monte Carlo tree search, and we use the results from Monte Carlo tree search to refine our parameterized policy and value function. As with the other actor critic methods, we apply gradient-based optimization of $\theta$ and $\phi$.

As we perform Monte Carlo tree search, we want to direct our exploration to some extent by our parameterized policy $\pi_\theta(a \mid s)$. One approach is to use an action that maximizes the probabilistic upper confidence bound:

$$a = \arg \max_a Q(s,a) + c \pi_\theta(a \mid s) \frac{\sqrt{N(s)}}{1 + N(s,a)}$$

(13.27)

where $Q(s,a)$ is the action value estimated through the tree search, $N(s,a)$ is the visit count as discussed in section 9.6, and $N(s) = \sum_a N(s,a)$.

After running tree search, we can use the statistics that we collect to obtain $\pi_{MCTS}(a \mid s)$. One way to define this is in terms of the counts:

$$\pi_{MCTS}(a \mid s) \propto N(s,a)^\eta$$

(13.28)

where $\eta \geq 0$ is a hyperparameter that controls the greediness of the policy. If $\eta = 0$, then $\pi_{MCTS}$ will generate actions at random. As $\eta \to \infty$, it will select the action that was selected the most from that state.

In our optimization of $\theta$, we want our model $\pi_\theta$ to match what we obtain through Monte Carlo tree search. One loss function that we can define is the expected cross entropy of $\pi_\theta(\cdot \mid s)$ relative to $\pi_{MCTS}(\cdot \mid s)$:

$$\ell(\theta) = -\mathbb{E}_s \left[ \sum_a \pi_{MCTS}(a \mid s) \log \pi_\theta(a \mid s) \right]$$

(13.29)

where the expectation is over states experienced during the tree exploration. The gradient is

$$\nabla \ell(\theta) = -\mathbb{E}_s \left[ \sum_a \frac{\pi_{MCTS}(a \mid s)}{\pi_\theta(a \mid s)} \nabla_\theta \pi_\theta(a \mid s) \right]$$

(13.30)

To learn $\phi$, we define a loss function in terms of a value function generated during the tree search:

$$U_{MCTS}(s) = \max_a Q(s,a)$$

(13.31)

which is defined at least at the states that we explore during tree search. The loss function aims to make $U_\phi$ agree with the estimates from the tree search:

$$\ell(\phi) = \frac{1}{2} \mathbb{E}_s \left[ (U_\phi(s) - U_{MCTS}(s))^2 \right]$$

(13.32)
The gradient is
\[
\nabla \ell(\phi) = \mathbb{E}_s \left[ \left( U_\phi(s) - U_{\text{MCTS}}(s) \right) \nabla U_\phi(s) \right] \tag{13.33}
\]

Like the actor-critic method in the first section, we need to be able to compute the gradient of our parameterized value function.

After performing some number of Monte Carlo tree search simulations, we update θ by stepping in the direction opposite to \( \nabla \ell(\theta) \) and φ by stepping in the direction opposite to \( \nabla \ell(\phi) \).

13.5 Summary

- In actor-critic methods, an actor attempts to optimize a parameterized policy with the help of a critic that provides a parameterized estimate of the value function.

- Generally, actor-critic methods use gradient-based optimization to learn the parameters of both the policy and value function approximation.

- The basic actor-critic method uses a policy gradient for the actor and minimizes the squared temporal difference residual for the critic.

- The generalized advantage estimate attempts to reduce the variance of its policy gradient at the expense of some bias by accumulating temporal difference residuals across multiple time steps.

- The deterministic policy gradient can be applied to problems with continuous action spaces and uses a deterministic policy actor and an action value critic.

- Online methods such as Monte Carlo tree search can be used to direct the optimization of the policy and value function estimate.

13.6 Exercises

Exercise 13.1. Consider applying Q-Learning with function approximation to a problem with a continuous action space by discretizing the action space. Suppose the continuous action space is in \( \mathbb{R}^n \), such as a robot with \( n \) actuators, and each dimension is discretized into \( m \) intervals. How many actions are in the resulting discrete action space? Is Q-Learning with functional approximation well-suited for use in continuous problems with many dimensions?
Solution: An action space with \( n \) dimensions and \( m \) intervals per dimension results in \( m^n \) discrete actions. The number of discrete actions increases exponentially in \( n \). Even if \( m \) is small, larger values of \( n \) can quickly result in very high action counts. Hence, Q-Learning with function approximation is not well-suited for use on continuous problems with many action dimensions.

Exercise 13.2. Would the actor-critic method with Monte Carlo tree search as presented in section 13.4 be a good method for solving the cart-pole problem (appendix F.3)?

Solution: The Monte Carlo tree search expands a tree based on visited states. The cart-pole problem has a continuous state space, leading to a search tree with an infinite branching factor. Use of this algorithm would require adjusting the problem, such as discretizing the state space.

Exercise 13.3. In the following expressions of advantage functions, determine which ones are correct and explain what they are referring to:

\[
\begin{align*}
(a) & \quad \mathbb{E}_{r,s'} \left[ r + \gamma U^{\pi_\theta}(s) - U^{\pi_\theta}(s') \right] \\
(b) & \quad \mathbb{E}_{r,s'} \left[ r + \gamma U^{\pi_\theta}(s') - U^{\pi_\theta}(s) \right] \\
(c) & \quad \mathbb{E}_{r_{1:d},s'} \left[ -U^{\pi_\theta}(s) + \gamma^k U^{\pi_\theta}(s') + \sum_{\ell=1}^{k} \gamma^{l-1} r_l \right] \\
(d) & \quad \mathbb{E}_{r_{1:d},s'} \left[ -U^{\pi_\theta}(s) + \gamma U^{\pi_\theta}(s') + \sum_{\ell=1}^{k} \gamma^{l-1} r_l \right] \\
(e) & \quad \mathbb{E} \left[ -U^{\pi_\theta}(s) + \sum_{\ell=1}^{d} \gamma^{l-1} r_l \right] \\
(f) & \quad \mathbb{E} \left[ -\gamma U^{\pi_\theta}(s') + \sum_{\ell=1}^{d+1} \gamma^{l-1} r_l \right] \\
(g) & \quad \mathbb{E} \left[ \sum_{\ell=1}^{k} \gamma^{l-1} \delta_{l-1} \right] \\
(h) & \quad \mathbb{E} \left[ \sum_{\ell=1}^{k} \gamma^{l-1} \delta_l \right] \\
(i) & \quad \mathbb{E} \left[ \sum_{k=1}^{\infty} (\gamma \lambda)^{k-1} \delta_k \right] \\
(j) & \quad \mathbb{E} \left[ \sum_{k=1}^{\infty} (\lambda)^{k-1} \delta_k \right]
\end{align*}
\]

Solution: The following table lists the correct expressions:
Exercise 13.4. What are the benefits of using a temporal difference residual over a sequence of rollouts reward and vice-versa?

Solution: Approximation using a temporal difference residual is more computationally efficient than using a sequence of rollouts. Temporal difference residual approximation has low variance but high bias due to using the critic value function $U_\phi$ as an approximator of the true value function $U^\pi_\theta$. On the other hand, rollout approximation has high variance but is unbiased. Obtaining an accurate estimate using a temporal difference residual approximation typically requires far fewer samples than when using a rollout approximation, at the cost of introducing bias into our estimate.

Exercise 13.5. Consider the action value function given in example 13.2, $Q_\phi(s,a) = \phi_1 + \phi_2 s + \phi_3 s^2 + \phi_4 (s + a)^2$. Calculate the gradients required for the deterministic policy gradient approach.

Solution: We need to calculate two gradients. For the actor we need to compute $\nabla_\phi Q_\phi(s,a)$, while for the critic we need to compute $\nabla_a Q_\phi(s,a)$.

\[
\nabla_\phi Q(s,a) = \begin{bmatrix} 1, s, s^2, (s + a)^2 \end{bmatrix}
\]
\[
\nabla_a Q(s,a) = 2\phi_4(s + a)
\]
14 Policy Validation

The methods presented in the earlier chapters show how to construct an optimal or approximately optimal solution with respect to a particular model of the dynamics and reward. However, before deploying a decision making system in the real world, it is generally desirable to validate in simulation that the behavior of the resulting policy is consistent with what is actually desired. This chapter discusses various analytical tools for validating decision strategies.¹ We will start by discussing how to go about evaluating different performance metrics. Accurately computing such metrics can be computationally challenging, especially when they pertain to rare events such as failures. We will discuss methods such as importance sampling that can help address computational efficiency. An important concern is that our systems be robust to differences between the models that we use for analysis and the real world. This chapter suggests methods for analyzing the robustness of our policies. Fundamental to the design of many decision making systems is the tradeoff between multiple objectives, and we will outline ways of analyzing these tradeoffs. The chapter concludes with a discussion of adversarial analysis, which can be used for finding the most likely failure trajectory for a given system.

14.1 Performance Metric Evaluation

Once we have a policy, we are often interested in evaluating it with respect to different performance metrics. For example, suppose we constructed a collision avoidance system—either through some form of optimization of a scalar reward function or just heuristically as discussed in example 14.1—and we want to assess its safety by computing the probability of collision when following our policy. Or, if we created a policy for constructing investment portfolios, we might be interested

in understanding the probability that our policy will result in an extreme loss or what the expected return may be.

For the moment, we will consider a single metric $f$ evaluated on a policy $\pi$. Often, this metric is defined as the expectation of a metric $f_{\text{traj}}$ evaluated on trajectories $\tau = (s_1, a_1, \ldots)$:

$$f(\pi) = \mathbb{E}_\tau[f_{\text{traj}}(\tau)] \quad (14.1)$$

This expectation is over the trajectory distribution. In order to define a trajectory distribution associated with an MDP, we need to specify an initial state distribution $b$. We have the probability of generating a trajectory $\tau$ as

$$P(\tau) = P(s_1, a_1, \ldots) = b(s_1) \prod_t T(s_{t+1} | s_t, a_t) \quad (14.2)$$

In the collision avoidance context, $f_{\text{traj}}$ may be 1 if the trajectory led to a collision and 0 otherwise. The expectation would correspond to the collision probability.

In some cases, we are interested in studying the distribution over the output of $f_{\text{traj}}$. Figure 14.1 shows an example of such a distribution. The expectation in equation (14.1) is just one of many different ways to convert a distribution over trajectory metrics to a single value. We will focus primarily on this expectation in our discussion, but examples of other transformations of the distribution to a value include variance, fifth percentile, and mean of the values below the fifth percentile.\(^2\)

The trajectory metric can sometimes be written in this form:

$$f_{\text{traj}}(\tau) = f_{\text{traj}}(s_1, a_1, \ldots) = \sum_t f_{\text{step}}(s_t, a_t) \quad (14.3)$$

where $f_{\text{step}}$ is some function that depends on the current state and action, much like the reward function in MDPs. If $f(\pi)$ is defined as the expectation of $f_{\text{traj}}$, the objective is the same as when solving an MDP, where $f_{\text{step}}$ is simply the reward function. It turns out that we can use the policy evaluation algorithms introduced in section 7.2 to evaluate our policy with respect to any performance metric of the form in equation (14.3).

Policy evaluation will output a value function that is a function of the state,\(^3\) corresponding to the expected value of the performance metric when starting from that state. Example 14.2 shows slices of this value function for the collision avoidance problem. The overall performance is given by

$$f(\pi) = \sum_b f_{\text{state}}(s)b(s) \quad (14.4)$$

\(^2\) Various risk measures have been discussed in the literature. An overview of some of them that have been used in the context of Markov decision processes are provided by A. Ruszczyński, “Risk-Averse Dynamic Programming for Markov Decision Processes,” Mathematical Programming, vol. 125, no. 2, pp. 235–261, 2010.

\(^3\) We used $U^\pi$ to represent the value function associated with policy $\pi$ in previous chapters.
In the aircraft collision avoidance problem, we need to decide when to issue a climb or descend advisory to our aircraft to avoid an intruder aircraft. The intruder is approaching us head-on with constant horizontal closing speed. The state is specified by the altitude $h$ of our aircraft measured relative to the intruder aircraft, our vertical rate $\dot{h}$, the previous action $a_{\text{prev}}$, and the time to potential collision $t_{\text{col}}$. There is a penalty of 1 when there is a collision, defined as when the intruder comes within 50 m when $t_{\text{col}} = 0$. Additionally, there is a penalty of 0.01 when $a \neq a_{\text{prev}}$.

We can use dynamic programming with linear interpolation (section 8.4) to derive an optimal policy. Alternatively, we can define a simple heuristic policy parameterized by thresholds on $t_{\text{col}}$ and $h$ that works as follows. If $|h| < h_{\text{thresh}}$ and $t_{\text{col}} < t_{\text{thresh}}$ then an advisory is generated. This advisory is a climb if $h > 0$ and a descend otherwise. By default, we use $h_{\text{thresh}} = 50$ m and $t_{\text{thresh}} = 30$ s. Below are plots of both the optimal and simple policies for two slices through the state space.

Example 14.1. Optimal and simple collision avoidance policies. Additional details of the problem are in appendix F.6.
Below is the result of applying policy evaluation to both an optimal policy and the simple policy introduced in example 14.1. Each point in the plot corresponds to the value of the metric conditioned on starting from the associated state. We define $f_{\text{state}}(s, a) = 1$ if $s$ is a collision and 0 otherwise. This plot shows where in the state space there is significant collision risk, indicated by “hotter” colors, when following the policy. We can see that the optimal policy is quite safe, especially if $t_{\text{col}} > 20$ s. When $t_{\text{col}}$ is low, even the optimal policy cannot avoid collision due to the physical acceleration constraints of the vehicle. The simple policy has a much higher level of risk associated with it compared to the optimal policy, especially when $t_{\text{col}} > 20$ s, $\dot{h} = 5$ m/s, and the intruder is below us—in part because the choice to produce an advisory in the simple strategy does not take $\dot{h}$ into account.

Example 14.2. Probability of collision when following the optimal and simple collision avoidance policies.
where \( f_{\text{state}} \) is the value function obtained through policy evaluation.

If the state space is discrete, then equation (14.4) can be computed analytically. However, if the state space is large or continuous, we may want to estimate \( f(\pi) \) through sampling. We can pull a sample from the initial state distribution and then rollout the policy and compute the trajectory metric. We can then estimate the value of the overall metric from the mean of the trajectory metrics. The quality of the estimate generally improves with more samples. Example 14.3 illustrates this process for estimating different metrics associated with collision avoidance policies.

We often use the standard error to measure the quality of our estimate, which is defined as

\[
\text{SE} = \frac{\hat{\sigma}}{\sqrt{n}}
\]

where \( \hat{\sigma} \) is the standard deviation of our samples and \( n \) is the number of samples. In example 14.3, our the standard deviation of our collision metric is 0.0173, making the standard error of our collision probability metric 0.000173.

We can convert the standard error to a confidence interval. For example, a 95% confidence interval would be \( \hat{\mu} \pm 1.96\text{SE} \), where \( \hat{\mu} \) is the mean of our samples. For our collision avoidance example, this interval is \((-3.94 \times 10^{-5}, 6.39 \times 10^{-4})\). Alternatively, we can take a Bayesian approach and represent our posterior as a beta distribution as discussed in section 4.2.

For small probabilities, such as failure probabilities in a relatively safe system, we are often interested in the relative standard error, which is given by

\[
\frac{\hat{\sigma}}{\hat{\mu} \sqrt{n}}
\]

This is equivalent to dividing the standard error by the mean. In our collision avoidance problem, our relative error is 0.578. Although the absolute error might be small, the relative error is quite high since we are trying to estimate a small probability.

### 14.2 Rare Event Simulation

As we see in example 14.3, we may need many samples to accurately estimate metrics where rare events are very influential, such as estimating collision probability. In the collision avoidance example, there were only three collisions out of our
We want to estimate the probability of collision and the probability of generating an advisory. Here, we will consider the optimal and simple policies introduced in example 14.1. To evaluate these metrics, we use $10^4$ samples from the initial state distribution used in figure 14.1 and then perform rollouts. The plots below show the convergence curves.

What we can see is that the optimal policy is much safer than the simple policy while producing advisories at approximately the same frequency. The advisory metric estimate converges much more quickly than the collision estimates. The reason for the faster convergence for the advisory metric is that advisories are more common than collisions. Collisions involving the optimal policy are so rare that even $10^4$ samples appear inadequate for an accurate estimate. The curve is very jagged, with large spikes at samples involving collisions, followed by a decay in the collision probability estimate as collision-free samples are simulated.
10^4 samples, as indicated by the three spikes in the plot. When we are designing algorithms for high-stakes systems, such as systems that trade our money or drive our cars, accurately estimating failure probabilities through direct sampling and simulation can be computationally challenging.

A common approach to improve efficiency is called importance sampling, which involves sampling from an alternative distribution and weighting the results appropriately to arrive at an unbiased estimate. We used this same kind of approach in the context of inference in Bayesian networks by the name of likelihood weighted sampling (section 3.7). The alternative sampling distribution is often called a proposal distribution, and we will use $P'(\tau)$ to represent the probability our proposal distribution assigns to trajectory $\tau$.

We will derive the appropriate way to weight samples from $P'$. If we have $\tau^{(1)}, \ldots, \tau^{(n)}$ drawn from the true distribution $P$, then we have

$$f(\tau) = \mathbb{E}_h[f_{\text{traj}}(\tau)]$$
$$= \sum_{\tau} f_{\text{traj}}(\tau) P(\tau)$$
$$\approx \sum_i f_{\text{traj}}(\tau^{(i)}) \text{ with } \tau^{(i)} \sim P$$

We can multiply equation (14.8) by $P'(\tau) / P'(\tau)$ and obtain the following:

$$f(\tau) = \sum_h f_{\text{traj}}(\tau) P(\tau) \frac{P'(\tau)}{P'(\tau)}$$
$$= \sum_{\tau} f_{\text{traj}}(\tau) P'(\tau) \frac{P(\tau)}{P'(\tau)}$$
$$\approx \sum_i f_{\text{traj}}(\tau^{(i)}) \frac{P(\tau^{(i)})}{P'(\tau^{(i)})} \text{ with } \tau^{(i)} \sim P'$$

In other words, we need to weight the outcomes of the samples from the proposal distribution, where the weight given to sample $i$ is $P(\tau^{(i)}) / P'(\tau^{(i)})$.

We want to choose the proposal distribution $P'$ to focus the generation of samples on those that are "important," in the sense that they are more likely to contribute to the overall performance estimate. In the case of collision avoidance, we will want this proposal distribution to encourage collisions so that we have more than just a few collision situations to estimate collision risk. However, we do not want all of our samples to result in collision. In general, assuming that the

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\[2021-03-07 23:12:00-08:00, revision 80bf4d6, comments to bugs@algorithmsbook.com\]
space of histories is discrete, the optimal proposal distribution is

\[ P^*(\tau) = \frac{|f_{\text{traj}}(\tau)|P(\tau)}{\sum_{\tau'}|f_{\text{traj}}(\tau')|P(\tau')} \]  

(14.13)

If \( f_{\text{traj}} \) is non-negative, then the denominator is exactly the same as the metric we are trying to estimate in equation (14.1).

Although equation (14.13) is generally not practical to compute exactly (this is why we are using importance sampling in the first place), it can provide some intuition as to how to use our domain expertise to construct a proposal distribution. It is common to bias the initial state distribution or the transition model slightly towards more important trajectories, such as towards collision.

To illustrate the construction of an importance distribution, we will use the optimal policy for the collision avoidance problem in example 14.1. Instead of starting at \( t_{\text{col}} = 40 \) s, we will start the aircraft closer with \( t_{\text{col}} = 20 \) s to make the collision avoidance problem more challenging. The true distribution has \( h \sim \mathcal{U}(-10, 10) \) (m) and \( \dot{h} \sim \mathcal{U}(-200, 200) \) (m/s). However, certain combinations of \( h \) and \( \dot{h} \) are more challenging for the optimal policy to resolve. We used dynamic programming on a discrete version of the problem to determine the probability of collision for different values for \( h \) and \( \dot{h} \). We can take these results and normalize them to turn them into the proposal distribution shown in figure 14.2.

Using the proposal distribution shown in figure 14.2 results in better estimates of the collision probability than direct sampling with the same number of samples. Figure 14.3 shows the convergence curves. By \( 5 \times 10^4 \) samples, both sampling methods converge to the same estimate. However, importance sampling converges closely to the true value within \( 10^4 \) samples. Using our proposal distribution, importance sampling generated 939 collisions, while direct sampling only generated 246. Even more collisions could be generated if we also biased the transition distribution, rather than solely the initial state distribution.

14.3 Robustness Analysis

Before deploying a system in the real world, it is important to study its robustness to modeling errors. We can use the tools mentioned in the previous sections, such as policy evaluation and importance sampling, but evaluate our policies on environments that deviate from the model assumed when optimizing the policy. Figure 14.4 shows how performance varies as the true model deviates from the
one used for optimization. We can also study the sensitivity of our metrics to modeling assumptions over the state space (example 14.4). If performance on the relevant metrics appears to be preserved under plausible perturbations of the environment model, then we can have greater confidence that our system will behave as planned when deployed.

We typically want our planning model, the model we use for optimizing our policies, to be relatively simple to prevent overfitting to potentially erroneous modeling assumptions that are not representative of the real world. A side benefit of simpler planning models is that they can make planning more computationally efficient. However, our evaluation model can be as complex as we can justify. For example, we may use a simple, low-dimensional, discrete model of aircraft dynamics when generating a collision avoidance policy, but then evaluate that policy in a continuous, high fidelity simulation. A simpler planning model is often more robust to perturbations in the evaluation model.

The process of evaluating our policies on a variety of evaluation models is sometimes referred to as stress testing, especially if the spectrum of evaluation models includes fairly extreme scenarios. In collision avoidance, extreme scenarios might include those where the aircraft are converging upon each other with extreme climb rates that may not be physically achievable. Understanding what categories of scenarios can lead to system failure can be useful during the design phase, even if we choose not to optimize the behavior of the system for these scenarios because they are deemed unrealistic.

If we find that our policies are overly sensitive to our modeling assumptions, we may consider using a method known as robust dynamic programming. Instead of committing to a particular transition model, we have a suite of transition models $T_{1:n}$ and reward models $R_{1:n}$. We can revise the Bellman update equation from equation (7.16) to provide robustness to different models:

$$U_{k+1}(s) = \max_a \min_i \left( R_i(s,a) + \gamma \sum_{s'} T_i(s' \mid s,a) U_k(s') \right)$$  \hspace{1cm} (14.14)

The update uses the action that maximizes expected utility when using the model that minimizes our utility.

---

We can plot collision probability when starting from different initial states, similar to example 14.2. Below, we use a policy optimized for the parameters in appendix F.6, but we vary the limit \( \ddot{h}_{\text{limit}} \) in the evaluation model.

We optimized the policy with \( \ddot{h}_{\text{limit}} = 1 \text{ m/s}^2 \). If it was actually \( 0.25 \text{ m/s}^2 \), then the policy performs poorly in some states since it takes longer to achieve a target vertical rate. If the limit was \( 1.25 \text{ m/s}^2 \), we are a bit safer.

Example 14.4. Probability of collision when following the optimal collision avoidance policies when there is a mismatch between the model used for planning and the model used for evaluation.
14.4 Trade Analysis

Many interesting tasks involve multiple, often competing, objectives. For autonomous systems, there is often a tradeoff between safety and efficiency. In designing a collision avoidance system, we want to be very safe without making too many unnecessary avoidance maneuvers. A trade analysis studies how the various performance metrics are traded as the design parameters are changed.

If we consider only two performance metrics, we can plot a tradeoff curve like the one discussed in example 14.5. By varying parameters in the policy, we obtain different values for the two metrics. These curves are useful when comparing different methodologies for generating policies. For example, the curves in example 14.5 suggests that a dynamic programming approach to generating policies can bring significant benefit over simple threshold-based policies—at least in the way we defined them.

For each of the curves in example 14.5, we vary only one parameter at a time, but to arrive at a satisfactory system, we may need to study the effects of varying multiple parameters. As we vary multiple parameters, we obtain a space of possible policies. Some of those policies may perform worse on all performance metrics relative to at least one other policy in that space. We can often eliminate from consideration those policies that are dominated by others. A policy is called Pareto optimal or Pareto efficient if it is not dominated by any other policy in that space. The set of Pareto optimal policies is called the Pareto frontier or (in two dimensions) the Pareto curve. Figure 14.5 shows an example Pareto curve.

14.5 Adversarial Analysis

It can be useful to study the robustness of a policy from the perspective of an adversarial analysis. At each time step, an adversary selects the state that results from applying the action specified by the policy from the current state. The adversary has two objectives to balance: minimizing our return and maximizing the likelihood of the resulting trajectory according to our transition model. We can transform our original problem into an adversarial problem. The adversarial state space is the same as in the original problem, but the adversarial action space is the state space of the original problem. The adversarial reward is

\[ R'(s, a) = -R(s, \pi(s)) + \lambda \log(T(a \mid s, \pi(s))) \]  

(14.15)
In our aircraft collision avoidance problem, we must balance safety in terms of collision probability with other metrics, such as the expected number of advisory changes. Both of these can be implemented using trajectory metrics that are additively decomposed by steps as done in equation (14.3), allowing us to compute them using exact policy evaluation.

The plot below shows three curves associated with different parameterized versions of the simple and optimal policies. The first curve shows the performance of the simple policy on the two metrics as the $h_{\text{thresh}}$ parameter (defined in example 14.1) is varied. The second curve shows the performance of the simple policy as $t_{\text{thresh}}$ is varied. The third curve shows the optimal policy as the parameter $\theta$ is varied, where the cost of collision is $-\theta$ and the cost of changing advisories is $-(1 - \theta)$.

We can see that the optimal policy dominates the curves generated by the parameterized simple policies. When $\theta$ is close to 1, then we are very safe, but we have to tolerate more advisory changes. As $\theta$ goes to 0, we are less safe but do not produce advisories. Given a particular threshold level of safety, we are able to create an optimized policy that has fewer advisory changes in expectation than either of the simple parametric policies.

Example 14.5. An analysis of the tradeoff between safety and operational efficiency when varying parameters of different collision avoidance systems.
where $\pi$ is our policy, $R$ is our original reward function, $T$ is our original transition model, and $\lambda \geq 0$ is a parameter that controls the importance of maximizing the resulting likelihood of the trajectory. Since an adversary attempts to maximize the sum of adversarial reward, it is maximizing our expected negative return plus $\lambda$ times the log-probability of the resulting trajectory. The adversarial transition model is deterministic; the state transitions to exactly what the adversary specifies as its action.

Algorithm 14.1 implements this conversion to an adversarial problem. It assumes a discrete state and action space, which can then be solved using one of the dynamic programming algorithms in chapter 7. The solution is an adversarial policy that maps states to states. Given an initial state, we can generate a trajectory that minimizes our reward given some level of probability. Since the problem is deterministic, it is actually a search problem, and any of the algorithms in appendix E can be used. If our problem is high-dimensional or continuous, we may use one of the approximate solution techniques discussed in chapters 8 and 9.

Algorithm 14.1. Conversion to an adversarial problem given a policy $\pi$. An adversarial agent tries to change the outcomes of our policy actions so as to optimize the dual objectives of minimizing our original utility and maximizing the likelihood of the trajectory. The parameter $\lambda$ controls how important it is to maximize the likelihood of the resulting trajectory. It returns an MDP whose transition and reward models are represented as matrices.

```
function adversarial(𝒫::MDP, π, λ)
    𝒮, 𝒴, 𝒯, 𝒴 = 𝒫.𝒮, 𝒫.𝒴, 𝒫.𝒯, 𝒫.𝒲, 𝒫.𝒲
    𝒮′ = 𝒮
    𝒫′ = zeros(length(𝒮′), length(𝒴′))
    𝒯′ = zeros(length(𝒮′), length(𝒴′), length(𝒮′))
    for s in 𝒮′
        for a in 𝒴′
            𝒫′[s, a] = -R(s, π(s)) + λ*log(𝒯(s, π(s), a))
            𝒯′[s, a, a] = 1
        end
    end
    return MDP(𝒯′, 𝒫′, 𝒲)
end
```

Sometimes we are interested in finding the most likely failure associated with a policy for some definition of failure. In some problems, failure can be defined as entering a particular state. For example, a collision may be considered a failure in our collision avoidance problem. Other problems may require a more complicated definition of failure that goes beyond just entering some subset of the state space. For example, we may want to specify failure using a temporal logic, which is a way to represent and reason about propositions qualified in terms of time. In many cases, however, we can use these failure specifications to create an augmented state space that we can then solve.\footnote{M. Bouton, J. Tumova, and M. J. Kochenderfer, “Point-Based Methods for Model Checking in Partially Observable Markov Decision Processes,” in AAAI Conference on Artificial Intelligence (AAAI), 2020.}
With a definition of failure states, we can solve for the most likely failure trajectory by changing the reward function in equation (14.15) to:

\[
R'(s, a) = \begin{cases} 
-\infty & \text{if } s \text{ is terminal and not a failure} \\
0 & \text{if } s \text{ is terminal and a failure} \\
\log(T(a \mid s, \pi(s))) & \text{otherwise}
\end{cases}
\] (14.16)

We can find these most likely failures using a variety of approximation methods. Depending on the approximation method, it may be important to relax the infinite penalty for not reaching a failure at termination so that the search can be guided to failures. If applying Monte Carlo tree search to collision avoidance, the penalty could be related to the miss distance.\(^9\)

We can play back the most likely failure trajectory and gauge whether that trajectory merits concern. If the trajectory is deemed extremely implausible, then we can feel more confident that our policy is safe. If the failure trajectory does merit concern, then we might have a few options:

1. **Change the action space.** We may add more extreme maneuvers to our action set for our collision avoidance problem.

2. **Change the reward function.** We may decrease the cost for changing advisories with the aim of lowering collision risk, as illustrated in the tradeoff curve in example 14.5.

3. **Change the transition function.** We may increase the acceleration limit so that the aircraft can achieve the target vertical rates more quickly when directed by our policy.

4. **Improve the solver.** We may have used a discretization of the state space that is too coarse to capture important features of the optimal policy. In exchange for additional computation time, we may be able to refine the discretization to obtain a better policy. Alternatively, we may adopt a different approximation technique.

5. **Do not deploy the system.** If the policy is unsafe, it may be better to not deploy it in the real world.

14.6 Summary

- Performance metrics for policies may be evaluated using the dynamic programming techniques discussed in earlier chapters or through sampling rollouts.

- We can assess our confidence in our performance metric evaluations using standard error, confidence intervals, or one of the Bayesian approaches discussed earlier.

- Estimating the probability of rare events can be done more efficiently using a method called importance sampling.

- Importance sampling involves sampling from an alternative distribution and weighting the results appropriately.

- Due to the fact that the model used for optimization may not be an accurate representation of the real world, it is important to study the sensitivity of our policy to modeling assumptions.

- Robust dynamic programming can help improve robustness to model uncertainty by optimizing with respect to a set of different transition and reward models.

- Trade analyses can help us determine how to balance multiple performance objectives when optimizing a policy.

- Adversarial analyses involve an adversary that chooses the state to which we transition at each step so as to minimize our objective while maximizing the likelihood of the trajectory.

14.7 Exercises

Exercise 14.1. We have a trajectory $\tau$ with

<table>
<thead>
<tr>
<th>$s_1$</th>
<th>$a_1$</th>
<th>$s_2$</th>
<th>$a_2$</th>
<th>$s_3$</th>
</tr>
</thead>
<tbody>
<tr>
<td>6.0</td>
<td>2.2</td>
<td>1.4</td>
<td>0.7</td>
<td>6.0</td>
</tr>
</tbody>
</table>

Our dynamics are linear Gaussian with $T(s' \mid s, a) = \mathcal{N}(s' \mid 2s + a, 5^2)$, and our initial state distribution is given by $\mathcal{N}(5, 6^2)$. What is the log-likelihood of the trajectory $\tau$?

**Solution:** The log-likelihood of the trajectory is

$$\log \mathcal{N}(6.0 \mid 5, 6^2) + \log \mathcal{N}(1.4 \mid 2 \cdot 6.0 + 2.2, 5^2) + \log \mathcal{N}(6.0 \mid 2 \cdot 1.4 + 0.7, 5^2) \approx -11.183$$
Exercise 14.2. We ran a million simulations and found that our collision avoidance system resulted in 10 collisions. What is our collision probability estimate and the relative standard error?

**Solution:** The collision probability estimate is

\[ \hat{\mu} = \frac{10}{10^6} = 10^{-5} \]

The \( i \)th sample \( x_i \) is 1 if there is a collision and 0 otherwise. The standard deviation is

\[ \hat{\sigma} = \sqrt{\frac{1}{10^6 - 1} \sum_{i=1}^{n} (x_i - \hat{\mu})^2} = \sqrt{\frac{1}{10^6 - 1} \left( 10(1 - \hat{\mu})^2 + (10^6 - 10)\hat{\mu}^2 \right)} \approx 0.00316 \]

The relative error is

\[ \frac{\hat{\sigma}}{\hat{\mu} \sqrt{n}} \approx \frac{0.00316}{10^{-5} \sqrt{10^6}} = 0.316 \]

Exercise 14.3. We want to compute the expectation \( E_{x \sim U(0,5)}[f(x)] \), where \( f(x) = -1 \) if \( |x| \leq 1 \) and 0 otherwise. What is the optimal proposal distribution?

**Solution:** The optimal proposal distribution is

\[ p^*(x) = \frac{|f(x)|p(x)}{\int |f(x)|p(x) \, dx} \]

which is equivalent to \( U(0,1) \) because \( f(x) \) is only non-zero for \( x \in [-1,1] \), \( U(0,5) \) only has support for \( x \in [0,5] \), and both \( f(x) \) and \( p(x) \) produce constant values when non-zero.

Exercise 14.4. Suppose we draw the sample 0.3 from the proposal distribution in the previous exercise. What is its weight? What is the estimate of \( E_{x \sim U(0,5)}[f(x)] \)?

**Solution:** The weight is \( p(x) / p^*(x) = 0.2 / 1 \). Since \( f(0.3) = -1 \), the estimate is \(-0.2\), which is the exact answer.

Exercise 14.5. Suppose we have the following four policies that have been evaluated on three metrics that we want to maximize:

<table>
<thead>
<tr>
<th>System</th>
<th>( f_1 )</th>
<th>( f_2 )</th>
<th>( f_3 )</th>
</tr>
</thead>
<tbody>
<tr>
<td>( \pi_1 )</td>
<td>2.7</td>
<td>1.1</td>
<td>2.8</td>
</tr>
<tr>
<td>( \pi_2 )</td>
<td>1.8</td>
<td>2.8</td>
<td>4.5</td>
</tr>
<tr>
<td>( \pi_3 )</td>
<td>9.0</td>
<td>4.5</td>
<td>2.3</td>
</tr>
<tr>
<td>( \pi_4 )</td>
<td>5.3</td>
<td>6.0</td>
<td>2.8</td>
</tr>
</tbody>
</table>

Which policies are on the Pareto frontier?

**Solution:** Only \( \pi_1 \) is dominated by other policies. Hence, \( \pi_2 \), \( \pi_3 \), and \( \pi_4 \) are on the Pareto frontier.
PART III:

MODEL UNCERTAINTY

In our discussion of sequential decision problems, we have assumed that the transition and reward models are known. In many problems, however, these models are not known exactly, and the agent must learn to act through experience. By observing the outcomes of its actions in the form of state transitions and rewards, the agent is to choose actions that maximize its long-term accumulation of rewards. Solving such problems in which there is model uncertainty is the subject of the field of reinforcement learning and the focus of this part of the book. We will discuss several challenges in addressing model uncertainty. First, the agent must carefully balance exploration of the environment with the exploitation of that knowledge gained through experience. Second, rewards may be received long after the important decisions have been made, so credit for later rewards must be assigned to earlier decisions. Third, the agent must generalize from limited experience. We will review the theory and some of the key algorithms for addressing these challenges.
15 Exploration and Exploitation

Reinforcement learning agents\(^1\) must balance exploration of the environment with exploitation of knowledge obtained through its interactions.\(^2\) Pure exploration will allow the agent to build a comprehensive model, but the agent will likely have to sacrifice the gathering of reward. Pure exploitation has the agent continually choosing the action it thinks best to accumulate reward, but there may be other, better actions that could be taken. This chapter introduces the challenges associated with the exploration-exploitation tradeoff by focusing on a problem with a single state. We conclude with introducing exploration in MDPs with multiple states.

15.1 Bandit Problems

Early analyses of the exploration-exploitation tradeoff were focused on slot machines, also called one-armed bandits.\(^3\) The name comes from older slot machines having a single pull lever and that the machine tends to take the gambler’s money. Many real world problems can be framed as multi-armed bandit problems,\(^4\) such as the allocation of clinical trials and adaptive network routing. Many different bandit problem formulations exist in the literature, but this chapter will focus on what is called a binary bandit, Bernoulli bandit, or binomial bandit. In these problems, arm \(a\) pays off 1 with probability \(\theta_a\) and 0 otherwise. Pulling an arm costs nothing, but we only have \(h\) pulls.

A bandit problem can be framed as an \(h\)-step Markov decision process with a single state, \(n\) actions, and an unknown, stochastic reward function \(R(s, a)\), as shown in figure 15.1. Recall that \(R(s, a)\) is the expected reward when taking action \(a\) in \(s\), but individual rewards realized in the environment may come from a probability distribution.


\(^{2}\)In some applications, we want to optimize a policy given a fixed set of trajectories. This context is known as batch reinforcement learning. This chapter assumes we have to collect our own data through interaction, which makes choosing an appropriate exploration strategy important.

\(^{3}\)These bandit problems were explored during World War II and proved exceptionally challenging to solve. According to Peter Whittle, “efforts to solve [bandit problems] so sapped the energies and minds of Allied analysts that the suggestion was made that the problem be dropped over Germany as the ultimate instrument of intellectual sabotage.” J.C. Gittins, “Bandit Processes and Dynamic Allocation Indices,” Journal of the Royal Statistical Society, Series B (Methodological), vol. 41, no. 2, pp. 148–177, 1979.

Algorithm 15.1 defines the simulation loop for a bandit problem. At each step, we evaluate our exploration policy $\pi$ on our current model of the payoff probabilities to generate an action $a$. The next section will discuss a way to model payoff probabilities, and the remainder of the chapter will outline several different exploration strategies. After obtaining $a$, we simulate a pull of that arm, returning binary reward $r$. The model is then updated using the observed $a$ and $r$. The simulation loop is repeated to horizon $h$.

```
struct BanditProblem
    θ # vector of payoff probabilities
    R # reward sampler
end

function BanditProblem(θ)
    R(a) = rand() < θ[a] ? 1 : 0
    return BanditProblem(θ, R)
end

function simulate(P::BanditProblem, model, π, h)
    for i in 1:h
        a = π(model)
        r = P.R(a)
        update!(model, a, r)
    end
end
```

Algorithm 15.1. Simulation of a bandit problem. A bandit problem is defined by a vector $\theta$ of payoff probabilities, one per action. We also define a function $R$ that simulates the generation of a stochastic binary reward in response to the selection of an action. Each step of a simulation involves generating an action $a$ from the exploration policy $\pi$. The exploration policy generally consults the model in the selection of the action. The selection of that action results in a randomly generated reward, which is then used to update the model. Simulations are run to horizon $h$. 

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15.2 Bayesian Model Estimation

We would like to track our belief over the win probability $\theta_a$ for arm $a$. The beta distribution (section 4.2) is often used for representing such a belief. Assuming a uniform prior of $\text{Beta}(1, 1)$, the posterior for $\theta_a$ after $w_a$ wins and $\ell_a$ losses is $\text{Beta}(w_a + 1, \ell_a + 1)$. The posterior probability of winning is

$$\rho_a = P(\text{win}_a \mid w_a, \ell_a) = \int_0^1 \theta \times \text{Beta}(\theta \mid w_a + 1, \ell_a + 1) \, d\theta = \frac{w_a + 1}{w_a + \ell_a + 2} \quad (15.1)$$

Algorithm 15.2 provides an implementation. Example 15.1 illustrates how to compute these posterior distributions from counts of wins and losses.

```plaintext
struct BanditModel
    B # vector of beta distributions
end

function update!(model::BanditModel, a, r)
    α, β = StatsBase.params(model.B[a])
    model.B[a] = Beta(α + r, β + (1-r))
return model
end
```

A greedy action is one that maximizes our expected immediate reward—or, in other words, the posterior probability of winning in the context of our binary bandit problem. There may be multiple greedy actions. We do not always want to select a greedy action because we may miss out on discovering another action that may actually provide higher reward in expectation. We can use the information from the beta distributions associated with the different actions to drive our exploration of non-greedy actions.

15.3 Undirected Exploration Strategies

There are several ad hoc exploration strategies that are commonly used to balance exploration with exploitation. This section discusses a type of ad hoc exploration called undirected exploration, where we do not use information from previous outcomes to guide exploration of non-greedy actions.
Suppose we have a two-armed bandit that we have played six times. The first arm has 1 win and 0 losses, and the other arm has 4 wins and 1 loss. Assuming a uniform prior, the posterior distribution for $\theta_1$ is Beta(2, 1) and the posterior distribution for $\theta_2$ is Beta(5, 2).

These posteriors assign non-zero likelihood to win probabilities between 0 and 1. The density at 0 is 0 for both arms because they both received at least one win. Similarly, the density at 1 for arm 2 is 0 because it received at least one loss. The payoff probabilities $\rho_1 = 2/3$ and $\rho_2 = 5/7$ are shown above with vertical lines. We believe that the second arm has the best chance of producing a payout.
One of the most common undirected exploration strategies is \( \epsilon \)-greedy exploration (algorithm 15.3). This strategy chooses a random arm with probability \( \epsilon \). Otherwise, we choose a greedy arm, \( \arg \max_a \rho_a \). This \( \rho_a \) is the posterior probability of a win with action \( a \) using the Bayesian model in the previous section. Alternatively, we can use the maximum likelihood estimate, but with enough pulls, the difference between the two approaches is small. Larger values of \( \epsilon \) lead to more exploration, thereby resulting in faster identification of the best arm, but more pulls are wasted on suboptimal arms. Example 15.2 demonstrates this exploration strategy and the evolution of our beliefs.

The \( \epsilon \)-greedy method maintains a constant amount of exploration, despite there being far more uncertainty early in the interaction with the bandit than later. One common adjustment is to decay \( \epsilon \) over time, such as with an exponential decay schedule with the following update:

\[
\epsilon \leftarrow \alpha \epsilon
\]

for some \( \alpha \in (0, 1) \) typically close to 1.

Another strategy is explore-then-commit exploration (algorithm 15.4), where we select actions uniformly at random for the first \( k \) timesteps. From that point on, we choose a greedy action.\(^5\) Large values for \( k \) reduce the risk of committing to a suboptimal action, but we waste more time exploring potentially suboptimal actions.

We would like to apply the $\epsilon$-greedy exploration strategy to a two-armed bandit. We can construct the model with a uniform prior and the exploration policy with $\epsilon = 0.3$ and $\alpha = 0.99$:

\[
\text{model(fill(Beta(),2))}
\]
\[
\pi = \text{EpsilonGreedyExploration(0.3, 0.99)}
\]

To obtain our first action, we call $\pi(\text{model})$, which returns 1 based on the current state of the random number generator. We observe a loss, with $r = 0$, and then call

\[
\text{update!(model, 1, 0)}
\]

which updates the beta distributions within the model to reflect that we took action 1 and received a reward of 0.

The plots below show the evolution of the payoff beliefs after each of six steps of execution using our exploration strategy. Blue corresponds to the first action, and red corresponds to the second action.

Example 15.2. Application of the $\epsilon$-greedy exploration strategy to a two-armed bandit problem.
15.4 Directed Exploration Strategies

Directed exploration uses information gathered from previous pulls to guide exploration of the non-greedy actions. For example, the softmax strategy (algorithm 15.5) pulls arm \( a \) with probability proportional to \( \exp(\lambda \rho_a) \), where the precision parameter \( \lambda \geq 0 \) controls the amount of exploration. We have uniform random selection as \( \lambda \to 0 \) and greedy selection as \( \lambda \to \infty \). As more data is accumulated, we may want to increase \( \lambda \) by a multiplicative factor to reduce exploration.

A variety of exploration strategies are grounded in the idea of optimism under uncertainty. If we are optimistic about the outcomes of our actions to the extent that our data statistically allows, we will be implicitly driven to balance exploration and exploitation. One such approach is quantile exploration (algorithm 15.6),\(^6\) where we choose the arm with the highest \( \alpha \)-quantile (section 2.2.2) for the payoff probability. Values for \( \alpha > 0.5 \) result in optimism under uncertainty, incentivizing the exploration of actions that have not been tried as often. Larger values for \( \alpha \)...

---

result in more exploration. Example 15.3 shows quantile estimation and compares it with the other exploration strategies.

```plaintext
mutable struct QuantileExploration
    α  # quantile (e.g. 0.95)
end

function (π::QuantileExploration)(model::BanditModel)
    return argmax([quantile(B, π.α) for B in model.B])
end
```

Algorithm 15.6. Quantile exploration, which returns the action with the highest $α$ quantile.

An alternative to computing the upper confidence bound for our posterior distribution exactly is to use UCB1 exploration (algorithm 15.7), originally introduced in section 9.6 for exploration in Monte Carlo tree search. In this strategy, we select the action $a$ that maximizes

$$ρ_a + c \sqrt{\frac{\log N}{N(a)}}$$

(15.3)

where $N(a)$ is the number of times we have taken action $a$ and $N = \sum_a N(a)$. The parameter $c \geq 0$ controls the amount of exploration that is encouraged through the second term. Larger values of $c$ lead to more exploration. This strategy is often used with maximum likelihood estimates of the payoff probabilities, but we can adapt it to the Bayesian context by having $N(a)$ be the sum of the beta distribution parameters associated with $a$.

Another general approach to exploration is to use posterior sampling (algorithm 15.8), also referred to as randomized probability matching or Thompson sampling. It is simple to implement and does not have any parameters to tune. The idea is to sample from the posterior distribution over the rewards associated with the different actions. The action with the largest sampled value is selected.

15.5 Optimal Exploration Strategies

The beta distribution associated with arm $a$ is parameterized by counts $(w_a, \ell_a)$. Together these counts $w_1, \ell_1, \ldots, w_n, \ell_n$ represent our belief about payoffs, and thus represent a belief state. These $2n$ numbers can describe $n$ continuous probability distributions over possible payoff probabilities.

---

Consider using exploration strategies given the information obtained in the two-armed bandit problem of example 15.1, where the posterior distribution for $\theta_1$ is Beta(2, 1) and the posterior distribution for $\theta_2$ is Beta(5, 2). The second arm has the higher payoff probability.

An $\epsilon$-greedy strategy with $\epsilon = 0.2$ has a 20\% chance of choosing randomly between the arms and an 80\% chance of choosing the second arm. Hence, the overall probability of choosing the first arm is 0.1 and the probability of choosing the second arm is 0.9.

A softmax strategy with $\lambda = 1$ assigns a weight of $\exp(\rho_1) = \exp(2/3) \approx 1.948$ to the first arm and a weight of $\exp(\rho_2) = \exp(5/7) \approx 2.043$ to the second. The probability of choosing the first arm is $1.948 / (1.948 + 2.043) \approx 0.488$, and the probability of choosing the second arm is 0.512. The plot below shows how the probability of choosing the first arm varies with $\lambda$.

Quantile exploration with $\alpha = 0.9$ computes the payoff probability that is greater than 90\% of the probability mass associated with each posterior distribution. The 0.9 quantile for $\theta_1$ is 0.949 and for $\theta_2$ is 0.907, as shown below. The first arm (blue) has the higher quantile and would be pulled next.
mutable struct UCB1Exploration
c # exploration constant
end

function bonus(\(\pi\)::UCB1Exploration, B, a)
N = sum(b.\(\alpha\) + b.\(\beta\) for b in B)
Na = B[a].\(\alpha\) + B[a].\(\beta\)
return \(\pi\).c * sqrt(log(N)/Na)
end

function (\(\pi\)::UCB1Exploration)(model::BanditModel)
B = model.B
\(\rho\) = mean.(B)
\(u\) = \(\rho\) .+ [bonus(\(\pi\), B, a) for a in eachindex(B)]
return argmax(\(u\))
end

struct PosteriorSamplingExploration end

(\(\pi\)::PosteriorSamplingExploration)(model::BanditModel) =
argmax(rand.(model.B))

We can construct an MDP whose states are vectors of length \(2n\) that represent the agent’s belief over the \(n\)-armed bandit problem. Dynamic programming can be used to solve this MDP to obtain an optimal policy \(\pi^*\) that specifies which arm to pull given the counts.

Let \(Q^*(w_1, \ell_1, \ldots, w_n, \ell_n, a)\) represent the expected payoff after pulling arm \(a\) and thereafter acting optimally. The optimal utility function and optimal policy can be written in terms of \(Q^*\):

\[
U^*(w_1, \ell_1, \ldots, w_n, \ell_n) = \max_a Q^*(w_1, \ell_1, \ldots, w_n, \ell_n, a) \tag{15.4}
\]

\[
\pi^*(w_1, \ell_1, \ldots, w_n, \ell_n) = \arg\max_a Q^*(w_1, \ell_1, \ldots, w_n, \ell_n, a) \tag{15.5}
\]

We can decompose \(Q^*\) into two terms:

\[
Q^*(w_1, \ell_1, \ldots, w_n, \ell_n, a) = \frac{w_a + 1}{w_a + \ell_a + 2} (1 + U^*(\ldots, w_a + 1, \ell_a, \ldots))
+ \left(1 - \frac{w_a + 1}{w_a + \ell_a + 2}\right) U^*(\ldots, w_a, \ell_a + 1, \ldots) \tag{15.6}
\]
15.6 Exploration with Multiple States

The first term is associated with a win for arm \( a \), and the second term is associated with a loss. The value \( (w_a + 1)/(w_a + \ell_a + 2) \) is the posterior probability of a win, which comes from equation (15.1).\(^8\) The first \( U^\ast \) in the equation above records a win, whereas the second \( U^\ast \) records a loss.

We can compute \( Q^\ast \) for the entire belief space, as we have assumed a finite horizon \( h \). We start with all terminal belief states with \( \sum_a (w_a + \ell_a) = h \), where \( U^\ast = 0 \). We can then work backward to states with \( \sum_a (w_a + \ell_a) = h - 1 \) and apply equation (15.6). This process is repeated until we reach our initial state. Such an optimal policy is computed in example 15.4.

Although this dynamic programming solution is optimal, the number of belief states is \( O(h^{2n}) \). We can formulate an infinite horizon, discounted version of the problem that can be solved efficiently using the Gittins allocation index.\(^9\) The allocation index can be stored as a lookup table that specifies a scalar allocation index value given the number of pulls and the number of wins associated with an arm.\(^10\) The arm that has the highest allocation index is the one that should be pulled next.

15.6 Exploration with Multiple States

In the general reinforcement learning context with multiple states, we must use observations about state transitions to inform our decisions. We can modify the simulation process in algorithm 15.1 to account for state transitions and update our model appropriately. Algorithm 15.9 provides an implementation. There are many ways to model the problem and perform exploration as we will discuss over the next few chapters, but the simulation structure is exactly the same.

15.7 Summary

- The exploration-exploitation tradeoff is a balance between exploring the state-action space for higher rewards and exploiting the already known favorable state-actions.
- Multi-armed bandit problems involve a single state where the agent receives stochastic rewards for taking different actions.
- A beta distribution can be used to maintain a belief over multi-armed bandit rewards.

\(^8\) This probability can be adjusted if we have a nonuniform prior.
Below we have constructed the state-action tree for a two-arm bandit problem with a two-step horizon. State vectors are shown as $[w_1, \ell_1, w_2, \ell_2]$, blue arrows indicate wins, and red arrows indicate losses.

Unsurprisingly, the policy is symmetric with respect to arms 1 and 2. We find that the first arm does not matter, and it is best to pull a winning arm twice and not pull a losing arm twice.

The optimal value functions were computed using:

\[
\begin{align*}
Q^*([2, 1, 1, 1], 1) &= \frac{3}{5}(1 + 0) + \frac{2}{5}(0) = 0.6 \\
Q^*([2, 1, 1, 1], 2) &= \frac{2}{4}(1 + 0) + \frac{2}{4}(0) = 0.5 \\
Q^*([1, 2, 1, 1], 1) &= \frac{2}{5}(1 + 0) + \frac{3}{5}(0) = 0.4 \\
Q^*([1, 2, 1, 1], 2) &= \frac{2}{4}(1 + 0) + \frac{2}{4}(0) = 0.5 \\
Q^*([1, 1, 1, 1], 1) &= \frac{2}{4}(1 + 0.6) + \frac{2}{4}(0.5) = 1.05
\end{align*}
\]
function simulate(\(\mathcal{P}:\text{MDP}\), model, \(\pi\), \(h\), \(s\))
    for \(i\) in 1:h
        \(a = \pi(\text{model}, s)\)
        \(s', r = \mathcal{P}.\text{TR}(s, a)\)
        update!(\text{model}, \(s, a, r, s'\))
        \(s = s'\)
    end
end

• Undirected exploration strategies, including \(\epsilon\)-greedy and explore-then-commit, are simple to implement but do not use information from previous outcomes to guide exploration of non-greedy actions.

• Directed exploration strategies, including softmax, quantile, UCB1, and posterior sampling exploration, use information from past actions to better explore promising actions.

• Dynamic programming can be used to derive optimal exploration strategies for finite horizons, but these strategies can be expensive to compute.

15.8 Exercises

Exercise 15.1. Consider 3-armed bandit problems where each arm has a win probability drawn uniformly between 0 and 1. Compare the \(\epsilon\)-greedy, \(\epsilon\)-greedy with \(\alpha\) decay, and explore-then-commit exploration strategies. Qualitatively, what values for \(\epsilon\), \(\alpha\), and \(k\) produce the highest expected reward on randomly generated bandit problems?

Solution: Below we plot the expected reward-per-step for the three different strategies. The effectiveness of the parameterization depends on the problem horizon, so several different depths are shown as well.
Purely greedy policies ($\epsilon = 0$) outperform other $\epsilon$-greedy strategies for shorter horizons. Having a small non-zero chance of choosing randomly is most valuable for very long horizons. Including a moderate $\alpha$ decay improves performance, especially for large values of $\epsilon$ on problems with longer horizons. These policies had the highest overall expected reward per iteration. The explore-then-commit policy behaves in a manner similar to $\epsilon$-greedy with $\alpha$ decay. It performs best with $k$ at roughly 20% of the horizon.

**Exercise 15.2.** Consider again the 3-armed bandit problems where each arm has a win probability drawn uniformly between 0 and 1. Compare the softmax, quantile, and UCB1 exploration strategies. Qualitatively, what values for $\lambda$, $\alpha$, and $\epsilon$ produce the highest expected reward on randomly generated bandit problems?
Solution: Below we plot the expected reward-per-step for the three different strategies. Again, the effectiveness of the parameterization depends on the problem horizon, so several different depths are shown as well.

The softmax strategy performs best for large values of $\lambda$, which prioritize pulling arms with higher expected reward according to the current belief. Upper confidence bound exploration performs better with longer horizons, independently of its parameterization. The size of the confidence bound $\alpha$ does not significantly affect performance except for values very close to 0 or 1. The UCB1 strategy performs best with small positive values of the exploration scalar $c$. The expected reward decays as $c$ increases. All three policies can be tuned to produce similar maximal expected rewards.
Exercise 15.3. Give an example of a multi-armed bandit problem.

Solution: There are many different multi-armed bandit problems. Consider, for example, a news company that would like to maximize interaction (clicks) on articles on its website. The company may have several articles to display, but must select on article to display at a given time. This problem is a multi-armed bandit problem because a user will either click article $i$ with probability $\theta_i$ or not click with probability $1 - \theta_i$. Exploration would consist of displaying different articles on the website and observing the number of clicks and exploitation would consist of displaying the article likely to lead to the highest number of clicks. This problem is related to A/B testing, where companies test different versions of a website to determine which version yields the most interaction.

Exercise 15.4. Given a single-armed bandit with a prior of $\theta \sim \text{Beta}(7, 2)$, provide bounds on the posterior probability of winning after 10 additional pulls.

Solution: A lower bound on our posterior probability of winning $\rho$ can be computed assuming all pulls result in a loss, e.g. $\ell = 10$ and $w = 0$. We can similarly compute an upper bound $\bar{\rho}$ assuming all pulls result in a win, e.g. $w = 10$ and $\ell = 0$. Thus, the bounds are

$$\rho = \frac{w + 7}{w + \ell + 9} = \frac{0 + 7}{0 + 10 + 9} = \frac{7}{19}$$

$$\bar{\rho} = \frac{w + 7}{w + \ell + 9} = \frac{10 + 7}{10 + 0 + 9} = \frac{17}{19}$$

Exercise 15.5. We have a bandit with arms $a$ and $b$, and we use an $\epsilon$-greedy exploration strategy with $\epsilon = 0.3$ and an exploration decay factor of $\alpha = 0.9$. We generate a random number $x$ between 0 and 1 to determine if we explore ($x < \epsilon$) or exploit ($x > \epsilon$). Given we have $\rho_a > \rho_b$, which arm is selected if $x = 0.2914$ in the first iteration? Which arm is selected if $x = 0.1773$ in the ninth iteration?

Solution: Since $x < \epsilon_1$ in the first iteration, we explore and choose $a$ with probability 0.5 and $b$ with probability 0.5. At the ninth iteration, $\epsilon_9 = a^8 \epsilon_1 \approx 0.129$. Since $x > \epsilon_9$, we exploit and select $a$.

Exercise 15.6. We have a four-armed bandit and we want to use a softmax exploration strategy with precision parameter $\lambda = 2$ and a prior belief $\theta_a \sim \text{Beta}(2, 2)$ for each arm $a$. Suppose we pull each arm four times with the result that arms 1, 2, 3, and 4 pay off 1, 2, 3 and 4 times, respectively. List the posterior distributions over $\theta_a$ and calculate the probability that we select arm 2.
Solution: The posterior distribution for each arm is respectively: Beta(3, 5), Beta(4, 4), Beta(5, 3), Beta(6, 2). The probability of selecting arm 2 can be computed in the following steps

\[
P(a = i) \propto \exp (\lambda \rho_i)
\]

\[
P(a = i) = \frac{\exp (\lambda \rho_i)}{\sum_a \exp (\lambda \rho_a)}
\]

\[
P(a = 2) = \frac{\exp (2 \times \frac{4}{8})}{\exp (2 \times \frac{3}{8}) + \exp (2 \times \frac{4}{8}) + \exp (2 \times \frac{5}{8}) + \exp (2 \times \frac{6}{8})}
\]

\[P(a = 2) \approx 0.2122\]

Exercise 15.7. Rewrite equation (15.6) for an arbitrary Beta(\(\alpha, \beta\)) prior.

Solution: We can rewrite the equation more generally as follows

\[
Q^*(w_1, \ell_1, \ldots, w_n, \ell_n, a) = \frac{w_a + \alpha}{w_a + \ell_a + \alpha + \beta} (1 + U^*(\ldots, w_a + 1, \ell_a, \ldots))
\]

\[+ \left(1 - \frac{w_a + \alpha}{w_a + \ell_a + \alpha + \beta}\right) U^*(\ldots, w_a, \ell_a + 1, \ldots)\]

Exercise 15.8. Recall example 15.4. Instead of having a payoff of 1 for each arm, let us assume arm 1 gives a payoff of 1 while arm 2 gives a payoff of 2. Calculate the new action value functions for both arms.

Solution: For arm 1, we have:

\[
Q^*([2, 1, 1, 1], 1) = \frac{3}{5} (1 + 0) + \frac{2}{5} (0) = 0.6
\]

\[
Q^*([2, 1, 1, 1], 2) = \frac{2}{4} (2 + 0) + \frac{2}{4} (0) = 1
\]

\[
Q^*([1, 2, 1, 1], 1) = \frac{2}{5} (1 + 0) + \frac{3}{5} (0) = 0.4
\]

\[
Q^*([1, 2, 1, 1], 2) = \frac{2}{4} (2 + 0) + \frac{2}{4} (0) = 1
\]

\[
Q^*([1, 1, 1, 1], 1) = \frac{2}{4} (1 + 1) + \frac{2}{4} (1) = 1.5
\]
And for arm 2, we have:

\[
Q^*([1, 1, 2, 1], 1) = \frac{2}{4}(1 + 0) + \frac{2}{4}(0) = 0.5 \\
Q^*([1, 1, 2, 1], 2) = \frac{3}{5}(2 + 0) + \frac{2}{5}(0) = 1.2 \\
Q^*([1, 1, 1, 2], 1) = \frac{2}{4}(1 + 0) + \frac{2}{4}(0) = 0.5 \\
Q^*([1, 1, 1, 2], 2) = \frac{2}{5}(2 + 0) + \frac{3}{5}(0) = 0.8 \\
Q^*([1, 1, 1, 1], 2) = \frac{2}{4}(2 + 1.2) + \frac{2}{4}(0.8) = 2
\]

**Exercise 15.9.** Prove that the number of belief states in an \(n\)-armed bandit problem with a horizon of \(h\) is \(O(h^{2n})\).

**Solution:** We begin by counting the number of solutions to \(w_1 + \ell_1 + \cdots + w_n + \ell_n = k\), where \(0 \leq k \leq h\). If \(n = 2\) and \(k = 6\), one solution is \(2 + 0 + 3 + 1 = 6\). For our counting argument, we will use tally marks to represent our integers. For example, we can write a solution like \(2 + 0 + 3 + 1 = |++|+|= 6\). For general values for \(n\) and \(k\), we would have \(k\) tally marks and \(2n - 1\) plus signs. Given that many tally marks and plus signs, we can arrange them in any order we want. We can represent a solution as a string of \(k + 2n - 1\) characters, where a character is either | or +, with \(k\) of those characters being |. To obtain the number of solutions, we count the number of ways we can choose \(k\) positions for | from the set of \(k + 2n - 1\) positions, resulting in

\[
\frac{(k + 2n - 1)!}{(2n - 1)!k!} = O(h^{2n-1})
\]

solutions. The number of belief states is this expression summed for \(k\) from 0 to \(h\), which is \(O(h \times h^{2n-1}) = O(h^{2n})\).

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2021-03-07 23:12:00-08:00, revision 80bf4d6, comments to bugs@algorithmsbook.com
This chapter discusses both maximum likelihood and Bayesian approaches for learning the underlying dynamics and reward through interaction with the environment. Maximum likelihood methods involve counting state transitions and recording the amount of reward received to estimate the model parameters. We will discuss a few approaches for planning using models that are continuously updated. Even if we solve the estimated problem exactly, we generally have to rely upon heuristic exploration strategies to arrive at a suitable solution. Bayesian methods involve computing a posterior distribution over model parameters. Solving for the optimal exploration strategy is generally intractable, but we can often obtain a sensible approximation through posterior sampling.

16.1 Maximum Likelihood Models

As introduced in section 15.6 and implemented in algorithm 15.9, reinforcement learning involves using information about past state transitions and rewards to inform decisions. This section describes how to obtain a maximum likelihood estimate of the underlying problem. This maximum likelihood estimate can be used to generate a value function estimate that can be used with an exploration strategy to generate actions.

We record the transition counts $N(s, a, s')$, indicating the number of times a transition from $s$ to $s'$ was observed when taking action $a$. The maximum likelihood estimate of the transition function given transition counts $N(s, a, s')$ is:

$$ T(s' \mid s, a) \approx \frac{N(s, a, s')}{N(s, a)} \quad (16.1) $$
where \( N(s, a) = \sum_{s'} N(s, a, s') \). If \( N(s, a) = 0 \), then the estimate of the transition probability is 0.

The reward function can also be estimated. As we receive rewards, we update \( \rho(s, a) \), the sum of all rewards obtained when taking action \( a \) in state \( s \). The maximum likelihood estimate of the reward function is the mean reward:

\[
R(s, a) \approx \frac{\rho(s, a)}{N(s, a)} \tag{16.2}
\]

If \( N(s, a) = 0 \), then our estimate of \( R(s, a) \) is 0. If we have prior knowledge about the transition probabilities or rewards, then we can initialize \( N(s, a, s') \) and \( \rho(s, a) \) to values other than 0.

Algorithm 16.1 updates \( N \) and \( \rho \) after observing the transition from \( s \) to \( s' \) after taking action \( a \) and receiving reward \( r \). Algorithm 16.2 converts the maximum likelihood model into a Markov decision process representation. Example 16.1 illustrates this process. We can use this maximum likelihood model to select actions while interacting with the environment and improving the model.

### 16.2 Update Schemes

As we update our maximum likelihood estimate of the model, we also need to update our plan. This section discusses several different update schemes in response to our continuously changing model. A major consideration is computational efficiency because we will want to perform these updates fairly frequently while interacting with the environment.

#### 16.2.1 Full Updates

Algorithm 16.3 solves the maximum likelihood model using the linear programming formulation from section 7.7, though we could have used value iteration or some other algorithm. After each step, we obtain a new model estimate, and we re-solve.

#### 16.2.2 Randomized Updates

Recomputing an optimal policy with each state transition is typically computationally expensive. An alternative is to perform a Bellman update on the estimated model at the previously visited state as well as a few randomly chosen states.\(^1\) Algorithm 16.4 implements this approach.

mutable struct MaximumLikelihoodMDP
    𝒮 # state space (assumes 1:nstates)
    𝒜 # action space (assumes 1:nactions)
    N # transition count N(s,a,s′)
    ρ # reward sum ρ(s, a)
    γ # discount
    U # value function
    planner
end

function lookahead(model::MaximumLikelihoodMDP, s, a)
    𝒮, U, γ = model.𝒮, model.U, model.γ
    n = sum(model.N[s,a,:])
    if n == 0
        return 0.0
    end
    r = model.ρ[s, a] / n
    T(s,a,s′) = model.N[s,a,s′] / n
    return r + γ * sum(T(s,a,s′)*U[s′] for s′ in 𝒮)
end

function backup(model::MaximumLikelihoodMDP, U, s)
    return maximum(lookahead(model, s, a) for a in model.𝒜)
end

function update!(model::MaximumLikelihoodMDP, s, a, r, s′)
    model.N[s,a,s′] += 1
    model.ρ[s,a] += r
    update!(model.planner, model, s, a, r, s′)
    return model
end

function MDP(model::MaximumLikelihoodMDP)
    N, ρ, 𝒮, 𝒜, γ = model.N, model.ρ, model.𝒮, model.𝒜, model.γ
    T, R = similar(N), similar(ρ)
    for s in 𝒮
        for a in 𝒜
            n = sum(N[s,a,:])
            if n == 0
                T[s,a,:] .= 0.0
                R[s,a] = 0.0
            else
                T[s,a,:] = N[s,a,:] / n
                R[s,a] = ρ[s,a] / n
            end
        end
    end
    return MDP(T, R, γ)
end

Algorithm 16.1. A method for updating the transition and reward model for maximum likelihood reinforcement learning with discrete state and action spaces. We increment $N(s,a,s')$ after observing a transition from $s$ to $s'$ after taking action $a$, and we add $r$ to $\rho(s,a)$. The model also contains an estimate of the value function $U$ and a planner. This algorithm block also includes methods for performing backup and lookahead with respect to this model.

Algorithm 16.2. A method for converting a maximum likelihood model to an MDP problem.
We would like to apply maximum likelihood model estimation to the hex world problem. The true transition matrices look like this:

There are six transition matrices, one for each action. The rows correspond to the current state and the columns correspond to the next state. There are 26 states. The intensity in the images relate to the probability of making the corresponding transition. In a reinforcement learning context, we do not know these transition probabilities ahead of time. However, we can interact with the environment and record the transitions we observe. After 10 simulations of 10 steps each from random initial states, maximum likelihood estimation results in the following matrices:

After 1000 simulations, our estimate becomes:
Algorithm 16.3. An method that performs a full update of the value function of \( U \) using the linear programming formulation from section 7.7.

Algorithm 16.4. Maximum likelihood model-based reinforcement learning with updates at randomized states. This approach performs a Bellman update at the previously visited state as well as at \( m \) additional states chosen randomly.

16.2.3 Prioritized Updates

An approach called prioritized sweeping\(^2\) (algorithm 16.5) uses a priority queue to help identify which states are most in need of updating. A transition from \( s \) to \( s' \) is followed by an update of \( U(s) \) based on our updated transition and reward models. We then iterate over all state-action pairs \( (s^-, a^-) \) that can immediately transition into \( s \). The priority of any such \( s^- \) is increased to \( T(s^+ | s^-, a^-) \times |U(s) - u| \), where \( u \) was the value of \( U(s) \) before the update. Hence, the larger the change in \( U(s) \) and the more likely the transition to \( s \), the higher the priority of states leading to \( s \). The process of updating the highest priority state in the queue continues for some fixed number of iterations or until the queue becomes empty.

16.3 Exploration

Regardless of the update scheme, some form of exploration strategy generally must be followed to avoid the pitfalls of pure exploitation mentioned in the previous chapter. We can adapt the exploration algorithms presented in that chapter

Algorithm 16.5. The prioritized sweeping algorithm maintains a priority queue \( pq \) of states that determines which are to be updated. With each update, we set the previous state to have infinite priority. We then perform \( m \) Bellman updates of the value function \( U \) at the highest priority states.

```plaintext
struct PrioritizedUpdate
    m # number of updates
    pq # priority queue
end

function update!(planner::PrioritizedUpdate, model, s)
    N, U, pq = model.N, model.U, planner.pq
    S, \( \Lambda \) = model.S, model.\( \Lambda \)
    u = U[s]
    U[s] = backup(model, U, s)
    for s⁻ in \( \mathcal{S} \)
        for a⁻ in \( \mathcal{A} \)
            n_sa = sum(N[s⁻, a⁻, s'] for s' in \( \mathcal{S} \))
            if n_sa > 0
                T = N[s⁻, a⁻, s] / n_sa
                priority = T * abs(U[s] - u)
                pq[s⁻] = max(get(pq, s⁻, -Inf), priority)
            end
        end
    end
    return planner
end

function update!(planner::PrioritizedUpdate, model, s, a, r, s')
    planner.pq[s] = Inf
    for i in 1:planner.m
        if isempty(planner.pq)
            break
        end
        update!(planner, model, dequeue!(planner.pq))
    end
    return planner
end
```
for use in multi-state problems. Algorithm 16.6 provides an implementation of the $\epsilon$-greedy exploration strategy.

```
function (π::EpsilonGreedyExploration)(model, s)
    $\lambda$, $\epsilon = model.\lambda$, $\pi.\epsilon$
    if rand() < $\epsilon$
        return rand($\lambda$)
    end
    $Q(s,a) = \text{lookahead}(model, s, a)$
    return argmax($a\rightarrow Q(s,a)$, $\lambda$)
end
```

Algorithm 16.6. The $\epsilon$-greedy exploration strategy for maximum likelihood model estimates. It chooses a random action with probability $\epsilon$, otherwise it uses the model to extract the greedy action.

A limitation of the exploration strategies discussed in the previous chapter is that they do not reason about exploring actions from states besides the current one. For instance, we might want to take actions that bring ourselves into an area of the state space that has not been explored. Several different algorithms have been suggested for addressing this issue that also provide probabilistic bounds on the quality of the resulting policy after a finite number of interactions.$^3$

One such algorithm is known as R-MAX (algorithm 16.7).$^4$ Its name comes from assigning maximal reward to state-action pairs that are underexplored as defined by a count threshold $m$. Instead of using the maximum likelihood estimate for the reward (equation (16.2)), we use

$$R(s,a) = \begin{cases} r_{\text{max}} & \text{if } N(s,a) < m \\ \rho(s,a)/N(s,a) & \text{otherwise} \end{cases}$$

where $r_{\text{max}}$ is the maximum achievable reward.

The transition model in R-MAX is also modified so that underexplored state-action pairs result in staying the same state:

$$T(s' | s,a) = \begin{cases} (s' = s) & \text{if } N(s,a) < m \\ N(s,a,s')/N(s,a) & \text{otherwise} \end{cases}$$

Hence, underexplored states have value $r_{\text{max}}/(1 - \gamma)$, providing incentive to explore them. This exploration incentive relieves us of needing a separate exploration mechanism. We simply choose our actions greedily with respect to the value function derived from our transition and reward estimates. Example 16.2 demonstrates $\epsilon$-greedy and R-MAX exploration.


mutable struct RmaxMDP
  S # state space (assumes 1:nstates)
  A # action space (assumes 1:nactions)
  N # transition count N(s,a,s')
  ρ # reward sum ρ(s, a)
  γ # discount
  U # value function
  planner
  m # count threshold
  rmax # maximum reward
end

function lookahead(model::RmaxMDP, s, a)
  S, U, γ = model.S, model.U, model.γ
  n = sum(model.N[s,a,:])
  if n < model.m
    return model.rmax / (1-γ)
  end
  r = model.ρ[s, a] / n
  T(s,a,s') = model.N[s,a,s'] / n
  return r + γ * sum(T(s,a,s')*U[s'] for s' in S)
end

function backup(model::RmaxMDP, U, s)
  return maximum(lookahead(model, s, a) for a in model.A)
end

function update!(model::RmaxMDP, s, a, r, s')
  model.N[s,a,s'] += 1
  model.ρ[s,a] += r
  update!(model.planner, model, s, a, r, s')
  return model
end

function MDP(model::RmaxMDP)
  T, R, m, rmax = similar(N), similar(ρ), model.m, model.rmax
  for s in S
    for a in A
      n = sum(N[s,a,:])
      if n < m
        T[s,a,:] .= 0.0
        T[s,a,s] = 1.0
        R[s,a] = rmax
      else
        T[s,a,:] = N[s,a,:] / n
        R[s,a] = ρ[s,a] / n
      end
    end
  end
  return MDP(T, R, γ)
end

Algorithm 16.7. The R-MAX exploration strategy modifies the transition and reward model from maximum likelihood estimation. It assigns maximum reward $r_{max}$ to any underexplored state-action pair, defined as being those that have been tried fewer than $m$ times, where $m$ is constrained to be at least 1. In addition, all underexplored state-action pairs are modeled as transitioning to the same state. This $RmaxMDP$ can be used as a replacement for the $MaximumLikelihoodMDP$ introduced in algorithm 16.1.
16.4 Bayesian Methods

In contrast with the maximum likelihood methods discussed so far, Bayesian methods balance exploration and exploitation without having to rely on heuristic exploration policies. This section describes a generalization of the Bayesian methods covered in section 15.5. In Bayesian reinforcement learning, we specify a prior distribution over all model parameters $\theta$. These model parameters may include the parameters governing the distribution over immediate rewards, but this section focuses on the parameters governing the state transition probabilities.

