

Multi-Agent Reinforcement Learning

Multi-Agent Reinforcement Learning: Foundations and Modern Approaches

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Preface

Multi-agent reinforcement learning (MARL) is a varied and highly active field of research. With the introduction of deep learning to MARL in the mid-2010s, the field has seen an explosive growth of activity and now all major artificial intelligence and machine learning conferences routinely feature papers that develop new MARL algorithms or apply MARL in some way. This steep growth is also documented by the increasing number of survey papers that have been published since, of which we list many in Appendix A.

In the wake of this growth, it became clear that the field needed a textbook to provide a principled introduction to MARL. The present book is in part based on, and largely follows the same structure as, the tutorial “*Multiagent Learning: Foundations and Recent Trends*” given by Stefano V. Albrecht and Peter Stone at the 2017 International Joint Conference on Artificial Intelligence in Melbourne, Australia. The book was written to provide a basic introduction to the models, solution concepts, algorithmic ideas, and technical challenges in MARL, and to describe modern approaches in MARL which integrate deep learning techniques to produce powerful new algorithms. In essence, we believe that the materials covered in this book should be known by every MARL researcher. In addition, the book aims to give practical guidance for researchers and practitioners when using MARL algorithms. To this end, the book comes with its own codebase written in the Python programming language, which contains implementations of several MARL algorithms discussed in this book. The primary purpose of the codebase is to provide algorithm code that is self-contained and easy to read, to aid the reader’s understanding.

This book assumes that readers have an undergraduate-level background in basic mathematics, including statistics, probability theory, linear algebra, and calculus. A basic familiarity with programming concepts is required to understand and use the codebase. In general, we recommend reading the book chapters in the given sequence. For readers unfamiliar with reinforcement learning and deep learning, we provide the basics in Chapters 2, 7 and 8,

respectively. Readers who are already familiar with reinforcement learning and deep learning, and who want to quickly get going with recent deep learning-based MARL algorithms, may read Chapter 3 and then skip to Chapter 9 and onward. To aid lecturers in adopting this book, we have developed lecture slides (available from the book’s website), which can be modified as required to suit the course’s needs.

MARL has become a large field of research, and this book does not cover all aspects of MARL. For instance, there is a growing body of work on using communication in MARL, which is not covered in this book. This includes questions such as how agents can learn to communicate robustly when communication channels are noisy and unreliable; and how agents may use MARL to learn specialised communication protocols or languages for a given task. While this book does not focus on communication in MARL, the models introduced in this book are general enough to also represent communication actions that can be observed by agents but do not affect the state of the environment. There has also been research on using evolutionary methods for multi-agent learning, in the sense of mutation and crossover in agent populations, which is not covered in this book. Finally, with the steep rise of activity in MARL in recent years, it would be futile to write a book that tries to keep up with new algorithms. We instead focus on the foundational concepts and ideas in MARL, and refer to survey papers (including those listed in Appendix A) for a more complete list of algorithm developments.

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Errata: Despite our best efforts, it is possible that some typos or imprecisions have gone unnoticed. If you detect any errors, we would be much obliged if you could report them to us via e-mail to issues@marl-book.com.

Book website, codebase, slides: The full PDF version of this book and links to accompanying materials, including the codebase and lecture slides, can be found on the book website at: **www.marl-book.com**

May, 2023

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Summary of Notation

Sets are denoted with capital letters.

Elements of sets are denoted with lower-case letters.

Time index t (or τ) is shown in superscript (e.g. s^t denotes state at time t).

Agent index is shown in subscript (e.g. a_i denotes action of agent i).

The most common symbols used in the book are listed below. Specific sections may use additional notation.

General

\mathbb{R}	set of real numbers
x^\top	transpose of a vector x
X^\top	transpose of a matrix X
\Pr	probability
$\Pr(x y)$	conditional probability of x given y
$\mathbb{E}_p[x]$	expectation of x under probability distribution p
$x \sim p$	x sampled according to probability distribution p
$x \leftarrow y$	assign value y to variable x
\mathcal{D}	training data set
$\frac{\partial f}{\partial x}$	derivative of function f wrt x
∇	gradient
$\langle a, b, c, \dots \rangle$	concatenation of inputs into tuple (a, b, c, \dots)
$[x]_1$	indicator function: returns 1 iff. x is true, otherwise returns 0

Game Model

I	set of agents
i, j	agent subscripts
$-i$	subscript to denote the tuple <all agents except agent i >
S	state space
s, \hat{s}	states
O, O_i	(joint) observation space, observation space of agent i
o, o_i	(joint) observation, observation of agent i
A, A_i	(joint) action space, action space of agent i
a, \hat{a}, a_i	(joint) actions, action of agent i
r, r_i	(joint) reward, reward of agent i

\Pr^0	initial state distribution
\mathcal{T}	state transition function
$\hat{\mathcal{T}}$	simulation/sampling model of state transitions
$\mathcal{O}, \mathcal{O}_i$	observation function (of agent i)
$\mathcal{R}, \mathcal{R}_i$	reward function (of agent i)
$\Gamma_s, \Gamma_{s,i}$	normal-form game for state s (and agent i)

Policies, Returns, Values

Π, Π_i	(joint) policy space, policy space of agent i
π, π_i	(joint) policy, policy of agent i
π^*	optimal policy, or equilibrium joint policy
H	set of histories
h	joint observation history (obs. type depends on game model)
h_i	local observation history of agent i
\hat{H}	set of full histories
\hat{h}	full history containing states, joint observations, joint actions
$\sigma(\hat{h})$	function returning joint observation history from full history \hat{h}
γ	discount factor
u, u_i	discounted return (for agent i)
U, U_i	expected discounted return (for agent i); also called “value”

(Multi-Agent) Reinforcement Learning

\mathbb{L}	learning algorithm
α	learning rate
ϵ	exploration rate
$\bar{\pi}_i$	averaged policy of agent i
$\hat{\pi}_j$	agent model for agent j
BR_i	set of best-response actions or policies for agent i
V^π, V_i^π	state value function (of agent i) under policy π
Q^π, Q_i^π	action value function (of agent i) under policy π
V^*, Q^*	optimal state/action value function
Value_i	equilibrium value of non-repeated normal-form game for agent i
AV_i	action value for agent i against a given agent model

Deep Learning

θ	network parameters
$f(x; \theta)$	function f over input x with parameters θ
\mathcal{L}	loss
\mathcal{B}	training data batch
B	batch size, i.e. number of samples in a batch

(Multi-Agent) Deep Reinforcement Learning

θ, θ_i	value function parameters (of agent i)
ϕ, ϕ_i	policy parameters (of agent i)
$\bar{\theta}$	target network parameters
$\mathcal{D}, \mathcal{D}_i$	experience replay buffer (of agent i)
\mathcal{H}	entropy

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1 Introduction

Imagine a scenario in which a collective of autonomous agents, each capable of making its own decisions, must interact in a shared environment to achieve certain goals. The agents may have a shared goal, such as a fleet of mobile robots whose task is to collect and deliver goods within a large warehouse, or a team of drones tasked with monitoring an oil rig in the sea. The agents may also have conflicting goals, such as agents trading goods in a virtual market in which each agent seeks to maximise its own gains. Since we may not know how the agents should interact to achieve their goals, we tell them to figure it out on their own. Thus, the agents begin to try actions in their environment and collect experiences about how the environment changes as a result of their actions, as well as how the other agents behave. In time, the agents begin to learn various concepts, such as skills needed to solve their task and, importantly, how to *coordinate* their actions with other agents. They may even learn to develop a shared language to enable communication between agents. Finally, the agents reach a certain level of proficiency and have become experts at interacting optimally to achieve their goals.

This exciting vision is, in a nutshell, what *multi-agent reinforcement learning* (MARL) aims to achieve. MARL is based on *reinforcement learning* (RL), in which agents learn optimal decision policies by trying actions and receiving rewards, with the goal of choosing actions to maximise the sum of received rewards over time. While in single-agent RL the focus is on learning an optimal policy for a single agent, in MARL the focus is on learning optimal policies for multiple agents and the unique challenges that arise in this learning process.

In this first chapter, we will begin to outline some of the underlying concepts and challenges in MARL. We begin by introducing the concept of a multi-agent system, which is defined by an environment, the agents in the environment, and their goals. We then discuss how MARL operates in such systems to learn optimal policies for the agents, which we illustrate with a number of examples of potential applications. Next we discuss some of the key challenges in MARL,

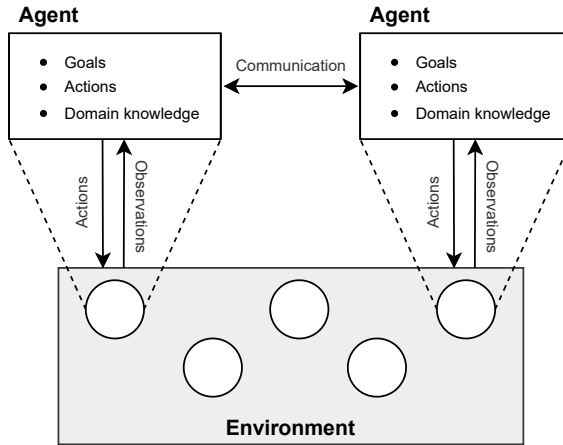


Figure 1.1: Schematic of a multi-agent system. A multi-agent system consists of an environment and multiple decision-making agents. The agents can observe information about the environment and take actions to achieve their goals.

such as the non-stationarity and equilibrium selection problems, as well as several “agendas” of MARL which describe different ways in which MARL can be used. At the end of this chapter, we give an overview of the topics covered in the two parts of this book.

1.1 Multi-Agent Systems

A multi-agent system consists of an *environment* and multiple decision-making *agents* that interact in the environment to achieve certain goals. Figure 1.1 shows a general schematic of a multi-agent system, and we describe the basic components below.

Environment An environment is a physical or virtual world whose state evolves over time and is influenced by the actions of the agents that exist within the environment. The environment specifies the actions that agents can take at any point in time, as well as the observations that individual agents receive about the state of the environment. The state of the environment may be defined as discrete or continuous quantities, or a combination of both. For example, in a 2D-maze environment, the state may be defined as the combination of the discrete integer positions of all agents together with their continuous orientations in radians. Similarly, actions may be

discrete or continuous, such as moving up/down/left/right in the maze and turning around by a specified continuous angle. Multi-agent environments are often characterised by the fact that agents only have a limited and imperfect view of the environment. This means that individual agents may only observe some partial information about the state of the environment, and different agents may receive different observations about the environment.

Agents An agent is an entity which receives information about the state of the environment and can choose different actions in order to influence the state. Agents may have different prior knowledge about the environment, such as the possible states that the environment can be in and how states are affected by the actions of the agents. Importantly, agents are goal-directed in the sense that agents have specified *goals* and choose their actions in order to achieve their goals. These goals could be to reach a certain environment state, or to maximise certain quantities such as monetary revenues. In MARL, such goals are defined by *reward functions* which specify scalar reward signals that agents receive after taking certain actions in certain states. The term *policy* refers to a function used by the agent to select actions (or assign probabilities to selecting each action) given the current state of the environment. If the environment is only partially observed by the agent, then the policy may be conditioned on the current and past observations of the agent.

As a concrete example of the above concepts, consider the level-based foraging example shown in Figure 1.2.¹ In this example, multiple robots are tasked with collecting items which are spatially distributed in a grid-world environment. Each robot and item has an associated skill level, and a group of one or more robots can collect an item if the robots are located next to the item and the sum of the robots' levels is greater than or equal to the item's level. The state of this environment at a given time is completely described by variables containing the x/y-positions of the robots and items, and binary variables for each item indicating whether the item still exists or not.² In this example, we use three independent agents to control each of the three robots. At any given time, each agent can observe the complete state of the environment, and choose an action from the set $\{up, down, left, right, collect, noop\}$ to control its robot. The first

1. The level-based foraging example will appear throughout this book. We provide an open-source implementation of this environment at <https://github.com/uoe-agents/lb-foraging>. See Section 11.3.1 for more details on this environment.

2. Note that the skill levels of robots and items are not included in the state, since these are assumed constant. However, if the skill levels can change between different episodes, then the state may additionally include the skill levels of the robots and items.

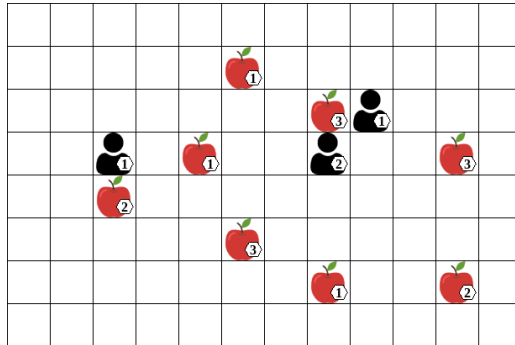


Figure 1.2: A level-based foraging task in which a group of three robots, each controlled by an agent, must collect all items (shown as apples). Each robot and item has an associated skill level, shown inset. A group of one or more robots can collect an item if they are located next to the item, and the sum of the robots' levels is greater than or equal to the item's level.

four actions in this action set modify the x/y-position of the robot in the state by moving the robot into the respective direction (unless the robot is already at the edge of the grid world, in which case the action has no effect). The *collect* action attempts to collect an item that is located adjacent to the robot; if the item is collected, the action modifies the binary existence variable corresponding to the item. The *noop* (no operation) action has no effect on the state.

Notice that in the above description, we used the terms “robot” and “agent” to refer to two distinct concepts. In level-based foraging, the word “robot” is a label for an object which is explicitly represented via the x/y-position variables. Similarly, the word “item” refers to an object in level-based foraging which is represented by its x/y-position variables and its binary existence variable. In contrast to these object labels, the term “agent” refers to an abstract decision-making entity which observes some information from the environment and chooses values for certain action variables; in this case, the action for a robot. If there is a direct one-to-one correspondence between agents and certain objects, such as agents and robots in level-based foraging, then it can be convenient to use the terms interchangeably. For example, in level-based foraging, we may say “the skill level of agent i ” when referring to the skill level of the robot controlled by agent i . Unless the distinction is relevant, in this book we will often use the term “agent” synonymously with the object that it controls.