The structure of the problem can be represented using the dynamic decision network shown in figure 16.1 with the model parameters made explicit. The shaded nodes indicate that the states are observed, but the model parameters are not. We generally assume that the model parameters are time invariant with $\theta_{t+1} = \theta_t$. However, our belief about $\theta$ evolves with time as we transition to new states.

The belief over transition probabilities can be represented using a collection of Dirichlet distributions, one for each source state and action. Each Dirichlet distribution represents the distribution over $s'$ for a given $s$ and $a'$. If $\theta_{(s,a)}$ is an

![Figure 16.1. A dynamic decision network for an MDP with model uncertainty.](image)

Example 16.2. Demonstration of $\epsilon$-greedy and R-MAX exploration.

$|S|$-element vector representing the distribution over the next state, then the prior distribution is given by

$$\text{Dir}(\theta_{(s,a)} \mid \mathbf{N}(s,a))$$

(16.5)

where $\mathbf{N}(s,a)$ is the vector of counts associated with transitions starting in state $s$ taking action $a$. It is common to use a uniform prior with all components set to 1, but prior knowledge of the transition dynamics can be used to initialize the counts differently. Example 16.3 illustrates how these counts are used by the Dirichlet distribution to represent the distribution over possible transition probabilities.

The distribution over $\theta$ is the result of the product of the Dirichlet distributions:

$$b(\theta) = \prod_s \prod_a \text{Dir}(\theta_{(s,a)} \mid \mathbf{N}(s,a))$$

(16.6)

Algorithm 16.8 provides an implementation of the Bayesian update for this type of posterior model. For problems with larger or continuous spaces, we can use other posterior representations.

16.5 Bayes-adaptive MDPs

We can formulate the problem of acting optimally in an MDP with an unknown model as a higher dimensional MDP with a known model. This higher dimensional MDP is known as a Bayes-adaptive Markov decision process, which is related to the partially observable Markov decision process discussed in the next part of the book.

The state space in the Bayes-adaptive MDP is the Cartesian product $S \times B$, where $B$ is the space of possible beliefs over the model parameters $\theta$. Though $S$ is discrete, $B$ is often a high-dimensional continuous state space. A state in a Bayes-adaptive MDP is a pair $(s,b)$ consisting of the current state $s$ in the base MDP and a belief state $b$. The action space and reward function are the same as in the base MDP.

The transition function in a Bayes-adaptive MDP is $T(s',b' \mid s,b,a)$, which is the probability of transitioning to some state $s'$ with a belief state $b'$, given that the agent starts in $s$ with belief $b$ and takes action $a$. The new belief state $b'$ can be deterministically computed according to Bayes’ rule. If we let this deterministic function be denoted $\tau$ so that $b' = \tau(s,b,a,s')$, then we can decompose the Bayes-
Suppose our agent randomly explores an environment with three states. The agent takes action $a_1$ from state $s_1$ five times. It transitions to $s_3$ four times, and remains in $s_1$ once. We end up with the following counts associated with $s_1$ and $a_1$:

$$N(s_1, a_1) = [1, 0, 4]$$

If we want to assume a uniform prior over resulting states, we would increment the counts by one to get $N(s_1, a_1) = [2, 1, 5]$. The transition function from $s_1$ taking action $a_1$ is a three-valued categorical distribution because there are three possible successor states. Each successor state has an unknown transition probability. The space of possible transition probabilities is the set of three-element vectors that sum to one. The Dirichlet distribution represents a probability distribution over these possible transition probabilities. Below is a plot of the density function.

Example 16.3. An example posterior Dirichlet distribution over transition probabilities from a particular state when taking a particular action. An agent learning the transition function in an unknown MDP may choose to maintain such a distribution over each state-action pair.
mutable struct BayesianMDP
    𝒮 # state space (assumes 1:nstates)
    𝒜 # action space (assumes 1:nactions)
    D # Dirichlet distributions D[s,a]
    R # reward function as matrix (not estimated)
    γ # discount
    U # value function
end

function lookahead(model::BayesianMDP, s, a)
    𝒮, U, γ = model.𝒮, model.U, model.γ
    n = sum(model.D[s,a].alpha)
    if n == 0
        return 0.0
    end
    r = model.R(s,a)
    T(s,a,s′) = model.D[s,a].alpha[s′] / n
    return r + γ * sum(T(s,a,s′)*U[s′] for s′ in 𝒮)
end

function update!(model::BayesianMDP, s, a, r, s′)
    α = model.D[s,a].alpha
    α[s′] += 1
    model.D[s,a] = Dirichlet(α)
    update!(model.planner, model, s, a, r, s′)
    return model
end

Algorithm 16.8. A Bayesian update method when the posterior distribution over transition models is represented as a product of Dirichlet distributions. We assume in this implementation that the reward model \( R \) is known, though we can use Bayesian methods to estimate expected reward from experience. The matrix \( D \) associates Dirichlet distributions with every state-action pair to model uncertainty in the transition to their successor states.
adaptive MDP transition function as:

\[ T(s', b' | s, b, a) = \delta_{\tau(s,b,a,s')}(b') P(s' | s, b, a) \]  \hspace{1cm} (16.7)

where \( \delta_x(y) \) is the Kronecker delta function such that \( \delta_x(y) = 1 \) if \( x = y \) and 0 otherwise.

The second term can be computed using integration:

\[ P(s' | s, b, a) = \int_\theta b(\theta)P(s' | s, b, a) \, d\theta = \int_\theta b(\theta)\theta_{(s,a,s')} \, d\theta \]  \hspace{1cm} (16.8)

This equation can be evaluated analytically in a manner similar to equation (15.1). In the case our belief \( b \) is represented by the factored Dirichlet in equation (16.6), we have

\[ P(s' | s, b, a) = N(s, a, s') / \sum_{s''} N(s, a, s'') \]  \hspace{1cm} (16.9)

We can generalize the Bellman equation (equation (7.16)) for MDPs with a known model to the case in which the model is unknown:

\[ U^*(s, b) = \max_a \left( R(s, a) + \gamma \sum_{s'} P(s' | s, b, a) U^*(s', \tau(s, b, a, s')) \right) \]  \hspace{1cm} (16.10)

Unfortunately, we cannot simply directly apply policy iteration or value iteration because \( b \) is continuous. We can, however use the approximation methods of chapter 8 or the online methods of chapter 9. The next part of the book presents methods that better use the structure of the Bayes-adaptive MDP.

### 16.6 Posterior Sampling

An alternative to solving for the optimal value function over the belief space is to use posterior sampling, which was originally introduced in the context of exploration in bandit problems in section 15.4. Here, we draw a sample \( \theta \) from the current belief \( b \) and then solve for the best action assuming \( \theta \) is the true model. We then update our belief, draw a new sample, and solve the corresponding MDP. Example 16.4 provides an example.

An advantage of posterior sampling is that we do not have to decide on heuristic exploration parameters. However, solving the MDP at every step can be expensive. A method for sampling a discrete MDP from the posterior is implemented in algorithm 16.9.

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Algorithm 16.9. The update method for posterior sampling. After updating the parameters of the Bayesian posterior, we sample an MDP problem from that posterior. This implementation assumes a discrete state and action space with a Dirichlet modeling our uncertainty in the transition probabilities from each state-action pair. To generate the transition model, we iterate over every state and action and sample from the associated Dirichlet distribution. Once we have a sampled problem $\mathcal{P}$, we solve it using the linear programming formulation and store the resulting value function $U$. 

We want to apply Bayesian model estimation to hex world. We start with associating uniform Dirichlet priors with every state-action pair. After 100 simulations of length 10 and adding our transition counts to our pseudo-counts in our prior, the parameters of our posterior distributions over our successor states appear as follows:

We can sample from this distribution to produce the model shown below. Notice that it has many more non-zero transition probabilities than the maximum likelihood models shown in example 16.1.
16.7 Summary

- Model-based methods learn the transition and reward models through interaction with the environment.

- Maximum likelihood models use transition counts to maintain an estimate of the transition probabilities to successor states and track the mean reward associated with state-action pairs.

- Maximum likelihood models must be paired with an exploration strategy, such as those introduced in the previous chapter in the context of bandits.

- Although we can replan with each step of experience, doing so exactly can be costly.

- Prioritized sweeping can focus replanning by updating the values of states that appear to need it the most in our evolving model of the environment.

- Bayesian model-based methods maintain a probability distribution over possible problems, allowing for principled reasoning about exploration.

- Bayes-adaptive MDPs are MDPs whose states augment the original MDP with the probability distribution over the possible MDP models.

- Posterior sampling reduces the high computational complexity of solving a Bayes-adaptive MDP by solving an MDP sampled from the belief state rather than reasoning about all possible MDPs.

16.8 Exercises

Exercise 16.1. Suppose we have an agent interacting in an environment with unknown transition and reward models. We perform one sequence of direct interaction with the environment. Table 16.1 tabulates the state, action, reward, and resulting state. Use maximum likelihood estimation to estimate the transition and reward functions from this data.

Solution: We first tabulate the number of transitions from each state and action \( N(s,a) \), the rewards received \( \rho(s,a) \), and the maximum likelihood estimate of the reward function \( \hat{R}(s,a) = \rho(s,a) / N(s,a) \):

<table>
<thead>
<tr>
<th>s</th>
<th>a</th>
<th>r</th>
<th>s'</th>
</tr>
</thead>
<tbody>
<tr>
<td>( s_2 )</td>
<td>( a_1 )</td>
<td>2</td>
<td>( s_1 )</td>
</tr>
<tr>
<td>( s_1 )</td>
<td>( a_2 )</td>
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<td>( s_2 )</td>
<td>( a_1 )</td>
<td>2</td>
<td>( s_3 )</td>
</tr>
</tbody>
</table>
Exercise 16.2. Provide a lower bound and an upper bound on the number of updates that could be performed during an iteration of prioritized sweeping.

Solution: A lower bound on the number of updates performed in an iteration of prioritized sweeping is 1. This could occur during our first iteration using a maximum likelihood model, where the only nonzero entry in our transition model is $T(s' | s, a)$. Since no state-action pairs $(s, a)$ transition to $s$, our priority queue would be empty and thus the only update performed would be for $U(s)$.
An upper bound on the number of updates performed in an iteration of prioritized sweeping is $|S|$. Suppose that we just transitioned to $s'$ and that $\hat{T}(s' \mid s, a) > 0$ for all $s$ and $a$. If we do not provide a maximum number of updates, we will perform $|S|$ updates. If we provide a maximum number of updates $m < |S|$, the upper bound is reduced to $m$.

**Exercise 16.3.** In performing Bayesian reinforcement learning of the transition model parameters for a discrete MDP with state space $S$ and action space $A$, how many independent parameters are there when using Dirichlet distributions to represent the uncertainty over the transition model?

*Solution:* For each state and action, we specify a Dirichlet distribution over the transition probability parameters, so we will have $|S||A|$ Dirichlet distributions. Each Dirichlet is specified using $|S|$ independent parameters. In total, we have $|S|^2|A|$ independent parameters.

**Exercise 16.4.** Consider the problem statement in exercise 16.1, but this time we want to use Bayesian reinforcement learning with a prior distribution represented by a Dirichlet distribution. Assuming a uniform prior, what is the posterior distribution over the next state given that we are in state $s_2$ and take action $a_1$?

*Solution:* $\text{Dir}(\theta_{(s_2,a_1)} \mid [2,1,2])$
17 Model-Free Methods

In contrast with model-based methods, model-free reinforcement learning does not require building explicit representations of the transition and reward models. The model-free methods discussed in this chapter model the action value function directly. Avoiding explicit representations is attractive, especially when the problem is high dimensional. This chapter begins by introducing incremental estimation of the mean of a function, which plays an important role in estimating the mean of returns. We then discuss some common model-free algorithms and methods for handling delayed reward more efficiently. Finally, we discuss how to use function approximation to generalize from our experience.

17.1 Incremental Estimation of the Mean

Many model-free methods incrementally estimate the action value function $Q(s, a)$ from samples. For the moment, suppose we are only concerned with the expectation of a single variable $X$ from $m$ samples:

$$\hat{x}_m = \frac{1}{m} \sum_{i=1}^{m} x^{(i)} \quad (17.1)$$

where $x^{(1)}, \ldots, x^{(m)}$ are our samples. We can derive an incremental update:

$$\hat{x}_m = \frac{1}{m} \left( x^{(m)} + \sum_{i=1}^{m-1} x^{(i)} \right) \quad (17.2)$$

$$= \frac{1}{m} \left( x^{(m)} + (m - 1)\hat{x}_{m-1} \right) \quad (17.3)$$

$$= \hat{x}_{m-1} + \frac{1}{m} \left( x^{(m)} - \hat{x}_{m-1} \right) \quad (17.4)$$


2 Although this part of the book has been focusing on problems where the model of the environment is unknown, reinforcement learning is often used for problems with known models. The model-free methods discussed in this chapter can be especially useful in complex environments as a form of approximate dynamic programming. They can be used to produce policies offline or as a means to generate the next action in an online context.
We can rewrite this equation with the introduction of a learning rate function $\alpha(m)$:

$$\hat{x}_m = \hat{x}_{m-1} + \alpha(m)\left(x^{(m)} - \hat{x}_{m-1}\right) \quad (17.5)$$

The learning rate can be a function other than $1/m$. To ensure convergence, we generally select $\alpha(m)$ such that we have $\sum_{m=1}^{\infty} \alpha(m) = \infty$ and $\sum_{m=1}^{\infty} \alpha^2(m) < \infty$. The first condition ensures that the steps are sufficiently large, and the second condition ensures that the steps are sufficiently small.\(^3\)

If the learning rate is constant, which is common in reinforcement learning applications, then the weights of older samples decay exponentially at the rate $(1 - \alpha)$.

With a constant learning rate, we can update our estimate after observing $x$ using the following rule:

$$\hat{x} \leftarrow \hat{x} + \alpha(x - \hat{x}) \quad (17.6)$$

Algorithm 17.1 provides an implementation. An example of several learning rates is shown in example 17.1.

The update rule above will appear again in later sections and is related to stochastic gradient descent. The magnitude of the update is proportional to the difference between the sample and the previous estimate. The difference between the sample and previous estimate is called the temporal difference error.

### 17.2 Q-Learning

Q-learning (algorithm 17.2) involves applying incremental estimation of the action value function $Q(s, a)$.\(^4\) The update is derived from the action value form of the Bellman equation:

$$Q(s, a) = R(s, a) + \gamma \sum_{s'} T(s' | s, a) U(s')$$

$$= R(s, a) + \gamma \sum_{s'} T(s' | s, a) \max_{a'} Q(s', a') \quad (17.7)$$

$$= R(s, a) + \gamma \sum_{s', a'} T(s' | s, a) \max_{a'} Q(s', a') \quad (17.8)$$

Instead of using $T$ and $R$, we can rewrite the equation above in terms of an expectation over samples of reward $r$ and the next state $s'$:

$$Q(s, a) = \mathbb{E}_{r, s'}[r + \gamma \max_{a'} Q(s', a')] \quad (17.9)$$


Consider estimating the expected value obtained when rolling a fair six-sided die. Below are *learning curves* that show the incremental estimates over 100 trials associated with different learning rate functions. As we can see, convergence is not guaranteed if \( \alpha(m) \) decays too quickly, and is slow if \( \alpha(m) \) does not decay quickly enough.

For constant values of \( \alpha \in (0, 1] \), the mean estimate will continue to fluctuate. Larger values of constant \( \alpha \) fluctuate wildly, whereas lower values take longer to converge.

### Example 17.1

The effect of decaying the learning rate with different functions for \( \alpha(m) \).

### Algorithm 17.1

A type for maintaining an incremental estimate of the mean of a random variable. The associated type maintains a current mean value \( \mu \), a learning rate function \( \alpha \), and an iteration count \( m \). Calling `update!` with a new value \( x \) updates the estimate.
We can use equation (17.6) to produce an incremental update rule to estimate the action value function:

\[ Q(s,a) \leftarrow Q(s,a) + \alpha \left( r + \gamma \max_{a'} Q(s',a') - Q(s,a) \right) \quad (17.10) \]

Our choice of actions affects which states we end up in, and therefore our ability to accurately estimate \( Q(s,a) \). In order to guarantee convergence of our action value function, we need to adopt some form of exploration policy, such as \( \epsilon \)-greedy or softmax, just as we did for our model-based methods in the previous chapter. Example 17.2 shows how to run a simulation with the \( Q \)-learning update rule and an exploration policy. Figure 17.1 illustrates this process on the hex world problem.

Algorithm 17.2. The \( Q \)-learning update for model-free reinforcement learning, which can be applied to problems with unknown transition and reward functions. The update modifies \( Q \), which is a matrix of state-action values. This update function can be used together with an exploration strategy, such as \( \epsilon \)-greedy in the simulate function in algorithm 15.9.

17.3 Sarsa

Sarsa (algorithm 17.3) is an alternative to \( Q \)-learning.\(^5\) It derives its name from the fact that it uses \((s,a,r,s',a')\) to update the \( Q \) function at each step. It uses the actual next action \( a' \) to update \( Q \) instead of maximizing over all possible actions:

\[ Q(s,a) \leftarrow Q(s,a) + \alpha \left( r + \gamma Q(s',a') - Q(s,a) \right) \quad (17.11) \]

When run with a suitable exploration strategy, the \( a' \) above will converge to \( \arg \max_{a'} Q(s',a') \), which is what is used in the \( Q \)-learning update.

\(^5\) The maximization in this equation can introduce a bias. Algorithms like double \( Q \)-learning attempt to correct for this bias and can lead to better performance. H. van Hasselt, “Double Q-Learning,” in Advances in Neural Information Processing Systems (NIPS), 2010.

Suppose we want to apply Q-learning to an MDP problem \( \mathcal{P} \). We can construct an exploration policy, such as the \( \epsilon \)-greedy policy implemented in algorithm 16.6 from the previous chapter. The Q-learning model comes from algorithm 17.2 and the simulate function is implemented in algorithm 15.9.

\[
\begin{align*}
Q &= \text{zeros}(\text{length}(\mathcal{P}.S), \text{length}(\mathcal{P}.A)) \\
\alpha &= 0.2 \ # \text{learning rate} \\
\text{model} &= \text{QLearning}(\mathcal{P}.S, \mathcal{P}.A, \mathcal{P}.\gamma, Q, \alpha) \\
\epsilon &= 0.1 \ # \text{probability of random action} \\
\alpha &= 1.0 \ # \text{exploration decay factor} \\
\pi &= \text{EpsilonGreedyExploration}(\epsilon, \alpha) \\
k &= 20 \ # \text{number of steps to simulate} \\
s &= 1 \ # \text{initial state} \\
\text{simulate}(\mathcal{P}, \text{model}, \pi, k, s)
\end{align*}
\]

Example 17.2. An example of how to use an exploration strategy with Q-learning in simulation. The parameter settings are notional.
Sarsa is referred to as a type of on-policy reinforcement learning method because it attempts to directly estimate the value of the exploration policy as it follows it. In contrast, Q-learning is an off-policy method because it attempts to find the value of the optimal policy while following the exploration strategy. Although Q-learning and Sarsa both converge to an optimal strategy, the speed of convergence depends on the application. Sarsa is run on the hex world problem in figure 17.2.

```
mutable struct Sarsa
    S # state space (assumes 1:nstates)
    A # action space (assumes 1:nactions)
    γ # discount
    Q # action value function
    α # learning rate
    ℓ # last experience tuple (s,a,r)
end

lookahead(model::Sarsa, s, a) = model.Q[s,a]

function update!(model::Sarsa, s, a, r, s′)
    if model.ℓ != nothing
        γ, Q, α, ℓ = model.γ, model.Q, model.α, model.ℓ
        model.Q[ℓ.s,ℓ.a] += α*(ℓ.r + γ*Q[s,a] - Q[ℓ.s,ℓ.a])
    end
    model.ℓ = (s=s, a=a, r=r)
    return model
end
```

Algorithm 17.3. The Sarsa update for model-free reinforcement learning. We update the matrix $Q$ containing the state-action values, $\alpha$ is a constant learning rate, and $\ell$ is the last experience tuple.

Figure 17.2. Sarsa used to iteratively learn an action value function for the hex world problem in a manner otherwise identical to figure 17.1. We find that Sarsa is slower to converge to the true action value function.
17.4 Eligibility Traces

One of the disadvantages of Q-learning and Sarsa is that learning can be very slow, especially with sparse rewards. For example, suppose the environment has a single goal state that provides a large reward, and the reward is zero at all other states. After some amount of random exploration in the environment, we reach the goal state. Regardless of whether we use Q-learning or Sarsa, we only update the action value of the state immediately preceding the goal state. The values at all other states leading up to the goal remain at zero. A large amount of exploration is required to slowly propagate non-zero values to the remainder of the state space.

Q-learning and Sarsa can be modified to propagate reward backward to the states and actions leading to the source of the reward using eligibility traces. The credit is decayed exponentially so that states closer to the reward are assigned larger values. It is common to use $0 \leq \lambda \leq 1$ as the exponential decay parameter. Versions of Q-learning and Sarsa with eligibility traces are often called Q($\lambda$) and Sarsa($\lambda$).

A version of Sarsa($\lambda$) is implemented in algorithm 17.4. The algorithm maintains an exponentially decaying visit count $N(s,a)$ for all state-action pairs. When action $a$ is taken in state $s$, $N(s,a)$ is incremented by 1. The Sarsa temporal difference update is then partially applied to every state-action pair according to this decaying visit count.

Let $\delta$ denote the Sarsa temporal difference update:

$$\delta = r + \gamma Q(s',a') - Q(s,a)$$  \hspace{1cm} (17.12)

Every entry in the action value function is then updated according to:

$$Q(s,a) \leftarrow Q(s,a) + \alpha \delta N(s,a)$$  \hspace{1cm} (17.13)

The visit counts are then decayed using both the discount factor and the exponential decay parameter:

$$N(s,a) \leftarrow \gamma \lambda N(s,a)$$  \hspace{1cm} (17.14)

Although the impact of eligibility traces is especially pronounced in environments with sparse reward, the algorithm can speed learning in general environments where reward is more distributed.
mutable struct SarsaLambda
  𝒮  # state space (assumes 1:nstates)
  𝒜  # action space (assumes 1:nactions)
  γ  # discount
  Q  # action value function
  N  # trace
  α  # learning rate
  λ  # trace decay rate
  ℓ  # last experience tuple (s,a,r)
end

lookahead(model::SarsaLambda, s, a) = model.Q[s,a]

function update!(model::SarsaLambda, s, a, r, s')
  if model.ℓ != nothing
    γ, λ, Q, α, ℓ = model.γ, model.λ, model.Q, model.α, model.ℓ
    model.N[ℓ.s,ℓ.a] += 1
    δ = ℓ.r + γ*Q[s,a] - Q[ℓ.s,ℓ.a]
    for s in model.𝒮
      for a in model.𝒜
        model.Q[s,a] += α*δ*model.N[s,a]
        model.N[s,a] *= γ*λ
      end
    end
  else
    model.N[:,] .= 0.0
  end
  model.ℓ = (s=s, a=a, r=r)
  return model
end

Algorithm 17.4. The Sarsa(λ) update, which uses eligibility traces to propagate reward back in time to speed learning of sparse rewards. The matrix Q contains the state-action values, the matrix N contains exponentially decaying state-action visit counts, α is a constant learning rate, λ is an exponential decay parameter, and ℓ is the last experience tuple.
Special care must be taken when applying eligibility traces to an off-policy algorithm like Q-learning that attempts to learn the value of the optimal policy. Eligibility traces propagate back values obtained from an exploration policy. This mismatch can result in learning instabilities.

17.5 Reward Shaping

Especially in problems where the reward function is sparse, we may want to augment the reward function to help guide our learning. For example, if we are trying to reach a single goal state, we could supplement the reward function by an amount inversely proportional to the distance to the goal. Or, alternatively, we could add an additional penalty based on how far away we are from the goal. If we are playing chess, we might add a penalty to our reward function when we lose a piece, even though we only care about winning or losing the game at the end and not about individual pieces.

Modifications to the reward function during training by incorporating domain knowledge to speed training is known as reward shaping. Suppose rewards in our problem are generated according to \( R(s, a, s') \), allowing rewards to depend on the resulting state. We will use \( F(s, a, s') \) to represent our shaping function. During training, instead of using \( R(s, a, s') \) as our reward, we use \( R(s, a, s') + F(s, a, s') \).

Adding \( F(s, a, s') \) to our reward can, of course, change the optimal policy. We are often interested in shaping reward without changing what is optimal. It turns out that a policy that is optimal under the original reward remains optimal under the shaped reward if and only if

\[
F(s, a, s') = \gamma \beta(s') - \beta(s) \tag{17.15}
\]

for some potential function \( \beta(s) \).

17.6 Action Value Function Approximation

The algorithms we have discussed so far in this chapter have assumed discrete state and action spaces where the action value function can be stored in a lookup table. We can adapt our algorithms to use global value function approximation, allowing us to apply them to problems with large or continuous spaces and generalize from limited experience. Similar to the approach taken in chapter 8

---


in the context of a known model, we will use \( Q_\theta(s,a) \) to represent a parametric approximation of our action value function when the model is unknown.\(^{11}\)

To illustrate the concept, we will derive a version of Q-learning that uses our parametric approximation. We want to minimize the loss between our approximation and the optimal action value function \( Q^*(s,a) \), which we define to be\(^{12}\)

\[
\ell(\theta) = \frac{1}{2} \mathbb{E}_{(s,a) \sim \pi^*} [(Q^*(s,a) - Q_\theta(s,a))^2] \tag{17.16}
\]

The expectation is over the state-action pairs experienced when following the optimal policy \( \pi^* \).

A common approach to minimizing this loss is to use some form of gradient descent. The gradient of the loss is

\[
\nabla \ell(\theta) = - \mathbb{E}_{(s,a) \sim \pi^*} [(Q^*(s,a) - Q_\theta(s,a)) \nabla_\theta Q_\theta(s,a)] \tag{17.17}
\]

We typically choose parametric representations of the action value function that are differentiable and where \( \nabla_\theta Q_\theta(s,a) \) is easy to compute, such as linear or neural network representations. If we apply gradient descent,\(^{13}\) our update rule is

\[
\theta \leftarrow \theta + \alpha \mathbb{E}_{(s,a) \sim \pi^*} [(Q^*(s,a) - Q_\theta(s,a)) \nabla_\theta Q_\theta(s,a)] \tag{17.18}
\]

where \( \alpha \) is our step factor or learning rate. We can approximate the update rule above using samples of our state-action \((s,a)\) pairs as we experience them:

\[
\theta \leftarrow \theta + \alpha (Q^*(s,a) - Q_\theta(s,a)) \nabla_\theta Q_\theta(s,a) \tag{17.19}
\]

Of course, we cannot compute equation (17.19) directly because that would require knowing the optimal policy, which is precisely what we are attempting to find. Instead, we attempt to estimate it from our observed transition and our action value approximation:

\[
Q^*(s,a) \approx r + \gamma \max_{a'} Q_\theta(s',a') \tag{17.20}
\]

which results in the following update rule:

\[
\theta \leftarrow \theta + \alpha (r + \gamma \max_{a'} Q_\theta(s',a') - Q_\theta(s,a)) \nabla_\theta Q_\theta(s,a) \tag{17.21}
\]
This update is implemented in algorithm 17.5 with the addition of a scaled gradient step (algorithm 12.2), which is often needed to ensure that the gradient steps do not become too large. Example 17.3 shows how to use this update with a linear action value approximation. Figure 17.3 demonstrates this algorithm on the mountain car problem.

```
struct GradientQLearning
    A # action space (assumes 1:nactions)
    γ # discount
    Q # parameterized action value function Q(θ,s,a)
    ∇Q # gradient of action value function
    θ # action value function parameter
    α # learning rate
end

function lookahead(model::GradientQLearning, s, a)
    return model.Q(model.θ, s,a)
end

function update!(model::GradientQLearning, s, a, r, s′)
    A, γ, Q, θ, α = model.A, model.γ, model.Q, model.θ, model.α
    u = maximum(Q(θ,s′,a′) for a′ in A)
    Δ = (r + γ*u - Q(θ,s,a)) * model.∇Q(θ,s,a)
    θ[:] += α * scale_gradient(Δ, 1)
    return model
end
```

Algorithm 17.5. The Q-learning update with action value function approximation. With each new experience tuple s, a, r, s′, we update our vector θ with constant learning rate α. Our parameterized action value function is given by Q(θ,s,a) and its gradient is ∇Q(θ,s,a).

17.7 Experience Replay

A major challenge of using global function approximation with reinforcement learning is catastrophic forgetting. For example, we might initially discover that our particular policy brings us to a low-reward region of the state space. We then refine our policy to avoid that area. However, after some amount of time, we may forget why it was important to avoid that region of the state space, and we may risk reverting back to a poorly performing policy.

Catastrophic forgetting can be mitigated with experience replay, where a fixed number of the most recent experience tuples are stored across training iterations. A batch of tuples are sampled uniformly from this replay memory to remind us to avoid strategies that we have already discovered are poor. Variations of this approach include prioritizing experiences. T. Schaul, J. Quan, I. Antonoglou, and D. Silver, “Prioritized Experience Replay,” in International Conference on Learning Representations (ICLR), 2016.
We are interested in applying $Q$-learning with a linear action value approximation to the simple regulator problem with $\gamma = 1$. Our action value approximation is $Q_\theta(s,a) = \theta^T \beta(s,a)$, where our basis function is

$$\beta(s,a) = [s, s^2, a, a^2, 1]$$

With this linear model,

$$\nabla_\theta Q_\theta(s,a) = \beta(s,a)$$

We can implement this as follows for problem $\mathcal{P}$:

$\beta(s,a) = [s, s^2, a, a^2, 1]$  
$Q(\theta, s, a) = \text{dot}(\theta, \beta(s,a))$  
$\nabla Q(\theta, s, a) = \beta(s,a)$  
$\theta = [0.1, 0.2, 0.3, 0.4, 0.5]$  
$\alpha = 0.5$ # initial parameter vector  
$\alpha = 0.5$ # learning rate  
$\text{model} = \text{GradientQLearning}(\mathcal{P}.\mathcal{A}, \mathcal{P}.\mathcal{V}, \mathcal{Q}, \nabla Q, \theta, \alpha)$  
$\epsilon = 0.1$ # probability of random action  
$\alpha = 1.0$ # exploration decay factor  
$\pi = \text{EpsilonGreedyExploration}(\epsilon, \alpha)$  
$k = 20$ # number of steps to simulate  
$s = 0.0$ # initial state  
$\text{simulate}(\mathcal{P}, \text{model}, \pi, k, s)$

Example 17.3. An example of how to use an exploration strategy with $Q$-learning with action value function approximation in simulation. The parameter settings are notional.
17.7. EXPERIENCE REPLAY

Figure 17.3. A utility function and policy obtained using linear approximation Q-learning applied to the mountain car problem. The basis functions are polynomials over position and velocity up to degree eight and are each duplicated three times for the three actions. The received reward is the reward received by an agent when run using a greedy policy with the approximate value function.
from equation (17.21) is modified to become

$$\theta \leftarrow \theta + \alpha \frac{1}{m_{\text{grad}}} \sum_i (r^{(i)} + \gamma \max_{a'} Q_\theta(s'^{(i)}, a') - Q_\theta(s^{(i)}, a^{(i)})) \nabla_\theta Q_\theta(s^{(i)}, a^{(i)})$$

(17.22)

where $s^{(i)}, a^{(i)}, r^{(i)}, s'^{(i)}$ is the $i$th experience tuple in a random batch of size $m_{\text{grad}}$.

Experience replay allows experience tuples to contribute to learning multiple times, thereby increasing data efficiency. Furthermore, sampling uniformly at random from the replay memory breaks apart otherwise correlated sequences that are obtained from rollouts, thereby reducing the variance of the gradient estimate. Experience replay stabilizes the learning process by retaining information from previous policy parameterizations.

Algorithm 17.6 shows how to incorporate experience replay into $Q$-learning with action value function approximation. Example 17.4 shows how to apply this approach to a simple regulator problem.

### 17.8 Summary

- Model-free methods seek to directly learn an action value function rather than transition and reward models.

- Simple techniques can be used to incrementally learn a mean from sequential updates.

- The $Q$-learning algorithm incrementally learns an action value function using an approximation of the Bellman equation.

- In contrast with $Q$-learning, Sarsa uses the action taken by the exploration policy rather than maximizing over all subsequent actions in its update.

- Eligibility traces can speed learning by propagating sparse rewards through the state-action space.

- $Q$-learning can be applied to approximate value functions using stochastic gradient descent.

- The catastrophic forgetting experienced by $Q$-learning and Sarsa can be mitigated using experience replay, which reuses past experience tuples.
Algorithm 17.6. Q-learning with function approximation and experience replay. The update depends upon a parameterized policy $Q(\theta, s, a)$ and gradient $\nabla Q(\theta, s, a)$. It updates the parameter vector $\theta$ and the circular memory buffer provided by DataStructures.jl. It updates $\theta$ every $m$ steps using a gradient estimated from $m_{-}\text{grad}$ samples from the buffer.
Suppose we want to add experience replay to example 17.3. When constructing the model, we need to provide a replay buffer with the desired capacity:

```
capacity = 100  # maximum size of the replay buffer
ExperienceTuple = Tuple{Float64,Float64,Float64,Float64}
M = CircularBuffer{ExperienceTuple}(capacity)  # replay buffer
m_grad = 20  # batch size
model = ReplayGradientQLearning(𝓚.𝓐, 𝓚.γ, Q, ∇Q, θ, α, M, m, m_grad)
```

We can vary the number of steps between gradient updates \( m \) and the depth of each simulation \( d \). In the plot below, we limit all training runs to \( md = 30 \) experience tuples with each iteration. The plot indicates that rollouts to a sufficient depth are necessary for training to succeed. In addition, very few rollouts to an excessive depth do not perform as well as a moderate number of rollouts to a moderate depth.

```
0 20 40 60 80 100
−100 −80 −60 −40 −20 0

iteration

expected value

\( m = 1, d = 30 \)
\( m = 2, d = 15 \)
\( m = 3, d = 10 \)
\( m = 5, d = 6 \)
\( m = 10, d = 3 \)
```

Example 17.4. Application of experience replay to the simple regulator problem with Q-learning and action value approximation.
17.9 Exercises

Exercise 17.1. Given the following set of samples, perform incremental estimation of the mean twice; once using a learning rate of $\alpha = 0.1$ and once using a learning rate of $\alpha = 0.5$. In both, use an initial mean equal to the first sample.

$$x^{(1:5)} = \{1.0, 1.8, 2.0, 1.6, 2.2\}$$

Solution: We set the mean at the first iteration equal to the first sample and proceed to incrementally estimate the mean using equation (17.6):

\[
\hat{x}_1 = 1.0 \\
\hat{x}_2 = 1.0 + 0.1(1.8 - 1.0) = 1.08 \\
\hat{x}_3 = 1.08 + 0.1(2.0 - 1.08) = 1.172 \\
\hat{x}_4 = 1.172 + 0.1(1.6 - 1.172) \approx 1.215 \\
\hat{x}_5 = 1.215 + 0.1(2.2 - 1.215) \approx 1.313
\]

\[
\hat{x}_1 = 1.0 \\
\hat{x}_2 = 1.0 + 0.5(1.8 - 1.0) = 1.4 \\
\hat{x}_3 = 1.4 + 0.5(2.0 - 1.4) = 1.7 \\
\hat{x}_4 = 1.7 + 0.5(1.6 - 1.7) = 1.65 \\
\hat{x}_5 = 1.65 + 0.5(2.2 - 1.65) = 1.925
\]

Exercise 17.2. Following the previous exercise, suppose once we have estimated the mean with five samples for both methods, we are provided with a single additional sample $x^{(6)}$ that we will use as the final sample in estimating our mean. Which of the two incremental estimation methods (i.e., $\alpha = 0.1$ or $\alpha = 0.5$) would be preferable?

Solution: While we do not know what the sample would be or what the underlying mean of the process is, we would likely prefer the second incrementally estimated mean that uses $\alpha = 0.5$. Since we only have one sample left, the first learning rate is too small to considerably change the mean, while the second learning rate is large enough to be responsive, without neglecting the past samples. Consider two cases:

1. If we assume the next sample is approximately equal to the incremental mean of all previous samples, then we have $x^{(6)} \approx \hat{x}_5$. Thus, performing an incremental update of the mean yields no change to our estimate. We have $\hat{x}_6 \approx 1.313$ for a learning rate of 0.1, and we have $\hat{x}_6 = 1.925$ for a learning rate of 0.5.

2. If we assume the next sample is approximately equal to the exact mean of all previous samples, then we have $x^{(6)} \approx 1.72$. The update using a learning rate of 0.1 yields $\hat{x}_6 \approx 1.354$, while the update using a learning rate of 0.5 yields $\hat{x}_6 \approx 1.823$.

In both of these cases, supposing the next sample is equal to the mean all previous samples, then the estimate using a learning rate of 0.5 is more accurate.

Exercise 17.3. Using incremental estimation of the mean, derive an update rule for a value function $U(s)$ for experience tuples of the form $(s, r, s')$. Is the resulting algorithm model-based or model-free?

Solution: The incremental estimation of the mean equation is

$$\hat{x} \leftarrow \hat{x} + \alpha(x - \hat{x})$$
Our value function can be written in the familiar form

\[ U(s) = \max_a \left( R(s, a) + \gamma \sum_{s'} T(s' | s, a) U(s') \right) \]

Applying incremental estimation to the value function where we receive experience tuples of the form \((s_t, r_t, s_{t+1})\), we can derive an update rule for the value function.

\[
U(s) \leftarrow U(s) + \alpha \left[ \max_a \left( R(s, a) + \gamma \sum_{s'} T(s' | s, a) U(s') \right) - U(s) \right]
\]

\[
U(s) \leftarrow U(s) + \alpha \left[ \max_a (r + \gamma U(s')) - U(s) \right]
\]

This algorithm is known as temporal difference (TD) learning. While the update rule is model-free, extracting an action or policy requires a transition and reward model:

\[
a^* = \pi^*(s) = \arg \max_a \left( R(s, a) + \gamma \sum_{s'} T(s' | s, a) U(s) \right)
\]

**Exercise 17.4.** What is the complexity of Q-learning if we interact with the environment for \(d\) time steps? What is the complexity of Sarsa if we interact with the environment for \(d\) time steps?

**Solution:** For Q-learning, our update rule is:

\[
Q(s, a) \leftarrow Q(s, a) + \alpha \left( r + \gamma \max_{a'} Q(s', a') - Q(s, a) \right)
\]

At each time step, we must perform a maximization over actions, so for \(d\) time steps, the complexity of Q-learning is \(O(d|A|)\). For Sarsa, our update rule is:

\[
Q(s, a) \leftarrow Q(s, a) + \alpha \left( r + \gamma Q(s', a') - Q(s, a) \right)
\]

(17.23)

At each time step, unlike Q-learning, we do not have to perform a maximization over actions, so for \(d\) time steps, the complexity of Sarsa is simply \(O(d)\).

**Exercise 17.5.** Is the computational complexity of Sarsa per experience tuple \((s_t, a_t, r_t, s_{t+1})\) more or less than that of Sarsa(\(\lambda\))?  

**Solution:** For Sarsa, our update rule is:

\[
Q(s, a) \leftarrow Q(s, a) + \alpha \left( r + \gamma Q(s', a') - Q(s, a) \right)
\]

(17.24)
So, for each experience tuple, we have \(O(1)\) complexity. For Sarsa(\(\lambda\)), our update rules are:

\[
\begin{align*}
\delta &\leftarrow r_t + \gamma Q(s_{t+1},a_{t+1}) - Q(s_t,a_t) \\
N(s_t,a_t) &\leftarrow N(s_t,a_t) + 1 \\
Q(s,a) &\leftarrow Q(s,a) + \alpha \delta N(s,a) \quad \text{for all } s,a \\
N(s,a) &\leftarrow \gamma \lambda N(s,a) \quad \text{for all } s,a
\end{align*}
\]

For each experience tuple, we need to compute \(\delta\) and increment the visit count at \((s_t,a_t)\), which are both \(O(1)\). However, we need to update both the action value function and the visit counts for all states and actions, which are both \(O(|\mathcal{S}| |\mathcal{A}|)\). Thus, the computational complexity per experience tuple is greater for Sarsa(\(\lambda\)). However, Sarsa(\(\lambda\)) often converges using fewer experience tuples.

**Exercise 17.6.** What is the behavior of \(Q(\lambda)\) in the limit as \(\lambda \to 0\)? What is the behavior of \(Q(\lambda)\) in the limit as \(\lambda \to 1\)?

**Solution:** For \(Q(\lambda)\), we perform the following update rules:

\[
\begin{align*}
\delta &\leftarrow r_t + \gamma \max_{a'} Q(s_{t+1},a') - Q(s_t,a_t) \\
N(s_t,a_t) &\leftarrow N(s_t,a_t) + 1 \\
Q(s,a) &\leftarrow Q(s,a) + \alpha \delta N(s,a) \quad \text{for all } s,a \\
N(s,a) &\leftarrow \gamma \lambda N(s,a) \quad \text{for all } s,a
\end{align*}
\]

In the limit as \(\lambda \to 0\), for our first iteration, we compute the temporal difference error \(\delta\) and we increment the visit count \(N(s_t,a_t)\). In the action value function update, the only non-zero \(N(s,a)\) is at \((s_t,a_t)\), so we perform \(Q(s_t,a_t) = Q(s_t,a_t) + \alpha \delta N(s_t,a_t)\). Finally, we reset all the visit counts to zero. From this, we can see that in the limit as \(\lambda \to 0\), we have no eligibility traces and we are performing a straightforward Q-learning update.

In the limit as \(\lambda \to 1\), our visit counts will accumulate and we have full eligibility traces, which will spread the reward the most possible over previously visited state-action pairs.

**Exercise 17.7.** Compute \(Q(s,a)\) using Sarsa(\(\lambda\)) after following the trajectory

\((s_1,a_R,0,s_2,a_R,0,s_3,a_L,10,s_2,a_R,4,s_1,a_R)\)

Use \(\alpha = 0.5\), \(\lambda = 1\), and initial action value function and visit counts equal to zero everywhere. Assume \(\mathcal{S} = \{s_1,s_2,s_3,s_4\}\) and \(\mathcal{A} = \{a_L,a_R\}\).

**Solution:** The Sarsa(\(\lambda\)) update rules are

\[
\begin{align*}
\delta &\leftarrow r_t + \gamma Q(s_{t+1},a_{t+1}) - Q(s_t,a_t) \\
N(s_t,a_t) &\leftarrow N(s_t,a_t) + 1 \\
Q(s,a) &\leftarrow Q(s,a) + \alpha \delta N(s,a) \quad \text{for all } s,a \\
N(s,a) &\leftarrow \gamma \lambda N(s,a) \quad \text{for all } s,a
\end{align*}
\]
For the first experience tuple, we have $\delta = 0 + 0.9 \times 0 - 0 = 0$, we increment the visit count at $N(s_1, a_R)$, the action value function does not change since $\delta = 0$, and we update our counts. After completing this, we have:

\[
\begin{array}{c|cccc}
Q(s, a) & s_1 & s_2 & s_3 & s_4 \\
\hline
a_L & 0 & 0 & 0 & 0 \\
a_R & 0 & 0 & 0 & 0 \\
\end{array}
\]  \quad \begin{array}{c|cccc}
N(s, a) & s_1 & s_2 & s_3 & s_4 \\
\hline
a_L & 0 & 0 & 0 & 0 \\
a_R & 0.9 & 0 & 0 & 0 \\
\end{array}
\]

For the second experience tuple, we have $\delta = 0$, we increment the visit count at $N(s_2, a_R)$, the action value function does not change since $\delta = 0$, and we update our counts. After completing this, we have:

\[
\begin{array}{c|cccc}
Q(s, a) & s_1 & s_2 & s_3 & s_4 \\
\hline
a_L & 0 & 0 & 0 & 0 \\
a_R & 0 & 0 & 0 & 0 \\
\end{array}
\]  \quad \begin{array}{c|cccc}
N(s, a) & s_1 & s_2 & s_3 & s_4 \\
\hline
a_L & 0 & 0 & 0 & 0 \\
a_R & 0.81 & 0.9 & 0 & 0 \\
\end{array}
\]

For the third experience tuple, we have $\delta = 10$, we increment the visit count at $N(s_3, a_L)$, we update the action value function, and we update our counts. After completing this, we have:

\[
\begin{array}{c|cccc}
Q(s, a) & s_1 & s_2 & s_3 & s_4 \\
\hline
a_L & 0 & 0 & 5 & 0 \\
a_R & 4.05 & 4.5 & 0 & 0 \\
\end{array}
\]  \quad \begin{array}{c|cccc}
N(s, a) & s_1 & s_2 & s_3 & s_4 \\
\hline
a_L & 0 & 0 & 0.9 & 0 \\
a_R & 0.729 & 0.81 & 0 & 0 \\
\end{array}
\]

For the fourth experience tuple, we have $\delta = 4 + 0.9 \times 4.05 - 4.5 = 3.145$, we increment the visit count at $N(s_2, a_R) = 0.81 + 1 = 1.81$, we update the action value function, and we update our counts. After completing this, we have:

\[
\begin{array}{c|cccc}
Q(s, a) & s_1 & s_2 & s_3 & s_4 \\
\hline
a_L & 0 & 0 & 6.415 & 0 \\
a_R & 5.196 & 7.346 & 0 & 0 \\
\end{array}
\]  \quad \begin{array}{c|cccc}
N(s, a) & s_1 & s_2 & s_3 & s_4 \\
\hline
a_L & 0 & 0 & 0.81 & 0 \\
a_R & 0.656 & 1.629 & 0 & 0 \\
\end{array}
\]
18 Imitation Learning

Previous chapters have assumed that a reward function is either known or that rewards are received while interacting with the environment. For some applications, it may be easier for an expert to demonstrate the desired behavior rather than specifying a reward function. This chapter discusses algorithms for imitation learning, where the desired behavior is learned from expert demonstration. We will cover a variety of methods ranging from very simple likelihood-maximization methods to more complicated iterative methods that involve reinforcement learning.\(^1\)

18.1 Behavioral Cloning

A simple form of imitation learning is to treat it as a supervised learning problem. This method, called behavioral cloning,\(^2\) trains a stochastic policy \(\pi_\theta\) parameterized by \(\theta\) to maximize the likelihood of actions from a dataset \(D\) of expert state-action pairs:

\[
\max_\theta \prod_{(s,a) \in D} \pi_\theta(a \mid s) \quad (18.1)
\]

As done in earlier chapters, we can transform the maximization over the product over \(\pi_\theta(a \mid s)\) to a sum over \(\log \pi_\theta(a \mid s)\).

Depending on how we want to represent the conditional distribution \(\pi_\theta(a \mid s)\), we may compute the maximum likelihood estimate of \(\theta\) analytically. For example, if we use a discrete conditional model (section 2.4), \(\theta\) would consist of all of the counts \(N(s,a)\) from \(D\) and \(\pi_\theta(a \mid s) = N(s,a) / \sum_a N(s,a)\). Example 18.1 applies a discrete conditional model to data from the mountain car problem.

If we have a factored representation of our policy, we can use a Bayesian network to represent the joint distribution over our state and action variables. Figure 18.1 shows an example. We can learn both the structure (chapter 5) and

---


Consider using behavioral cloning on expert demonstrations for the mountain car problem (appendix F.4). We are given 10 rollouts from an expert policy. We fit a conditional distribution and plot the results. The continuous trajectories were discretized with 10 bins each for position and for speed.

\[ P(a = -1 \mid s) \quad P(a = 0 \mid s) \]

The state space is not fully covered by expert demonstrations, which is typical for imitation learning problems. The resulting policy may perform well when used in regions with coverage, but it assigns a uniform distribution to actions in regions without coverage. Even if we start in a region with coverage, we may transition to regions without coverage due to stochasticity in the environment.

Example 18.1. A demonstration of behavioral cloning applied to the mountain car problem. The light blue regions are areas without training data, resulting in poor policy performance when the agent encounters those states.
parameters (chapter 4) from the data $D$. Given the current state, we can then infer the distribution over actions using one of the inference algorithms discussed earlier (chapter 3).

![Figure 18.1. Bayesian networks can be used to represent a joint distribution over the state and action variables. We can apply an inference algorithm to generate a distribution over actions given the current values of the state variables.](image)

We can use many other representations for $\pi_\theta$. For example, we might want to use a neural network, where the input corresponds to the values of the state variables and the output corresponds to parameters of a distribution over the action space. If our representation is differentiable, which is the case with neural networks, we can attempt to optimize equation (18.1) using gradient ascent. This approach is implemented in algorithm 18.1.

```plaintext
struct BehavioralCloning
    α  # step size
    k_max # number of iterations
    ∇logπ # log likelihood gradient
end

function optimize(M::BehavioralCloning, D, θ)
    α, k_max, ∇logπ = M.α, M.k_max, M.∇logπ
    for k in 1:k_max
        ∇ = mean(∇logπ(θ, a, s) for (s,a) in D)
        θ += α*∇
    end
    return θ
end
```

Algorithm 18.1. A method for learning a parameterized stochastic policy from expert demonstrations in the form of a set of state-action tuples $D$. The policy parameterization vector $\theta$ is iteratively improved by maximizing the log likelihood of the actions given the states. Behavioral cloning requires a step size $\alpha$, an iteration count $k_{\text{max}}$, and a log likelihood gradient $\nabla \log \pi$.

The closer the expert demonstrations are to optimal, the better the resulting behavioral cloning policy will perform. However, behavioral cloning suffers from cascading errors. As discussed in example 18.2, small inaccuracies compound during a rollout and eventually lead to states that are poorly represented in the training data, thereby leading to worse decisions, and ultimately to invalid or unseen situations. Though behavioral cloning is attractive due to its simplicity, cascading errors cause the method to perform poorly on many problems, especially when policies must be used for long time horizons.

---

Consider applying behavioral cloning to train a policy for driving an autonomous race car. A human race car driver provides expert demonstrations. Being an expert, the driver never drifts onto the grass or too close to a railing. A model trained with behavioral cloning would have no information to go off of when near a railing or when drifting onto the grass, and would not know how to recover.

### 18.2 Dataset Aggregation

One way to address the problem of cascading errors is to correct a trained policy using additional expert input. *Sequential interactive demonstration* methods alternate between collecting data from an expert in situations generated by a trained policy and using this data to improve this policy.

One type of sequential interactive demonstration method is called *dataset aggregation (DAgger)* (algorithm 18.2). It starts by training a stochastic policy using behavioral cloning. The policy is then used to run several rollouts from an initial state distribution $b$, which are then given to an expert to provide the correct actions for each state. The new data is aggregated with the previous dataset and a new policy is trained. Example 18.3 illustrates this process.

These interactive demonstrations iteratively builds a dataset covering the regions of the state space that the agent is likely to encounter, based on previous learning iterations. With each iteration, newly added examples compose a smaller fraction of the dataset, thereby leading to smaller policy changes. While sequential interactive demonstration can work well in practice, it is not guaranteed to converge. It can be shown that mixing in influence from the expert policy can guarantee convergence, which is the subject of the next section.

### 18.3 Stochastic Mixing Iterative Learning

Sequential interactive methods can also iteratively build up a policy by stochastically mixing in newly trained policies. One such method is *stochastic mixing iterative learning (SMiLe)* (algorithm 18.3). It uses behavioral cloning in every iteration but mixes the newly trained policy in with the previous ones.

We start with the expert policy, $\pi^{(1)} = \pi_E$. In each iteration, we execute...

---

Example 18.2. A brief example of the generalization issue inherent to behavioral cloning approaches.

---

4 S. Ross, G. J. Gordon, and J. A. Bagnell, “A Reduction of Imitation Learning and Structured Prediction to No-Regret Online Learning,” in *International Conference on Artificial Intelligence and Statistics (AISTATS)*, vol. 15, 2011.


6 We do not have an explicit representation of $\pi_E$. Evaluating $\pi_E$ requires interactively querying the expert as done in the previous section.
Algorithm 18.2. The DAgger method of dataset aggregation for learning a stochastic parameterized policy from expert demonstrations. This method takes an initial dataset of state-action tuples $D$, a stochastic parameterized policy $\pi_\theta(\theta, s)$, an MDP $\mathcal{P}$ that defines a transition function, and an initial state distribution $b$. Behavioral cloning, algorithm 18.1, is used in each iteration to improve the policy.

An expert policy $\pi_E$ labels trajectories sampled from the latest learned policy to augment the dataset. The original paper generated trajectories by stochastically mixing in the expert policy. This implementation is thus the original DAgger with an extreme mixing value of zero.

In practice an expert policy would not exist, and calls to this policy would be replaced with queries to a human expert.
Consider using dataset aggregation to train a policy on the mountain car problem where the reward is not observed. We use an expert policy that accelerates in the direction of travel. In this example we train a policy using the features:

$$f(s) = [1[v > 0], 1[v < 0], x, x^2, v, v^2, xv]$$

where $x$ and $v$ are the position and speed of the car.

Trajectories begin in red and proceed to blue. In the first iteration, the agent behaves randomly, unable to make progress toward the goal ($x \geq 0.6$). With additional iterations, the agent learns to mimic the expert policy of accelerating in the direction of travel. This behavior is apparent in the new trajectories, which spiral outward, and the policy, which assigns high likelihood to $a = 1$ when $v > 0$ and $a = -1$ when $v < 0$.
the latest policy \( \pi^{(k)} \) to generate a new dataset, querying the expert to provide the correct actions. Behavioral cloning is applied only to this new dataset to train a new component policy \( \hat{\pi}^{(k)} \). This component policy is mixed together with component policies from the previous iterations to produce a new policy \( \pi^{(k+1)} \).

The mixing of component policies to generate \( \pi^{(k+1)} \) is governed by a mixing scalar \( \beta \in (0, 1) \). The probability of acting according to the expert policy is \((1 - \beta)^k\), and the probability of acting according to \( \hat{\pi}^{(i)} \) is \( \beta(1 - \beta)^{i-1} \). This scheme assigns more weight to older policies under the hypothesis that older policy components were trained on the states more likely to be encountered.\(^7\) With each iteration, the probability of acting according to the original expert policy decays to zero. The mixing scalar is typically small such that the agent does not abandon the expert’s policy too quickly. Example 18.4 demonstrates this approach on mountain car.

18.4 Maximum Margin Inverse Reinforcement Learning

In many application settings, we do not have an expert that can be interactively queried, but instead have a batch of expert demonstration trajectories. We will assume that the expert demonstration data \( D \) consists of \( m \) trajectories. Each trajectory \( \tau \) in \( D \) involves a rollout to depth \( d \). In inverse reinforcement learning, we assume that the expert is optimizing some unknown reward function. From \( D \), we attempt to derive that reward function. With that reward function, we can use the methods discussed in prior chapters to derive an optimal policy.

There are different approaches to inverse reinforcement learning. We generally need to define a parameterization of the reward function. A common assumption is that this parameterization is linear with \( R_\phi(s, a) = \phi^\top \beta(s, a) \), where \( \beta(s, a) \) is a feature vector and \( \phi \) is a vector of weightings. In this section, we will focus on an approach known as maximum margin inverse reinforcement learning,\(^8\) where the features are assumed to be binary. Since optimal policies remain optimal with positive scaling of the reward function, this method additionally constrains the weight vector such that \( \| \phi \|_2 \leq 1 \). The expert data activates each binary feature with different frequencies, perhaps pursuing some and avoiding others. This approach attempts to learn this pattern of activation and trains an agent to mimic these activation frequencies.

\(^7\) In SMILe, we are acting according to our latest learned policy. We expect that this learned policy will match the expert fairly well and to primarily mispredict when we deviate from the expert policy. The learned component policies generally only need to make smaller and smaller contributions with each iteration to make up the difference in what has not already been learned.

\(^8\) P. Abbeel and A. Y. Ng, “Apprenticeship Learning via Inverse Reinforcement Learning,” in International Conference on Machine Learning (ICML), 2004.
Algorithm 18.3. The SMILe algorithm for training a stochastic parameterized policy from expert demonstrations for an MDP $\mathcal{P}$. SMILe successively mixes in new component policies with smaller and smaller weight, while simultaneously reducing the probability of acting according to the expert policy. The method returns the probabilities $P_s$ and parameterizations $\theta_s$ for the component policies.
Consider using SMILe to train a policy on the mountain car problem where the reward is not observed. We use the same features that were used for dataset aggregation in example 18.3. Both dataset aggregation and SMILe receive a new expert-labeled dataset with each iteration. Instead of accumulating a larger dataset of expert-labeled data, SMILe trains a new policy component using only the most recent data, mixing the new policy component with the previous policy components.

Example 18.4. An example of using SMILe to learn a policy for the mountain car problem. In contrast with dataset aggregation in example 18.3, SMILe mixes the expert into the policy during rollouts. This expert component, whose influence wanes with each iteration, causes the initial rollouts to better progress toward the goal.
An important part of this algorithm involves reasoning about the expected return under a policy $\pi$ for a weighting $\phi$ and initial state distribution $b$:

$$
\mathbb{E}_{s \sim b} [U(s)] = \mathbb{E}_\tau \left[ \sum_{k=1}^{d} \gamma^{k-1} R_{\phi^{(k)}}(s^{(k)}, a^{(k)}) \right] = \mathbb{E}_\tau \left[ \sum_{k=1}^{d} \gamma^{k-1} \phi^{(k)} \beta(s^{(k)}, a^{(k)}) \right] = \phi^\top \left( \mathbb{E}_\tau \left[ \sum_{k=1}^{d} \gamma^{k-1} \beta(s^{(k)}, a^{(k)}) \right] \right) = \phi^\top \mu_{\pi}
$$

(18.2) (18.3) (18.4) (18.5)

where $\tau$ corresponds to trajectories generated by $\pi$ to depth $d$. Here, we introduce the feature expectations $\mu_{\pi}$, which is the expected discounted accumulated feature values. These feature expectations can be estimated from $m$ rollouts, as implemented in algorithm 18.4.

```python
struct InverseReinforcementLearning
    \# problem
    P
    \# initial state distribution
    b
    \# depth
    d
    \# number of samples
    m
    \# parameterized policy
    \pi
    \# binary feature mapping
    \beta
    \# expert feature expectations
    \mu_E
    \# reinforcement learning method
    RL
    \# tolerance
    \epsilon
end

function feature_expectations(M::InverseReinforcementLearning, \pi)
    \P, \b, \m, \d, \beta, \gamma = M.P, M.b, M.m, M.d, M.beta, M.gamma
    \mu(\tau) = \sum(\gamma^{(k-1)} \beta(s, a)) \text{ for } (k, (s, a)) \text{ in enumerate}(\tau)
    \tau_s = [\text{simulate}(\P, \text{rand}(\b), \pi, \d) \text{ for } i \text{ in } 1:m]
    \text{return mean}(\mu(\tau) \text{ for } \tau \text{ in } \tau_s)
end
```


We can use the expert demonstrations to estimate the expert feature expectations $\mu_E$, and we want to find a policy that matches these feature expectations as closely as possible. At the first iteration, we begin with a randomized policy $\pi^{(1)}$ and estimate its feature expectations that we denote $\mu^{(1)}$. At iteration $k$, we find a new $\phi^{(k)}$, corresponding to a reward function $R_{\phi^{(k)}}(s, a) = \phi^{(k)}^\top \beta(s, a)$, such
that the expert outperforms all previously found policies by the greatest margin $t$:

$$\begin{align*}
\max_{t, \phi} & \quad t \\
\text{subject to} & \quad \phi^T \mu_E \geq \phi^T \mu^{(i)} + t \quad \text{for} \quad i = 1, \ldots, k - 1 \\
& \quad \|\phi\|_2 \leq 1
\end{align*}$$

Equation (18.6) is a quadratic program that can be easily solved. We then solve for a new policy $\pi^{(k)}$ using the reward function $R(s, a) = \phi^{(k)} \beta(s, a)$ and produce a new vector of feature expectations. Figure 18.2 illustrates this margin maximization process.

We iterate until the margin is sufficiently small with $t \leq \epsilon$. At convergence, we can solve for a mixed policy that attempts to have feature expectations as close as possible to that of the expert policy:

$$\begin{align*}
\min_{\lambda} & \quad \|\mu_E - \mu_{\lambda}\|_2 \\
\text{subject to} & \quad \lambda \geq 0 \\
& \quad \|\lambda\|_1 = 1
\end{align*}$$

where $\mu_{\lambda} = \sum \lambda_i \mu^{(i)}$. The mixture weights $\lambda$ combine the policies found at each iteration. With probability $\lambda_i$, we follow policy $\pi^{(i)}$. Inverse reinforcement learning is implemented in algorithm 18.5.

18.5 Maximum Entropy Inverse Reinforcement Learning

The inverse reinforcement learning approach from the previous section is underspecified, meaning there are often multiple policies that can produce the same feature expectations as the expert demonstrations. This section introduces maximum entropy inverse reinforcement learning, which avoids this ambiguity by preferring the policy that results in the distribution over trajectories that has maximum entropy (appendix A.8). The problem can be transformed into one of finding the best reward function parameters $\phi$ in a maximum likelihood estimation problem given the expert data $\mathcal{D}$.

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2021-03-07 23:12:00-08:00, revision 80bf4d6, comments to bugs@algorithmsbook.com

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Figure 18.2. A geometric visualization of three example iterations of the maximum-margin inverse reinforcement learning algorithm, going top to bottom. In each iteration, the new weight vector points in the direction perpendicular to the hyperplane that separates the expert feature expectation vector from the those of the previous policy with the largest possible margin. The margin decreases with each iteration.
Algorithm 18.5. Maximum margin inverse reinforcement learning, which computes a mixed policy whose feature expectations match those of given expert demonstrations. We use JuMP.jl to solve our constrained optimization problems. This implementation requires that the provided reinforcement learning struct has a weight vector \( \phi \) that can be updated with new values. The method returns the stochastic weightings \( \lambda \) and parameterizations \( \theta s \) for the component policies.

```julia
function calc_weighting(M::InverseReinforcementLearning, \( \mu s \))
    \( \mu E = M.\mu E \)
    \( \mu E \)
    \( k = \text{length}(\mu E) \)
    \( \text{model} = \text{Model}(\text{Ipopt.Optimizer}) \)
    \( \text{@variable}(\text{model}, t) \)
    \( \text{@variable}(\text{model}, \phi[1:k] \geq 0) \)
    \( \text{@objective}(\text{model}, \text{Max}, t) \)
    for \( \mu \) in \( \mu s \)
        \( \text{@constraint}(\text{model}, \phi \cdot \mu E \geq \phi \cdot \mu + t) \)
    end
    \( \text{@constraint}(\text{model}, \phi \cdot \phi \leq 1) \)
    \( \text{optimize!}(\text{model}) \)
    \( \text{return} \ (\text{value}(t), \text{value}(\phi)) \)
end

function calc_policy_mixture(M::InverseReinforcementLearning, \( \mu s \))
    \( \mu E = M.\mu E \)
    \( k = \text{length}(\mu s) \)
    \( \text{model} = \text{Model}(\text{Ipopt.Optimizer}) \)
    \( \text{@variable}(\text{model}, \lambda[1:k] \geq 0) \)
    \( \text{@objective}(\text{model}, \text{Min}, (\mu E - \text{sum}(\lambda[i] \cdot \mu s[i] \text{ for } i \text{ in } 1:k)).) \)
    \( (\mu E - \text{sum}(\lambda[i] \cdot \mu s[i] \text{ for } i \text{ in } 1:k))) \)
    \( \text{@constraint}(\text{model}, \text{sum}(\lambda) == 1) \)
    \( \text{optimize!}(\text{model}) \)
    \( \text{return} \ \text{value}(\lambda) \)
end

function optimize(M::InverseReinforcementLearning, \( \theta \))
    \( \pi, \epsilon, RL = M.\pi, M.\epsilon, M.RL \)
    \( \theta s = [\theta] \)
    \( \mu s = \text{[feature_expectations}(M, s \to \pi(\theta, s))] \)
    while true
        \( t, \phi = \text{calc_weighting}(M, \mu s) \)
        if \( t \leq \epsilon \)
            \( \text{break} \)
        end
        copyto!(RL.\phi, \phi) \# R(s,a) = \phi \cdot \beta(s,a)
        \( \theta = \text{optimize}(RL, \pi, \theta) \)
        push!(\( \theta s, \theta \))
        push!(\( \mu s, \text{feature_expectations}(M, s \to \pi(\theta, s)) \))
    end
    \( \lambda = \text{calc_policy_mixture}(M, \mu s) \)
    \( \text{return} \ \lambda, \theta s \)
end
```

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Any policy \( \pi \) induces a distribution over trajectories\(^{10} \) \( P_\pi(\tau) \). Different policies produce different trajectory distributions. We are free to choose any of these distributions over trajectories that match the expert feature expectations. The principle of maximum entropy chooses the least informative distribution, which corresponds to the one with maximum entropy.\(^{11} \) It can be shown that the least informative trajectory distribution has the form:

\[
P_\phi(\tau) = \frac{1}{Z(\phi)} \exp(R_\phi(\tau)) \tag{18.8}
\]

where \( P_\phi(\tau) \) is the likelihood of a trajectory \( \tau \) given reward parameter \( \phi \), and

\[
R_\phi(\tau) = \sum_{k=1}^{d} \gamma^{k-1} R_\phi(s^{(k)}, a^{(k)}) \tag{18.9}
\]

is the discounted trajectory reward. We make no assumption on the parameterization of \( R_\phi(s^{(k)}, a^{(k)}) \) other than that it is differentiable, allowing for representations such as neural networks. The normalization scalar \( Z(\phi) \) ensures that the probabilities sum to 1:

\[
Z(\phi) = \sum_{\tau} \exp(R_\phi(\tau)) \tag{18.10}
\]

The summation is over all possible trajectories.

We seek to maximize the likelihood of the expert demonstrations under this distribution:

\[
\max \phi f(\phi) = \max \phi \sum_{\tau \in D} \log P_\phi(\tau) \tag{18.11}
\]

We can rewrite the objective function \( f(\phi) \) from equation (18.11):

\[
f(\phi) = \sum_{\tau \in D} \log \frac{1}{Z(\phi)} \exp(R_\phi(\tau)) \tag{18.12}
\]

\[
= \left( \sum_{\tau \in D} R_\phi(\tau) \right) - |D| \log Z(\phi) \tag{18.13}
\]

\[
= \left( \sum_{\tau \in D} R_\phi(\tau) \right) - |D| \log \sum_{\tau} \exp(R_\phi(\tau)) \tag{18.14}
\]

\(^{10} \) For simplicity, this section assumes a finite horizon and that the state and action spaces are discrete, making \( P_\phi(\tau) \) a probability mass. To extend maximum entropy inverse reinforcement learning both to problems with continuous state and action spaces where the dynamics may be unknown, consider guided cost learning: C. Finn, S. Levine, and P. Abbeel, “Guided Cost Learning: Deep Inverse Optimal Control via Policy Optimization,” in *International Conference on Machine Learning (ICML)*, 2016.

We can attempt to optimize this objective function through gradient ascent. The gradient of $f$ is:

$$
\nabla_{\phi} f = \left(\sum_{\tau \in D} \nabla_{\phi} R_{\phi}(\tau)\right) - \frac{|D|}{\sum_{\tau} \exp(R_{\phi}(\tau))} \sum_{\tau} \exp(R_{\phi}(\tau)) \nabla_{\phi} R_{\phi}(\tau)
$$

(18.15)

$$
= \left(\sum_{\tau \in D} \nabla_{\phi} R_{\phi}(\tau)\right) - |D| \sum_{\tau} P_{\phi}(\tau) \nabla_{\phi} R_{\phi}(\tau)
$$

(18.16)

$$
= \left(\sum_{\tau \in D} \nabla_{\phi} R_{\phi}(\tau)\right) - |D| \sum_{s} b_{\gamma,\phi}(s) \sum_{a} \pi_{\phi}(a | s) \nabla_{\phi} R_{\phi}(s, a)
$$

(18.17)

If the reward function is linear with $R_{\phi}(s, a) = \phi^\top \beta(s, a)$ as in the previous section, then $\nabla_{\phi} R_{\phi}(s, a)$ is simply $\beta(s, a)$.

Updating the parameter vector $\phi$ thus requires both the discounted state visitation frequency $b_{\gamma,\phi}$ and the optimal policy under the current parameter vector, $\pi_{\phi}(a | s)$. We can obtain the optimal policy by running reinforcement learning. To compute the discounted state visitation frequencies we can use rollouts or take a dynamic programming approach.

If we take a dynamic programming approach to compute the discounted state visitation frequencies, we can start with the initial state distribution $b_{\gamma,\phi}^{(1)}(s) = b(s)$ and iteratively work forward in time:

$$
b_{\gamma,\phi}^{(k+1)}(s) = \gamma \sum_{a} \sum_{s'} b_{\gamma,\phi}^{(k)}(s') \pi(a | s) T(s' | s, a)
$$

(18.18)

This version of maximum entropy inverse reinforcement learning is implemented in algorithm 18.6.

### 18.6 Generative Adversarial Imitation Learning

In generative adversarial imitation learning (GAIL),\(^{12}\) we optimize a differentiable parameterized policy $\pi_{\theta}$, often represented by a neural network. Rather than provide a reward function, we use adversarial learning (appendix D.7). We also train a discriminator $C_{\phi}(s, a)$, typically also a neural network, to return the probability it assigns to the state-action pair coming from the learned policy. The process involves alternating between training this discriminator to become better

Algorithm 18.6. Maximum entropy inverse reinforcement learning, which finds a stochastic policy that maximizes the likelihood of the expert demonstrations under a maximum-entropy trajectory distribution. This implementation computes the expected visitations using dynamic programming over all states, which requires that the problem be discrete.
at distinguishing, and training the policy to look indistinguishable from the expert demonstrations. The process is sketched in figure 18.3.

![Diagram](image)

Figure 18.3. Instead of inferring a reward function, generative adversarial imitation learning optimizes a discriminator to distinguish between simulated and expert state-action pairs, and it optimizes a policy to appear indistinguishable to the discriminator. The aim is to eventually produce a policy that resembles the expert.

The discriminator and policy have opposite objectives. Generative adversarial imitation learning seeks to find a saddle-point \((\theta, \phi)\) of the negative log loss of the discriminator’s binary classification problem:\(^{13}\)

\[
\max_{\phi} \min_{\theta} \mathbb{E}_{(s,a) \sim D} \left[ \log \left( C_{\phi}(s,a) \right) \right] + \mathbb{E}_{(s,a) \sim D} \left[ \log \left( 1 - C_{\phi}(s,a) \right) \right]
\]

where we use \((s,a) \sim D\) to represent samples from the distribution represented by the expert dataset \(D\). We can alternate between gradient ascent on \(\phi\) to increase the objective and trust region policy optimization (section 12.4) on \(\theta\) to reduce the objective, generating the necessary trajectory samples from the policy to conduct each of these steps. The discriminator provides a learning signal to the policy similar to how a reward signal would if it were known.

### 18.7 Summary

- Imitation learning involves learning the desired behavior from expert demonstration without the use of a reward function.

- One type of imitation learning is behavioral cloning, which produces a stochastic policy that maximizes the conditional likelihood of the actions in the dataset.
• When an expert can be queried multiple times, we can use iterative approaches like dataset aggregation or stochastic mixing iterative learning.

• Inverse reinforcement learning involves inferring a reward function from expert data and then using traditional methods for finding an optimal policy.

• Maximum margin inverse reinforcement learning attempts to find a policy that matches the frequency of binary features found in the expert dataset.

• Maximum entropy inverse reinforcement learning frames the problem of finding the best reward parameter as a maximum likelihood estimation problem, which it tries to solve using gradient ascent.

• Generative adversarial imitation learning iteratively optimizes a discriminator and a policy; the discriminator tries to discriminate between decisions made by the policy and decisions made by the expert, and the policy attempts to deceive the discriminator.

18.8 Exercises

Exercise 18.1. Consider applying behavioral cloning to a discrete problem where we have been given expert demonstrations. An alternative could be to define a feature function $\beta(s)$ and represent the policy with a softmax distribution

$$\pi(a \mid s) \propto \exp(\theta_a^\top \beta(s))$$

We would then learn the parameters $\theta_a$ for each action from the expert data. Why might we want to use this approach over one where we directly estimate a discrete distribution for each state, with one parameter per state-action pair?

Solution: In imitation learning, we are generally limited to a relatively small set of expert demonstrations. The distribution $P(a \mid s)$ has $(|A| - 1)|S|$ independent parameters that must be learned, which is often prohibitively large. Expert demonstrations typically only cover a small portion of the state-space. Even if $P(a \mid s)$ can be reliably trained for the states covered in the provided dataset, the resulting policy would be untrained in other states. Using a feature function allows for generalization to unseen states.
**Exercise 18.2.** The section on behavioral cloning suggested using a maximum likelihood approach for training a policy from expert data. This approach attempts to find the parameters of the policy that maximizes the likelihood assigned to the training examples. In some problems, however, we know that assigning high probability to one incorrect action is not as bad as assigning high probability to another incorrect action. For example, predicting an acceleration of $-1$ in the mountain car problem when the expert dictates an acceleration of $1$ is worse than predicting an acceleration of $0$. How might behavioral cloning be modified to allow different penalties to be given to different misclassifications?