The defining characteristic of a multi-agent system is that the agents must coordinate their actions with (or against) each other in order to achieve their goals. In a fully cooperative scenario, the agents’ goals are perfectly aligned so

that the agents need to collaborate towards achieving a shared goal. For example, in level-based foraging, all agents may receive a reward of +1 whenever an item is successfully collected by any of the agents. In a competitive scenario, the agents' goals may be diametrically opposed so that the agents are in direct competition with each other. An example of such a competitive scenario is two agents playing a game of chess, in which the winning player gets a reward of +1 and the losing player gets a reward of -1 (or both get 0 for a drawn outcome). In between these two extremes, the agents' goals may align in some respects while differing in other respects, which can lead to complex multi-agent interaction problems that involve both cooperation and competition to varying degrees. For example, in the actual implementation of level-based foraging we use in this book (described in Section 11.3.1), only those agents which were involved in the collection of an item (rather than all agents) will receive a positive normalised reward. Thus, the agents have a motivation to maximise their own returns (sum of rewards), which can lead them to try and collect items before other agents can do so; but they may also need to collaborate with other agents at certain times in order to collect an item.

The above concepts of states, actions, observations, and rewards are formally defined within *game models*. Different types of game models exist, and Chapter 3 introduces the most common game models used in MARL, including normal-form games, stochastic games, and partially observable stochastic games. A solution for a game model consists of a set of policies for the agents which satisfies certain desired properties. As we will see in Chapter 4, there exist a range of solution concepts in the general case. Most solution concepts are anchored in some notion of *equilibrium*, which means that no individual agent can deviate from its policy in the solution to improve its outcome.

Research in multi-agent systems has a long history in artificial intelligence and spans a vast range of technical problems (Shoham and Leyton-Brown 2008; Wooldridge 2009). These include questions such as how to design algorithms which enable agents to choose optimal actions towards their specified goals; how to design environments to incentivise certain long-term behaviours in agents; how information is communicated and propagated among agents; and how norms, conventions and roles may emerge in a collective of agents. This book is concerned with the first of these questions, with a focus on using RL techniques to optimise and coordinate the policies of agents in order to maximise the rewards they accumulate over time.

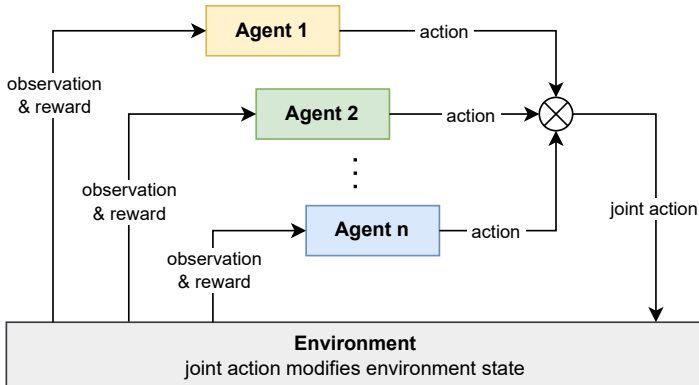


Figure 1.3: Schematic of multi-agent reinforcement learning. A set of n agents receive individual observations about the state of the environment, and choose actions to modify the state of the environment. After taking an action, each agent receives a scalar reward and a new observation, and the loop repeats.

1.2 Multi-Agent Reinforcement Learning

Multi-agent reinforcement learning (MARL) algorithms learn optimal policies for a set of agents in a multi-agent system.³ As in the single-agent counterpart, the policies are learned via a process of trial-and-error to maximise the agents' cumulative rewards, or *returns*. Figure 1.3 shows a basic schematic of the MARL training loop. A set of n agents choose individual actions, which together are referred to as the *joint action*. The joint action changes the state of the environment according to the environment dynamics, and the agents receive individual rewards as a result of this change, as well as individual observations about the new environment state. This loop continues until a terminal criterion is satisfied (such as one agent winning a game of chess) or indefinitely. A complete run of this loop from the initial state to the terminal state is called an *episode*. The generated data produced from multiple independent episodes – i.e. the experienced observations, actions, and rewards in each episode – are used to continually improve the agents' policies.

In the level-based foraging environment introduced in the previous section, each agent $i \in \{1, 2, 3\}$ observes the full environment state and chooses an action $a_i \in \{up, down, left, right, collect, noop\}$. Given the joint action (a_1, a_2, a_3) , the

3. This book uses a literal definition of the MARL term in which we learn policies for *multiple* agents. This is in contrast to learning a policy for a single agent that operates in a multi-agent system, in which we have no control over the other agents.

environment state transitions into a new state by modifying the x/y-positions of the robots and the binary existence variables of the items, depending on the chosen actions in the joint action. Then each agent receives a reward, such as +1 if any of the items has been collected and 0 otherwise, and observes the new state of the environment. An episode in level-based foraging terminates once all items have been collected, and after a maximum number of allowed time steps. Initially, each agent starts with a random policy which selects actions randomly. As the agents keep trying different actions in different states and observe the resulting rewards and new states, they will change their policies to select actions in each state which will maximise the sum of received rewards.

The MARL loop shown in Figure 1.3 is analogous to the single-agent RL loop (which will be covered in Chapter 2) and extends it to multiple agents. There are several important use cases in which MARL can have significant benefits over single-agent RL. One use case for MARL is to decompose a large, intractable decision problem into smaller, more tractable decision problems. To illustrate this idea, consider again the level-based foraging example shown in Figure 1.2. If we view this as a single-agent RL problem, then we have to train a single central agent which selects actions for each of the three robots. Thus, an action of the central agent is defined by the tuple (a_1, a_2, a_3) , in which a_i specifies what robot i does. This results in a decision problem with $6^3 = 216$ possible actions for the central agent in every time step. Even in this toy example, most standard single-agent RL algorithms do not scale easily to action spaces this large. However, we can decompose this decision problem by introducing three independent agents, one for each robot, such that each agent faces a decision problem with only 6 possible actions in each time step. Of course, this decomposition introduces a new challenge, which is that the agents need to coordinate their actions in order to be successful. MARL algorithms may use various approaches to facilitate the learning of coordinated agent policies.

Even if we were able to successfully solve the above example using single-agent RL to train a single central agent, this approach rests on the implicit assumption that the environment allows for centralised control. However, in many applications of multi-agent systems, it may not be possible to control and coordinate the actions of multiple agents from a central place, such as due to non-existent communication links to and between agents. Examples include autonomous driving in urban environments where each car requires its own local policy to drive; or a team of mobile robots used in search-and-rescue missions in which each agent (robot) may need to act fully independently. In such applications, the agents may need to learn *decentralised* policies, where each agent executes its own policy locally based on its own observations. For

such applications, MARL algorithms can learn agent policies which can be executed in a decentralised fashion.

MARL algorithms can be categorised based on various properties, such as assumptions about the agents' rewards (e.g. fully cooperative, competitive, or mixed), and what type of solution concept the algorithm is designed to achieve (e.g. Nash equilibrium), both of which will be discussed in detail in Chapters 3 and 4. Another important property is based on what assumptions are made during the learning of agent policies ("training") versus assumptions made after learning ("execution"). Centralised training and execution assumes that both stages have access to some centrally shared mechanism or information between agents, such as sharing all observations between agents. For example, a single central agent may receive information from all other agents and dictate the actions to the agents. Such centralisation can help to improve coordination between agents and alleviate issues such as non-stationarity (discussed in Section 1.4). In contrast, decentralised training and execution assumes no such centrally shared information, and instead requires that the learning of an agent's policy as well as the policy itself only use the local information of that agent. The third major category, centralised training with decentralised execution, aims to combine the benefits of the two aforementioned approaches, by assuming that centralisation is feasible during training (e.g. in simulation) while producing policies that can be executed in a fully decentral way. These ideas are further discussed in Chapter 9.

1.3 Application Examples

We provide several examples to illustrate the MARL training loop shown in Figure 1.3 and its different constituent elements, such as agents, observations, actions, and rewards. Each example is based on a potential real-world application, and we give pointers to works which have used MARL to develop such applications.

1.3.1 Multi-Robot Warehouse Management

Imagine a large warehouse consisting of many aisles of storage racks that contain all manner of items. There is a constant stream of orders, which specify certain items and quantities to be picked up from the storage racks and delivered to a work station for further processing. Suppose we have 100 mobile robots which can move along the aisles and pick items from the storage racks. We can use MARL to train these robots to collaborate optimally to service the incoming orders, with the goal of completing the orders as quickly and efficiently as

possible. In this application, each robot could be controlled by an independent agent, so we would have 100 agents. Each agent might observe information about its own location and current heading within the warehouse, the items it is carrying, and the current order it is servicing. It might also observe information about other agents, such as their locations, items, and orders. The actions of an agent may include physical movement actions, such as rotating toward a certain direction and accelerating/breaking, as well as picking items. Actions might also include sending communication messages to other robots, which could for example contain information about the travel plans of the communicating agent. Lastly, each agent might receive an individual positive reward when completing an order, which includes picking all items at the specified quantities in the order. Alternatively, the agents may all receive a collective reward when any order has been completed by any robot. The latter case, when all agents receive identical rewards, is also called *common-reward* and is an important special case in MARL (discussed further in Chapter 3). A simple simulator of such a multi-robot warehouse is described in Section 11.3.4. Krnjaic et al. (2022) and Sultana et al. (2020) applied MARL algorithms to such warehouse applications.

1.3.2 Competitive Play in Board Games and Video Games

MARL can be used to train agents to achieve strong competitive play in board and card games (e.g. Backgammon, Chess, Go, Poker) and multi-player video games (e.g. shooting games, racing games, etc.). Each agent assumes the role of one of the players in the game. Agents may have actions available to move individual pieces or units to specific positions, placing specific cards, shooting target units, etc. Agents may observe the full game state, such as the entire game board with all pieces; or they may receive only a partial observation such as their own cards but not the cards of other players, or a partial view of the game map. Depending on the rules and mechanics of the game, the agents may or may not observe the chosen actions of other agents. In fully competitive games with two agents, one agent's reward is the negative of the other agent's reward. Thus, an agent may receive a reward of +1 for winning the game, in which case the other losing agent will receive a reward of -1, and vice versa. This property is referred to as *zero-sum reward*, and is another important special case in MARL. With this setup, during MARL training, the agents will learn to exploit each other's weaknesses and improve their play to eliminate their own weaknesses, leading to strong competitive play. Many different types of board games, card games, and video games have been tackled using MARL approaches (Tesauro 1994; Silver et al. 2018; Vinyals et al. 2019; Bard et al. 2020; Meta Fundamental AI

Research Diplomacy Team (FAIR) et al. 2022; Perolat et al. 2022; Wurman et al. 2022).

1.3.3 Autonomous Driving

Autonomous driving in urban environments and highways involves frequent interactions with other vehicles. Using MARL, we could train control policies for multiple vehicles to navigate through complicated interaction scenarios, such as driving through busy junctions and roundabouts, and merging onto highways. The actions of an agent might be the continuous controls for a vehicle, such as steering and acceleration/breaking; or discrete actions such as deciding between different maneuvers to execute (e.g. change lane, turning, overtaking). An agent may receive observations about its own controlled vehicle (e.g. position on lane, orientation, speed) as well as observations about other nearby vehicles. Observations about other vehicles may be uncertain due to sensor noise, and may be *incomplete* due to partial observability caused by occlusions (e.g. other vehicles blocking the agent's view). The reward of each agent can involve multiple factors. At a basic level, agents must avoid collisions and so any collision would result in a large negative reward. In addition, we want the agents to produce efficient and natural driving behaviour, so there may be positive rewards for minimising driving times, and negative rewards for abrupt acceleration/breaking and frequent lane changes. Therefore, in contrast to the multi-robot warehouse (agents have the same goal) and game playing (agents have opposed goals), here we have a mixed-motive scenario in which agents collaborate to avoid collisions but are also self-interested based on their desire to minimise driving times and drive smoothly. This case is referred to as *general-sum reward* and is among the most challenging tasks in MARL. Several works have used MARL for autonomous driving tasks (e.g. Shalev-Shwartz, Shammah, and Shashua 2016; Peake et al. 2020; Zhou, Luo, et al. 2020).

1.3.4 Automated Trading in Electronic Markets

Software agents can be developed to assume the roles of traders in electronic markets (Wellman, Greenwald, and Stone 2007). The typical objective of agents in a market is to maximise their own returns by placing buying and selling orders. Thus, agents have actions to buy or sell commodities at specified times, prices, and quantities. Agents receive observations about price developments in the market and other key performance indicators, and possibly some information about the current state of the order book. In addition, the agents may need to model and monitor external events and processes based on diverse types of observed information, such as news pertaining to certain companies, or energy

demand and usage of own managed households in peer-to-peer energy markets. The reward of an agent could be defined as a function of gains and losses made over a certain period of time, for example at the end of each trading day, quarter, or year. Thus, trading in electronic markets is another example of a mixed-motive scenario, since the agents need to collaborate in some sense to agree on sell-buy prices while aiming to maximise their own individual gains. MARL algorithms have been proposed for different types of electronic markets, including financial markets and energy markets (Roesch et al. 2020; Qiu et al. 2021; Shavandi and Khedmati 2022).

1.4 Challenges of MARL

Various challenges exist in multi-agent reinforcement learning which stem from aspects such as that agents may have conflicting goals, that agents may have different partial views of their environment, and that agents are learning concurrently to optimise their policies. Below we outline some of the main challenges, which will be discussed in more detail in Chapter 5.

Non-stationarity caused by learning agents An important characteristic of MARL is non-stationarity caused by the continually changing policies of the agents during their learning processes. This non-stationarity can lead to a moving target problem, because each agent adapts to the policies of other agents whose policies in turn also adapt to changes in other agents, thereby potentially causing cyclic and unstable learning dynamics. This problem is further exacerbated by the fact that the agents may learn different behaviours at different rates as a result of their different rewards and local observations. Thus, the ability to handle such non-stationarity in a robust way is often a crucial aspect in MARL algorithms, and has been the subject of much research.