*Solution:* We can instead supply a cost function $C(s, a_{\text{true}}, a_{\text{pred}})$ that defines the cost of predicting action $a_{\text{pred}}$ for state $s$ when the expert’s action is $a_{\text{true}}$. For example, with the mountain-car problem we might use:

$$C(s, a_{\text{true}}, a_{\text{pred}}) = -|a_{\text{true}} - a_{\text{pred}}|$$

which penalizes greater deviations more than smaller deviations. The cost associated with the expert’s action is typically zero.

If we have a stochastic policy $\pi(a \mid s)$, then we seek to minimize the cost over our dataset:

$$\min_{\theta} \sum_{(s,a_{\text{true}},a_{\text{pred}}) \in D} \sum_{a_{\text{pred}}} C(s, a_{\text{true}}, a_{\text{pred}}) \pi(a_{\text{pred}} \mid s)$$

This technique is called *cost-sensitive classification*.\(^{14}\) One benefit of cost-sensitive classification is that we can use a wide variety of off-the-shelf classification models like $k$-nearest neighbors, support vector machines, or decision trees to train a policy.

**Exercise 18.3.** Provide an example of where maximum margin inverse reinforcement learning does not uniquely define an optimal policy.

*Solution:* Maximum margin inverse reinforcement learning extracts binary features from the expert data and seeks a reward function whose optimal policy produces trajectories with the same frequencies of these binary features. There is no guarantee that multiple policies do not produce the same feature expectations. For example, an autonomous car that only makes left lane changes could have the same lane change frequencies as an autonomous car that only makes right lane changes.

**Exercise 18.4.** Maximum margin inverse reinforcement learning measures how similar a policy is to expert demonstrations using feature expectations. How is this similarity measure affected if non-binary features are used?

**Solution:** If we use non-binary features, then it is possible that some features can get larger than others, incentivizing the agent to match those features over those that tend to be smaller. Scale is not the only issue. Even if all features are constrained to lie within $[0, 1]$, then a policy that consistently produces $\phi(s, a)_1 = 0.5$ will have the same feature expectations as one that produces $\phi(s, a)_1 = 0$ half the time and $\phi(s, a)_1 = 1$ half the time. Depending on what the feature encodes, this can result in very different policies. Any set of continuous features can be discretized, and thus approximated by a set of binary features.

**Exercise 18.5.** Suppose we are building a system for an elevator in a high-rise that has to choose which floor to send an elevator. We have trained several policies to match the feature expectations of expert demonstrations, such as how long a customer has to wait for an elevator or how long they have to wait to get to their destination. We run multiple rollouts for each policy and plot the relative duration spent on each floor. Which policy should we prefer according to the principle of maximum entropy, assuming each policy matches the feature expectations equally?

**Solution:** These distributions over relative duration are analogous to distributions over trajectories for this elevator problem. In applying the principle of maximum entropy, we prefer the distribution with most entropy. Hence, we would choose policy B, which in being most uniform, has the greatest entropy.

**Exercise 18.6.** Consider the policy optimization step in generative adversarial imitation learning. Rewrite the objective in the form of a reward function so that traditional reinforcement learning techniques can be applied.

**Solution:** We rewrite equation (18.19), dropping the terms dependent on the expert dataset, and flip the sign to change from minimization over $\theta$ to a maximization over $\theta$ of the reward, producing the surrogate reward function:

$$\tilde{R}_\phi(s, a) = -\log C_\phi(s, a)$$
Although \( \hat{R}_\phi(s, a) \) may be quite different from the unknown true reward function, it can be used to drive the learned policy into regions of the state-action space similar to those covered by the expert.

**Exercise 18.7.** Explain how generative adversarial imitation learning could be changed such that the discriminator takes in trajectories rather than state-action pairs. Why might this be useful?

**Solution:** Changing generative adversarial imitation learning such that the discriminator takes trajectories is straightforward, especially if the trajectories are of fixed length. The expert dataset is split into trajectories, and the learned policy is used to produce trajectories, just as it was before. Rather than operating on state-action pairs, the discriminator takes in trajectories using a representation such as a recurrent neural network (appendix D.5) and produces a classification probability. The objective function remains largely unchanged:

\[
\max_{\phi} \min_{\theta} \mathbb{E}_{\tau \sim \pi^0} \left[ \log (C_{\phi}(\tau)) \right] + \mathbb{E}_{\tau \sim \mathcal{D}} \left[ \log (1 - C_{\phi}(\tau)) \right]
\]

The advantage of running the discriminator over entire trajectories is that it can help the discriminator capture features that are not apparent from individual state-action pairs, which can result in better policies. For example, when looking at individual accelerations and turn rates for an autonomous driving policy, there is very little for a discriminator to learn. A discriminator trained to look at longer trajectories can see more of the vehicle’s behavior, such as lane change aggressiveness and smoothness, to better match expert driving demonstrations.\(^{15}\)

\(^{15}\) This approach was used in A. Kuefler, J. Morton, T.A. Wheeler, and M.J. Kochenderfer, “Imitating Driver Behavior with Generative Adversarial Networks,” in *IEEE Intelligent Vehicles Symposium (IV)*, 2017.
PART IV:

STATE UNCERTAINTY

Previous chapters have included uncertainty in the transition function, both in terms of the uncertainty in the resulting state as well as in the model. In this part, we extend uncertainty to include the state. Instead of observing the state exactly, we receive observations that have only a probabilistic relationship with the state. Such problems can be modeled as a *partially observable Markov decision process* (POMDP). A common approach to solving POMDPs involves inferring a belief distribution over the underlying state at the current time step and then applying a policy that maps beliefs to actions. Chapter 19 shows how to update our belief distribution given a past sequence of observations and actions. Chapter 20 outlines exact methods for optimizing policies. Chapter 21 reviews a variety of offline approximation methods that tend to scale much better than exact methods to larger problems. Chapter 22 extends some of the online approximation discussed earlier in this book to accommodate partial observability. Chapter 23 introduces finite state controllers to represent policies and methods for optimizing them.
A POMDP is an MDP with state uncertainty. The agent receives a potentially imperfect observation of the current state rather than the true state. From the past sequence of observations and actions, the agent develops an understanding of the world. This chapter discusses how the belief of the agent can be represented by a probability distribution over the underlying state. Various algorithms are presented for updating our belief based on the observation and action taken by the agent. We can perform exact belief updates if the state space is discrete or if certain linear Gaussian assumptions are met. In cases where these assumptions do not hold, we can use approximations based on linearization or sampling.

19.1 Belief Initialization

There are different ways to represent our beliefs. In this chapter, we will discuss parametric representations where the belief distribution is represented by a set of parameters of a fixed distribution family, such as the categorical or the multivariate normal distribution. We will also discuss non-parametric representations where the belief distribution is represented by particles, or points sampled from the state space. Associated with different belief representations are different procedures for updating the belief based on the action taken by the agent and the observation.

Before the agent takes any actions or makes any observations, we start with an initial belief distribution. If we have some prior information about where the agent might be in the state space, we can encode this in the initial belief. We generally want to use diffuse initial beliefs in the absence of information to avoid being overly confident in the agent being in a region of the state space where it might not actually be. A strong initial belief focused on states that are far from the true state can lead to poor state estimates even after many observations.
A diffuse initial prior can cause difficulties, especially for non-parametric representations of the belief, where the state space can only be very sparsely sampled. In some cases, it may be useful to wait until an informative observation is made to initialize our beliefs. For example, in robot navigation problems, we might want to wait until the sensors detect a known landmark and then initialize the belief appropriately. The landmark can help narrow down the relevant region of the state space so we can focus our sampling of the space in the area consistent with the landmark observation. Example 19.1 illustrates this concept.

Consider an autonomous car equipped with a localization system that uses camera, radar, and lidar data to track its position. The car is able to identify a unique landmark at a range $r$ and bearing $\theta$ from its current pose.

![Diagram of car and landmark](image)

The range and bearing measurements have zero-mean Gaussian noise with variance $\nu_r$ and $\nu_\theta$, respectively, and the landmark is known to be at $(x, y)$. Given a measurement $r$ and $\theta$, we can produce a distribution over the car’s position $(\hat{x}, \hat{y})$ and orientation $\hat{\psi}$:

$$
\hat{r} \sim N(r, \nu_r) \quad \hat{\theta} \sim N(\theta, \nu_\theta) \quad \hat{\phi} \sim U(0, 2\pi)
$$

$$
\hat{x} \leftarrow x + \hat{r} \cos \hat{\phi} \quad \hat{y} \leftarrow y + \hat{r} \sin \hat{\phi} \quad \hat{\psi} \leftarrow \hat{\phi} - \hat{\theta} + \pi
$$

where $\hat{\phi}$ is the angle of the car from the landmark in the global frame.

### 19.2 Discrete State Filter

In a POMDP, the agent does not directly observe the underlying state of the environment. Instead, the agent receives an observation, which belongs to some observation space $O$, at each time step. The probability of observing $o$ given the agent took action $a$ and transitioned to state $s'$ is given by $O(o \mid a, s')$. If $O$...
is continuous, then \( O(o \mid a, s') \) is a probability density. Figure 19.1 shows the dynamic decision network associated with POMDPs. Algorithm 19.1 provides an implementation of the POMDP data structure.

```
struct POMDP
    γ # discount factor
    𝒮 # state space
    𝒜 # action space
    𝒪 # observation space
    T # transition function
    R # reward function
    O # observation function
    TRO # sample transition, reward, and observation
end
```

Algorithm 19.1. A data structure for POMDPs. We will use the TRO field to sample the next state, reward, and observation given the current state and action: \( s', r, o = TRO(s, a) \).

A kind of inference known as recursive Bayesian estimation can be used to update our belief distribution over the current state given the most recent action and observations. We use \( b(s) \) to represent the probability (or probability density for continuous state spaces) assigned to state \( s \). A particular belief \( b \) belongs to a belief space \( B \), which contains all possible beliefs.

When the state and observation spaces are finite, we can use a discrete state filter to perform this inference exactly. Beliefs for problems with discrete state spaces can be represented using categorical distributions, where a probability mass is assigned to each state. This categorical distribution can be represented as a vector of length \(|S|\) and is often called a belief vector. In cases where \( b \) can be treated as a vector, we will use \( b \). In this case, \( B \subseteq \mathbb{R}^{|S|} \). Sometimes \( B \) is referred to as a probability simplex or belief simplex.

Because a belief vector represents a probability distribution, the elements must be strictly non-negative and must sum to one:

\[
b(s) \geq 0 \text{ for all } s \in S \quad \sum_s b(s) = 1 \tag{19.1}\]

In vector form, we have
\[
b \geq 0 \quad 1^\top b = 1 \tag{19.2}\]

The belief space for a POMDP with three states is given in figure 19.2. An example discrete POMDP problem is given in example 19.2.

If an agent with belief \( b \) takes an action \( a \) and receives an observation \( o \), the new belief \( b' \) can be calculated as follows due to the independence assumptions

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The *crying baby problem* is a simple POMDP with two states, three actions, and two observations. Our goal is to care for a baby, and we do so by choosing at each time step whether to feed the baby, sing to it, or ignore it.

The baby becomes hungry over time. One does not directly observe whether the baby is hungry, but instead receives a noisy observation in the form of whether or not the baby is crying. A hungry baby cries 80% of the time, whereas a sated baby cries 10% of the time. Singing to the baby yields a perfect observation. The state, action, and observation spaces are:

\[
S = \{\text{hungry, sated}\} \\
A = \{\text{feed, sing, ignore}\} \\
O = \{\text{crying, quiet}\}
\]

The transition dynamics are:

\[
T(\text{sated} \mid \text{hungry, feed}) = 100\% \\
T(\text{hungry} \mid \text{hungry, sing}) = 100\% \\
T(\text{hungry} \mid \text{hungry, ignore}) = 100\% \\
T(\text{sated} \mid \text{sated, feed}) = 100\% \\
T(\text{hungry} \mid \text{sated, sing}) = 10\% \\
T(\text{hungry} \mid \text{sated, ignore}) = 10\%
\]

The reward function assigns $-10$ reward if the baby is hungry and $-5$ reward for feeding the baby. Thus, feeding a hungry baby results in $-15$ reward. Singing to a sated baby makes it happy to receive the attention, and thus yields 5 reward. Singing to a hungry baby takes extra effort, and thus incurs $-2$ reward. As baby caretakers, we seek the optimal infinite horizon policy with discount factor $\gamma = 0.9$. 

Example 19.2. The crying baby problem is a simple POMDP used to demonstrate decision making with state uncertainty.
in figure 19.1:

\[ b'(s') = P(s' | b, a, o) \]  \hspace{1cm} (19.3)
\[ \propto P(o | b, a, s')P(s' | b, a) \]  \hspace{1cm} (19.4)
\[ \propto O(o | a, s')P(s' | b, a) \]  \hspace{1cm} (19.5)
\[ \propto O(o | a, s') \sum_s P(s' | a, b, s)P(s | b, a) \]  \hspace{1cm} (19.6)
\[ \propto O(o | a, s') \sum_s T(s' | s, a)b(s) \]  \hspace{1cm} (19.7)

An example of updating discrete beliefs is given in example 19.3, and the belief update is implemented in algorithm 19.2. The success of the belief update depends upon having accurate observation and transition models. In cases where these models are not known well, it is generally advisable to use simplified models with more diffuse distributions to help prevent overconfidence that leads to brittleness in the state estimates.

19.3 Linear Gaussian Filter

We can adapt equation (19.7) to handle continuous state spaces as follows:

\[ b'(s') \propto O(o | a, s') \int T(s' | s, a)b(s) \, ds \]  \hspace{1cm} (19.8)

The integration above can be challenging unless we make some assumptions about the form of \( T, O, \) and \( b. \) A special type of filter known as the Kalman filter (algorithm 19.3)\(^2\) provides an exact update under the assumption that \( T \) and \( O \) are linear Gaussian and \( b \) is Gaussian:\(^3\)

\[ T(s' | s, a) = \mathcal{N}(s' | T_s s + T_a a, \Sigma_s) \]  \hspace{1cm} (19.9)
\[ O(o | s') = \mathcal{N}(o | O_s s', \Sigma_o) \]  \hspace{1cm} (19.10)
\[ b(s) = \mathcal{N}(s | \mu_b, \Sigma_b) \]  \hspace{1cm} (19.11)

The Kalman filter begins with a predict step, which uses the transition dynamics to get a predicted distribution with the following mean and covariance:

\[ \mu_p \leftarrow T_s \mu_b + T_a a \]  \hspace{1cm} (19.12)
\[ \Sigma_p \leftarrow T_s \Sigma_b T_s^\top + \Sigma_s \]  \hspace{1cm} (19.13)

\(^2\) Named after the Hungarian-American electrical engineer Rudolf E. Kálmán (1930–2016) who was involved in the early development of this filter.

The crying baby problem (example 19.2) assumes a uniform initial belief state: \([b(\text{sated}), b(\text{hungry})] = [0.5, 0.5]\).

Suppose we ignore the baby and the baby cries. We update our belief according to equation (19.7):

\[
b'(\text{sated}) \propto O(\text{crying} \mid \text{ignore}, \text{sated}) \sum_s T(\text{sated} \mid s, \text{ignore}) b(s)
\]
\[
\propto 0.1(0.0 \cdot 0.5 + 0.9 \cdot 0.5)
\]
\[
\propto 0.045
\]

\[
b'(\text{hungry}) \propto O(\text{crying} \mid \text{ignore}, \text{hungry}) \sum_s T(\text{hungry} \mid s, \text{ignore}) b(s)
\]
\[
\propto 0.8(1.0 \cdot 0.5 + 0.1 \cdot 0.5)
\]
\[
\propto 0.440
\]

After normalizing, our new belief is approximately \([0.0928, 0.9072]\). A crying baby is likely to be hungry.

Suppose we then feed the baby and the crying stops. Feeding deterministically caused the baby to be sated, so the new belief is \([1, 0]\).

Finally, we sing to the baby and the baby is quiet. Equation (19.7) is used again to update the belief, resulting in \([0.9759, 0.0241]\). A sated baby only becomes hungry 10% of the time, and this percentage is further reduced by not observing any crying.

```python
function update(b::Vector{Float64}, ℙ, a, o)
    S, T, O = ℙ.S, ℙ.T, ℙ.O
    b' = similar(b)
    for (i', s') in enumerate(S)
        po = O(a, s', o)
        b'[i'] = po * sum(T(s, a, s') * b[i] for (i, s) in enumerate(S))
    end
    if sum(b') ≈ 0.0
        fill!(b', 1)
    end
    return normalize!(b', 1)
end
```

Example 19.3. Discrete belief updating in the crying baby problem.

Algorithm 19.2. A method that updates a discrete belief based on equation (19.7), where \(b\) is a vector and \(ℙ\) is the POMDP model. If the given observation has a zero likelihood, a uniform distribution is returned.
In the *update step*, we use this predicted distribution with the current observation to update our belief:

\[
K \leftarrow \Sigma_p O_s^\top \left( O_s \Sigma_p O_s^\top + \Sigma_o \right)^{-1} \\
\mu_b \leftarrow \mu_p + K \left( o - O_s \mu_p \right) \\
\Sigma_b \leftarrow (I - KO_s) \Sigma_p
\]

where \( K \) is called the *Kalman gain*.

```
struct KalmanFilter
    µb # mean vector
    Σb # covariance matrix
end

function update(b::KalmanFilter, ℙ, a, o)
    µb, Σb = b.µb, b.Σb
    Ts, Ta, Os = ℙ.Ts, ℙ.Ta, ℙ.Os
    Σs, Σo = ℙ.Σs, ℙ.Σo
    # predict
    µp = Ts*µb + Ta*a
    Σp = Ts*Σb*Ts' + Σs
    # update
    K = Σp*Os'/\left( Os*Σp*Os' + Σo \right)
    µb' = µp + K*(o - Os*µp)
    Σb' = (I - K*Os)*Σp
    return KalmanFilter(µb', Σb')
end
```

Kalman filters are often applied to systems that do not actually have linear Gaussian dynamics and observations. A variety of different modifications to the basic Kalman filter have been proposed to better accommodate such systems.\(^4\)

### 19.4 Extended Kalman Filter

The *extended Kalman filter (EKF)*, is a simple extension of the Kalman filter to problems whose dynamics are nonlinear with Gaussian noise:

\[
T(s' \mid s, a) = \mathcal{N}(s' \mid f_T(s, a), \Sigma_s) \tag{19.17}
\]
\[
O(o \mid s') = \mathcal{N}(o \mid f_O(s'), \Sigma_o) \tag{19.18}
\]

where \( f_T(s, a) \) and \( f_O(s') \) are differentiable functions.

Consider the following linear Gaussian transition and observation functions:

\[
T(s' \mid s, a) = \mathcal{N}\left(s' \mid s + a \begin{bmatrix} 1 & 1/2 \\ 1/2 & 1 \end{bmatrix}ight)
\]

\[
O(o \mid s') = \mathcal{N}\left(o \mid s' \begin{bmatrix} 1/2 & -1/2 \\ -1/2 & 2 \end{bmatrix} \right)
\]

Suppose our initial belief is \( b = \mathcal{N}([-0.75, 1], I) \). We take action \( a = [0.5, -0.5] \) and observe \( o = [0.3, 0.5] \).

Using the Kalman filter update equations, our new belief is:

\[
b' = \mathcal{N}\left(\begin{bmatrix} 0.184 \\ 0.571 \end{bmatrix}, \begin{bmatrix} 0.037 & -0.011 \\ -0.011 & 0.050 \end{bmatrix}\right)
\]

using the Kalman gain:

\[
K = \begin{bmatrix} 0.789 & 0.110 \\ 0.128 & 0.716 \end{bmatrix}
\]

Note that because the observation dynamics are more certain, the new belief is centered closer to the observation than to the ideal \( s' \) predicted by the transition dynamics applied to the previous belief mean and action.
19.5. Unscented Kalman Filter

Exact belief updates through nonlinear dynamics are not guaranteed to produce new Gaussian beliefs, as shown in figure 19.3. The extended Kalman filter uses a local linear approximation to the nonlinear dynamics, thereby producing a new Gaussian belief that approximates the true updated belief. We can use similar update equations as the Kalman filter, but must compute the matrices $T_s$ and $O_s$ at every iteration based on our current belief.

The local linear approximation to the dynamics, or linearization, is given by first-order Taylor expansions in the form of Jacobians. For the state matrix, the Taylor expansion is conducted at $\mu_b$ and the current action, whereas for the observation matrix, it is computed at the predicted mean $\mu_p = f_T(\mu_b)$.

The extended Kalman filter is implemented in algorithm 19.4. Although it is an approximation, it is fast and performs well on a variety of real-world problems. The EKF does not generally preserve the true mean and variance of the posterior, and it does not model multimodal posterior distributions.

```
struct ExtendedKalmanFilter
    μb # mean vector
    Σb # covariance matrix
end

import ForwardDiff: jacobian

function update(b::ExtendedKalmanFilter, ℙ, a, o)
    μb, Σb = b.μb, b.Σb
    fT, fO = ℙ.fT, ℙ.fO
    Σs, Σo = ℙ.Σs, ℙ.Σo
    # predict
    μp = fT(μb, a)
    Ts = jacobian(s→fT(s, a), μb)
    Os = jacobian(fO, μp)
    Σp = Ts*Σb*Ts' + Σs
    # update
    K = Σp*Os'/(Σs*Σp*Os' + Σo)
    μb' = μp + K*(o - fO(μp))
    Σb' = (I - K*Os)*Σp
    return ExtendedKalmanFilter(μb', Σb')
end
```

Algorithm 19.4. The extended Kalman filter, an extension of the Kalman filter to problems with nonlinear Gaussian dynamics. The current belief is represented by mean $\mu_b$ and covariance $\Sigma_b$. The problem $\mathcal{P}$ specifies the nonlinear dynamics using the mean transition dynamics function $f_T$ and mean observation dynamics function $f_O$. The Jacobians are obtained using the ForwardDiff.jl package.

The Jacobian of a multivariate function $f$ with $n$ inputs and $m$ outputs is an $m \times n$ matrix where the $(i, j)$th entry is $\frac{\partial f_i}{\partial x_j}$.


7 According to Uhlmann, the term “unscented” comes from a label on a deodorant container that he saw on someone’s desk. He used that term to avoid calling it the Uhlmann filter. IEEE History Center Staff, “Proceedings of the IEEE Through 100 Years: 2000–2009,” *Proceedings of the IEEE*, vol. 100, no. 11, pp. 3131–3145, 2012.
filter, the unscented Kalman filter is derivative-free, and relies on a deterministic sampling strategy to approximate the effect of a distribution undergoing a (typically nonlinear) transformation.

The unscented Kalman filter was developed to estimate the effect of passing a random variable $x$ through a nonlinear distribution $f(x)$, producing the random variable $x'$. We would like to estimate the mean $\mu'$ and covariance $\Sigma'$ of the distribution over $x'$. The unscented transform allows for more information of $p(x)$ to be used than the mean $\mu$ and covariance $\Sigma$ of the distribution over $x$.\(^8\)

An unscented transform passes a set of sigma points $S$ through $f$ and uses the transformed points to approximate the transformed mean $\mu'$ and covariance $\Sigma'$. The mean and covariance are reconstructed using a vector of weights $w$:

$$\mu' = \sum_i w_i s_i$$  \hspace{1cm} (19.19)

$$\Sigma' = \sum_i w_i (s_i - \mu')(s_i - \mu')^\top$$  \hspace{1cm} (19.20)

These weights must sum to 1 in order to provide an unbiased estimate, but need not all be positive.
The updated mean and covariance matrix given by the unscented transform through $f$ are thus:

$$
\mu' = \sum_i w_i f(s_i) \tag{19.21}
$$

$$
\Sigma' = \sum_i w_i (f(s_i) - \mu') (f(s_i) - \mu')^\top \tag{19.22}
$$

An example set of sigma points and associated weight vector is given in example 19.5.

A simple set of $2n$ sigma points for an $n$-dimensional distribution is given by:

$$
s_{2i} = \mu + \sqrt{n} \Sigma_i \text{ for } i \text{ in } 1 : n
$$

$$
s_{2i-1} = \mu - \sqrt{n} \Sigma_i \text{ for } i \text{ in } 1 : n
$$

where the $i$ subscript indicates the $i$th column of the square root matrix.

If we use the weights $w_i = 1/2n$, the reconstructed mean is:

$$
\sum_i w_i s_i = \sum_{i=1}^n \frac{1}{2n} (\mu + \sqrt{n} \Sigma_i) + \frac{1}{2n} (\mu - \sqrt{n} \Sigma_i) = \sum_{i=1}^n \frac{1}{n} \mu = \mu
$$

and the reconstructed covariance is:

$$
\sum_i w_i (s_i - \mu') (s_i - \mu')^\top = 2 \sum_{i=1}^n \frac{1}{2n} (\sqrt{n} \Sigma_i) (\sqrt{n} \Sigma_i)^\top
$$

$$
= \frac{1}{n} \sum_{i=1}^n (\sqrt{n} \Sigma_i) (\sqrt{n} \Sigma_i)^\top
$$

$$
= \sqrt{\Sigma} \sqrt{\Sigma}^\top
$$

$$
= \Sigma
$$

Sigma points and weights can be selected to incorporate additional information about the prior distribution. Suppose we always have a set of sigma points and weights that produce a mean $\mu'$ and covariance $\Sigma'$. We can construct a new set of sigma points by additionally including the mean $\mu'$ as a sigma point. We scale the sigma points such that the covariance is preserved. This results in a scaled set...
of sigma points with different higher-order moments, but the same mean and covariance. Example 19.6 derives a set of sigma points by applying this technique to the sigma points in example 19.5.

We can include the mean \( \mu \) in the sigma points from example 19.5 to obtain a new set of \( 2n + 1 \) sigma points:

\[
\begin{align*}
\mathbf{s}_1 &= \mu \\
\mathbf{s}_{2i} &= \mu + \left( \sqrt{\frac{n}{1 - w_1}} \Sigma \right)_i \quad \text{for } i \text{ in } 1 : n \\
\mathbf{s}_{2i+1} &= \mu - \left( \sqrt{\frac{n}{1 - w_1}} \Sigma \right)_i \quad \text{for } i \text{ in } 1 : n
\end{align*}
\]

where \( w_1 \) is the weight on the first sigma point. The weights on the remaining sigma points are \((1 - w_1) / 2n\). The reconstructed mean is still \( \mu \), and the reconstructed covariance is still \( \Sigma \).

We can vary \( w_1 \) to produce different sets of sigma points. Setting \( w_1 > 0 \) causes the sigma points to spread away from the mean. Setting \( w_1 < 0 \) moves the sigma points closer to the mean.

A common set of sigma points include the mean \( \mu \in \mathbb{R}^n \) and an additional \( 2n \) points formed from perturbations of \( \mu \) in directions determined by the covariance matrix \( \Sigma \):

\[
\begin{align*}
\mathbf{s}_1 &= \mu \\
\mathbf{s}_{2i} &= \mu + \left( \sqrt{(n + \lambda) \Sigma} \right)_i \quad \text{for } i \text{ in } 1 : n \\
\mathbf{s}_{2i+1} &= \mu - \left( \sqrt{(n + \lambda) \Sigma} \right)_i \quad \text{for } i \text{ in } 1 : n
\end{align*}
\]

These sigma points are associated with the weights:

\[
\begin{align*}
w_i &= \begin{cases} 
\frac{\lambda}{n + \lambda} & \text{for } i = 1 \\
\frac{1}{2(n + \lambda)} & \text{otherwise}
\end{cases}
\end{align*}
\]
The scalar *spread parameter* $\lambda$ determines how far the sigma points are spread from the mean. It is recommended to use $\lambda = 2$, which is optimal for matching the fourth moment of Gaussian distributions. Several sigma point sets for different values of $\lambda$ are shown in figure 19.4.

![Figure 19.4. The effect of varying $\lambda$ on the sigma points from equation (19.23) generated for a Gaussian distribution with zero mean and covariance $\Sigma = \begin{bmatrix} 1 & 1/2 \\ 1/2 & 2 \end{bmatrix}$.](image)

The unscented Kalman filter performs two unscented transformations, one for the prediction step and one for the observation update. The original Kalman filter computes the Kalman gain using the observation covariance $O_s$. The unscented Kalman filter does not have this observation matrix, and instead calculates the Kalman gain using a *cross covariance matrix* using both prediction values and observation values. Algorithm 19.5 provides an implementation.

### 19.6 Particle Filter

Discrete problems with large state spaces or continuous problems with dynamics that are not well approximated by the linear Gaussian assumption of the Kalman filter must often resort to approximation techniques to represent the belief and to perform the belief update. One common approach is to use a *particle filter*, which represents the belief state as a collection of states. Each state in the approximate belief is called a *particle*.

A particle filter is initialized by selecting or randomly sampling a collection of particles that represent the initial belief. The belief update for a particle filter with $m$ particles begins by propagating each state $s_i$ by sampling from the transition distribution to obtain a new state $s'_i$ with probability $T(s'_i \mid s_i, a)$. The new belief is constructed by drawing $m$ particles from the propagated states weighted according to the observation function $w_i = O(o \mid a, s')$. This procedure is given in algorithm 19.6. Example 19.7 illustrates an application of a particle filter.

---

9 A covariance matrix measures the variance between components of the same multi-dimensional variable. A cross covariance matrix measures the variance between two multi-dimensional variables. In algorithm 19.5, we compute a cross covariance matrix $\Sigma_{po}$.

Algorithm 19.5. The unscented Kalman filter, an extension of the Kalman filter to problems with nonlinear Gaussian dynamics. The current belief is represented by mean $\mu_b$ and covariance $\Sigma_b$. The problem $\mathcal{P}$ specifies the nonlinear dynamics using the mean transition dynamics function $f_T$ and mean observation dynamics function $f_O$. The sigma points used in the unscented transforms are controlled by the scalars $\alpha$, $\beta$, and $\kappa$. 

```plaintext
struct UnscentedKalmanFilter
    μb # mean vector
    Σb # covariance matrix
    λ # spread parameter
end

function unscented_transform(μ, Σ, f, λ, ws)
    n = length(μ)
    Δ = sqrt((n + λ) * Σ)
    S = [μ]
    for i in 1:n
        push!(S, μ + Δ[:, i])
        push!(S, μ - Δ[:, i])
    end
    S′ = f(S)
    μ′ = sum(w * S for (w, s) in zip(ws, S′))
    Σ′ = sum(w * (s - μ′) * (s - μ′)′ for (w, s) in zip(ws, S′))
    return (μ′, Σ′, S, S′)
end

function update(b::UnscentedKalmanFilter, P, a, o)
    μb, Σb, λ = b.μb, b.Σb, b.λ
    fT, fO = P.fT, P.fO
    n = length(μb)
    ws = [λ / (n + λ); fill(1/(2(n + λ)), 2n)]
    # predict
    μp, Σp, Sp, Sp′ = unscented_transform(μb, Σb, s→fT(s,a), λ, ws)
    Σp += P.Σs
    # update
    μo, Σo, So, So′ = unscented_transform(μp, Σp, fO, λ, ws)
    Σo += P.Σo
    Σo = sum(w*(s - μp)*(s′ - μo))′ for (w,s,s′) in zip(ws, So, So′)
    K = Σo / Σo
    μb′ = μp + K*(o - μo)
    Σb′ = Σp - K*Σo*K′
    return UnscentedKalmanFilter(μb′, Σb′, λ)
end
```
Suppose we want to determine our position based on imperfect distance measurements to radio beacons whose locations are known. We remain approximately still for a few steps to collect independent measurements. The particle filter states are our potential locations. We can compare the ranges we would expect to measure for each particle to the observed ranges.

We assume individual range observations from each beacon are observed with zero-mean Gaussian noise. Our particle transition function adds zero-mean Gaussian noise since we remain only approximately still.

The images below show the evolution of the particle filter. The rows correspond to different numbers of beacons. The red dot indicates our true location and the blue dots are particles. The circles indicate the positions consistent with noiseless distance measurements from each sensor.

Three beacons are required to accurately identify our location. A strength of the particle filter is that it is able to represent the multimodal distributions that are especially apparent when there are only one or two beacons.
struct ParticleFilter
    states # vector of state samples
end

function update(b::ParticleFilter, 𝒫, a, o)
    T, O = 𝒫.T, 𝒫.O
    states = [rand(T(s, a)) for s in b.states]
    weights = [O(a, s′, o) for s′ in states]
    D = SetCategorical(states, weights)
    return ParticleFilter(rand(D, length(states)))
end

In problems with discrete observations, we can also perform particle belief updates with rejection. Here, each state $s_i$ in the filter is propagated to obtain successor states $s'_i$ as before. However, observations $o_i$ are simultaneously sampled for each successor state. Any $o_i$ that does not equal the true observation $o$ is rejected, and a new $s'_i$ and $o_i$ are sampled until the observations match. This particle filter with rejection is implemented in algorithm 19.7.

struct RejectionParticleFilter
    states # vector of state samples
end

function update(b::RejectionParticleFilter, 𝒫, a, o)
    T, O = 𝒫.T, 𝒫.O
    states = similar(b.states)
    i = 1
    while i ≤ length(states)
        s = rand(b.states)
        s′ = rand(T(s, a))
        if rand(O(a, s′)) == o
            states[i] = s′
            i += 1
        end
    end
    return RejectionParticleFilter(states)
end

As the number of particles in a particle filter increases, the distribution represented by the particles approaches the true posterior distribution. Unfortunately, particle filters can fail in practice. Low particle coverage and the stochastic nature of the resampling procedure can cause there to be no particles near the true
state. This problem of particle deprivation can be somewhat mitigated by several strategies. A motivational example is given in example 19.8.

Spelunker Joe is lost in a grid-based maze. He lost his lantern, so can only observe his surroundings by touch. At any given moment, Joe can tell whether his location in the maze has walls in each cardinal direction. Joe is fairly confident in his ability to feel walls, so he assumes his observations are perfect.

Joe uses a particle filter to track his belief over time. At some point he stops to rest. He continues to run his particle filter to update his belief.

The initial belief has one particle in each grid location that matches his current observation of a wall to the north and south. Spelunker Joe does not move and does not gain new information, so his belief should not change over time. Due to the stochastic nature of resampling, subsequent beliefs may not contain all of the initial states. Over time, his belief will continue to lose states until it only contains a single state. It is possible that this state is not where Spelunker Joe is located.

19.7 Particle Injection

Particle injection involves injecting random particles to protect against particle deprivation. Algorithm 19.8 injects a fixed number of particles from a broader distribution, such as a uniform distribution over the state space.\footnote{For robotic localization problems, it is common practice to inject particles from a uniform distribution over all possible robot poses, weighted by the current observation.} While particle injection can help prevent particle deprivation, it also reduces the accuracy of the posterior belief represented by the particle filter.

Instead of using a fixed number of injected particles at each update, we can take a more adaptive approach. When the particles are all being given very low weight, then we generally want to inject more particles. It might be tempting to choose the number of injected particles based solely on the mean weight of
the current set of particles. However, doing so can make the success of the filter sensitive to naturally low observation probabilities in the early periods when the filter is still converging or in moments of high sensor noise.\textsuperscript{12}

Algorithm 19.9 presents an \textit{adaptive injection} algorithm that keeps track of two exponential moving averages of the mean particle weight and bases the number of injections on their ratio.\textsuperscript{13} If $w_{\text{mean}}$ is the current mean particle weight, the two moving averages are updated according to

\begin{align}
    w_{\text{fast}} &\leftarrow w_{\text{fast}} + \alpha (w_{\text{mean}} - w_{\text{fast}}) \\
    w_{\text{slow}} &\leftarrow w_{\text{slow}} + \alpha (w_{\text{mean}} - w_{\text{slow}})
\end{align}

where $0 \leq \alpha_{\text{slow}} < \alpha_{\text{fast}} \leq 1$.

The number of injected samples in a given iteration is obtained by comparing the fast and slow mean particle weights:\textsuperscript{14}

\[
m_{\text{inject}} = \left\lfloor m \max \left(0, 1 - \nu \frac{w_{\text{fast}}}{w_{\text{slow}}} \right) \right\rfloor
\]

The scalar $\nu \geq 1$ allows us to tune the injection rate.

\section*{19.8 Summary}

- Partially observable Markov decision processes (POMDPs) extend MDPs to include state uncertainty.
mutable struct AdaptiveInjectionParticleFilter
    states  # vector of state samples
    w_slow  # slow moving average
    w_fast  # fast moving average
    α_slow  # slow moving average parameter
    α_fast  # fast moving average parameter
    ν       # injection parameter
    D_inject # injection distribution
end

function update(b::AdaptiveInjectionParticleFilter, ℙ, a, o)
    T, O = ℙ.T, ℙ.O
    w_slow, w_fast, α_slow, α_fast, ν, D_inject =
        b.w_slow, b.w_fast, b.α_slow, b.α_fast, b.ν, b.D_inject
    states = [rand(T(s, a)) for s in b.states]
    weights = [O(a, s', o) for s' in states]
    w_mean = mean(weights)
    w_slow += α_slow*(w_mean - w_slow)
    w_fast += α_fast*(w_mean - w_fast)
    m = length(states)
    m_inject = round(Int, m * max(0, 1.0 - ν*w_fast / w_slow))
    D = SetCategorical(states, weights)
    states = vcat(rand(D, m - m_inject), rand(D_inject, m_inject))
    b.w_slow, b.w_fast = w_slow, w_fast
    return AdaptiveInjectionParticleFilter(states,
                                            w_slow, w_fast, α_slow, α_fast, ν, D_inject)
end

Algorithm 19.9. A particle filter with adaptive injection, which maintains fast and slow exponential moving averages $w_{fast}$ and $w_{slow}$ of the mean particle weight with smoothness factors $α_{fast}$ and $α_{slow}$, respectively. Particles are only injected if the fast moving average of the mean particle weight is less than $1/ν$ of the slow moving average. Recommended values from the original paper are $α_{fast} = 0.1$, $α_{slow} = 0.001$, and $ν = 2$. 

Sample code:
```
Spelunker Joe from example 19.9 now moves one tile east and moves all particles in his particle filter one tile to the east as well. He now only senses walls to the north and east, and unfortunately, this observation does not agree with any of the updated particles in his filter. He decides to use adaptive injection to fix his particle deprivation problem. Below we see how his filter injects particles from a uniform random distribution, along with the values for the fast and slow filters.

Example 19.9. A particle filter with adaptive injection $\alpha_{\text{slow}} = 0.01, \alpha_{\text{fast}} = 0.3$, and $\nu = 2.0$ starting from a deprived state with 16 identical particles. The moving averages are initialized to 1 to reflect a long period of observations that perfectly match every particle in the filter. Over the next iterations, these moving averages change at different rates based on the quantity of particles that match the observation.
- The uncertainty requires agents in a POMDP to maintain a belief over their state.

- Beliefs for POMDPs with discrete state spaces can be represented using categorical distributions and can be updated analytically.

- Beliefs for linear Gaussian POMDPs can be represented using Gaussian distributions and can also be updated analytically.

- Beliefs for nonlinear continuous POMDPs can also be represented using Gaussian distributions but cannot typically be updated analytically. In this case the extended Kalman filter and the unscented Kalman filter can be used.

- Continuous problems can sometimes be modeled assuming they are linear Gaussian.

- Particle filters approximate the belief with a large collection of state particles.

19.9 Exercises

Exercise 19.1. Can every MDP be framed as a POMDP?

Solution: Yes. The POMDP formulation extends the MDP formulation by introducing state uncertainty in the form of the observation distribution. Any MDP can be framed as a POMDP with \( \mathcal{O} = \mathcal{S} \) and \( O(o \mid a, s') = (o = s') \).

Exercise 19.2. What is the belief update for a discrete POMDP with no observation? What is the belief update for a POMDP with linear Gaussian dynamics with no observation?

Solution: If an agent in a POMDP without an observation with belief \( b \) takes an action \( a \), the new belief \( b' \) can be calculated as follows:

\[
b'(s') = P(s' \mid b, a) = \sum_s P(s' \mid a, b, s)P(s \mid b, a) = \sum_s T(s' \mid s, a)b(s)
\]

This belief update is equivalent to having a uniform observation distribution. A POMDP with linear Gaussian dynamics that has no observation will update its belief using only the Kalman filter predict step in equation (19.12).

Exercise 19.3. An autonomous vehicle represents its belief over its position using a multivariate normal distribution. It comes to a rest at a traffic light, and the belief updater continues to run while it sits. Over time, the belief concentrates and becomes extremely confident in a particular location. Why might this be a problem? How might this extreme confidence be avoided?
Solution: Yes, overconfidence in a belief can be a problem when the models or belief updates do not perfectly represent reality. The overconfident belief may have converged on a state that does not match the true state. Once the vehicle moves again, new observations may be inconsistent with the belief and result in poor estimates. To help address this issue, we can require that the values of the diagonal elements of the covariance matrix be above some threshold.

Exercise 19.4. Consider tracking our belief over the dud rate for widgets produced at a factory. We use a Poisson distribution to model the probability that \( k \) duds are produced in one day of factory operation given that the factory has a dud rate of \( \lambda \): \[
P(k \mid \lambda) = \frac{1}{k!} \lambda^k e^{-\lambda} ...
\]

Suppose that our initial belief over the dud rate follows a gamma distribution: \[
p(\lambda \mid \alpha, \beta) = \frac{\beta^\alpha}{\Gamma(\alpha)} \lambda^{\alpha-1} e^{-\beta \lambda}
\]
where \( \lambda \in (0, \infty) \) and the belief is parameterized by the shape \( \alpha > 0 \) and the rate \( \beta > 0 \). After a day of factory operation, we observe that \( d \geq 0 \) duds were produced. Show that our updated belief over the dud rate is also a gamma distribution.\(^{15}\)

Solution: We seek the posterior distribution \( p(\lambda \mid d, \alpha, \beta) \), which we can obtain through Bayes’ rule:

\[
p(\lambda \mid d, \alpha, \beta) \propto p(d \mid \lambda) p(\lambda \mid \alpha, \beta)
\]
\[
\propto \frac{1}{d!} \lambda^d e^{-\lambda} \frac{\beta^\alpha}{\Gamma(\alpha)} \lambda^{\alpha-1} e^{-\beta \lambda}
\]
\[
\propto \lambda^{d+\alpha-1} e^{-(\beta+1)\lambda}
\]

This is a gamma distribution:

\[
p(\lambda \mid \alpha + d, \beta + 1) = \frac{(\beta + 1)^{a+d}}{\Gamma(a+d)} \lambda^{a+d-1} e^{-(\beta+1)\lambda}
\]

Exercise 19.5. Why are particle filters with rejection not used for updating beliefs in POMDPs with continuous observations?

Solution: Rejection sampling requires repeatedly sampling the transition and observation functions until the sampled observation matches the true observation. The probability of sampling any particular value in a continuous probability distribution is zero, making rejection sampling run forever. In practice, we would use a finite representation for continuous values such as 64-bit floating point numbers, but rejection sampling can run for an extremely long time for each particle.
Exercise 19.6. Explain why Spelunker Joe would not benefit from switching to a particle filter with adaptive injection with $\nu \geq 1$ in example 19.8.

Solution: The particle filter with adaptive injection injects new particles when $\nu \frac{w_{\text{fast}}}{w_{\text{slow}}}$ is less than 1. Spelunker Joe assumes perfect observations and has a belief with particles that match his current observation. Thus, every particle has a weight of 1, and so both $w_{\text{fast}}$ and $w_{\text{slow}}$ are 1. It follows that $\frac{w_{\text{fast}}}{w_{\text{slow}}}$ is always 1, leading to no new particles.

Exercise 19.7. Why is the injection rate scalar $\nu$ in a particle filter with adaptive injection typically not set to a value less than 1?

Solution: Particle injection was designed to inject particles when the current observations have lower likelihood than a historic trend over the observation likelihood. Thus, injection typically only occurs when the short-term estimate of the mean particle weight $w_{\text{fast}}$ is less than the long-term estimate of the mean particle weight $w_{\text{slow}}$. If $\nu < 1$, then particles can still be generated even if $w_{\text{fast}} \geq w_{\text{slow}}$, despite indicating that current observations have higher likelihood than the past average.

Exercise 19.8. Suppose we are dropped into a rectangular forest at an initial location chosen uniformly at random. We do not know which direction we are facing. Fortunately, we do know the dimensions of the forest (it has width $w$ and length much greater than its width). We can move in a continuous path, continuously observing whether we are still in the forest. How can we apply belief updating to this problem? Below are three possible policies, each defining a different path. Which of these policies are guaranteed to escape the forest? Which policy is best?

Solution: Our initial belief is a uniform distribution over all two-dimensional locations and orientations (states) in the forest. We can represent an updated belief using the path we have traveled thus far. If we are still in the forest, our belief consists of all states that can be reached from a state within the forest by following our path, while remaining entirely in the forest. As soon as we exit the forest, our belief consists of all states that reach the edge by following our path, while remaining entirely in the forest.

---

16 This problem was motivated by Richard Bellman’s “Lost in a Forest Problem,” in which we start off at a random location and orientation in a forest with known geometry and must find a policy that minimizes the average (or maximum) time to exit. R. Bellman, “Minimization Problem,” Bulletin of the American Mathematical Society, vol. 62, no. 3, p. 270, 1956.
Of the given policies, only the last two are guaranteed to escape the forest. The path formed by the two perpendicular segments and by the two sides of the equilateral triangle will always intersect with the forest’s border. The straight segment, however, may not leave the forest. We prefer the shorter of the two escaping policies, which is the equilateral triangle.

**Exercise 19.9.** Algorithm 19.2 checks whether the updated belief is a zero vector. When can a belief update yield a zero vector? Why might this arise in real-world applications?

**Solution:** A zero belief vector can result from an observation \( o \) that is considered impossible. This situation can arise after taking action \( a \) from belief \( b \) when \( O(o \mid a, s') = 0 \) for all possible next states \( s' \) according to \( b \) and our transition model. Algorithm 19.2 handles this case by returning a uniform belief. In practical applications, there may be a mismatch between the model and the real world. We generally want to be careful to avoid assigning zero probability to observations, just in case our belief, transition, or observations models are incorrect.

**Exercise 19.10.** Suppose we are performing in-flight monitoring of an aircraft. The aircraft is either in a state of normal operation \( s^0 \) or a state of malfunction \( s^1 \). We receive observations through the absence of a warning \( w^0 \) or the presence of a warning \( w^1 \). We can choose to allow the plane to continue to fly \( m^0 \) or send the plane in for maintenance \( m^1 \). We have the following transition and observation dynamics, where we assume the warnings are independent of the actions given the status of the plane:

\[
T(s^0 \mid s^0, m^0) = 0.95 \quad O(w^0 \mid s^0) = 0.99
\]
\[
T(s^0 \mid s^0, m^1) = 1 \quad O(w^1 \mid s^0) = 0.7
\]
\[
T(s^1 \mid s^1, m^0) = 1 \quad O(w^0 \mid s^1) = 0.99
\]
\[
T(s^0 \mid s^1, m^1) = 0.98 \quad O(w^1 \mid s^1) = 0.7
\]

Given the initial belief \( b = [0.95, 0.05] \), compute the updated belief \( b' \) given that we allow the plane to continue to fly and we observe a warning.

**Solution:** Using equation (19.7), we update the belief for \( s^0 \):

\[
b'(s^0) \propto O(w^1 \mid s^0) \sum_s T(s^0 \mid s, m^0) b(s)
\]
\[
b'(s^0) \propto O(w^1 \mid s^0) (T(s^0 \mid s^0, m^0) b(s^0) + T(s^0 \mid s^1, m^0) b(s^1))
\]
\[
b'(s^0) \propto (1 - 0.99)(0.95 \times 0.95 + (1 - 1) \times 0.05) = 0.009025
\]
We repeat the update for $s^1$:

\[
b'(s^1) \propto O(w^1 \mid s^1) \sum_s T(s^1 \mid s, m^0) b(s)
\]

\[
b'(s^1) \propto O(w^1 \mid s^1)(T(s^1 \mid s^0, m^0)b(s^0) + T(s^1 \mid s^1, m^0)b(s^1))
\]

\[
b'(s^1) \propto 0.7((1 - 0.95) \times 0.95 + 1 \times 0.05) = 0.06825
\]

After normalization, we obtain the updated belief:

\[
b'(s^0) = \frac{b'(s^0)}{b'(s^0) + b'(s^1)} \approx 0.117
\]

\[
b'(s^1) = \frac{b'(s^1)}{b'(s^0) + b'(s^1)} \approx 0.883
\]

\[
b' \approx [0.117, 0.883]
\]

**Exercise 19.11.** Consider a robot moving along a line with position $x$, velocity $v$, and acceleration $a$. At each timestep, we directly control the acceleration and we observe the velocity. The equations of motion for the robot are

\[
x' = x + v \Delta t + \frac{1}{2} a \Delta t^2
\]

\[
v' = v + a \Delta t
\]

where $\Delta t$ is the duration of each step. Suppose we would like to implement a Kalman filter to update our belief. The state vector is $s = [x, v]$. Determine $T_s$, $T_a$, and $O_s$.

**Solution:** The transition and observation dynamics can be written in linear form as follows

\[
\begin{bmatrix}
x' \\
v'
\end{bmatrix} = 
\begin{bmatrix}
1 & \Delta t \\
0 & 1
\end{bmatrix}
\begin{bmatrix}
x \\
v
\end{bmatrix} + 
\begin{bmatrix}
\frac{1}{2} \Delta t^2 \\
\Delta t
\end{bmatrix}
\begin{bmatrix}
a
\end{bmatrix}
\]

\[
o = 
\begin{bmatrix}
0 & 1
\end{bmatrix}
\begin{bmatrix}
x \\
v
\end{bmatrix}
\]

Through these equations, we can identify $T_s$, $T_a$, and $O_s$

\[
T_s = 
\begin{bmatrix}
1 & \Delta t \\
0 & 1
\end{bmatrix}
\quad T_a = 
\begin{bmatrix}
\frac{1}{2} \Delta t^2 \\
\Delta t
\end{bmatrix}
\quad O_s = 
\begin{bmatrix}
0 & 1
\end{bmatrix}
\]

**Exercise 19.12.** Consider a robot with a differential drive moving in two dimensions at a constant speed $v$. The robot’s state is its position $(x, y)$ and its heading $\theta$. At each timestep, we control the robot’s turn rate $\omega$. The equations of motion for the robot are

\[
x' = x + v \cos(\theta) \Delta t
\]

\[
y' = y + v \sin(\theta) \Delta t
\]

\[
\theta' = \theta + \omega \Delta t
\]
This transition function is nonlinear. What is its linearization, $T_s$, as a function of the state $s = [x, y, \theta]$?

Solution: The linearization is given by the Jacobian:

$$T_s = \begin{bmatrix}
\frac{\partial x'}{\partial x} & \frac{\partial x'}{\partial y} & \frac{\partial x'}{\partial \theta} \\
\frac{\partial y'}{\partial x} & \frac{\partial y'}{\partial y} & \frac{\partial y'}{\partial \theta} \\
\frac{\partial \theta'}{\partial x} & \frac{\partial \theta'}{\partial y} & \frac{\partial \theta'}{\partial \theta}
\end{bmatrix} = \begin{bmatrix}
1 & 0 & -v \sin(\theta) \Delta t \\
0 & 1 & v \cos(\theta) \Delta t \\
0 & 0 & 1
\end{bmatrix}$$

This linearization can be used in an extended Kalman filter to maintain a belief.

Exercise 19.13. Compute the set of sigma points and weights with $\lambda = 2$ for a multivariate Gaussian distribution with

$$\mu = \begin{bmatrix} 1 \\ 2 \end{bmatrix}, \quad \Sigma = \begin{bmatrix} 4 & 0 \\ 0 & 2.25 \end{bmatrix}$$

Solution: Since we have a two-dimensional Gaussian distribution and we are given $\lambda = 2$, we need to compute $2n + 1 = 5$ sigma points. We need to compute the square-root matrix $B = \sqrt{(n+\lambda)\Sigma}$, such that $BB^T = (n+\lambda)\Sigma$. Since the scaled covariance matrix is diagonal, the square-root matrix is simply the element-wise square root of $(n+\lambda)\Sigma$

$$\sqrt{(n+\lambda)\Sigma} = \sqrt{(2+2) \begin{bmatrix} 4 & 0 \\ 0 & 2.25 \end{bmatrix}} = \begin{bmatrix} 2 & 0 \\ 0 & 3 \end{bmatrix}$$

Now, we can compute the sigma points and weights:

$$s_1 = \begin{bmatrix} 1 \\ 2 \end{bmatrix}, \quad w_1 = \frac{2}{2+2} = \frac{1}{2}$$

$$s_2 = \begin{bmatrix} 1 \\ 2 \end{bmatrix} + \begin{bmatrix} 4 \\ 0 \end{bmatrix} = \begin{bmatrix} 5 \\ 2 \end{bmatrix}, \quad w_2 = \frac{1}{2(2+2)} = \frac{1}{8}$$

$$s_3 = \begin{bmatrix} 1 \\ 2 \end{bmatrix} - \begin{bmatrix} 4 \\ 0 \end{bmatrix} = \begin{bmatrix} -3 \\ 2 \end{bmatrix}, \quad w_3 = \frac{1}{2(2+2)} = \frac{1}{8}$$

$$s_4 = \begin{bmatrix} 1 \\ 2 \end{bmatrix} + \begin{bmatrix} 0 \\ 3 \end{bmatrix} = \begin{bmatrix} 1 \\ 5 \end{bmatrix}, \quad w_4 = \frac{1}{2(2+2)} = \frac{1}{8}$$

$$s_5 = \begin{bmatrix} 1 \\ 2 \end{bmatrix} - \begin{bmatrix} 0 \\ 3 \end{bmatrix} = \begin{bmatrix} 1 \\ -1 \end{bmatrix}, \quad w_5 = \frac{1}{2(2+2)} = \frac{1}{8}$$

Exercise 19.14. Using the sigma points and weights from the previous exercise, compute the updated mean and covariance given by the unscented transform through $f(x) = [2x_1, x_1 x_2]$. 

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Solution: The transformed sigma points are

\[
\begin{align*}
    f(s_1) &= \begin{bmatrix} 2 \\ 2 \end{bmatrix} & f(s_2) &= \begin{bmatrix} 10 \\ 10 \end{bmatrix} & f(s_3) &= \begin{bmatrix} -6 \\ -6 \end{bmatrix} & f(s_4) &= \begin{bmatrix} 2 \\ 5 \end{bmatrix} & f(s_5) &= \begin{bmatrix} 2 \\ -1 \end{bmatrix}
\end{align*}
\]

We can reconstruct the mean as the weighted sum of transformed sigma points

\[
\mu' = \sum_i w_i f(s_i)
\]

\[
\mu' = \frac{1}{2} \begin{bmatrix} 2 \\ 2 \end{bmatrix} + \frac{1}{8} \begin{bmatrix} 10 \\ 10 \end{bmatrix} + \frac{1}{8} \begin{bmatrix} -6 \\ -6 \end{bmatrix} + \frac{1}{8} \begin{bmatrix} 2 \\ 5 \end{bmatrix} + \frac{1}{8} \begin{bmatrix} 2 \\ -1 \end{bmatrix} = \begin{bmatrix} 2 \\ 2 \end{bmatrix}
\]

The covariance matrix can be reconstructed from the weighted sum of pointwise covariance matrices

\[
\Sigma' = \sum_i w_i (f(s_i) - \mu') (f(s_i) - \mu')^T
\]

\[
\Sigma' = \frac{1}{2} \begin{bmatrix} 0 & 0 \\ 0 & 0 \end{bmatrix} + \frac{1}{8} \begin{bmatrix} 64 & 64 \\ 64 & 64 \end{bmatrix} + \frac{1}{8} \begin{bmatrix} 64 & 64 \\ 64 & 64 \end{bmatrix} + \frac{1}{8} \begin{bmatrix} 0 & 0 \\ 0 & 9 \end{bmatrix} + \frac{1}{8} \begin{bmatrix} 0 & 0 \\ 0 & 9 \end{bmatrix} = \begin{bmatrix} 16 & 16 \\ 16 & 18.25 \end{bmatrix}
\]

Exercise 19.15. What are some advantages and disadvantages of using a particle filter instead of a Kalman filter?

Solution: A Kalman filter can provide an exact belief update when the system is linear Gaussian. Particle filters can work better when the system is nonlinear and the uncertainty is multi-modal. Particle filters are generally more computationally expensive and may suffer from particle deprivation.

Exercise 19.16. Consider using a particle filter to maintain a belief in a problem where observations are very reliable, with observations having either high or low likelihood. For example, in the Spelunker Joe problem, we can reliably determine which of the four walls are present, allowing us to immediately discount any states that do not match the observation. Why might a particle filter with rejection be a better match than a traditional particle filter for such problems?

Solution: A traditional particle filter produces a set of particles and assigns weights to them according to their observation likelihoods. In problems like the one with Spelunker Joe, many particles may end up with little to no weight. Having many particles with low weight makes the belief vulnerable to particle deprivation. A particle filter with rejection ensures that each particle’s successor state is compatible with the observation, thus mitigating the issue of particle deprivation.
20 Exact Belief State Planning

Because states are not directly observable, the agent must use its past history of actions and observations to inform its belief, which can be represented as a probability distribution over states as discussed in the previous chapter. The objective in a POMDP is to choose actions that maximize the accumulation of reward while interacting with the environment. There are different approaches for computing an optimal policy that maps beliefs to actions given models of the transitions, observations, and rewards.\(^1\) One approach is to convert a POMDP into an MDP and apply dynamic programming. Other approaches include representing policies as conditional plans or as piecewise linear value functions over the belief space. The chapter concludes with an algorithm for computing an optimal policy that is analogous to value iteration for MDPs.

20.1 Belief-State Markov Decision Processes

Any POMDP can be viewed as an MDP that uses beliefs as states, also called a belief-state MDP.\(^2\) The state space of a belief-state MDP is the set of all beliefs \(B\). The action space is identical to that of the POMDP. If the state and observation spaces are discrete, the belief-state transition function for a belief-state MDP is


given by:

\[ T(b' \mid b, a) = P(b' \mid b, a) \]

\[ = \sum_o P(b' \mid b, a, o) P(o \mid b, a) \]

\[ = \sum_o P(b' \mid b, a, o) \sum_s P(o \mid b, a, s) P(s \mid b, a) \]

\[ = \sum_o P(b' \mid b, a, o) \sum_s P(o \mid b, a, s) b(s) \]

\[ = \sum_o (b' = \text{Update}(b, a, o)) \sum_{s'} O(o \mid a, s') \sum_s T(s' \mid s, a) b(s) \]

Above, \( \text{Update}(b, a, o) \) returns the updated belief using the deterministic process discussed in the previous chapter. For continuous problems, we replace the summations with integrals.

The reward function for a belief-state MDP depends on the belief and action taken. It is simply the expected value of the reward:

\[ R(b, a) = \sum_s R(s, a) b(s) \]

Solving belief-state MDPs is challenging because the state space is continuous. We can use the approximate dynamic programming techniques presented in earlier chapters, but we can often do better by taking advantage of the structure of the belief-state MDP, as will be discussed in the remainder of this chapter.

### 20.2 Conditional Plans

There are different ways to represent policies for POMDPs. One approach is to use a **conditional plan** represented as a tree. Figure 20.1 shows an example of a three-step conditional plan with binary action and observation spaces. The nodes correspond to belief states. The edges are annotated with observations, and the nodes are annotated with actions. If we have a plan \( \pi \), the action associated with the root is denoted \( \pi() \) and the subplan associated with observation \( o \) is denoted \( \pi(o) \). Algorithm 20.1 provides an implementation.

A conditional plan tells us what to do in response to our observations up to the horizon represented by the tree. To execute a conditional plan, we start with

![Figure 20.1. An example 3-step conditional plan.](image)
the root node and execute the action associated with it. We proceed down the tree according to our observations, taking the actions associated with the nodes through which we pass.

Suppose we have a conditional plan \( \pi \), and we want to compute its expected utility when starting from state \( s \). This computation can be done recursively:

\[
U^\pi(s) = R(s, \pi()) + \gamma \sum_{s'} T(s' \mid s, \pi()) \sum_{o} O(o \mid \pi(), s') U^{\pi(o)}(s')
\] (20.8)

An implementation for this procedure is given in algorithm 20.2.

**Algorithm 20.2.** The conditional plan data structure consisting of an action and a mapping from observations to subplans. The \texttt{subplans} field is a \texttt{Dict} from observations to conditional plans. For convenience, we have created a special constructor for plans consisting of a single node.

If we do not know the current state exactly, we can compute the utility of our belief \( b \) as follows:

\[
U^\pi(b) = \sum_{s} b(s) U^\pi(s)
\] (20.9)

Example 20.1 shows how to compute the utility associated with a three-step conditional plan.

Now that we have a way to evaluate conditional plans up to some horizon \( h \), we can compute the optimal \( h \)-step value function:

\[
U^*(b) = \max_{\pi} U^\pi(b)
\] (20.10)
Consider the following 3-step conditional plan for the crying baby problem:

\[
\begin{align*}
\text{ignore} & \quad \text{crying} \quad \text{feed} \quad \text{quiet} \quad \text{ignore} \\
\text{quiet} & \quad \text{ignore} \quad \text{crying} \quad \text{feed} \quad \text{quiet} \quad \text{ignore}
\end{align*}
\]

In this plan, we begin by ignoring the baby. If we observe any crying, we feed the baby. If we do not observe any crying, we ignore the baby. Our third action again feeds if there is crying.

The expected utility for this plan in belief space is plotted alongside a 3-step plan that always feeds the baby and one that always ignores the baby.

We find that the given plan is not universally better than either always ignoring or always feeding the baby.
An optimal action can be generated from the action associated with the root of a maximizing \( \pi \).

Solving an \( h \)-step POMDP by directly enumerating all \( h \)-step conditional plans is generally computationally intractable as shown in figure 20.2. There are \( (|O|^h - 1)/(|O| - 1) \) nodes in an \( h \)-step plan. In general, any action can be inserted into any node, resulting in \( |A|(|O|^{h-1})/(|O|-1) \) possible \( h \)-step plans. This exponential growth means enumerating over all plans is intractable even for modest values of \( h \). As will be discussed later in this chapter, there are alternatives to explicitly enumerating over all possible plans.

## 20.3 Alpha Vectors

We can rewrite equation (20.9) in vector form:

\[
U^\pi(b) = \sum_s b(s)U^\pi(s) = \alpha^\top \pi b \tag{20.11}
\]

The vector \( \alpha_\pi \) is called an alpha vector and contains the expected utility under plan \( \pi \) for each state. As with belief vectors, alpha vectors have dimension \( |S| \). Unlike beliefs, the components in alpha vectors represent utilities and not probability masses. Algorithm 20.3 shows how to compute an alpha vector.

```plaintext
function alphavector(\( \mathcal{P} ::\text{POMDP}, \pi ::\text{ConditionalPlan} \))
return [evaluate_plan(\( \mathcal{P}, \pi, s \)) for s in \( \mathcal{P}.\mathcal{S} \)]
end
```

Each alpha vector defines a hyperplane in belief space. The optimal value function equation (20.11) is the maximum over these hyperplanes,

\[
U^*(b) = \max_\pi \alpha^\top_\pi b \tag{20.12}
\]

making the value function piecewise-linear and convex.\(^4\)

An alternative to using a conditional plan to represent a policy is to use a set of alpha vectors \( \Gamma \), each annotated with an action. Although not practical, one way to generate the set \( \Gamma \) is to enumerate the set of \( h \)-step conditional plans and compute their alpha vectors. The action associated with an alpha vector is the action at the root of the associated conditional plan. We execute a policy represented by \( \Gamma \) by updating our belief state and executing the action associated with the dominating

\(^4\) The optimal value function for continuous-state POMDPs is also convex, as can be seen by approximating the POMDP through state space discretization and taking the limit as the number of discrete states approaches infinity.
alpha vector at the new belief $b$. The dominating alpha vector $\alpha$ at $b$ is the one that maximizes $\alpha^T b$. This strategy can be used to select actions beyond the horizon of the original conditional plans. Algorithm 20.4 provides an implementation.

Algorithm 20.4. An alpha vector policy is defined in terms of a set of alpha vectors $\Gamma$ and an array of associated actions $a$. Given the current belief $b$, it will find the alpha vector that gives the highest value at that belief point. It will return the associated action.

If we use one-step lookahead, we do not have to keep track of the actions associated with the alpha vectors in $\Gamma$. The one-step lookahead action from belief $b$ using the value function represented by $\Gamma$, denoted $U^\Gamma$, is

$$\pi^\Gamma(b) = \arg \max_a \left[ R(b,a) + \gamma \sum_{a} P(o \mid b,a) U^\Gamma(\text{Update}(b,a,o)) \right]$$

(20.13)

where

$$P(o \mid b,a) = \sum_{s} P(o \mid s,a)b(s)$$

(20.14)

$$P(o \mid s,a) = \sum_{s'} T(s' \mid s,a)O(o \mid s',a)$$

(20.15)

Algorithm 20.5 provides an implementation. Example 20.2 demonstrates one-step lookahead on the crying baby problem.

20.4 Pruning

If we have a collection of alpha vectors $\Gamma$, we may want to prune alpha vectors that do not contribute to our representation of the value function or plans that are not optimal for any belief. Removing such alpha vectors or plans can improve computational efficiency. We can check whether an alpha vector $\alpha$ is dominated by
Algorithm 20.5. A policy represented by a set of alpha vectors $\Gamma$. It uses one-step lookahead to produce an optimal action and associated utility. Equation (20.13) is used to compute the lookahead.

function lookahead($\mathcal{P}$::POMDP, $U$, $b$::Vector, $a$)
    $S$, $O$, $T$, $O$, $R$, $\gamma$ = $\mathcal{P}$.S, $\mathcal{P}$.O, $\mathcal{P}$.T, $\mathcal{P}$.O, $\mathcal{P}$.R, $\mathcal{P}$.$\gamma$
    $r$ = $\sum(R(s,a) \cdot b[i])$ for $(i,s)$ in enumerate($S$)
    $\text{Posa}(o,s,a)$ = $\sum(O(a,s',o) \cdot T(s,a,s')$ for $s'$ in $S$)
    $\text{Poba}(o,b,a)$ = $\sum(b[i] \cdot \text{Posa}(o,s,a))$ for $(i,s)$ in enumerate($S$)
    return $r + \gamma \cdot \sum(\text{Poba}(o,b,a) \cdot U(\text{update}(b, \mathcal{P}, a, o))$ for $o$ in $O$)
end

function greedy($\mathcal{P}$::POMDP, $U$, $b$::Vector)
    $u$, $a$ = findmax($a$->lookahead($\mathcal{P}$, $U$, $b$, $a$), $\mathcal{P}$.\mathcal{A})
    return $(a=a, u=u)$
end

struct LookaheadAlphaVectorPolicy
    $\mathcal{P}$ # POMDP problem
    $\Gamma$ # alpha vectors
end

function utility($\pi$::LookaheadAlphaVectorPolicy, $b$)
    return $\max(\alpha \cdot b$ for $\alpha$ in $\pi.\Gamma$)
end

function greedy($\pi$, $b$)
    $U(b)$ = utility($\pi$, $b$)
    return greedy($\pi.\mathcal{P}$, $U$, $b$)
end

($\pi$::LookaheadAlphaVectorPolicy)($b$) = greedy($\pi$, $b$).$a$
Consider using one-step lookahead on the crying baby problem with a value function given by the alpha vectors \([-3.7, -15]\) and \([-2, -21]\). Suppose our current belief is \(b = [0.5, 0.5]\), meaning that we believe it is equally likely the baby is hungry as not hungry. We apply equation (20.13):

\[
\begin{align*}
R(b, \text{feed}) &= -10 \\
\gamma P(\text{crying} \mid b, \text{feed}) U(\text{Update}(b, \text{feed}, \text{crying})) &= -0.18 \\
\gamma P(\text{quiet} \mid b, \text{feed}) U(\text{Update}(b, \text{feed}, \text{quiet})) &= -1.62 \\
\rightarrow Q(b, \text{feed}) &= -11.8 \\
R(b, \text{ignore}) &= -5 \\
\gamma P(\text{crying} \mid b, \text{ignore}) U(\text{Update}(b, \text{ignore}, \text{crying})) &= -6.09 \\
\gamma P(\text{quiet} \mid b, \text{ignore}) U(\text{Update}(b, \text{ignore}, \text{quiet})) &= -2.81 \\
\rightarrow Q(b, \text{ignore}) &= -13.9 \\
R(b, \text{sing}) &= -5.5 \\
\gamma P(\text{crying} \mid b, \text{sing}) U(\text{Update}(b, \text{sing}, \text{crying})) &= -6.09 \\
\gamma P(\text{quiet} \mid b, \text{sing}) U(\text{Update}(b, \text{sing}, \text{quiet})) &= -1.85 \\
\rightarrow Q(b, \text{sing}) &= -14.0
\end{align*}
\]

We use \(Q(b, a)\) to represent the action value function from a belief state. The policy predicts that feeding the baby will result in the highest expected utility, so it takes that action.
the alpha vectors in a set $\Gamma$ by solving a linear program to maximize the utility gap $\delta$ that vector achieves over all other vectors:\footnote{Constraints of the form $a \geq b$ are element-wise. That is, we mean $a_i \geq b_i$ for all $i$.}

$$\begin{align*}
\text{maximize} & \quad \delta \\
\text{subject to} & \quad b \geq 0 \\
& \quad 1^\top b = 1 \\
& \quad \alpha^\top b \geq \alpha'^\top b + \delta, \quad \alpha' \in \Gamma
\end{align*}$$

(20.16)

The first two constraints ensure that $b$ is a categorical distribution, and the final set of constraints ensures that we find a belief vector for which $\alpha$ has a higher expected reward than all alpha vectors in $\Gamma$. If, after solving the linear program, the utility gap $\delta$ is negative, then $\alpha$ is dominated. If $\delta$ is positive, then $\alpha$ is not dominated and $b$ is a belief at which $\alpha$ is not dominated. Algorithm 20.6 provides an implementation for solving equation (20.16) to determine a belief, if one exists, where $\delta$ is most positive.