Multi-agent credit assignment Temporal credit assignment in RL is the problem of determining which past actions contributed to a received reward. In MARL, this problem is compounded by the additional problem of determining *whose* action contributed to the reward. To illustrate, consider the level-based foraging example shown in Figure 1.2 and assume all agents choose the “collect” action, following which they receive a collective reward of +1. Given only this state/action/reward information, it can be highly non-trivial to disentangle the contribution of each agent to the received reward, in particular that the agent on the left did not contribute to the reward since its action had no effect (the agent’s level was not large enough). While ideas based on counterfactual reasoning can address this

problem in principle, it is still an open problem how to resolve multi-agent credit assignment in an efficient and scalable way.

Optimality of policies and equilibrium selection When are the policies of agents in a multi-agent system *optimal*? In single-agent RL, a policy is optimal if it achieves maximum expected returns in each state. However, in MARL, the returns of one agent’s policy also depend on the other agents’ policies, and thus we require more sophisticated notions of optimality. Chapter 4 presents a range of solution concepts, such as equilibrium-type solutions in which each agent’s policy is in some specific sense optimal with respect to the other agents’ policies. In addition, while in the single-agent case all optimal policies yield the same expected return to the agent, in a multi-agent system (where agents may receive different rewards) there may be multiple equilibrium solutions, and each equilibrium may entail different returns to different agents. Thus, there is an additional challenge of agents having to essentially negotiate during learning which equilibrium to converge to (Harsanyi and Selten 1988a). A central goal of MARL research is to develop learning algorithms which can learn agent policies that robustly converge to a particular solution type.

Scaling in number of agents In a multi-agent system, the total number of possible action combinations between agents may grow exponentially with the number of agents. This is in particular the case if each added agent comes with its own additional action variables. For example, in level-based foraging each agent controls a robot, and adding another agents comes with its own associated action variable to control a robot. (But see Section 5.4.4 for a counter-example without exponential growth.) In the early days of MARL research, it was common to use only two agents to avoid issues with scaling. Even with today’s deep learning-based MARL algorithms, it is common to use a number of agents between 2 and 10. How to handle many more agents in an efficient and robust way is an important goal in MARL research.

1.5 Agendas of MARL

An influential article by Shoham, Powers, and Grenager (2007), titled “*If multi-agent learning is the answer, what is the question?*”, describes several distinct

agendas which have been pursued in MARL research.⁴ The agendas differ in their motivations and goals for using MARL, as well as the criteria by which progress and success are measured. In their article, Shoham et al. made the important point that one must be clear about the purpose and goals when using MARL. We describe the main agendas as follows:

Computational The computational agenda uses MARL as an approach to compute solutions for game models. A solution consists of a collection of decision policies for the agents which satisfy certain properties (e.g. Nash equilibrium and other solution concepts, discussed in Chapter 4). Once computed, a solution could be deployed in an application of the game to control the agents (see Section 1.3 for some examples), or it may be used to conduct further analysis of the game. Thus, in the computational agenda, MARL algorithms compete with other direct methods to compute game solutions which do not rely on iterative learning (Nisan et al. 2007). Such direct methods can be significantly more efficient than MARL algorithms for certain types of games, but they typically require full knowledge of the game (including the reward functions of all agents). In contrast, most MARL algorithms can learn without full knowledge of the game.

Descriptive The descriptive agenda uses MARL to study the behaviours of natural agents, such as humans and animals, when learning in a population. This agenda typically begins by proposing a certain MARL algorithm which is designed to mimic how humans or animals adapt their actions based on past interactions. Methods from social sciences and behavioural economics can be used to test how closely a MARL algorithm matches the behaviour of a natural agent, such as via controlled experimentation in a laboratory setting. This is then followed by analysing whether a population of such natural agents converges to a certain kind of equilibrium solution if all agents use the proposed MARL algorithm.

Prescriptive The prescriptive agenda focuses specifically on the behaviours and performance of agents *during learning*, and asks how they should learn to achieve a given set of criteria. Different criteria have been proposed in this context. One possible criterion is that the average reward received by a learning agent should not fall below a certain threshold during learning, regardless of how other agents may be learning. An additional criterion

4. The article of Shoham, Powers, and Grenager (2007) was part of a special issue called “Foundations of Multi-Agent Learning” which was published in the *Artificial Intelligence* journal (Vohra and Wellman 2007). This special issue contains many interesting articles from some of the early contributors in MARL research, including responses to Shoham et al.’s article.

might be that the learning agent must learn optimal actions if the other agents come from a certain class of agents (such as static, non-learning agents). Thus, mimicking the behaviours of natural agents is not necessarily the goal in this agenda, nor is convergence to a particular solution concept such as equilibrium solutions.

Our perspective in this book is to view MARL as a method to optimise the decision policies of agents toward defined criteria. As such, the book primarily covers ideas and algorithms from the computational and prescriptive agendas. In particular, the computational agenda is closest to our perspective, which is reflected in the structure of the book by first introducing game models and solution concepts, followed by algorithms designed to learn such solutions. Using MARL to study the learning behaviours of natural agents, i.e. the descriptive agenda, is outside the scope of this book.

1.6 Book Contents and Structure

This book provides an introduction to the theory and practice of multi-agent reinforcement learning, suitable for university students, researchers, and practitioners. Following this introductory chapter, the remainder of the book is divided into two parts.

Part I of the book provides foundational knowledge about the basic models and concepts used in MARL. Specifically, Chapter 2 gives an introduction to the theory and tabular algorithms of single-agent RL. Chapter 3 introduces the basic game models to define concepts such as states, actions, observations, and rewards in a multi-agent environment. Chapter 4 then introduces a series of solution concepts which define what it means to solve these game models; that is, what it means for agents to act optimally. The final two chapters in this part of the book explore a range of MARL approaches for computing solutions in games: Chapter 5 presents basic concepts such as central and independent learning, and discusses core challenges in MARL. Chapter 6 introduces different classes of foundation algorithms developed in MARL research and discusses their learning properties.

Part II of the book focuses on contemporary research in MARL which leverages deep learning techniques to create new powerful MARL algorithms. We start by providing introductions to deep learning and deep reinforcement learning in Chapters 7 and 8, respectively. Building on the previous two chapters, Chapter 9 introduces several of the most important MARL algorithms developed in recent years, including ideas such as centralised training with decentralised execution, value decomposition, and parameter sharing. Chapter 10 provides

practical guidance when implementing and using MARL algorithms and how to evaluate the learned policies. Finally, Chapter 11 describes examples of multi-agent environments which have been developed in MARL research.

One of the goals of this book is to provide a starting point for readers who want to use the MARL algorithms discussed in this book in practice, as well as develop their own algorithms. Thus, the book comes with its own MARL codebase (downloadable from the book’s website) which was developed in the Python programming language, providing implementations of many existing MARL algorithms which are self-contained and easy to read. Chapter 10 uses code snippets from the codebase to explain implementation details of the important concepts underlying the algorithms presented in the earlier chapters. We hope that the provided code will be useful to readers in understanding MARL algorithms as well as getting started with using them in practice.

I FOUNDATIONS OF MULTI-AGENT REINFORCEMENT LEARNING

Part I of this book covers the foundations of multi-agent reinforcement learning. The chapters in this part focus on basic questions, including how to represent the mechanics of a multi-agent system via game models; how to define a learning objective in games to specify optimal agent behaviours; and how reinforcement learning methods can be used to learn optimal agent behaviours, as well as the complexities and challenges involved in multi-agent learning.

Chapter 2 provides an introduction to the basic models and algorithmic concepts of reinforcement learning, including Markov decision processes, dynamic programming, and temporal-difference learning. Chapter 3 then introduces game models to represent interaction processes in a multi-agent system, including basic normal-form games, stochastic games, and partially observable stochastic games. Chapter 4 introduces a range of solution concepts from game theory to define optimal agent policies in games, including equilibrium-type solutions such as minimax, Nash, and correlated equilibrium, as well as other concepts such as Pareto optimality, welfare/fairness, and no-regret. We provide examples for each solution concept and also discuss important conceptual limitations. Together, a game model and a solution concept define a learning problem in multi-agent reinforcement learning.

Building on the previous chapters, Chapters 5 and 6 look at how to use reinforcement learning techniques to learn optimal agent policies in a game. Chapter 5 begins by defining the general learning process in games and different convergence types, and introduces the basic concepts of central learning and independent learning that reduce the multi-agent learning problem to a single-agent learning problem. The chapter then discusses the central challenges in multi-agent reinforcement learning, including non-stationarity, multi-agent credit assignment, equilibrium selection, and scaling to many agents. Chapter 6 introduces several classes of foundation algorithms for multi-agent reinforcement learning that go beyond the basic approaches introduced in the prior chapter, and discusses their convergence properties.

2 Reinforcement Learning

Multi-agent reinforcement learning (MARL) is, in essence, reinforcement learning (RL) applied to multi-agent game models to learn optimal policies for the agents. Thus, MARL is deeply rooted in both RL theory and game theory. This chapter provides a basic introduction to the theory and algorithms of RL when there is only a single agent for which we want to learn an optimal policy. We will begin by providing a general definition of RL, following which we will introduce the Markov decision process (MDP) as the foundational model used in RL to represent single-agent decision processes. Based on the MDP model, we will define basic concepts such as expected returns, optimal policies, value functions, and Bellman equations. The goal in an RL problem is to learn an optimal policy that chooses actions to achieve some objective, such as maximising the expected (discounted) return in each state of the environment (Figure 2.1).

We will then introduce two basic families of algorithms to compute optimal policies for MDPs: dynamic programming and temporal-difference learning. Dynamic programming requires complete knowledge of the MDP specification and uses this knowledge to compute optimal value functions and policies. In contrast, temporal-difference learning does not require complete knowledge of the MDP, instead learning optimal value functions and policies by interacting with the environment and generating experiences. Most of the MARL algorithms introduced in Chapter 6 build on these families of algorithms and essentially extend them to game models.

Part I of this book focuses on the basic models and concepts used in MARL, and as such this chapter focuses only on the basic RL concepts which are required to understand the following chapters of this part of the book. In particular, topics such as value and policy function approximation are not covered in this chapter. These latter topics will be covered in Chapters 7 and 8 in Part II of the book.

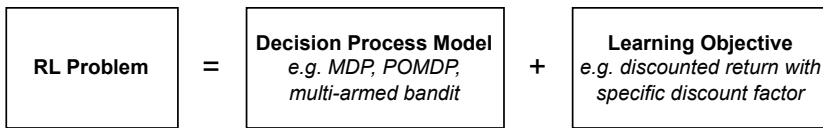


Figure 2.1: An RL problem is defined by the combination of a decision process model which defines the mechanics of the agent-environment interaction, and a learning objective which specifies the properties of the optimal policy to be learned (e.g. maximise expected discounted return in each state).

2.1 General Definition

We begin by providing a general definition of reinforcement learning:

Reinforcement learning (RL) algorithms learn solutions for sequential decision processes via repeated interaction with an environment.

This definition raises three main questions:

- What is a sequential decision process?
- What is a solution to the process?
- What is learning via repeated interaction?

A sequential decision process is defined by an agent which makes decisions over multiple time steps within an environment to achieve a specified goal. In each time step, the agent receives an observation from the environment and chooses an action. In a fully observable setting, as we assume in this chapter, the agent observes the full state of the environment; but in general, observations may be incomplete and noisy. Given the chosen action, the environment may change its state according to some transition dynamics and send a scalar reward signal to the agent. Figure 2.2 summarises this process.

A solution to the decision process is an optimal decision policy for the agent, which chooses actions in each state to achieve some specified learning objective. Typically, the learning objective is to maximise the expected return for the agent in each possible state.⁵ The return in a state when following a given policy is defined as the sum of rewards received over time from that state onward. Thus, RL assumes that the goal in the decision process can in principle be framed as the maximisation of expected returns.

5. Other learning objectives can be specified, such as maximising the average reward (Sutton and Barto 2018) and different types of “risk-sensitive” objectives (Mihatsch and Neuneier 2002).

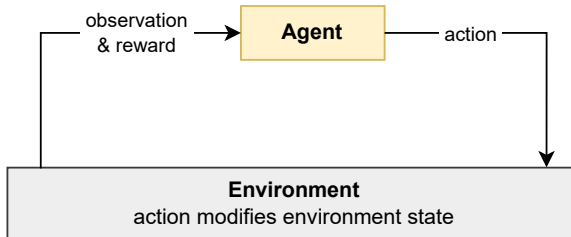


Figure 2.2: Basic reinforcement learning loop for a single-agent system.

Finally, RL algorithms learn such optimal policies by trying different actions in different states and observing the outcomes. This way of learning is sometimes described as “trial and error”, since the actions may lead to positive or negative outcomes which are not known beforehand and, therefore, must be discovered by trying the actions. A central problem in this learning process, often called the *exploration-exploitation dilemma*, is how to balance exploring the outcomes of different actions versus sticking with actions which are currently believed to be best. Exploration may discover better actions but can accrue low rewards in the process, while exploitation achieves a certain level of returns but may not discover the optimal actions.

RL is a type of machine learning which differs from other types such as supervised learning and unsupervised learning. In supervised learning, we have access to a set of labelled input-output pairs $\{x_i, y_i\}$ of some unknown function $f(x_i) = y_i$, and the goal is to learn this function using the data. In unsupervised learning, we have access to some unlabelled data $\{x_i\}$ and the goal is to identify some useful structure within the data. RL is neither of these: RL is not supervised learning because the reward signals do not tell the agent which action to take in each state x_i , and thus do not act as a supervision signal y_i . This is because some actions may give lower immediate reward, but may lead to states from which the agent can eventually receive higher rewards. RL also differs from unsupervised learning because the rewards, while not a supervised signal, act as a proxy from which to learn an optimal policy.

In the following sections, we will formally define these concepts – sequential decision processes, optimal policies, and learning by interaction – within a framework called the Markov decision process.