```plaintext
function find_maximal_belief(α, Γ)
    m = length(α)
    if isempty(Γ)
        return fill(1/m, m) # arbitrary belief
    end
    model = Model(GLPK.Optimizer)
    @variable(model, δ)
    @variable(model, b[i=1:m] ≥ 0)
    @constraint(model, sum(b) == 1.0)
    for a in Γ
        @constraint(model, (α-a)⋅b ≥ δ)
    end
    @objective(model, Max, δ)
    optimize!(model)
    return value(δ) > 0 ? value.(b) : nothing
end
```

Algorithm 20.6. A method for finding the belief vector $b$ for which the alpha vector $\alpha$ improves the most compared to the set of alpha vectors $\Gamma$. Nothing is returned if no such belief exists. The packages JuMP.jl and GLPK.jl provide a mathematical optimization framework and a solver for linear programs, respectively.

Algorithm 20.7 provides a procedure that uses algorithm 20.6 for finding the dominating alpha vectors in a set $\Gamma$. Initially, all of the alpha vectors are candidates for being dominating. We then choose one of the candidates and determine the belief $b$ where the candidate leads to the greatest improvement in value compared to all other alpha vectors in the dominating set. If the candidate does not bring improvement, we remove it as a candidate. If it does bring improvement, we move an alpha vector from the candidate set that brings the greatest improvement
at \( b \) to the dominating set. The process continues until there are no longer any candidates. We can prune away any alpha vectors and associated conditional plans that are not dominating at any belief point. Example 20.3 demonstrates pruning on the crying baby problem.

Algorithm 20.7. A method for pruning dominated alpha vectors and associated plans. The \texttt{find_dominating} function identifies all of the dominating alpha vectors in the set \( \Gamma \). It uses binary vectors \texttt{candidates} and \texttt{dominating} to track which alpha vectors are candidates for inclusion in the dominating set and which are currently in the dominating set, respectively.

\begin{verbatim}
function find_dominating(\Gamma)
    n = length(\Gamma)
    candidates, dominating = trues(n), falses(n)
    while any(candidates)
        i = findfirst(candidates)
        b = find_maximal_belief(\Gamma[i], \Gamma[dominating])
        if b == nothing
            candidates[i] = false
        else
            k = argmax([candidates[j] ? b \cdot \Gamma[j] : -Inf for j in 1:n])
            candidates[k], dominating[k] = false, true
        end
    end
    return dominating
end

function prune(plans, \Gamma)
    d = find_dominating(\Gamma)
    return (plans[d], \Gamma[d])
end
\end{verbatim}

20.5 Value Iteration

The value iteration algorithm for MDPs can be adapted for POMDPs.\(^6\) POMDP value iteration (algorithm 20.8) begins by constructing all one-step plans. We prune any plans that are never optimal for any initial belief. Then, we expand all combinations of one-step plans to produce two-step plans. Again, we prune any suboptimal plans from consideration. This procedure of alternating between expansion and pruning is repeated until the desired horizon is reached. Figure 20.3 demonstrates value iteration on the crying baby problem.

---

\(^6\) This section describes a version of value iteration in terms of conditional plans and alpha vectors. For a version that only uses alpha vectors, see A. R. Cassandra, M. L. Littman, and N. L. Zhang, “Incremental Pruning: A Simple, Fast, Exact Method for Partially Observable Markov Decision Processes,” in Conference on Uncertainty in Artificial Intelligence (UAI), 1997.
We can construct all two-step plans for the crying baby problem. There are $3^3 = 27$ such plans.

The expected utility for each plan in belief space is plotted below. We find that two plans dominate all others. These dominating plans are the only ones that need be considered as subplans for optimal three-step plans.

Example 20.3. The expected utility over the belief space for all two-step plans for the crying baby problem (appendix F.7). The thick lines are optimal for some beliefs, whereas the thin lines are dominated.

Algorithm 20.8. Value iteration for POMDPs, which finds the dominating $h$-step plans for a finite-horizon POMDP of horizon $k_{\text{max}}$ by iteratively constructing optimal subplans. The ValueIteration structure is the same as what was defined in algorithm 7.8 in the context of MDPs.
Figure 20.3. POMDP value iteration used to find the optimal value function for the crying baby problem to various horizons.
The expansion step (algorithm 20.9) in this process constructs all possible \( k + 1 \)-step plans from a set of \( k \)-step plans. New plans can be constructed using a new first action and all possible combinations of the \( k \)-step plans as subplans, as shown in figure 20.4. While plans can also be extended by adding additional actions to the ends of subplans, top-level expansion allows alpha vectors constructed for the \( k \)-step plans to be used to efficiently construct alpha vectors for the \( (k + 1) \)-step plans.

Computing the alpha vector associated with a plan \( \pi \) from a set of alpha vectors associated with its subplans can be done as follows. We use \( \alpha_o \) to represent the alpha vector associated with subplan \( \pi(o) \). The alpha vector associated with \( \pi \) is then:

\[
\alpha(s) = R(s, \pi()) + \gamma \sum_{s'} T(s' | s, \pi()) \sum_o O(o | \pi(), s') \alpha_o(s')
\]  

(20.17)

Even for relatively simple problems to shallow depths, computing alpha vectors from subplans in this way is much more efficient than computing them from scratch as in algorithm 20.2.

20.6 Linear Policies

As discussed in section 19.3, the belief state in a problem with linear Gaussian dynamics can be represented by a Gaussian distribution \( \mathcal{N}(\mu_b, \Sigma_b) \). If the reward function is quadratic, then it can be shown that the optimal policy can be computed exactly offline using a process that is often called linear quadratic Gaussian (LQG) control. The optimal action is obtained in an identical manner as in section 7.8, but the \( \mu_b \) computed using the linear Gaussian filter is treated as the true state. With each observation, we simply use the filter to update our \( \mu_b \) and obtain an optimal action by multiplying \( \mu_b \) with the policy matrix from algorithm 7.11. Example 20.4 demonstrates this process.

20.7 Summary

- Exact solutions for POMDPs can typically only be obtained for finite-horizon discrete POMDPs.
function ConditionalPlan(\(\mathcal{P}\)::POMDP, a, plans)
    subplans = Dict(o => π for (o, π) in zip(\(\mathcal{P}\).𝒪, plans))
    return ConditionalPlan(a, subplans)
end

function combine_lookahead(\(\mathcal{P}\)::POMDP, s, a, Γ₀)
    \(\mathcal{S}, \mathcal{O}, \mathcal{T}, \mathcal{O}, \mathcal{R}, \gamma = \mathcal{P}.\mathcal{S}, \mathcal{P}.\mathcal{O}, \mathcal{P}.\mathcal{T}, \mathcal{P}.\mathcal{O}, \mathcal{P}.\mathcal{R}, \mathcal{P}.\gamma \)
    \(U'(s',i) = \sum O(a,s',o) * \alpha[i] \) for (o,α) in zip(\(\mathcal{O}, \Gamma₀\))
    return \( R(s,a) + \gamma * \sum T(s,a,s') * U'(s',i) \) for (i,s') in enumerate(\(\mathcal{S}\))
end

function combine_alphavector(\(\mathcal{P}\)::POMDP, a, Γ₀)
    return [combine_lookahead(\(\mathcal{P}\), s, a, Γ₀) for s in \(\mathcal{P}\).𝒮]
end

function expand(plans, Γ, \(\mathcal{P}\))
    \(\mathcal{S}, \mathcal{A}, \mathcal{O}, \mathcal{T}, \mathcal{O}, \mathcal{R} = \mathcal{P}.\mathcal{S}, \mathcal{P}.\mathcal{A}, \mathcal{P}.\mathcal{O}, \mathcal{P}.\mathcal{T}, \mathcal{P}.\mathcal{O}, \mathcal{P}.\mathcal{R} \)
    plans', Γ' = [], []
    for a in \(\mathcal{A}\)
        # iterate over all possible mappings from observations to plans
        for inds in product([eachindex(plans) for o in \(\mathcal{O}\)])
            π₀ = plans[[inds...]]
            Γ₀ = Γ[[inds...]]
            π = ConditionalPlan(\(\mathcal{P}\), a, π₀)
            α = combine_alphavector(\(\mathcal{P}\), a, Γ₀)
            push!(plans', π)
            push!(Γ', α)
        end
    end
    return (plans', Γ')
end

Algorithm 20.9. The expansion step in value iteration, which constructs all \(k + 1\)-step conditional plans and associated alpha vectors from a set of \(k\)-step conditional plans and alpha vectors. The way we combine alpha vectors of sub-plans follows equation (20.17).
Consider a satellite navigating in two dimensions, neglecting gravity, drag, or other external forces. The satellite can use its thrusters to accelerate in any direction with linear dynamics:

$$\begin{bmatrix}
    x \\
    y \\
    \dot{x} \\
    \dot{y}
\end{bmatrix} \leftarrow \begin{bmatrix}
    1 & 0 & \Delta t & 0 \\
    0 & 1 & 0 & \Delta t \\
    0 & 0 & 1 & 0 \\
    0 & 0 & 0 & 1
\end{bmatrix} \begin{bmatrix}
    x \\
    y \\
    \dot{x} \\
    \dot{y}
\end{bmatrix} + \begin{bmatrix}
    \frac{1}{2} \Delta t^2 & 0 \\
    0 & \frac{1}{2} \Delta t^2 \\
    \Delta t & 0 \\
    0 & \Delta t
\end{bmatrix} \begin{bmatrix}
    \ddot{x} \\
    \ddot{y}
\end{bmatrix} + \epsilon$$

where $\Delta t$ is the duration of a time-step and $\epsilon$ is zero-mean Gaussian noise with covariance $\Delta t / 20I$.

We seek to place the satellite in its orbital slot at the origin, while minimizing fuel use. Our quadratic reward function is:

$$R(s, a) = -s^\top \begin{bmatrix}
    I_{2 \times 2} & 0_{2 \times 2} \\
    0_{2 \times 2} & 0_{2 \times 2}
\end{bmatrix} s - 2a^\top a$$

The satellite’s sensors only estimate its position with the linear observation:

$$o = \begin{bmatrix}
    I_{2 \times 2} & 0_{2 \times 2}
\end{bmatrix} s + \epsilon$$

where $\epsilon$ is zero-mean Gaussian noise with covariance $\Delta t / 10I$.

Below are 50 trajectories from 10-step rollouts using the optimal policy for $\Delta t = 1$ and a Kalman filter to track the belief. In each case, the satellite was started at $s = \mu_b = [-5, 2, 0, 1]$ with $\Sigma_b = [I_0; 0 0.25I]$. 

Example 20.4. An example of an optimal policy used for a POMDP with linear Gaussian dynamics and quadratic reward.
• Policies for these problems can be represented as conditional plans, which are trees that describe the actions to take based on the observations.

• Alpha vectors contain the expected utility when starting from different states and following a particular conditional plan.

• Alpha vectors can also serve as an alternative representation of a POMDP policy.

• POMDP value iteration can avoid the computational burden of enumerating all conditional plans by iteratively computing subplans and pruning those that are suboptimal.

• Linear Gaussian problems with quadratic reward can be solved exactly using methods very similar to those derived for the fully observable case.

20.8 Exercises

Exercise 20.1. Can every POMDP be framed as an MDP?

Solution: Yes. Any POMDP can equivalently be viewed as a belief-state MDP whose state space is the space of beliefs in the POMDP, whose action space is the same as that of the POMDP, and whose transition function is given by equation (20.1).

Exercise 20.2. What are the alpha vectors for the one-step crying baby problem (appendix F.7)? Are all actions dominant?

Solution: There are three one-step conditional plans, one for each action, resulting in three alpha vectors. The optimal one-step policy must choose between these actions given the current belief. The one-step alpha vectors for a POMDP can be obtained from the optimal one-step belief value function:

\[ U^*(b) = \max_a \sum_s b(s) R(s, a) \]

Feeding the baby yields an expected reward:

\[ R(\text{hungry, feed})P(\text{hungry}) + R(\text{sated, feed})P(\text{sated}) \]
\[ = -15P(\text{hungry}) - 5(1 - P(\text{hungry})) \]
\[ = -10P(\text{hungry}) - 5 \]
Singing to the baby yields an expected reward:

\[
R(\text{hungry}, \text{sing})P(\text{hungry}) + R(\text{sated}, \text{sing})P(\text{sated}) \\
= -10.5P(\text{hungry}) - 0.5(1 - P(\text{hungry})) \\
= -10P(\text{hungry}) - 0.5
\]

Ignoring the baby yields an expected reward:

\[
R(\text{hungry}, \text{ignore})P(\text{hungry}) + R(\text{sated}, \text{ignore})P(\text{sated}) \\
= -10P(\text{hungry})
\]

The expected reward for each action is plotted below over the belief space:

We find that under a one-step horizon it is never optimal to feed or sing to the baby. The ignore action is dominant.

**Exercise 20.3.** Why does the implementation of value iteration in algorithm 20.8 call expand plans in algorithm 20.9 rather than evaluating the plan in algorithm 20.2 to obtain alpha vectors for each new conditional plan?

**Solution:** The plan evaluation method applies equation (20.8) recursively to evaluate the expected utility for a conditional plan. Conditional plans grow very large as the horizon increases. POMDP value iteration can save computation by using the alpha vectors for the subplans from the previous iteration:

\[
U^\pi(s) = R(s, \pi()) + \gamma \left[ \sum_{s'} T(s' \mid s, \pi()) \sum_o O(o \mid \pi(), s') \alpha_{s'}^{\pi(o)} \right]
\]

**Exercise 20.4.** Does the number of conditional plans increase faster with the number of actions or with the number of observations?
Solution: Recall that there are $|A|^{(|O|^h-1)/(|O|-1)}$ possible $h$-step plans. Exponential growth ($n^x$) is faster than polynomial growth ($x^n$), and we have better-than exponential growth in $|O|$ and polynomial growth in $|A|$. The number of plans thus increases faster with respect to the number of observations. To demonstrate, let us use $|A| = 3$, $|O| = 3$, and $h = 3$ as a baseline. The baseline has 1,594,323 plans. Incrementing the number of actions results in 67,108,864 plans, whereas incrementing the number of observations results in 10,460,353,203 plans.

Exercise 20.5. Suppose we have a patient, and we are unsure whether or not they have a particular disease. We do have three diagnostic tests, each with different probabilities that they will correctly indicate whether or not the disease is present. While the patient is in our office, we have the option to administer potentially multiple diagnostic tests in sequence. We observe the outcome of each diagnostic test immediately. In addition, we can repeat any diagnostic test multiple times, with the outcomes of all tests being conditionally independent of each other given the presence or absence of the disease. When we are done with the tests, we decide whether to treat the disease or send the patient home without treatment. Explain how you would define the various components of a POMDP formulation.

Solution: We have three states:
1. $s_{\text{healthy}}$: the patient does not have the disease
2. $s_{\text{disease}}$: the patient has the disease
3. $s_{\text{terminal}}$: the interaction is over (terminal state)
We have five actions:
1. $a_1$: administer test 1
2. $a_2$: administer test 2
3. $a_3$: administer test 3
4. $a_{\text{treat}}$: administer treatment and send patient home
5. $a_{\text{stop}}$: send patient home without treatment
We have three observations:
1. $o_{\text{healthy}}$: the outcome of the test (if administered) indicates the patient is healthy
2. $o_{\text{disease}}$: the outcome of the test (if administered) indicates the patient has the disease
3. $o_{\text{terminal}}$: a test was not administered
The transition model would be deterministic with:

$$
T(s' | s, a) = \begin{cases} 
1 & \text{if } a \in \{a_{\text{treat}}, a_{\text{stop}}\} \land s' = s_{\text{stop}} \\
1 & \text{if } s = s' \\
0 & \text{otherwise}
\end{cases}
$$
The reward function would be a function of the cost of administering treatment and each test as well as the cost of not treating the disease if it is indeed present. The reward available from \( s_{\text{terminal}} \) is 0. The observation model assigns probabilities to correct and incorrect observations of the disease state as a result of a diagnostic test from one of the nonterminal states. The initial belief would assign our prior probability to whether or not the patient has the disease, with zero probability assigned to the terminal state.

**Exercise 20.6.** Why might we want to perform the same test multiple times in the previous exercise?

**Solution:** Depending on the probability of incorrect results, we may want to perform the same test multiple times to improve our confidence in whether the patient has the disease or is healthy. The results of the tests are independent given the disease state.

**Exercise 20.7.** Suppose we have three alpha vectors: \([1, 0], [0, 1], \) and \([\theta, \theta], \) for some constant \( \theta \). Under what conditions on \( \theta \) can we prune alpha vectors?

**Solution:** We can prune alpha vectors if \( \theta < 0.5 \) or \( \theta > 1 \). If \( \theta < 0.5 \), then \([\theta, \theta]\) is dominated by the other two alpha vectors. If \( \theta > 1 \), then \([\theta, \theta]\) dominates the other two alpha vectors.

**Exercise 20.8.** We have \( \Gamma = \{[1, 0], [0, 1]\} \) and \( \alpha = [0.7, 0.7] \). What belief \( \mathbf{b} \) maximizes the utility gap \( \delta \) as defined by the linear program in equation (20.16)?

**Solution:** The alpha vectors in \( \Gamma \) are shown in blue and the alpha vector \( \alpha \) is shown in red. We care only about the region where \( 0.3 \leq b_2 \leq 0.7 \) where \( \alpha \) dominates the alpha vectors in \( \Gamma \); in other words, where the red line is above the blue lines. The point where the gap between the red line and the maximum of the blue lines occurs at \( b_2 = 0.5 \) with a gap of \( \delta = 0.2 \). Hence, the belief that maximizes this gap is \( \mathbf{b} = [0.5, 0.5] \).
21 Offline Belief State Planning

In the worst case, an exact solution for a general finite-horizon POMDP is PSPACE-complete, which is a complexity class that includes NP-complete problems and is suspected to include problems even more difficult.\(^1\) General infinite-horizon POMDPs have been shown to be uncomputable.\(^2\) Hence, there has been a tremendous amount of research recently on approximation methods. This chapter discusses different offline POMDP solution methods, which involve performing all or most of the computation prior to execution. We focus on methods that represent the value function as alpha vectors and different forms of interpolation.

21.1 Fully Observable Value Approximation

One of the simplest offline approximation techniques is QMDP, which derives its name from the action value function associated with a fully observed MDP.\(^3\) This approach and several others discussed in this chapter involve iteratively updating a set \(\Gamma\) of alpha vectors as shown in algorithm 21.1. The resulting set \(\Gamma\) defines a value function and a policy that can be used directly or with one-step lookahead.

```
function alphavector_iteration(P::POMDP, M, \(\Gamma\))
    for k in 1:M.k_max
        \(\Gamma\) = update(P, M, \(\Gamma\))
    end
    return \(\Gamma\)
end
```

Algorithm 21.1. Iteration structure for updating a set of alpha vectors \(\Gamma\) used by several of the methods in this chapter. The various methods, including QMDP, differ in their implementation of update. After \(k_{\text{max}}\) iterations, this function returns a policy represented by the alpha vectors in \(\Gamma\).

---


QMDP (algorithm 21.2) constructs a single alpha vector $\alpha_d$ for each action $a$ using value iteration. Each alpha vector is initialized to zero, and then we iterate:

$$
\alpha_d^{(k+1)}(s) = R(s,a) + \gamma \sum_{s'} T(s'|s,a) \max_{a'} \alpha_d^{(k)}(s')
$$

(21.1)

Each iteration requires $O(|A|^2|S|^2)$ operations. Figure 21.1 illustrates the process.

Algorithm 21.2. The QMDP algorithm, which finds an approximately optimal policy for an infinite horizon POMDP with a discrete state and action space, where $k_{max}$ is the number of iterations. QMDP assumes perfect observability.

When QMDP is run to the horizon in finite horizon problems or to convergence for infinite horizon problems, the resulting policy is equivalent to assuming that there will be full observability after taking the first step. Because we can only do
better if we have full observability, QMDP will produce an upper bound on the true optimal value function $U^*(b)$. In other words, $\max_a \alpha_{a}^\top b \geq U^*(b)$ for all $b$.\footnote{Although the value function represented by the QMDP alpha vectors upper bounds the optimal value function, the utility realized by a QMDP policy will not exceed that of an optimal policy in expectation, of course.}

If QMDP is not run to convergence for infinite horizon problems, it might not provide an upper bound. One way to guarantee that QMDP will provide an upper bound after a finite number of iterations is to initialize the value function to some upper bound. One rather loose upper bound is the best-action best-state upper bound, which is the utility obtained from taking the best action from the best state forever:

$$\overline{U}(b) = \max_{s,a} \frac{R(s,a)}{1-\gamma}$$

The assumption of full observability after the first step can cause QMDP to poorly approximate the value of information gathering actions, which are actions that significantly reduce the uncertainty in the state. For example, looking over our shoulder before changing lanes when driving is an information gathering action. QMDP can perform well in problems where the optimal policy does not include costly information gathering.

We can generalize the QMDP approach to problems that may not have a small, discrete state space. In such problems, the iteration in equation (21.1) may not be feasible, but we may use one of the many methods discussed in earlier chapters for obtaining an approximate action value function $Q(s,a)$. This value function might be defined over a high-dimensional, continuous state space using, for example, a neural network representation. The value function evaluated at a belief point is then

$$U(b) = \max_a \int Q(s,a)b(s) \, ds$$

The integral above may be approximated through sampling.

21.2 Fast Informed Bound

As with QMDP, the fast informed bound computes one alpha vector for each action. However, the fast informed bound takes into account, to some extent, the observation model.\footnote{© 2022 Massachusetts Institute of Technology, shared under a under a Creative Commons CC-BY-NC-ND license. 2021-03-07 23:12:00-08:00, revision 80bf4d6, comments to bugs@algorithmsbook.com} The iteration is:

$$\alpha_{d}^{(k+1)}(s) = R(s,a) + \gamma \sum_o \sum_{a'} \sum_{s'} O(o | a, s') T(s' | s, a) \alpha_{d'}^{(k)}(s')$$

which requires $O(|A|^2|S|^2|O|)$ operations per iteration.
The fast informed bound provides an upper bound on the optimal value function. That upper bound is guaranteed to be no looser than that provided by QMDP, and it tends to be tighter. The fast informed bound is implemented in algorithm 21.3 and is used to compute optimal value functions in figure 21.2.

Algorithm 21.3. The fast informed bound algorithm, which finds an approximately optimal policy for an infinite horizon POMDP with discrete state, action, and observation spaces, where \( k_{\text{max}} \) is the number of iterations.

21.3 Fast Lower Bounds

The previous two sections introduced methods that can be used to produce upper bounds on the value function represented as alpha vectors. This section
introduces a couple methods for quickly producing lower bounds represented as alpha vectors without any planning in the belief space. Although the upper bound methods can often be used directly to produce sensible policies, the lower bounds discussed in this section are generally only used to seed other planning algorithms. Figure 21.3 plots the two lower-bound methods discussed in this section.

A common lower bound is the best-action worst-state lower bound (algorithm 21.4). It is the discounted reward obtained by taking the best action in the worst state forever:

\[ r_{\text{baws}} = \max_a \sum_{k=1}^{\infty} \gamma^{k-1} \min_s R(s, a) = \frac{1}{1 - \gamma} \max_a \min_s R(s, a) \]  

(21.5)

This lower bound is represented by a single alpha vector. This bound is typically very loose, but it can be used to seed other algorithms that can tighten the bound, as we will discuss shortly.

```
function baws_lowerbound(\mathcal{P}::POMDP)
    S, \mathcal{A}, R, \gamma = \mathcal{P}.S, \mathcal{P}.A, \mathcal{P}.R, \mathcal{P}.\gamma
    r = \text{maximum}(\text{minimum}(R(s, a) \text{ for } s \text{ in } S) \text{ for } a \text{ in } \mathcal{A}) / (1-\gamma)
    \alpha = \text{fill}(r, \text{length}(S))
    \text{return } \alpha
end
```

The blind lower bound (algorithm 21.5) also represents a lower bound with one alpha vector per action. It makes the assumption that we are forced to commit to a single action forever, blind to what we observe in the future. To compute these alpha vectors, we start with some other lower bound, typically the best-action worst-state lower bound, and then perform some number of iterations:

\[ \alpha_a^{(k+1)}(s) = R(s, a) + \gamma \sum_{s'} T(s' | s, a) \alpha_a^{(k)}(s') \]  

(21.6)

This iteration is similar to the QMDP update in equation (21.1) except that it does not have a maximization over the alpha vectors on the right-hand side.

21.4 Point-Based Value Iteration

QMDP and the fast informed bound generates one alpha vector for each action, but the optimal value function is often better approximated by many more
chapter 21. offline belief state planning

function blind_lowerbound(\( \mathcal{P} \), \( k_{\text{max}} \))

\( \mathcal{S}, \mathcal{A}, T, R, \gamma = \mathcal{P}.\mathcal{S}, \mathcal{P}.\mathcal{A}, \mathcal{P}.T, \mathcal{P}.R, \mathcal{P}.\gamma \)

\( Q(s,a,\alpha) = R(s,a) + \gamma \sum_{(j,s')} (T(s,a,s')|\alpha[j]) \) for \( (j,s') \) in enumerate(\( \mathcal{S} \))

\( \Gamma = [\text{baws_lowerbound}(\mathcal{P}) \text{ for } a \text{ in } \mathcal{A}] \)

for \( k \) in 1: \( k_{\text{max}} \)

\( \Gamma = [[[Q(s,a,\alpha) \text{ for } s \text{ in } \mathcal{S}] \text{ for } (\alpha,a) \text{ in } \text{zip}(\Gamma, \mathcal{A})]] \)

end

return \( \Gamma \)
end

Algorithm 21.5. Implementation of the blind lower bound represented as a set of alpha vectors.

alpha vectors. Point-based value iteration\(^6\) computes \( m \) different alpha vectors \( \Gamma = \{\alpha_1, \ldots, \alpha_m\} \), each associated with different belief points \( B = \{b_1, \ldots, b_m\} \). Methods for selecting these beliefs will be discussed in section 21.7. As before, these alpha vectors define an approximately optimal value function:

\[
U^\Gamma(b) = \max_{\alpha \in \Gamma} \alpha^\top b
\] (21.7)

The algorithm maintains a lower bound on the optimal value function, \( U_\Gamma(b) \leq U^*(b) \) for all \( b \). We initialize our alpha vectors to start with a lower bound and then perform a backup to update the alpha vectors at each point in \( B \). The backup operation (algorithm 21.6) takes a belief \( b \) and a set of alpha vectors \( \Gamma \) and constructs a new alpha vector. The algorithm iterates through every possible action \( a \) and observation \( o \) and extracts the alpha vector from \( \Gamma \) that is maximal at the resulting belief state:

\[
\alpha_{a,o} = \arg \max_{\alpha \in \Gamma} \alpha^\top \text{Update}(b, a, o)
\] (21.8)

Then, for each available action \( a \), we construct a new alpha vector based on these \( \alpha_{a,o} \) vectors:

\[
\alpha_a(s) = R(s,a) + \gamma \sum_{s',o} O(o | a, s') T(s' | s, a) \alpha_{a,o}(s')
\] (21.9)

The alpha vector that is ultimately produced by the backup operation is

\[
\alpha = \arg \max_{\alpha_a} \alpha_a^\top b
\] (21.10)

If \( \Gamma \) is a lower bound, the backup operation will only produce alpha vectors that are still a lower bound.


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2021-03-07 23:12:00-08:00, revision 80bf4d6, comments to bugs@algorithmsbook.com
Repeated application of the backup operation over the beliefs in $B$ gradually increases the lower bound on the value function represented by the alpha vectors until convergence. The converged value function will not necessarily be optimal because $B$ typically does not include all beliefs reachable from the initial belief. However, so long as the beliefs in $B$ are well distributed across the reachable belief space, the approximation may be acceptable. In any case, the resulting value function is guaranteed to provide a lower bound that can be used with other algorithms, potentially online, to further improve the policy.

Point-based value iteration is implemented in algorithm 21.7. Figure 21.4 shows several iterations on an example problem.

Algorithm 21.6. A method for backing up a belief for a POMDP with discrete state and action spaces, where $\Gamma$ is a vector of alpha vectors and $b$ is a belief vector at which to apply the backup. The update method for vector beliefs is defined in algorithm 19.2.

21.5 Randomized Point-Based Value Iteration

Randomized point-based value iteration (algorithm 21.8) is a variation of the point-based value iteration approach from the previous section. The primary difference is in the fact that we are not forced to maintain an alpha vector at every belief in $B$. We initialize the algorithm with a single alpha vector in $\Gamma$ and then update it at every iteration, potentially increasing or decreasing the number of alpha vectors in $\Gamma$ as appropriate. This modification of the update step can improve efficiency.
Algorithm 21.7. Point-based value iteration, which finds an approximately optimal policy for an infinite horizon POMDP with discrete state, action, and observation spaces, where $\mathcal{B}$ is a vector of beliefs and $k_{\text{max}}$ is the number of iterations.

Figure 21.4. Approximate value functions obtained using point-based value iteration on the crying baby problem with belief vectors $[1/4, 3/4]$ and $[3/4, 1/4]$. Unlike QMDP and the fast informed bound, point-based value iteration’s value function is always a lower bound of the true value function.
Each update takes a set of alpha vectors \( \Gamma \) as input and outputs a set of alpha vectors \( \Gamma' \) that improves on the value function represented by \( \Gamma \) at the beliefs in \( B \). In other words, it outputs a \( \Gamma' \) such that \( U_{\Gamma'}(b) \geq U_{\Gamma}(b) \) for all \( b \in B \). We begin by initializing \( \Gamma' \) to the empty set and initializing \( B' \) to \( B \). We then remove a point \( b \) randomly from \( B' \) and perform a belief backup (algorithm 21.6) on \( b \) using \( \Gamma \) to get a new alpha vector \( \alpha \). We then find the alpha vector in \( \Gamma' \cup \{ \alpha \} \) that dominates at \( b \) and add it to \( \Gamma' \). As the algorithm progresses, \( B' \) becomes smaller and contains the set of points that have not been improved by \( \Gamma' \). The update finishes when \( B' \) is empty. Figure 21.5 illustrates this process on the crying baby problem.

```plaintext
struct RandomizedPointBasedValueIteration
    B   # set of belief points
    k_max # maximum number of iterations
end

function update(\( \mathcal{P} \)::POMDP, M::RandomizedPointBasedValueIteration, \( \Gamma \))
    \( \Gamma' \), B' = [], copy(M.B)
    while !isempty(B')
        b = rand(B')
        \( \alpha = \text{argmax}(\alpha \rightarrow \alpha \cdot b, \Gamma) \)
        \( \alpha' = \text{backup}(\mathcal{P}, \Gamma, b) \)
        if \( \alpha' \cdot b \geq \alpha \cdot b \)
            push!(\( \Gamma' \), \( \alpha' \))
        else
            push!(\( \Gamma' \), \( \alpha \))
        end
    end
    filter!(b -> maximum(\( \alpha \cdot b \) for \( \alpha \) in \( \Gamma' \)) <
    maximum(\( \alpha \cdot b \) for \( \alpha \) in \( \Gamma \)), B')
    end

function solve(M::RandomizedPointBasedValueIteration, \( \mathcal{P} \))
    \( \Gamma = [\text{baws_lowerbound}(\mathcal{P})] \)
    \( \Gamma = \text{alphavector_iteration}(\mathcal{P}, M, \Gamma) \)
    return LookaheadAlphaVectorPolicy(\( \mathcal{P} \), \( \Gamma \))
end
```

Algorithm 21.8. Randomized point-based backup, which updates the alpha vectors \( \Gamma \) associated with beliefs \( B \) for the \( \mathcal{P} \). This backup can be used in place of \( \text{point_based_update} \) in algorithm 21.7.
21.6 Sawtooth Upper Bound

The sawtooth upper bound is an alternative way to represent the value function. Instead of storing a set of alpha vectors $\Gamma$, we store a set of belief-utility pairs:

$$V = \{(b_1, U(b_1)), \ldots, (b_m, U(b_m))\}$$  \hspace{1cm} (21.11)

with the requirement that $V$ contains all of the standard basis beliefs:

$$E = \{e_1 = [1, 0, \ldots, 0], \ldots, e_n = [0, 0, \ldots, 1]\}$$  \hspace{1cm} (21.12)

such that

$$\{(e_1, U(e_1)), \ldots, (e_n, U(e_n))\} \subseteq V$$  \hspace{1cm} (21.13)

If these utilities are upper bounds (for example, as obtained from the fast informed bound), then the way we use $V$ to estimate $U(b)$ at arbitrary beliefs $b$ will result in an upper bound.\(^8\)

The “sawtooth” name comes from the way we estimate $U(b)$ by interpolating points in $V$. For each belief-utility pair $(b, U(b))$ in $V$, we form a single pointed “tooth.” If the belief space is $n$-dimensional, each tooth is an inverted $n$-dimensional pyramid. When multiple pairs are considered, it forms a “sawtooth” shape. The bases of the pyramids are formed by the standard basis beliefs $(e_i, U(e_i))$. The apex point of each tooth corresponds to each belief-utility pair $(b, U(b)) \in V$. Since these are pyramids in general, each tooth has walls equivalently defined by $n$-hyperplanes with bounded regions. These hyperplanes can also be interpreted as alpha vectors that act over a bounded region of the belief...
space, not the entire belief space as in normal alpha vectors. The combination of multiple pyramids forms the \( n \)-dimensional sawtooth. The sawtooth upper bound at any belief is similarly the minimum value among these pyramids at that belief.

Consider the sawtooth representation in a two-state POMDP, such as in the crying baby problem shown in figure 21.6. The corners of each tooth are the values \( U(e_1) \) and \( U(e_2) \) for each standard basis belief \( e_i \). The sharp lower point of each tooth is the value \( U(b) \), since each tooth is a point-set pair \( (b, U(b)) \). The linear interpolation from \( U(e_1) \) to \( U(b) \) and again from \( U(b) \) to \( U(e_2) \) forms the tooth. To combine multiple teeth and form the upper bound, we take the minimum interpolated value at any belief, forming the distinctive sawtooth shape.

To compute the sawtooth at any belief \( b \), we iterate over each belief-utility pair \((b', U(b'))\) in \( V \). The key idea is to compute the utility \( U'(b) \) for this hyperpyramid, first by finding the farthest basis point, then using this to determine the matching hyperplane from the hyperpyramid, and finally computing a utility using a rescaled version of the hyperplane. The farthest basis belief \( e_i \) is computed using

\[
i \leftarrow \arg \max_j \|b - e_j\|_1 - \|b' - e_j\|_1
\]  

(21.14)

This \( e_i \) uniquely identifies the particular hyperplane among those forming the hyperpyramid for \( U(b') \). Specifically, this hyperplane is defined by all corners \( e_j \neq e_i \) and using \( b' \) as a replacement for \( e_i \). At this point, we know this is the hyperplane for the region of the utility that \( b \) is contained within. The hyperplane’s utilities are \( U(e_j) \) for \( e_j \neq e_i \) and \( U(b') \) as a replacement for \( U(e_i) \). However, we cannot directly compute the desired utility \( U'(b) \) using a dot product because this is not the standard simplex. We instead compute the weight \( w \) of \( b \) in terms of the weighted distance from the hyperplane’s corners \( e_j \neq e_i \) and \( b' \). This allows us to be able to compute \( U'(b) \), essentially creating a simplex amenable to a dot product with \( U(e_j) \) and \( U(b') \):

\[
U'(b) = w_i U(b') + \sum_{j \neq i} w_j U(e_j)
\]  

(21.15)

This entire process is done among all \((b', U(b'))\), resulting in

\[
U(b) = \min_{(b', U(b')) \in V} U'(b)
\]  

(21.16)
Algorithm 21.9 provides an implementation. We can also derive a policy using greedy one-step lookahead.

Algorithm 21.9. The sawtooth upper bound representation for value functions and policies. It is defined using a dictionary $V$ that maps belief vectors to upper bounds on their utility obtained, for example, from the fast informed bound. A requirement of this representation is that $V$ contain belief-utility pairs at the standard basis beliefs, which can be obtained from the basis function. We can use one-step lookahead to obtain greedy action-utility pairs from arbitrary beliefs $b$.

We can iteratively apply greedy one-step lookahead at a set of beliefs $B$ to tighten our estimates of the upper bound. The beliefs in $B$ can be a superset of the beliefs in $V$. Algorithm 21.10 provides an implementation. Example 21.1 shows the effect of multiple iterations of the sawtooth approximation on the crying baby problem.
Suppose we want to maintain an upper bound of the value for the crying baby problem with regularly spaced beliefs points with a step size of 0.2. To obtain an initial upper bound, we use the fast informed bound. We can then run sawtooth iteration for three steps as follows:

\[
\begin{align*}
& n = \text{length}(\mathcal{P}.S) \\
& \pi_{\text{fib}} = \text{solve}(\text{FastInformedBound}(1), \mathcal{P}) \\
& V = \text{Dict}(e \Rightarrow \text{utility}(\pi_{\text{fib}}, e) \text{ for } e \text{ in } \text{basis}(\mathcal{P})) \\
& B = [[p, 1 - p] \text{ for } p \text{ in } 0.0:0.2:1.0] \\
& \pi = \text{solve}(\text{SawtoothIteration}(V, B, 2), \mathcal{P})
\end{align*}
\]

The sawtooth upper bound improves as follows:
21.7 Point Selection

Algorithms like point-based value iteration and sawtooth iteration require a set of beliefs \( B \). We want to choose \( B \) so that there are more points in the relevant areas of the belief space; we do not want to waste computation on beliefs that we are not likely to reach under our (hopefully approximately optimal) policy. One way to explore the potentially reachable space is to take steps (algorithm 21.11) in the belief space. The outcome of the step will be random because the observation is generated according to our probability model.

```
function randstep(\( \mathcal{P} : \text{POMDP} \), \( b \), \( a \))
    s = rand(SetCategorical(\( \mathcal{P}.S \), \( b \)))
    s', r, o = \( \mathcal{P}.TRO(s, a) \)
    b' = update(b, \( \mathcal{P} \), a, o)
    return b', r
end
```

Algorithm 21.11. A function for randomly sampling the next belief \( b' \) and reward \( r \) given the current belief \( b \) and action \( a \) in problem \( \mathcal{P} \).

We can create \( B \) from the belief states reachable from some initial belief under a random policy. This random belief expansion procedure (algorithm 21.12) may explore much more of the belief space than might be necessary; the belief space reachable by a random policy can be much larger than the space reachable by an optimal policy. Of course, computing the belief space reachable by an optimal policy generally requires knowing the optimal policy, which is what we want to compute in the first place. One approach that can be taken is to use successive approximations of the optimal policy to iteratively generate \( B \).

In addition to wanting our belief points to be focused on the reachable belief space, we also want those points to be spread out to allow for better value function approximation. The quality of the approximation provided by the alpha vectors associated with the points in \( B \) degrades as we evaluate points further from \( B \). We can take an exploratory belief expansion approach (algorithm 21.13), where we try every action for every belief in \( B \) and add the resulting belief states that are furthest away from the beliefs already in the set. Distance in belief space can be measured in different ways. This algorithm uses the \( L_1 \)-norm. Figure 21.7 shows an example of the belief points added to \( B \) using this approach.

\[ \sum_s |b(s) - b'(s)| \]


The \( L_1 \) distance between \( b \) and \( b' \) is \( \sum_s |b(s) - b'(s)| \) and is denoted \( \|b - b'\|_1 \). See appendix A.4.
Algorithm 21.12. An algorithm for randomly expanding a finite set of beliefs $B$ used in point-based value iteration based on reachable beliefs.

```plaintext
function random_belief_expansion($P$, $B$)
    $B'$ = copy($B$)
    for $b$ in $B$
        $a$ = rand($P$.A)
        $b'$, $r$ = randstep($P$, $b$, $a$)
        push!(B', $b'$)
    end
    return unique!(B')
end
```

Algorithm 21.13. An algorithm for expanding a finite set of beliefs $B$ used in point-based value iteration by exploring the reachable beliefs and adding those that are furthest from the current beliefs.

```plaintext
function exploratory_belief_expansion($P$, $B$)
    $B'$ = copy($B$)
    for $b$ in $B$
        best = ($b$=copy($b$), d=0.0)
        for $a$ in $P$.A
            $b'$, $r$ = randstep($P$, $b$, $a$)
            $d$ = minimum(norm($b$ - $b'$, 1) for $b$ in $B'$)
            if $d$ > best.d
                best = ($b$=$b'$, d=$d$)
            end
        end
        push!(B', best.b)
    end
    return unique!(B')
end
```
Chapter 9 introduced the concept of heuristic search as an online method in the fully observable context. This section discusses sawtooth heuristic search (algorithm 21.14) as an offline method that produces a set of alpha vectors that can be used to represent an offline policy. However, like the online POMDP methods discussed in the next chapter, the computational effort is focused on beliefs that are reachable from some specified initial belief. The heuristic that drives the exploration of reachable belief space is the gap between the upper and lower bounds of the value function.\footnote{The heuristic search value iteration (HSV) algorithm introduced the concept of using the sawtooth-based action heuristic and gap-based observation heuristic. T. Smith and R.G. Simmons, “Heuristic Search Value Iteration for POMDPs,” in Conference on Uncertainty in Artificial Intelligence (UAI), 2004. The SARSOP algorithm built upon this work. H. Kurniawati, D. Hsu, and W.S. Lee, “SARSOP: Efficient Point-Based POMDP Planning by Approximating Optimally Reachable Belief Spaces,” in Robotics: Science and Systems, 2008.}

The algorithm is initialized with an upper bound on the value function represented by a set of sawtooth belief-utility pairs \( V \), together with a lower bound on the value function represented by a set of alpha vectors \( \Gamma \). The belief-utility pairs defining the sawtooth upper bound can be obtained from the fast informed bound. The lower bound can be obtained from the best-action worst-state bound as shown in algorithm 21.14, or some other method such as point-based value iteration.
At each iteration, we explore beliefs that are reachable from our initial belief to some maximum depth. As we do these explorations, we update the set of belief-action pairs forming our sawtooth upper bound and the set of alpha vectors forming our lower bound. We stop exploring after a certain number of iterations or until the gap at our initial state is below a threshold $\delta > 0$.

When we encounter a belief $b$ along our path from the initial node during our exploration, we check whether the gap at $b$ is below a threshold $\delta / \gamma^d$, where $d$ is our current depth. If we are below that threshold, then we can stop exploring along that branch. As the depth $d$ increases, so does the threshold—in other words, the deeper we are the more tolerant we are of the gap. The reason for this is that the gap at $b$ after an update is at most $\gamma$ times the weighted average of the gap at the beliefs that are immediately reachable.

If the gap at $b$ is above the threshold and we have not reached our maximum depth, then we can explore the next belief $b'$, which is selected as follows. First, we determine the action $a$ recommended by our sawtooth policy. Then, we choose the observation $o$ that maximizes the gap at the resulting belief. We recursively explore down the tree. After exploring the descendants of $b'$, we add $(b', u)$ to $V$, where $u$ is the one-step lookahead value of $b'$. We also add to $\Gamma$ the alpha vector that results from a backup at $b'$. Figure 21.8 shows how the upper and lower bounds become tighter with each iteration.

### 21.9 Triangulated Value Functions

As discussed in section 20.1, a POMDP can be converted to a belief-state MDP. The state space in that belief-state MDP is continuous, corresponding to the space of possible beliefs in the original POMDP. We can approximate the value function in a way similar to what was discussed in chapter 8 and then apply a dynamic programming algorithm such as value iteration to the approximation. This section discusses a particular kind of local value function approximation that involves Freudenthal triangulation over a discrete set of belief points $B$. This triangulation allows us to interpolate the value function at arbitrary points in the belief space. As with the sawtooth representation, we use a set of belief-utility pairs $V = \{(b, U(b)) | b \in B\}$ to represent our value function. This approach can be used to obtain an upper bound on the value function.

The Freudenthal triangulation produces a set of vertices defined by integer components, which can later be converted to real-valued belief vectors for use

---

12 Some variants simply sample the next observations. Others select the observation that maximizes the gap weighted by its likelihood.

Algorithm 21.14. The sawtooth heuristic search policy. The solver starts from belief $b$ and explores to a depth $d$ for no more than $k_{\text{max}}$ iterations. It uses an upper bound obtained through the fast informed bound computed with $k_{\text{fib}}$ iterations. The lower bound is obtained from the best-action worst-state bound. The gap threshold is $\delta$. 

```python
struct SawtoothHeuristicSearch
    b # initial belief
    δ # gap threshold
    d # depth
    k_{\text{max}} # maximum number of iterations
    k_{\text{fib}} # number of iterations for fast informed bound
end

function explore!(M::SawtoothHeuristicSearch, 𝒫, \pihi, \pilo, b, d=0)
    S, λ, 0, γ = 𝒫.S, 𝒫.λ, 𝒫.0, 𝒫.γ
    ε(b') = utility(\pihi, b') - utility(\pilo, b')
    if d ≥ M.d || ε(b) ≤ M.δ / γ^d
        return
    end
    a = \pihi(b)
    o = argmax(\ o \rightarrow ε(update(b, 𝒫, a, o)), 0)
    b' = update(b, 𝒫, a, o)
    explore!(M, 𝒫, \pihi, \pilo, b', d+1)
    if b' ∉ basis(𝒫)
        \pihi.V[b'] = greedy(\pihi, b').u
    end
    push!(\pilo.Γ, backup(𝒫, \pilo.Γ, b'))
end

function solve(M::SawtoothHeuristicSearch, 𝒫::POMDP)
    π_{\text{fib}} = solve(FastInformedBound(M.k_{\text{fib}}, 𝒫)
    Vhi = Dict(e => utility(π_{\text{fib}}, e) for e in basis(𝒫))
    \pihi = SawtoothPolicy(𝒫, Vhi)
    πlo = LookaheadAlphaVectorPolicy(𝒫, [baws_lowerbound(𝒫)])
    for i in 1:M.k_{\text{max}}
        explore!(M, 𝒫, \pihi, \pilo, M.b)
        if utility(\pihi, M.b) - utility(πlo, M.b) < M.δ
            break
        end
    end
    return πlo
end
```
in our value function approximation. The number of vertices depends upon the dimensionality $n$ and granularity $m$. Given a granularity $m > 0$, the vertices in the Freudenthal triangulation are the $n$-dimensional vectors $v$ with integer components that satisfy:

$$m = v_1 \geq v_2 \geq v_3 \geq \cdots \geq v_n \geq 0$$  \hspace{1cm} (21.17)

Algorithm 21.15 generates this set of vertices. Figure 21.9 shows how the number of vertices varies with $n$ and $m$. 

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2021-03-07 23:12:00-08:00, revision 80bf4d6, comments to bugs@algorithmsbook.com
To be used in belief-state planning, these integer vertices must be transformed to form a triangulation of the belief over \( n \) discrete states. This transformation changes vertex \( \mathbf{v} \) into:

\[
\mathbf{v}' = \frac{1}{m} [v_1 - v_2, v_2 - v_3, v_3 - v_4, \ldots, v_{n-1} - v_n, v_n]
\]

where \( \mathbf{v}' \) represents a valid probability distribution that sums to 1. The probability assigned to the \( i \)th state is given by \( v_i' \). Example 21.2 shows how to generate the vertices and transform them into points in the belief-space.

This transformation is equivalent to the matrix multiplication \( \mathbf{v}' = \mathbf{Bv} \) with

\[
\mathbf{B} = \frac{1}{m} \begin{bmatrix}
1 & -1 & 0 & 0 & \cdots & 0 \\
0 & 1 & -1 & 0 & \cdots & 0 \\
0 & 0 & 1 & -1 & \cdots & 0 \\
\vdots & \vdots & \vdots & \vdots & \ddots & \vdots \\
0 & 0 & \cdots & 1 & -1 \\
0 & 0 & \cdots & 0 & 1
\end{bmatrix}
\]

This matrix \( \mathbf{B} \) is invertible, so the opposite transformation can be done. Hence, any belief \( \mathbf{b} \) can thus be mapped into the Freudenthal space to obtain its counter-
part \( \mathbf{x} = \mathbf{B}^{-1} \mathbf{b} \). It turns out that the \( i \)th component of \( \mathbf{B}^{-1} \mathbf{b} \) is simply \( m \sum_{k=i}^{n} b_k \). Algorithm 21.16 converts between these two forms.

Algorithm 21.16. The method to_belief takes a vector \( \mathbf{x} \) and a granularity integer \( m \) and returns \( \mathbf{Bx} \), a vector in belief space. The method to_freudenthal takes a belief vector \( \mathbf{b} \) and a granularity integer \( m \) and returns \( \mathbf{B}^{-1} \mathbf{b} \), a vector in the space of the Freudenthal triangulation.

If we know the value function \( U \) at a set of discrete belief points \( B \), we can use the triangulation to estimate the value function at arbitrary belief points through interpolation. We will first discuss how to interpolate a general function \( f \) in Freudenthal space at a point \( \mathbf{x} \). To interpolate in belief space, we use the transformation we just discussed to map between the Freudenthal space to belief space.

If we know the values of a function \( f \) at the integer Freudenthal vertices, we can use triangulation to estimate the value at an arbitrary point \( \mathbf{x} \) from the \( n + 1 \) vertices of the simplex enclosing \( \mathbf{x} \):

\[
f(\mathbf{x}) = \sum_{i=1}^{n+1} \lambda_i f(\mathbf{v}(i)) \tag{21.20}
\]

The vertices of the simplex containing \( \mathbf{x} \) are given by \( \mathbf{v}(1), \ldots, \mathbf{v}(n+1) \). The scalars \( \lambda_{1:n+1} \) are the barycentric coordinates of \( \mathbf{x} \) with respect to the simplex vertices. They are non-negative weights that sum to 1, such that \( \mathbf{x} = \sum_{i=1}^{n+1} \lambda_i \mathbf{v}(i) \).

The first vertex consists of the componentwise floor of \( \mathbf{x} \):\(^{14}\)

\[
\mathbf{v}(1) = [\lfloor x_1 \rfloor, \lfloor x_2 \rfloor, \ldots, \lfloor x_n \rfloor] \tag{21.21}
\]

We then compute the difference \( \mathbf{d} = \mathbf{x} - \mathbf{v}(1) \), and sort the components in descending order:

\[
d_{p_1} \geq d_{p_2} \geq \ldots \geq d_{p_n} \tag{21.22}
\]

where \( \mathbf{p} \) is a permutation of \( 1 : n \).

The remaining \( n \) simplex vertices can be constructed according to:

\[
\mathbf{v}(k+1) = \mathbf{v}(k) + \mathbf{e}_{p_k} \tag{21.23}
\]

\(^{14}\) Recall that \( \lfloor a \rfloor \) returns the greatest integer less than or equal to \( a \).
The Freudenthal discretization of a belief space over 3 states with \( m = 3 \) has 10 vertices:

\[
\begin{align*}
\{ & [3, 0, 0] [3, 1, 0] [3, 2, 0] [3, 3, 0] \\
& [3, 1, 1] [3, 2, 1] [3, 3, 1] \\
& [3, 2, 2] [3, 3, 2] [3, 3, 3] \\
\}
\end{align*}
\]

The first component in every vertex is \( m \). The other two dimensions, along with the projected triangulation, are shown in the left image below.

In the right image, we have the same triangulation mapped to the 3-state belief space. These vertices are:

\[
\begin{align*}
\{ & [1, 0, 0] [2/3, 1/3, 0] [1/3, 2/3, 0] [0, 1, 0] \\
& [2/3, 0, 1/3] [1/3, 1/3, 1/3] [0, 2/3, 1/3] \\
& [1/3, 0, 2/3] [0, 1/3, 2/3] [0, 0, 1] \\
\}
\end{align*}
\]
Suppose we want to find the simplex vertices for the Freudenthal triangulation of \( \mathbf{x} = [1.2, -3.4, 2] \). The first vertex is \( \mathbf{v}^{(1)} = [1, -4, 2] \), resulting in a difference \( \mathbf{d} = [0.2, 0.6, 0.0] \). We arrange the components of \( \mathbf{d} \) in descending order, \( 0.6 \geq 0.2 \geq 0.0 \) according to the permutation \( \mathbf{p} = [2, 1, 3] \). The remaining simplex vertices are:

\[
\begin{align*}
\mathbf{v}^{(2)} &= [1, -4, 2] + [0, 1, 0] = [1, -3, 2] \\
\mathbf{v}^{(3)} &= [1, -3, 2] + [1, 0, 0] = [2, -3, 2] \\
\mathbf{v}^{(4)} &= [2, -3, 2] + [0, 0, 1] = [2, -3, 3]
\end{align*}
\]
where \( \mathbf{e}_i \) represents the \( i \)th standard basis vector. Algorithm 21.17 computes the Freudenthal simplex containing a vector, and example 21.3 illustrates how it works for an arbitrary vector.

The barycentric coordinates \( \lambda \) are obtained using:

\[
\begin{align*}
\lambda_{n+1} &= d_{p_n} \\
\lambda_n &= d_{p_{n-1}} - d_{p_n} \\
\lambda_{n-1} &= d_{p_{n-2}} - d_{p_{n-1}} \\
&\vdots \\
\lambda_2 &= d_{p_1} - d_{p_2} \\
\lambda_1 &= 1 - \sum_{i=1}^{n+1} \lambda_i
\end{align*}
\]  

(21.24)

Algorithm 21.18 computes barycentric coordinates. Example 21.4 illustrates how to perform this computation.

```python
function barycentric_coordinates(x, vertices)
    d = x - vertices[1]
    p = sortperm(d, rev=true)
    n = length(x)
    λ = zeros(n+1)
    λ[n+1] = d[p[n]]
    for i in n:-1:2
        λ[i] = d[p[i-1]] - d[p[i]]
    end
    λ[1] = 1 - sum(λ[2:end])
    return λ
end
```

Algorithm 21.18. A method for computing the barycentric coordinates for a vector \( x \) in a Freudenthal simplex given by vertices \( V \). These vertices can be computed using algorithm 21.17.

So far we have discussed how to interpolate in Freudenthal space with integer simplex vertices. To interpolate in belief space, we simply use the transformation to the belief space. Suppose we know the values \( U(b) \) at all of the vertices \( \mathbf{v}' \) in the belief triangulation. Given any new belief \( b \), we can compute \( x = B^{-1}b \), find the vertices \( \mathbf{v}^{(1)}, \ldots, \mathbf{v}^{(n+1)} \) of the corresponding simplex in the original Freudenthal triangulation, and then compute the barycentric coordinates \( \lambda \) for \( x \). The interpolated value is then:

\[
U(b) = \sum_{i=1}^{n+1} \lambda_i U(\mathbf{v}^{(i)}) = \sum_{i=1}^{n+1} \lambda_i U(B\mathbf{v}^{(i)})
\]  

(21.25)

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2021-03-07 23:12:00-08:00, revision 80bf4d6, comments to bugs@algorithmsbook.com
Suppose we want to find the barycentric coordinates for the Freudenthal triangulation of \( x = [1.2, -3.4, 2] \). Recall that the vertices for \( x \) were obtained in example 21.3. We compute:

\[
\begin{align*}
\lambda_4 &= d_3 = 0.0 \\
\lambda_3 &= d_1 - d_3 = 0.2 \\
\lambda_2 &= d_2 - d_1 = 0.4 \\
\lambda_1 &= 1 - 0.0 - 0.2 - 0.4 = 0.4
\end{align*}
\]

This interpolation is implemented in algorithm 21.19.

Algorithm 21.20 applies a variation of approximate value iteration (introduced in algorithm 8.1) to our triangulated policy representation. We simply iteratively apply backups over our beliefs in \( B \) using one-step lookahead with our value function interpolation. If \( U \) is initialized with an upper bound, value iteration will result in an upper bound even after a finite number of iterations.\(^{16}\) Figure 21.10 shows an example policy and utility function.

### 21.10 Summary

- The QMDP algorithm assumes perfect observability after the first step, resulting in an upper-bound on the true value function.

- The fast informed bound provides a tighter upper bound on the value function than QMDP by accounting for the observation model.

- Point-based value iteration provides a lower bound on the value function using alpha vectors at a finite set of beliefs.

- Randomized point-based value iteration performs updates at randomly selected points in the belief set until the values at all points in the set are improved.

- The sawtooth upper bound allows for iterative improvement of the fast informed bound using an efficient point-set representation.

- Carefully selecting which belief points to use in point-based value iteration can improve the quality of the resulting policies.

---

\(^{16}\) This property holds because value functions are convex and the linear interpolation between vertices on the value function must lie on or above the underlying convex function. See lemma 4 of W.S. Lovejoy, “Computationally Feasible Bounds for Partially Observed Markov Decision Processes,” *Operations Research*, vol. 39, no. 1, pp. 162–175, 1991.
Algorithm 21.19. A policy representation using Freudenthal triangulation with granularity $m$. As with the sawtooth method, we maintain a dictionary $V$ that maps belief vectors in $B$ to utilities. At construction, we initialize the utilities to 0 in this implementation, but if we want to use this policy representation for an upper bound, then we would need to initialize those utilities appropriately. We define a utility function to perform the necessary interpolation to compute the utility for an arbitrary belief $b$. We can extract a policy from this representation using greedy lookahead, as with the other policy representations introduced in this chapter.

Algorithm 21.20. Approximate value iteration with $k_{\text{max}}$ iterations using a Triangulated Policy with granularity $m$. At each iteration, we update the utilities associated with the beliefs in $B$ using greedy one-step lookahead with triangulated utilities.

```plaintext
struct TriangulatedPolicy
    𝒫 # POMDP problem
    V # dictionary mapping beliefs to utilities
    B # beliefs
    m # granularity
end

function TriangulatedPolicy(𝒫::POMDP, m)
    vertices = freudenthal_vertices(length(𝒫.$𝒮$), m)
    B = to_belief.(vertices)
    V = Dict(b => 0.0 for b in B)
    return TriangulatedPolicy(𝒫, V, B, m)
end

function utility(π::TriangulatedPolicy, b)
    x = to_freudenthal(b, π.m)
    vertices = freudenthal_simplex(x)
    B = to_belief.(vertices)
    λ = barycentric_coordinates(x, vertices)
    return sum(λi < √eps() ? 0.0 : λi*π.V[b] for (λi, b) in zip(λ, B))
end

(π::TriangulatedPolicy)(b) = greedy(π, b).

struct TriangulatedIteration
    m      # granularity
    k_max # maximum number of iterations
end

function solve(M::TriangulatedIteration, 𝒫)
    π = TriangulatedPolicy(𝒫, M.m)
    U(b) = utility(π, b)
    for k in 1:M.k_max
        U′ = [greedy(𝒫, U, b).u for b in π.B]
        for (b, u′) in zip(π.B, U′)
            π.V[b] = u′
        end
    end
    return π
end
```
• Sawtooth heuristic search attempts to tighten the upper and lower bounds of the value function represented by sawtooth pairs and alpha vectors, respectively.

• One approach to approximately solving POMDPs is to discretize the belief space and then to apply dynamic programming to extract an upper bound on the value function and a policy.

21.11 Exercises

Exercise 21.1. Suppose we are in a variation of the straight-line hex world problem (appendix F.1) consisting of four cells, corresponding to states $s_{1:4}$. There are two actions: move left ($\ell$) and move right ($r$). The effects of those actions are deterministic. Moving left in $s_1$ or moving right in $s_4$ gives a reward of 100 and ends the game. With a discount factor of 0.9, compute alpha vectors using QMDP. Then, using the alpha vectors, compute the approximately optimal action given the belief $b = [0.3, 0.1, 0.5, 0.1]$.

Solution: We denote the alpha vector associated with moving left as $\alpha_\ell$ and the alpha vector associated with moving right as $\alpha_r$. We initialize the alpha vectors to zero:

$$
\alpha_\ell^{(1)} = [R(s_1, \ell), R(s_2, \ell), R(s_3, \ell), R(s_4, \ell)] = [0, 0, 0, 0] \\
\alpha_r^{(1)} = [R(s_1, r), R(s_2, r), R(s_3, r), R(s_4, r)] = [0, 0, 0, 0]
$$
In the first iteration, since all of the entries in the alpha vectors are zero, only the reward term contributes to the QMDP update (equation (21.1)):

\[
\alpha^{(2)}_L = [100, 0, 0, 0] \\
\alpha^{(2)}_R = [0, 0, 0, 100]
\]

In the next iteration, we apply the update, which leads to new values for \( s_2 \) for the left alpha vector and for \( s_3 \) for the right alpha vector. The updates for the left alpha vector are as follows (with the right alpha vector updates being symmetric):

\[
\alpha^{(3)}_L (s_1) = 100 \quad \text{(terminal state)} \\
\alpha^{(3)}_L (s_2) = 0 + 0.9 \times \max(\alpha^{(2)}_L (s_1), \alpha^{(2)}_R (s_1)) = 90 \\
\alpha^{(3)}_L (s_3) = 0 + 0.9 \times \max(\alpha^{(2)}_L (s_2), \alpha^{(2)}_R (s_2)) = 0 \\
\alpha^{(3)}_L (s_4) = 0 + 0.9 \times \max(\alpha^{(2)}_L (s_3), \alpha^{(2)}_R (s_3)) = 0
\]

This leads to:

\[
\alpha^{(3)}_L = [100, 90, 0, 0] \\
\alpha^{(3)}_R = [0, 0, 90, 100]
\]

In the third iteration, the updates for the left alpha vector are:

\[
\alpha^{(4)}_L (s_1) = 100 \quad \text{(terminal state)} \\
\alpha^{(4)}_L (s_2) = 0 + 0.9 \times \max(\alpha^{(3)}_L (s_1), \alpha^{(3)}_R (s_1)) = 90 \\
\alpha^{(4)}_L (s_3) = 0 + 0.9 \times \max(\alpha^{(3)}_L (s_2), \alpha^{(3)}_R (s_2)) = 81 \\
\alpha^{(4)}_L (s_4) = 0 + 0.9 \times \max(\alpha^{(3)}_L (s_3), \alpha^{(3)}_R (s_3)) = 81
\]

Our alpha vectors are then:

\[
\alpha^{(4)}_L = [100, 90, 81, 81] \\
\alpha^{(4)}_R = [81, 81, 90, 100]
\]

At this point, our alpha vector estimates have converged. We now determine the optimal action by maximizing the utility associated with our belief over all actions.

\[
\alpha^\top_L b = 100 \times 0.3 + 90 \times 0.1 + 81 \times 0.5 + 81 \times 0.1 = 87.6 \\
\alpha^\top_R b = 81 \times 0.3 + 81 \times 0.1 + 90 \times 0.5 + 100 \times 0.1 = 87.4
\]

Thus, we find that moving left is the optimal action for this belief state, despite higher probability of being on the right half of the grid world. This is due to the relatively high likelihood we assign to being in state \( s_1 \), where we would receive a large immediate reward by moving left.
Exercise 21.2. Recall the simplified hex world problem from exercise 21.1. Compute alpha vectors for each action using the blind lower bound. Then, using the alpha vectors, compute the value at the belief $b = [0.3, 0.1, 0.5, 0.1]$.

Solution: The blind lower bound, equation (21.6), is like the QMDP update, but lacks the maximization. We initialize the components of the alpha vectors to zero and run to convergence:

$$\alpha_{\ell}^{(2)} = [100, 0, 0, 0]$$

$$\alpha_{r}^{(2)} = [0, 0, 0, 100]$$

$$\alpha_{\ell}^{(3)} = [100, 90, 0, 0]$$

$$\alpha_{r}^{(3)} = [0, 0, 90, 100]$$

$$\alpha_{\ell}^{(4)} = [100, 90, 81, 0]$$

$$\alpha_{r}^{(4)} = [0, 81, 90, 100]$$

$$\alpha_{\ell}^{(5)} = [100, 90, 81, 72.9]$$

$$\alpha_{r}^{(5)} = [72.9, 81, 90, 100]$$

At this point, our alpha vector estimates have converged. We now determine the value by maximizing the utility associated with our belief over all actions:

$$\alpha_{\ell}^T b = 100 \times 0.3 + 90 \times 0.1 + 81 \times 0.5 + 72.9 \times 0.1 = 86.79$$

$$\alpha_{r}^T b = 72.9 \times 0.3 + 81 \times 0.1 + 90 \times 0.5 + 100 \times 0.1 = 84.97$$

Thus, the lower bound at $b$ is 86.79.

Exercise 21.3. What is the complexity of a single iteration of point-based value iteration over all belief points in $B$ under the assumption that $|B| < |S|$?

Solution: Point-based value iterations performs an alpha vector backup for every belief point in $B$. Each backup operation performs a belief update for each action and observation. A belief update, as shown in equation (19.7), is $O(|S|^2)$ because it iterates over all initial and successor states. Together, these operations are $O(|B||A||O||S|^2)$. Each backup operation also computes a successor alpha vector for each action by iterating over all initial and successor states, and each observation. Together, these operations are also $O(|B||A||O||S|^2)$. As a result, the complexity of point-based value iteration is $O(|B||A||O||S|^2)$. 

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Exercise 21.4. Consider the set of belief-utility pairs given by:

\[ V = \{ ([1, 0], 0), ([0, 1], -10), ([0.8, 0.2], -4), ([0.4, 0.6], -6) \} \]

Using weights \( w_i = 0.5 \) for all \( i \), determine the utility for belief \( b = [0.5, 0.5] \) using the sawtooth upper bound.

Solution: We interpolate with the belief-utility pairs. For each non-basis belief, we start by finding the farthest basis belief \( e_i \). Starting with \( b_3 \), we compute:

\[
i_3 = \arg \max_j \| b - e_j \|_1 - \| b_3 - e_j \|_1
\]

\[
\| b - e_1 \|_1 - \| b_3 - e_1 \|_1 = \left\| \begin{bmatrix} 0.5 \\ 0.5 \end{bmatrix} - \begin{bmatrix} 1 \\ 0 \end{bmatrix} \right\|_1 - \left\| \begin{bmatrix} 0.8 \\ 0.2 \end{bmatrix} - \begin{bmatrix} 1 \\ 0 \end{bmatrix} \right\|_1
\]

\[
= \left\| \begin{bmatrix} -0.5 \\ 0.5 \end{bmatrix} \right\|_1 - \left\| \begin{bmatrix} -0.2 \\ 0.2 \end{bmatrix} \right\|_1
\]

\[
= 0.6
\]

\[
\| b - e_2 \|_1 - \| b_3 - e_2 \|_1 = \left\| \begin{bmatrix} 0.5 \\ 0.5 \end{bmatrix} - \begin{bmatrix} 0 \\ 1 \end{bmatrix} \right\|_1 - \left\| \begin{bmatrix} 0.8 \\ 0.2 \end{bmatrix} - \begin{bmatrix} 0 \\ 1 \end{bmatrix} \right\|_1
\]

\[
= \left\| \begin{bmatrix} 0.5 \\ -0.5 \end{bmatrix} \right\|_1 - \left\| \begin{bmatrix} 0.8 \\ -0.8 \end{bmatrix} \right\|_1
\]

\[
= -0.6
\]

\[
i_3 = 1
\]

Thus, \( e_1 \) is the farthest basis belief from \( b_3 \).
Thus, \( e_2 \) is the farthest basis belief from \( b_4 \).

We can compute \( U(b) \) using our weights along with the appropriate corners and utility pairs \((e_2, b_3)\) and \((e_1, b_4)\):

\[
U_3(b) = 0.5 \times -4 + 0.5 \times (-10) = -7 \\
U_4(b) = 0.5 \times -6 + 0.5 \times 0 = -3
\]

Finally, we compute \( U(b) \) by taking the minimum of \( U_3(b) \) and \( U_4(b) \). Thus, \( U(b) = -7 \).

**Exercise 21.5.** Suppose we have a valid lower bound represented as a set of alpha vectors \( \Gamma \). Is it possible for a backup at a belief state \( b \) to result in an alpha vector \( \alpha' \) such that \( \alpha'^T b \) is lower than the utility function represented by \( \Gamma \)? In other words, can a backup at a belief \( b \) result in an alpha vector that assigns lower utility to \( b \) than the value function represented by \( \Gamma \)?

**Solution:**

It is possible. Suppose we have only one action, observations are perfect, there is no discounting, and the state space is \( \{s^0, s^1\} \). The reward is \( R(s^i) = i \) for all \( i \), and states transition deterministically to \( s^0 \). We start with a valid lower bound, \( \Gamma = \{[-1, +1]\} \), as shown in red in figure 21.11. We choose \( b = [0.5, 0.5] \) for the belief where we do the backup. Using equation (21.9), we obtain

\[
\alpha(s^0) = R(s^0) + U^\Gamma(s^0) = 0 + (-1) = -1 \\
\alpha(s^1) = R(s^1) + U^\Gamma(s^0) = 1 + (-1) = 0
\]
Hence, the alpha vector we get after a backup is $[-1, 0]$, shown in blue in figure 21.11. The utility at $b$ with that alpha vector is 0. However, $U^\Gamma(b) = 0.5$, showing that backing up a belief can result in an alpha vector that, on its own, represents a lower utility at that belief. This fact motivates the use of the if-statement in randomized point-based value iteration (algorithm 21.8). That if statement will use either the alpha vector from the backup or the dominating alpha vector in $\Gamma$ at belief $b$, whichever gives the greatest utility estimate.

**Exercise 21.6.** Derive equation (21.24), used to obtain the barycentric coordinates $\lambda$ for a vector $x \in \mathbb{R}^n$ in a Freudenthal triangulation simplex with $n + 1$ vertices $v^{(i)}$.

**Solution:** The barycentric coordinates are a convex combination of the simplex vertices that produce $x$:

$$x = \sum_{i=1}^{n+1} \lambda_i v^{(i)}$$

The components of $\lambda$ are non-negative and must sum to 1.

Recalling the definition of the vertices for the Freudenthal triangulation, we expand:

$$x = \sum_{i=1}^{n+1} \lambda_i v^{(i)} = \sum_{i=1}^{n+1} \lambda_i \left( \sum_{j=1}^{i-1} e_{p_j} \right)$$

$$= \left( \sum_{i=1}^{n+1} \lambda_i \right) v^{(1)} + \left( \sum_{i=2}^{n+1} \lambda_i \right) e_{p_1} + \left( \sum_{i=3}^{n+1} \lambda_i \right) e_{p_2} + \ldots + \left( \sum_{i=n}^{n+1} \lambda_i \right) e_{p_{n-1}} + \lambda_{n+1} e_{p_n}$$

$$= v^{(1)} + \left( \sum_{i=2}^{n+1} \lambda_i \right) e_{p_1} + \left( \sum_{i=3}^{n+1} \lambda_i \right) e_{p_2} + \ldots + \left( \sum_{i=n}^{n+1} \lambda_i \right) e_{p_{n-1}} + \lambda_{n+1} e_{p_n}$$

where in the last step we used the fact that the components sum to 1.

Recall that $d = x - v^{(1)}$. We thus have:

$$d = \left( \sum_{i=2}^{n+1} \lambda_i \right) e_{p_1} + \left( \sum_{i=3}^{n+1} \lambda_i \right) e_{p_2} + \ldots + \left( \sum_{i=n}^{n+1} \lambda_i \right) e_{p_{n-1}} + \lambda_{n+1} e_{p_n}$$

This can be solved starting with $\lambda_{n+1}$ to obtain equation (21.24).

**Exercise 21.7.** Consider Freudenthal value interpolation for the belief $b = [4/6, 1/6, 1/6]$ with a granularity constant $m = 3$. Is there a unique permutation of the components of the direction $d$ associated with this belief? Geometrically, what does this say about $b$? Which permutation of $d$ should be used?

**Solution:** We begin by converting the belief vector to the space of the Freudenthal triangulation, resulting in:

$$x = B^{-1}b = [3, 1, 0.5]$$
The first vertex in the Freudenthal triangulation is formed by the floor of the components of \( x \), and is thus \( v^{(1)} = [3, 1, 0] \). We can calculate \( d \):

\[
\mathbf{d} = \mathbf{x} - v^{(1)} = [0, 0, 0.5]
\]

The components of \( d \) are non-unique, so both permutations \( p = [3, 1, 2] \) and \( p = [3, 2, 1] \) are valid. Geometrically, this means that \( b \) lies on an edge shared by two simplexes. Both permutations are equally valid for value interpolation, resulting in the same barycentric coordinates that zero out the problematic components. Here, the permutations both produce \( \lambda = [0.5, 0.5, 0, 0] \).
Online methods determine the optimal policy by planning from the current belief state. The belief space reachable from the current state is typically small compared with the full belief space. As introduced in the fully observable context, many online methods use variations of tree-based search up to some horizon. Various strategies can be used to try to avoid the exponential computational growth with the tree depth. Although online methods require more computation per decision step during execution than offline approaches, online methods are sometimes easier to apply to high-dimensional problems.

### 22.1 Lookahead with Rollouts

Section 9.2 introduced lookahead with rollouts as an online method in fully observed problems. The algorithm provided in that section can be used directly for partially observed problems. It uses a function for randomly sampling the next state, which corresponds to a belief state in the context of partial observability. This function was already introduced in algorithm 21.11. Because we can use a generative model rather than an explicit model for transitions, rewards, and observations, we can accommodate problems with high dimensional state and observation spaces.

### 22.2 Forward Search

We can apply the forward search strategy from algorithm 9.2 to partially observed problems without modification. The difference between MDPs and POMDPs is encapsulated by the one-step lookahead, which branches on actions and observations as shown in figure 22.1. The value of taking action $a$ from belief $b$ can be

---

defined recursively to some depth $d$:

$$Q_d(b,a) = \begin{cases} R(b,a) + \gamma \sum_o P(o \mid b,a)U_{d-1}(\text{Update}(b,a,o)) & \text{if } d > 0 \\ U(b) & \text{otherwise} \end{cases}$$

where $U_d(b) = \max_a Q_d(b,a)$. When $d = 0$, we have reached maximum depth and return the utility using the approximate value function $U(b)$. This approximate value function may be obtained from one of the methods discussed in the previous chapter, heuristically chosen, or estimated from one or more rollouts. When $d > 0$, we continue to search deeper, recursing down another level. Example 22.1 shows how to combine QMDP with forward search for the machine replacement problem. Example 22.2 demonstrates forward search on the crying baby problem.

Example 22.1. An example of applying forward search to a POMDP problem

Consider applying forward search to the machine replacement problem. We can first obtain an approximate value function through QMDP (algorithm 21.2). We can then construct a ForwardSearch object, which was originally defined in algorithm 9.2. The call to lookahead within that function will use the one defined for POMDPs in algorithm 20.5. The following code applies forward search to the problem $\mathcal{P}$ from belief state $[0.5, 0.2, 0.3]$ to depth 5 using our estimate of the utility obtained from QMDP at the leaf nodes.

```python
k_max = 10 # maximum number of iterations of QMDP
πQMDP = solve(QMDP(k_max), ρ)
d = 5 # depth
U(b) = utility(πQMDP, b)
π = ForwardSearch(ρ, d, U)
π([0.5,0.2,0.3])
```

The computation associated with the recursion in equation (22.1) grows exponentially with depth, $O(|A|^d|O|^d)$. Hence, forward search is generally limited to a relatively shallow depth. To go deeper, we can limit the action or observation branching. For example, if we have some domain knowledge, we may restrict the action set either at the root or further down the tree. For the observation branching, we may restrict our consideration to a small set of likely observations—or even just the most likely observation.² Branching can be avoided entirely by adopting the open loop or hindsight optimization methods described in section 9.9.3 with states sampled from the current belief.

Consider forward search in the crying baby problem with an approximate value function given by the alpha vectors $[-3.7, -15]$ and $[-2, -21]$. Running forward search to depth 2 from the initial belief $b = [0.5, 0.5]$ proceeds as follows:

$Q_2(b, a_{feed}) = R(b, a_{feed}) + \gamma (P(\text{crying} \mid b, \text{feed}) U_1([1.0, 0.0])$

$+ P(\text{quiet} \mid b, \text{feed}) U_1([1.0, 0.0]))$

$= -10 + 0.9(0.1 \times -3.2157 + 0.9 \times -3.2157)$

$= -12.894$

$Q_2(b, a_{ignore}) = R(b, a_{ignore}) + \gamma (P(\text{crying} \mid b, \text{ignore}) U_1([0.093, 0.907])$

$+ P(\text{quiet} \mid b, \text{ignore}) U_1([0.786, 0.214]))$

$= -5 + 0.9(0.485 \times -15.872 + 0.515 \times -7.779)$

$= -15.534$

$Q_2(b, a_{sing}) = R(b, a_{sing}) + \gamma (P(\text{crying} \mid b, \text{sing}) U_1([0.0, 1.0])$

$+ P(\text{quiet} \mid b, \text{sing}) U_1([0.891, 0.109]))$

$= -5.5 + 0.9(0.495 \times -16.8 + 0.505 \times -5.543)$

$= -15.503$

Recall that feeding the baby always results in a sated baby ($b = [1, 0]$), and singing to the baby ensures that it only cries if it is hungry ($b = [0, 1]$). Each $U_1$ value must be evaluated by recursively looking ahead and eventually evaluating the approximate value function. The policy predicts that feeding the baby will result in the highest expected utility, so it recommends that action.
22.3 Branch and Bound

The branch and bound technique originally introduced in the context of MDPs can be extended to POMDPs. The same algorithm in section 9.4 can be used without modification (see example 22.3), relying on the appropriate look-ahead implementation to account for the observations and updating beliefs. The efficiency of the algorithm still depends upon the quality of the upper and lower bounds for pruning.

Although we can use domain-specific heuristics for the upper and lower bounds as we did in the fully observed case, we can alternatively use one of the methods introduced in the previous chapter for discrete state spaces. For example, we can use the fast informed bound for the upper bound and point-based value iteration for the lower bound. So long as the lower bound $U$ and upper bound $Q$ are true lower and upper bounds, the result of the branch and bound algorithm will be the same as the forward search algorithm with $U$ as the approximate value function.

Example 22.3. An application of branch and bound to the crying baby problem.

```
In this example, we apply branch and bound to the crying baby problem with a depth of 5. The upper bound comes from the fast informed bound and the lower bound comes from point-based value iteration. We compute the action from belief $[0.4, 0.6]$.