2.2 Markov Decision Processes

The standard model used in RL to define the sequential decision process is the *Markov decision process*:

Definition 1 (Markov decision process (MDP)) A finite Markov decision process (MDP) consists of:

- Finite set of states S
- Finite set of actions A
- Reward function $\mathcal{R} : S \times A \times S \mapsto \mathbb{R}$
- State transition probability function $\mathcal{T} : S \times A \times S \mapsto [0, 1]$
- Initial state distribution $\Pr^0 : S \mapsto [0, 1]$

An MDP starts in an initial state $s^0 \in S$ which is sampled from \Pr^0 . At time t , the agent observes the current state $s^t \in S$ of the MDP and chooses an action $a^t \in A$ with probability given by its policy, $\pi(a^t | s^t)$, which is conditioned on the state. Given the state s^t and action a^t , the MDP transitions into a next state $s^{t+1} \in S$ with probability given by $\mathcal{T}(s^t, a^t, s^{t+1})$, and the agent receives a reward $r^t = \mathcal{R}(s^t, a^t, s^{t+1})$. We will also write this probability as $\mathcal{T}(s^{t+1} | s^t, a^t)$ to emphasise that it is conditioned on the state-action pair s^t, a^t . These steps are repeated until some termination criterion is satisfied, such as reaching a maximum number of time steps, or reaching a terminal state; or the steps may be repeated for an infinite number of time steps if the MDP never terminates. Each independent run of the above process is referred to as an *episode*.

A finite MDP can be compactly shown as a finite state machine. Figure 2.3 shows an example MDP, in which a Mars rover has collected some samples and must return to the base station. From the *Start* state, there are two paths which lead to the *Base* target state. The rover can travel down a steep mountain slope (action *right*) which will take it directly to the base station. However, there is a high probability of 0.5 that the rover will fall down the cliff and be destroyed (-10 reward). Or the rover can travel through a longer path (action *left*, then two times action *right*) which passes by two sites (*Site A* and *Site B*) before arriving at the base station. However, it takes a day to arrive at each site (-1 reward) and there is a probability of 0.3 that the rover will get stuck on the rocky ground and be immobilised (-3 reward). Arriving at the base station gives a reward of +10.

The term “Markov” comes from the *Markov property*, which states that the future state and reward are conditionally independent of past states and actions, given the current state and action:

$$\Pr(s^{t+1}, r^t | s^t, a^t, s^{t-1}, a^{t-1}, \dots, s^0, a^0) = \Pr(s^{t+1}, r^t | s^t, a^t) \quad (2.1)$$

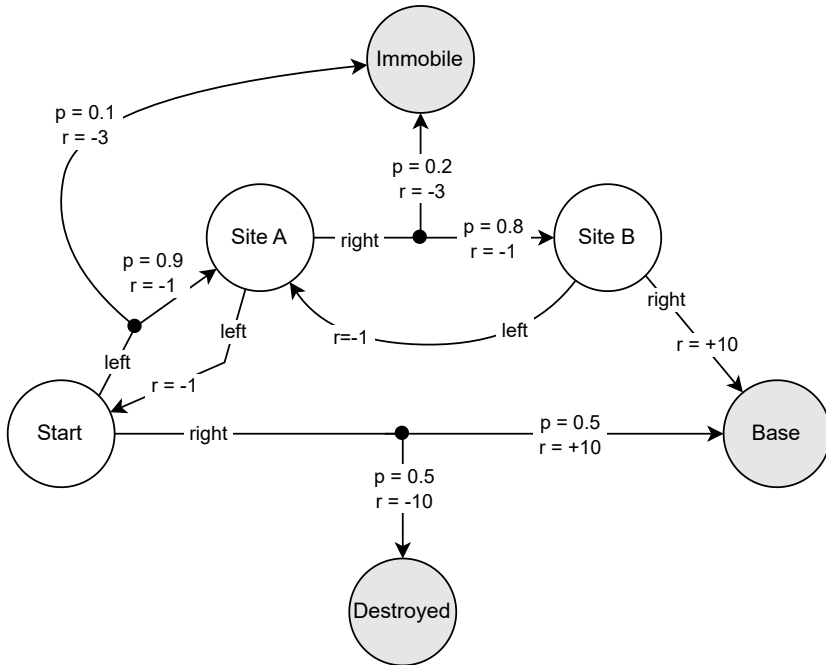


Figure 2.3: Mars Rover MDP. States are shown as circles, where *Start* is the initial state. Each non-terminal state (white) has two possible actions, *right* and *left*, which are shown as directed edges with associated transition probabilities (p) and rewards (r). Grey shaded circles mark terminal states.

MDPs satisfy this Markov property, which means that the current state provides sufficient information to choose optimal actions – past states and actions are not relevant.

The most common assumption in RL is that the dynamics of the MDP, in particular the transition and reward functions \mathcal{T} and \mathcal{R} , are *a priori* unknown to the agent. Typically, the only parts of the MDP which are assumed to be known are the action space A and the state space S .

While Definition 1 defines MDPs with finite, i.e. *discrete*, states and actions, MDPs can also be defined with continuous states and actions, or a mixture of both discrete and continuous elements. Moreover, while the reward function \mathcal{R} as defined here is deterministic, MDPs can also define a probabilistic reward function such that $\mathcal{R}(s, a, s')$ gives a distribution over possible rewards.

The *multi-armed bandit problem* is an important special case of the MDP in which the reward function is probabilistic and unknown to the agent, and

there is only a single state in S . Thus, in such a bandit problem, each action produces a reward from some unknown reward distribution, and the goal is effectively to find the action which yields the highest expected reward as quickly as possible. Bandit problems have been used as a basic model to study the exploration-exploitation dilemma (Lattimore and Szepesvari 2020).

MDPs assume that the agent can fully observe the state of the environment in each time step. In many applications, an agent only observes partial and noisy information about the environment. *Partially observable Markov decision processes* (POMDPs) generalise MDPs by defining a decision process in which the agent receives observations o^t rather than directly observing the state s^t , and these observations depend in a probabilistic or deterministic way on the state. Thus, in general, the agent will need to take into account the history of past observations o^0, o^1, \dots, o^t in order to infer the possible current states s^t of the environment. POMDPs are a special case of the partially observable stochastic game (POSG) model introduced in Chapter 3; namely, a POMDP is a POSG in which there is only one agent. We refer to Section 3.4 for a more detailed definition and discussion of partial observability in decision processes.

We have defined the MDP to model a decision process, but we have not yet specified the learning objective in the MDP. In the next section, we will introduce *discounted returns* as the learning objective in MDPs. Together, an MDP and learning objective specify an RL problem.

2.3 Expected Discounted Returns and Optimal Policies

Given a policy π which specifies action probabilities in each state, and assuming that each episode in the MDP terminates in time step T , the total *return* in an episode is the cumulative reward received over time

$$r^0 + r^1 + \dots + r^{T-1}. \quad (2.2)$$

Due to the stochastic nature of the MDP, it may not be possible to maximise this return in all episodes. This is because some actions may lead to different outcomes with certain probabilities, and these outcomes are outside the control of the agent. Therefore, the agent can maximise the *expected return* given by

$$\mathbb{E}_\pi [r^0 + r^1 + \dots + r^{T-1}] \quad (2.3)$$

where the expectation assumes that the initial state is sampled from the initial state distribution (i.e. $s^0 \sim \Pr^0$), that the agent follows policy π to select actions (i.e. $a^t \sim \pi(s^t)$), and that successor states are governed by the state transition probabilities (i.e. $s^{t+1} \sim \mathcal{T}(\cdot | s^t, a^t)$).

The above definition of total returns is guaranteed to be finite for a terminating MDP. However, in non-terminating MDPs the total return may be infinite, in which case returns may not be informative to distinguish the performance of different policies that achieve infinite returns. The standard approach to ensuring finite returns in non-terminating MDPs is to use a *discount factor* $\gamma \in [0, 1]$, based on which we define the *discounted return*

$$\mathbb{E}_\pi[r^0 + \gamma r^1 + \gamma^2 r^2 + \dots] \quad (2.4)$$

$$= \mathbb{E}_\pi \left[\sum_{t=0}^{\infty} \gamma^t r^t \right]. \quad (2.5)$$

For $\gamma < 1$ and assuming that rewards are constrained to lie in a finite range $[r_{\min}, r_{\max}]$, the discounted return is guaranteed to be finite

$$\sum_{t=0}^{\infty} \gamma^t r^t \leq r_{\max} \sum_{t=0}^{\infty} \gamma^t = r_{\max} \frac{1}{1-\gamma} \quad (2.6)$$

where the right-hand fraction in the above equation is the closed form of the geometric series given by $\sum_{t=0}^{\infty} \gamma^t$.

The discount factor has two equivalent interpretations. One interpretation is that $(1 - \gamma)$ is the probability with which the MDP terminates after each time step. Thus, the probability that the MDP terminates after a total of $l > 0$ time steps is $\gamma^{l-1}(1 - \gamma)$, where γ^{l-1} is the probability of continuing (not terminating) in the first $l - 1$ time steps, multiplied by the probability $(1 - \gamma)$ of terminating in the following time step. For example, in the Mars Rover MDP shown in Figure 2.3, we can specify a discount factor of $\gamma = 0.95$ to model the fact that the rover's battery may fail with a probability of 0.05 after each state transition. The second interpretation is that the agent gives "weight" γ^t to reward r^t received at time t . Thus, a γ close to 0 leads to a myopic agent which cares more about near-term rewards, while a γ close to 1 leads to a farsighted agent which also values distant rewards. In either case, it is important to note that the discount rate is part of the learning objective, and not a tunable algorithm parameter; γ is a fixed parameter specified by the learning objective. In the remainder of this book, whenever we refer to returns, we specifically mean discounted returns with some discount factor γ .

Finally, the above definition of discounted returns can be made to work for both non-terminating and terminating MDPs via the convention of *absorbing states*. If the MDP reaches a terminal state, we define this state to be absorbing in that any subsequent actions in this state will transition the MDP into the same absorbing state with probability 1, and will give a reward of 0 to the agent.

Thus, once the MDP reaches an absorbing state, it will forever remain in that state and the agent will not accrue any more rewards.

Based on the above definition of discounted returns and absorbing states, we can now define the solution to the MDP as the *optimal policy* π^* that maximises the expected discounted return.

2.4 Value Functions and Bellman Equation

Based on the Markov property defined in Equation 2.1, we know that given the current state and action, the future states and rewards are independent of the past states and actions. This implies independence of past rewards, since rewards are a function of the states and actions, $r^t = \mathcal{R}(s^t, a^t, s^{t+1})$. Therefore, in an MDP, the expected return can be defined individually for each state $s \in S$. This gives rise to the concept of *value functions* which are central to much of RL theory and many algorithms.

First, note that the discounted reward sequence can be written in recursive form as

$$u^t = r^t + \gamma r^{t+1} + \gamma^2 r^{t+2} + \dots \quad (2.7)$$

$$= r^t + \gamma u^{t+1} \quad (2.8)$$

Given a policy π , the *state-value function* $V^\pi(s)$ gives the “value” of state s under π , which is the expected return when starting in state s and following policy π to select actions, formally

$$V^\pi(s) = \mathbb{E}_\pi [u^t | s^t = s] \quad (2.9)$$

$$= \mathbb{E}_\pi [r^t + \gamma u^{t+1} | s^t = s] \quad (2.10)$$

$$= \sum_{a \in A} \pi(a | s) \sum_{s' \in S} \mathcal{T}(s' | s, a) [\mathcal{R}(s, a, s') + \gamma \mathbb{E}_\pi [u^{t+1} | s^{t+1} = s']] \quad (2.11)$$

$$= \sum_{a \in A} \pi(a | s) \sum_{s' \in S} \mathcal{T}(s' | s, a) [\mathcal{R}(s, a, s') + \gamma V^\pi(s')]. \quad (2.12)$$

This recursive equation is called the *Bellman equation* in honour of Richard Bellman and his pioneering work (Bellman 1957).

The Bellman equation for V^π defines a system of m linear equations with m variables, where $m = |S|$ is the number of states in the finite MDP:

$$V^\pi(s_1) = \sum_{a \in A} \pi(a | s_1) \sum_{s' \in S} \mathcal{T}(s' | s_1, a) [\mathcal{R}(s_1, a, s') + \gamma V^\pi(s')] \quad (2.13)$$

$$V^\pi(s_2) = \sum_{a \in A} \pi(a | s_2) \sum_{s' \in S} \mathcal{T}(s' | s_2, a) [\mathcal{R}(s_2, a, s') + \gamma V^\pi(s')] \quad (2.14)$$

⋮

$$V^\pi(s_m) = \sum_{a \in A} \pi(a | s_m) \sum_{s' \in S} \mathcal{T}(s' | s_m, a) [\mathcal{R}(s_m, a, s') + \gamma V^\pi(s')] \quad (2.15)$$

where $V^\pi(s_k)$ for $k = 1, \dots, m$ are the variables in the equation system, and $\pi(a | s_k)$, $\mathcal{T}(s' | s_k, a)$, $\mathcal{R}(s_k, a, s')$, and γ are constants. This equation system has a unique solution given by the value function V^π for policy π . If all elements of the MDP are known, then one can solve this equation system to obtain V^π using any method to solve linear equation systems (such as Gauss elimination).

Analogous to state-value functions, we can define *action-value functions* $Q^\pi(s, a)$ which give the expected return when selecting action a in state s and then following policy π to select actions subsequently,

$$Q^\pi(s, a) = \mathbb{E}_\pi [u^t | s^t = s, a^t = a] \quad (2.16)$$

$$= \mathbb{E}_\pi [r^t + \gamma u^{t+1} | s^t = s, a^t = a] \quad (2.17)$$

$$= \sum_{s' \in S} \mathcal{T}(s' | s, a) [\mathcal{R}(s, a, s') + \gamma V^\pi(s')] \quad (2.18)$$

$$= \sum_{s' \in S} \mathcal{T}(s' | s, a) [\mathcal{R}(s, a, s') + \gamma \sum_{a' \in A} \pi(a' | s') Q^\pi(s', a')]. \quad (2.19)$$

This admits an analogous system of linear equations that has a unique solution given by Q^π .