$k_\text{max} = 10$  # maximum number of iterations for bounds
$\pi_{\text{FIB}} = \text{solve}([\text{FastInformedBound}(k_\text{max}), \mathcal{P})$  
$d = 5$  # depth
$U_{\text{lo}}(b) = \text{utility}(\pi_{\text{FIB}}, b)$  
$B = [[p, 1 - p] \text{ for } p \text{ in } 0.0:0.2:1.0]$  
$\pi_{\text{PBVI}} = \text{solve}([\text{PointBasedValueIteration}(B, k_\text{max}), \mathcal{P})$
$U_{\text{hi}}(b) = \text{utility}(\pi_{\text{PBVI}}, b)$  
$Q_{\text{hi}}(b, a) = \text{lookahead}([\mathcal{P}, U_{\text{hi}}, b, a)$  
$\pi = \text{BranchAndBound}(\mathcal{P}, d, U_{\text{lo}}, Q_{\text{hi}})$  
$\pi([0.4, 0.6])$
```

22.4 Sparse Sampling

Forward search sums over all possible observations, resulting in a runtime exponential in $|\mathcal{O}|$. As introduced in section 9.5, we can can use sampling to avoid exhaustive summation. We can generate $m$ observations for each action and then
compute

\[
Q_d(b, a) = \begin{cases} 
\frac{1}{m} \sum_{i=1}^{m} \left( r_d^{(i)} + \gamma U_{d-1} \left( \text{Update}(b, a, o_d^{(i)}) \right) \right) & \text{if } d > 0 \\
U(\text{Update}(b, a, o_d^{(i)})) & \text{otherwise}
\end{cases}
\]  \tag{22.2}

where \( r_d^{(i)} \) and \( o_d^{(i)} \) are the \( i \)th sampled observation and reward associated with action \( a \) from belief \( b \). We may use algorithm 9.4 without modification. The resulting complexity is \( O(|A|^d m^d) \).

### 22.5 Monte Carlo Tree Search

The Monte Carlo tree search approach for MDPs can be extended to POMDPs, though we cannot use the same exact implementation.\(^3\) The input to the algorithm is a belief state \( b \), depth \( d \), exploration factor \( c \), and a rollout policy \( \pi \).\(^4\) The main difference between the POMDP algorithm (algorithm 22.1) and the MDP algorithm is that the counts and values are associated with histories instead of states. A history is a sequence of past actions and observations. For example, if we have two actions \( a_1 \) and \( a_2 \) and two observations \( o_1 \) and \( o_2 \), then a possible history could be the sequence \( h = a_1 o_2 a_2 o_2 a_1 o_1 \). During the execution of the algorithm, we update the value estimates \( Q(h, a) \) and counts \( N(h, a) \) for a set of history-action pairs.\(^5\)

The histories associated with \( Q \) and \( N \) may be organized in a tree similar to the one in figure 22.2. The root node represents the empty history starting from the initial belief state \( b \). During the execution of the algorithm, the tree structure expands. The layers of the tree alternate between action nodes and observation nodes. Associated with each action node are values \( Q(h, a) \) and \( N(h, a) \), where the history is determined by the path from the root node. As with the MDP version, when searching down the tree, the algorithm takes the action that maximizes

\[
Q(h, a) + c \sqrt{\frac{\log N(h)}{N(h)}}
\]  \tag{22.3}

where \( N(h) = \sum_a N(h, a) \) is the total visit count for history \( h \) and \( c \) is an exploration parameter. Importantly, \( c \) augments the value of actions that are unexplored and underexplored, thus representing the relative tradeoff between exploration and exploitation.


\(^4\) Monte Carlo tree search can be implemented with a POMDP rollout policy that operates on beliefs or on an MDP rollout policy that operates on states. Random policies are commonly used.

\(^5\) There are many variations of the basic algorithm introduced here, including some that incorporate aspects of double progressive widening, discussed in section 9.6. Z.N. Sunberg and M.J. Kochenderfer, “Online Algorithms for POMDPs with Continuous State, Action, and Observation Spaces,” in *International Conference on Automated Planning and Scheduling (ICAPS)*, 2018.
Algorithm 22.1. Monte Carlo tree search for POMDPs from belief \( b \). The initial history \( h \) is optional. This implementation is similar to the one in algorithm 9.5.

```plaintext
struct HistoryMonteCarloTreeSearch
  \( \mathcal{P} \)  # problem
  \( N \)  # visit counts
  \( Q \)  # action value estimates
  \( d \)  # depth
  \( k_{\text{max}} \)  # number of simulations
  \( c \)  # exploration constant
  \( \pi \)  # rollout policy
end

function explore(\( \pi \)::HistoryMonteCarloTreeSearch, \( h \))
  \( \mathcal{A}, N, Q, c = \pi.\mathcal{P}.\mathcal{A}, \pi.N, \pi.Q, \pi.c \)
  \( Nh = \text{sum}(\text{get}(N, (h,a), 0) \text{ for } a \text{ in } \mathcal{A}) \)
  return argmax(a→\( Q[(h,a)] + c*\text{bonus}(N[(h,a)], Nh), \mathcal{A} \))
end

function simulate(\( \pi \)::HistoryMonteCarloTreeSearch, \( s, h, d \))
  if \( d \leq 0 \)
    return 0.0
  end
  \( \mathcal{P}, N, Q, c = \pi.\mathcal{P}, \pi.N, \pi.Q, \pi.c \)
  \( S, A, TRO, \gamma = \mathcal{P}.S, \mathcal{P}.A, \mathcal{P}.TRO, \mathcal{P}.\gamma \)
  if !\( \text{haskey}(N, (h, \text{first}(A))) \)
    for \( a \text{ in } A \)
      \( N[(h,a)] = 0 \)
      \( Q[(h,a)] = 0.0 \)
    end
    \( b = [s == s' \ ? 1.0 : 0.0 \text{ for } s' \text{ in } S] \)
  end
  \( a = \text{explore}(\pi, h) \)
  \( s', r, o = \text{TRO}(s,a) \)
  \( q = r + \gamma*\text{simulate}(\pi, s', \text{vcat}(h, (a,o)), d-1) \)
  \( N[(h,a)] += 1 \)
  \( Q[(h,a)] += (q-Q[(h,a)])/N[(h,a)] \)
  return q
end

function (\( \pi \)::HistoryMonteCarloTreeSearch)(\( b \), \( h=[] \))
  for i in 1:\( \pi.k_{\text{max}} \)
    \( s = \text{rand}(\text{SetCategorical}(\pi.\mathcal{P}.S, b)) \)
    simulate(\( \pi \), \( s, h, \pi.d \))
  end
  return argmax(a→\( \pi.Q[(h,a)], \pi.\mathcal{P}.A \))
end
```

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2021-03-07 23:12:00-08:00, revision 80bf4d6, comments to bugs@algorithmsbook.com
As with the MDP version, the Monte Carlo tree search algorithm is an anytime algorithm. The loop in algorithm 22.1 can be terminated at any time, and the best solution found up to that point will be returned. With a sufficient number of iterations, the algorithm converges to the optimal action.

Prior knowledge can be incorporated into Monte Carlo tree search in how we initialize $N$ and $Q$. Our implementation uses zero, but other choices are possible, including having the initialization of the action values be a function of history. The rollout policy can also incorporate prior knowledge, with better rollout policies tending to produce better value estimates.

The algorithm does not need to be reinitialized with each decision. The history tree and associated counts and value estimates can be maintained between calls. The observation node associated with the selected action and actual observation becomes the root node at the next time step.

### 22.6 Determinized Sparse Tree Search

Determinized sparse tree search strives to reduce the overall amount of sampling in both sparse sampling and Monte Carlo tree search by making the observation resulting from performing an action deterministic. It does so by building a determinized belief tree from a special particle belief representation to form a sparse tree.

Ye et al. present a determinized sparse tree search algorithm for POMDPs called Determinized Sparse Partially Observable Tree, or DESPOT. N. Ye, A. Somani, D. Hsu, and W.S. Lee, “DESPOT: Online POMDP Planning with Regularization,” Journal of Artificial Intelligence Research, vol. 58, pp. 231–266, 2017. In addition, the algorithm includes branch and bound, heuristic search, and regularization techniques.

![Figure 22.2. A search tree containing all histories covered when running Monte Carlo tree search with 100 samples on the machine replacement problem. Visitations given beneath each action node, and color indicates node values with high values in blue and low values in red. Expanded nodes with zero visitations are not shown. This search used an exploration constant $c = 0.5$, a maximum depth $d = 5$, and a uniform random rollout policy. The initial belief is certainty in a fully working system. Monte Carlo tree search is able to avoid certain actions and instead focus samples on more promising paths.](image)
approximation of the true belief tree. Each particle refers to one of \( m \) \textit{scenarios}, each of depth \( d \). A scenario represents a fixed history that the particle will follow for any given sequence of actions \( a^{(1)}, a^{(2)}, \ldots, a^{(d)} \). Every distinct action sequence produces a distinct history under a particular scenario.\(^7\) This determinization reduces the size of the search tree to \( O(|A|^d m) \). An example of a history is given in example 22.4. A determinized tree is shown in figure 22.3.

Suppose we have two states \( s_1 \) and \( s_2 \), two actions \( a_1 \) and \( a_2 \), and two observations \( o_1 \) and \( o_2 \). A possible history of depth \( d = 2 \) for the particle with initial state \( s_2 \) is the sequence \( h = s_2 a_1 o_2 s_1 a_2 o_1 \). If this history is used as a scenario, then this history is returned every time the belief tree is traversed from the initial state with the action sequence \( a^{(1)} = a_1 \) and \( a^{(2)} = a_2 \).

A search tree with \( m \) scenarios up to depth \( d \) can be fully specified by a compact \( m \times d \) \textit{determinizing matrix} \( \Phi \) containing probability masses. The element \( \Phi_{ij} \) contains the information needed for a particle following the \( i \)th scenario at depth \( j \) to identify its successor state and observation. Specifically, \( \Phi_{ij} \) is a uniformly distributed random number that can generate the successor pair \( (s', o) \) from a state-action pair \( (s, a) \), following the distribution \( P(s', o | s, a) = T(s' | s, a)O(o | a, s') \).

We can generate a determinizing matrix by filling it with values sampled uniformly between 0 and 1.

Beliefs are represented as vectors of belief particles. Each belief particle \( \phi \) contains a state \( s \) and indices \( i \) and \( j \) into the determinizing matrix \( \Phi \) corresponding to a scenario \( i \) and current depth \( j \). Given a particular action \( a \), \( \Phi_{ij} \) is used to deterministically transition to successor state \( s' \) and observation \( o \). The successor particle \( \phi' = (s', i, j + 1) \) receives \( s' \) as its state and increments \( j \) by 1. Example 22.5 demonstrates this tree traversal process. The particle belief representation is implemented in algorithm 22.2 and is used in forward search in algorithm 22.3.

### 22.7 Gap Heuristic Search

Similar to the offline heuristic search presented in section 21.8, we can use the gap between the upper and lower bounds to guide our search toward beliefs that have uncertainty in their associated value and as an indication of when we can stop exploring. The gap \( \epsilon(b) \) at a belief \( b \) is the difference between the upper bound
Suppose we generate a determinizing matrix $\Phi$ for a problem with 4 histories up to depth 3:

$$
\Phi = \begin{bmatrix}
0.393 & 0.056 & 0.369 \\
0.313 & 0.749 & 0.273 \\
0.078 & 0.262 & 0.009 \\
0.969 & 0.598 & 0.095
\end{bmatrix}
$$

Suppose we take action $a_3$ in state $s_2$ when at depth 2 while following history 3. The corresponding belief particle is $\phi = (2, 3, 2)$, and the determinizing value in $\Phi$ is $\Phi_{3,2} = 0.262$.

The deterministic successor action and observation are given by iterating over all successor state-observation pairs and accumulating their transition probabilities. We begin with $p = 0$ and evaluate $s' = s_1, o = o_1$. Suppose we get $T(s_1 \mid s_2, a_3)O(o_1 \mid a_3, s_1) = 0.1$. We increase $p$ to 0.1, which is less than $\Phi_{3,2}$, so we continue.

Next we evaluate $s' = s_1, o = o_2$. Suppose we get $T(s_1 \mid s_2, a_3)O(o_2 \mid a_3, s_2) = 0.17$. We increase $p$ to 0.27, which is greater than $\Phi_{3,2}$. We thus deterministically proceed to $s' = s_1, o = o_2$ as our successor state, resulting in a new particle $\phi' = (1, 3, 3)$. 

Example 22.5. Determinized sparse tree search uses a matrix to make tree traversal deterministic for a given particle.
struct DeterminizedParticle
    s # state
    i # scenario index
    j # depth index
end

function successor(ℙ, Φ, ϕ, a)
    ℡, ℡₀, ℙ, ℙ₀ = ℙ.s, ℙ.o, ℙ.t, ℙ₀
    p = 0.0
    for (s', o) in product(℡, ℡₀)
        p += ℙ₁(Φ.s, a, s') * ℙ₀(a, s', o)
        if p ≥ Φ[ϕ.i, ϕ.j]
            return (s', o)
        end
    end
    return last(℡), last(℡₀)
end

function possible_observations(ℙ, Φ, b, a)
    ℡₀ = []
    for ϕ in b
        s', o = successor(ℙ, Φ, ϕ, a)
        push!(℡₀, o)
    end
    return unique(℡₀)
end

function update(b, Φ, ℙ, a, o)
    b' = []
    for ϕ in b
        s', o' = successor(ℙ, Φ, ϕ, a)
        if o == o'
            push!(b', DeterminizedParticle(s', ϕ.i, ϕ.j + 1))
        end
    end
    return b'
end

Algorithm 22.2. The determinized particle belief update produces a determinized sparse tree search for POMDPs. Each belief \( b \) consists of particles \( \phi \) that each encode a particular scenario and depth along the scenario. Their scenario's trajectory is determined through a matrix \( \Phi \) containing random values in \([0, 1]\). Each particle \( \phi \) represents a particular scenario \( i \) at a particular depth \( j \), referring to the \( i \)th row and \( j \)th column of \( \Phi \).
Algorithm 22.3. An implementation of determinized sparse tree search, a modification of forward search, for POMDPs. The policy takes a belief $b$ in the form of a vector of probabilities, which is approximated by a vector of determinized particles by `determinized_approximate_belief`.

```plaintext
struct DeterminizedSparseTreeSearch
    P # problem
    d # depth
    Φ # m×d determinizing matrix
    U # value function to use at leaf nodes
end

function determinized_sparse_tree_search(P, b, d, Φ, U)
    if d == 0
        return (a=nothing, u=U(b))
    end
    best = (a=nothing, u=-Inf)
    for a in A
        u = sum(R(ϕ.s, a) for ϕ in b) / length(b)
        for o in possible_observations(P, Φ, b, a)
            Poba = sum(sum(O(a,s′,o)*T(ϕ.s,a,s′) for s′ in S) for ϕ in b) / length(b)
            b′ = update(b, Φ, P, a, o)
            u′ = determinized_sparse_tree_search(P,b′,d-1,Φ,U).u
            u += γ*Poba*u'
        end
        if u > best.u
            best = (a=a, u=u)
        end
    end
    return best
end

function determinized_approximate_belief(b, P, m)
    particles = []
    for i in 1:m
        s = rand(SetCategorical(P.S, b))
        push!(particles, DeterminizedParticle(s, i, 1))
    end
    return particles
end

function (π::DeterminizedSparseTreeSearch)(b)
    particles = determinized_approximate_belief(b, π.P, size(π.Φ,1))
end
```
and lower bound values $\epsilon(b) = \overline{U}(b) - \underline{U}(b)$. Search algorithms with the gap heuristic select the observation that maximizes the gap because they are more likely to benefit from a belief backup. Actions are often selected according to a lookahead using an approximate value function.

The initial upper and lower bound values used in heuristic search play an important role in the algorithm's performance. Algorithm 22.4 uses the best-action best-state upper bound from equation (21.2). It uses a rollout policy for the lower bound $\underline{U}(b)$. A rollout, of course, is not guaranteed to produce a lower bound because it is based on a single trial up to a fixed depth. As the number of samples of increases, then it will converge to a true lower bound. Many other forms of upper and lower bounds exist that can provide faster convergence at the cost of run time and implementation complexity. For example, using the fast informed bound (algorithm 21.3) for the upper bound can improve exploration and help reduce the gap. For the lower bound, we can use a problem-specific rollout policy to better guide the search. Example 22.6 demonstrates this process.

### 22.8 Summary

- A simple online strategy is to perform a one-step lookahead, which considers each action taken from the current belief and estimates its expected value using an approximate value function.
- Forward search is a generalization of lookahead to arbitrary horizons, which can lead to better policies but its computational complexity grows exponentially with the horizon.
- Branch and bound is a more efficient version of forward search that can avoid searching certain paths by using upper and lower bounds on the value function.
- Sparse sampling is an approximation method that can reduce the computational burden of iterating over the space of all possible observations.
- Monte Carlo tree search can be adapted to POMDPs by operating over histories rather than states.
- Determinized sparse tree search uses a special form of particle belief that ensures observations are determinized, greatly reducing the search tree.
Algorithm 22.4. An implementation of heuristic search that uses bounds, a gap criterion, and a rollout policy as its heuristic. The upper and lower bounds are maintained by $\text{Uhi}$ and $\text{Ulo}$, respectively. The rollout policy $\pi$ must be a function that returns an action given a belief $\pi(b)$. At belief $b$, the gap is $\text{Uhi}[b] - \text{Ulo}[b]$. Exploration stops when the gap is smaller than the threshold $\delta$ or the maximum depth $d_{\text{max}}$ is reached. A maximum number of iterations $k_{\text{max}}$ is allotted to search.

```
struct GapHeuristicSearch
    \(\mathcal{P}\) # problem
    Uhi # upper bound on value function
    Ulo # lower bound on value function
    \(\pi\) # rollout policy
    \(\delta\) # gap threshold
    k_max # maximum number of simulations
    d_max # maximum depth
end

function heuristic_search(\(\pi::\text{GapHeuristicSearch}\), b, d)
    \(\mathcal{P}, \mathcal{A}, \mathcal{O}, \mathcal{R}, \gamma = \mathcal{P}.\mathcal{S}, \mathcal{P}.\mathcal{A}, \mathcal{P}.\mathcal{O}, \mathcal{P}.\mathcal{R}, \mathcal{P}.\gamma\)
    \(B = \text{Dict}((a, o) \Rightarrow \text{update}(b, \mathcal{P}, a, o) \text{ for } (a, o) \text{ in product}(\mathcal{A}, \mathcal{O}))\)
    \(B = \text{merge}(B, \text{Dict}()) \Rightarrow \text{copy}(b))\)
    Rmax = \text{maximum}(R(s, a) \text{ for } (s, a) \text{ in product}(\mathcal{S}, \mathcal{A}))
    for \((ao, b')\) in B
        if !\text{haskey}(\text{Uhi}[b'], b')
            \text{Uhi}[b'], \text{Ulo}[b'] = Rmax/(1.0-\gamma), \text{rollout}(\mathcal{P}, b', \pi, \mathcal{P}, d)
        end
    end
    if d == 0 || \text{Uhi}[b] - \text{Ulo}[b] \leq \delta
        return
    end
    a = \text{argmax}(a \rightarrow \text{lookahead}(\mathcal{P}, b' \rightarrow \text{Uhi}[b'], b, a), \mathcal{A})
    o = \text{argmax}(o \rightarrow \text{Uhi}[\mathcal{B}[(a, o)] - \text{Ulo}[\mathcal{B}[(a, o)]], o)
    b' = \text{update}(b, \mathcal{P}, a, o)
    \text{heuristic_search}(\pi, b', d-1)
    \text{Uhi}[b] = \text{maximum}(\text{lookahead}(\mathcal{P}, b' \rightarrow \text{Uhi}[b'], b, a) \text{ for } a \text{ in } \mathcal{A})
    \text{Ulo}[b] = \text{maximum}(\text{lookahead}(\mathcal{P}, b' \rightarrow \text{Ulo}[b'], b, a) \text{ for } a \text{ in } \mathcal{A})
end

function (\(\pi::\text{GapHeuristicSearch}\))(b)
    \(\mathcal{Uhi}, \mathcal{Ulo}, k_{\text{max}}, d_{\text{max}}, \delta = \pi.\mathcal{Uhi}, \pi.\mathcal{Ulo}, \pi.k_{\text{max}}, \pi.d_{\text{max}}, \pi.\mathcal{M}, \pi.\delta\)
    for i in 1:k_{\text{max}}
        \text{heuristic_search}(\pi, b, d_{\text{max}})
        if \text{Uhi}[b] - \text{Ulo}[b] < \delta
            break
        end
    end
    return \text{argmax}(a \rightarrow \text{lookahead}(\mathcal{P}, b' \rightarrow \text{Ulo}[b'], b, a), \mathcal{P}, \mathcal{A})
end
```
Consider applying online heuristic search to the crying baby problem. On each iteration of heuristic search, it explores the belief space following the bounds and improves. Below we show six iterations of heuristic search with an initial belief $b$ of $[0.5, 0.5]$, maximum depth $d_{\text{max}}$ of 10, and a random action rollout policy. In each iteration, the upper bound is shown in green and the lower bound is shown in blue.

We see that not all beliefs are explored in the first iteration. The iteration starts at $[0.5, 0.5]$ and heuristic search explores to the right first, towards belief $[0.0, 1.0]$. As the iterations progress, it slowly explores more towards $[1.0, 0.0]$. The jagged bounds are due to some beliefs not being re-explored based on the action and observation selection. There are computational savings due to pruning of the search tree. On the bottom row, we see that it has explored many of the beliefs once, but the bounds are still loose. Heuristic search guides itself towards the goal of reducing the gap.

Example 22.6. An example illustrating the use of heuristic search upper and lower bounds for the crying baby problem over iterations of heuristic search.
• Heuristic search intelligently selects action-observation pairs to explore regions with a high gap between the upper and lower bounds on the value function that it maintains.

22.9 Exercises

Exercise 22.1. Suppose we have \( A = \{a^1, a^2\} \) and a belief \( b = [0.5, 0.5] \). The reward is always 1. The observation function is given by \( P(o^1 \mid a^1) = 0.8 \) and \( P(o^1 \mid a^2) = 0.4 \). We have an approximate value function represented by an alpha vector \( \alpha = [-3, 4] \). With \( \gamma = 0.9 \), use forward search to a depth of 1 to compute \( U(b) \). Use the following updated beliefs in the calculation:

<table>
<thead>
<tr>
<th>( a )</th>
<th>( o )</th>
<th>Update(( b, a, o ))</th>
</tr>
</thead>
<tbody>
<tr>
<td>( a^1 )</td>
<td>( o^1 )</td>
<td>([0.3, 0.7])</td>
</tr>
<tr>
<td>( a^2 )</td>
<td>( o^1 )</td>
<td>([0.2, 0.8])</td>
</tr>
<tr>
<td>( a^1 )</td>
<td>( o^2 )</td>
<td>([0.5, 0.5])</td>
</tr>
<tr>
<td>( a^2 )</td>
<td>( o^2 )</td>
<td>([0.8, 0.2])</td>
</tr>
</tbody>
</table>

Solution: We need to calculate the action value function at depth 1 according to equation (22.1):

\[
Q_d(b, a) = R(b, a) + \gamma \sum_o P(o \mid b, a) U_{d-1}(\text{Update}(b, a, o))
\]

We first calculate the utility for the updated beliefs.

\[
U_0(\text{Update}(b, a^1, o^1)) = \alpha^\top b' = 0.3 \times -3 + 0.7 \times 4 = 1.9
\]

\[
U_0(\text{Update}(b, a^2, o^1)) = 0.2 \times -3 + 0.8 \times 4 = 2.6
\]

\[
U_0(\text{Update}(b, a^1, o^2)) = 0.5 \times -3 + 0.5 \times 4 = 0.5
\]

\[
U_0(\text{Update}(b, a^2, o^2)) = 0.8 \times -3 + 0.2 \times 4 = -1.6
\]

Second, we compute the action value function for both actions:

\[
Q_1(b, a^1) = 1 + 0.9((P(o^1 \mid b, a^1)U_0(\text{Update}(b, a^1, o^1)) + (P(o^2 \mid b, a^1)U_0(\text{Update}(b, a^1, o^2)))
= 1 + 0.9(0.8 \times 1.9 + 0.2 \times 0.5) = 2.458
\]

\[
Q_1(b, a^2) = 1 + 0.9((P(o^1 \mid b, a^2)U_0(\text{Update}(b, a^2, o^1)) + (P(o^2 \mid b, a^2)U_0(\text{Update}(b, a^2, o^2)))
= 1 + 0.9(0.4 \times 2.6 + 0.6 \times -1.6) = 1.072
\]

Finally, we have \( U_1(b) = \max_a Q_1(b, a) = 2.458 \)
Exercise 22.2. Using the following trajectory samples, compute the action value function for belief $b$ and actions $a^1$ and $a^2$ based on sparse sampling to depth 1. Use the following updated beliefs, discount factor $\gamma = 0.9$, and approximate value function represented by an alpha vector $\alpha = [10, 1]$.

<table>
<thead>
<tr>
<th>$a$</th>
<th>$o$</th>
<th>$r$</th>
<th>Update($b,a,o$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>1</td>
<td>0</td>
<td>[0.47, 0.53]</td>
</tr>
<tr>
<td>2</td>
<td>1</td>
<td>1</td>
<td>[0.22, 0.78]</td>
</tr>
<tr>
<td>1</td>
<td>2</td>
<td>1</td>
<td>[0.49, 0.51]</td>
</tr>
<tr>
<td>2</td>
<td>1</td>
<td>1</td>
<td>[0.22, 0.78]</td>
</tr>
<tr>
<td>2</td>
<td>2</td>
<td>1</td>
<td>[0.32, 0.68]</td>
</tr>
<tr>
<td>1</td>
<td>2</td>
<td>1</td>
<td>[0.49, 0.51]</td>
</tr>
</tbody>
</table>

Solution: We first calculate the utility for the updated beliefs:

<table>
<thead>
<tr>
<th>$a$</th>
<th>$o$</th>
<th>$r$</th>
<th>Update($b,a,o$)</th>
<th>$U_0$(Update($b,a,o$))</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>1</td>
<td>0</td>
<td>[0.47, 0.53]</td>
<td>5.23</td>
</tr>
<tr>
<td>2</td>
<td>1</td>
<td>1</td>
<td>[0.22, 0.78]</td>
<td>2.98</td>
</tr>
<tr>
<td>1</td>
<td>2</td>
<td>1</td>
<td>[0.49, 0.51]</td>
<td>5.41</td>
</tr>
<tr>
<td>2</td>
<td>1</td>
<td>1</td>
<td>[0.22, 0.78]</td>
<td>2.98</td>
</tr>
<tr>
<td>2</td>
<td>2</td>
<td>1</td>
<td>[0.32, 0.68]</td>
<td>3.88</td>
</tr>
<tr>
<td>1</td>
<td>2</td>
<td>1</td>
<td>[0.49, 0.51]</td>
<td>5.41</td>
</tr>
</tbody>
</table>

Then, we can compute the action value function over all actions using equation (22.2):

$$Q_1(b,a^1) = \frac{1}{3}(0 + 1 + 1 + 0.9(5.23 + 5.41 + 5.41)) = 5.48$$

$$Q_1(b,a^2) = \frac{1}{3}(1 + 1 + 1 + 0.9(2.98 + 2.98 + 3.88)) = 3.95$$

Exercise 22.3. Consider example 22.5. Give the new particle associated with $\phi = (1, 4, 2)$. Suppose we take action $a_3$, and we have the following transition functions:

$$T(s_2 \mid s_1,a_3) = 0.4 \quad O(o_1 \mid s_1,a_3) = 0.6$$

$$T(s_3 \mid s_1,a_3) = 0.45 \quad O(o_2 \mid s_1,a_3) = 0.5$$

Solution: From the determinizing matrix, our determinizing value is $\Phi_{4,2} = 0.598$ and we are in state $s_1$. Then, we calculate $p$:

$$p \leftarrow T(s_2 \mid s_1,a_3)O(o_1 \mid s_1,a_3) = 0.4 \times 0.6 = 0.24 \quad (22.4)$$

$$p \leftarrow p + T(s_2 \mid s_1,a_3)O(o_2 \mid s_1,a_3) = 0.24 + 0.4 \times 0.5 = 0.44 \quad (22.5)$$

$$p \leftarrow p + T(s_3 \mid s_1,a_3)O(o_1 \mid s_1,a_3) = 0.44 + 0.45 \times 0.6 = 0.71 \quad (22.6)$$

We stop our iteration because $p > 0.598$. Thus, from our final iteration, we proceed to $(s_3,o_1)$.  

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2021-03-07 23:12:00-08:00, revision 80bf4d6, comments to bugs@algorithmsook.com
Exercise 22.4. Summarize the techniques covered to reduce branching over actions.

Solution: Branch and bound can reduce action branching by using an upper bound on the value function. It skips actions that cannot improve upon the value obtained from actions it explored earlier. Gap heuristic search and Monte Carlo tree search use approximations of action values to guide the selection of actions during exploration.

Exercise 22.5. Summarize the techniques covered to reduce branching over observations.

Solution: Sparse sampling reduces observation branching by sampling only a small number of observations. Observations are sampled from $P(o \mid b, a)$, which means that observations that have greater probability are more likely to be sampled. Determinized sparse tree search uses a similar approach, but the sampling occurs once and is then fixed. Branching over observations can also be reduced based on the lookahead value $U(b')$. Gap heuristic search evaluates the gap and avoids branching on observations for which we have high confidence in the value function.
23 Controller Abstractions

This chapter introduces controller representations for POMDP policies, which allow policies to maintain their own internal state. These representations can improve scalability over previous methods that enumerate over belief points. This chapter presents algorithms that construct controllers using policy iteration, mathematical programming, and gradient ascent.

23.1 Controllers

A controller is a policy representation that maintains its own internal state. This controller is represented as a graph consisting of a finite set of nodes $X$. The active node changes as actions are taken and new observations are made. Having a finite set of nodes makes these controllers more computationally tractable than belief-point methods that must consider the reachable belief space.

Actions are selected according to an action distribution $\psi(a \mid x)$ that depends on the current node. When selecting an action, in addition to transitioning to an unobserved $s'$ and receiving an observation $o$, the control state also advances according to a successor distribution $\eta(x' \mid x, a, o)$. Figure 23.1 shows how these distributions are used as a controller policy is followed. Algorithm 23.1 provides an implementation, and example 23.1 shows a controller for the crying baby problem.

Controllers generalize conditional plans, which were introduced in section 20.2. Conditional plans represent policies as trees, with each node deterministically assigning an action and each edge specifying a unique successor node. Controllers represent policies as directed graphs, and actions may have stochastic transitions to multiple successor nodes. Example 23.2 compares these two representations.

Such a policy representation is also called a finite state controller. We will endeavor to refer to the controller states as nodes rather than states to reduce ambiguity with the environment state.
We can construct a simple controller for the crying baby problem (appendix F.7). This policy is shown below as a graph with two nodes, $x^1$ and $x^2$. When in $x^1$, the controller always ignores the baby. When in $x^2$, the controller always feeds the baby. If the baby cries, we always transition to $x^2$, and if the baby is quiet, we always transition to $x^1$.

Example 23.1. A two-node example controller for the crying baby problem. This compact representation captures a straightforward solution to the crying baby problem, namely to react immediately to the most recent observation.

Algorithm 23.1. A finite state controller policy representation for a POMDP $\mathcal{P}$. The nodes in $x$ are an abstract representation of reachable beliefs. Actions and controller successor nodes are selected stochastically. Given a node $x$, actions are selected following the distribution $\psi$. The function $\pi(x)$ implements this mechanism to stochastically select actions. After performing action $a$ in node $x$ and observing observation $o$, the successor is selected following the distribution $\eta$. The function $\text{update}$ implements this mechanism to stochastically select successor nodes.
Consider a three-step conditional plan (below left) compared with the more general two-node finite state controller (below right) from example 23.1. In this case, actions and successors are selected deterministically. The deterministic action is marked in the center of a node, and the outgoing edges represent the deterministic successor nodes. This problem has two actions ($a^1$ and $a^2$) and two observations ($o^1$ and $o^2$).

The conditional plan performs action $a^1$ first, toggles the previously chosen action if it observes $o^1$, and preserves the previously chosen action if it observes $o^2$. The controller performs the same logic, with five fewer controller nodes. Moreover, the controller represents the described infinite horizon policy perfectly with only two nodes (compared to seven). The conditional plan cannot capture this infinite horizon policy because it would require a tree of infinite depth.
Controllers have several advantages over conditional plans. First, controllers can provide a more compact representation. The number of nodes in a conditional plan grows exponentially with depth, but this need not be the case with finite state controllers. The approximation methods from previous chapters might also not be as efficient because they must maintain a large set of beliefs and corresponding alpha vectors. Controllers can be much more compact, considering infinitely many possible reachable beliefs with a small, finite number of nodes. Another advantage of controllers is that they do not require that a belief be maintained.\footnote{Each controller node corresponds to a subset of the belief space. A controller transitions between these subsets that together cover the reachable belief space.}

The controller itself selects a new node based on each observation, rather than relying on a belief update, which can be expensive for some domains.

The utility when following a controller can be computed by forming a product of \(\pi\) and \(X\) for \(s\) and \(s'\) following equation (23.1).

\[
U(x, s) = \sum_a \psi(a | x) \left( R(s, a) + \gamma \sum_{s'} T(s' | s, a) \sum_o O(o | a, s') \sum_{x'} \eta(x' | x, a, o) U(x', s') \right)
\]  

(23.1)

Policy evaluation involves solving the system of linear equations in equation (23.1). Alternatively, we can apply iterative policy evaluation as shown in algorithm 23.2.

\begin{verbatim}
function utility(\pi :: ControllerPolicy, U, x, s)
    \(\mathcal{S}, \mathcal{X}, O = \pi.\mathcal{T.S}, \pi.\mathcal{T.X}, \pi.\mathcal{T.O}\)
    \(T, O, R, \gamma = \pi.\mathcal{T.T}, \pi.\mathcal{T.O}, \pi.\mathcal{T.R}, \pi.\mathcal{T.\gamma}\)
    \(X, \psi, \eta = \pi.\mathcal{X}, \pi.\psi, \pi.\eta\)
    \(U'(a, s', o) = \text{sum}(\eta[x, a, o, x'] \cdot U[x', s'] \text{ for } x \text{ in } X)\)
    \(U'(a, s') = T(s, a, s') \cdot \text{sum}(O(a, s', o) \cdot U'(a, s', o) \text{ for } o \text{ in } O)\)
    \(U'(a) = R(s, a) + \gamma \text{sum}(U'(a, s') \text{ for } s' \text{ in } \mathcal{S})\)
    return \text{sum}(\psi[x, a] \cdot U'(a) \text{ for } a \text{ in } \mathcal{X})
end

function iterative_policy_evaluation(\pi :: ControllerPolicy, k_max)
    \(\mathcal{S}, \mathcal{X} = \pi.\mathcal{T.S}, \pi.\mathcal{X}\)
    \(U = \text{Dict}((x, s) => 0.0 \text{ for } x \text{ in } \mathcal{X}, s \text{ in } \mathcal{S})\)
    for k in 1:k_max
        \(U = \text{Dict}((x, s) => \text{utility}(\pi, U, x, s) \text{ for } x \text{ in } \mathcal{X}, s \text{ in } \mathcal{S})\)
    end
    return U
end
\end{verbatim}

If a belief is known, then the current value is

\[
U(x, b) = \sum_s b(s) U(x, s)
\]  

(23.2)
We can think of $U(x, s)$ as defining a set of alpha vectors, one for each node $x$ in $X$. Each alpha vector $\alpha_x$ is defined by $\alpha_x(s) = U(x, s)$. The current value for a given alpha vector is $U(x, b) = b^T \alpha_x$.

Given a controller and an initial belief, we can select an initial node by maximizing:

$$x^* = \arg \max_x U(x, b) = \arg \max_x b^T \alpha_x \quad (23.3)$$

### 23.2 Policy Iteration

Section 20.5 showed how to incrementally add nodes in a conditional plan to arrive at optimal finite-horizon policy (algorithm 20.8). This section shows how to incrementally add nodes to a controller to optimize for infinite horizon problems. Although the policy representation is different, the version of policy iteration for partially observable problems introduced in this section has some similarities with the policy iteration algorithm for fully observed problems (section 7.4).

Policy iteration (algorithm 23.3) begins with any initial controller, and then iterates between policy evaluation and policy improvement. In policy evaluation, we evaluate the utilities $U(x, s)$ by solving equation (23.1). In policy improvement, we introduce new nodes to our controller. Specifically, we introduce a new node $x'$ for every combination of deterministic action assignments $\psi(a_i \mid x') = 1$ and deterministic successor selection distributions $\eta(x \mid x', a, o)$. This process adds $|A||X(k)||O|$ new controller nodes to the set of nodes $X(k)$ at iteration $k$. An improvement step is demonstrated in example 23.3.

Policy improvement cannot worsen the expected value of the controller policy. The value of any nodes in $X(k)$ remain unchanged, as they and their reachable successor nodes remain unchanged. It is guaranteed that if $X(k)$ is not an optimal controller, then at least one of the new nodes introduced in policy improvement will have better expected values for some states, and thus the overall controller must be improved.

Many of the nodes added during policy improvement tend to not improve the policy. Pruning is conducted after policy evaluation to eliminate unnecessary nodes. Pruning does not degrade the optimal value function of the controller and can help reduce the exponential growth in nodes that comes with the improvement step.

---

3 The policy iteration method given here was given by E. A. Hansen, “Solving POMDPs by Searching in Policy Space,” in Conference on Uncertainty in Artificial Intelligence (UAI), 1998.

4 Adding all possible combinations is often not feasible. An alternative algorithm called bounded policy iteration adds only one node. P. Poupart and C. Boutilier, “Bounded Finite State Controllers,” in Advances in Neural Information Processing Systems (NIPS), 2003. Algorithms can also add a number in between. Monte Carlo value iteration, for example, adds $O(n|A||X^{(k)}|)$ new nodes at each iteration $k$, where $n$ is a parameter. H. Bai, D. Hsu, W.S. Lee, and V.A. Ngo, “Monte Carlo Value Iteration for Continuous-State POMDPs,” in International Workshop on the Algorithmic Foundations of Robotics (WAFFR), 2011.
Algorithm 23.3. Policy iteration for a POMDP $\mathcal{P}$ given a fixed number of iterations $k_{\text{max}}$ and number of policy evaluation iterations $\text{eval}_{\text{max}}$. The algorithm iteratively applies policy evaluation from algorithm 23.2 and policy improvement. Pruning is implemented in algorithm 23.4.

```
struct ControllerPolicyIteration
    $k_{\text{max}}$  # number of iterations
    $\text{eval}_{\text{max}}$  # number of evaluation iterations
end

function solve($M$::ControllerPolicyIteration, $\mathcal{P}$::POMDP)
    $\mathcal{X}$, $\mathcal{O}$, $k_{\text{max}}$, $\text{eval}_{\text{max}}$ = $\mathcal{P}$.\mathcal{X}, $\mathcal{P}$.\mathcal{O}, $M$.\text{eval}_{\text{max}}
    $X$ = [1]
    $\psi$ = Dict((x, a) $\Rightarrow$ $1.0 / \text{length}(A)$ for x in $X$, a in $\mathcal{A}$)
    $\eta$ = Dict((x, a, o, x') $\Rightarrow$ $1.0$ for x in $X$, a in $\mathcal{A}$, o in $\mathcal{O}$, x' in $X$)
    $\pi$ = ControllerPolicy($\mathcal{P}$, $X$, $\psi$, $\eta$)
    for i in 1:$k_{\text{max}}$
        prevX = copy($\pi$.\mathcal{X})
        $U$ = iterative_policy_evaluation($\pi$, $\text{eval}_{\text{max}}$)
        policy_improvement!($\pi$, $U$, prevX)
        prune!($\pi$, $U$, prevX)
    end
    return $\pi$
end

function policy_improvement!($\pi$::ControllerPolicy, $U$, prevX)
    $\mathcal{S}$, $\mathcal{A}$, $\mathcal{O}$ = $\pi$.\mathcal{S}, $\pi$.\mathcal{A}, $\pi$.\mathcal{O}
    $X$, $\psi$, $\eta$ = $\pi$.\mathcal{X}, $\pi$.\psi, $\pi$.\eta
    repeatX$\mathcal{O}$ = fill($X$, length($\mathcal{O}$))
    assign$\mathcal{AX}'$ = vec(collect(product($\mathcal{A}$, repeatX$\mathcal{O}$...)))
    for ax' in assign$\mathcal{AX}'$
        x, a = maximum($X$) + 1, ax'[1]
        push!(X, x)
        successor(o) = ax'[findfirst(isequal(o), $\mathcal{O}$) + 1]
        $U'(o,s') = U[\text{successor}(o), s']$
        for s in $\mathcal{S}$
            $U[x, s] = $\text{lookahead}($\pi$.\mathcal{P}, $U'$, s, a)
        end
        for a' in $\mathcal{A}$
            $\psi[x, a'] = a' == a ? 1.0 : 0.0$
            for (o, x') in product($\mathcal{O}$, prevX)
                $\eta[x, a', o, x'] = x' == \text{successor}(o) ? 1.0 : 0.0$
            end
        end
    end
    for (x, a, o, x') in product($X$, $\mathcal{A}$, $\mathcal{O}$, $X$)
        if !haskey($\eta$, (x, a, o, x'))
            $\eta[x, a, o, x'] = 0.0$
        end
    end
end
```
We can apply policy improvement to the crying baby controller from example 23.1. The actions are $A = \{\text{feed, sing, ignore}\}$ and observations are $O = \{\text{crying, quiet}\}$. The policy improvement backup step results in $|A||X^{(1)}||O| = 3 \times 2^2 = 12$ new nodes. The new controller policy has nodes $\{x^1, \ldots, x^{14}\}$ and distributions as follows:

<table>
<thead>
<tr>
<th>Node</th>
<th>Action</th>
<th>Successors (for all $a$ below)</th>
</tr>
</thead>
<tbody>
<tr>
<td>$x^3$</td>
<td>$\psi(\text{feed}</td>
<td>x^3) = 1$</td>
</tr>
<tr>
<td>$x^4$</td>
<td>$\psi(\text{feed}</td>
<td>x^4) = 1$</td>
</tr>
<tr>
<td>$x^5$</td>
<td>$\psi(\text{feed}</td>
<td>x^5) = 1$</td>
</tr>
<tr>
<td>$x^6$</td>
<td>$\psi(\text{feed}</td>
<td>x^6) = 1$</td>
</tr>
<tr>
<td>$x^7$</td>
<td>$\psi(\text{sing}</td>
<td>x^7) = 1$</td>
</tr>
<tr>
<td>$x^8$</td>
<td>$\psi(\text{sing}</td>
<td>x^8) = 1$</td>
</tr>
</tbody>
</table>

We have the following controller, with new nodes in blue and the original two nodes in black:

![Diagram showing the new controller policy](image-url)
We prune any new nodes that are identical to existing nodes. We also prune any new nodes that are dominated by other nodes. A node \( x \) is dominated by another node \( x' \) when:

\[
U(x, s) \leq U(x', s) \quad \text{for all } s
\]

(23.4)

Existing nodes can be pruned as well. Whenever a new node dominates an existing node, we prune the existing node from the controller. Any transitions to the deleted node are instead rerouted to the dominating node.

Example 23.4

This process is identical to pruning the new node instead, and updating the dominated node’s action and successor links to those of the new node.

23.3 Nonlinear Programming

The policy improvement problem can be framed as a single large nonlinear programming formulation (algorithm 23.5) that involves simultaneously optimizing for \( \psi \) and \( \eta \) across all nodes. This formulation allows general purpose solvers to be applied. The nonlinear programming method directly searches the space of controllers to maximize the utility of a given initial belief while satisfying the Bellman equation, equation (23.1). There is no alternating between policy evaluation and policy improvement steps, and the controller node count remains fixed.

We use \( x^1 \) to denote the initial node corresponding to the given initial belief \( b \). The optimization problem is then:

\[
\text{maximize}_{U, \psi, \eta} \sum_s b(s)U(x^1, s)
\]

subject to

\[
U(x, s) = \sum_a \psi(a | x) \left( R(s, a) + \gamma \sum_{s'} T(s' | s, a) \sum_o O(o | a, s') \sum_{x'} \eta(x' | x, a, o) U(x', s') \right)
\]

for all \( x, s \)

(23.5)

\[
\psi(a | x) \geq 0 \quad \text{for all } x, a
\]

\[
\sum_a \psi(a | x) = 1 \quad \text{for all } x
\]

\[
\eta(x' | x, a, o) \geq 0 \quad \text{for all } x, a, o, x'
\]

\[
\sum_{x'} \eta(x' | x, a, o) = 1 \quad \text{for all } x, a, o
\]
function prune!(π::ControllerPolicy, U, prevX)
    newX, removeX = setdiff(X, prevX), []
    # prune dominated from previous nodes
    dominated(x,x′) = all(U[x,s] ≤ U[x′,s] for s in S)
    for (x,x′) in product(prevX, newX)
        if x′ ∉ removeX && dominated(x, x′)
            for s in S
                U[x,s] = U[x′,s]
            end
            for a in A
                ψ[x,a] = ψ[x′,a]
                for (o,x′′) in product(O, X)
                    η[x,a,o,x′′] = η[x′,a,o,x′′]
                end
            end
            push!(removeX, x′)
        end
    end
    # prune identical from previous nodes
    identical_action(x,x′) = all(ψ[x,a] ≈ ψ[x′,a] for a in A)
    identical_successor(x,x′) = all(η[x,a,o,x′′] ≈ η[x′,a,o,x′′] for a in A, o in O, x′′ in X)
    identical(x,x′) = identical_action(x,x′) && identical_successor(x,x′)
    for (x,x′) in product(prevX, newX)
        if x′ ∉ removeX && identical(x, x′)
            push!(removeX, x′)
        end
    end
    # prune dominated from new nodes
    for (x,x′) in product(X, newX)
        if x′ ∉ removeX && dominated(x′,x) && x ≠ x′
            push!(removeX, x′)
        end
    end
    # update controller
    π.X = setdiff(X, removeX)
    π.ψ = Dict(k => v for (k,v) in ψ if k[1] ∉ removeX)
    π.η = Dict(k => v for (k,v) in η if k[1] ∉ removeX)
end

Algorithm 23.4. The pruning step of policy iteration. It reduces the number of nodes in the current policy π, using the utilities U computed by policy evaluation, and the previous node list prevX. Its first step replaces any pointwise dominated previous nodes by their improved nodes, marking the redundant node as now dominated. The second step marks any newly added nodes that are identical to previous nodes. The third step marks any pointwise dominated new nodes. Finally, all marked nodes are pruned.
Recall example 23.3. Below we show the first iteration of policy iteration using the same initial controller. It consists of the two main steps: policy evaluation (left) and policy improvement (center), as well as the optional pruning step (right).

The second iteration of policy iteration follows the same pattern:

The utility has greatly improved after the second iteration to near optimal values. We see that the prune step removes dominated and duplicate nodes from previous iterations as well as the current iteration’s new nodes.
Nonlinear Programming

This problem can be written as a quadratically constrained linear program (QCLP), which can be solved efficiently using a dedicated solver. Example 23.5 demonstrates this approach.

```
struct NonlinearProgramming
    b # initial belief
    ℓ # number of nodes
end

function tensorform(ℙ::POMDP)
    ℳ, ℬ, ℰ, ℓ, ℛ, ℓ = ℙ.ℳ, ℙ.ℬ, ℙ.ℰ, ℙ.ℓ, ℙ.ℛ, ℙ.ℓ
    ℳ′ = eachindex(ℳ)
    ℬ′ = eachindex(ℬ)
    ℰ′ = eachindex(ℰ)
    ℓ′ = eachindex(ℓ)
    ℛ′ = [ℛ(ℳ,a) for ℳ in ℳ, ℬ in ℬ]
    ℓ′ = ℛ(ℳ, ℬ, ℰ′) for ℳ in ℳ, ℬ in ℬ, ℰ′ in ℰ
    return ℳ′, ℬ′, ℰ′, ℛ′, ℓ′
end

function solve(M::NonlinearProgramming, ℙ::POMDP)
    ℇ₁, ℇ = 1, collect(1:M.ℓ)
    ℙ, ℰ, ℏ, ℒ = ℙ, ℙ.ℰ, ℙ.ℏ, ℙ.ℒ
    ℳ, ℬ, ℰ, ℓ, ℛ, ℓ = tensorform(ℙ)
    model = Model(Ipopt.Optimizer)
    @variable(model, ℛ[ℳ, ℬ])
    @variable(model, ℏ[ℳ, ℬ])
    @variable(model, ℓ[ℳ, ℬ, ℰ, ℓ])
    @objective(model, Max, ℏ * ℒ[
        ℇ₁, ℇ]
    @NLconstraint(model, ℳ[ℳ, ℬ], ℛ[ℳ, ℬ], ℰ[ℳ, ℬ, ℰ, ℓ], ℓ[ℳ, ℬ, ℰ, ℓ])
    @constraint(model, ℳ[ℳ, ℬ], ℏ[ℳ, ℬ])
    @constraint(model, ℳ[ℳ, ℬ, ℰ, ℓ], ℓ[ℳ, ℬ, ℰ, ℓ])
    optimize!(model)
    ℏ, ℓ = value.(ℏ), value.(ℓ)
    return ControllerPolicy(ℙ, ℳ, Dict(( ℙ.ℳ[ℳ, ℬ] ⇒ ℏ[ℳ, ℬ] for ℳ in ℳ, ℬ in ℬ)),
    Dict(( ℙ.ℳ[ℳ, ℬ, ℰ, ℓ] ⇒ ℓ[ℳ, ℬ, ℰ, ℓ], ℙ.ℰ[ℰ]) ⇒ ℇ for ℳ in ℳ, ℬ in ℬ, ℰ in ℰ, ℓ in ℓ))
end
```

Algorithm 23.5. A nonlinear programming approach to compute the optimal fixed-size controller policy for POMDP ℙ starting at initial belief ℏ. The size of the finite state controller is specified by number of nodes ℓ.
Below are optimal fixed-size controllers computed using nonlinear programming for the crying baby problem with $b_0 = [0.5, 0.5]$. The top node is $x_1$. controller utility ($k = 1$)

Example 23.5. An example of the nonlinear programming algorithm for controllers with a fixed size of $k$ set to 1, 2, and 3. Each row shows the policy and its corresponding utilities (alpha vectors) on the left and right, respectively. The stochastic controllers are shown as circles, with the most likely action in the middle. The outgoing edges show successor node selections given an observation. The stochasticity in node actions and successors are shown as opacity (more opaque is higher probability, more transparent is lower probability).

With $k = 1$, the optimal policy is to simply ignore forever. With $k = 2$, the optimal policy is to ignore until crying is observed, at which point the best action is to feed the child, and then return to ignoring. This policy is close to optimal for the infinite horizon crying baby POMDP. With $k = 3$, the optimal policy essentially remains unchanged from when $k = 2$. 

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2021-03-07 23:12:00-08:00, revision 80bf4d6, comments to bugs@algorithmsbook.com
23.4 Gradient Ascent

A fixed-size controller policy can be iteratively improved using gradient ascent (covered in appendix A.11). Though the gradient is challenging to compute, this opens up controller optimization to a wide variety of gradient-based optimization techniques. Algorithm 23.6 implements controller gradient ascent using algorithm 23.7.

Consider an explicit description of the nonlinear problem from section 23.3. For initial belief $b$ and an arbitrary initial controller node $x^1$, we seek to maximize:

$$\sum_s b(s) U(x^1, s) \tag{23.6}$$

with the utility $U(x, s)$ defined by the Bellman equation for all $x$ and $s$:

$$U(x, s) = \sum_a \psi(a | x) \left( R(s, a) + \gamma \sum_{s'} T(s' | s, a) \sum_o O(o | a, s') \sum_{x'} \eta(x' | x, a, o) U(x', s') \right) \tag{23.7}$$

In addition, $\psi$ and $\eta$ must also be proper probability distributions. To apply gradient ascent, it is more convenient to rewrite this problem using linear algebra.

We define the transition function with a controller, which has a state space $X \times S$. For any fixed-size controller policy parameterized by $\theta = (\psi, \eta)$, the transition matrix $T_\theta \in \mathbb{R}^{|X| \times |S| \times |X| \times |S|}$ is:

$$T_\theta((x, s), (x', s')) = \sum_a \psi(x, a) T(s, a, s') \sum_o O(a, s', o) \eta(x, a, o, x') \tag{23.8}$$

The reward for a parameterized policy is represented as a vector $r_\theta \in \mathbb{R}^{|X| \times |S|}:

$$r_\theta((x, s)) = \sum_a \psi(x, a) R(s, a) \tag{23.9}$$

The Bellman equation for utility $u_\theta \in \mathbb{R}^{|X| \times |S|}$ is then:

$$u_\theta = r_\theta + \gamma T_\theta u_\theta \tag{23.10}$$

We can consider an initial node-belief vector $\beta \in \mathbb{R}^{|X| \times |S|}$ with $\beta_{xs} = b(s)$ if $x = x^1$ and $\beta_{xs} = 0$ otherwise. A utility vector $u_\theta \in \mathbb{R}^{|X| \times |S|}$ also is defined over the nodes $X$ and states $S$ for any of these fixed-size parameterized controller policies $\theta = (\psi, \eta)$. We now seek to maximize:

$$\beta^T u_\theta \tag{23.11}$$
We begin by rewriting equation (23.10):

\[ u_\theta = r_\theta + \gamma T_\theta u_\theta \]  
\[ (I - \gamma T_\theta)u_\theta = r_\theta \]  
\[ u_\theta = (I - \gamma T_\theta)^{-1}r_\theta \]  
\[ u_\theta = Z^{-1}r_\theta \]

with \( Z = I - \gamma T_\theta \) for convenience. In order to perform gradient ascent, we need to know the partial derivatives of equation (23.15) with respect to the policy parameters:

\[
\frac{\partial u_\theta}{\partial \theta} = \frac{\partial Z^{-1}}{\partial \theta} r_\theta + Z^{-1} \frac{\partial r_\theta}{\partial \theta} \\
= -Z^{-1} \frac{\partial Z}{\partial \theta} Z^{-1} r_\theta + Z^{-1} \frac{\partial r_\theta}{\partial \theta} \\
= Z^{-1} \left( \frac{\partial r_\theta}{\partial \theta} - \frac{\partial Z}{\partial \theta} Z^{-1} r_\theta \right)
\]

with \( \partial \theta \) referring to both \( \partial \psi(\hat{x}, \hat{a}) \) and \( \partial \eta(\hat{x}, \hat{a}, \hat{\delta}, \hat{x}') \) for convenience.

Computing the partial derivatives of \( Z \) and \( r_\theta \) results in four equations:

\[
\frac{\partial r_\theta((x, s))}{\partial \psi(\hat{x}, \hat{a})} = \begin{cases} 
R(s, a) & \text{if } x = \hat{x} \\
0 & \text{otherwise}
\end{cases}
\]

(23.19)

\[
\frac{\partial r_\theta((x, s))}{\partial \eta(\hat{x}, \hat{a}, \hat{\delta}, \hat{x}')} = 0
\]

(23.20)

\[
\frac{\partial Z((x, s), (x', s'))}{\partial \psi(\hat{x}, \hat{a})} = \begin{cases} 
-\gamma T(s, \hat{a}, s') \sum_o O(\hat{a}, s', o) \eta(\hat{x}, \hat{a}, o, x') & \text{if } x = \hat{x} \\
0 & \text{otherwise}
\end{cases}
\]

(23.21)

\[
\frac{\partial Z((x, s), (x', s'))}{\partial \eta(\hat{x}, \hat{a}, \hat{\delta}, \hat{x}')} = \begin{cases} 
-\gamma \psi(\hat{x}, \hat{a}) T(s, \hat{a}, s') O(\hat{a}, s', \hat{\delta}) \eta(\hat{x}, \hat{a}, \hat{\delta}, x') & \text{if } x = \hat{x} \text{ and } x' = \hat{x}' \\
0 & \text{otherwise}
\end{cases}
\]

(23.22)

Finally, these four gradients are used in the utility gradients from equation (23.18):

\[
\frac{\partial u_\theta}{\partial \psi(\hat{x}, \hat{a})} = Z^{-1} \left( \frac{\partial r_\theta}{\partial \psi(\hat{x}, \hat{a})} - \frac{\partial Z}{\partial \psi(\hat{x}, \hat{a})} Z^{-1} r_\theta \right)
\]

(23.23)

\[
\frac{\partial u_\theta}{\partial \eta(\hat{x}, \hat{a}, \hat{\delta}, \hat{x}')} = Z^{-1} \left( \frac{\partial r_\theta}{\partial \eta(\hat{x}, \hat{a}, \hat{\delta}, \hat{x}')} - \frac{\partial Z}{\partial \eta(\hat{x}, \hat{a}, \hat{\delta}, \hat{x}')} Z^{-1} r_\theta \right)
\]

(23.24)
We finally can return to the original objective in equation (23.11). Controller gradient ascent starts with a fixed number of nodes in $X$ and an arbitrary policy $\psi$ and $\eta$. At iteration $k$, it updates these parameters as follows:

$$\psi^{k+1}(x, a) = \psi^k(x, a) + \alpha \beta^\top \frac{\partial u^k_\theta}{\partial \psi^k(\hat{x}, \hat{a})}$$ (23.25)

$$\eta^{k+1}(x, a, o, x') = \eta^k(x, a, o, x') + \alpha \beta^\top \frac{\partial u^k_\theta}{\partial \eta^k(\hat{x}, \hat{a}, \hat{o}, \hat{x'})}$$ (23.26)

with gradient step size $\alpha > 0$. After this update, $\psi^{k+1}$ and $\eta^{k+1}$ may no longer be valid distributions. To make them valid distributions, we project them onto the probability simplex. One approach to projecting a vector $y$ onto the probability simplex is to solve the following quadratic program:

$$\begin{align*}
\text{minimize} & \quad \frac{1}{2} \| y - b \|^2_2 \\
\text{subject to} & \quad b \geq 0 \\
& \quad 1^\top b
\end{align*}$$ (23.27)

This optimization can be solved exactly through a simple algorithm included in algorithm 23.6. Example 23.6 demonstrates the process of updating the controller.

The optimization objective in equation (23.6) is not necessarily convex. Hence, normal gradient ascent can converge to a local optimum depending on the initial controller. Adaptive gradient algorithms can be applied to help smooth and speed convergence.

23.5 Summary

- Controllers are a policy representation that do not rely on exploring or maintaining beliefs.

- Controllers consist of nodes, an action selection function, and a successor selection function.

- Nodes and the controller graph are abstract, however, they can be interpreted as sets of the countably infinite reachable beliefs; the value function for a node can also be interpreted as an alpha vector.

---


\* This objective is distinct from the utility $U(x, b) = \sum_s b(s) U(x, s)$, which is guaranteed to be piece-wise linear and convex with respect to the belief state $b$ as discussed in section 20.3.
struct ControllerGradient
b  # initial belief
ℓ  # number of nodes
α  # gradient step
k_max  # maximum iterations
end

function solve(M::ControllerGradient, P::POMDP)
    ℳ, Ω, ℓ, k_max = ℳ.ℳ, ℳ.Ω, ℳ.ℓ, ℳ.k_max
    X = collect(1:ℓ)
    ψ = Dict((x, a) => rand() for x in X, a in Ω)
    η = Dict((x, a, o, x') => rand() for x in X, a in Ω, o in Ω, x' in X)
    π = ControllerPolicy(ℳ, X, ψ, η)
    for i in 1:k_max
        improve!(π, ℳ, P)
    end
    return π
end

function improve!(π::ControllerPolicy, M::ControllerGradient, P::POMDP)
    ℳ, Ω, X, x1, ψ, η = ℳ.ℳ, ℳ.Ω, ℳ.X, ℳ.x1, ℳ.ψ, ℳ.η
    n, m, z, b, ℓ, α = length(s), length(Ω), length(ℳ.ℳ), ℳ.b, ℳ.ℓ, ℳ.α
    ∂U'∂ψ, ∂U'∂η = gradient(π, ℳ, P)
    UIndex(x, s) = (s - 1) * ℓ + (x - 1) + 1
    E(U, x1, b) = sum(b[s]*ℳ.Μ(x1, s)) for s in 1:n
    ψ' = Dict((x, a) => 0.0 for x in X, a in Ω)
    η' = Dict((x, a, o, x') => 0.0 for x in X, a in Ω, o in Ω, x' in X)
    for x in X
        ψ'x = [ψ[x, a] + α * E(∂U'∂ψ(x, a), x1, b) for a in Ω]
        ψ'x = project_to_simplex(ψ'x)
        for (aIndex, a) in enumerate(Ω)
            ψ'[x, a] = ψ'x[aIndex]
        end
        for (a, o) in product(Ω, Ω)
            η'x = [η[x, a, o, x'] + α * E(∂U'∂η(x, a, o, x'), x1, b) for x' in X]
            η'x = project_to_simplex(η'x)
            for (x'Index, x') in enumerate(X)
                η'[x, a, o, x'] = η'x[x'Index]
            end
        end
    end
    π.ψ, π.η = ψ', η'
end

function project_to_simplex(y)
    u = sort(copy(y), rev=true)
    i = maximum([j for j in eachindex(u)
        if u[j] + (1 - sum(u[1:j])/j) > 0.0])
    δ = (1 - sum(u[j] for j = 1:i))/i
    return [max(y[j] + δ, 0.0) for j in eachindex(u)]
end

Algorithm 23.6. An implementation of a controller gradient ascent algorithm for POMDP P at initial belief b. The controller itself has a fixed-size of ℓ nodes. It is improved over k_max iterations by following the gradient of the controller, with a step size of α, to maximally improve the value of the initial belief.
Unlike tree-based conditional plans, controllers can represent policies that can be executed indefinitely. They do not have to grow exponentially in size with the horizon.

23.6 Exercises

Exercise 23.1. List any advantages that a controller policy representation has over conditional plan and belief-based representations.

Solution: Unlike tree-based conditional plans, controllers can represent policies that can be executed indefinitely. They do not have to grow exponentially in size with the horizon.
Consider the catch problem (appendix F.9) with a uniform initial belief $b_1$. The figures below show the utility of the policy over gradient ascent iteration applied to the catch problem with $k = 3$ nodes. The left node is $x_1$.

At iteration 1, the policy is essentially random, both in action selection and successor selection.

At iteration 50, the agent has determined a reasonable distance to throw the ball (50) but still has not used its three nodes to remember anything useful.

At iteration 500, the policy has constructed a reasonable plan, given its fixed three nodes of memory. It first tries throwing the ball at a distance of 40. If the child catches the ball, then it increases the range to 50. It uses the final node to remember how many times the child caught the ball (up to twice) to choose the distance.

Example 23.6. An example of the controller gradient algorithm for controllers with a fixed size of $\ell = 3$. The policy is shown to refine itself over the algorithm’s iterations. The agent incrementally determines how to best use its fixed number of nodes, resulting in a reasonable and interpretable policy upon convergence.
Compared to belief-based representations, the number of parameters in a controller representation tends to be far less than the number of alpha vectors for larger problems. We can also more easily optimize controllers for a fixed amount of memory.

Controllers will never divide by zero the way that belief-based policies can during execution. Belief-based methods require maintaining a belief. The discrete state filter from equation (19.7) will divide by zero if an impossible observation is made. This can happen when a noisy observation from a sensor returns an observation that the models of \( T(s,a,s') \) and \( O(o \mid a,s') \) does not accurately capture.

**Exercise 23.2.** Controller policy iteration only adds nodes with deterministic action selection functions and successor distributions. Does this mean that the resulting controller is necessarily suboptimal?

**Solution:** Controller policy iteration is guaranteed to converge on an optimal policy in the limit. However, the method cannot find more compact representations of optimal controller policies that may require stochastic nodes.

**Exercise 23.3.** Prove that in policy iteration, any pruned node does not affect the utility.

**Solution:** Let \( x' \) be the new node from some iteration \( i \), and \( x \) be a previous node from iteration \( i - 1 \).

By construction, \( \eta(x',a,o,x) \) defines all new nodes \( x' \) to only have a successor \( x \) from the previous iteration. Thus for each state \( s \), \( U^{(i)}(x',s) \) only sums over the successors \( U^{(i-1)}(x,s') \) in equation (23.1). This means that the other utilities in iteration \( i \), including a self-loop to \( x \) itself, do not affect the utility \( U^{(i)}(x',s) \). Since the initial node is chosen by equation (23.3), we must ensure the utility with and without the pruned node at all beliefs is the same. A node is pruned in one of two ways.

First, \( x' \) obtains a higher utility over all states than its pruned successor \( x \). Formally, \( U^{(i)}(x,s) \leq U^{(i-1)}(x',s) \) for all \( s \). The prune step replaces \( x \) with \( x' \), including \( U \), \( \psi \), and \( \eta \). By construction, \( U \) has not decreased at any state \( s \).

Second, \( x \) is identical to an existing previous node \( x' \). Note that this means the transition \( \eta(x,a,o,x') = \eta(x',a,o,x') \). This means that the utility is identical except that \( x \) is reduced by \( \gamma \); in other words, \( \gamma U^{(i)}(x,s) = U^{(i-1)}(x,s) \) by equation (23.1). It does not affect the final utility to prune \( x \).

**Exercise 23.4.** Devise an algorithm that uses the nonlinear program algorithm to find the minimum fixed-sized controller required to obtain the optimality of a large fixed-sized controller of size \( \ell \). You can assume the nonlinear optimizer returns the optimal policy in this case.

**Solution:** The idea is to create an outer loop that increments the fixed-size of the controller, after knowing the utility of the large fixed-sized controller. First, we must compute the large fixed-sized controller’s utility \( U^* = \sum_s b_1(s)U(x_1,s) \) at initial node \( x_1 \) and initial belief \( b_1 \). Next, we create a loop that increments the size \( \ell \) of the controller. At each step, we
evaluate the policy and compute the utility $U^\ell$. By our assumption, the controller returned produces a globally optimal utility for the fixed size $\ell$. Once we arrive at a utility $U^\ell$, if we see that $U^\ell = U^*$, then we stop and return the policy.

**Exercise 23.5.** Analyze the controller gradient ascent algorithm’s gradient step. Assume that $|S|$ is larger than $|A|$ and $|O|$. What is the most computationally expensive part of the gradient step? How might this be improved?

**Solution:** Computing the inverse $Z^{-1} = (I - \gamma T_\theta)$ is the most computationally expensive part of the gradient step, as well as the entire gradient algorithm. The matrix $Z$ is of size $|X \times S|$. Gauss–Jordan elimination requires $O(|X \times S|^3)$ operations, though the 3 in the exponent can be reduced to 2.3728639 using a state-of-the-art matrix inversion algorithm.\(^\text{10}\) The creation of the temporary matrix $T_\theta$ also requires $O(|X \times S|^2 |A \times O|)$ operations to support computing the inverse. All other loops and other temporary array creations require far fewer operations. This can be improved using an approximate inverse technique.
PART V:

MULTIAGENT SYSTEMS

Up to this point, we have focused on decision making from the perspective of a single agent. We now extend the core concepts we have discussed so far to problems involving multiple agents. In multiagent systems, we can model other agents as potential allies or adversaries and adapt accordingly over time. These problems are inherently challenging due to complexities surrounding agent interactions and agents reasoning about other agents who reason about the agent, and so on. Chapter 24 introduces multiagent reasoning in games and outlines how to compute equilibria from simple interactions. Chapter 25 discusses how to design algorithms for multiple agents interacting over time, describing learning algorithms that favor rational adaptation over convergence to equilibria. Chapter 26 shows that state uncertainty significantly increases the complexity of the problem, emphasizing the distinct challenges in such domains. Chapter 27 focuses on the various models and algorithms for collaborative agents that strive to work together toward a common goal.
24 Multiagent Reasoning

So far, we have focused on making rational decisions for a single agent. These models have natural extensions to multiple agents. New challenges emerge as agents interact; agents can aid each other or act in their own best interests. Multiagent reasoning is a subject of game theory. This chapter builds upon the concepts introduced earlier, extending them to multiagent contexts. We will discuss the foundational game theoretic approaches to compute decision strategies and multiagent equilibria.

24.1 Simple Games

A simple game (algorithm 24.1) is a fundamental model for multiagent reasoning. Each agent \( i \in \mathcal{I} \) selects an action \( a^i \) to maximize their own accumulation of reward \( r^i \). The joint action space \( \mathcal{A} = \mathcal{A}^1 \times \cdots \times \mathcal{A}^k \) consists of all possible permutations of the actions \( \mathcal{A}^i \) available to each agent. The actions selected simultaneously across agents can be combined to form a joint action \( \mathbf{a} = (a^1, \ldots, a^k) \) from this joint action space. The joint reward function \( \mathbf{R}(\mathbf{a}) = (R^1(\mathbf{a}), \ldots, R^k(\mathbf{a})) \) represents the reward produced by the joint action \( \mathbf{a} \). The joint reward is written \( \mathbf{r} = (r^1, \ldots, r^k) \). Simple games do not include states or transition functions. Example 24.1 introduces a simple game.

```
struct SimpleGame
    γ  # discount factor
    I  # agents
    \mathcal{A}  # joint action space
    R  # joint reward function
end
```


3 A joint action is also called an action profile.
The prisoner’s dilemma is a two-agent, two-action simple game involving two prisoners that are on trial. They can choose to cooperate and remain silent about their shared crime, or defect and blame the other for their crime. If they both cooperate, they both serve time for one year. If agent $i$ cooperates and the other agent defects, then $i$ serves no time and the other serves four years. If both defect, then they both serve three years.

Two-agent simple games can be represented by a table. Rows represent actions for agent 1. Columns represent actions for agent 2. The rewards for agent 1 and 2 are shown in each cell.

<table>
<thead>
<tr>
<th></th>
<th>cooperate</th>
<th>defect</th>
</tr>
</thead>
<tbody>
<tr>
<td>cooperate</td>
<td>$-1, -1$</td>
<td>$-4, 0$</td>
</tr>
<tr>
<td>defect</td>
<td>$0, -4$</td>
<td>$-3, -3$</td>
</tr>
</tbody>
</table>

A joint policy $\pi$ specifies a probability distribution over joint actions taken by the agents. Joint policies can be decomposed into individual agent policies. The probability agent $i$ selects action $a$ is given by $\pi_i(a)$. In game theory, a deterministic policy is called a pure strategy and a stochastic policy is called a mixed strategy. The utility of a joint policy $\pi$ from the perspective of agent $i$ is

$$U^i(\pi) = \sum_{a \in A} R^i(a) \prod_{j \in I} \pi^j(a^j)$$

Algorithm 24.2 implements routines for representing policies and computing their utility.

A zero-sum game is a type of simple game where the sum of rewards across agents is zero. Here, any gain of an agent results at a loss to the other agents. A zero-sum game with two agents $I = \{1, 2\}$ has opposing reward functions $R^1(a) = -R^2(a)$. They are typically solved with algorithms specialized for this reward structure. Example 24.2 describes such a game.

### 24.2 Response Models

Before discussing different concepts for solving for a joint policy, we will begin by discussing how to model the response of a single agent $i$ given fixed policies of the other agents. We will use the notation $-i$ as shorthand for $(1, \ldots, i-1, i+1, \ldots, k)$. Using this notation, a joint action is written $a = (a^i, a^{-i})$, a joint reward is written $R(a^i, a^{-i})$, and a joint policy $\pi = (\pi^i, \pi^{-i})$. This section discusses different approaches for computing a response to a known $\pi^{-i}$.
Rock-paper-scissors is a zero-sum game for two agents. Each agent selects either rock, paper, or scissors. Rock wins against scissors, paper wins against rock, and scissors wins against paper, with a reward of 1 for the winner and −1 from the loser. If the agents select the same action, both receive 0 reward. Generally, two-agent repeated games can be represented as a sequence of payoff matrices, as shown below.

<table>
<thead>
<tr>
<th></th>
<th>rock</th>
<th>paper</th>
<th>scissors</th>
</tr>
</thead>
<tbody>
<tr>
<td>agent 1</td>
<td>0,0</td>
<td>−1,1</td>
<td>1,−1</td>
</tr>
<tr>
<td>agent 2</td>
<td>1,−1</td>
<td>0,0</td>
<td>−1,1</td>
</tr>
<tr>
<td></td>
<td>−1,1</td>
<td>1,−1</td>
<td>0,0</td>
</tr>
</tbody>
</table>

24.2.1 Best Response

A best response of agent $i$ to the policies of the other agents $\pi^{-i}$ is a policy $\pi^i$ that satisfies

$$U^i(\pi^i, \pi^{-i}) \geq U^i(\pi'^i, \pi^{-i})$$

(24.2)

for all other policies $\pi'^i \neq \pi^i$. In other words, a best response for an agent is a policy where there is no incentive for them to change their policy given a fixed set of policies of the other agents. There may be multiple such best responses.

If we restrict ourselves to deterministic policies, a deterministic best response to opponent policies $\pi^{-i}$ is straightforward to compute. We simply iterate over all of the actions of agent $i$ and return the one that maximizes utility:

$$\arg \max_{a^i \in A^i} U^i(a^i, \pi^{-i})$$

(24.3)

Algorithm 24.3 provides an implementation.
Algorithm 24.2. A policy associated with an agent is represented by a dictionary that maps actions to probabilities. There are different ways to construct a policy. One way is to pass in a dictionary directory, in which case the probabilities are normalized. Another way is to pass in a generator that creates this dictionary. We can also construct a policy by passing in an action, in which case it assigns probability 1 to that action. If we have an individual policy $\pi_i$, we can call $\pi_i(ai)$ to compute the probability the policy associates with action $ai$. If we call $\pi_i()$, then it will return a random action according to that policy.

We can use $\text{joint}(\mathcal{A})$ to construct the joint action space from $\mathcal{A}$. We can use $\text{utility}(\mathcal{P}, \pi, i)$ to compute the utility associated with executing joint policy $\pi$ in the game $\mathcal{P}$ from the perspective of agent $i$.

Algorithm 24.3. For a simple game $\mathcal{P}$, we can compute a deterministic best response for agent $i$ given that the other agents are playing policies in $\pi$. 