A policy π is optimal in the MDP if the policy's (state or action) value function is the *optimal value function* of the MDP, defined as

$$V^*(s) = \max_{\pi'} V^{\pi'}(s), \quad \forall s \in S \quad (2.20)$$

$$Q^*(s, a) = \max_{\pi'} Q^{\pi'}(s, a), \quad \forall s \in S, a \in A \quad (2.21)$$

We use π^* to denote any optimal policy with optimal value function V^* or Q^* .

Because of the Bellman equation, this means that for any optimal policy π^* we have

$$\forall \pi' \forall s : V^*(s) \geq V^{\pi'}(s). \quad (2.22)$$

Thus, maximising the expected return in an MDP amounts to maximising the expected return in each possible state $s \in S$.

In fact, we can write the optimal value functions without reference to the policy, using the *Bellman optimality equations*:

$$V^*(s) = \max_{a \in A} \sum_{s' \in S} \mathcal{T}(s' | s, a) [\mathcal{R}(s, a, s') + \gamma V^*(s')] \quad (2.23)$$

$$Q^*(s, a) = \sum_{s' \in S} \mathcal{T}(s' | s, a) \left[\mathcal{R}(s, a, s') + \gamma \max_{a' \in A} Q^*(s', a') \right] \quad (2.24)$$

The Bellman optimality equations define a system of m non-linear equations, where m is again the number of states in the finite MDP. The non-linearity is due to the max-operator used in the equations. The unique solution to the system is the optimal value function V^*/Q^* .

Once we know the optimal action-value function Q^* , the optimal policy π^* is simply derived by choosing actions with maximum value (i.e. expected return) in each state,

$$\pi^*(s) = \arg \max_{a \in A} Q^*(s, a) \quad (2.25)$$

where we use the notation in Equation 2.25 as a convenient shorthand to denote the deterministic policy which assigns probability 1 to the arg max-action in state s . If multiple actions have the same maximum value under Q^* , then the optimal policy can assign arbitrary probabilities to these actions (the probabilities must sum to 1). Thus, while the optimal value function of the MDP is always unique, there may be multiple⁶ optimal policies which have the same optimal (unique) value function.

In the following sections, we will present two families of algorithms to compute optimal value functions and optimal policies. The first, dynamic programming, uses the Bellman (optimality) equations as an iterative operator to estimate value functions, and requires knowledge of the complete MDP including the reward function and state transition probabilities. The second, temporal-difference learning, does not need complete knowledge of the MDP and instead uses sampled experiences from interactions with the environment to update value estimates.

6. In fact, if multiple actions have maximum value, then there will be an *infinite* number of optimal policies since the set of possible probability assignments to these actions spans a continuous (infinite) space.

2.5 Dynamic Programming

Dynamic programming (DP)⁷ is a family of algorithms to compute value functions and optimal policies in MDPs (Bellman 1957; Puterman 2014). DP algorithms use the Bellman equations as operators to iteratively estimate the value function and optimal policy. Thus, DP algorithms require complete knowledge of the MDP model, including the reward function \mathcal{R} and state transition probabilities \mathcal{T} . While DP algorithms do not “interact” with the environment as per our definition in Section 2.1, the basic concepts underlying DP are an important building block in RL theory, including temporal-difference learning presented in Section 2.6.

The basic DP approach is *policy iteration*, which alternates between two tasks:

- **Policy evaluation:** compute value function V^π for current policy π
- **Policy improvement:** improve current policy π with respect to V^π

Thus, policy iteration produces a sequence of policy and value function estimates, starting with some initial policy π^0 (e.g. uniform-random policy) and value function V^0 (e.g. zero vector):

$$\pi^0 \rightarrow V^{\pi^0} \rightarrow \pi^1 \rightarrow V^{\pi^1} \rightarrow \pi^2 \rightarrow \dots \rightarrow V^* \rightarrow \pi^* \quad (2.26)$$

This sequence converges to the optimal value function, V^* , and optimal policy, π^* .

Recall that the Bellman equation for V^π defines a system of linear equations, which can be solved to obtain V^π . However, using Gauss elimination (the de-facto standard solution approach for linear equation systems) has time complexity $O(m^3)$, where m is the number of states of the MDP. *Iterative policy evaluation* instead applies the Bellman equation for V^π iteratively to produce successive estimates of V^π . The algorithm first initialises a vector $V^0(s)=0$ for all $s \in S$. It then repeatedly performs update sweeps for all states $s \in S$:

$$V^{k+1}(s) \leftarrow \sum_{a \in A} \pi(a|s) \sum_{s' \in S} \mathcal{T}(s'|s, a) [\mathcal{R}(s, a, s') + \gamma V^k(s')] \quad (2.27)$$

This sequence V^0, V^1, V^2, \dots converges to the value function V^π . Hence, in practice, we may stop the updates once there are no more changes between V^k, V^{k+1} after performing an update sweep.

7. The name “dynamic programming” does not refer to a type of software programming. The term “programming” here refers to a mathematical optimisation problem, analogous to how the term is used in linear programming and non-linear programming. Regarding the adjective “dynamic”, Bellman (1957) states in his preface that it “indicates that we are interested in processes in which time plays a significant role, and in which the order of operations may be crucial.”

Notice that Equation 2.27 updates the value estimate for a state s using value estimates for other states s' . This property is called *bootstrapping* and is a core property of many RL algorithms.

As an example, we apply iterative policy evaluation in the Mars Rover problem from Section 2.2 with $\gamma=0.95$. First, consider a policy π which in the initial state *Start* chooses action *right* with probability 1. (We ignore the other states since they will never be visited under this policy.) After we run iterative policy evaluation to convergence, we obtain a value of $V^\pi(\text{Start})=0$. From the MDP specification in Figure 2.3, it can be easily verified that this value is correct, since $V^\pi(\text{Start})=-10 * 0.5 + 10 * 0.5 = 0$. Now, consider a policy π which chooses both actions with equal probability 0.5 in the initial state, and action *right* with probability 1 in states *Site A* and *Site B*. In this case, the converged values are $V^\pi(\text{Start})=2.05$, $V^\pi(\text{Site A})=6.2$, and $V^\pi(\text{Site B})=10$. For both policies, the values for the terminal states are all 0, since these are absorbing states as defined in Section 2.3.

To see why this process converges to the value function V^π , one can show that the Bellman operator defined in Equation 2.27 is a *contraction mapping*. A mapping $f : \mathcal{X} \mapsto \mathcal{X}$ on a $\|\cdot\|$ -normed complete vector space \mathcal{X} is a γ -contraction, for $\gamma \in [0, 1)$, if for all $x, y \in \mathcal{X}$:

$$\|f(x) - f(y)\| \leq \gamma \|x - y\| \quad (2.28)$$

By the Banach fixed-point theorem, if f is a contraction mapping, then for any initial vector $x \in \mathcal{X}$ the sequence $f(x), f(f(x)), f(f(f(x))), \dots$ converges to a unique fixed point $x^* \in \mathcal{X}$ such that $f(x^*)=x^*$. Using the max-norm $\|x\|_\infty = \max_i |x_i|$, it can be shown that the Bellman equation indeed satisfies Equation 2.28. First, rewrite the Bellman equation as follows:

$$V^\pi(s) = \sum_{a \in A} \pi(a|s) \sum_{s' \in S} \mathcal{T}(s'|s, a) [\mathcal{R}(s, a, s') + \gamma V^\pi(s')] \quad (2.29)$$

$$= \sum_{a \in A} \sum_{s' \in S} \pi(a|s) \mathcal{T}(s'|s, a) \mathcal{R}(s, a, s') + \sum_{a \in A} \sum_{s' \in S} \pi(a|s) \mathcal{T}(s'|s, a) \gamma V^\pi(s') \quad (2.30)$$

This can be written as an operator $f^\pi(v)$ over a value vector $v \in \mathbb{R}^{|S|}$

$$f^\pi(v) = r^\pi + \gamma M^\pi v \quad (2.31)$$

where $r^\pi \in \mathbb{R}^{|S|}$ is a vector with elements

$$r_s^\pi = \sum_{a \in A} \sum_{s' \in S} \pi(a|s) \mathcal{T}(s'|s, a) \mathcal{R}(s, a, s') \quad (2.32)$$

and $M^\pi \in \mathbb{R}^{|S| \times |S|}$ is a matrix with elements

$$M_{s,s'}^\pi = \sum_{a \in A} \pi(a|s) \mathcal{T}(s'|s, a). \quad (2.33)$$

Then, we have for any two value vectors v, u :

$$\|f^\pi(v) - f^\pi(u)\|_\infty = \|(r^\pi + \gamma M^\pi v) - (r^\pi + \gamma M^\pi u)\|_\infty \quad (2.34)$$

$$= \gamma \|M^\pi(v - u)\|_\infty \quad (2.35)$$

$$\leq \gamma \|v - u\|_\infty. \quad (2.36)$$

Therefore, the Bellman operator is a γ -contraction under the max-norm, and repeated application of the Bellman operator converges to a unique fixed point, which is V^π .⁸

Once we have computed the value function V^π , the policy improvement task modifies the policy π by making it *greedy* with respect to V^π for all $s \in S$:

$$\pi' = \arg \max_{a \in A} \mathcal{T}(s'|s, a) [\mathcal{R}(s, a, s') + \gamma V^\pi(s')] \quad (2.37)$$

$$= \arg \max_{a \in A} Q^\pi(s, a) \quad (2.38)$$

By the *Policy Improvement Theorem* (Sutton and Barto 2018), we know that if for all $s \in S$

$$\sum_{a \in A} \pi'(a|s) Q^\pi(s, a) \geq \sum_{a \in A} \pi(a|s) Q^\pi(s, a) \quad (2.39)$$

$$= V^\pi(s) \quad (2.40)$$

then π' must be as good as or better than π :

$$\forall s : V^{\pi'}(s) \geq V^\pi(s). \quad (2.41)$$

If after the policy improvement task, the greedy policy π' did not change from π , then we know that $V^{\pi'} = V^\pi$ and it follows for all $s \in S$:

$$V^{\pi'}(s) = \max_{a \in A} \mathbb{E}_\pi [r^t + \gamma V^\pi(s^{t+1}) | s^t = s, a^t = a] \quad (2.42)$$

$$= \max_{a \in A} \mathbb{E}_{\pi'} [r^t + \gamma V^{\pi'}(s^{t+1}) | s^t = s, a^t = a] \quad (2.43)$$

$$= \max_{a \in A} \sum_{s' \in S} \mathcal{T}(s'|s, a) [\mathcal{R}(s, a, s') + \gamma V^{\pi'}(s')] \quad (2.44)$$

$$= V^*(s) \quad (2.45)$$

8. The inequality in (2.36) holds because for each $s \in S$ we have $\sum_{s'} M_{s,s'}^\pi = 1$ and, therefore, $\|M^\pi x\|_\infty \leq \|x\|_\infty$ for any value vector x .

Algorithm 1 Value Iteration for MDP

1: Initialise: $V(s) = 0$ for all $s \in S$ 2: Repeat until V converged:

$$V(s) \leftarrow \max_{a \in A} \sum_{s' \in S} \mathcal{T}(s'|s, a) [\mathcal{R}(s, a, s') + \gamma V(s')], \quad \forall s \in S \quad (2.47)$$

3: Return optimal policy π^* with

$$\pi^*(s) = \arg \max_{a \in A} \sum_{s' \in S} \mathcal{T}(s'|s, a) [\mathcal{R}(s, a, s') + \gamma V(s')] \quad (2.48)$$

Therefore, if π' did not change from π after policy improvement, we know that π' must be the optimal policy of the MDP, and policy iteration is complete.

The above version of policy iteration, which uses iterative policy evaluation, makes multiple complete update sweeps over the entire state space S . *Value iteration* is a DP algorithm which combines one sweep of iterative policy evaluation and policy improvement in a single update equation, by using the Bellman optimality equation as an update operator:

$$V^{k+1}(s) \leftarrow \max_{a \in A} \sum_{s' \in S} \mathcal{T}(s'|s, a) [\mathcal{R}(s, a, s') + \gamma V^k(s')], \quad \forall s \in S \quad (2.46)$$

Following a similar argument as for the Bellman operator, it can be shown that the Bellman optimality operator defined in Equation 2.46 is a γ -contraction mapping. Thus, repeated application of value iteration converges to the unique fixed point which is the optimal value function V^* . Algorithm 1 provides the complete pseudocode for the value iteration algorithm.

Returning to the Mars Rover example, if we run value iteration to convergence, we obtain the optimal state values $V^*(Start) = 4.1$, $V^*(Site\ A) = 6.2$, and $V^*(Site\ B) = 10$. The optimal policy π^* chooses action *left* with probability 1 in the initial state, and action *right* with probability 1 in states *Site A* and *Site B*.

2.6 Temporal-Difference Learning

Temporal-difference learning (TD) is a family of RL algorithms which learn value functions and optimal policies based on experiences from interactions with the environment. The experiences are generated by following the agent-environment interaction loop shown in Figure 2.2: in state s^t , sample an action $a^t \sim \pi(\cdot | s^t)$ from policy π , then observe reward $r^t = \mathcal{R}(s^t, a^t, s^{t+1})$ and the new state $s^{t+1} \sim \mathcal{T}(\cdot | s^t, a^t)$. Like DP algorithms, TD algorithms learn value functions based on the Bellman equations and bootstrapping – estimating the value of a

state or action using value estimates of other states/actions. However, unlike DP algorithms, TD algorithms do not require complete knowledge of the MDP, such as the reward function and state transition probabilities. Instead, TD algorithms perform the policy evaluation and policy improvement tasks based solely on the experiences collected while interacting with the environment.

TD algorithms use the following general update rule to learn action-value functions:

$$Q(s^t, a^t) \leftarrow Q(s^t, a^t) + \alpha [\mathcal{X} - Q(s^t, a^t)] \quad (2.49)$$

where \mathcal{X} is the update *target*, and $\alpha \in (0, 1]$ is the learning rate (or step size). The target \mathcal{X} is constructed based on experience samples (s^t, a^t, r^t, s^{t+1}) collected from interactions with the environment. A wide range of options exist to specify the target, and here we present two basic variants.

Recall the Bellman equation for the action-value function Q^π for policy π :

$$Q^\pi(s, a) = \sum_{s' \in S} \mathcal{T}(s' | s, a) \left[\mathcal{R}(s, a, s') + \gamma \sum_{a' \in A} \pi(a' | s') Q^\pi(s', a') \right] \quad (2.50)$$

Sarsa (Sutton and Barto 2018) is a TD algorithm which constructs an update target based on Equation 2.50 by replacing the summations over successor states s' and actions a' as well as the reward function $\mathcal{R}(s, a, s')$ with their corresponding elements from the experience tuple $(s^t, a^t, r^t, s^{t+1}, a^{t+1})$ (hence the name Sarsa):

$$\mathcal{X} = r^t + \gamma Q(s^{t+1}, a^{t+1}) \quad (2.51)$$

Here, the action a^{t+1} is sampled from the policy π in the successor state s^{t+1} , $a^{t+1} \sim \pi(\cdot | s^{t+1})$. The complete Sarsa update rule is specified as

$$Q(s^t, a^t) \leftarrow Q(s^t, a^t) + \alpha [r^t + \gamma Q(s^{t+1}, a^{t+1}) - Q(s^t, a^t)]. \quad (2.52)$$

If π stays fixed, then it can be shown under certain conditions that Sarsa will learn the action-value function $Q = Q^\pi$. The first condition is that all state-action combinations $(s, a) \in S \times A$ must be tried an infinite number of times during learning. The second condition is given by the “standard stochastic approximation conditions” which state that the learning rate α must be reduced over time in a way that satisfies the following conditions:

$$\forall s \in S, a \in A : \quad \sum_{k=0}^{\infty} \alpha_k(s, a) \rightarrow \infty \quad \text{and} \quad \sum_{k=0}^{\infty} \alpha_k(s, a)^2 < \infty \quad (2.53)$$

where $\alpha_k(s, a)$ denotes the learning rate used when applying the update rule in Equation 2.52 after the k -th selection of action a in state s . The left-hand sum in Equation 2.53 ensures that the learning rate is large enough to overcome initial learning conditions, while the right-hand sum ensures that the sequence will converge at a certain rate. Therefore, $\alpha_k(s, a) = \frac{1}{k}$ satisfies these conditions on the

Algorithm 2 Sarsa for MDPs

- 1: Initialise: $Q(s, a) = 0$ for all $s \in S, a \in A$
- 2: Repeat for every episode:
- 3: $a' \leftarrow \emptyset$
- 4: **for** $t = 0, 1, 2, \dots$ **do**
- 5: Observe current state s^t
- 6: **if** $a' = \emptyset$ **then**
- 7: With probability ϵ : choose random action $a^t \in A$
- 8: Else: choose action $a^t \in \arg \max_a Q(s^t, a)$
- 9: **else**
- 10: $a^t \leftarrow a'$
- 11: Apply action a^t , observe reward r^t and next state s^{t+1}
- 12: With probability ϵ : choose random action $a^{t+1} \in A$
- 13: Else: choose action $a^{t+1} \in \arg \max_a Q(s^{t+1}, a)$
- 14: $Q(s^t, a^t) \leftarrow Q(s^t, a^t) + \alpha [r^t + \gamma Q(s^{t+1}, a^{t+1}) - Q(s^t, a^t)]$
- 15: $a' \leftarrow a^{t+1}$

learning rate, while a constant learning rate $\alpha_k(s, a) = c$ does not. Nonetheless, in practice it is common to use a constant learning rate since learning rates that meet the above conditions, while theoretically sound, can lead to slow learning.

In order for Sarsa to learn the optimal value function Q^* and optimal policy π^* , it must gradually modify the policy π in a way that brings it closer to the optimal policy. To achieve this, π can be made greedy with respect to the value estimates Q , similarly to the DP policy improvement task defined in Equation 2.38. However, making π fully greedy and deterministic would violate the first condition of trying all state-action combinations infinitely often. Thus, a common technique in TD algorithms is to instead use an ϵ -greedy policy which uses a parameter $\epsilon \in [0, 1]$,⁹

$$\pi(a | s) = \begin{cases} 1 - \epsilon + \frac{\epsilon}{|A|}, & a \in \arg \max_{a' \in A} Q(s, a') \\ \frac{\epsilon}{|A|}, & \text{else} \end{cases} . \quad (2.54)$$

Thus, the ϵ -greedy policy chooses the greedy action with probability $1 - \epsilon$, and with probability ϵ chooses a random other action. In this way, if $\epsilon > 0$, the requirement to try all state-action combinations an infinite number of times is

9. Equation 2.54 assumes that there is a single greedy action with maximum value under Q . If multiple actions with maximum value exist for a given state, then the definition can be modified to distribute the probability mass $(1 - \epsilon)$ among these actions.

Algorithm 3 Q-Learning for MDPs

- 1: Initialise: $Q(s, a) = 0$ for all $s \in S, a \in A$
 - 2: Repeat for every episode:
 - 3: **for** $t = 0, 1, 2, \dots$ **do**
 - 4: Observe current state s^t
 - 5: With probability ϵ : choose random action $a^t \in A$
 - 6: Else: choose action $a^t \in \arg \max_a Q(s^t, a)$
 - 7: Apply action a^t , observe reward r^t and next state s^{t+1}
 - 8: $Q(s^t, a^t) \leftarrow Q(s^t, a^t) + \alpha [r^t + \gamma \max_{a'} Q(s^{t+1}, a') - Q(s^t, a^t)]$
-

ensured. Now, in order to learn the optimal policy π^* , we can gradually reduce the value of ϵ to 0 during learning, such that π will gradually converge to π^* . Pseudocode for Sarsa using ϵ -greedy policies is given in Algorithm 2.

While Sarsa is based on the Bellman equation for Q^π , one can also construct a TD update target based on the Bellman optimality equation for Q^* ,

$$Q^*(s, a) = \sum_{s' \in S} \mathcal{T}(s' | s, a) \left[\mathcal{R}(s, a, s') + \gamma \max_{a' \in A} Q^*(s', a') \right]. \quad (2.55)$$

Q-learning (Watkins and Dayan 1992) is a TD algorithm which constructs an update target based on the above Bellman optimality equation, again by replacing the summation over states s' and the reward function $\mathcal{R}(s, a, s')$ with corresponding elements from the experience tuple (s^t, a^t, r^t, s^{t+1}) :

$$\mathcal{X} = r^t + \gamma \max_{a' \in A} Q(s^{t+1}, a') \quad (2.56)$$

The complete Q-learning update rule is specified as

$$Q(s^t, a^t) \leftarrow Q(s^t, a^t) + \alpha \left[r^t + \gamma \max_{a' \in A} Q(s^{t+1}, a') - Q(s^t, a^t) \right]. \quad (2.57)$$

Pseudocode for Q-learning using ϵ -greedy policies is given in Algorithm 3.

Q-learning is guaranteed to converge to the optimal policy π^* under the same conditions as Sarsa (i.e. trying all state-action pairs infinitely often, and standard stochastic approximation conditions in Equation 2.53). However, in contrast to Sarsa, Q-learning does not require that the policy π used to interact with the environment be gradually made closer to the optimal policy π^* . Instead, Q-learning may use *any* policy to interact with the environment, so long as the convergence conditions are upheld. For this reason, Q-learning is called an *off-policy* TD algorithm while Sarsa is an *on-policy* TD algorithm, and this distinction has a number of implications which we will discuss in more detail in

Chapter 8. In Section 2.7, we will use learning curves to evaluate and compare the performance of Q-learning and Sarsa on an RL problem.

2.7 Evaluation with Learning Curves

The standard approach to evaluate the performance of an RL algorithm on a learning problem is via *learning curves*. A learning curve shows the performance of the learned policy over increasing training time, where performance can be measured in terms of the learning objective (e.g. discounted returns) as well as other secondary metrics.

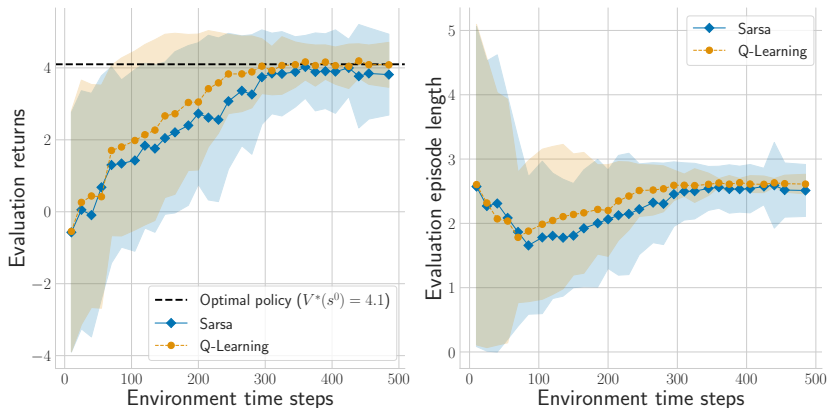
Figure 2.4 shows various learning curves for the Sarsa and Q-learning algorithms applied to the Mars Rover problem from Section 2.2 with discount factor $\gamma = 0.95$. In these figures, the x-axis shows the environment time steps across episodes, and the y-axis shows the average discounted *evaluation* returns achieved from the initial state, i.e. $V^\pi(s^0)$.¹⁰

The term “evaluation return” means that the shown returns are for the greedy policy¹¹ with respect to the learned action values after T learning time steps. Thus, the plots answer the question: if we finish learning after T time steps and extract the greedy policy, what expected returns can we expect to achieve with this policy? The results shown here are averaged over 100 independent training runs, each using a different random seed. Specifically, each point on a line is produced by taking the greedy policy from each training run at that time, running 100 independent episodes with the policy, averaging over the 100 resulting returns, and finally averaging over the 100 average returns from the 100 training runs. The shaded area shows the standard deviation over the averaged returns from each training run. In Figure 2.4b, we show the average episode lengths of the greedy policy instead of the average returns.

In this example, Sarsa and Q-learning both learn the optimal policy π^* which chooses action *left* in state *Start* and action *right* in states *Site A* and *Site B*. Moreover, the two algorithms basically have the same learning curves for the evaluation returns. Figures 2.4c and 2.4d (for Q-learning) show that the choice of the learning rate α and exploration rate ϵ can have an important impact on the learning. In this case, the “average” learning rate $\alpha_k(s, a) = \frac{1}{k}$ performs best for Q-learning, but for more complex MDPs with larger state and action sets

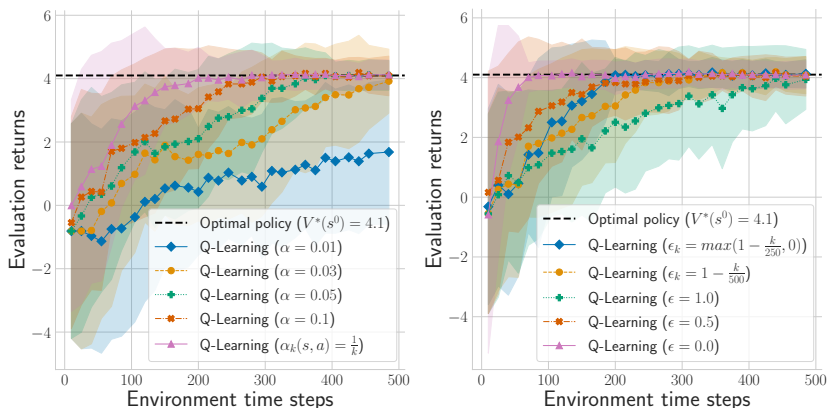
10. Recall that we use the term “return” as a shorthand for discounted return when the learning objective is to maximise discounted returns.

11. In policy gradient RL algorithms, discussed in Chapter 8, the algorithm learns a probabilistic policy for which there usually is no “greedy” version. In this case, the evaluation return just uses the current policy without modification.



(a) Average evaluation returns for Sarsa and Q-learning

(b) Average episode length for Sarsa and Q-learning



(c) Q-learning with different learning rates α (ϵ linearly decayed from 1 to 0)

(d) Q-learning with different exploration rates ϵ ($\alpha = 0.1$)

Figure 2.4: Results when applying the Sarsa and Q-learning algorithms to the Mars Rover problem (Figure 2.3, page 23) with $\gamma = 0.95$, averaged over 100 independent training runs. The shaded area shows the standard deviation over the averaged discounted returns from each training run. The dashed horizontal line marks the optimal value of the initial state ($V^*(s^0)$), computed via value iteration. In Figures 2.4a and 2.4b, the algorithms use a constant learning rate $\alpha = 0.1$ and an exploration rate which is linearly decayed from $\epsilon = 1$ to $\epsilon = 0$ over time steps $t = 0, \dots, 500$ (i.e. $\epsilon^t = 1 - t \frac{1}{500}$).

this choice of learning rate usually leads to much slower learning compared to appropriately chosen constant learning rates.

Note that the x-axis of the plots show the cumulative training time steps *across* episodes. This is not to be confused with the time steps t *within* episodes. The reason that we show cumulative time steps instead of the number of completed episodes on the x-axis is that the latter can potentially skew the comparison of algorithms. For instance, suppose we show training curves for algorithms A and B with number of episodes on the x-axis. Suppose both algorithms eventually reach the same performance with their learned policies, but algorithm A learns much faster than algorithm B (i.e. the curve for A rises more quickly). In this case, it could be that in the early episodes, algorithm A explored many more actions (i.e. time steps) than algorithm B before finishing the episode, which results in a larger number of collected experiences (or training data) for algorithm A. Thus, although both algorithms completed the same number of episodes, algorithm A will have done more learning updates (assuming updates are done after each time step, as in the TD methods presented in this chapter), which would explain the seemingly faster growth in the learning curve. If we instead showed the learning curve for both algorithms with cumulative time steps across episodes, then the learning curve for A might not grow faster than the curve for B.

Recall that at the beginning of this chapter (Figure 2.1, page 20), we defined an RL problem as the combination of a decision process model (e.g. MDP) and a learning objective for the policy (e.g. maximising the expected discounted return in each state for a specific discount factor). When evaluating a learned policy, we want to evaluate its ability to achieve this learning objective. For example, our Mars Rover problem specifies a discounted return objective with the discount factor $\gamma = 0.95$. Therefore, our learning curves in Figure 2.4a show exactly this objective on the y-axis.