```plaintext
struct SimpleGamePolicy
    p # dictionary mapping actions to probabilities

    function SimpleGamePolicy(p::Base.Generator)
        return SimpleGamePolicy(Dict(p))
    end

    function SimpleGamePolicy(p::Dict)
        vs = collect(values(p))
        vs /= sum(vs)
        return new(Dict(k => v for (k, v) in zip(keys(p), vs)))
    end

    SimpleGamePolicy(ai) = new(Dict(ai => 1.0))
end

(\pi_i::SimpleGamePolicy)(ai) = get(\pi_i, ai, 0.0)

function (\pi_i::SimpleGamePolicy)()
    D = SetCategorical(collect(keys(\pi_i.p)), collect(values(\pi_i.p)))
    return rand(D)
end

joint(X) = vec(collect(product(X...)))

joint(\pi, \pi_i, i) = [i == j ? \pi_i : \pi_j for (j, \pi_j) in enumerate(\pi)]

function utility(\mathcal{P}::SimpleGame, \pi, i)
    \mathcal{A}, R = \mathcal{P}.\mathcal{A}, \mathcal{P}.R
    p(a) = prod(\pi_j(a_j) for (j, a_j) in zip(\pi, a))
    return sum(R[a][i]*p(a) for a in joint(\mathcal{A}))
end

function best_response(\mathcal{P}::SimpleGame, \pi, i)
    U(ai) = utility(\mathcal{P}, joint(\pi, SimpleGamePolicy(ai), i), i)
    ai = argmax(U, \mathcal{P}.\mathcal{A}[i])
    return SimpleGamePolicy(ai)
end
```
24.2.2 Softmax Response

We can use a softmax response to model how agent $i$ will select their action. As discussed in section 6.7, humans are often not perfectly rational optimizers of expected utility. The principle underlying the softmax response model is that (typically human) agents are more likely to make errors in their optimization when those errors are less costly. Given a precision parameter $\lambda \geq 0$, this model selects action $a^i$ according to

$$\pi^i(a^i) \propto \exp(\lambda U^i(a^i, \pi^{-i})) \quad (24.4)$$

As $\lambda \to 0$, the agent is insensitive to differences in utility, and they select actions uniformly at random. As $\lambda \to \infty$, the policy converges to a deterministic best response. We can treat $\lambda$ as a parameter that can be learned from data using, for example, maximum likelihood estimation (section 4.1). This learning-based approach aims to be predictive of behavior rather than prescriptive of behavior, though having a predictive model of other human agents can be useful in building a system that prescribes optimal behavior. Algorithm 24.4 provides an implementation of softmax response.

```
function softmax_response(\mathcal{P}::SimpleGame, \pi, i, \lambda)
    Ai = \mathcal{P}.A[i]
    U(ai) = utility(\mathcal{P}, joint(\pi, SimpleGamePolicy(ai), i), i)
    return SimpleGamePolicy(ai \Rightarrow \exp(\lambda \times U(ai)) \text{ for } ai \text{ in } Ai)
end
```

Algorithm 24.4. For a simple game $\mathcal{P}$ and a particular agent $i$, we can compute the softmax response policy $\pi_i$ given the other agents are playing policies in $\pi$. This computation requires specifying the precision parameter $\lambda$.

24.3 Nash Equilibrium

Computing a best response for an agent requires that we hold the policies of all other agents constant. When we do not constrain the policies of the other agents, the meaning of an optimal joint policy becomes less clear. In the context of games, there are several notions for optimality or solution concepts, but the Nash equilibrium is perhaps the most widely known. A Nash equilibrium is a joint policy $\pi$ in which all agents are following a best response. In other words, a Nash equilibrium is a joint policy in which no agent has an incentive to unilaterally switch their policy.

All games with a finite action space have at least one Nash equilibrium.\footnote{Exercise 24.1 explores the case where the action space is infinite.} Multiple Nash equilibria can exist in a single game (exercise 24.2). Sometimes Nash equilibria may involve deterministic policies, but this is not always the case (see example 24.3). Computing a Nash equilibrium is PPAD-complete, a class that is distinct from NP-complete (appendix C.2) but also has no known polynomial time algorithm.\footnote{C. Daskalakis, P.W. Goldberg, and C.H. Papadimitriou, “The Complexity of Computing a Nash Equilibrium,” Communications of the ACM, vol. 52, no. 2, pp. 89–97, 2009.}

Suppose we wish to find a Nash equilibrium for the prisoner’s dilemma from example 24.1. If both agents always defect, both receive \(-3\) reward. Any deviation by any agent will result in a \(-4\) reward for that agent; hence, there is no incentive to deviate. Having both agents defect is thus a Nash equilibrium for the prisoner’s dilemma.

Suppose we now wish to find a Nash equilibrium for rock-paper-scissors from example 24.2. Any deterministic strategy by one agent can be easily countered by the other agent. For example, if agent 1 plays rock then agent 2’s best response is paper. Because there is no deterministic Nash equilibrium for rock-paper-scissors, we know there must be one involving stochastic policies. Suppose each agent selects from the actions uniformly at random. This solution produces an expected utility of 0 for both agents:

\[
U^i(\pi) = 0 \cdot \frac{1}{3} \cdot \frac{1}{3} - \frac{1}{3} \cdot \frac{1}{3} + \frac{1}{3} \cdot \frac{1}{3} \\
+ \frac{1}{3} \cdot \frac{1}{3} + 0 \cdot \frac{1}{3} - \frac{1}{3} \cdot \frac{1}{3} \\
- \frac{1}{3} \cdot \frac{1}{3} + \frac{1}{3} \cdot \frac{1}{3} + 0 \cdot \frac{1}{3} \\
= 0
\]

Any deviation by an agent would decrease their expected payoff, meaning that we have found a Nash equilibrium.
The problem of finding a Nash equilibrium can be framed as an optimization problem:

\[
\min_{\pi, U} \sum_i \left( U^i - U^i(\pi) \right)
\]

subject to \( U^i \geq U^i(a^i, \pi^{-i}) \) for all \( i, a^i \)

\[
\sum_{a^i} \pi^i(a^i) = 1 \text{ for all } i
\]

\[
\pi^i(a^i) \geq 0 \text{ for all } i, a^i
\]

(24.5)

The optimization variables correspond to the parameters of \( \pi \) and \( U \). At convergence, the objective will be 0 with \( U^i \) matching the utilities associated with policy \( \pi \) as computed in equation (24.1) for each agent \( i \). The first constraint ensures that no agent will do better by unilaterally changing their action. Like the objective, this first constraint in nonlinear because it involves a product of the parameters in the optimization variable \( \pi \). The last two constraints are linear, ensuring that \( \pi \) represents a valid set of probability distributions over actions.

```
struct NashEquilibrium end

function tensorform(𝒫::SimpleGame)
    ℐ, ℋ, R = 𝒫.ℐ, 𝒫.ℋ, 𝒫.R
    ℐ' = eachindex(ℐ)
    ℋ' = [eachindex(ℋ[i]) for i in ℐ]
    R' = [R(a) for a in joint(ℋ)]
    return ℐ', ℋ', R'
end

function solve(M::NashEquilibrium, 𝒫::SimpleGame)
    ℐ, ℋ, R = tensorform(𝒫)
    model = Model(Ipopt.Optimizer)
    @variable(model, U[ℐ])
    @variable(model, π[i=ℐ, ℋ[i]] ≥ 0)
    @NLobjective(model, Min,
                  sum(U[i] - sum(prod(π[j,a[j]] for j in ℐ) * R[y][i]
                        for (y,a) in enumerate(joint(ℋ))) for i in ℐ))
    @NLconstraint(model, [i=ℐ, ai=ℋ[i]],
                   U[i] ≥ sum(
                        prod(j==i ? (a[j]==ai ? 1.0 : 0.0) : π[j,a[j]] for j in ℐ) * R[y][i] for (y,a) in enumerate(joint(ℋ))))
    @constraint(model, [i=ℐ, sum(π[i,ai] for ai in ℋ[i]) == 1])
    optimize!(model)
    ⋆π'(i) = SimpleGamePolicy(𝒫.ℋ[i][ai] ⇒ value(π[i,ai]) for ai in ℋ[i])
    return [π'(i) for i in ℐ]
end
```

Algorithm 24.5. This nonlinear program computes a Nash equilibrium for a simple game \( 𝒫 \).
24.4 Correlated Equilibrium

The correlated equilibrium generalizes the Nash equilibrium concept by relaxing the assumption that the agents act independently. The joint action in this case comes from a full joint distribution. A correlated joint policy $\pi(a)$ is a single distribution over the joint actions of all agents. Consequently, the actions of the various agents may be correlated, preventing the policies from being decoupled into individual policies $\pi^i(a^i)$. Algorithm 24.6 shows how to represent such a policy.

A correlated equilibrium is a correlated joint policy where no agent $i$ can increase their expected utility by deviating from their current action $a^i$ to another action $a^{i'}$:

$$\sum_{a^{-i}} R^i(a^i, a^{-i}) \pi(a^i, a^{-i}) \geq \sum_{a^{-i}} R^i(a^{i'}, a^{-i}) \pi(a^{i'}, a^{-i})$$

(24.6)

Example 24.4 demonstrates this concept.

Every Nash equilibrium is a correlated equilibrium because we can always form a joint policy from independent policies:

$$\pi(a) = \prod_{i=1}^{k} \pi^i(a^i)$$

(24.7)

If the individual policies satisfy equation (24.2), then the joint policy will satisfy equation (24.6). Not all correlated equilibria, however, are Nash equilibria.

A correlated equilibrium can be computed using linear programming (algorithm 24.7):
Consider again rock-paper-scissors from example 24.2. In example 24.3, we found that a Nash equilibrium is for both agents select their actions uniformly at random. In correlated equilibria, we use a correlated joint policy \( \pi(a) \), meaning we need to find a distribution over (rock, rock), (rock, paper), (rock, scissors), (paper, rock), and so on. There are nine possible joint actions.

First, consider the joint policy in which agent 1 selects rock and agent 2 selects scissors. The utilities are

\[
U^1(\pi) = 0 \cdot \frac{0}{9} - 1 \cdot \frac{0}{9} + 1 \cdot \frac{9}{9} + 1 \cdot \frac{0}{9} + \cdots = 1
\]

\[
U^2(\pi) = 0 \cdot \frac{0}{9} + 1 \cdot \frac{0}{9} - 1 \cdot \frac{9}{9} - 1 \cdot \frac{0}{9} + \cdots = -1
\]

If agent 2 switched its action to paper, they would receive a utility of 1. Hence, this is not a correlated equilibrium.

Consider instead a correlated joint policy in which the joint action was chosen uniformly at random, with \( \pi(a) = 1/9 \).

\[
U^1(\pi) = 0 \cdot \frac{1}{9} - 1 \cdot \frac{1}{9} + 1 \cdot \frac{1}{9} + 1 \cdot \frac{1}{9} + \cdots = 0
\]

\[
U^2(\pi) = 0 \cdot \frac{1}{9} + 1 \cdot \frac{1}{9} - 1 \cdot \frac{1}{9} - 1 \cdot \frac{1}{9} + \cdots = 0
\]

Any deviation from this results in one agent gaining utility and the other losing utility. This is a correlated equilibrium for rock-paper-scissors.
maximize  \[ \sum_i \sum_a R^i(a) \pi(a) \] 
subject to  \[ \sum_a R^i(a^i, a^{-i}) \pi(a^i, a^{-i}) \geq \sum_a R^i(a'^i, a^{-i}) \pi(a'^i, a^{-i}) \] for all \( a, a^i, a'^i \) 
\[ \sum_a \pi(a) = 1 \] 
\[ \pi(a) \geq 0 \] for all \( a \)

Although linear programs can be solved in polynomial time, the size of the joint action space grows exponentially with the number of agents. The constraints enforce a correlated equilibrium. The objective, however, can be used to select between different valid correlated equilibria. Table 24.1 provides several common choices for the objective function. Not all of these are linear.

<table>
<thead>
<tr>
<th>Name</th>
<th>Description</th>
<th>Objective Function</th>
</tr>
</thead>
<tbody>
<tr>
<td>Utilitarian</td>
<td>Maximize the net utility.</td>
<td>maximize_\pi \sum_i \sum_a R^i(a) \pi(a)</td>
</tr>
<tr>
<td>Egalitarian</td>
<td>Maximize the minimum of all agents’ utilities.</td>
<td>maximize_\pi \min \sum_a R^i(a) \pi(a)</td>
</tr>
<tr>
<td>Plutocratic</td>
<td>Maximize the maximum of all agents’ utilities.</td>
<td>maximize_\pi \max \sum_a R^i(a) \pi(a)</td>
</tr>
<tr>
<td>Dictatorial</td>
<td>Maximize agent ( i )'s utility.</td>
<td>maximize_\pi \sum_a R^i(a) \pi(a)</td>
</tr>
</tbody>
</table>

Algorithm 24.7. Correlated equilibria are a more general notion of optimality for a simple game \( \mathcal{P} \) than a Nash equilibrium. They can be computed using a linear program. The resulting policies are correlated, meaning that the agents stochastically select their joint actions.

Table 24.1. Alternative objective functions for equation (24.8), which select for different correlated equilibria. These descriptions were adapted from A. Greenwald and K. Hall, “Correlated Q-Learning,” in International Conference on Machine Learning (ICML), 2003.
24.5 Iterated Best Response

Because computing a Nash equilibrium can be computationally expensive, an alternative approach is to iteratively apply best responses in a series of repeated games. In iterated best response (algorithm 24.8), we randomly cycle between agents, solving for each agent’s best response policy in turn. This process may converge to a Nash equilibrium, but there are guarantees only for certain classes of games.\(^8\) In many problems, it is common to observe cycles.

Algorithm 24.8. Iterated best response involves cycling through the agents and applying their best response to the other agents. The algorithm starts with some initial policy and stops after \(k_{\text{max}}\) iterations. For convenience, we have a constructor that takes as input a simple game and creates an initial policy that has each agent select actions uniformly at random. The same solve function will be reused in the next chapter in the context of more complicated forms of games.

24.6 Hierarchical Softmax

An area known as behavioral game theory aims to model human agents. When building decision-making systems that must interact with humans, computing the Nash equilibrium is not always helpful. Humans often do not play a Nash equilibrium strategy. First of all, it may be unclear which equilibrium to adopt if there are many different equilibria in the game. For games with only one equilibrium, it may be difficult for a human to compute the Nash equilibrium because of cognitive limitations. Even if human agents can compute the Nash equilibrium, they may doubt that their opponents can perform that computation.
There are many different behavioral models in the literature, but one approach is to combine the iterated approach from the previous section with a softmax model. This hierarchical softmax approach (algorithm 24.9) models the depth of rationality of an agent by a level $k \geq 0$. A level-0 agent plays its actions uniformly at random. A level-1 agent assumes the other players adopt level-0 strategies and selects actions according to a softmax response with precision $\lambda$. A level-$k$ agent selects actions according to a softmax model of the other players playing level-$(k-1)$. Figure 24.1 illustrates this approach on a simple game.

Algorithm 24.9. The hierarchical softmax model with precision parameter $\lambda$ and level $k$. By default, it starts with an initial joint policy that assigns uniform probability to all individual actions.

We can learn the $k$ and $\lambda$ parameters of this behavioral model from data. If we have a collection of joint actions played by different agents, we can compute the associated likelihood for a given $k$ and $\lambda$. We can then use an optimization algorithm to attempt to find a $k$ and $\lambda$ that maximizes likelihood. This optimization typically cannot be done analytically, but we can use numerical methods to perform this optimization. Alternatively, we can use a Bayesian approach to parameter learning.

24.7 Fictitious Play

An alternative approach for computing policies for different agents is to have them play each other in simulation and learn how to best respond. Algorithm 24.10 provides an implementation of the simulation loop. At each iteration, we evaluate

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Figure 24.1. The hierarchical softmax model applied to the traveler’s dilemma (appendix F.12) for different depths of rationality $k$ and precision parameters $\lambda$. People tend to select actions between $97$ and $100$, even though the Nash equilibrium is only $2$. 
the various policies to obtain a joint action, and then this joint action is used to by the agents to update their policies. We can use different ways to update the policies in response to observed joint actions. This section focuses on fictitious play, where the agents use maximum likelihood estimates (section 16.1) of the policies followed by the other agents. Each agent follows its own best response assuming the other agents act according to those estimates.13

<table>
<thead>
<tr>
<th>function simulate((\mathcal{P}::\text{SimpleGame}, \pi, k_{\text{max}}))</th>
</tr>
</thead>
<tbody>
<tr>
<td>for (k = 1:k_{\text{max}})</td>
</tr>
<tr>
<td>(a = [\pi_i() \text{ for } \pi_i \text{ in } \pi])</td>
</tr>
<tr>
<td>for (\pi_i \text{ in } \pi)</td>
</tr>
<tr>
<td>update!((\pi_i, a))</td>
</tr>
<tr>
<td>end</td>
</tr>
<tr>
<td>end</td>
</tr>
<tr>
<td>return (\pi)</td>
</tr>
</tbody>
</table>

To compute a maximum likelihood estimate, agent \(i\) tracks the number of times agent \(j\) takes action \(a^j\), storing it in a table \(N^i(j, a^j)\). These counts can be initialized to any value, but they are often initialized to 1 to create initial uniform uncertainty. Agent \(i\) computes their best response assuming that each agent \(j\) follows the stochastic policy:

\[
\pi^i(a^j) \propto N^i(j, a^j)
\]  

(24.9)

At each iteration, we have each agent act according to a best response assuming these stochastic count-based policies for the other agents. We then update the action counts for the actions taken. Algorithm 24.11 implements this simple adaptive procedure. Figure 24.2 and figure 24.3 show how agents’ policies evolve over time using fictitious play. Fictitious play is not guaranteed to converge to a Nash equilibrium.14

There are many variants of fictitious play. One variant, called smooth fictitious play,15 selects a best response using expected utility plus a smoothing function such as the entropy of the policy. Another variant is called rational learning or Bayesian learning. Rational learning expands the model of fictitious play to be any belief over other agent’s actions, formulated as a Bayesian prior. Bayes’ rule is then used to update the beliefs given the history of joint actions. Traditional fictitious play can be seen as rational learning with a Dirichlet prior (section 4.2.2).


mutable struct FictitiousPlay
    ℋ # simple game
    i # agent index
    N # array of action count dictionaries
    πi # current policy
end

function FictitiousPlay(ℋ::SimpleGame, i)
    N = [Dict(aj => 1 for aj in ℋ.𝒜[j]) for j in ℋ.ℐ]
    πi = SimpleGamePolicy(ai => 1.0 for ai in ℋ.𝒜[i])
    return FictitiousPlay(ℋ, i, N, πi)
end

(πi::FictitiousPlay)() = πi.πi()

(πi::FictitiousPlay)(ai) = πi.πi(ai)

function update!(πi::FictitiousPlay, a)
    N, ℋ, ℐ, i = πi.N, πi.ℋ, πi.ℐ, πi.i
    for (j, aj) in enumerate(a)
        N[j][aj] += 1
    end
    p(j) = SimpleGamePolicy(aj => u/sum(values(N[j])) for (aj, u) in N[j])
    π = [p(j) for j in ℐ]
    πi.πi = best_response(ℋ, π, i)
end

Algorithm 24.11. Fictitious play is a simple learning algorithm for an agent i of a simple game ℋ that maintains counts of other agent action selections over time and averages them assuming this is their stochastic policy. It then computes a best response to this policy and performs the corresponding utility-maximizing action.
Figure 24.2. Two fictitious play agents learning and adapting to one another in a prisoner’s dilemma game. The first row illustrates agent 1’s learned model of 2 (left) and agent 1’s policy (right) over iteration. The second row follows the same pattern but for agent 2. To illustrate variation in learning behavior, the initial counts for each agent’s model over the other agent’s action were assigned to a random number between 1 and 10.

Figure 24.3. A visualization of two fictitious play agents learning and adapting to one another in a rock-paper-scissors game. The first row illustrates agent 1’s learned model of 2 (left) and agent 1’s policy (right) over time. The second row follows the same pattern but for agent 2. To illustrate variation in learning behavior, the initial counts for each agent’s model over the other agent’s action were assigned to a random number between 1 and 10. In this zero-sum game, fictitious play agents approach convergence to their stochastic policy Nash equilibrium.
24.8 Gradient Ascent

Gradient ascent (algorithm 24.12) incrementally adjusts the agent’s policy in the gradient with respect to its utility. At time $t$, the gradient for agent $i$ is

$$\frac{\partial U^i(\pi_i)}{\partial \pi_i(a^i)} = \frac{\partial}{\partial \pi_i(a^i)} \left( \sum_a R^i(a) \prod_j \pi^j(a^j) \right) = \sum_{a \neq i} R^i(a^i, a^{-i}) \prod_{j \neq i} \pi^j(a^j)$$

(24.10)

We can then use standard gradient ascent with

$$\pi_{t+1}^i(a^i) = \pi_t^i(a^i) + \alpha_t^i \frac{\partial U^i(\pi_t)}{\partial \pi_i(a^i)}$$

(24.11)

with learning rate $\alpha_t^i$. This $\pi_{t+1}^i$ may need to be projected back to a valid probability distribution, just as we did in section 23.4 for POMDP policies.

In practice, however, an agent $i$ knows only their own policy $\pi^i_t$, not the policies of the others, making the computation of the gradient difficult. Agents do observe the joint actions $a_t$ that are performed. Although we could try to estimate their policies as done in fictitious play, one simple approach is to assume the policy of the other agents is to replay their most recent action. The gradient then simplifies to

$$\frac{\partial U^i(\pi_t)}{\partial \pi^i_t(a^i)} = R^i(a^i, a^{-i})$$

(24.12)

Figure 24.4 demonstrates this approach on a simple game.

24.9 Summary

- In simple games, multiple agents compete to maximize expected reward.
- Optimality is not as straightforward in the multiagent setting, with multiple possible solution concepts for extracting policies from a reward specification.
- A best response of an agent to a fixed set of policies of the other agents is one where there is no incentive to deviate.
- A Nash equilibrium is a joint policy where all agents follow a best response.
- A correlated equilibrium is the same as a Nash equilibrium, except all agents follow a single joint action distribution that allows for correlation between agents.

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mutable struct GradientAscent
    ℙ # simple game
    i # agent index
    t # time step
    πi # current policy
end

function GradientAscent(ℙ::SimpleGame, i)
    uniform() = SimpleGamePolicy(ai => 1.0 for ai in ℙ.𝒜[i])
    return GradientAscent(ℙ, i, 1, uniform())
end

(πi::GradientAscent)() = πi.πi()

(πi::GradientAscent)(ai) = πi.πi(ai)

function update!(πi::GradientAscent, a)
    ℙ, ℐ, Ai, i, t = πi.ℙ, πi.ℐ, πi.ℙ.𝒜[πi.i], πi.i, πi.t
    jointπ(ai) = [SimpleGamePolicy(j == i ? ai : a[j]) for j in ℙ]
    r = [utility(ℙ, jointπ(ai), i) for ai in Ai]
    π′ = [πi.πi(ai) for ai in Ai]
    π = project_to_simplex(π′ + r / sqrt(t))
    πi.t = t + 1
    πi.πi = SimpleGamePolicy(ai => p for (ai, p) in zip(Ai, π))
end

Algorithm 24.12. An implementation of gradient ascent for an agent i of a simple game ℙ. The algorithm updates its distribution over actions incrementally following gradient ascent to improve the expected utility. The projection function from algorithm 23.6 is used to ensure that the resulting policy remains a valid probability distribution.

Figure 24.4. Two gradient ascent agents with randomly initialized policies in a rock-paper-scissors game. We use a variation of algorithm 24.12 with a learning rate of 0.1/√t. Shown are 20 policy updates. Although different simulation traces will converge because the step size goes to 0, different samples from the stochastic policies may result in convergence to different policies.

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2021-03-07 23:12:00-08:00, revision 80bf4d6, comments to bugs@algorithmsbook.com
• Iterated best response can quickly optimize a joint policy by iteratively applying best responses, but there are no general guarantees of convergence.

• Hierarchical softmax attempts to model agents in terms of their depth of rationality and precision, which can be learned from past joint actions.

• Fictitious play is a learning algorithm that uses maximum-likelihood action models for other agents to find best response policies, with the potential to converge to a Nash equilibrium.

• Gradient ascent followed by projection onto the probability simplex can be used to learn policies.

24.10 Exercises

Exercise 24.1. Give an example of a game with two agents and an infinite number of actions such that a Nash equilibrium does not exist.

Solution: Suppose the action space of each agent consists of the negative real numbers and their reward is equal to their action. Since no greatest negative number exists, a Nash equilibrium cannot exist.

Exercise 24.2. Give an example of a game with two agents, two actions, and two Nash equilibria involving deterministic policies.

Solution: Here is one example. Suppose we have two aircraft on a collision course, and the pilots of each aircraft must choose between climb or descend to avoid collision. If the pilots both choose the same maneuver, then there is a crash with utility $-4$ to both pilots. Because climbing requires more fuel than descending, there is an additional penalty of $-1$ to any pilot who decides to climb.

Exercise 24.3. Given a stationary joint policy $\pi$ that is a Nash equilibrium for a simple game with horizon of 1, prove that it is also a Nash equilibrium for the same simple game repeated to any finite or infinite horizon.

Solution: By definition of the Nash equilibrium in equation (24.2), all agents $i$ are performing a best response $\pi^i$:

$$U^i(\pi^i, \pi^{-i}) \geq U^i(\pi^{i'}, \pi^{-i'})$$

to all other policies $\pi^{i'} \neq \pi^i$. By definition of $U^i$, we have

$$U^i(\pi) = \sum_{a \in A} R^i(a) \prod_{j=1}^{k} \pi^j(a^j)$$

The joint policy remains constant over time for all agents. Apply any horizon $n$, with any discount factor ($\gamma = 1$ for $n < \infty$; $\gamma < 1$ for $n \to \infty$). The utility of agent $i$ after $n$ steps is

$$U^{i,n}(\pi) = \sum_{t=1}^{n} \gamma^{t-1} \sum_{a \in A} R^i(a) \prod_{j=1}^{k} \pi^j(a^j)$$

$$= \sum_{a \in A} R^i(a) \prod_{j=1}^{k} \pi^j(a^j) \sum_{t=1}^{n} \gamma^{t-1}$$

$$= U^i(\pi) \sum_{t=1}^{n} \gamma^{t-1}$$

$$= U^i(\pi) c$$

The discount factor becomes a constant multiplier $c > 0$. Therefore, any constant multiplication of equation (24.2) on both sides results in the same inequality, completing the proof:

$$U^i(\pi^i, \pi^{-i}) \geq U^i(\pi^{i'}, \pi^{-i'})$$

$$U^i(\pi^i, \pi^{-i}) c \geq U^i(\pi^{i'}, \pi^{-i'}) c$$

$$U^i(\pi^i, \pi^{-i}) \sum_{t=1}^{n} \gamma^{t-1} \geq U^i(\pi^{i'}, \pi^{-i'}) \sum_{t=1}^{n} \gamma^{t-1}$$

$$\sum_{t=1}^{n} \gamma^{t-1} U^i(\pi^i, \pi^{-i}) \geq \sum_{t=1}^{n} \gamma^{t-1} U^i(\pi^{i'}, \pi^{-i'})$$

$$U^{i,n}(\pi^i, \pi^{-i}) \geq U^{i,n}(\pi^{i'}, \pi^{-i'})$$

Exercise 24.4. Prove that a Nash equilibrium is a correlated equilibrium.

Solution: Given any Nash equilibrium $\pi$, define a correlated joint policy $\pi(a)$ such that

$$\pi(a) = \prod_{i=1}^{k} \pi^i(a^i)$$
By definition of a Nash equilibrium, for any agent $i$ and any other policy $\pi^i \neq \pi_i$, we have a best response. Applying the definition of best response from equation (24.2), utility from equation (24.1), and this correlated joint policy $\pi(a)$, we obtain:

$$U^i(\pi^i, \pi^{-i}) \geq U^i(\pi'^i, \pi^{-i})$$

$$\sum_a R^i(a) \pi^i(a) \prod_{j \neq i} \pi^j(a^j) \geq \sum_a R^i(a) \pi'^i(a) \prod_{j \neq i} \pi'^j(a^j)$$

Consider each element in the summation over $a^i$, and consider any action with positive likelihood $\pi^i(a^i) > 0$ and divide each side by it. The left side is 1. The right side is either $< 1$ or $\pi^i(a^i) < \pi'^i(a^i)$, in which case the inequality can be preserved by simply $\pi^i(a^i) \geq 1$. The result is that the term falls out.

Consider the following equation, for any deterministically chosen action $a^i$ such that $\pi^i(a^i) = 1$ and any other action $a'^i$:

$$\sum_a R^i(a^i, a^{-i}) \pi^{-i}(a^{-i}) \geq \sum_a R^i(a'^i, a^{-i}) \pi^{-i}(a^{-i})$$

$$\pi^i(a^i) \sum_a R^i(a^i, a^{-i}) \pi^{-i}(a^{-i}) \geq \pi^i(a'^i) \sum_a R^i(a'^i, a^{-i}) \pi^{-i}(a^{-i})$$

$$\sum_{a^{-i}} R^i(a^i, a^{-i}) \pi(a^{-i}) \geq \sum_{a^{-i}} R^i(a'^i, a^{-i}) \pi(a^{-i})$$

Any mixing of these equations over actions trivially holds true. This satisfies equation (24.6).

**Exercise 24.5.** Give an example two-agent game, each with two actions, for which the correlated equilibria cannot be represented as a Nash equilibrium.

**Solution:** Consider the following game in which two people want to go on a date but have a conflicting preference on what kind of date, in this case a dinner or a movie.

<table>
<thead>
<tr>
<th></th>
<th>dinner</th>
<th>movie</th>
</tr>
</thead>
<tbody>
<tr>
<td>dinner</td>
<td>2, 1</td>
<td>0, 0</td>
</tr>
<tr>
<td>movie</td>
<td>0, 0</td>
<td>1, 2</td>
</tr>
</tbody>
</table>
A Nash equilibrium here is to simply uniformly at random choose dinner or movie,
\[ \pi^i(\text{dinner}) = \pi^i(\text{movie}) = 0.5 \] for both agents \( i \). This results in an expected utilities of \( \frac{2}{3} \):

\[
U^1(\pi) = 0.5 \cdot 0.5 \cdot 2 + 0.5 \cdot 0.5 \cdot 0 + 0.5 \cdot 0.5 \cdot 0 + 0.5 \cdot 0.5 \cdot 1 = 0.25 \cdot 2 + 0.25 \cdot 1 = \frac{2}{3} \\
U^2(\pi) = 0.5 \cdot 0.5 \cdot 1 + 0.5 \cdot 0.5 \cdot 0 + 0.5 \cdot 0.5 \cdot 0 + 0.5 \cdot 0.5 \cdot 2 = 0.25 \cdot 1 + 0.25 \cdot 2 = \frac{2}{3}
\]

This is of course a correlated equilibrium as well.

However, if the two agents correlated their actions on a fair coin flip \( \pi(\text{movie, movie}) = \pi(\text{dinner, dinner}) = 0.5 \), then they could coordinate either both going to the dinner or both going to the movies.

\[
U^1(\pi) = 0.5 \cdot 2 + 0.0 \cdot 0 + 0.0 \cdot 0 + 0.5 \cdot 1 = 0.5 \cdot 2 + 0.5 \cdot 1 = \frac{3}{2} \\
U^2(\pi) = 0.5 \cdot 1 + 0.5 \cdot 0 + 0.5 \cdot 0 + 0.5 \cdot 2 = 0.5 \cdot 1 + 0.5 \cdot 2 = \frac{3}{2}
\]

This is not possible with a Nash equilibrium. Intuitively, in this example, this is because the probabilistic weight is spread out over each row, independently for each player. Conversely, a correlated equilibrium can be targeted toward a specific cell, in this case able to obtain a higher payoff.

**Exercise 24.6.** Algorithms such as iterated best response and fictitious play do not converge in every game. Construct a game that demonstrates this non-convergence.

*Solution:* Iterated best response diverges in rock-paper-scissors. Below is an example of the first 10 iterations with a random initialization:

<table>
<thead>
<tr>
<th>Iteration</th>
<th>Agent 1’s Action</th>
<th>Agent 2’s Action</th>
<th>Rewards</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>scissors</td>
<td>rock</td>
<td>-1.0, 1.0</td>
</tr>
<tr>
<td>2</td>
<td>paper</td>
<td>rock</td>
<td>1.0, -1.0</td>
</tr>
<tr>
<td>3</td>
<td>paper</td>
<td>scissors</td>
<td>-1.0, 1.0</td>
</tr>
<tr>
<td>4</td>
<td>rock</td>
<td>scissors</td>
<td>1.0, -1.0</td>
</tr>
<tr>
<td>5</td>
<td>rock</td>
<td>paper</td>
<td>-1.0, 1.0</td>
</tr>
<tr>
<td>6</td>
<td>scissors</td>
<td>paper</td>
<td>1.0, -1.0</td>
</tr>
<tr>
<td>7</td>
<td>scissors</td>
<td>rock</td>
<td>-1.0, 1.0</td>
</tr>
<tr>
<td>8</td>
<td>paper</td>
<td>rock</td>
<td>1.0, -1.0</td>
</tr>
<tr>
<td>9</td>
<td>paper</td>
<td>scissors</td>
<td>-1.0, 1.0</td>
</tr>
<tr>
<td>10</td>
<td>rock</td>
<td>scissors</td>
<td>1.0, -1.0</td>
</tr>
</tbody>
</table>

Fictitious play also will not converge in almost-rock-paper-scissors:19

---

Here is an example of fictitious play agents playing this game for 60 iterations:

Exercise 24.7. What does iterated best response converge to in the traveler’s dilemma (appendix F.12)?

Solution: It converges to the Nash equilibrium of $2.$
25  Sequential Problems

This chapter extends simple games to a sequential context with multiple states. A Markov game (MG) can be viewed as a Markov decision process involving multiple agents with their own reward functions. In this formulation, transitions depend on the joint action and all agents seek to maximize their own reward. We generalize the response models and the Nash equilibrium solution concept from simple games to take into account the state transition model. The last part of this chapter discusses learning-based models where the agents adapt their policies based on information from observed interactions and knowledge of the reward and transition functions.

25.1  Markov Games

An MG (algorithm 25.1) is an extension of simple games to include a shared state \( s \in S \). The likelihood of transitioning from a state \( s \) to a state \( s' \) under a joint action \( a \) is given by the transition distribution \( T(s' | s, a) \). Each agent \( i \) receives a reward according to its own reward function \( R^i(s, a) \), which now additionally depends on the state. Example 25.1 sketches how traffic routing can be framed as a Markov game.

```plaintext
struct MG
    γ    # discount factor
    ℐ    # agents
    ℳ    # state space
    ℳ️    # joint action space
    T    # transition function
    R    # joint reward function
end
```

Example 25.1. Data structure for a Markov game.
Consider commuters headed to work by car. Each car has a starting position and a destination. Each car can take any of several available roads to get to their destination, but these roads vary in the time it takes to drive them. The more cars that drive on a given road, the slower they all move.

This problem is a Markov game. The agents are the commuters in their cars, the states are the locations of all the cars on the roads, and the actions correspond to decisions on which road to take next. The state transition moves all car agents forward following their joint action. The negative reward is proportional to the time spent driving on a road.

The joint policy $\pi$ in a Markov game specifies a probability distribution over joint actions given the current state. As with Markov decision processes, we will focus on policies that depend on the current state rather than the past history because future states and rewards are conditionally independent of the history given the current state. In addition, we will focus on stationary policies, which do not depend on time. The probability agent $i$ selects action $a$ at state $s$ is given by $\pi^i(a \mid s)$. We will often use $\pi(s)$ to represent a distribution over joint actions.

The utility of a joint policy $\pi$ from the perspective of agent $i$ can be computed using a variation of policy evaluation introduced in section 7.2 for MDPs. The reward to agent $i$ from state $s$ when following joint policy $\pi$ is

$$ R^i(s, \pi(s)) = \sum_a R^i(s, a) \prod_{j \in I} \pi^j(a^j \mid s) $$

(25.1)

The probability of transitioning from state $s$ to $s'$ when following $\pi$ is

$$ T(s' \mid s, \pi(s)) = \sum_a T(s' \mid s, a) \prod_{j \in I} \pi^j(a^j \mid s) $$

(25.2)

In an infinite horizon, discounted game, the utility for agent $i$ from state $s$ is

$$ U^{\pi,i}(s) = R^i(s, \pi(s)) + \gamma \sum_{s'} T(s' \mid s, \pi(s)) U^{\pi,i}(s') $$

(25.3)

which can be solved exactly (algorithm 25.2).
struct MGPolicy
    p # dictionary mapping states to simple game policies
    MGPolicy(p::Base.Generator) = new(Dict(p))
end

(πi::MGPolicy)(s, ai) = πi.p[s](ai)
(πi::SimpleGamePolicy)(s, ai) = πi(ai)

probability(P::MG, s, π, a) = prod(πj(s, aj) for (πj, aj) in zip(π, a))
reward(P::MG, s, π, i) =
    sum(P.R(s,a)[i]*probability(P,s,π,a) for a in joint(P.𝒜))
transition(P::MG, s, π, s′) =
    sum(P.T(s,a,s′)*probability(P,s,π,a) for a in joint(P.𝒜))

function policy_evaluation(P::MG, π, i)
    p(s,a) = prod(πj(s, aj) for (πj, aj) in zip(π, a))
    R′ = [sum(R(s,a)[i]*p(s,a) for a in joint(A)) for s in S]
    T′ = [sum(T(s,a,s′)*p(s,a) for a in joint(A)) for s in S, s′ in S]
    return (1 - γ*T′)\R′
end

25.2 Response Models

We can generalize the response models introduced in the previous chapter to Markov games. Doing so requires taking into account the state transition model.

25.2.1 Best Response

A response policy for agent i is a policy π^i that maximizes expected utility given the fixed policies of other agents π^{-i}. If the policies of the other agents are fixed, then the problem reduces to an MDP. This MDP has state space S and action space A^i. We can define the transition and reward functions as follows:

\[ T'(s' | s, a^i) = T(s' | s, a^i, \pi^{-i}(s)) \] \hspace{1cm} (25.4)

\[ R'(s, a^i) = R^i(s, a^i, \pi^{-i}(s)) \] \hspace{1cm} (25.5)

Because this is a best response for agent i, the MDP only uses reward R^i. Solving this MDP results in a best response policy for agent i. Algorithm 25.3 provides an implementation.
25.2.2 Softmax Response

Similar to what was done in the previous chapter, we can define a softmax response policy, which assigns a stochastic response to the policies of the other agents at each state. As we did in the construction of a deterministic best response policy, we solve an MDP where the agents with the fixed policies $\pi^{-i}$ are folded into the environment. We then extract the action value function $Q(s,a)$ using one-step lookahead. The softmax response is then

$$\pi^i(a^i | s) \propto \exp(\lambda Q(s,a^i)) \quad (25.6)$$

with precision parameter $\lambda \geq 0$. Algorithm 25.4 provides an implementation. This approach can be used to generate hierarchical softmax solutions (section 24.6). In fact, we can use algorithm 24.9 directly.

Algorithm 25.4. The softmax response of agent $i$ to joint policy $\pi$ with precision parameter $\lambda$.

25.3 Nash Equilibrium

The Nash equilibrium concept can be generalized to Markov games. As with simple games, all agents perform a best response to one another, and have no incentive to deviate. All finite Markov games with a discounted infinite horizon have a Nash equilibrium.\(^3\)

\(^2\) Because we assume policies are stationary, in that they do not vary over time, the Nash equilibria covered here are stationary Markov perfect equilibria.

We can find a Nash equilibrium by solving a nonlinear optimization problem similar to the one we solved in the context of simple games. This problem minimizes the sum of the lookahead utility deviations and constrains the policies to be valid distributions:

\[
\begin{align*}
\text{minimize} & \quad \sum_{i \in I} \sum_s \left( U^i(s) - Q^i(s, \pi(s)) \right) \\
\text{subject to} & \quad U^i(s) \geq Q^i(s, a^i, \pi^{-i}(s)) \text{ for all } i, s, a^i \\
& \quad \sum_{a^i} \pi^i(a^i | s) = 1 \text{ for all } i, s \\
& \quad \pi^i(a^i | s) \geq 0 \text{ for all } i, s, a^i
\end{align*}
\] (25.7)

where

\[
Q^i(s, \pi(s)) = R^i(s, \pi(s)) + \gamma \sum_{s'} T(s' | s, \pi(s)) U^i(s')
\] (25.8)

This nonlinear optimization problem is implemented and solved in algorithm 25.5.4

25.4  Fictitious Play

As we did in the context of simple games, we can take a learning-based approach to arrive at joint policies by running agents in simulation. Algorithm 25.6 generalizes the simulation loop introduced in the previous chapter to handle state transitions. The various policies run in simulation update themselves based on the state transitions and the actions taken by the various agents.

One approach for updating policies is to use a generalization of fictitious play (algorithm 25.7) from the previous chapter.5 It involves maintaining a maximum-likelihood model over the policies of the other agents. The maximum likelihood model tracks the state in addition to the action being taken by each agent. We track the number of times agent \( j \) takes action \( a^j \) in state \( s \), storing it in a table \( N(j, a^j, s) \), typically initialized to 1. Then, we can compute the best response assuming that each agent \( j \) follows the state-dependent stochastic policy:

\[
\pi^j(a^j | s) \propto N^j(j, a^j, s)
\] (25.9)

After observing joint action \( a \) in states \( s \), we update

\[
N(j, a^j, s) \leftarrow N(j, a^j, s) + 1
\] (25.10)

for each agent \( j \).
function tensorform(P::MG)
    ℋ' = eachindex(ℋ)
    S' = eachindex(S)
    A' = [eachindex(A[i]) for i in ℋ]
    R' = [R(s,a) for s in S, a in joint(A)]
    T' = [T(s,a,s') for s in S, a in joint(A), s' in S]
    return ℋ', S', A', R', T'
end

function solve!(M::NashEquilibrium, P::MG)
    ℋ, S, A, R, T = tensorform(P)
    S', A', γ = P.S, P.A, P.γ
    model = Model(Ipopt.Optimizer)
    @variable(model, U[i=ℋ, s=S], ai=A[i] ≥ 0)
    @NLobjective(model, Min,
        sum(U[i,s] - sum(prod(π[j,s,a[j]] for j in ℋ)
            * (R[s,y][i] + γ*sum(T[s,y,s']*U[i,s'] for s' in S))
            for (y,a) in enumerate(joint(A))) for i in ℋ, s in S))
    @NLconstraint(model, [i=ℋ, s=S, ai=A[i]],
        U[i,s] ≥ sum(
            prod(j==i ? (a[j]==ai ? 1.0 : 0.0) : π[j,s,a[j]] for j in ℋ)
            * (R[s,y][i] + γ*sum(T[s,y,s']*U[i,s'] for s' in S))
            for (y,a) in enumerate(joint(A))))
    @constraint(model, [i=ℋ, s=S], sum(π[i,s,ai] for ai in A[i]) == 1)
    optimize!(model)
    π' = value.(π)
    πi'(i,s) = SimpleGamePolicy(A'[i][ai] ⇒ π'[i,s,ai] for ai in A[i])
    πi'(i) = MGPolicy(S'[s] ⇒ πi'(i,s) for s in S)
    return [πi'(i) for i in ℋ]
end

Algorithm 25.5. This nonlinear program computes a Nash equilibrium for a Markov game P.
Algorithm 25.6. Functions for taking a random step and running full simulations in Markov games. The simulate function will simulate the joint policy \( \pi \) for \( k_{\text{max}} \) steps starting from a state randomly sampled from \( b \).

As the distributions of the other agents’ actions change, we must update the utilities. The utilities in Markov games are significantly more difficult to compute compared to simple games because of the state dependency. As described in section 25.2.1, for any assignment of fixed policies of others \( \pi^{-i} \) determined from equation (25.9) induces an MDP. Instead of solving an MDP at each update, it is common to apply the update periodically, a strategy adopted from asynchronous value iteration. An example of fictitious play is given in example 25.2.

Our policy \( \pi^i(s) \) for a state \( s \) is derived from a given opponent model \( \pi^{-i} \) and computed utility \( U^i \). We then select a best response:

\[
\arg \max_a Q^i(s, a, \pi^{-i})
\]  

(25.11)

In the implementation here, we use the property that each state of a Markov game policy is a simple game policy whose reward is the corresponding \( Q^i \).

25.5 Gradient Ascent

We can use gradient ascent (algorithm 25.8) to learn policies in a way similar to what was done in the previous chapter for simple games. The state must now be considered and requires learning the action value function. At each time step \( t \), all agents perform joint actions \( a_t \) in a state \( s_t \). As in gradient ascent for simple
mutable struct MGFictitiousPlay
    ℐ # Markov game
    i # agent index
    Qi # state-action value estimates
    Ni # state-action counts
end

function MGFictitiousPlay(T::MG, i)
    Qi = Dict((s, a) => R(s, a)[i] for s in S for a in joint(A))
    Ni = Dict(((j, s, aj)) => 1.0 for j in T for s in S for aj in A[j])
    return MGFictitiousPlay(T, i, Qi, Ni)
end

function (πi::MGFictitiousPlay)(s)
    ℐ, i, Qi = πi.ℐ, πi.i, πi.Qi
    πi'(i, s) = SimpleGamePolicy(ai => πi.Ni[i, s, ai] for ai in joint(A))
    πi'(i) = MGPolicy(s => πi'(i, s) for s in S)
    π = [πi'(i) for i in T]
    U(s, π) = sum(πi.Qi[s, a]*probability(T, s, π, a) for a in joint(A))
    Q(s, π) = reward(T, s, π, i) + γ*sum(transition(T, s, π, s')*U(s', π) for s' in S)
    Q(ai) = Q(s, joint(π, SimpleGamePolicy(ai), i))
    ai = argmax(Q, T.A[πi.i])
    return SimpleGamePolicy(ai)
end

function update!(πi::MGFictitiousPlay, s, a, s')
    ℐ, i, Qi = πi.ℐ, πi.i, πi.Qi
    for (j, aj) in enumerate(A)
        πi.Ni[j, s, aj] += 1
    end
    πi'(i, s) = SimpleGamePolicy(ai => πi.Ni[i, s, ai] for ai in joint(A))
    πi'(i) = MGPolicy(s => πi'(i, s) for s in S)
    π = [πi'(i) for i in T]
    U(π, s) = sum(πi.Qi[s, a]*probability(T, s, π, a) for a in joint(A))
    Q(s, a) = R(s, a)[i] + γ*sum(transition(T, s, a, s')*U(π, s') for s' in S)
    for a in joint(A)
        πi.Qi[s, a] = Q(s, a)
    end
end

Algorithm 25.7. Fictitious play for agent i in an MG T that maintains counts Ni of other agent action selections over time for each state and averages them assuming this is their stochastic policy. It then computes a best response to this policy and performs the corresponding utility-maximizing action.
The predator-prey hex world Markov game (appendix F.13) has one predator (red) and one prey (blue). If the predator catches the prey, it receives a reward of 10 and the prey receives a reward of $-100$. Otherwise, both agents receive a $-1$ reward. The agents move simultaneously. We apply fictitious play with resets to the initial state every 10 steps.

We observe that the predator learns to chase the prey and the prey learns to flee. Interestingly, the predator also learns that the prey runs to the right corner and waits. Here, the prey learns that if it waits at this corner, it can flee from the predator immediately as it jumps towards the prey. In this case, it escapes the predator and runs to the other side of the map.

Below is a plot of the learned opponent model of the highlighted state (both predator and prey hex locations) for both the predator and the prey.
games, an agent $i$ assumes the agents’ policies $\pi_t^{-i}$ are the observed actions $a_t^{-i}$. The gradient is then

$$\frac{\partial U^{\pi_t(s_t)}}{\partial \pi_t(a^i | s_t)} = \frac{\partial}{\partial \pi_t(a^i | s_t)} \left( \sum_a \prod_j \pi_t^j(a^j | s_t) Q_t^{\pi_t}(s_t, a_t) \right)$$  \hspace{1cm} (25.12)

$$= Q_t^{\pi_t}(s_t, a_t, a_t^{-i})$$  \hspace{1cm} (25.13)

The gradient step follows a similar pattern as in the previous chapter, except the state $s$ is included and the expected utility estimate $Q_t^i$ is used:

$$\pi_{t+1}^i(a^i | s_t) = \pi_t^i(a^i | s_t) + \alpha_t Q_t^i(s_t, a_t, a_t^{-i})$$  \hspace{1cm} (25.14)

Again, this update may require projection to ensure a valid probability distribution.

As with fictitious play in the previous section, we must estimate $Q_t^i$. We can use Q-learning:

$$Q_{t+1}^i(s_t, a_t) = Q_t^i(s_t, a_t) + \alpha_t \left( R^i(s_t, a_t) + \gamma \max_{a^i} Q_t^i(s_{t+1}, a^i, a_t^{-i}) - Q_t^i(s_t, a_t) \right)$$  \hspace{1cm} (25.15)

We can use the inverse square-root learning rate $\alpha_t = 1/\sqrt{t}$. Exploration is also necessary. We can use an $\epsilon$-greedy strategy, perhaps also with $\epsilon_t = 1/\sqrt{t}$.

### 25.6 Nash Q-Learning

Another learning-based approach is Nash Q-learning (algorithm 25.9), which borrows inspiration from Q-learning (section 17.2).\textsuperscript{6} The method maintains an estimate of the action value function, which is adapted as the agents react to each other’s changing policies. In the process of updating the action value function, it computes a Nash equilibrium to model the behavior of the other agents.

An agent following Nash Q-learning maintains an estimate of a joint action value function $Q(s, a)$. This action value function is updated after every state transition using a Nash equilibrium computed from a simple game constructed from this value function. After a transition from $s$ to $s'$ following the joint action $a$, we construct a simple game with the same number of agents and the same joint action space, but have the reward function equal to the estimated value of $s'$ such that $R(a') = Q(s', a')$. The agent computes a Nash equilibrium policy $\pi'$ over the

Algorithm 25.8. Gradient ascent for an agent $i$ of an MG $\mathcal{P}$. The algorithm incrementally updates its distributions of actions at visited states following gradient ascent to improve the expected utility. The projection function from algorithm 23.6 is used to ensure that the resulting policy remains a valid probability distribution.

```plaintext
mutable struct MGGradientAscent
    ℙ  # Markov game
    i  # agent index
    t  # time step
    Qi # state-action value estimates
    πi # current policy
end

function MGGradientAscent(ℙ::MG, i)
    ℱ, ℳ = ℙ.ℱ, ℙ.ℳ
    Qi = Dict((s, a) => 0.0 for s in ℳ, a in joint(ℳ))
    uniform() = Dict(s => SimpleGamePolicy(ai => 1.0 for ai in ℙ.ℳ[i]) for s in ℳ)
    return MGGradientAscent(ℙ, i, 1, Qi, uniform())
end

function (πi::MGGradientAscent)(s)
    ℳi, t = πi.ℳ[πi.i], πi.t
    ε = 1 / sqrt(t)
    πi′(ai) = ε/length(ℳi) + (1-ε)*πi.ℳi[s](ai)
    return SimpleGamePolicy(ai => πi′(ai) for ai in ℳi)
end

function update!(πi::MGGradientAscent, s, a, s′)
    ℛ, i, t, Qi = πi.ℛ, πi.i, πi.t, πi.Qi
    ℱ, ℳ, ℳi, R, γ = ℙ.ℱ, ℙ.ℳ, ℙ.ℳ[πi.i], ℙ.ℛ, ℙ.ℛ
gointn(ai) = Tuple(j => i ? ai : a[j] for j in ℱ)
    α = 1 / sqrt(t)
    Qmax = maximum(Qi[s′, jointn(ai)] for ai in ℳi)
    πi.Qi[s, a] += α * (R(s, a)[i] + γ * Qmax - Qi[s, a])
    u = [Qi[s, jointn(ai)] for ai in ℳi]
    π′ = [πi.ℳi[s](ai) for ai in ℳi]
    π = project_to_simplex(π′ + u / sqrt(t))
    πi.t = t + 1
    πi.ℳi[s] = SimpleGamePolicy(ai => p for (ai, p) in zip(ℳi, π))
end
```
next action \(a'\). Under the derived policy, the expected utility of the successor state is:

\[
U(s') = \sum_{a'} Q(s', a') \prod_{j \in I} \pi_j'(a_j')
\]  

(25.16)

The agent then updates its value function:

\[
Q(s, a) \leftarrow Q(s, a) + \alpha (R(s, a) + \gamma U(s') - Q(s, a))
\]  

(25.17)

where the learning rate \(\alpha\) is typically a function of the state-action count \(\alpha = 1/\sqrt{N(s, a)}\).

As with regular Q-learning, we need to adopt an exploration strategy to ensure that all states and actions are tried sufficiently often. In algorithm 25.9, the agent follows an \(\epsilon\)-greedy policy. With probability \(\epsilon = 1/\sum_a N(s, a)\), it selects an action uniformly at random. Otherwise, it will use the result from the Nash equilibrium.

### 25.7 Summary

- Markov games are an extension of MDPs to multiple agents, or an extension of simple games to sequential problems. In these problems, multiple agents compete and individually receive reward over time.
- The Nash equilibrium can be formulated for Markov games, but must now consider all actions for all agents in all states.
- The problem of finding a Nash equilibrium can be formulated as a nonlinear optimization problem.
- We can generalize fictitious play to Markov games by using a known transition function and incorporating estimates of action values.
- Policy hill climbing does not assume a model, but instead uses a gradient ascent approach to iteratively improve a stochastic policy.
- Nash Q-learning adapts traditional Q-learning to multiagent problems and involves solving for a Nash equilibrium of a simple game constructed from models of the other players.
mutable struct NashQLearning
    ℋ # Markov game
    ℌ # agent index
    Q # state-action value estimates
    N # history of actions performed
end

function NashQLearning(ℋ::MG, ℌ)
    ℋ, ℋ, ℋ, ℋ
    Q = Dict((j, s, a) => 0.0 for j in ℋ, s in ℋ, a in joint(ℋ))
    N = Dict((s, a) => 1.0 for s in ℋ, a in joint(ℋ))
    return NashQLearning(ℋ, ℌ, Q, N)
end

function (πi::NashQLearning)(s)
    πi, ℋ, πi, ℋ, ℋ
    ℋ, ℋ, ℋ, ℋ
    M = NashEquilibrium()
    ℋ = SimpleGame(ℋ, ℋ, ℋ, ℋ) for j in ℋ]
    π = solve(M, ℋ)
    ε = 1 / sum(N[s, a] for a in joint(ℋ))
    π′(ai) = ε/length(ℋi) + (1-ε)*π[i](ai)
    return SimpleGamePolicy(ai => π′(ai) for ai in ℋi)
end

function update!(πi::NashQLearning, s, a, s′)
    πi, ℋ, πi, ℋ, ℋ, ℋ
    ℋ, ℋ, ℋ, ℋ
    M = NashEquilibrium()
    ℋ = SimpleGame(ℋ, ℋ, ℋ, ℋ) for j in ℋ]
    π = solve(M, ℋ)
    π.N[s, a] += 1
    α = 1 / sqrt(N[s, a])
    for j in ℋ
        πi.Q[j,s,a] += α*(R(s,a)[j] + γ*utility(ℋ,π,j) - Q[j,s,a])
    end
end

Algorithm 25.9. Nash Q-learning for an agent ℌ in an MG ℋ. The algorithm performs joint-action Q-learning to learn a state-action value function for all agents. A simple game is built with Q, and we compute a Nash equilibrium using algorithm 24.5. The equilibrium is then used to update the value function. This implementation also uses a variable learning rate proportional to the number of times state-joint-action pairs are visited, which is stored in N. Additionally, it uses ε-greedy exploration to ensure all states and actions are explored.

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2021-03-07 23:12:00-08:00, revision 80bf4d6, comments to bugs@algorithmsbook.com
25.8 Exercises

Exercise 25.1. Show how Markov games are extensions of both MDPs and simple games. Show this by formulating an MDP as a Markov game and by formulating a simple game as a Markov game.

Solution: Markov games generalize simple games. For any simple game with $I, A, R$, we can construct a Markov game by just having a single state that self-loops. In other words, this Markov game has $S = \{s^1\}$, $T(s^1 | s^1, a) = 1$, and $R(s^1, a) = R(a)$.

Markov games generalize MDPs. For any MDP with $S, A, T, R$, we can construct a Markov game by just assigning the agents to be this single agent. In other words, this Markov game has $I = \{1\}$, $A^1 = A$, $T(s' | s, a) = T(s' | s', a)$, and $R(s, a) = R(s, a)$.

Exercise 25.2. For an agent $i$, given the fixed policies of other agents $\pi^{-i}$, can there exist a stochastic best response that yields a greater utility than a deterministic best response? Why then do we consider stochastic policies in a Nash equilibrium?

Solution: No, if given fixed policies of other agents $\pi^{-i}$, a deterministic best response is sufficient to obtain the highest utility. The best response can be formulated as solving an MDP as described in section 25.2. It has been shown that deterministic policies are sufficient to provide optimal utility-maximization. Hence, the same is true for a best response in a Markov game.

In a Nash equilibrium, a best response has to hold for all agents. Although a deterministic best response might be equal in utility to a stochastic one, an equilibrium may require stochastic responses to prevent other agents from wanting to deviate.

Exercise 25.3. This chapter discussed only stationary Markov policies. What other categories of policies are there?

Solution: A so-called behavioral policy $\pi^i(h_t)$ is one that has a dependence on complete history $h_t = (s_{1:t}, a_{1:t-1})$. Such policies depend on the history of play of other agents. A non-stationary Markov policy $\pi^i(s, t)$ is one that depends on the time step $t$, but not on the complete history. For example, in the predator prey hex world domain, for the first 10 time steps, the action at a hex might be to go east and after 10 time steps to go west.

There can exist Nash equilibria that are in the space of non-stationary non-Markov joint policies, stationary non-Markov joint policies, and so forth. However, it is proven that every (stationary) Markov game has a stationary Markov Nash equilibrium.

Exercise 25.4. In Markov games, fictitious play requires the utilities to be estimated. List different approaches to compute utilities with their benefits and drawbacks.

Solution: The algorithm presented in this chapter performs a single backup for the visited state $s$ and all joint actions $a$. This approach has the benefit of being relatively efficient because it is a single backup. Updating all joint actions at that state results in exploring actions that were not observed. The drawback of this approach is that we may need to do this update at all states many times to obtain a suitable policy.
An alternative is to only update the visited state and the joint action that was actually taken, which results in a faster update step. The drawback is that it requires many more steps to explore the full range of joint actions.

Another alternative is to perform value iteration at all states \( s \) until convergence at every update step. Recall that the model of the opponent changes on each update. This induces a new MDP, as described for deterministic best response in section 25.2.1. Consequently, we would need to rerun value iteration after each update. The benefit of this approach is that it can result in the most informed decision at each step because the utilities \( Q^i \) consider all states over time. The drawback is that the update step is very computationally expensive.
The multiagent models discussed so far have assumed that all agents can observe the true state. Just as an MDP can be extended to include partial observability, so too can a Markov game be extended to produce a partially observable Markov game (POMG). In fact, a POMG generalizes all others presented in this book. These complex problems can be used to represent domains in which multiple agents receive partial or noisy observations of the environment. This generality makes modeling and solving POMGs computationally challenging. This chapter defines the POMG, outlines policy representations, and presents solution methods.

### 26.1 Partially Observable Markov Games

A POMG (algorithm 26.1) can be seen as either an extension of Markov games to partial observability or as an extension of POMDPs to multiple agents. Each agent \( i \in I \) selects an action \( a^i \in A^i \) based only on local observations \( o^i \) made of a shared state \( s \). The true state of the system \( s \in S \) is shared by all agents, but is not necessarily fully observed. The initial state is drawn from a known initial state distribution \( b \). The likelihood of transitioning from a state \( s \) to a state \( s' \) under their joint action \( a \) follows \( T(s' \mid s, a) \). A joint reward \( r \) is generated following \( R^j(s, a) \), as in Markov games. Each agent strives to maximize its own accumulated reward. After all agents perform their joint action \( a \), a joint observation is emitted by the environment \( o = (o^1, \ldots, o^k) \) from a joint observation space \( O = O^1 \times \cdots \times O^k \). Each agent then receives their individual observation \( o^i \) from this joint observation. The crying baby problem is extended multiple agents in example 26.1.

In POMDPs, we were able to maintain a belief state as discussed in chapter 19, but this approach is not possible in POMGs. Individual agents cannot perform the same kind of belief updates as in POMDPs because the joint action and...
joint observations are not observed. Inferring a probability distribution over joint actions requires that each agent reason about the other agents reasoning about each other, who in turn reasoning about each other—and so on. Inferring a distribution over the other observations is just as complicated because the observations depend upon the actions of the other agents.²

Because of the difficulty of explicitly modeling beliefs in POMGs, we will focus on policy representations that do not require a belief to determine an action. We can use the tree-based conditional plan representation and the graph-based controller representation introduced in the earlier chapters on POMDPs. As in Markov games, each agent in a POMG acts according to a policy \( \pi^i \), or equivalently, the agents together act according to a joint policy \( \pi = (\pi^1, \ldots, \pi^k) \).

Algorithm 26.1. Data structure for a partially observable Markov game.

Example 26.1. The multi-caregiver crying baby problem as a POMG. Appendix F.14 provides additional detail.

Consider a multiagent POMG generalization of the crying baby problem. We have two caregivers taking care of a baby. As in the POMDP version, the states are the baby being hungry or sated. Each caregiver’s actions are to feed, sing, or ignore the baby. If both caregivers choose to perform the same action, the cost is halved. For example, if both caregivers feed the baby, then the reward is only \(-2.5\) instead of \(-5\). However, the caregivers do not perfectly observe the state of the child. Instead they rely on the noisy observations of the child crying, both receiving the same observation. As a consequence of the reward structure, there is a trade-off between helping each other and greedily choosing a less costly action.

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2021-03-07 23:12:00-08:00, revision 80bf4d6, comments to bugs@algorithmsbook.com
26.2 Policy Evaluation

This section discusses how to evaluate joint policies represented as either tree-based conditional plans or graph-based controllers. As in the context of POMDPs, we use conditional plans to represent deterministic policies and controllers to represent stochastic policies.

26.2.1 Evaluating Conditional Plans

Recall that a conditional plan (section 20.2) is a tree where actions are associated with nodes and observations are associated with edges. Each agent has its own tree and initially selects the action associated with its root. After making an observation, each agent proceeds down the tree, taking the edge associated with their observation. The process of taking actions and selecting edges based on observations continues until reaching the end of the tree. Example 26.2 shows an example joint policy consisting of a conditional plan for each agent.

We can compute the joint utility function $U^\pi$ recursively, similar to what was done in equation (20.8) for POMDPs when starting in state $s$:

$$U^\pi(s) = R(s, \pi()) + \gamma \left[ \sum_{s'} T(s' \mid s, \pi()) \sum_{o} O(o \mid \pi(), s') U^\pi(o)(s') \right] \quad (26.1)$$

where $\pi()$ is the vector of actions at the root of the tree associated with $\pi$ and $\pi(o)$ is the vector of subplans associated with the different agents observing their components of the joint observation $o$.

The utility associated with policy $\pi$ from initial state distribution $b$ is given by

$$U^\pi(b) = \sum_{s} b(s) U^\pi(s) \quad (26.2)$$

Algorithm 26.2 provides an implementation.