However, in some cases, it can be useful to show *undiscounted returns* (i.e. $\gamma = 1$) for a learned policy, even if the RL problem specified a discounted return objective with a specific discount factor $\gamma < 1$. The reason is that undiscounted returns can sometimes be easier to interpret than discounted returns. For example, suppose we want to learn an optimal policy for a video game in which the agent controls a space ship and receives a +1 score (reward) for each destroyed enemy. If in an episode the agent destroys 10 enemies at various points in time, the undiscounted return will be 10 but the discounted return will be less than 10, making it more difficult to understand how many enemies the agent destroyed. More generally, when analysing a learned policy, it is often useful to show additional metrics besides the returns, such as win rates in competitive games, or episode lengths as we did in Figure 2.4b.

When showing undiscounted returns and additional metrics such as episode lengths, it is important to keep in mind that the evaluated policy was not actually trained to maximise these objectives – it was trained to maximise the expected discounted return for a specific discount factor. In general, two RL problems which differ only in their discount factor but are otherwise identical (i.e. same MDP) may have different optimal policies. In our Mars Rover example, a discounted return objective with discount factor $\gamma = 0.95$ leads to an optimal policy which chooses action *left* in the initial state, achieving a value of $V^*(Start) = 4.1$. In contrast, a discount factor of $\gamma = 0.5$ leads to an optimal policy which chooses action *right* in the initial state, achieving a value of $V^*(Start) = 0$. Both policies are *optimal*, but they are for two different learning problems that use different discount factors. Both learning problems are valid, and the choice of γ is part of designing the learning problem.

2.8 Equivalence of $\mathcal{R}(s, a, s')$ and $\mathcal{R}(s, a)$

Our definition of MDPs uses the most general definition of reward functions, $\mathcal{R}(s^t, a^t, s^{t+1})$, by conditioning on the current state s^t and action a^t as well as the successor state s^{t+1} . Another definition for reward functions commonly used in the RL literature is to condition only on s^t and a^t , i.e. $\mathcal{R}(s^t, a^t)$. This may be a point of confusion for newcomers to RL. However, these two definitions are in fact equivalent, in that for any MDP using $\mathcal{R}(s^t, a^t, s^{t+1})$ we can construct an MDP using $\mathcal{R}(s^t, a^t)$ (all other components are identical to the original MDP) such that a given policy π produces the same expected returns in both MDPs.

Recall the Bellman equation for the state-value function V^π that we have been using so far for an MDP with reward function $\mathcal{R}(s, a, s')$:

$$V^\pi(s) = \sum_{a \in A} \pi(a | s) \sum_{s' \in S} \mathcal{T}(s' | s, a) [\mathcal{R}(s, a, s') + \gamma V^\pi(s')] \quad (2.58)$$

Assuming that $\mathcal{R}(s, a, s')$ depends only on s, a , we can rewrite the above equation as

$$V^\pi(s) = \sum_{a \in A} \pi(a | s) \sum_{s' \in S} \mathcal{T}(s' | s, a) [\mathcal{R}(s, a) + \gamma V^\pi(s')] \quad (2.59)$$

$$= \sum_{a \in A} \pi(a | s) \left[\sum_{s' \in S} \mathcal{T}(s' | s, a) \mathcal{R}(s, a) + \sum_{s' \in S} \mathcal{T}(s' | s, a) \gamma V^\pi(s') \right] \quad (2.60)$$

$$= \sum_{a \in A} \pi(a | s) [\mathcal{R}(s, a) + \gamma \sum_{s' \in S} \mathcal{T}(s' | s, a) V^\pi(s')]. \quad (2.61)$$

This is the simplified Bellman equation for MDPs using a reward function $\mathcal{R}(s, a)$. Going the reverse direction from $\mathcal{R}(s, a)$ to $\mathcal{R}(s, a, s')$, we know that by

defining $\mathcal{R}(s, a)$ to be the expected reward under the state transition probabilities

$$\mathcal{R}(s, a) = \sum_{s' \in S} \mathcal{T}(s' | s, a) \mathcal{R}(s, a, s') \quad (2.62)$$

and plugging this into Equation 2.61, we recover the original Bellman equation that uses $\mathcal{R}(s, a, s')$, given in Equation 2.58. Analogous transformations can be done for the Bellman equation of the action value function Q^π , as well as the Bellman optimality equations for V^*/Q^* . Therefore, one may use either $\mathcal{R}(s, a, s')$ or $\mathcal{R}(s, a)$.

In this book, we use $\mathcal{R}(s, a, s')$ for the following pedagogical reasons:

1. Using $\mathcal{R}(s, a, s')$ can be more convenient when specifying examples of MDPs, since it is useful to show the differences in reward for multiple possible outcomes of an action. In our Mars Rover MDP shown in Figure 2.3, when using $\mathcal{R}(s, a)$ we would have to show the expected reward (Equation 2.62) on the transition arrow for action *right* from the initial state *Start* (rather than showing the different rewards for the two possible outcomes), which would be less intuitive to read.
2. The Bellman equations when using $\mathcal{R}(s, a, s')$ show a helpful visual congruence with the update targets used in TD algorithms. For example, the Q-learning target, $r^t + \gamma \max_{a'} Q(s^{t+1}, a')$, appears in a corresponding form in the Bellman optimality equation (within the [] brackets):

$$Q^*(s, a) = \sum_{s' \in S} \mathcal{T}(s' | s, a) \left[\mathcal{R}(s, a, s') + \gamma \max_{a' \in A} Q^*(s', a') \right] \quad (2.63)$$

And similarly for the Sarsa target and the Bellman equation for Q^π (see Section 2.6).

Finally, we mention that most of the original MARL literature presented in Chapter 6 in fact defined reward functions as $\mathcal{R}(s, a)$. To stay consistent in our notation, we will continue to use $\mathcal{R}(s, a, s')$ in the remainder of this book, noting that equivalent transformations from $\mathcal{R}(s, a)$ to $\mathcal{R}(s, a, s')$ always exist.

3 Games: Models of Multi-Agent Interaction

Chapter 1 introduced the general idea of agents which interact in a shared environment to achieve specified goals. In this chapter, we will define this idea formally via models of multi-agent interaction. These models are rooted in game theory, and are hence called *games*. The game models we will introduce define a hierarchy of increasingly complex models, shown in Figure 3.1.

The most basic model is the normal-form game, in which there are multiple agents but there is no evolving environment state. Further up the hierarchy are stochastic games, which define environment states that change over time as a result of the agents' actions and probabilistic state transitions. At the top of the game hierarchy are partially observable stochastic games, in which agents do not directly observe the full environment but instead observe incomplete and/or noisy information about the environment. Games may also use different assumptions about what the agents know about the game. We will introduce each of the game models in turn and provide examples.¹²

Note that this chapter focuses on defining models of multi-agent interaction, but does not define what it means to *solve* the games – that is, what it means for agents to act optimally. There are many possible solution concepts which define optimal policies for agents in games. We will introduce a range of solution concepts for games in Chapter 4.

12. There exists another type of game model, not covered in this chapter, called “extensive-form game”. The primary distinction is that in extensive-form games, agents choose actions in turns (one after another) while in the games we introduce, agents choose actions simultaneously (at the same time). We focus on simultaneous-move games since most MARL research uses this type of game model. Transformations between extensive-form games and (sequential) simultaneous-move games are possible (Shoham and Leyton-Brown 2008).

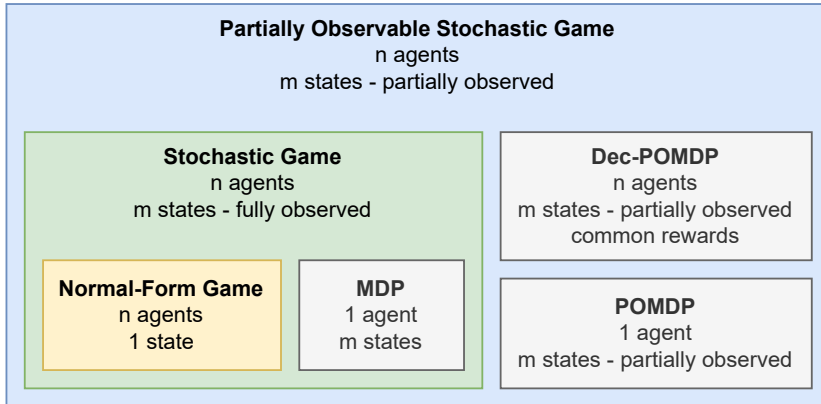


Figure 3.1: Hierarchy of game models. Partially observable stochastic games (POSGs) include stochastic games as a special case in which the states and agents’ chosen actions are fully observable by all agents. Stochastic games include normal-form games as a special case in which there is only a single environment state, and they include Markov decision processes (MDPs) as a special case in which there is only a single agent. POSGs also include partially observable Markov decision processes (POMDPs) as a special case in which there is only a single agent, and decentralised POMDPs (Dec-POMDPs) as a special case of POSGs in which the agents receive common rewards.

3.1 Normal-Form Games

A *normal-form game* (also known as “strategic-form” game) defines a single interaction between two or more agents. Similar to how the multi-armed bandit problem can be seen as the basic kernel of the MDP (Section 2.2), the normal-form game can be seen as the basic building block of all game models introduced in this chapter.

Definition 2 (Normal-form game) *A normal-form game consists of:*

- *Finite set of agents $I = \{1, \dots, n\}$*
- *For each agent $i \in I$:*
 - *Finite set of actions A_i*
 - *Reward function $\mathcal{R}_i : A \mapsto \mathbb{R}$, where $A = A_1 \times \dots \times A_n$*

A normal-form game proceeds as follows: First, each agent i selects a policy $\pi_i : A_i \mapsto [0, 1]$, which assigns probabilities to the actions available to the agent. Each agent then samples an action $a_i \in A_i$ with probability $\pi_i(a_i)$ given by its

policy. The resulting actions of all agents form a *joint action*, $a = (a_1, \dots, a_n)$. Finally, each agent i receives a reward based on its reward function and the joint action, $r_i = \mathcal{R}_i(a)$.

Normal-form games can be classified based on the relationship between the reward functions of agents:

- In a *zero-sum* game, the sum of the agents' rewards is always 0, i.e. $\sum_{i \in I} \mathcal{R}_i(a) = 0$ for all $a \in A$.¹³ In zero-sum games with two agents, i and j ,¹⁴ one agent's reward function is simply the negative of the other agent's reward function, i.e. $\mathcal{R}_i = -\mathcal{R}_j$.
- In a *common-reward* game, all agents receive the same reward, i.e. $\mathcal{R}_i = \mathcal{R}_j$ for all $i, j \in I$.
- In a *general-sum* game, there are no restrictions on the relationship of reward functions.

Normal-form games with two agents are also referred to as *matrix games* because, in this case, the reward function can be compactly represented as a matrix.¹⁵ Figure 3.2 shows three example matrix games. Agent 1's action is to choose the row position, and agent 2's action is to choose the column position. The values (r_1, r_2) in the matrix cells show the agents' rewards for playing a particular joint action. Figure 3.2a shows the Rock-Paper-Scissors game in which each agent chooses one of three possible actions (R,P,S). This is a zero-sum game (i.e. $r_1 = -r_2$) since for each action combination, one agent wins (+1 reward) and the other agent loses (-1 reward), or there is a draw (0 reward to both). Figure 3.2b shows a “coordination” game with two actions (A,B) for each agent, and agents must select the same action to receive a positive reward. This game is common-reward (i.e. $r_1 = r_2$), hence it suffices to show a single reward in the matrix cells which both of the agents receive. Lastly, Figure 3.2c shows a widely-studied game called the Prisoner’s Dilemma, which is a general-sum game. Here, each agent chooses to either cooperate (C) or defect (D). While mutual cooperation would give both agents the second-highest reward, each agent is individually incentivised to defect since this is the “dominant action”, meaning that D always achieves higher rewards compared to C. What makes

13. Zero-sum games are a special case of *constant-sum* games, in which the agents' rewards sum to a constant.

14. For games with two agents, we often use i and j to refer to the two agents in formal descriptions, and we use 1 and 2 to refer to the agents in specific examples.

15. Some game theory literature uses the term “bimatrix game” to specifically refer to general-sum games that require two matrices to define the agents' reward functions, and “matrix game” for games that require only a single matrix to define the reward functions. This book uses “matrix game” in both cases.

	R	P	S		A	B		C	D
R	0,0	-1,1	1,-1		10	0		-1,-1	-5,0
P	1,-1	0,0	-1,1		0	10		0,-5	-3,-3
S	-1,1	1,-1	0,0						
(a) Rock-Paper-Scissors	(b) Coordination Game	(c) Prisoner's Dilemma							

Figure 3.2: Examples of normal-form games for two agents (i.e. matrix games).

these and many other games interesting is that one agent’s reward depends on the choices of the other agents, which are not known in advance.

Section 11.2 provides a comprehensive listing of all structurally distinct and strictly ordinal 2×2 normal-form games (meaning games with two agents and two actions each), of which there are 78 in total. In the remainder of the book, we will use “normal-form game” when making general statements about n -agent normal-form games, and we will sometimes use “matrix game” when discussing specific examples of normal-form games for two agents.

3.2 Repeated Normal-Form Games

A normal-form game, as presented in Section 3.1, defines a single interaction between multiple agents. The most basic way to extend this to *sequential* multi-agent interactions is by repeating the same normal-form game for a finite or infinite number of times, giving rise to the *repeated normal-form game*. This type of game model is among the most widely-studied models in game theory. For example, the repeated Prisoner’s Dilemma has been extensively studied in the game theory literature and still serves as an important example of a sequential social dilemma.