```
function lookahead(P::POMG, U, s, a)
    u' = sum(T(s,a,s')*sum(O(a,s',o)*U(o,s') for o in O) for s' in S)
    return R(s,a) + γ*u'
end

function evaluate_plan(P::POMG, π, s)
    a = Tuple(πi() for πi in π)
    U(o,s') = evaluate_plan(P, [πi oi for (πi, oi) in zip(π, o)], s')
    return isempty(first(π).subplans) ? P.R(s,a) : lookahead(P, U, s, a)
end

function utility(P::POMG, b, π)
    u = [evaluate_plan(P, π, s) for s in P.S]
    return sum(bs * us for (bs, us) in zip(b, u))
end
```

**Algorithm 26.2.** Conditional plans represent policies in a finite horizon POMG. They are defined for a single agent in algorithm 20.1. We can compute the utility associated with executing a joint policy $\pi$ represented by conditional plans when starting from a state $s$. Computing the utility from an initial state distribution $b$ involves taking a weighted average of utilities when starting from different states.

### 26.2.2 Evaluating Stochastic Controllers

A controller (section 23.1) is represented as a stochastic graph. The controller associated with agent $i$ is defined by the action distribution $\psi^i(a^i \mid x^i)$ and successor distribution $\eta^i(x'^i \mid x^i, a^i, o^i)$. The utility of being in state $s$ with joint node $x$ active and following joint policy $\pi$ is:

$$U^\pi(x, s) = \sum_{a} \prod_{i} \psi^i(a^i \mid x^i) \left( R(s,a) + \gamma \sum_{s'} T(s' \mid s, a) \sum_{o} O(o \mid a, s') \sum_{x'} \prod_{i} \eta^i(x'^i \mid x^i, a^i, o^i) U^\pi(x', s') \right) \quad (26.3)$$
Policy evaluation in this context involves solving this system of linear equations. Alternatively, we can use iterative policy evaluation similar to algorithm 23.2 for POMDPs. The utility when starting from an initial state distribution \( b \) and joint controller state \( x \) is:

\[
U^\pi(x, b) = \sum_s b(s)U(x, s)
\]  

Equation (26.4)

Example 26.3 shows an example joint stochastic controller.

Example 26.3. An example of a two-agent joint policy using controllers for the multi-caregiver crying baby problem.

Below is a joint controller policy \( \pi = (\pi^1, \pi^2) \) for the two caregivers in the crying baby problem. Each controller has two nodes, \( X^i = \{x^i_1, x^i_2\} \).

<table>
<thead>
<tr>
<th>agent 1’s policy ( \pi^1 )</th>
<th>agent 2’s policy ( \pi^2 )</th>
</tr>
</thead>
<tbody>
<tr>
<td>( o^1 ) ( a^1 ) ( o^2 )</td>
<td>( o^1 ) ( a^1 ) ( o^2 )</td>
</tr>
<tr>
<td>( o^1 = \text{quiet} ) ( a^1 = \text{ignore} )</td>
<td>( o^1 = \text{quiet} ) ( a^1 = \text{ignore} )</td>
</tr>
<tr>
<td>( o^2 = \text{hungry} ) ( a^2 = \text{feed} )</td>
<td>( o^2 = \text{hungry} ) ( a^2 = \text{feed} )</td>
</tr>
</tbody>
</table>

26.3 Nash Equilibrium

As with simple games and Markov games, a Nash equilibrium for a POMG is when all agents act according to a best response policy to each other, such that no agent has an incentive to deviate from their policy. Nash equilibria for POMGs tend to be incredibly computationally difficult to solve. Algorithm 26.3 computes a \( d \)-step Nash equilibrium for a POMG is to enumerate all of its possible \( d \)-step joint conditional plans, and to use them to construct a simple game as shown in example 26.4. A Nash equilibrium for this simple game is also a Nash equilibrium for the POMG.

The simple game has the same agents as the POMG. There is a joint action in the simple game for every joint conditional plan in the POMG. The reward received for each action is equal to the utilities under the joint conditional plan in
the POMG. A Nash equilibrium of this constructed simple game can directly be applied as a Nash equilibrium of the POMG.

26.4 Dynamic Programming

The approach taken in the previous section for computing a Nash equilibrium is typically extremely computationally expensive because the actions correspond to all possible conditional plans to some depth. We can adapt the value iteration approach for POMDPs (section 20.5), where we iterated between expanding the depth of the set of considered conditional plans and pruning suboptimal plans. While the worst-case computational complexity is the same as that of the full expansion of all policy trees, this incremental approach can lead to significant savings.

Algorithm 26.4 implements this dynamic programming approach. It begins by constructing all one-step plans. We prune any plans that are dominated by another plan, and we then expand all combinations of one-step plans to produce two-step plans. This procedure of alternating between expansion and pruning is repeated until the desired horizon is reached.

The pruning step eliminates all dominated policies. A policy \( \pi^i \) belonging to an agent \( i \) can be pruned if there exists another policy \( \pi''^i \) that always performs at least as well as \( \pi^i \). Though computationally expensive, this condition can be checked by solving a linear program. This process is related to controller node pruning in POMDP (algorithm 23.4).

It would be computational intractable to solve a separate linear program for every possible combination of the other agent’s policies \( \pi^{-i} \). Instead, we can take a much more efficient approach that will never prune an optimal policy but may under prune. A policy \( \pi^i \) is dominated by \( \pi''^i \) if there do not exist \( b(\pi^{-i}, s) \) between other joint policies \( \pi^{-i} \) and states \( s \) such that:

\[
\sum_{\pi^{-i}} \sum_{s} b(\pi^{-i}, s) U^{\pi'', \pi^{-i}, i}(s) \geq \sum_{\pi^{-i}} \sum_{s} b(\pi^{-i}, s) U^{\pi', \pi^{-i}, i}(s)
\]  

(26.5)

Here, \( b \) is a joint distribution over the policies of other agents and the state. As mentioned at the start of this chapter, it is generally infeasible to compute a belief state, but the equation above checks the space of beliefs for individual policy domination.
Consider the multi-caregiver crying baby problem with a two-step horizon. Recall that for each agent $i$, there are three actions $A^i = \{a^i_1, a^i_2, a^i_3\} = \{\text{feed, sing, ignore}\}$ and two observations $O^i = \{o^i_1, o^i_2\} = \{\text{cry, silent}\}$. Converting this POMG to a simple game results in the following game table. Each caregiver selects simple game actions that correspond to a complete conditional plan. The simple game reward for each agent is the utility associated with the joint policy.

Example 26.4. Computing a Nash equilibrium for the multi-caregiver crying baby problem by converting it into a simple game where the actions correspond to conditional plans.
Algorithm 26.3. A Nash equilibrium is computed for a POMG $\mathcal{P}$ with initial state distribution $b$ by creating a simple game of all conditional plans to some depth $d$. We solve for a Nash equilibrium in this simple game using algorithm 24.5. For simplicity, we select the most probable joint policy. Alternatively, we can randomly select the joint policy at the start of execution.
Algorithm 26.4. Dynamic programming computes a Nash equilibrium $\pi$ for a POMG $\mathcal{P}$, given an initial belief $b$ and horizon depth $d$. It iteratively computes the policy trees and their expected utilities at each step. The pruning phase at each iteration removes dominated policies, which are policy trees that result in a lower expected utility than at least one other available policy tree.

struct POMGDynamicProgramming
    b  # initial belief
    d  # depth of conditional plans
end

function solve($M$::POMGDynamicProgramming, $\mathcal{P}$::POMG)
    $I$, $S$, $A$, $R$, $\gamma$, $b$, $d$ = $\mathcal{P}$.I, $\mathcal{P}$.S, $\mathcal{P}$.A, $\mathcal{P}$.R, $\mathcal{P}$.V, $M$.b, $M$.d
    $\Pi$ = [[ConditionalPlan($a_i$) for $a_i$ in $A[i]$] for $i$ in $I$]
    for $t$ in 1:d
        $\Pi$ = expand_conditional_plans($\mathcal{P}$, $\Pi$)
    end
    $G$ = SimpleGame($\gamma$, $I$, $\Pi$, $\pi$ → utility($\mathcal{P}$, $b$, $\pi$))
    $\pi$ = solve(NashEquilibrium(), $G$)
    return Tuple(argmax($\pi_i$.p) for $\pi_i$ in $\pi$)
end

function prune_dominated!(($\Pi$, $\mathcal{P}$::POMG))
    done = false
    while !done
        done = true
        for $i$ in shuffle($\mathcal{P}$.I)
            for $\pi_i$ in shuffle($\Pi$[$i$])
                if length($\Pi$[$i$]) > 1 && is_dominated($\mathcal{P}$, $\Pi$, $i$, $\pi_i$)
                    filter!(($\pi_i'$$\rightarrow$$\pi_i' \neq \pi_i$, $\Pi$[$i$])
                    done = false
                    break
                end
            end
        end
    end
end

function is_dominated($\mathcal{P}$::POMG, $\Pi$, $i$, $\pi_i$)
    $I$, $S = \mathcal{P}$.I, $\mathcal{P}$.S
    joint$\Pi$noti = joint([[$\Pi$[$j$] for $j$ in $I$ if $j \neq i$])
    $\pi$(\pi_i', $\Pi$noti) = [\j=i ? \pi_i': $\Pi$noti[j>i ? j-1 : j] for $j$ in $I$]
    $U_i$ = Dict((\pi_i', $\Pi$noti, $s$) => evaluate_plan($\mathcal{P}$, $\pi$(\pi_i', $\Pi$noti), $s$)[i]
            for $\pi_i'$ in $\Pi$[$i$], $\Pi$noti in joint$\Pi$noti, $s$ in $S$)
    model = Model(Ipopt.Optimizer)
    @variable(model, $b$)
    @objective(model, Max, $\delta$)
    @constraint(model, [\i'=\Pi[i]],
                  sum($b$[$\Pi$noti, $s$] * (\U_i[\pi_i', $\Pi$noti, $s$] - \U_i[\pi_i, $\Pi$noti, $s$])
                    for $\Pi$noti in joint$\Pi$noti for $s$ in $S$) ≥ $\delta$)
    @constraint(model, sum($b$) == 1)
    optimize!(model)
    return value($\delta$) ≥ 0
end
We can construct a single linear program to check equation (26.5). If the linear program is feasible, then it means $\pi^i$ is not dominated by any other $\pi^{i'}$. A similar linear program was created to prune alpha vectors in POMDPs in equation (20.16).

\[
\begin{align*}
\text{maximize} & \quad \delta \\
\text{subject to} & \quad b(\pi^{-i}, s) \geq 0 \text{ for all } \pi^{-i}, s \\
& \quad \sum_{\pi^{-i}} \sum_s b(\pi^{-i}, s) = 1 \\
& \quad \sum_{\pi^{-i}} \sum_s b(\pi^{-i}, s) \left( U^{\pi^{i'}, \pi^{-i}, i} - U^{\pi^{i}, \pi^{-i}, i} \right) \geq \delta \text{ for all } \pi^{i'}
\end{align*}
\]  

(26.6)

The pruning step removes dominated policies by randomly selecting an agent $i$ and checking for domination of each of its policies. This process repeats until a pass over all agents fails to find any dominated policies. Example 26.5 shows this process on the multi-caregiver crying baby problem.

26.5 Summary

- Partially observable Markov games (POMGs) generalize POMDPs to multiple agents and MGs to partial observability.
- Because agents generally cannot maintain beliefs in POMGs, policies typically take the form of conditional plans or finite state controllers.
- Nash equilibria in the form of $d$-step conditional plans for POMGs can be obtained by finding Nash equilibria for simple games whose joint actions consist of all possible POMG joint policies.
- Dynamic programming approaches can be used to more efficiently compute Nash equilibria by iteratively constructing sets of deeper conditional plans, while pruning dominated plans to restrict the search space.

26.6 Exercises

Exercise 26.1. Show that a POMG generalizes both a POMDP and an MG.
Consider the multi-caregiver crying baby problem solved by dynamic programming. Initially, the policies at depth $d = 2$ are:

After the pruning step, the agent policies are:

In this case, the pruning step finds the best joint policy. This approach significantly reduces the number of possible joint policies that the next iteration of the algorithm needs to consider.
**Solution:** For any POMDP, we can define a POMG with one agent $I = \{1\}$. The states $S$ are identical, as are the actions $A = (A^1)$ and observations $O = (O^1)$. Thus, the state transition, observation function, and rewards of the POMG directly follow. The Nash equilibrium optimization only has one agent, so it results in a simple maximization of expected value, identical to a POMDP.

For any MG, we can define a POMG with the same agents $I$, states $S$, joint actions $A$, transitions $T$, and joint rewards $R$. The individual observations is assigned to be the states $O^i = S$. The observation function then deterministically provides each agent with the true state $O(o \mid a, s') = 1$ if $o = (s', \ldots, s')$ and 0 otherwise.

**Exercise 26.2.** How can we incorporate communication between agents into the POMG framework?

**Solution:** The action space for the agents can be augmented to include communication actions. The other agents can observe these communication actions according to their observation model.

**Exercise 26.3.** Do agents always have an incentive to communicate?

**Solution:** Agents in POMGs are often competitive, in which case there would be no incentive to communicate with others. If their rewards are aligned to some degree, they may be inclined to communicate.

**Exercise 26.4.** How many possible joint conditional plans are there of depth $d$?

**Solution:** Recall that there are $|A|^{|O|^d-1}/(|O|-1)$ possible $d$-step single-agent conditional plans. We can construct a joint policy of conditional plans using every combination of these single-agent conditional plans across agents. The number of $d$-step multiagent conditional plans is

$$
\prod_{i \in I} |A^i|^{|O^i|^d-1}/(|O^i|-1)
$$

**Exercise 26.5.** Define best response for a POMG in terms of an agent $i$’s utilities $U^{\pi,i}$. Propose iterated best response for POMGs.

**Solution:** The best response $\pi^i$ of agent $i$ to other agents’ policies $\pi^{-i}$ is defined following equation (24.2) for an initial belief $b$:

$$
U^{\pi^i, \pi^{-i}}(b) \geq U^{\pi'^i, \pi^{-i}}(b)
$$

with any other policy $\pi'^i$. For conditional plans, $U^{\pi,i}$ is defined by equation (26.1) and equation (26.2).

The implementation of iterated best response follows from section 24.2.1. First, the conditional plans and simple game can be created, as in algorithm 26.3. Then, we can iterate best response using algorithm 24.8.
27 Collaborative Agents

Many multiagent domains are collaborative, where all agents act independently in an environment while working towards a common shared objective. Applications range from robotic search and rescue to interplanetary exploration rovers. The decentralized partially observable Markov decision process (Dec-POMDP) captures the generality of POMGs while focusing on such collaborative agent settings. The model is more amenable to scalable approximate algorithms because of its single shared objective, as opposed to finding an equilibrium among multiple individual agent objectives. This chapter presents the Dec-POMDP model, highlights its subclasses, and describes algorithms that solve them optimally and approximately.

27.1 Decentralized Partially Observable Markov Decision Processes

A Dec-POMDP (algorithm 27.1) is a POMG with all agents sharing the same objective. Each agent $i \in \mathcal{I}$ selects a local action $a^i \in A^i$ and based on a history of local observations $o^i \in \mathcal{O}^i$. The true state of the system $s \in S$ is shared by all agents. A single reward is generated by $R(s, a)$ based on the state $s$ and the joint action $a$. The goal of all agents is to maximize the shared expected reward over time under local partial observability. Example 27.1 describes a Dec-POMDP version of the predator-prey problem.

Example 27.1. The collaborative predator-prey problem as a Dec-POMDP. Additional detail is provided in appendix F.15.
Many of the same challenges of POMGs persist in Dec-POMDPs, such as the general inability of agents to maintain a belief state. We focus on policies represented as conditional plans or controllers. The same algorithms introduced in the previous chapter can be used to evaluate policies. All that is required is to create a POMG with \( R^i(s, a) \) for each agent \( i \) equal to the \( R(s, a) \) from the Dec-POMDP.

Algorithm 27.1. Data structure for a decentralized partially observable Markov decision process (Dec-POMDP). The joint function from algorithm 24.2 allows the creation of all permutations of a set provided, such as \( A \) or \( O \). The tensorform function converts the Dec-POMDP to a tensor representation.

27.2 Subclasses

There are many notable subclasses of Dec-POMDPs. Categorizing these subclasses is useful when designing algorithms that take advantage of their specific structure. Table 27.1 summarizes some of these subclasses. Figure 27.1 illustrates the relationships between the models discussed in this book.

<table>
<thead>
<tr>
<th>Agents</th>
<th>Observability</th>
<th>Communication</th>
<th>Model</th>
</tr>
</thead>
<tbody>
<tr>
<td>Single</td>
<td>Full</td>
<td>—</td>
<td>MDP</td>
</tr>
<tr>
<td>Single</td>
<td>Partial</td>
<td>—</td>
<td>POMDP</td>
</tr>
<tr>
<td>Multiple</td>
<td>Full</td>
<td>Free</td>
<td>MMDP</td>
</tr>
<tr>
<td>Multiple</td>
<td>Full</td>
<td>General</td>
<td>MMDP</td>
</tr>
<tr>
<td>Multiple</td>
<td>Joint full</td>
<td>Free</td>
<td>MMDP</td>
</tr>
<tr>
<td>Multiple</td>
<td>Joint full</td>
<td>General</td>
<td>Dec-MDP</td>
</tr>
<tr>
<td>Multiple</td>
<td>Partial</td>
<td>Free</td>
<td>MPOMDP</td>
</tr>
<tr>
<td>Multiple</td>
<td>Partial</td>
<td>General</td>
<td>Dec-POMDP</td>
</tr>
</tbody>
</table>

One attribute of interest is joint full observability, which is when each agent observes an aspect of the state, such that if they were to combine their observations,
it would uniquely reveal the true state. The agents, however, do not share their observations. This property ensures that if $O(o | a, s') > 0$ then $P(s' | o) = 1$. A Dec-POMDP with joint full observability is called a decentralized Markov decision process (Dec-MDP).

In many settings, the state space of a Dec-POMDP is factored, one for each agent and one for the environment. This is called a factored Dec-POMDP. We have $S = S^0 \times S^1 \times S^k$, where $S^i$ is the factored state component associated with agent $i$ and $S^0$ is the factored state component associated with the general environment. For example, in collaborative predator-prey, each agent has their own a state factor for their location and the position of the prey is associated with the environment component of the state space.

In some problems, a factored Dec-POMDP may have one or more of the following properties:

- **Transition independence**, where agents may not affect each other’s state:
  \[
  T(s' | s, a) = T^0(s'^0 | s^0) \prod_i T^i(s'^i | s^i, a^i) \tag{27.1}
  \]
Independence | Complexity
--- | ---
Transitions, observations, and rewards | P-complete
Transitions and observations | NP-complete
Any other subset | NEXP-complete

- **Observation independence**, where the observations of agents depend only on their local state and actions:

\[
O(o \mid a, s') = \prod_i O^i(o^i \mid a^i, s'^i)
\]  

(27.2)

- **Reward independence**, where the reward can be decomposed into multiple independent pieces that are combined together:

\[
R(s, a) = R^0(s^0) + \sum_i R^i(s^i, a^i)
\]

(27.3)

Depending on which of these independence properties are satisfied, the resulting computational complexity can vary significantly as summarized in Table 27.2. It is important to take these independences into account when modeling a problem to improve scalability.

A network distributed partially observable Markov decision process (ND-POMDP) is a Dec-POMDP with transition and observation independence and a special reward structure. The reward structure is represented by a coordination graph. In contrast with the graphs used earlier in this book, a coordination graph is a type of hypergraph, which allows edges to connect any number of nodes. The nodes in the ND-POMDP hypergraph correspond to the various agents. The edges relate to interactions between the agents in the reward function. An ND-POMDP associates with each edge \( j \) in the hypergraph a reward component \( R_j \) that depends on the state and action components to which the edge connects. The reward function in an ND-POMDP is simply the sum of the reward components associated with the edges. Figure 27.2 provides an example coordination graph that results in a reward function that can be decomposed as follows:

\[
R_{123}(s_1, s_2, s_3, a_1, a_2, a_3) + R_{34}(s_3, s_4, a_3, a_4) + R_5(s_5, a_5)
\]

(27.4)

Sensor network and target tracking problems are often framed as ND-POMDPs.

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2021-03-07 23:12:00-08:00, revision 80bf4d6, comments to bugs@algorithmsbook.com
The ND-POMDP model is similar to the transition and observation independent Dec-MDP model, but it does not make the joint full observability assumption. Even if all observations are shared, the true state of the world may not be known. Furthermore, even with factored transitions and observations, a policy in an ND-POMDP is a mapping from observation histories to actions, unlike the transition and observation Dec-MDP case in which policies are mappings from local states to actions. The worst-case complexity remains the same as a full Dec-POMDP (NEXP-complete), but algorithms for ND-POMDPs are typically much more scalable in the number of agents. Scalability can increase as the coordination graph becomes less connected.

If agents are able to communicate their actions and observations perfectly without penalty, the agents are then able to maintain a collective belief state. This model is called a multiagent MDP (MMDP) or a multiagent POMDP (MPOMDP). MMDPs and MPOMDPs can also result when there is transition, observation, and reward independence. Any MDP or POMDP algorithm from earlier chapters can be applied to solve these problems.

### 27.3 Dynamic Programming

The dynamic programming algorithm for Dec-POMDPs applies the Bellman equation at each step and prunes dominated policies. This process is identical to dynamic programming for POMGs, except that each agent shares the same reward. Algorithm 27.2 implements this procedure.

```plaintext
struct DecPOMDPDynamicProgramming
    b  # initial belief
    d # depth of conditional plans
end

function solve(M::DecPOMDPDynamicProgramming, P::DecPOMDP)
    R′(s, a) = [R(s, a) for i in ℐ]
    P′ = POMG(γ, ℐ, S, A, O, T, O, R′)
    M′ = POMGDynamicProgramming(M.b, M.d)
    return solve(M′, P′)
end
```

Algorithm 27.2. Dynamic programming computes the optimal joint policy π for a Dec-POMDP P, given an initial belief b and horizon depth d. Since Dec-POMDPs are a special collaborative class of POMGs, we can directly use the POMG algorithm.
27.4 Iterated Best Response

Instead of exploring joint policies directly, we can perform a form of itera
ted best response (algorithm 27.3). In this approach, we iteratively select an agent and compute a best response policy assuming that the other agents are following a fixed policy.\(^3\) This approximate algorithm is typically fast because it is only choosing the best policy for one agent at a time. Moreover, since all agents share the same reward, it tends to terminate after relatively few iterations.

Iterated best response begins with a random initial joint policy \(\pi_1\). The process randomly iterates over the agents. If agent \(i\) is selected, its policy \(\pi^i\) is updated with a best response to the other agents’ fixed policies \(\pi^{-i}\) with initial belief distribution \(b\):

\[
\pi^i \leftarrow \arg \max_{\pi''} U^{\pi'', \pi^{-i}}(b) \tag{27.5}
\]

with ties favoring the current policy. This process can terminate when agents stop changing their policy.

While this algorithm is fast and is guaranteed to converge, it does not always find the best joint policy. It relies on iterated best response to find a Nash equilibrium, but there may be many Nash equilibria with different utilities associated with them. This approach will only find one of them.

27.5 Heuristic Search

Instead of expanding all joint policies, heuristic search (algorithm 27.4) explores a fixed number of policies.\(^4\) The fixed number of policies stored over iterations prevents exponential growth. The heuristic exploration guides the search by attempting to only expand the best joint policies until depth \(d\) is reached.

Each iteration \(k\) of the algorithm keeps a set of joint policies \(\Pi_k\). Initially this is the one-step conditional plans. Subsequent iterations begin by fully expanding the conditional plans. The goal is to add a fixed number of these for the next iteration.

To decide which among the possible conditional plans to add the set, we want to prioritize the policies that are more likely to maximize utility. However, since we expand the conditional plans from the bottom up, we cannot simply evaluate the policies from the initial belief state \(b\). Instead, we need an estimate of the belief \(d - k\) steps into the future, which we compute by taking random actions and

\[^3\text{This type of algorithm is also called joint equilibrium-based search for policies (JESP). R. Nair, M. Tambe, M. Yokoo, D. Pynadath, and S. Marsella, “Taming Decentralized POMDPs: Towards Efficient Policy Computation for Multiagent Settings,” in International Joint Conference on Artificial Intelligence (IJCAI), 2003. It can be further improved by performing dynamic programming.}\]

\[^4\text{This approach is also known as memory bounded dynamic programming (MBDP) S. Seuken and S. Zilberstein, “Memory-Bounded Dynamic Programming for Decentralized POMDPs,” in International Joint Conference on Artificial Intelligence (IJCAI), 2007. There are other heuristic search algorithms such as multiagent A* (MMA*). D. Szer, F. Charpillet, and S. Zilberstein, “MAA*: A Heuristic Search Algorithm for Solving Decentralized POMDPs,” in Conference on Uncertainty in Artificial Intelligence (UAI), 2005.}\]
27.6. Nonlinear Programming

We can use nonlinear programming (NLP) (algorithm 27.5) to find an optimal joint controller policy representation of a fixed-size. This method generalizes the NLP approach for POMDPs from section 23.3. Given a fixed set of nodes $X_i$ for each agent $i$, initial belief $b$, and initial joint nodes $x_1$, the optimization problem is:

$$\text{minimize } U^\pi(b_k)$$

subject to:

$$\pi \in \pi_k$$

where $U^\pi(b_k)$ is the utility at iteration $k$. Example 27.2 demonstrates the process.

Algorithm 27.3. Iterated best response for collaborative DecPOMDP $\mathcal{P}$ iteratively performs a deterministic best response to rapidly search the conditional plan policies space. The solve function implements this procedure up to $k_{\text{max}}$ steps, maximizing the value at an initial belief $b$ for conditional plans of depth $d$.

Algorithm 27.4. Memory bounded heuristic search uses a heuristic function to search the space of conditional plans for a Dec-POMDP $\mathcal{P}$. The $\text{solve}$ function tries to maximize the value at an initial belief $b$ for joint conditional plans of depth $d$. The $\text{explore}$ function generates a belief $t$ steps into the future by taking random actions and simulating actions and observations. The algorithm is memory-bounded, only keeping $\pi_{\text{max}}$ conditional plans per agent.

```math
\text{struct DecPOMDPHeuristicSearch}
\begin{align*}
& \text{b} \quad \# \text{initial belief} \\
& \text{d} \quad \# \text{depth of conditional plans} \\
& \pi_{\text{max}} \quad \# \text{number of policies}
\end{align*}
\text{end}

\text{function solve}(M::\text{DecPOMDPHeuristicSearch}, \mathcal{P}::\text{DecPOMDP})
\begin{align*}
& \mathcal{I}, \mathcal{S}, \mathcal{A}, \mathcal{O}, \mathcal{T}, \mathcal{O}, \mathcal{R}, \mathcal{V} = \mathcal{P}.I, \mathcal{P}.S, \mathcal{P}.A, \mathcal{P}.O, \mathcal{P}.T, \mathcal{P}.O, \mathcal{P}.R, \mathcal{P}.V \\
& b, d, \pi_{\text{max}} = M.b, M.d, M.\pi_{\text{max}} \\
& R'(s, a) = [R(s, a) \text{ for } i \text{ in } \mathcal{I}] \\
& \mathcal{P}' = \text{POMG}(\mathcal{V}, \mathcal{I}, \mathcal{S}, \mathcal{A}, \mathcal{O}, \mathcal{T}, \mathcal{O}, \mathcal{R}) \\
& \Pi = \[[\text{ConditionalPlan}(a_i) \text{ for } a_i \text{ in } \mathcal{A}[i]] \text{ for } i \text{ in } \mathcal{I}] \\
& \text{for } t \text{ in } 1:d \\
& \quad \text{all}\Pi = \text{expand}_\text{conditional}_\text{plans}(\mathcal{P}, \Pi) \\
& \quad \Pi = \[[] \text{ for } i \text{ in } \mathcal{I}] \\
& \quad \text{for } z \text{ in } 1:\pi_{\text{max}} \\
& \quad \quad b' = \text{explore}(M, \mathcal{P}, t) \\
& \quad \quad \pi = \text{argmax}(\pi \to \text{first}(\text{utility}(\mathcal{P}', b', \pi)), \text{joint}(\text{all}\Pi)) \\
& \quad \quad \text{for } i \text{ in } \mathcal{I} \\
& \quad \quad \quad \text{push!}(\Pi[i], \pi[i]) \\
& \quad \quad \quad \text{filter!}(\pi_i \to \pi_i != \pi[i], \text{all}\Pi[i]) \\
& \quad \text{end} \\
& \quad \text{return } \text{argmax}(\pi \to \text{first}(\text{utility}(\mathcal{P}', b, \pi)), \text{joint}(\Pi)) \\
\end{align*}
\text{end}

\text{function explore}(M::\text{DecPOMDPHeuristicSearch}, \mathcal{P}::\text{DecPOMDP}, t)
\begin{align*}
& \mathcal{I}, \mathcal{S}, \mathcal{A}, \mathcal{O}, \mathcal{T}, \mathcal{O}, \mathcal{R}, \mathcal{V} = \mathcal{P}.I, \mathcal{P}.S, \mathcal{P}.A, \mathcal{P}.O, \mathcal{P}.T, \mathcal{P}.O, \mathcal{P}.R, \mathcal{P}.V \\
& b = \text{copy}(M.b) \\
& b' = \text{similar}(b) \\
& s = \text{rand}(\text{SetCategorical}(\mathcal{S}, b)) \\
& \text{for } t \in 1:t \\
& \quad a = \text{Tuple}(\text{rand}(\mathcal{A}[i]) \text{ for } a_i \text{ in } \mathcal{A}) \\
& \quad s' = \text{rand}(\text{SetCategorical}(\mathcal{S}, [\mathcal{T}(s, a, s') \text{ for } s' \text{ in } \mathcal{S}])) \\
& \quad o = \text{rand}(\text{SetCategorical}(\text{joint}(0), [\mathcal{O}(a, s', o) \text{ for } o \text{ in } \text{joint}(0)])) \\
& \quad \text{for } (i', s') \text{ in enumerate}(\mathcal{S}) \\
& \quad \quad \text{po} = \mathcal{O}(a, s', o) \\
& \quad \quad b'[i'] = \mathcal{P} \times \text{sum}(\mathcal{T}(s, a, s') \times b[i] \text{ for } (i, s) \text{ in enumerate}(\mathcal{S})) \\
& \quad \quad \text{normalize!}(b', 1) \\
& \quad \quad b, s = b', s' \\
& \text{end} \\
& \text{return } b'
\end{align*}
\text{end}
```
Consider the collaborative predator-prey problem shown in the margin. We apply heuristic search to a depth of $d = 3$ with 3 policies retained at each iteration. After iteration $k = 1$, the policies are:

At the next iteration $k = 2$, heuristic search again starts at the initial belief and takes $d - k = 3 - 2 = 1$ steps following the heuristic exploration. The explored beliefs used to select the next three conditional plans are:

The policies after iteration $k = 2$ are:

The beliefs were used to determine the root node’s action and the two subtrees below it. These subtrees are built from the prior iteration’s trees.
maximize \[ U_{\psi, \eta} \sum_s b(s) U(x_1, s) \]
subject to \[ U(x, s) = \sum_a \prod_i \psi_i(a^i | x^i) \left( R(s, a) + \gamma \sum_{s'} T(s' | s, a) \sum_o O(o | a, s') \sum_{x'} \prod_i \eta_i(x^{ii} | x^i, a^i, o^i) U(x', s') \right) \]
for all \( x, s \)
\[ \psi_i(a^i | x^i) \geq 0 \quad \text{for all } i, x^i, a^i \]
\[ \sum_a \psi_i(a^i | x^i) = 1 \quad \text{for all } i, x^i \]
\[ \eta_i(x^{ii} | x^i, a^i, o^i) \geq 0 \quad \text{for all } i, x^i, a^i, o^i, x^{ii} \]
\[ \sum_{x^{ii}} \eta_i(x^{ii} | x^i, a^i, o^i) = 1 \quad \text{for all } i, x^i, a^i, o^i \]

(27.6)

27.7 Summary

- Decentralized partially observable Markov decision processes (Dec-POMDPs) are fully cooperative POMGs that model a team of agents working together towards a shared goal, each acting individually using only local information.

- Because determining a belief state is infeasible as in POMGs, policies are generally represented as conditional plans or controllers, allowing each agent to map individual sequences of observations to individual actions.

- Many subclasses of Dec-POMDPs exist with different degrees of computational complexity.

- Dynamic programming computes the value function iteratively, pruning dominated policies as it iterates using a linear program.

- Iterated best response computes a best utility-maximizing response policy for a single agent at a time, iteratively converging to a joint equilibrium.

- Heuristic search searches a fixed subset of policies at each iteration, guided by a heuristic.

- Nonlinear programming can be used to generate controllers of a fixed size.
Algorithm 27.5. Nonlinear programming (NLP) computes the optimal joint controller policy \( \pi \) for a Dec-POMDP \( \mathcal{P} \), given an initial belief \( b \) and number of controller nodes \( \ell \) for each agent. This generalizes the NLP solution in algorithm 23.5.

```python
struct DecPOMDPNonlinearProgramming
    b # initial belief
    \( \ell \) # number of nodes for each agent
end

function tensorform(\( \mathcal{P} :: \text{DecPOMDP} \))
    \( \mathcal{T}, \mathcal{S}, \mathcal{A}, 0, R, T, O = \mathcal{P}.T, \mathcal{P}.S, \mathcal{P}.A, \mathcal{P}.0, \mathcal{P}.R, \mathcal{P}.T, \mathcal{P}.O \)
    \( \mathcal{T}' = \text{eachindex}(\mathcal{T}) \)
    \( \mathcal{S}' = \text{eachindex}(\mathcal{S}) \)
    \( \mathcal{A}' = [\text{eachindex}(\mathcal{A}_i) \text{ for } \mathcal{A}_i \text{ in } \mathcal{A}] \)
    \( \mathcal{O}' = [\text{eachindex}(\mathcal{O}_i) \text{ for } \mathcal{O}_i \text{ in } \mathcal{O}] \)
    \( \mathcal{R}' = [R(s,a) \text{ for } s \in \mathcal{S}, \text{ a in } \text{joint}(\mathcal{A})] \)
    \( \mathcal{T}' = [T(s,a,s') \text{ for } s \in \mathcal{S}, \text{ a in } \text{joint}(\mathcal{A}), s' \in \mathcal{S}] \)
    \( \mathcal{O}' = [O(a,s',o) \text{ for } a \in \text{joint}(\mathcal{A}), s' \in \mathcal{S}, o \in \text{joint}(\mathcal{O})] \)
    return \( \mathcal{T}', \mathcal{S}', \mathcal{A}', \mathcal{O}', \mathcal{R}', \mathcal{T}', \mathcal{O}' \)
end

function solve(M::DecPOMDPNonlinearProgramming, \( \mathcal{P} :: \text{DecPOMDP} \), \( \mathcal{T}, \mathcal{S}, \mathcal{A}, 0, R, T, O = \text{tensorform}(\mathcal{P}) \)
    \( X = [\text{collect}(1:M.\ell) \text{ for } i \text{ in } \mathcal{T}] \)
    jointX, jointA, jointO = joint(\( X \)), joint(\( \mathcal{A} \)), joint(\( \mathcal{O} \))
    \( x1 = \text{jointX}[1] \)
    model = Model(Iopt.Optimizer)
    \( @variable(model, U[jointX, :]) \)
    \( @variable(model, \Psi[i=T,X[i],[\mathcal{A}[i]]] \geq 0) \)
    \( @variable(model, \eta[i=T,X[i],[\mathcal{A}[i], \mathcal{O}[i], X[i]]] \geq 0) \)
    \( @objective(model, \text{Max}, \text{b} \cdot U(x1,:)) \)
    \( @NLconstraint(model, [x=jointX, s=S], \) \( U[x,s] = (\sum \prod (\Psi[i,x1,[\mathcal{A}[i]]) \text{ for } i \text{ in } \mathcal{T}) \)
    \( \ast (R[s,y] \ast \sum \text{prod}(T[s,y,s'] \ast \text{sum}(O[y,s',z])) \ast \text{prod}(\eta[i,x1,[\mathcal{A}[i],o[i],x'[i]]) \text{ for } i \text{ in } \mathcal{T}) \)
    \( \ast U[x',s'] \text{ for } x' \text{ in } \text{jointX} \)
    \( \text{for } (z, o) \text{ in } \text{enumerate}((\text{jointO})) \text{ for } s' \text{ in } \mathcal{S}) \)
    for (y, a) \text{ in } \text{enumerate}((\text{jointA})) \)
    \( @constraint(model, [i=T,xi=X[i]], \) sum(\( \Psi[i,xi,ai] \) \text{ for } ai \text{ in } \mathcal{A}[i]) = 1) \)
    \( @constraint(model, [i=T,xi=X[i],ai=\mathcal{A}[i],0i=\mathcal{O}[i]], \) sum(\( \eta[i,xi,ai,0i,xi'] \) \text{ for } xi' \text{ in } X[i]) = 1) \)
    optimize!(model)
    \( \Psi', \eta' = \text{value}(\Psi), \text{value}(\eta) \)
    return [ControllerPolicy(\( \mathcal{P}, X[i], \)) for i \text{ in } \mathcal{T}]
end
```
27.8 Exercises

Exercise 27.1. Compared to a POSG, why might a Dec-POMDP be more useful in practice, especially in robotic applications?

Solution: Dec-POMDPs are fully-cooperative. Multi-robot domains, such as search and rescue, require the robots to work together as a team. The engineers have full control of the executable policies of each robot. Conversely, in a competitive POMG (or even MG), the algorithms compute policies for all agents. However, in competitive settings, we likely only have control over one agent’s policy. Additionally, the model realistically captures the limited sensors and sensor models in robotics, as in POMDPs, except in the multiagent setting.

Exercise 27.2. Why is a Dec-MDP with joint full observability different from agents knowing the state?

Solution: Full joint observability means if agents were to share their individual observations, then the team would know the true state. This can be done offline during planning. Thus in Dec-MDPs, the true state is essentially known during planning. The issue is that it requires agents to share their individual observations, which cannot be done online during execution. Therefore, planning still needs to reason about the uncertain observations made by the other agents.

Exercise 27.3. Propose a fast algorithm for a Dec-MDP with transition, observation, and reward independence. Prove that it is correct.

Solution: If a factored Dec-MDP satisfies all three independence assumptions, then we can solve it as \(|I|\) separate MDPs. The resulting policy \(\pi^i\) for each agent \(i\)'s MDP can then be combined to form the optimal joint policy. To prove this fact, consider the utility of each agent’s individual MDP:

\[
U^{\pi^i}(s^i) = R(s^i, \pi^i()) + \gamma \left[ \sum_{s'^i} T^i(s'^i | s^i, \pi^i()) \sum_{o^i} O^i(o^i | \pi^i(), s'^i) U^{\pi^i(o^i)}(s'^i) \right]
\]

(27.7)

We sum of each of their individual contributions:

\[
\sum_i U^{\pi^i}(s) = \sum_i \left[ R(s^i, \pi^i()) + \gamma \left[ \sum_{s'^i} T^i(s'^i | s^i, \pi^i()) \sum_{o^i} O^i(o^i | \pi^i(), s'^i) U^{\pi^i(o^i)}(s'^i) \right] \right]
\]

(27.8)

We can combine \(T^i\) and \(O^i\) into a single probability distribution \(P\), move the summation, and apply the definition of reward independence:
This is the Dec-MDP utility function derived from equation (26.1), completing the proof.

Exercise 27.4. How can we use an MMDP or MPOMDP as a heuristic in Dec-POMDP heuristic search?
**Solution:** We can assume free communication for planning. At each time step $t$, all agents know $a_t$ and $o_t$, allowing us to maintain a multiagent belief $b_t$, resulting in an MPOMDP. This MPOMDP solution can be used as a heuristic to guide the search of policy trees. Alternatively, we create a heuristic where we assume that the true state and joint actions known. This results in an MMDP, and can also be used as a heuristic. These assumptions are only used for planning. Execution is still a Dec-POMDP wherein agents receive individual observations without free communication among themselves. Either heuristic results in a joint policy $\hat{\pi}$ for heuristic exploration.

**Exercise 27.5.** How can we compute a best response controller? Describe how this could be used in iterated best response.

**Solution:** For an agent $i$, the best response controller $X^i$, $\psi^i$, and $\eta^i$ can be computed by a nonlinear program. The program is similar to section 27.6, except that $X^{-i}$, $\psi^{-i}$, and $\eta^{-i}$ are now given and no longer variables:

\[
\begin{align*}
\text{maximize } & \sum_s b(s) U(x_1, s) \\
\text{subject to } & U(x, s) = \sum_a \prod_i \psi^i(a^i | x^i) \left( R(s, a) + \gamma \sum_{s'} T(s' | s, a) \sum_o O(o | a, s') \sum_{x'} \prod_i \eta^i(x'^i | x^i, a^i, o^i) U(x', s') \right) \\
& \forall x, s \\
& \psi^i(a^i | x^i) \geq 0 \quad \forall x^i, a^i \\
& \sum_a \psi^i(a^i | x^i) = 1 \quad \forall x^i \\
& \eta^i(x'^i | x^i, a^i, o^i) \geq 0 \quad \forall x^i, a^i, o^i, x'^i \\
& \sum_{x'^i} \eta^i(x'^i | x^i, a^i, o^i) = 1 \quad \forall x^i, a^i, o^i \quad (27.19)
\end{align*}
\]

Adapting algorithm 27.3 for controller policies, this program replaces the inner best response operation.
APPENDICES
A Mathematical Concepts

This appendix provides a brief overview of some of the mathematical concepts used in this book.

A.1 Measure Spaces

Before introducing the definition of a measure space, we will first introduce the notion of a sigma-algebra over a set $\Omega$. A sigma-algebra is a collection $\Sigma$ of subsets of $\Omega$ such that:

1. $\Omega \in \Sigma$.
2. If $E \in \Sigma$, then $\Omega \setminus E \in \Sigma$ (closed under complementation).
3. If $E_1, E_2, E_3, \ldots \in \Sigma$, then $E_1 \cup E_2 \cup E_2 \ldots \in \Sigma$ (closed under countable unions).

An element $E \in \Sigma$ is called a measurable set.

A measure space is defined by a set $\Omega$, a sigma-algebra $\Sigma$, and a measure $\mu : \Omega \to \mathbb{R} \cup \{\infty\}$. For $\mu$ to be a measure, the following properties must hold:

1. If $E \in \Sigma$, then $\mu(E) \geq 0$ (non-negativity).
2. $\mu(\emptyset) = 0$.
3. If $E_1, E_2, E_3, \ldots \in \Sigma$ are pairwise disjoint, then $\mu(E_1 \cup E_2 \cup E_2 \ldots) = \mu(E_1) + \mu(E_2) + \mu(E_2) + \cdots$ (countable additivity).
A.2 Probability Spaces

A probability space is a measure space \((\Omega, \Sigma, \mu)\) with the requirement that \(\mu(\Omega) = 1\).

In the context of probability spaces, \(\Omega\) is called the sample space, \(\Sigma\) is called the event space, and \(\mu\) (or more commonly \(P\)) is the probability measure. The probability axioms\(^1\) refer to the non-negativity and countable additivity properties of measure spaces together with the requirement that \(\mu(\Omega) = 1\).

A.3 Metric Spaces

A set with a metric is called a metric space. A metric \(d\), sometimes called a distance metric, is a function that maps pairs of elements in \(X\) to non-negative real numbers such that for all \(x, y, z \in X\):

1. \(d(x, y) = 0\) if and only if \(x = y\) (identity of indiscernibles).
2. \(d(x, y) = d(y, x)\) (symmetry).
3. \(d(x, y) \leq d(x, z) + d(z, y)\) (triangle inequality).

A.4 Normed Vector Spaces

A normed vector space consists of a vector space \(X\) and a norm \(\| \cdot \|\), which maps elements of \(X\) to non-negative real numbers such that for all scalars \(\alpha\) and vectors \(x, y \in X\):

1. \(\|x\| = 0\) if and only if \(x = 0\).
2. \(\|\alpha x\| = |\alpha| \|x\|\) (absolutely homogeneous).
3. \(\|x + y\| \leq \|x\| + \|y\|\) (triangle inequality).

The \(L_p\) norms are a commonly used set of norms parameterized by a scalar \(p \geq 1\). The \(L_p\) norm of vector \(x\) is

\[
\|x\|_p = \lim_{\rho \to p} \left( |x_1|^\rho + |x_2|^\rho + \cdots + |x_n|^\rho \right)^{\frac{1}{\rho}} \tag{A.1}
\]
$L_1: \|x\|_1 = |x_1| + |x_2| + \cdots + |x_n|$  
This metric is often referred to as the *taxicab norm*.

$L_2: \|x\|_2 = \sqrt{x_1^2 + x_2^2 + \cdots + x_n^2}$  
This metric is often referred to as the *Euclidean norm*.

$L_\infty: \|x\|_\infty = \max(|x_1|, |x_2|, \cdots, |x_n|)$  
This metric is often referred to as the *max norm, Chebyshev norm, or chessboard norm*.  
The latter name comes from the minimum number of moves a chess king needs to move between two chess squares.

Table A.1. Common $L_p$ norms.  
The illustrations show the shape of the norm contours in two dimensions. All points on the contour are equidistant from the origin under that norm.
where the limit is necessary for defining the infinity norm, $L_\infty$. Several $L_p$ norms are shown in table A.1.

Norms can be used to induce distance metrics in vector spaces by defining the metric $d(x, y) = \|x - y\|$. We can then, for example, use an $L_p$ norm to define distances.

### A.5 Positive Definiteness

A symmetric matrix $A$ is **positive definite** if $x^T Ax$ is positive for all points other than the origin. In other words, $x^T Ax > 0$ for all $x \neq 0$. A symmetric matrix $A$ is **positive semidefinite** if $x^T Ax$ is always non-negative. In other words, $x^T Ax \geq 0$ for all $x$.

### A.6 Convexity

A **convex combination** of two vectors $x$ and $y$ is the result of

$$ax + (1 - \alpha)y$$

for some $\alpha \in [0, 1]$. Convex combinations can be made from $m$ vectors,

$$w_1v^{(1)} + w_2v^{(2)} + \cdots + w_mv^{(m)}$$

with nonnegative weights $w$ that sum to one.

A **convex set** is a set for which a line drawn between any two points in the set is entirely within the set. Mathematically, a set $S$ is convex if we have

$$ax + (1 - \alpha)y \in S.$$ 

for all $x, y$ in $S$ and for all $\alpha$ in $[0, 1]$. A convex and a nonconvex set are shown in figure A.1.

A **convex function** is a bowl-shaped function whose domain is a convex set. By bowl-shaped, we mean it is a function such that any line drawn between two points in its domain does not lie below the function. A function $f$ is convex over a convex set $S$ if, for all $x, y$ in $S$ and for all $\alpha$ in $[0, 1],$

$$f(ax + (1 - \alpha)y) \leq \alpha f(x) + (1 - \alpha)f(y)$$

Convex and concave regions of a function are shown in figure A.2.

A function $f$ is **strictly convex** over a convex set $S$ if, for all $x, y$ in $S$ and $\alpha$ in $(0, 1),$

$$f(ax + (1 - \alpha)y) < \alpha f(x) + (1 - \alpha)f(y)$$
Strictly convex functions have at most one minimum, whereas a convex function can have flat regions. Examples of strict and nonstrict convexity are shown in figure A.3.

A function $f$ is concave if $-f$ is convex. Furthermore, $f$ is strictly concave if $-f$ is strictly convex.
A.7 Information Content

If we have a discrete distribution that assigns probability \( P(x) \) to value \( x \), the information content\(^3 \) of observing \( x \) is given by

\[
I(x) = -\log P(x)
\]  \hspace{1cm} (A.7)

The unit of information content depends on the base of the logarithm. We generally assume natural logarithms (with base \( e \)), making the unit nat, short for natural. In information theoretic contexts, the base is often 2, making the unit bit. We can think of this quantity as the number of bits required to transmit the value \( x \) according to an optimal message encoding when the distribution over messages follows the specified distribution.

A.8 Entropy

Entropy is an information theoretic measure of uncertainty. The entropy associated with a discrete random variable \( X \) is the expected information content:

\[
H(X) = \mathbb{E}_x[I(x)] = \sum_x P(x)I(x) = -\sum_x P(x) \log P(x)
\]  \hspace{1cm} (A.8)

where \( P(x) \) is the mass assigned to \( x \).

For a continuous distribution where \( p(x) \) is the density assigned to \( x \), the differential entropy or continuous entropy is defined to be

\[
h(X) = \int p(x)I(x) \, dx = -\int p(x) \log p(x) \, dx
\]  \hspace{1cm} (A.9)

A.9 Cross Entropy

The cross entropy of one distribution relative to another can be defined in terms of expected information content. If we have one discrete distribution with mass function \( P(x) \) and another with mass function \( Q(x) \), then the cross entropy of \( P \) relative to \( Q \) is given by

\[
H(P, Q) = -\mathbb{E}_{x \sim P} \log Q(x) = -\sum_x P(x) \log Q(x)
\]  \hspace{1cm} (A.10)

For continuous distributions with density functions \( p(x) \) and \( q(x) \), we have

\[
H(p, q) = -\int p(x) \log q(x) \, dx
\]  \hspace{1cm} (A.11)

A.10 Relative Entropy

The relative entropy or Kullback–Leibler (KL) divergence is a measure of how one probability distribution is different from a reference distribution.\footnote{Named for the two American mathematicians who introduced this measure, Solomon Kullback (1907–1994) and Richard A. Liebler (1914–2003). S. Kullback, Information Theory and Statistics. Wiley, 1959.} If $P(x)$ and $Q(x)$ are mass functions, then the KL divergence from $Q$ to $P$ is the expectation of the logarithmic differences, with the expectation using $P$:

$$D_{KL}(P \parallel Q) = \sum_x P(x) \log \frac{P(x)}{Q(x)} = -\sum_x P(x) \log \frac{Q(x)}{P(x)} \quad (A.12)$$

This quantity is only defined if the support of $P$ is a subset of that of $Q$. The summation is over the support of $P$ to avoid division by zero.

For continuous distributions with density functions $p(x)$ and $q(x)$, we have

$$D_{KL}(p \parallel q) = \int p(x) \log \frac{p(x)}{q(x)} \, dx \quad (A.13)$$

Again, this quantity is only defined if the support of $p$ is a subset of that of $q$. The integral is over the support of $p$ to avoid division by zero.

A.11 Gradient Ascent

Gradient ascent is a general approach for attempting to maximize a function $f(x)$ when $f$ is a differentiable function. We begin at a point $x$ and iteratively apply the following update rule:

$$x \leftarrow x + \alpha \nabla f(x) \quad (A.14)$$

where $\alpha > 0$ is called a step factor. The idea of this optimization approach is that we take steps in the direction of the gradient until reaching a local maximum. There is no guarantee that we will find a global maximum using this method. Small values for $\alpha$ will generally require more iterations to come close to a local maximum. Large values for $\alpha$ will often result in bouncing around the local optimum without quite reaching it. If $\alpha$ is constant over iterations, it is sometimes called a learning rate. Many applications involve a decaying step factor where, in addition to updating $x$ at each iteration, we also update $\alpha$ according to

$$\alpha \leftarrow \gamma \alpha \quad (A.15)$$

where $0 < \gamma < 1$ is the decay factor.
A.12 Taylor Expansion

The Taylor expansion, also called the Taylor series, of a function is important to many approximations used in this book. From the first fundamental theorem of calculus, we know that

\[ f(x + h) = f(x) + \int_0^h f'(x + a) \, da \]  \hspace{1cm} (A.16)

Nesting this definition produces the Taylor expansion of \( f \) about \( x \):

\[
\begin{align*}
f(x + h) &= f(x) + \int_0^h \left( f'(x) + \int_0^a f''(x + b) \, db \right) \, da \\
&= f(x) + f'(x)h + \int_0^h \int_0^a f''(x + b) \, db \, da \\
&= f(x) + f'(x)h + \int_0^h \left( f''(x) + \int_0^b f'''(x + c) \, dc \right) \, db \, da \\
&= f(x) + f'(x)h + \frac{f''(x)}{2!} h^2 + \int_0^h \int_0^a \int_0^b f'''(x + c) \, dc \, db \, da \\
&\vdots \\
&= f(x) + \frac{f'(x)}{1!} h + \frac{f''(x)}{2!} h^2 + \frac{f'''(x)}{3!} h^3 + \ldots \\
&= \sum_{n=0}^{\infty} \frac{f^{(n)}(x)}{n!} h^n \hspace{1cm} (A.22)
\end{align*}
\]

In the formulation above, \( x \) is typically fixed and the function is evaluated in terms of \( h \). It is often more convenient to write the Taylor expansion of \( f(x) \) about a point \( a \) such that it remains a function of \( x \):

\[ f(x) = \sum_{n=0}^{\infty} \frac{f^{(n)}(a)}{n!} (x - a)^n \hspace{1cm} (A.24) \]

The Taylor expansion represents a function as an infinite sum of polynomial terms based on repeated derivatives at a single point. Any analytic function can be represented by its Taylor expansion within a local neighborhood.

A function can be locally approximated by using the first few terms of the Taylor expansion. Figure A.4 shows increasingly better approximations for \( \cos(x) \) about \( x = 1 \). Including more terms increases the accuracy of the local approximation, but error still accumulates as one moves away from the expansion point.
### A.13 Monte Carlo Estimation

Monte Carlo estimation allows us to evaluate the expectation of a function $f$ when its input $x$ follows a probability density function $p$:

$$\mathbb{E}_{x \sim p}[f(x)] = \int f(x)p(x) \, dx \approx \frac{1}{n} \sum_i f(x^{(i)}) \quad \text{(A.28)}$$

A linear Taylor approximation uses the first two terms of the Taylor expansion:

$$f(x) \approx f(a) + f'(a)(x - a) \quad \text{(A.25)}$$

A quadratic Taylor approximation uses the first three terms:

$$f(x) \approx f(a) + f'(a)(x - a) + \frac{1}{2} f''(a)(x - a)^2 \quad \text{(A.26)}$$

and so on.

In multiple dimensions, the Taylor expansion about $a$ generalizes to

$$f(x) = f(a) + \nabla f(a)^\top (x - a) + \frac{1}{2} (x - a)^\top \nabla^2 f(a)(x - a) + \ldots \quad \text{(A.27)}$$

The first two terms form the tangent plane at $a$. The third term incorporates local curvature. This text will use only the first three terms shown here.
where $x^{(1)}, \ldots, x^{(n)}$ are drawn from $p$. The variance of the estimate is equal to $\text{Var}_{x \sim p}[f(x)]/n$.

### A.14 Importance Sampling

Importance sampling allows us to compute $E_{x \sim p}[f(x)]$ from samples drawn from a different distribution $q$:

$$E_{x \sim p}[f(x)] = \int f(x) p(x) \, dx \quad (A.29)$$

$$= \int f(x) p(x) \frac{q(x)}{q(x)} \, dx \quad (A.30)$$

$$= \int f(x) p(x) \frac{p(x)}{q(x)} q(x) \, dx \quad (A.31)$$

$$= E_{x \sim q} \left[ f(x) \frac{p(x)}{q(x)} \right] \quad (A.32)$$

The equation above can be approximated using samples $x^{(1)}, \ldots, x^{(n)}$ drawn from $q$:

$$E_{x \sim p}[f(x)] = E_{x \sim q} \left[ f(x) \frac{p(x)}{q(x)} \right] \approx \frac{1}{n} \sum_i f(x^{(i)}) \frac{p(x^{(i)})}{q(x^{(i)})} \quad (A.33)$$

### A.15 Contraction Mappings

A contraction mapping $f$ is defined with respect to a function over a metric space such that

$$d(f(x), f(y)) \leq \alpha d(x, y) \quad (A.34)$$

where $d$ is the distance metric associated with the metric space and $0 \leq \alpha < 1$. A contraction mapping thus reduces the distance between any two members of a set. Such a function is sometimes referred to as a contraction or contractor.

A consequence of repeatedly applying a contraction mapping is that the distance between any two members of the set is driven to 0. The contraction mapping theorem or the Banach fixed-point theorem states that every contraction mapping on a complete, non-empty metric space has a unique fixed point. Furthermore, for any element $x$ in that set, repeated application of a contraction mapping to that element results in convergence to that fixed point.

---

6 A complete metric space is one where every Cauchy sequence in that space converges to a point in that space. A sequence $x_1, x_2, \ldots$ is Cauchy, if for every positive real number $\epsilon > 0$ there is a positive integer $n$ such that for all positive integers $i, j > n$, we have $d(x_i, x_j) < \epsilon$. 
Showing that a function $f$ is a contraction mapping on a metric space is useful in various convergence proofs associated with the concepts presented earlier. For example, we can show that the Bellman operator is a contraction mapping on the space of value functions with the max-norm. Application of the contraction mapping theorem allows us to prove that repeated application of the Bellman operator results in convergence to a unique value function. Example A.1 shows a simple example of a contraction mapping.

Consider the function $f(x) = [x_2/2 + 1, x_1/2 + 1/2]$. We can show $f$ is a contraction mapping for the set $\mathbb{R}^2$ and the Euclidean distance function:

$$
\begin{align*}
    d(f(x), f(y)) &= \|f(x) - f(y)\|_2 \\
    &= \|[x_2/2 + 1, x_1/2 + 1/2] - [y_2/2 + 1, y_1/2 + 1/2]\|_2 \\
    &= \|\left[\frac{1}{2}(x_2 - y_2), \frac{1}{2}(x_1 - y_1)\right]\|_2 \\
    &= \frac{1}{2} \|\left[(x_2 - y_2), (x_1 - y_1)\right]\|_2 \\
    &= \frac{1}{2} d(x, y)
\end{align*}
$$

We can plot the effect of repeated applications of $f$ to points in $\mathbb{R}^2$ and show how they converge toward $[5/3, 4/3]$.
A.16  Graphs

A graph $G = (V, E)$ is defined by a set of nodes (also called vertices) $V$ and edges $E$. Figure A.5 shows an example of a graph. An edge $e \in E$ is a pair of nodes $(v_i, v_j)$. We focus primarily on directed graphs, where edges are directed and define parent-child relationships. An edge $e = (v_i, v_j)$ is often represented graphically as an arrow from $v_i$ to $v_j$ with $v_i$ as the parent and $v_j$ as the child. If there is an edge connecting $v_i$ and $v_j$, then we say that $v_i$ and $v_j$ are neighbors. The set of all parents of a node $v_i$ is denoted $\text{Pa}(v_i)$.

A path from node $v_i$ to node $v_j$ is a sequence of edges connecting $v_i$ to $v_j$. If this path can be followed from node to node along the direction of the edges, then we can call this path a directed path. A directed path is a path without regard to the direction of the edges. A node $v_j$ is a descendant of $v_i$ if there exists a directed path from $v_i$ to $v_j$. A cycle is a directed path from a node to itself. If a graph does not contain any cycles, it is acyclic.

Figure A.5. An example graph. Here, $\text{Pa}(C) = \{A, B\}$. The sequence $(A, C, E, F)$ is a directed path, and $(A, C, B)$ is an undirected path. Node $A$ is a parent of $C$ and $D$. Node $E$ is a descendant of $B$. Neighbors of $C$ include $A$, $B$, and $E$. 
B Probability Distributions

This appendix summarizes several different families of probability distributions relevant to the topics introduced in this book. The distributions are represented by either probability mass functions or probability density functions, and the relevant functions are provided along with the parameters that govern each distribution. Plots show how the various parameters influence the distribution. The index includes page references to where these distributions are used in the body of the book. Some distributions are univariate, meaning they are distributions over a scalar variable; others are multivariate, meaning they are distributions over multiple variables.

<table>
<thead>
<tr>
<th>Name</th>
<th>Parameters</th>
<th>Distribution Function</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Uniform</strong></td>
<td>$a$</td>
<td>$b$</td>
</tr>
<tr>
<td>$U(a, b)$</td>
<td>lower bound</td>
<td>upper bound</td>
</tr>
</tbody>
</table>

$\begin{array}{cc}
\hline
a & b \\
\hline
-1 & 1 \\
0 & 3 \\
5 & 8 \\
-6 & -5 \\
\end{array}$

\[ p(x) = \frac{1}{b-a} \]

\[ x \in (a, b) \]

<table>
<thead>
<tr>
<th>Name</th>
<th>Parameters</th>
<th>Distribution Function</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Gaussian</strong></td>
<td>$\mu$</td>
<td>$\sigma^2$</td>
</tr>
<tr>
<td>$(\text{univariate})$</td>
<td></td>
<td></td>
</tr>
<tr>
<td>$N(\mu, \sigma^2)$</td>
<td>mean</td>
<td>variance</td>
</tr>
</tbody>
</table>

$\begin{array}{cc}
\hline
\mu & \sigma^2 \\
\hline
0 & 1 \\
0 & 3 \\
5 & 4 \\
-3 & 2 \\
\end{array}$

\[ p(x) = \frac{1}{\sqrt{2\pi}} \phi \left( \frac{x-\mu}{\sigma} \right) \]

\[ x \in \mathbb{R} \]

\[ \phi(x) = \frac{1}{\sqrt{2\pi}} \exp(-x^2/2) \]

**Appendix B. Probability Distributions**

**Beta**

\[ \text{Beta}(\alpha, \beta) \]

- \(\alpha > 0\) shape
- \(\beta > 0\) shape

\[
p(x) = \frac{\Gamma(\alpha + \beta)}{\Gamma(\alpha) \Gamma(\beta)} x^{\alpha-1} (1 - x)^{\beta-1}
\]

with \(x \in (0, 1)\)

\[ p(x) \]

\[ x \]

![Beta Distribution](image)

\[ \alpha = 1, \beta = 1 \]

\[ \alpha = 5, \beta = 5 \]

\[ \alpha = 2, \beta = 5 \]

\[ \alpha = 1, \beta = 2 \]

---

**Gaussian (multivariate)**

\[ \mathcal{N}(\mu, \Sigma) \]

- \(\mu\) mean
- \(\Sigma\) covariance

\[
p(x) = \frac{1}{(2\pi)^{n/2} |\Sigma|^{1/2}} \exp \left( -\frac{1}{2} (x - \mu)^\top \Sigma^{-1} (x - \mu) \right) \quad \text{where} \ n = \dim(x)
\]

with \(x \in \mathbb{R}^n\)

\[ x_1 \]

\[ x_2 \]

\[ -10 -5 0 5 10 \]

\[ -10 -5 0 5 10 \]

\[ -10 -5 0 5 10 \]

\[ \mu = [0, 0], \Sigma = [1 0; 0 1] \]

\[ \mu = [0, 5], \Sigma = [3 0; 0 3] \]

\[ \mu = [3, 3], \Sigma = [4 2; 2 4] \]

---

**Dirichlet**

\[ \text{Dir}(\alpha) \]

- \(\alpha > 0\) concentration

\[
p(x) = \frac{\Gamma(\alpha_0)}{\prod_{i=1}^{n} \Gamma(\alpha_i)} \prod_{i=1}^{n} x_i^{\alpha_i - 1} \quad \text{where} \ \alpha_0 = \sum_{i} \alpha_i
\]

with \(x_i \in (0, 1)\) and \(\sum x_i = 1\)

\[ x_1 \]

\[ x_2 \]

\[ x_3 \]

\[ 0 0.2 0.4 0.6 0.8 1 \]

\[ 0 0.2 0.4 0.6 0.8 1 \]

\[ 0 0.2 0.4 0.6 0.8 1 \]

\[ \alpha = [1, 1, 1] \]

\[ \alpha = [5, 5, 5] \]

\[ \alpha = [2, 1, 5] \]

---

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2021-03-07 23:12:00-08:00, revision 80bf4d6, comments to bugs@algorithmsbook.com
C Computational Complexity

When discussing various algorithms, it is useful to analyze their computational complexity, which refers to the resources required to run them to completion.\(^1\) We are generally interested in either time or space complexity. This appendix reviews asymptotic notation, which is what is generally used to characterize complexity. We then review a few of the complexity classes that are relevant to the algorithms in the book and discuss the problem of decidability.

C.1 Asymptotic Notation

Asymptotic notation is often used to characterize the growth of a function. This notation is sometimes called big-Oh notation, since the letter $O$ is used because the growth rate of a function is often called its order. This notation can be used to describe the error associated with a numerical method or the time or space complexity of an algorithm. This notation provides an upper bound on a function as its argument approaches a certain value.

Mathematically, if $f(x) = O(g(x))$ as $x \to a$ then the absolute value of $f(x)$ is bounded by the absolute value of $g(x)$ times some positive and finite $c$ for values of $x$ sufficiently close to $a$:

$$|f(x)| \leq c|g(x)| \quad \text{for } x \to a \quad (C.1)$$

Writing $f(x) = O(g(x))$ is a common abuse of the equal sign. For example, $x^2 = O(x^2)$ and $2x^2 = O(x^2)$, but, of course, $x^2 \neq 2x^2$. In some mathematical texts, $O(g(x))$ represents the set of all functions that do not grow faster than $g(x)$. For example, $5x^2 \in O(x^2)$. Example C.1 demonstrates asymptotic notation.

If $f(x)$ is a linear combination\(^2\) of terms, then $O(f)$ corresponds to the order of the fastest growing term. Example C.2 compares the orders of several terms.

\(^1\) The analysis of algorithms represents a large field within computer science. For an introductory textbook, see O. Goldreich, Computational Complexity: A Conceptual Perspective. Cambridge University Press, 2008. A rigorous treatment requires the introduction of concepts and computational models, such as Turing machines, that we will bypass in our discussion here.

\(^2\) A linear combination is a weighted sum of terms. If the terms are in a vector $x$, then the linear combination is $w_1x_1 + w_2x_2 + \cdots = w^Tx$. 

Consider \( f(x) = 10^6 e^x \) as \( x \to \infty \). Here, \( f \) is a product of constant \( 10^6 \) and \( e^x \). The constant can simply be incorporated into the bounding constant \( c \):

\[
|f(x)| \leq c|g(x)| \\
10^6|e^x| \leq c|g(x)| \\
|e^x| \leq c|g(x)|
\]

Thus, \( f = O(e^x) \) as \( x \to \infty \).

Consider \( f(x) = \cos(x) + x + 10x^{3/2} + 3x^2 \). Here, \( f \) is a linear combination of terms. The terms \( \cos(x) \), \( x \), \( x^{3/2} \), \( x^2 \) are arranged in order of increasing value as \( x \) approaches infinity. We plot \( f(x) \) along with \( c|g(x)| \), where \( c \) has been chosen for each term such that \( c|g(x = 2)| \) exceeds \( f(x = 2) \).

There is no constant \( c \) such that \( f(x) \) is always less than \( c|x^{3/2}| \) for sufficiently large values of \( x \). The same is true for \( \cos(x) \) and \( x \).

We find that \( f(x) = O(x^3) \), and in general \( f(x) = O(x^m) \) for \( m \geq 2 \), along with other function classes like \( f(x) = e^x \). We typically discuss the order that provides the tightest upper bound. Thus, \( f = O(x^2) \) as \( x \to \infty \).
C.2 Time Complexity Classes

The difficulty of solving certain problems can be grouped into different time complexity classes. Important classes that appear frequently throughout this book include:

- **P**: problems that can be solved in polynomial time,
- **NP**: problems whose solutions can be verified in polynomial time,
- **NP-hard**: problems that are at least as hard as the hardest problems in NP, and
- **NP-complete**: problems that are both NP-hard and in NP.

Formal definitions of these complexity classes are rather involved. It is generally believed that \( P \neq NP \), but it has not been proven and remains one of the most important open problems in mathematics. In fact, modern cryptography depends on the fact that there are no known efficient (i.e., polynomial time) algorithms for solving NP-hard problems. Figure C.1 illustrates the relationship between the complexity classes under the assumption that \( P \neq NP \).

A common approach to proving whether a particular problem \( Q \) is NP-hard is to come up with a polynomial transformation from a known NP-complete problem \( Q' \) to an instance of \( Q \). The 3SAT problem is the first known NP-complete problem and is discussed in example C.3.

C.3 Space Complexity Classes

Another set of complexity classes pertain to space, referring to the amount of memory required to execute an algorithm to completion. The complexity class \( \text{PSPACE} \) contains the set of all problems that can be solved with a polynomial amount of space, without any considerations about time. There is a fundamental difference between time and space complexity in that time cannot be reused, but space can be reused. We know that \( P \) and \( NP \) are subsets of \( \text{PSPACE} \). It is not yet known, but it is suspected, that \( \text{PSPACE} \) includes problems not in \( NP \). Through polynomial time transformations, we can define the \( \text{PSPACE-hard} \) and \( \text{PSPACE-complete} \) classes, just as we did with the NP-hard and NP-complete classes.

---

*Figure C.1. Complexity classes.*

3 There are many well-known NP-complete problems as surveyed in the paper by R.M. Karp, “Reducibility Among Combinatorial Problems,” in Complexity of Computer Computations, 1972.
The problem of Boolean satisfiability involves determining whether a Boolean formula is satisfiable. The Boolean formula consists of conjunctions (\( \land \)), disjunctions (\( \lor \)), and negations (\( \neg \)) involving \( n \) Boolean variables \( x_1, \ldots, x_n \). A literal is a variable \( x_i \) or its negation \( \neg x_i \). A 3SAT clause is a disjunction of up to three literals; for example, \( x_3 \lor \neg x_5 \lor x_6 \). A 3SAT formula is a conjunction of 3SAT clauses like

\[
F(x_1, x_2, x_3, x_4) = (x_1 \lor x_2 \lor x_3) \land
\]

\[
(\neg x_1 \lor \neg x_2 \lor x_3) \land
\]

\[
(x_2 \lor \neg x_3 \lor x_4)
\]

The challenge in 3SAT is to determine whether a possible assignment of truth values to variables exists that makes the formula true. In the formula above,

\[
F(\text{true}, \text{false}, \text{false}, \text{true}) = \text{true}
\]

Hence, the formula is satisfiable. Although a satisfying assignment can be easily found for some 3SAT problems, sometimes just by quick inspection, they are difficult to solve in general. One way to determine whether a satisfying assignment can be made is to enumerate the \( 2^n \) possible truth values of all the variables. Although determining whether a satisfying truth assignment exists is difficult, verification of whether a truth assignment leads to satisfaction can be done in linear time.
C.4 Decideability

An *undecidable* problem is a problem that cannot always be solved in finite time. Perhaps one of the most famous undecidable problems is the *halting problem*. The halting problem involves taking any program written in a sufficiently expressive language\(^4\) as input and deciding whether it will terminate. It was proven that there does not exist any algorithm that can perform such an analysis in general. Although algorithms exist that can correctly determine whether some programs terminate or not, there does not exist any algorithm that can determine whether any arbitrary program will terminate or not.

\(^4\) The technical requirement is that the language is *Turing complete* or *computationally universal*, meaning that it can be used to simulate any Turing machine.
Neural networks are parametric representations of nonlinear functions.\(^1\) The function represented by a neural network is differentiable, allowing gradient-based optimization algorithms such as stochastic gradient descent to optimize their parameters to better approximate desired input-output relationships.\(^2\) Neural representations can be helpful in a variety of contexts related to decision making, such as representing probabilistic models, utility functions, and decision policies. This appendix outlines several relevant architectures.

### D.1 Neural Networks

A neural network is a differentiable function \(y = f_\theta(x)\), which maps inputs \(x\) to produce outputs \(y\) and is parameterized by \(\theta\). Modern neural networks may have millions of parameters and can be used to convert inputs in the form of high-dimensional images or video into high-dimensional outputs like multidimensional classifications or speech.

The parameters of the network \(\theta\) are generally tuned to minimize a scalar loss function \(\ell(f_\theta(x), y)\) that is related to how far the network output is from the desired output. Both the loss function and the neural network are differentiable, allowing us to use the gradient of the loss function with respect to the parameterization \(\nabla_\theta \ell\) to iteratively improve the parameterization. This process is often referred to as neural network training or parameter tuning. It is demonstrated in example D.1.

Neural networks are typically trained on a dataset of input-output pairs \(D\). In this case, we tune the parameters to minimize the aggregate loss over the dataset:

\[
\arg \min_\theta \sum_{(x,y) \in D} \ell(f_\theta(x), y) \tag{D.1}
\]
Consider a very simple neural network, \( f_\theta(x) = \theta_1 + \theta_2 x \). We wish our neural network to take the square footage \( x \) of a home and predict its price \( y_{\text{pred}} \). We want to minimize the square deviation between the predicted housing price and the true housing price by the loss function \( \ell(y_{\text{pred}}, y_{\text{true}}) = (y_{\text{pred}} - y_{\text{true}})^2 \). Given a training pair, we can compute the gradient:

\[
\nabla_\theta \ell(f(x), y_{\text{true}}) = \nabla_\theta ((\theta_1 + \theta_2 x) - y_{\text{true}})^2 = \begin{bmatrix} 2(\theta_1 + \theta_2 x - y_{\text{true}}) \\ 2(\theta_1 + \theta_2 x - y_{\text{true}}) x \end{bmatrix}
\]

If our initial parameterization were \( \theta = [10,000, 123] \) and we had the input-output pair \( (x = 2,500, y_{\text{true}} = 360,000) \), then the loss gradient would be \( \nabla_\theta \ell = [-85,000, -2.125 \times 10^8] \). We would take a small step in the opposite direction to improve our function approximation.

Datasets for modern problems tend to be very large, making the gradient of equation (D.1) expensive to evaluate. It is common to sample random subsets of the training data in each iteration, using these batches to compute the loss gradient. In addition to reducing computation, computing gradients with smaller batch sizes also introduces some stochasticity to the gradient, which helps training avoid getting stuck in local minima.

### D.2 Feedforward Networks

Neural networks are typically constructed to pass the input through a series of layers.\(^3\) Networks with many layers are often called deep. In feedforward networks, each layer applies an affine transform followed by a nonlinear activation function applied elementwise:\(^4\)

\[
x' = \phi(Wx + b)
\]

where matrix \( W \) and vector \( b \) are parameters associated with the layer. A fully connected layer is shown in figure D.1. The dimension of the output layer is different from that of the input layer when \( W \) is non-square. Figure D.2 shows a more compact depiction of the same network.

---


Without activation functions between them, multiple successive affine transformations can be collapsed into a single, equivalent affine transform:

$$W_2(W_1x + b_1) + b_2 = W_2W_1x + (W_2b_1 + b_2) \quad (D.3)$$

These nonlinearities are necessary to allow neural networks to adapt to fit arbitrary target functions. To illustrate, figure D.3 shows the output of a neural network trained to approximate a nonlinear function.

There are many different types of activation functions that are commonly used. Similar to their biological inspiration, they tend to be close to zero when their input is low and large when their input is high. Some common activation functions are given in figure D.5.

Sometimes special layers are incorporated to achieve certain effects. For example, in figure D.4, we used a softmax layer at the end in order to force the output to represent a two-element categorical distribution. The softmax function applies...
Figure D.4. A simple two-layer fully connected network trained to classify whether a given coordinate lies within a circle (shown in white). The nonlinearities allow neural networks to form complicated, nonlinear decision boundaries.

Figure D.5. Several common activation functions.
the exponential function to each element, which ensures they are positive, and then renormalizes the resulting values:

$$\text{softmax}(x)_i = \frac{\exp(x_i)}{\sum_j \exp(x_j)}$$ \hspace{1cm} (D.4)

Gradients for neural networks are typically computed using reverse accumulation.\(^5\) The method begins with a forward step, in which the neural network is evaluated using all input parameters. In the backward step, the gradient of each term of interest is computed working from the output back to the input. Reverse accumulation makes use of the chain rule for derivatives:

$$\frac{\partial f(g(h(x)))}{\partial x} = \frac{\partial f(g(h))}{\partial h} \frac{\partial h(x)}{\partial x} = \left( \frac{\partial f(g)}{\partial g} \frac{\partial g(h)}{\partial h} \right) \frac{\partial h(x)}{\partial x}$$ \hspace{1cm} (D.5)

Example D.2 demonstrates this process. Many deep learning packages compute gradients using such automatic differentiation techniques.\(^6\) Users rarely have to provide their own gradients.

### D.3 Parameter Regularization

Neural networks are typically underdetermined, meaning there are multiple parameter instantiations that can result in the same optimal training loss.\(^7\) It is common to use parameter regularization, also called weight regularization, to introduce an additional term to the loss function that penalizes large parameter values. Regularization also helps prevent overfitting, which occurs when a network over-specializes to the training data but fails to generalize to unseen data.

Regularization typically has the form of an \(L^2\)-norm of the parameterization vector:

$$\arg\min_{\theta} \sum_{(x,y) \in D} \ell(f_\theta(x), y) - \beta \|\theta\|^2$$ \hspace{1cm} (D.6)

where the positive scalar \(\beta\) controls the strength of the parameter regularization. The scalar is often quite small, with values as low as \(10^{-6}\), to minimize the degree to which matching the training set is sacrificed by introducing regularization.


\(^7\) For example, suppose we have a neural network with a final softmax layer. The inputs to that layer can be scaled while producing the same output, and therefore the same loss.
Recall the neural network and loss function from example D.1. Below we have drawn the computational graph for the loss calculation.

Reverse accumulation begins with a forward pass in which the computational graph is evaluated. We will again use \( \theta = [10,000, 123] \) and the input-output pair \((x = 2,500, y_{\text{true}} = 360,000)\):

The gradient is then computed by working back up the tree:

Finally, we compute:

\[
\frac{\partial \ell}{\partial \theta_1} = \frac{\partial \ell}{\partial c_2} \frac{\partial c_2}{\partial y_{\text{pred}}} \frac{\partial y_{\text{pred}}}{\partial c_1} \frac{\partial c_1}{\partial \theta_1} = -85,000 \cdot 1 \cdot 1 = -85,000
\]

\[
\frac{\partial \ell}{\partial \theta_2} = \frac{\partial \ell}{\partial c_2} \frac{\partial c_2}{\partial y_{\text{pred}}} \frac{\partial y_{\text{pred}}}{\partial c_1} \frac{\partial c_1}{\partial \theta_2} = -85,000 \cdot 1 \cdot 1 \cdot 2500 = -2.125 \times 10^8
\]
D.4 Convolutional Neural Networks

Neural networks may have images or other multi-dimensional structures such as lidar scans as inputs. Even a relatively small $256 \times 256$ RGB image (similar to figure D.6) has $256 \times 256 \times 3 = 196,608$ entries. Any fully connected layer taking an $m \times m \times 3$ image as input and producing a vector of $n$ outputs would have a weight matrix with $3m^2n$ values. The large number of parameters to learn is not only computationally expensive, it is also wasteful. Information in images is typically translation-invariant; an object in an image that is shifted right by one pixel should produce a similar, if not identical, output.

Convolutional layers\(^8\) both significantly reduce the amount of computation and support translation-invariance by sliding a smaller fully connected window to produce their output. Significantly fewer parameters need to be learned. These parameters tend to be receptive to local textures in much the same way that the neurons in the visual cortex respond to stimuli in their receptive fields.

The convolutional layer consists of a set of features, or kernels, each of which is equivalent to a fully connected layer to which one can input a smaller region of the input tensor. A single kernel is shown being applied once in figure D.7. These features have full depth, meaning if an input tensor is $n \times m \times d$, the features will also have a third-dimension of $d$. The features are applied many times by sliding them over the input in both the first and second dimensions. If the stride is $1 \times 1$, then all $k$ filters are applied to every possible position and the output dimension will be $n \times m \times k$. If the stride is $2 \times 2$, then the filters are shifted by 2 in the first and second dimensions with every application, resulting in an output of size $n/2 \times m/2 \times k$. It is common for convolutional neural networks to increase in the third dimension and reduce in the first two dimensions with each layer.
Convolutional layers are translation-invariant because each filter behaves the same independently of where in the input is applied. This property is especially useful in spatial processing because shifts in an input image can yield similar outputs, making it easier for neural networks to extract common features. Individual features tend to learn how to recognize local attributes such as colors and textures.

The MNIST dataset contains handwritten digits in the form of $28 \times 28$ monochromatic images. It is often used to test image classification networks. To the right, we have a sample convolutional neural network that takes an MNIST image as input and produces a categorical probability distribution over the ten possible digits. Convolutional layers are used to efficiently extract features. The model shrinks in the first two dimensions and expands in the third dimension (the number of features) as the network depth increases. Eventually reaching a first and second dimension of 1 ensures that information from across the entire image can affect every feature. The flatten operation takes the $1 \times 1 \times 32$ input and flattens it into a 32-component output. Such operations are common when transitioning between convolutional and fully connected layers. This model has 19,722 parameters. The parameters can be tuned to maximize the likelihood of the training data.

D.5 Recurrent Networks

The neural network architectures we have discussed so far are ill-suited for temporal or sequential inputs. Operations on sequences occur when processing images from videos, when translating a sequence of words, or when tracking time-series data. In such cases, the outputs depend on more than just the most recent input. In addition, the neural network architectures discussed so far do not naturally produce variable-length outputs. For example, a neural network that writes an essay would be difficult to train using a conventional fully connected neural network.

When a neural network has sequential input, sequential output, or both (figure D.8), we can use a recurrent neural network to act over multiple iterations. These neural networks maintain a recurrent state $r$, sometimes called its memory, to retain information over time. For example, in translation, a word used early in a sentence may be relevant to the proper translation of words later in the sentence. Figure D.9 shows the structure of a basic recurrent neural network, and how the same neural network can be understood to be a larger network unrolled in time.

This unrolled structure can be used to produce a rich diversity of sequential neural networks, as shown in figure D.10. Many-to-many structures come in multiple forms. In one form, the output sequence begins with the input sequence. In another form, the output sequence does not begin with the input sequence. When using variable-length outputs, the neural network output itself often indicates when a sequence begins or ends. The recurrent state is often initialized to zero, as are extra inputs after the input sequence has been passed in, but this need not be the case.
Figure D.9. A recurrent neural network (left) and the same recurrent neural network unrolled in time (right). These networks maintain a recurrent state $r$ that allows the network to develop a sort of memory, transferring information across iterations.

Figure D.10. A recurrent neural network can be unrolled in time to produce different relationships. Unused or default inputs and outputs are grayed out.
Recurrent neural networks with many layers, unrolled over multiple time steps, effectively produce a very deep neural network. During training, gradients are computed with respect to the loss function. The contribution of layers farther from the loss function tends to be smaller than that of layers close to the loss function. This leads to the \textit{vanishing gradient} problem, in which deep neural networks have vanishingly small gradients in their upper layers. These small gradients slow training.

Very deep neural networks can also suffer from \textit{exploding gradients}, in which successive gradient contributions through the layers combine to produce very large values. Such large values make learning unstable. Example D.4 shows an example of both exploding and vanishing gradients.

To illustrate vanishing and exploding gradients, consider a deep neural network made of one-dimensional fully connected layers with relu activations. For example, if the network has three layers, its output is:

$$f_\theta(x) = \text{relu}(w_3 \text{ relu}(w_2 \text{ relu}(w_1 x_1 + b_1) + b_2) + b_3)$$

The gradient with respect to a loss function depends on the gradient of $f_\theta$.

We can get vanishing gradients in the parameters of the first layer, $w_1$ and $b_1$, if the gradient contributions in successive layers are less than one. For example, if any of the layers has a negative input to its relu function, the gradient of its inputs will be zero, so the gradient vanishes entirely. In a less extreme case, suppose the weights are all $w = 0.5 \mathbf{1}$, the offsets are all $b = 0$, and our input $x$ is positive. In this case, the gradient with respect to $w_1$ is

$$\frac{\partial f}{\partial w_1} = x_1 \cdot w_2 \cdot w_3 \cdot w_4 \cdot w_5 \ldots$$

The deeper the network, the smaller the gradient will be.

We can get exploding gradients in the parameters of the first layer if the gradient contributions in successive layers are greater than one. If we merely increase our weights to $w = 2 \mathbf{1}$, the very same gradient is suddenly doubling every layer.
While exploding gradients can often be handled with gradient clipping, regularization, and initializing parameters to small values, these solutions merely shift the problem toward that of vanishing gradients. Recurrent neural networks often use layers specifically constructed to mitigate the vanishing gradients problem. They function by selectively choosing whether to retain memory or not, and these gates help regulate the memory and the gradient. Two common recurrent layers are long-short term memory (LSTM) and gated recurrent units (GRU).

D.6 Autoencoder Networks

Neural networks are often used to process high-dimensional inputs such as images or point clouds. These high-dimensional inputs are often highly structured, with the actual information content being much lower dimensional than the high-dimensional space in which it is presented. Pixels in images tend to be highly correlated with their neighbors, and point clouds often have many regions of continuity. Sometimes we wish to build an understanding of the information content of our datasets by converting them to a (much) smaller set of features, or an embedding. This compression, or representation learning, has many advantages. Lower-dimensional representations can help facilitate the application of traditional machine learning techniques like Bayesian networks to what would otherwise be intractable. The features can be inspected to develop an understanding of the information content of the dataset, and these features can be used as inputs to other models.

An autoencoder is a neural network trained to discover a low-dimensional feature representation of a higher-level input. An autoencoder network takes in a high-dimensional input $x$ and produces an output $x'$ with the same dimensionality. We design the network architecture to pass through a lower-dimensional intermediate representation called a bottleneck. The activations $z$ at this bottleneck are our low-dimensional features, which exist in a latent space that is not explicitly observed. Such an architecture is shown in figure D.11.

We train the autoencoder to reproduce its input. For example, to encourage the output $x'$ to match $x$ as closely as possible, we may simply minimize the $L_2$-norm,

$$\min_{\theta} \mathbb{E}_{x \in D} [\|f_{\theta}(x) - x\|_2]$$  \hspace{1cm} (D.7)


\[11\] Such dimensionality reduction can also be done using traditional machine learning techniques, such as principle components analysis. Neural representations allows for more flexibility and can handle nonlinear representations.

```
x
\[\downarrow\]
\text{encoder}
\[\downarrow\]
\text{encoding } z
\[\downarrow\]
\text{decoder}
\[\downarrow\]
x'
```

Figure D.11. An autoencoder passes a high-dimensional input through a low-dimensional bottleneck and then reconstructs the original input. Minimizing reconstruction loss can result in an efficient low-dimensional encoding.
Noise is often added to the input in order to produce a more robust feature embedding:

$$\text{minimize } \theta \quad \mathbb{E}_{x \in D} [ \| f_\theta (x + \epsilon) - x \|_2 ] \quad (D.8)$$

Training to minimize the reconstruction loss forces the autoencoder to find the most efficient low-dimensional encoding sufficient to accurately reconstruct the original input. Furthermore, training is \textit{unsupervised} in that we do not need to guide the training to a particular feature set.

After training, the upper portion of the autoencoder above the bottleneck can be used as an \textit{encoder} that transforms an input into the feature representation. The lower portion of the autoencoder can be used as a \textit{decoder} that transforms the feature representation into the input representation. Decoding is useful when training neural networks to generate images or other high-dimensional outputs. Example D.5 shows an embedding learned for handwritten digits.

A \textit{variational autoencoder}, shown in figure D.12, extends the autoencoder framework to learn a probabilistic encoder.\textsuperscript{12} Rather than outputting a deterministic sample, the encoder produces a distribution over the encoding, which allows the model to assign confidence to its encoding. Multivariate Gaussian distributions with diagonal covariance matrices are often used for their mathematical convenience. In such a case, the encoder outputs both an encoding mean and diagonal covariance matrix.

Variational autoencoders are trained to both minimize the expected reconstruction loss while keeping the encoding components close to unit Gaussian. The former is achieved by taking a single sample from the encoding distribution with each passthrough, $z \sim \mathcal{N}(\mu, \sigma^T \sigma)$. For backpropagation to work, we typically include random noise $w \sim \mathcal{N}(0, I)$ as an additional input to the neural network and obtain our sample according to $z = \mu + w \odot \sigma$.

The components are kept close to unit Gaussian by additionally minimizing the KL divergence (appendix A.10).\textsuperscript{13} This objective encourages smooth latent space representations. The network is penalized for spreading out the latent representations (large values for $\| \mu \|$) and for focusing each representation into a very small encoding space (small values for $\| \sigma \|$), ensuring better coverage of the latent space. As a result, smooth variations into the decoder can result in smoothly varying outputs. This property allows decoders to be used as \textit{generative models}, where samples from a unit multivariate Gaussian can be input to the decoder to


\textsuperscript{13} The KL divergence for two unit Gaussians is

$$\log \left( \frac{\sigma_2}{\sigma_1} \right) + \frac{\sigma_1^2 + (\mu_1 - \mu_2)^2}{2\sigma_2^2} - \frac{1}{2}$$
produce realistic samples in the original space. The combined loss function is:

\[
\min_\theta \mathbb{E}_{x \in D} \left[ \|x' - x\|_2 + c \sum_{i=1}^{\mid \mu \mid} D_{KL}(\mathcal{N}(\mu_i, \sigma_i^2) || \mathcal{N}(0, 1)) \right]
\]

subject to \( \mu, \sigma = \text{encoder}(x + \epsilon) \)
\( x' = \text{decoder}(\mu + w \odot \sigma) \)

where the tradeoff between the two losses is tuned by the scalar \( c > 0 \). Example D.6 demonstrates this process on a latent space learned from handwritten digits.

Variational autoencoders are derived by representing the encoder as a conditional distribution \( q(z \mid x) \), where \( x \) belongs to the observed input space and \( z \) is in the unobserved embedding space. The decoder performs inference in the other direction, representing \( p(x \mid z) \), in which case it also outputs a probability distribution. We seek to minimize the KL divergence between \( q(z \mid x) \) and \( p(z \mid x) \), which is the same as minimizing \( \mathbb{E}[\log p(x \mid z)] - D_{KL}(q(z \mid x) || p(z)) \), where \( p(z) \) is our prior, the unit multivariate Gaussian to which we bias our encoding distribution. We thus obtain our reconstruction loss and our KL divergence.

### D.7 Adversarial Networks

We often want to train neural networks to produce high-dimensional outputs, such as images or sequences of helicopter control inputs. When the output space is large, the training data may only cover a very small region of the state space. Hence, training purely on the available data can cause unrealistic results or overfitting.

We generally want the output of a neural network to produce plausible outputs. For example, when producing images, we want the images to look realistic. When mimicking human driving such as in imitation learning (chapter 18), we want the vehicle to typically stay within its lane and to react appropriately to other drivers.
We can use an autoencoder to train an embedding for the MNIST dataset. In this example, we use an encoder similar to the convolutional network in example D.3, except with a two-dimensional output and no softmax layer. We construct a decoder that mirrors the encoder and train the full network to minimize the reconstruction loss. Below are the encodings for 10,000 images from the MNIST dataset after training. Each encoding is colored according to the corresponding digit.

![Example D.5. A visualization of a two-dimensional embedding learned for the MNIST digits.](image)

We find that the digits tend to be clustered into regions that are roughly radially distributed from the origin. Note how the encodings for 1 and 7 are similar, as the two digits look alike. Recall that training is unsupervised, and the network is not given any information about the digit values. Nevertheless these clusterings are produced.
In example D.5, we trained an autoencoder on the MNIST dataset. We can adapt the same network to produce two-dimensional mean and variance vectors at the bottleneck instead of a two-dimensional embedding, and then train it to minimize both the reconstruction loss and the KL divergence. Below we show the mean encodings for the same 10,000 images for the MNIST dataset. Each encoding is again colored according to the corresponding digit.

The variational autoencoder also produces clusters in the embedding space for each digit, but this time they are roughly distributed according to a zero-mean, unit variance Gaussian distribution. We again see how some encodings are similar, such as the significant overlap for 4 and 9.
One common approach to penalize off-nominal outputs or behavior is to use \textit{adversarial learning} by including a \textit{discriminator} as shown in figure D.13.\footnote{These techniques were introduced by I. Goodfellow, J. Pouget-Abadie, M. Mirza, B. Xu, D. Warde-Farley, S. Ozair, et al., “Generative Adversarial Nets,” in \textit{Advances in Neural Information Processing Systems (NIPS), 2014.}} A discriminator is a neural network that acts as a binary classifier that takes in neural network outputs and learns to distinguish between real outputs from the training set and the outputs from the primary neural network. The primary neural network, also called a \textit{generator}, is then trained to deceive the discriminator, thereby naturally producing outputs that are more difficult to distinguish from the dataset. The primary advantage of this technique is that we do not need to architect special features to identify or quantify how the output fails to match the training data, but can allow the discriminator to naturally learn such differences.

Learning is adversarial in the sense that we have two neural networks: the primary neural network that we would like to produce realistic outputs and the discriminator network that distinguishes between primary network outputs and real examples. They are each training to outperform the other. Training is an iterative process where each network is improved in turn. It can sometimes be challenging to balance their relative performance; if one network becomes too good, the other can become stuck.

![Figure D.13. A generative adversarial network causes a primary network’s output to be more realistic by using a discriminator to force the primary network to produce more realistic output.](image)
E  Search Algorithms

A search problem is concerned with finding an appropriate sequence of actions to maximize obtained reward over subsequent deterministic transitions. Search problems are Markov decision processes (covered in part II) with deterministic transition functions. Some well known search problems include sliding tile puzzles, the Rubik’s Cube, Sokoban, and finding the shortest path to a destination.

E.1 Search Problems

In a search problem, we choose action $a_t$ at time $t$ based on observing state $s_t$, and then receive a reward $r_t$. The action space $A$ is the set of possible actions, and the state space $S$ is the set of possible states. Some of the algorithms assume these sets are finite, but this is not required in general. The state evolves deterministically and depends only on the current state and action taken. We use $A(s)$ to denote the set of valid actions from state $s$. When there are no valid actions, the state is considered to be absorbing and yields zero reward for all future timesteps. Goal states, for example, are typically absorbing.

The deterministic state transition function $T(s, a)$ gives the successor state $s'$. The reward function $R(s, a)$ gives the reward received when executing action $a$ from state $s$. Search problems typically do not include a discount factor $\gamma$ that penalizes future rewards. The objective is to choose a sequence of actions that maximizes the sum of rewards or return. Algorithm E.1 provides a data structure for representing search problems.
Appendix E. Search Algorithms

### Struct Search

```plaintext
struct Search
    S  # state space
    A  # valid action function
    T  # transition function
    R  # reward function
end
```

#### E.2 Search Graphs

A search problem with finite state and action spaces can be represented as a search graph. The nodes correspond to states, and edges correspond to transitions between states. Associated with each edge from a source to a destination state is both an action that results in that state transition and the expected reward when taking that action from the source state. Figure E.1 depicts a subset of such a search graph for a $3 \times 3$ sliding tile puzzle.

Many graph search algorithms conduct a search from an initial state and fan out from there. In so doing, these algorithms trace out a search tree. The initial state is the root node, and any time we transition from $s$ to $s'$ during search, an edge from $s$ to a new node $s'$ is added to the search tree. A search tree for the same sliding tile puzzle is shown in figure E.2.

#### E.3 Forward Search

Perhaps the simplest graph search algorithm is forward search (algorithm E.2), which determines the best action to take from an initial state $s$ by looking at all possible action-state transitions up to a depth (or horizon) $d$. At depth $d$, the algorithm uses an estimate of the value of the state $U(s)$.

The approximate value functions in this chapter are expected to return 0 when in a state with no available actions.
E.4 Branch and Bound

A general method known as branch and bound (algorithm E.3) can significantly reduce computation by using domain information about the upper and lower
bounds on expected reward. The upper bound on the return from taking action $a$ from state $s$ is $\overline{Q}(s,a)$. The lower bound on the return from state $s$ is $\underline{U}(s)$. Branch and bound follows the same procedure as depth-first-search, but iterates over the actions according to their upper bound, and only proceeds to a successor node if the best possible value it could return is lower than what has already been discovered by following an earlier action. Branch and bound search is compared to forward search in example E.1.

Algorithm E.3. The branch and bound search algorithm for finding an approximately optimal action for a discrete search problem $\mathcal{P}$ from a current state $s$. The search is performed to depth $d$ with value function lower bound $\underline{U}$ and action value function upper bound $\overline{Q}$. The returned named tuple consists of the best action $a$ and its finite-horizon expected value $u$. 

function branch_and_bound($\mathcal{P}::\text{Search}, s, d, \underline{U}, \overline{Q}$) 
  $A, T, R = \mathcal{P}.A(s), \mathcal{P}.T, \mathcal{P}.R$
  if isempty($A$) || $d \leq 0$
    return (a=nothing, u=$\underline{U}(s)$)
  end
  best = (a=nothing, u=-Inf)
  for $a$ in sort($A$, by=$a$->$\overline{Q}(s,a)$, rev=true)
    if $\overline{Q}(s,a) \leq$ best.$u$
      return best # safe to prune
    end
    $u = R(s,a) + \text{branch_and_bound}(\mathcal{P},T(s,a),d-1,\underline{U},\overline{Q}).u$
    if $u >$ best.$u$
      best = ($a=a$, $u=u$)
    end
  end
  return best
end
Consider using branch and bound on a hex world search problem. Actions in search problems cause deterministic transitions, so unlike the hex-world MDP, we always correctly transition between neighboring tiles when the corresponding action is taken.

The circle indicates the start state. All transitions incur a reward of $-1$. The blue colored tile is terminal, and produces reward $5$ when entered.

Above we show the search trees for both forward search and branch and bound to depth $4$. For branch and bound we used a lower bound $\underline{U}(s) = -6$ and an upper bound $\overline{Q}(s,a) = 5 - \delta(T(s,a))$, where the function $\delta(s)$ is the minimum number of steps from the given state to the terminal reward state. The search tree of branch and bound is a subset of that of forward search because branch and bound can ignore portions it knows are not optimal.

Due to the upper bound, branch and bound evaluates moving right first, and because that happens to be optimal, it is able to immediately identify the optimal sequence of actions and avoid exploring other actions. If the start and goal states were reversed, the search tree would be larger. In the worst case, it can be as large as forward search.

Example E.1. A comparison of the savings that branch and bound can have over forward search. Branch and bound can be significantly more efficient with appropriate bounds.
Branch and bound is not guaranteed to reduce computation over forward search. Both approaches have the same worst-case time complexity. The efficiency of the algorithm greatly depends on the heuristic.

### E.5 Dynamic Programming

Neither forward search nor branch and bound remember whether a state has been previously visited; they waste computational resources by evaluating these states multiple times. Dynamic programming (algorithm E.4) avoids duplicate effort by remembering when a particular subproblem has been previously solved. Dynamic programming can be applied to problems in which an optimal solution can be constructed from optimal solutions of its subproblems, a property called *optimal substructure*. For example, if the optimal sequence of actions from $s_1$ to $s_3$ goes through $s_2$, then the subpaths from $s_1$ to $s_2$ and from $s_2$ to $s_3$ are also optimal. This substructure is shown in figure E.4.

![Diagram](image.png)

In the case of graph search, when evaluating a state we first check a *transposition table* to see whether the state has been previously visited, and if it has, we return its stored value. Otherwise we evaluate the state as normal and store the result in the transposition table. A comparison to forward search is shown in figure E.5.

### E.6 Heuristic Search

Heuristic search (algorithm E.5) improves upon branch and bound by ordering its actions based on a provided *heuristic* function $\overline{U}(s)$, which is an upper bound of the return. Like dynamic programming, heuristic search has a mechanism by which state evaluations can be cached in order to avoid redundant computation. Furthermore, heuristic search does not require the lower bound value function needed by branch and bound.\(^3\)

\(^3\) Heuristic search is also known as *informed search* or *best-first search*.

\(^4\) Our implementation does use two value functions: the heuristic for guiding the search and an approximate value function for evaluating terminal states.
Algorithm E.4. Dynamic programming applied to forward search, which includes a transposition table \( M \). Here, \( M \) is a dictionary that stores depth-state tuples from previous evaluations, allowing the method to return previously computed results. The search is performed to depth \( d \), at which point the terminal value is estimated with an approximate value function \( U \). The returned named tuple consists of the best action \( a \) and its finite-horizon expected value \( u \).

Function `dynamic_programming(P::Search, s, d, U, M=Dict())`

```plaintext
def function dynamic_programming(P::Search, s, d, U, M=Dict())
    if haskey(M, (d,s))
        return M[(d,s)]
    end
    \( \mathcal{A}, T, R = P.A(s), P.T, P.R \)
    if isempty(\( \mathcal{A} \)) \&\& d ≤ 0
        best = (a=nothing, u=U(s))
    else
        best = (a=first(\( \mathcal{A} \)), u=-Inf)
        for a in \( \mathcal{A} \)
            s' = T(s,a)
            u = R(s,a) + dynamic_programming(P, s', d-1, U, M).u
            if u > best.u
                best = (a=a, u=u)
            end
        end
    end
    M[(d,s)] = best
    return best
end
```

Figure E.5. A comparison of the number of state evaluations for pure forward search and forward search augmented with dynamic programming on the hex-world search problem of example E.1. Dynamic programming is able to avoid the exponential growth in state visitation by caching results.
Actions are sorted based on the immediate reward plus a heuristic estimate of the future return:

\[ R(s,a) + \overline{U}(T(s,a)) \]  

(E.1)

In order to guarantee optimality, the heuristic must be both admissible and consistent. An admissible heuristic is an upper bound of the true value function. A consistent heuristic is never less than the expected reward gained by transitioning to a neighboring state:

\[ \overline{U}(s) \geq R(s,a) + \overline{U}(T(s,a)) \]  

(E.2)

The method is compared to branch and bound search in example E.2.

---

**Algorithm E.5.** The heuristic search algorithm for solving a search problem \( \mathcal{P} \) starting from state \( s \) and searching to a maximum depth \( d \). A heuristic \( \text{Uhi} \) is used to guide the search, the approximate value function \( \text{U} \) is evaluated at terminal states, and a transposition table \( M \) in the form of a dictionary containing depth-state tuples allows the algorithm to cache values from previously explored states.
We can apply heuristic search to the same hex world search problem as in example E.1. We use the heuristic $H(s) = 5 - \delta(s)$, where $\delta(s)$ is the number of steps from the given state to the terminal reward state. Below we show the number of states visited when running either branch and bound (left) or heuristic search (right) from each starting state. Branch and bound is just as efficient in states near and to the left of the goal state, whereas heuristic search is able to efficiently search from any initial state.

Example E.2. A comparison of the savings that heuristic search can have over branch and bound search. Heuristic search automatically orders actions according to their lookahead heuristic value.
F Problems

Decision making researchers use several test problems to evaluate decision making algorithms. This section covers some of the problems used throughout this book. Table F.1 summarizes some of the important properties of these problems.

| Problem                           | |S| |A| |O| |γ|
|----------------------------------|-----------------|-------------|------|------|---|---|
| Hex world                        | varies          | 6           | —    | —    | 0.9 |
| 2048                             | —               | 4           | —    | 1    |
| Cart-pole                        | (⊂ R₄)          | 2           | —    | 1    |
| Mountain car                     | (⊂ R³)          | 3           | —    | 1    |
| Simple regulator                 | (⊂ R)           | (⊂ R)       | —    | 1 or 0.9 |
| Aircraft collision avoidance     | —               | 3           | —    | 1    |
| Crying baby                      | 2               | 3           | 2    | 0.9 |
| Machine replacement              | 3               | 4           | 2    | 1    |
| Catch                            | 4               | 10          | 2    | 0.9 |
| Prisoner’s dilemma               | 2               | 2 per agent | —    | 1    |
| Rock-paper-scissors              | 2               | 3 per agent | —    | 1    |
| Traveler’s dilemma               | 2               | 99 per agent| —    | 1    |
| Predator-prey hex world          | varies          | varies      | 6 per agent | —    | 0.9 |
| Multiagent crying Baby           | 2               | 2           | 3 per agent | 2 per agent | 0.9 |
| Collaborative predator-prey hex world | varies | varies | 6 per agent | 2 per agent | 0.9 |

Table F.1. Problem summary.

F.1 Hex World

The hex world problem is a simple MDP in which we must traverse a tile map to reach a goal state. Each cell in the tile map represents a state in the MDP. We can attempt to move in any of the 6 directions. The effects of these actions are stochastic. As shown in figure F.1, we move 1 step in the specified direction with probability 0.7, and we move 1 step in one of the neighboring directions, each with probability 0.15. If we bump against the outer border of the grid, then we do not move at all, at cost 1.0.

Figure F.1. Actions in the hex world problem have probabilistic effects.
Certain cells in the hex world problem are terminal states. Taking any action in these cells gives us a specified reward and then transports us to a terminal state. No further reward is received in the terminal state. The total number of states in the hex world problem is thus the number of tiles plus 1, for the terminal state. Figure F.2 shows an optimal policy for two hex world problem configurations used throughout this text. We refer to the larger instance as hex world and to the smaller, simpler instance as straight-line hex world. The straight-line hex world formulation is used to illustrate how reward is propagated from its single reward-bearing state on the rightmost cell.

Figure F.2. The standard hex world and straight-line hex world problems. The top row shows the base problem setup and colors hexes with terminal rewards. The bottom row shows an optimal policy for each problem, colored according to the expected value, with arrows indicating the action to take in each state.

F.2 2048

The 2048 problem is based on a popular tile game. It has discrete state and action spaces. The game is played on a $4 \times 4$ board. The board is initially empty except for two tiles, each of which can have value 2 or 4. A randomly selected starting state is shown in figure F.3.

The agent can move all tiles left, down, right, or up. Choosing a direction pushes all tiles in that direction. A tile stops when it hits a wall or another tile of a different value. A tile that hits another tile of the same value merges with that tile, forming a new tile with their combined value. After shifting and merging, a new tile of value 2 or 4 is spawned in a random open space. This process is shown in figure F.4.

Figure F.3. A random starting state in the 2048 problem consists of two tiles, each with value 2 or 4.
The game ends when we can no longer shift tiles to produce an empty space. Rewards are only obtained when merging two tiles and are equal to the merge tile’s value. An example state-action transition with a merge is shown in figure F.5.

A common strategy is to choose a corner and alternate between the two actions that lead in that direction. This tends to stratify the tiles such that the larger-valued ones are in the corner and the newly spawned tiles are in the periphery.

F.3 Cart-Pole

The cart-pole problem, also sometimes called the pole balancing problem, has the agent move a cart back and forth. As shown in figure F.6, this cart has a rigid pole attached to it by a swivel, such that as the cart moves back and forth, the pole begins to rotate. The objective is to keep the pole vertically balanced while keeping the cart near within the allowed lateral bounds. As such, 1 reward is obtained each time step in which these conditions are met, and transition to a terminal zero-reward state occurs whenever they are not.

The actions are to either apply a left or right force $F$ on the cart. The state space is defined by four continuous variables: the lateral position of the cart $x$, its lateral velocity $v$, the angle of the pole $\theta$, and the pole’s angular velocity $\omega$. The problem involves a variety of parameters including the mass of the cart $m_{\text{cart}}$, the mass of the pole $m_{\text{pole}}$, the pole length $\ell$, the force magnitude $|F|$, gravitational acceleration $g$, the timestep $\Delta t$, the maximum $x$ deviation, the maximum angular deviation, and friction losses between the cart and pole or between the cart and its track.\footnote{4 We use the parameters implemented in the OpenAI Gym. G. Brockman, V. Cheung, L. Pettersson, J. Schneider, J. Schulman, J. Tang, et al., “OpenAI Gym,” 2016. arXiv: 1606.01540v1.}

Given an input force $F$, the angular acceleration of the pole is

$$\alpha = \frac{g \sin(\theta) - \tau \cos(\theta)}{\frac{\ell}{2} \left( \frac{4}{3} - \frac{m_{\text{pole}}}{m_{\text{cart}} + m_{\text{pole}}} \cos(\theta)^2 \right)}$$ (F.1)

where

$$\tau = \frac{F + \omega^2 \ell \sin \theta / 2}{m_{\text{cart}} + m_{\text{pole}}}$$ (F.2)

and the lateral cart acceleration is

$$a = \tau - \frac{\ell}{2} \alpha \cos(\theta) \frac{m_{\text{pole}}}{m_{\text{cart}} + m_{\text{pole}}}$$ (F.3)

The state is updated with Euler integration:

$$x \leftarrow x + v \Delta t$$
$$v \leftarrow v + a \Delta t$$
$$\theta \leftarrow \theta + \omega \Delta t$$
$$\omega \leftarrow \omega + \alpha \Delta t$$ (F.4)

The cart-pole problem is typically initialized with each random value drawn from $U(-0.05, 0.05)$. Rollouts are run until the lateral or angular deviations are exceeded.

---

### F.4 Mountain Car

In the mountain car problem, a vehicle must drive to the right, out of a valley. The valley walls are steep enough that blindly accelerating toward the goal with insufficient speed causes the vehicle to come to a halt and slide back down. The agent must learn to accelerate left first, in order to gain enough momentum on the return to make it up the hill.

---

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The state is the vehicle’s horizontal position \( x \in [-1.2, 0.6] \) and speed \( v \in [-0.07, 0.07] \). At any given time step, the vehicle can accelerate left (\( a = -1 \)), accelerate right (\( a = 1 \)), or coast (\( a = 0 \)). We receive \(-1\) reward every turn, and terminate when the vehicle makes it up the right side of the valley past \( x = 0.6 \). A visualization of the problem is given in figure F.7.

![Figure F.7. In the mountain car problem, a vehicle must alternate between accelerating left and right in order to power itself up a hill. The goal region is shown in blue.](image)

Transitions in the mountain car problem are deterministic:

\[
\begin{align*}
v' &\leftarrow v + 0.001a - 0.0025 \cos(3x) \\
x' &\leftarrow x + v'
\end{align*}
\]

The gravitational term in the speed update is what drives the under-powered vehicle back toward the valley floor. Transitions are clamped to the bounds of the state-space.

The mountain car problem is a good example of a problem with delayed return. Many actions are required to get to the goal state, making it difficult for an untrained agent to receive anything other than consistent unit penalties. The best learning algorithms are able to efficiently propagate knowledge from trajectories that reach the goal back to the rest of the state space.

### F.5 Simple Regulator

The simple regulator problem is a simple linear quadratic regulator problem with a single state. It is an MDP with a single real-valued state and a single real-valued
action. Transitions are linear Gaussian such that a successor state $s'$ is drawn from the Gaussian distribution $\mathcal{N}(s + a, 0.1^2)$. Rewards are quadratic, $R(s, a) = -s^2$, and do not depend on the action. The examples in this text use the initial state distribution $\mathcal{N}(0.3, 0.1^2)$.

Optimal finite-horizon policies cannot be derived using the methods from section 7.8. In this case, $T_s = [1]$, $T_a = [1]$, $R_s = [-1]$, $R_a = [0]$ and $w$ is drawn from $\mathcal{N}(0, 0.1^2)$. Applications of the Riccati equation require that $R_a$ be negative definite, which it is not.

The optimal policy is $\pi(s) = -s$, resulting in a successor state distribution centered at the origin. In the policy gradient chapters we often learn parameterized policies of the form $\pi_\theta(s) = \mathcal{N}(\theta_1 s, \theta_2^2)$. In such cases, the optimal parameterization for the simple regulator problem is $\theta_1 = -1$ and $\theta_2$ is asymptotically close to zero.

The optimal value function for the simple regulator problem is also centered about the origin, with reward decreasing quadratically:

$$U(s) = -s^2 + \frac{\gamma}{1 - \gamma} \mathbb{E}_{s \sim \mathcal{N}(0, 0.1^2)}[-s^2]$$

$$\approx -s^2 - 0.010 \frac{\gamma}{1 - \gamma}$$

### F.6 Aircraft Collision Avoidance

The aircraft collision avoidance problem involves deciding when to issue a climb or descend advisory to our aircraft to avoid an intruder aircraft. There are three actions corresponding to no advisory, commanding a $5 \text{ m/s}$ descend, and commanding a $5 \text{ m/s}$ climb. The intruder is approaching us head-on with constant horizontal closing speed. The state is specified by the altitude $h$ of our aircraft measured relative to the intruder aircraft, our vertical rate $\dot{h}$, the previous action $a_{\text{prev}}$, and the time to potential collision $t_{\text{col}}$. Figure F.8 illustrates the problem scenario.

Given action $a$, the state variables are updated as follows:

$$h \leftarrow h + \dot{h} \Delta t \quad (F.5)$$

$$\dot{h} \leftarrow \dot{h} + (\dot{h} + \nu) \Delta t \quad (F.6)$$

$$a_{\text{prev}} \leftarrow a \quad (F.7)$$

$$t_{\text{col}} \leftarrow t_{\text{col}} - \Delta t \quad (F.8)$$

---

6 This formulation is a highly simplified version of the problem described by M.J. Kochenderfer and J.P. Chryssanthacopoulos, “Robust Airborne Collision Avoidance Through Dynamic Programming,” Massachusetts Institute of Technology, Lincoln Laboratory, Project Report ATC-371, 2011.
where $\Delta t = 1$ s and $\nu$ is selected from a discrete distribution over $-2, 0, 2$ m/s$^2$ with associated probabilities 0.25, 0.5, and 0.25. The value $\ddot{h}$ is given by

$$
\ddot{h} = \begin{cases} 
0 & \text{if } a = \text{no advisory} \\
\frac{a}{\Delta t} & \text{if } |a - \dot{h}| / \Delta t < \ddot{h}_{\text{limit}} \\
\text{sign}(a - \dot{h})\ddot{h}_{\text{limit}} & \text{otherwise}
\end{cases} \quad (F.9)
$$

where $\ddot{h}_{\text{limit}} = 1$ m/s$^2$.

The episode terminates when taking an action when $t_{\text{col}} < 0$. There is a penalty of 1 when the intruder comes within 50 m when $t_{\text{col}} = 0$, and there is a penalty of 0.01 when $a \neq a_{\text{prev}}$.

The aircraft collision avoidance problem can be efficiently solved over a discretized grid using backwards induction value iteration (section 7.6) because the dynamics deterministically reduce $t_{\text{col}}$. Slices of the optimal value function and policy are depicted in figure F.9.

### F.7 Crying Baby

The *crying baby problem* is a simple POMDP with two states, three actions, and two observations. Our goal is to care for a baby, and we do so by choosing at each time step whether to feed the baby, sing to the baby, or ignore the baby.

The baby becomes hungry over time. We do not directly observe whether the baby is hungry, but instead receive a noisy observation in the form of whether or not the baby is crying. The state, action, and observation spaces are as follows:

- $S = \{\text{hungry}, \text{sated}\}$
- $A = \{\text{feed}, \text{sing}, \text{ignore}\}$
- $O = \{\text{crying}, \text{quiet}\}$

---

\[ \dot{h} = 0.0 (\text{m/s}) \quad \dot{h} = 5.0 (\text{m/s}) \]

Figure F.9. Optimal value function slices (top) and policy slices (bottom) for the aircraft collision avoidance problem. The value function and policy is symmetric about 0 when the vertical separation rate is 0, but is skewed when the vertical separation rate is non-zero. Overall our aircraft need not take action until the intruder aircraft is close.
Feeding will always sate the baby. Ignoring the baby risks a sated baby becoming hungry, and ensures that a hungry baby remains hungry. Singing to the baby is an information gathering action with the same transition dynamics as ignoring, but without the potential for crying when sated (not hungry) and with an increased chance of crying when hungry.

The transition dynamics are:

\[
T(\text{sated} \mid \text{hungry, feed}) = 100%
\]

\[
T(\text{hungry} \mid \text{hungry, sing}) = 100%
\]

\[
T(\text{hungry} \mid \text{hungry, ignore}) = 100%
\]

\[
T(\text{sated} \mid \text{sated, feed}) = 100%
\]

\[
T(\text{hungry} \mid \text{sated, sing}) = 10%
\]

\[
T(\text{hungry} \mid \text{sated, ignore}) = 10%
\]

The observation dynamics are:

\[
O(\text{cry} \mid \text{feed, hungry}) = 80%
\]

\[
O(\text{cry} \mid \text{sing, hungry}) = 90%
\]

\[
O(\text{cry} \mid \text{ignore, hungry}) = 80%
\]

\[
O(\text{cry} \mid \text{feed, sated}) = 10%
\]

\[
O(\text{cry} \mid \text{sing, sated}) = 0%
\]

\[
O(\text{cry} \mid \text{ignore, sated}) = 10%
\]

The reward function assigns $-10$ reward if the baby is hungry independent of the action taken. The effort of feeding the baby adds a further $-5$ reward, whereas singing adds $-0.5$ reward. As baby caregivers, we seek the optimal infinite horizon policy with discount factor $\gamma = 0.9$. Figure F.10 shows the optimal value function and associated policy.

![Figure F.10. The optimal policy for the crying baby problem.](image)

F.8 Machine Replacement

The machine replacement problem is a discrete POMDP in which we maintain a machine that produces products. This problem is used for its relative simplicity and the varied size and shape of the optimal policy regions. The optimal policy for certain horizons even has disjoint regions in which the same action is optimal, as shown in figure F.12.

---

8 R. D. Smallwood and E. J. Sondik, “The Optimal Control of Partially Observable Markov Processes Over a Finite Horizon,” *Operations Research*, vol. 21, no. 5, pp. 1071–1088, 1973. The original problem formulation includes *salvage values*, or *terminal rewards* that are equal to the number of working parts. We do not separately model terminal rewards in this text. Terminal rewards could be included in our framework by explicitly including the horizon in the problem state.
The machine produces products for us when it is working properly. Over time, the two primary components in the machine may break down, together or individually, leading to defective product. We can indirectly observe whether the machine is faulty by examining the products, or by directly examining the components in the machine.

The problem has states $S = \{0, 1, 2\}$, corresponding to the number of faulty internal components. There are four actions, used prior to each production cycle:

1. *manufacture*, manufacture product and do not examine the product,
2. *examine*, manufacture product and examine the product,
3. *interrupt*, interrupt production, inspect, and replace failed components, and
4. *replace* replace both components after interrupting production.

When we examine the product, we can observe whether or not it is defective. All other actions only observe non-defective products.

The components in the machine independently have a 10% chance of breaking down with each production cycle. Each failed component contributes a 50% chance of producing a defective product. A nondefective product nets 1 reward, whereas a defective product nets 0 reward. The transition dynamics assume that component breakdown is determined before a product is made, so the manufacture action on a fully-functional machine does not have a 100% chance of producing 1 reward.

The manufacture action incurs no penalty. Examining the product costs 0.25. Interrupting the line costs 0.5 to inspect the machine, causes no product to be produced, and incurs 1 for each broken component. Simply replacing both components always incurs 2, but does not have an inspection cost.

The transition, observation, and reward functions are given in table F.2. Optimal policies for increasing horizons are shown in figure F.11.

### F.9 Catch

In the *catch problem*, Johnny would like to successfully catch throws from his father, and he prefers catching longer-distance throws. However, he is uncertain about the relationship between the distances of a throw and the probability of a successful catch. He does know that the probability of a successful catch is the
Figure F.11. Optimal policies for the machine replacement problem for increasing horizons. Each polygon corresponds to the region in which a particular alpha vector dominates.

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**Table F.2.** The transition, observation, and reward functions for the machine replacement problem.

| Action   | $T(s' | s, a)$          | $O(o | a, s')$   | $R(s, a)$ |
|----------|------------------------|-----------------|-----------|
| manufacture | $\begin{bmatrix} 0.81 & 0.18 & 0.01 \\ 0 & 0.9 & 0.1 \\ 0 & 0 & 1 \end{bmatrix}$ | $\begin{bmatrix} 1 \\ 1 \\ 1 \end{bmatrix}$ | $\begin{bmatrix} 0.9025 \\ 0.475 \\ 0.25 \end{bmatrix}$ |
| examine     | $\begin{bmatrix} 0.81 & 0.18 & 0.01 \\ 0 & 0.9 & 0.1 \\ 0 & 0 & 1 \end{bmatrix}$ | $\begin{bmatrix} 0.5 & 0.5 \\ 0.25 & 0.75 \end{bmatrix}$ | $\begin{bmatrix} 0.6525 \\ 0.225 \end{bmatrix}$ |
| interrupt   | $\begin{bmatrix} 1 & 0 & 0 \\ 1 & 0 & 0 \\ 1 & 0 & 0 \end{bmatrix}$ | $\begin{bmatrix} 1 \\ 1 \\ 1 \end{bmatrix}$ | $\begin{bmatrix} -0.5 \\ -1.5 \\ -2.5 \end{bmatrix}$ |
| replace     | $\begin{bmatrix} 1 & 0 & 0 \\ 1 & 0 & 0 \\ 1 & 0 & 0 \end{bmatrix}$ | $\begin{bmatrix} 1 \\ 1 \\ 1 \end{bmatrix}$ | $\begin{bmatrix} -2 \\ -2 \end{bmatrix}$ |

Figure F.12. The 14-step optimal policy for the machine replacement problem has disjoint regions where manufacturing is optimal. Each polygon corresponds to the region in which a particular alpha vector dominates.
same regardless of whether he is throwing or catching, and he has a finite number of attempted catches to maximize his expected utility before he has to go home.

As shown in figure F.13, Johnny models the probability of successfully catching a ball thrown a distance \( d \) as

\[
P(\text{catch} \mid d) = 1 - \frac{1}{1 + \exp\left(-\frac{d - s}{15}\right)} \tag{F.10}
\]

where the proficiency \( s \) is unknown and does not change over time. To keep things manageable, he assumes \( s \) belongs to the discrete set \( S = \{20, 40, 60, 80\} \).

The reward for a successful catch is equal to the distance. If the catch is unsuccessful, then the reward is zero. Johnny wants to maximize the reward over a finite number of attempted throws. With each throw, Johnny chooses a distance from a discrete set \( \mathcal{A} = \{10, 20, \ldots, 100\} \). Johnny begins with a uniform distribution over \( S \).

\section*{F.10 Prisoner’s Dilemma}

The prisoner’s dilemma is a classic problem in game theory involving agents with conflicting objectives. There are two prisoners that are on trial. They can choose to cooperate, remaining silent about their shared crime, or defect, blaming the other for their crime. If they both cooperate, they both serve time for one year. If agent \( i \) cooperates and the other agent \(-i\) defects, then \( i \) serves no time and \(-i\) serves four years. If both defect, then they both serve three years.\footnote{A W. Tucker gave the name and formulated the story. It was based on the original problem formulation of M. Flood and M. Dresher at RAND in 1950. A history is provided by W. Poundstone, \textit{Prisoner’s Dilemma}. Doubleday, 1992.}

The game has \( I = \{1, 2\} \) and \( \mathcal{A} = \mathcal{A}^1 \times \mathcal{A}^2 \) with each \( \mathcal{A}^i = \{\text{cooperate}, \text{defect}\} \). or two agents, we use the table in figure F.14 to express the individual rewards. Rows represent actions for agent 1. Columns represent actions for agent 2. The rewards for agent 1 and 2 are shown in each cell: \( R^1(a^1, a^2), R^2(a^1, a^2) \). The game can be played once or repeated any number of times. In the infinite horizon case, we use a discount factor of \( \gamma = 0.9 \).

\section*{F.11 Rock-Paper-Scissors}

One common game played around the world is rock-paper-scissors. There are two agents who can choose either rock, paper, or scissors. Rock beats scissors, resulting in a unit reward for the agent playing rock and a unit penalty for the agent playing scissors. Scissors beats paper, resulting in a unit reward for the agent playing
scissors and a unit penalty for the agent playing paper. Finally, paper beats rock, resulting in a unit reward for the agent playing paper and a unit penalty for the agent playing rock.

We have $I = \{1, 2\}$ and $A = A^1 \times A^2$ with each $A^i = \{\text{rock, paper, scissors}\}$. Figure F.15 shows the rewards associated with the game, with each cell denoting $R^1(a^1, a^2), R^2(a^1, a^2)$. The game can be played once or repeated any number of times. In the infinite horizon case, we use a discount factor of $\gamma = 0.9$.

F.12 Traveler’s Dilemma

The traveler’s dilemma is a game where an airline loses two identical suitcases from two travelers.\textsuperscript{10} The airline asks the travelers to write down the value of their

suitcases, which can be between $2 and $100, inclusive. If both put down the same value, then they both get that value. The traveler with the lower value gets their value plus $2. The traveler with the higher value gets the lower value minus $2. In other words, the reward function is as follows:

\[
R_i(a_i, a_{-i}) = \begin{cases} 
    a_i & \text{if } a_i = a_{-i} \\
    a_i + 2 & \text{if } a_i < a_{-i} \\
    a_{-i} - 2 & \text{otherwise}
\end{cases}
\] (F.11)

Most people tend to put down between $97 and $100. However, somewhat counter-intuitively, there is a unique Nash equilibrium of only $2.

\section{F.13 Predator-Prey Hex World}

The \textit{predator-prey hex world} problem expands the hex world dynamics to include multiple agents consisting of predators and prey. A predator tries to capture a prey as quickly as possible, and a prey tries to escape the predators as long as possible. The initial state of the hex world is shown in figure F.16. There are no terminal states in this game.

There is a set of predators \( \mathcal{I}_{\text{pred}} \) and a set of prey \( \mathcal{I}_{\text{prey}} \), with \( \mathcal{I} = \mathcal{I}_{\text{pred}} \cup \mathcal{I}_{\text{prey}} \). The states contain the locations of each agent: \( S = S^1 \times \cdots \times S^{\mid \mathcal{I} \mid} \), with each \( S^i \) equal to all hex locations. The joint action space is \( \mathcal{A} = \mathcal{A}^1 \times \cdots \times \mathcal{A}^{\mid \mathcal{I} \mid} \), where each \( \mathcal{A}^i \) consists of all six hex directions of movement.

If a predator \( i \in \mathcal{I}_{\text{pred}} \) and prey \( j \in \mathcal{I}_{\text{prey}} \) share the same hex with \( s_i = s_j \), then the prey is devoured. The prey \( j \) is then transported to a random hex cell, representing its offspring appearing in the world. Otherwise, the state transitions are independent and are as described in the original hex world.

One or more predators can capture one or more prey if they all happen to be in the same cell. If \( n \) predators and \( m \) prey all share the same cell, the predators receive a reward of \( m/n \). For example, if two predators capture one prey together, they each get a reward of 1/2. If three predators capture five prey together, they each get a reward of 5/3. Moving predators receive unit penalty. Prey can move with no penalty, but they receive a penalty of 100 for being devoured.
**F.14 Multi-Caregiver Crying Baby**

The *multi-caregiver crying baby problem* is a multiagent extension of the crying baby problem. For each caregiver $i \in \mathcal{I} = \{1, 2\}$, the states, actions, and observations are as follows:

$$S = \{\text{hungry}, \text{sated}\} \quad (\text{F.12})$$

$$A^i = \{\text{feed}, \text{sing}, \text{ignore}\} \quad (\text{F.13})$$

$$O^i = \{\text{crying}, \text{quiet}\} \quad (\text{F.14})$$

The transition dynamics are similar to the original crying baby problem, except that either caregiver can feed to satisfy the baby:

$$T(\text{sated} \mid \text{hungry}, (\text{feed}, \ast)) = T(\text{sated} \mid \text{hungry}, (\ast, \text{feed})) = 100\% \quad (\text{F.15})$$

where $\ast$ indicates all possible other variable assignments. Otherwise, if the actions are not feed, then the baby transitions between sated to hungry as before:

$$T(\text{hungry} \mid \text{hungry}, (\ast, \ast)) = 100\% \quad (\text{F.16})$$

$$T(\text{sated} \mid \text{hungry}, (\ast, \ast)) = 50\% \quad (\text{F.17})$$

The observation dynamics are also similar to the single agent version, but the model ensures both caregivers make the same observation of the baby, but not necessarily of each other’s choice of caregiving action:

$$O((\text{cry, cry}) \mid (\text{sing, \ast}), \text{hungry}) = O((\text{cry, cry}) \mid (\ast, \text{sing}), \text{hungry}) = 90\% \quad (\text{F.18})$$

$$O((\text{quiet, quiet}) \mid (\text{sing, \ast}), \text{hungry}) = O((\text{quiet, quiet}) \mid (\ast, \text{sing}), \text{hungry}) = 10\% \quad (\text{F.19})$$

$$O((\text{cry, cry}) \mid (\text{sing, \ast}), \text{sated}) = O((\text{cry, cry}) \mid (\ast, \text{sing}), \text{sated}) = 0\% \quad (\text{F.20})$$

If the actions are not sing, then the observations are as follows:

$$O((\text{cry, cry}) \mid (\ast, \ast), \text{hungry}) = O((\text{cry, cry}) \mid (\ast, \ast), \text{hungry}) = 90\% \quad (\text{F.21})$$

$$O((\text{quiet, quiet}) \mid (\ast, \ast), \text{hungry}) = O((\text{quiet, quiet}) \mid (\ast, \ast), \text{hungry}) = 10\% \quad (\text{F.22})$$

$$O((\text{cry, cry}) \mid (\ast, \ast), \text{sated}) = O((\text{cry, cry}) \mid (\ast, \ast), \text{sated}) = 0\% \quad (\text{F.23})$$

$$O((\text{quiet, quiet}) \mid (\ast, \ast), \text{sated}) = O((\text{quiet, quiet}) \mid (\ast, \ast), \text{sated}) = 100\% \quad (\text{F.24})$$
Both caregivers want to help the baby when the baby is hungry, assigning the same penalty of $-10.0$ for both. However, the first caregiver favors feeding and the second caregiver favors singing. For feeding, the first caregiver receives an extra penalty of only $-2.5$, while the second caregiver receives an extra penalty of $-5.0$. For signing, the first caregiver is penalized by $-0.5$, while the second caregiver is penalized by only $-0.25$.

**F.15 Collaborative Predator-Prey Hex World**

The *collaborative predator-prey hex world* is an variant of the predator-prey hex world in which a team of predators chase a single moving prey. The predators must work together to capture a prey. The prey moves randomly to a neighboring cell that is not occupied by a predator.

Predators also only make noisy local observations of the environment. Each predator $i$ detects whether a prey is within a neighboring cell $O^i = \{\text{prey, nothing}\}$. The predators are penalized with a $-1$ reward for movement. They receive a reward of $10$ if one or more of them capture the prey, meaning they are in the same cell as the prey. At this point, the prey is randomly assigned a new cell, signifying the arrival of a new prey for the predators to begin hunting.
G Julia

Julia is a scientific programming language that is free and open source.¹ It is a relatively new language that borrows inspiration from languages like Python, MATLAB, and R. It was selected for use in this book because it is sufficiently high level² so that the algorithms can be compactly expressed and readable while also being fast. This book is compatible with Julia version 1.5. This appendix introduces the concepts necessary for understanding the included code, omitting many of the advanced features of the language.

G.1 Types

Julia has a variety of basic types that can represent data given as truth values, numbers, strings, arrays, tuples, and dictionaries. Users can also define their own types. This section explains how to use some of the basic types and how to define new types.

G.1.1 Booleans

The Boolean type in Julia, written Bool, includes the values true and false. We can assign these values to variables. Variable names can be any string of characters, including Unicode, with a few restrictions.

```
α = true
done = false
```

The variable name appears on the left-hand side of the equal sign; the value that variable is to be assigned is on the right-hand side.

¹ Julia may be obtained from http://julialang.org.

² In contrast with languages like C++, Julia does not require programmers to worry about memory management and other lower-level details, yet it allows low-level control when needed.
We can make assignments in the Julia console. The console, or REPL (for read, eval, print, loop), will return a response to the expression being evaluated. The # symbol indicates that the rest of the line is a comment.

```
 julia> x = true
 true
 julia> y = false; # semicolon suppresses the console output
 julia> typeof(x)
 Bool
 julia> x == y # test for equality
 false
```

The standard Boolean operators are supported.

```
 julia> !x      # not
 false
 julia> x && y # and
 false
 julia> x || y # or
 true
```

### G.1.2 Numbers

Julia supports integer and floating point numbers as shown here:

```
 julia> typeof(42)
 Int64
 julia> typeof(42.0)
 Float64
```

Here, `Int64` denotes a 64-bit integer, and `Float64` denotes a 64-bit floating point value.\(^3\) We can perform the standard mathematical operations:

```
 julia> x = 4
 4
 julia> y = 2
 2
 julia> x + y
 6
 julia> x - y
 2
 julia> x * y
 8
 julia> x / y
 2.0
```

\(^3\) On 32-bit machines, an integer literal like 42 is interpreted as an `Int32`. 
```julia
julia> x ^ y # exponentiation
16
julia> x % y # remainder from division
0
julia> div(x, y) # truncated division returns an integer
2
```

Note that the result of \( x / y \) is a Float64, even when \( x \) and \( y \) are integers. We can also perform these operations at the same time as an assignment. For example, \( x += 1 \) is shorthand for \( x = x + 1 \).

We can also make comparisons:

```julia
julia> 3 > 4
false
julia> 3 >= 4
false
julia> 3 ≥ 4 # unicode also works, use \ge[tab] in console
false
julia> 3 < 4
true
julia> 3 <= 4
true
julia> 3 ≤ 4 # unicode also works, use \le[tab] in console
true
julia> 3 == 4
false
julia> 3 < 4 < 5
true
```

### G.1.3 Strings

A string is an array of characters. Strings are not used very much in this textbook except for reporting certain errors. An object of type String can be constructed using " characters. For example:

```julia
julia> x = "optimal"
"optimal"
julia> typeof(x)
String
```
G.1.4 Symbols

A symbol represents an identifier. It can be written using the `:` operator or constructed from strings.

```julia
julia> :A
:A
julia> :Battery
:Battery
julia> Symbol("Failure")
:Failure
```

G.1.5 Vectors

A vector is a one-dimensional array that stores a sequence of values. We can construct a vector using square brackets, separating elements by commas.

```julia
julia> x = []
# empty vector
julia> x = trues(3)
# Boolean vector containing three trues
julia> x = ones(3)
# vector of three ones
julia> x = zeros(3)
# vector of three zeros
julia> x = rand(3)
# vector of three random numbers between 0 and 1
julia> x = [3, 1, 4]
# vector of integers
julia> x = [3.1415, 1.618, 2.7182]
# vector of floats
```

An array comprehension can be used to create vectors.

```julia
julia> [sin(x) for x = 1:5]
5-element Array{Float64,1}:
0.8414709848078965
0.9092974268256817
0.1411200080598672
-0.7568024953079282
-0.9589242746631385
```

We can inspect the type of vectors:

```julia
julia> typeof([3, 1, 4])
# 1-dimensional array of Int64s
Array{Int64,1}
julia> typeof([3.1415, 1.618, 2.7182])
# 1-dimensional array of Float64s
Array{Float64,1}
julia> Vector{Float64}
# alias for a 1-dimensional array
Array{Float64,1}
```

We index into vectors using square brackets.
We can pull out a range of elements from an array. Ranges are specified using a colon notation.

```
julia> x = [1, 2, 5, 3, 1]
5-element Array{Int64,1}:
 1
 2
 5
 3
 1
```

```
julia> x[1:3]  # pull out the first three elements
3-element Array{Int64,1}:
 1
 2
 5
```

```
julia> x[1:2:end]  # pull out every other element
3-element Array{Int64,1}:
 1
 5
 1
```

```
julia> x[end:-1:1]  # pull out all the elements in reverse order
5-element Array{Int64,1}:
 1
 3
 5
 2
 1
```

We can perform a variety of different operations on arrays. The exclamation mark at the end of function names is used to indicate that the function mutates (i.e., changes) the input.

```
julia> length(x)
5
```

```
julia> [x, x]  # concatenation
2-element Array{Array{Int64,1},1}:
```

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[1, 2, 5, 3, 1]
[1, 2, 5, 3, 1]

**julia**> push!(x, -1)  # add an element to the end
6-element Array{Int64,1}:
1
2
5
3
1
-1

**julia**> pop!(x)  # remove an element from the end
-1

**julia**> append!(x, [2, 3])  # append [2, 3] to the end of x
7-element Array{Int64,1}:
1
2
5
3
1
2
3

**julia**> sort!(x)  # sort the elements, altering the same vector
7-element Array{Int64,1}:
1
1
2
2
3
3
5

**julia**> sort(x);  # sort the elements as a new vector

**julia**> x[1] = 2; print(x)  # change the first element to 2
[2, 1, 2, 2, 3, 3, 5]

**julia**> x = [1, 2];

**julia**> y = [3, 4];

**julia**> x + y  # add vectors
2-element Array{Int64,1}:
4
6

**julia**> 3x - [1, 2]  # multiply by a scalar and subtract
2-element Array{Int64,1}:
2
4

**julia**> using LinearAlgebra
It is often useful to apply various functions elementwise to vectors. This is a form of broadcasting. With infix operators (e.g., +, *, and ^), a dot is prefixed to indicate elementwise broadcasting. With functions like \( \sqrt \) and \( \sin \), the dot is postfixed.

\[
\begin{align*}
\texttt{julia> } x \cdot y & \quad \# \text{ elementwise multiplication} \\
& \quad \text{2-element Array\{Int64,1\}:} \\
& \quad 3 \quad 8 \\
\texttt{julia> } x \cdot^2 & \quad \# \text{ elementwise squaring} \\
& \quad \text{2-element Array\{Int64,1\}:} \\
& \quad 1 \quad 4 \\
\texttt{julia> } \sin.(x) & \quad \# \text{ elementwise application of } \sin \\
& \quad \text{2-element Array\{Float64,1\}:} \\
& \quad 0.8414709848078965 \quad 0.9092974268256817 \\
\texttt{julia> } \sqrt.(x) & \quad \# \text{ elementwise application of } \sqrt \\
& \quad \text{2-element Array\{Float64,1\}:} \\
& \quad 1.0 \quad 1.4142135623730951
\end{align*}
\]

G.1.6 Matrices

A matrix is a two-dimensional array. Like a vector, it is constructed using square brackets. We use spaces to delimit elements in the same row and semicolons to delimit rows. We can also index into the matrix and output submatrices using ranges.

\[
\begin{align*}
\texttt{julia> } X = [1 \ 2 \ 3; \ 4 \ 5 \ 6; \ 7 \ 8 \ 9; \ 10 \ 11 \ 12]; \\
\texttt{julia> } \text{typeof}(X) & \quad \# \text{ a 2-dimensional array of Int64s} \\
& \quad \text{Array\{Int64,2\}} \\
\texttt{julia> } X[2] & \quad \# \text{ second element using column-major ordering} \\
& \quad 4 \\
\texttt{julia> } X[3,2] & \quad \# \text{ element in third row and second column} \\
& \quad 8
\end{align*}
\]
julia> X[1,:]
# extract the first row
3-element Array{Int64,1}:
    1
    2
    3
julia> X[:,2]
# extract the second column
4-element Array{Int64,1}:
    2
    5
    8
   11
julia> X[:,1:2]
# extract the first two columns
4×2 Array{Int64,2}:
    1  2
    4  5
    7  8
   10 11
julia> X[1:2,1:2]
# extract a 2x2 submatrix from the top left of x
2×2 Array{Int64,2}:
    1  2
    4  5
julia> Matrix{Float64}
# alias for a 2-dimensional array
Array{Float64,2}

We can also construct a variety of special matrices and use array comprehensions:

julia> Matrix(1.0I, 3, 3)
# 3x3 identity matrix
3×3 Array{Float64,2}:
    1.0  0.0  0.0
    0.0  1.0  0.0
    0.0  0.0  1.0
julia> Matrix(Diagonal([3, 2, 1]))
# 3x3 diagonal matrix with 3, 2, 1 on diagonal
3×3 Array{Int64,2}:
    3  0  0
    0  2  0
    0  0  1
julia> zeros(3,2)
# 3x2 matrix of zeros
3×2 Array{Float64,2}:
    0.0  0.0
    0.0  0.0
    0.0  0.0
julia> rand(3,2)
# 3x2 random matrix
3×2 Array{Float64,2}:
    0.823025  0.209922
0.892967 0.955656
0.980616 0.340586

```julia
julia> [sin(x + y) for x = 1:3, y = 1:2]  # array comprehension
3x2 Array{Float64,2}:
  0.909297  0.14112
  0.14112  -0.756802
-0.756802  -0.958924
```

Matrix operations include the following:

```julia
julia> X'  # complex conjugate transpose
3x4 LinearAlgebra.Adjoint{Int64,Array{Int64,2}}:
  1  4  7  10
  2  5  8  11
  3  6  9  12

julia> 3X.+ 2  # multiplying by scalar and adding scalar
4x3 Array{Int64,2}:
  5  8  11
 14 17  20
 23 26  29
 32 35  38

julia> X = [1 3; 3 1];  # create an invertible matrix
julia> inv(X)  # inversion
2x2 Array{Float64,2}:
  -0.125  0.375
  0.375  -0.125

julia> det(X)  # determinant (requires LinearAlgebra)
-8.0

julia> [X X]  # horizontal concatenation, same as hcat(X, X)
2x4 Array{Int64,2}:
  1 3 1 3
  3 1 3 1

julia> [X; X]  # vertical concatenation, same as vcat(X, X)
4x2 Array{Int64,2}:
  1 3
  3 1
  1 3
  3 1

julia> sin.(X)  # elementwise application of sin
2x2 Array{Float64,2}:
  0.841471  0.14112
  0.14112  0.841471

julia> map(sin, X)  # elementwise application of sin
2x2 Array{Float64,2}:
  0.841471  0.14112
0.14112 0.841471
julia> vec(X)  # reshape an array as a vector
4-element Array{Int64,1}:
 1
 3
 3
 1

G.1.7 Tuples

A tuple is an ordered list of values, potentially of different types. They are constructed with parentheses. They are similar to vectors, but they cannot be mutated.

julia> x = ()  # the empty tuple
()

julia> isempty(x)
true

julia> x = (1,)  # tuples of one element need the trailing comma
(1,)

julia> typeof(x)
Tuple{Int64}

julia> x = (1, 0, [1, 2], 2.5029, 4.6692)  # third element is a vector
(1, 0, [1, 2], 2.5029, 4.6692)

julia> typeof(x)
Tuple{Int64,Int64,Array{Int64,1},Float64,Float64}

julia> x[2]
0

julia> x[end]
4.6692

julia> x[4:end]
(2.5029, 4.6692)

julia> length(x)
5

julia> x = (1, 2)
(1, 2)

julia> a, b = x;

julia> a
1

julia> b
2

G.1.8 Named Tuples

A named tuple is like a tuple but where each entry has its own name.
```julia
x = (a=1, b=-Inf)
(a = 1, b = -Inf)
x isa NamedTuple
true
x.a
1
a, b = x;
a
1
(; :a=>10)
(a = 10,)
(; :a=>10, :b=>11)
(a = 10, b = 11)
x merge(x, (d=3, e=10)) # merge two named tuples
(a = 1, b = -Inf, d = 3, e = 10)
```

**G.1.9 Dictionaries**

A *dictionary* is a collection of key-value pairs. Key-value pairs are indicated with a double arrow operator `=>`. We can index into a dictionary using square brackets just as with arrays and tuples.

```julia
x = Dict(); # empty dictionary
x[3] = 4 # associate key 3 with value 4
4
x = Dict(3=>4, 5=>1) # create a dictionary with two key-value pairs
Dict{Int64,Int64} with 2 entries:
  3 => 4
  5 => 1
x[5] # return value associated with key 5
1
haskey(x, 3) # check whether dictionary has key 3
true
haskey(x, 4) # check whether dictionary has key 4
false
```

**G.1.10 Composite Types**

A *composite type* is a collection of named fields. By default, an instance of a composite type is immutable (i.e., it cannot change). We use the `struct` keyword and then give the new type a name and list the names of the fields.
struct A
    a
    b
end

Adding the keyword `mutable` makes it so that an instance can change.

mutable struct B
    a
    b
end

Composite types are constructed using parentheses, between which we pass in values for each field. For example,

\[ x = A(1.414, 1.732) \]

The double-colon operator can be used to specify the type for any field.

struct A
    a::Int64
    b::Float64
end

These type annotations require that we pass in an `Int64` for the first field and a `Float64` for the second field. For compactness, this text does not use type annotations, but it is at the expense of performance. Type annotations allow Julia to improve runtime performance because the compiler can optimize the underlying code for specific types.

G.1.11 Abstract Types

So far we have discussed *concrete types*, which are types that we can construct. However, concrete types are only part of the type hierarchy. There are also *abstract types*, which are supertypes of concrete types and other abstract types.

We can explore the type hierarchy of the `Float64` type shown in figure G.1 using the `supertypes` and `subtypes` functions.

![Type Hierarchy Diagram](#)
We can define our own abstract types.

```julia
abstract type C end
abstract type D <: C end  # D is an abstract subtype of C
struct E <: D  # E is composite type that is a subtype of D
    a
end
```

### G.1.12 Parametric Types

Julia supports *parametric types*, which are types that take parameters. The parameters to a parametric type are given within braces and delimited by commas. We have already seen a parametric type with our dictionary example.

```julia
julia> x = Dict(3=>1.4, 1=>5.9)
Dict{Int64,Float64} with 2 entries:
  3 => 1.4
  1 => 5.9
```

For dictionaries, the first parameter specifies the key type, and the second parameter specifies the value type. The example has `Int64` keys and `Float64` values, making the dictionary of type `Dict{Int64,Float64}`. Julia was able to infer these types based on the input, but we could have specified it explicitly.
While it is possible to define our own parametric types, we do not need to do so in this text.

### G.2 Functions

A function maps its arguments, given as a tuple, to a result that is returned.

#### G.2.1 Named Functions

One way to define a named function is to use the `function` keyword, followed by the name of the function and a tuple of names of arguments.

```julia
function f(x, y)
    return x + y
end
```

We can also define functions compactly using assignment form.

```julia
julia> f(x, y) = x + y;

julia> f(3, 0.1415)
3.1415
```

#### G.2.2 Anonymous Functions

An anonymous function is not given a name, though it can be assigned to a named variable. One way to define an anonymous function is to use the arrow operator.

```julia
julia> h = x -> x^2 + 1 # assign anonymous function with input x to a variable h
#1 (generic function with 1 method)

julia> h(3)
10

julia> g(f, a, b) = [f(a), f(b)]; # applies function f to a and b and returns array
julia> g(h, 5, 10)
2-element Array{Int64,1}:
   26
   101

julia> g(x->sin(x)+1, 10, 20)
2-element Array{Float64,1}:
0.4559788891106302
1.9129452507276277
```
G.2.3 Callable Objects

We can define a type and associate functions with it, allowing objects of that type to be callable.

```julia
julia> (x::A)() = x.a + x.b  # adding a zero-argument function to the type A defined earlier
julia> (x::A)(y) = y*x.a + x.b # adding a single-argument function
julia> x = A(22, 8);
julia> x()
30
julia> x(2)
52
```

G.2.4 Optional Arguments

We can assign a default value to an argument, making the specification of that argument optional.

```julia
julia> f(x=10) = x^2;
julia> f()  
100
julia> f(3)
9
julia> f(x, y, z=1) = x*y + z;
julia> f(1, 2, 3)
5
julia> f(1, 2)
3
```

G.2.5 Keyword Arguments

Functions may use keyword arguments, which are arguments that are named when the function is called. Keyword arguments are given after all the positional arguments. A semicolon is placed before any keywords, separating them from the other arguments.

```julia
julia> f(; x = 0) = x + 1;
julia> f()
1
julia> f(x = 10)
11
julia> f(x, y = 10; z = 2) = (x + y)*z;
julia> f(1)
```

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G.2.6 Dispatch

The types of the arguments passed to a function can be specified using the double colon operator. If multiple methods of the same function are provided, Julia will execute the appropriate method. The mechanism for choosing which method to execute is called dispatch.

```
 julia> f(x::Int64) = x + 10;
 julia> f(x::Float64) = x + 3.1415;
 julia> f(1)
 11
 julia> f(1.0)
 4.141500000000001
 julia> f(1.3)
 4.4415000000000004
```

The method with a type signature that best matches the types of the arguments given will be used.

```
 julia> f(x) = 5;
 julia> f(x::Float64) = 3.1415;
 julia> f([3, 2, 1])
 5
 julia> f(0.00787499699)
 3.1415
```

G.2.7 Splatting

It is often useful to *splat* the elements of a vector or a tuple into the arguments to a function using the ... operator.
G.3 Control Flow

We can control the flow of our programs using conditional evaluation and loops. This section provides some of the syntax used in the book.

G.3.1 Conditional Evaluation

Conditional evaluation will check the value of a Boolean expression and then evaluate the appropriate block of code. One of the most common ways to do this is with an `if` statement.

```julia
if x < y
    # run this if x < y
elseif x > y
    # run this if x > y
else
    # run this if x == y
end
```

We can also use the *ternary operator* with its question mark and colon syntax. It checks the Boolean expression before the question mark. If the expression evaluates to true, then it returns what comes before the colon; otherwise it returns what comes after the colon.

```julia
f(x) = x > 0 ? x : 0;
```

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G.3.2 Loops

A loop allows for repeated evaluation of expressions. One type of loop is the while loop. It repeatedly evaluates a block of expressions until the specified condition after the while keyword is met. The following example sums the values in the array \( x \).

```julia
X = [1, 2, 3, 4, 6, 8, 11, 13, 16, 18]
s = 0
while !isempty(X)
    s += pop!(X)
end
```

Another type of loop is the for loop. It uses the for keyword. The following example will also sum over the values in the array \( x \) but will not modify \( x \).

```julia
X = [1, 2, 3, 4, 6, 8, 11, 13, 16, 18]
s = 0
for i = 1:length(X)
    s += X[i]
end
```

The `=` can be substituted with `in` or `∈`. The following code block is equivalent.

```julia
X = [1, 2, 3, 4, 6, 8, 11, 13, 16, 18]
s = 0
for y in X
    s += y
end
```

G.3.3 Iterators

We can iterate over collections in contexts such as for loops and array comprehensions. To demonstrate various iterators, we will use the `collect` function, which returns an array of all items generated by an iterator:

```julia
julia> X = ["feed", "sing", "ignore"];
julia> collect(enumerate(X)) # return the count and the element
3-element Array{Tuple{Int64,String},1}:
(1, "feed")
(2, "sing")
(3, "ignore")
julia> collect(eachindex(X)) # equivalent to 1:length(X)
3-element Array{Int64,1}:
```

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G.4 Packages

A *package* is a collection of Julia code and possibly other external libraries that can be imported to provide additional functionality. This section briefly reviews a few of the key packages that we build upon. To add a registered package like Distributions.jl, we can run:

```julia
using Pkg
Pkg.add("Distributions")
```

To update packages, we use:

```
julia> import IterTools: subsets
julia> collect(subsets(X)) # iterate over all subsets
8-element Array{Array{String,1},1}:
    []
    ["feed"]
    ["sing"]
    ["feed", "sing"]
    ["ignore"]
    ["feed", "ignore"]
    ["sing", "ignore"]
    ["feed", "sing", "ignore"]
```

```julia
julia> collect(eachindex(X)) # iterate over indices into a collection
3-element Array{Int64,1}:
    1
    2
    3
```

```
julia> Z = [1  2;  3  4;  5  6];
julia> import Base.Iterators: product
julia> collect(product(X,Y)) # iterate over Cartesian product of multiple iterators
3×3 Array{Tuple{String,Float64},2}:
     ("feed", -5.0) ("feed", -0.5) ("feed", 0.0)
     ("sing", -5.0) ("sing", -0.5) ("sing", 0.0)
     ("ignore", -5.0) ("ignore", -0.5) ("ignore", 0.0)
```

G.4 Packages

A *package* is a collection of Julia code and possibly other external libraries that can be imported to provide additional functionality. This section briefly reviews a few of the key packages that we build upon. To add a registered package like Distributions.jl, we can run:

```julia
using Pkg
Pkg.add("Distributions")
```

To update packages, we use:
To use a package, we use the keyword `using`:

```julia
using Distributions
```

### G.4.1 LightGraphs.jl

We use the LightGraphs.jl package (version 1.3) to represent graphs and perform operations on them:

```julia
julia> using LightGraphs
julia> G = SimpleDiGraph(3); # create a directed graph with three nodes
julia> add_edge!(G, 1, 3);  # add edge from node 1 to 3
julia> add_edge!(G, 1, 2);  # add edge from node 1 to 2
julia> rem_edge!(G, 1, 3); # remove edge from node 1 to 3
julia> add_edge!(G, 2, 3);  # add edge from node 2 to 3
julia> typeof(G)
LightGraphs.SimpleGraphs.SimpleDiGraph{Int64}
```

```julia
julia> nv(G) # number of nodes (also called vertices)
3
julia> outneighbors(G, 1) # list of outgoing neighbors for node 1
1-element Array{Int64,1}:
2
julia> inneighbors(G, 1) # list of incoming neighbors for node 1
Int64[]
```

### G.4.2 Distributions.jl

We use the Distributions.jl package (version 0.24) to represent, fit, and sample from probability distributions:

```julia
julia> using Distributions
julia> μ, σ = 5.0, 2.5;
julia> dist = Normal(μ, σ) # create a normal distribution
Distributions.Normal{Float64}(μ=5.0, σ=2.5)
julia> rand(dist) # sample from the distribution
4.576185798799843
julia> data = rand(dist, 3) # generate three samples
3-element Array{Float64,1}:
2.572834058347085
6.122558160059754
6.182668194105785
julia> data = rand(dist, 1000); # generate many samples
```

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julia> Distributions.fit(Normal, data)  # fit a normal distribution to the samples
Distributions.Normal{Float64}(μ=5.01500600151507, σ=2.533059172253862)

julia> μ = [1.0, 2.0];
julia> Σ = [1.0 0.5; 0.5 2.0];
julia> dist = MvNormal(μ, Σ)  # create a multivariate normal distribution
FullNormal(
  dim: 2
  μ: [1.0, 2.0]
  Σ: [1.0 0.5; 0.5 2.0]
)

julia> rand(dist, 3)  # generate three samples
2×3 Array{Float64,2}:
  1.54004  1.59641  2.27612
  1.50691  2.62729  1.09536

julia> dist = Dirichlet(ones(3))  # create a Dirichlet distribution Dir(1,1,1)
Distributions.Dirichlet{Float64}(alpha=[1.0, 1.0, 1.0])

julia> rand(dist)  # sample from the distribution
3-element Array{Float64,1}:
  0.34454105328617957
  0.5868641887928219
  0.06859475792099853

G.4.3 JuMP.jl

We use the JuMP.jl package (version 0.21) to specify optimization problems that we can then solve using a variety of different solvers, such as those included in GLPK.jl and Ipopt.jl:

julia> using JuMP

julia> using GLPK

julia> model = Model(GLPK.Optimizer)  # create model and use GLPK as solver
A JuMP Model
Feasibility problem with:
Variables: 0
Model mode: AUTOMATIC
CachingOptimizer state: EMPTY_OPTIMIZER
Solver name: GLPK

julia> @variable(model, x[1:3])  # define variables x[1], x[2], and x[3]
3-element Array{JuMP.VariableRef,1}:
  x[1]
  x[2]
  x[3]

julia> @objective(model, Max, sum(x) - x[2])  # define maximization objective
julia> @constraint(model, x[1] + x[2] ≤ 3)  # add constraint
julia> @constraint(model, x[2] + x[3] ≤ 2)  # add another constraint
julia> @constraint(model, x[2] ≥ 0)  # add another constraint
x[2] ≥ 0.0
julia> optimize!(model)  # solve
julia> value.(x)  # extract optimal values for elements in x
3-element Array{Float64,1}:
  3.0
  0.0
  2.0

G.5 Convenience Functions

There are a few functions that allow us to more compactly specify the algorithms in the body of this book. Julia 1.7 will support a two-argument version of findmax, where we can pass in a function and a collection. It returns the maximum of the function when evaluated on the elements of the collection along with the first maximizing element. The argmax function is similar, but it only returns the first maximizing element. To support this in Julia 1.5, we manually extend these functions.

```julia
function Base.findmax(f::Function, xs)
    f_max = -Inf
    x_max = first(xs)
    for x in xs
        v = f(x)
        if v > f_max
            f_max, x_max = v, x
        end
    end
    return f_max, x_max
end

Base.argmax(f::Function, xs) = findmax(f, xs)[2]
```

julia> findmax(x→x^2, [0, -10, 3])
(100, -10)
julia> argmax(abs, [0, -10, 3])
-10
The following functions are useful when working with dictionaries and named
tuples:

```
Base.Dict{Symbol,V}(a::NamedTuple) where V =
   Dict{Symbol,V}(n↦v for (n,v) in zip(keys(a), values(a)))
Base.convert(::Type{Dict{Symbol,V}}, a::NamedTuple) where V =
   Dict{Symbol,V}(a)
Base.isequal(a::Dict{Symbol,<:Any}, nt::NamedTuple) =
   length(a) == length(nt) &&
   all(a[n] == v for (n,v) in zip(keys(nt), values(nt)))
```

```
julia> a = Dict{Symbol,Integer}((a=1, b=2, c=3))
Dict{Symbol,Integer} with 3 entries:
    :a => 1
    :b => 2
    :c => 3
julia> isequal(a, (a=1, b=2, c=3))
true
julia> isequal(a, (a=1, c=3, b=2))
true
julia> Dict{Dict{Symbol,Integer},Float64}((a=1, b=1)↦0.2, (a=1, b=2)↦0.8)
Dict{Dict{Symbol,Integer},Float64} with 2 entries:
    Dict{Symbol,Integer}(:a↦1,:b↦1) => 0.2
    Dict{Symbol,Integer}(:a↦1,:b↦2) => 0.8
```

We define `SetCategorical` to represent distributions over discrete sets.

```
struct SetCategorical{S}
    elements::Vector{S} # Set elements (could be repeated)
    distr::Categorical    # Categorical distribution over set elements

    function SetCategorical(elements::AbstractVector{S}) where S
        weights = ones(length(elements))
        return new{S}(elements, Categorical(normalize(weights, 1)))
    end

    function SetCategorical(elements::AbstractVector{S}, weights::AbstractVector{Float64}) where S
        ℓ₁ = norm(weights,1)
        if ℓ₁ < 1e-6 || isinf(ℓ₁)
            return SetCategorical(elements)
        end
        distr = Categorical(normalize(weights, 1))
        return new{S}(elements, distr)
    end
end
```

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2021-03-07 23:12:00-08:00, revision 80bf4d6, comments to bugs@algorithmsbook.com
Distributions.rand(D::SetCategorical) = D.elements[rand(D.distr)]
Distributions.rand(D::SetCategorical, n::Int) = D.elements[rand(D.distr, n)]
function Distributions.pdf(D::SetCategorical, x)
    sum(e == x ? w : 0.0 for (e,w) in zip(D.elements, D.distr.p))
end

julia> D = SetCategorical(["up", "down", "left", "right"],[0.4, 0.2, 0.3, 0.1]);
julia> rand(D)
"up"

julia> rand(D, 5)
5-element Array{String,1}:
  "left"
  "up"
  "right"
  "right"
  "left"

julia> pdf(D, "up")
0.3999999999999999
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