Given a normal-form game defined by $(I, \{A_i\}_{i \in I}, \{\mathcal{R}_i\}_{i \in I})$, a repeated normal-form game plays the same normal-form game repeatedly over time steps $t = 0, 1, 2, \dots, T$, where T is either finite or goes to infinity. At each time step t , each agent i samples an action a_i^t with probability $\pi_i(a_i^t | h^t)$ given by its policy. The policy is now conditioned on the *joint-action history* $h^t = (a^0, \dots, a^{t-1})$, which contains all joint actions before the current time step t (for $t=0$, the history is the empty set). Given the joint action $a^t = (a_1^t, \dots, a_n^t)$, each agent i receives a reward $r_i^t = \mathcal{R}_i(a^t)$.

A repeated normal-form game may terminate after a finite number of time steps T which is known a-priori to all agents, or it may continue for an infinite number of time steps. It is important to note that a game with finite repetitions

is not in general equivalent to the same game with infinite repetitions. In finite games, there can be “end-game” effects: if the agents know that a game will finish after T time steps, they may choose different actions closer to the end of the game compared to earlier in the game (see Section 6.3.3 for an example in Prisoner’s Dilemma). For infinite games, one may specify a probability with which each time step terminates the game. This termination probability is related to the discount factor $\gamma \in [0, 1]$ used in the discounted-return learning objective in RL, in that $1 - \gamma$ specifies the probability of termination in each time step (see the discussion in Section 2.3). For $\gamma < 1$, the game still counts as “infinite” since any finite number $T > 0$ of time steps will have a non-zero probability of occurring.

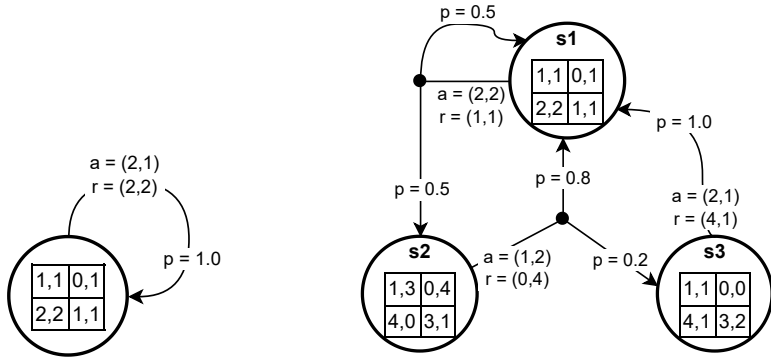
Besides the time index t , the important addition in repeated normal-form games is that policies can now make decisions based on the entire history of past joint actions. This gives rise to a very complex space of possible policies that agents can use. Typically, policies are conditioned on an *internal state* which is a function of the history, $f(h^t)$. For example, a policy may be conditioned on just the most recent joint action a^{t-1} , or on summary statistics such as the action counts of other agents in the history. “Tit-for-Tat” is a famous policy for the game of repeated Prisoner’s Dilemma which simply conditions on the most recent action of the other agent, choosing to cooperate if the other agent cooperated and defecting if the other agent instead defected (Axelrod and Hamilton 1981).

3.3 Stochastic Games

While the relative simplicity of normal-form games is useful to study interactions between agents, they lack the notion of an environment state which is affected by the actions of the agents. Moving closer to the full multi-agent system described in Section 1.1, *stochastic games* define a state-based environment in which the state evolves over time based on the agents’ actions and probabilistic state transitions (Shapley 1953).

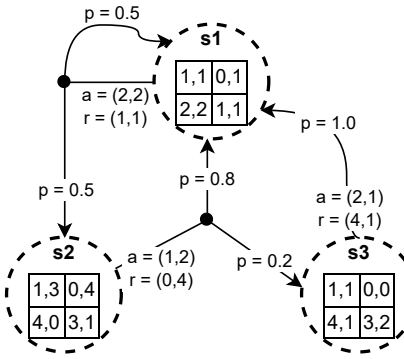
Definition 3 (Stochastic game) A stochastic game consists of:

- Finite set of agents $I = \{1, \dots, n\}$
- Finite set of states S
- For each agent $i \in I$:
 - Finite set of actions A_i
 - Reward function $\mathcal{R}_i : S \times A \times S \mapsto \mathbb{R}$, where $A = A_1 \times \dots \times A_n$
- State transition probability function $\mathcal{T} : S \times A \times S \mapsto [0, 1]$



(a) Repeated normal-form game

(b) Stochastic game



(c) POSG

Figure 3.3: Normal-form games are the basic building block of all game models described in Chapter 3. This figure shows one example game for each type of game model, shown as a directed cyclic graph. Each game is for two agents with two actions each. Each node in the graph corresponds to a state and shows a normal-form game being played in that state. We show one possible joint action choice for each state, along with the rewards received and the probabilities of reaching the respective next state in the outgoing arrows. For POSG, the states are dashed to represent that agents do not directly observe the current state of the game; instead, agents receive partial/noisy observations about the state.

- *Initial state distribution* $\Pr^0 : S \mapsto [0, 1]$

A stochastic game proceeds as follows: The game starts in an initial state $s^0 \in S$ sampled from \Pr^0 . At time t , each agent i observes the current state s^t and chooses an action a_i^t with probability given by its policy, $\pi_i(a_i^t | h^t)$, resulting in the joint action $a^t = (a_1^t, \dots, a_n^t)$. The policy is conditioned on the *state-action history*, $h^t = (s^0, a^0, s^1, a^1, \dots, s^t)$, which is observed by all agents – this property is also known as *full observability*. Given the state s^t and joint action a^t , the game transitions into a next state s^{t+1} with probability given by $\mathcal{T}(s^t, a^t, s^{t+1})$, and each agent i receives a reward $r_i^t = \mathcal{R}_i(s^t, a^t, s^{t+1})$. We will also write this probability as $\mathcal{T}(s^{t+1} | s^t, a^t)$ to emphasise that it is conditioned on the state-action pair s^t, a^t . These steps are repeated for a finite or infinite number of time steps, or until some terminal state is reached.

Similar to MDPs, stochastic games have the Markov property in that the probability of the next state and reward is conditionally independent of the past states and joint actions, given the current state and joint action

$$\Pr(s^{t+1}, r^t | s^t, a^t, s^{t-1}, a^{t-1}, \dots, s^0, a^0) = \Pr(s^{t+1}, r^t | s^t, a^t) \quad (3.1)$$

where $r^t = (r_1^t, \dots, r_n^t)$ is the joint reward at time t . For this reason, stochastic games are also sometimes called *Markov games* (e.g. Littman 1994).¹⁶

As a concrete example of a stochastic game, we can model the level-based foraging environment shown in Figure 1.2 (page 4). Each state is a vector which specifies the x-y integer positions of all agents and items, as well as binary flags for each item to indicate whether it has been collected. The agents' action spaces include actions for moving up/down/left/right, collecting an item, and doing nothing (noop). The effect of joint actions is specified in the transition probability function \mathcal{T} . For example, two agents that together collect an existing item will modify the state by switching the binary flag associated with that item (meaning that the item has been collected and no longer exists). In a common-reward version of the game, every agent will receive a reward of +1 whenever any of the items has been collected by any agents. A general-sum version may specify individual rewards for agents, such as +1 reward for agents which were actually involved in the collection of an item and 0 reward for all other agents. The game terminates after a fixed number of time steps, or when a terminal state has been reached in which all items have been collected.

Stochastic games include repeated normal-form games as a special case in which there is only a single state in S . More generally, if we define reward

16. For a comment on the naming, see <https://agents.inf.ed.ac.uk/blog/multiagent-rl-inaccuracies>.

functions as $\mathcal{R}_i(s, a)$,¹⁷ then each state $s \in S$ in the stochastic game can be viewed as a non-repeated normal-form game with rewards given by $\mathcal{R}_i(s, \cdot)$, as shown in Figure 3.3. It is in this sense that normal-form games form the basic building block of stochastic games. Stochastic games also include MDPs as the special case in which there is only a single agent in I . And, like MDPs, while the state and action sets in our definition of stochastic games are finite (as per the original definition of Shapley (1953)), a stochastic game can be analogously defined for continuous states and actions. It is also possible to define different action sets for different states, but this adds more notation and does not make the model more general.

Finally, the classification of normal-form games into zero-sum games, common-reward games, and general-sum games also carry over to stochastic games (Section 3.1). That is, stochastic games may specify zero-sum rewards, common rewards, or general-sum rewards.

Chapter 11 gives many more examples of state-based multi-agent environments, many of which can be modelled as a stochastic game. See Section 11.3 for details.

3.4 Partially Observable Stochastic Games

Sitting at the top of the game model hierarchy shown in Figure 3.1, the most general model we use in this book is the *partially observable stochastic game*, or POSG for short. While in stochastic games the agents can directly observe the environment state and the chosen actions of all agents, in a POSG the agents receive “observations” which carry some incomplete information about the environment state and agents’ actions. This allows POSGs to represent decision processes in which agents have limited ability to sense their environment, such as in autonomous driving and other robot control tasks, or strategic games where players have private information not seen by other players (e.g. card games).

In its most general form, a POSG defines state-observation probabilities $\Pr(s^t, o^t | s^{t-1}, a^{t-1})$, where $o^t = (o_1^t, \dots, o_n^t)$ is the joint observation at time t which contains the agents’ individual observations o_i^t . However, it is often the case that observations only depend on the new environment state s^t and the joint action a^{t-1} that led to this state (and not on the previous state s^{t-1}). Thus, it is common to define for each agent i an individual *observation function* \mathcal{O}_i which specifies probabilities over the agent’s possible observations o_i^t given the state s^t and joint action a^{t-1} . We give the full definition of POSGs below:

17. For a comment on the equivalence of $\mathcal{R}_i(s, a, s')$ and $\mathcal{R}_i(s, a)$, see Section 2.8.

Definition 4 (Partially observable stochastic game) A partially observable stochastic game (POSG) is defined by the same elements of a stochastic game (Definition 3) and additionally defines for each agent $i \in I$:

- Finite set of observations O_i
- Observation function $\mathcal{O}_i : A \times S \times O_i \mapsto [0, 1]$

A POSG proceeds similarly to a stochastic game: The game starts in an initial state $s^0 \in S$ sampled from \Pr^0 . At each time t , each agent i receives an observation $o_i^t \in O_i$ with probability given by its observation function, $\mathcal{O}_i(a^{t-1}, s^t, o_i^t)$, where we assume that the joint action a^{t-1} for $t=0$ is the empty set. We will also write this as $\mathcal{O}_i(o_i^t | a^{t-1}, s^t)$ to emphasise that the probability is conditioned on the state s^t and joint action a^{t-1} . Each agent then chooses an action a_i^t based on action probabilities given by its policy, $\pi_i(a_i^t | h_i^t)$, resulting in the joint action $a^t = (a_1^t, \dots, a_n^t)$. The policy π_i is conditioned on agent i 's *observation history* $h_i^t = (o_i^0, \dots, o_i^t)$, which includes all of the agent's past observations up to and including the most recent observation. Note that an agent's observation may or may not include the actions of other agents, as we will further discuss below. Given the joint action a^t , the game transitions into the next state s^{t+1} with probability $\mathcal{T}(s^{t+1} | s^t, a^t)$, and each agent i receives a reward $r_i^t = \mathcal{R}_i(s^t, a^t, s^{t+1})$. These steps are repeated for a finite or infinite number of time steps, or until some terminal state is reached.

Again, the classification of zero-sum reward, common reward, and general-sum reward also applies to the POSG model. POSGs with common rewards, in which all agents have the same reward function, are also known as “Decentralised POMDP” (Dec-POMDP) and have been widely studied in the area of multi-agent planning (see, for example, the textbook by Oliehoek and Amato (2016)). POSGs can also be defined analogously with continuous-valued (or mixed discrete-continuous) observations.

POSGs include stochastic games as the special case in which $o_i^t = (s^t, a^{t-1})$. They also include POMDPs as the special case in which there is only a single agent in I . In general, the observation functions in POSGs can be used to represent diverse observability conditions of interest. Examples include:

Unobserved actions of other agents Agents may observe the state and their own previous action, but not the previous actions of other agents, i.e. $o_i^t = (s^t, a_i^{t-1})$. An example of this scenario is robot soccer, in which the robots may observe the full playing field but do not directly communicate their chosen actions to other players (especially players from the opponent team). In this case, agents may need to infer with some uncertainty the possible actions of other agents based on changes in the observed environment

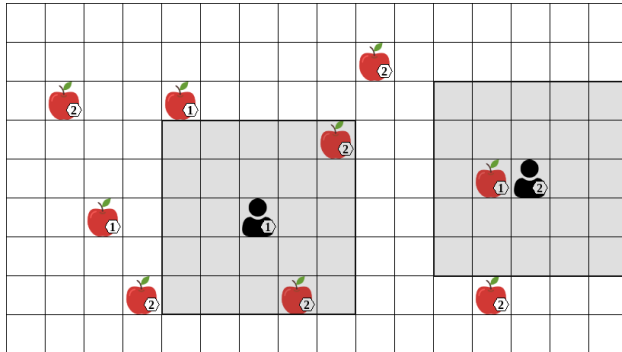


Figure 3.4: Level-based foraging environment with partial observability. Agents observe the world through their local vision fields (shown as grey rectangles around the agents).

state (e.g. inferring a pass action between two players based on location, direction, and velocity of ball). Another example is a market setting in which the asset prices are observed by all agents but the agents' buy/sell actions are private.

Limited view region Agents may observe a subset of the state and joint action, i.e. $o_i^t = (\bar{s}^t, \bar{a}^t)$ where $\bar{s}^t \subset s^t$ and $\bar{a}^t \subset a^t$. Such a scenario may arise if agents have a limited view of their surrounding environment, such that an agent may only see those parts of the state s^t and joint action a^{t-1} which are (or took place) within its view region. For example, in the partially observable version of level-based foraging shown in Figure 3.4, the agents have local vision fields and can only see items, other agents, and their actions within their own vision fields. Another example is agents playing a real-time strategy game in which parts of the game map are concealed by “fog of war” (e.g. Vinyals et al. 2019).

Observation functions can model uncertainty in observations (i.e. *noise*) by assigning non-zero probability to multiple possible observations. For example, such uncertainty may be caused by imperfect sensors in robotics applications. Observation functions can also model communication between agents which may be limited by range and can be unreliable. A communication message can be modelled as a multi-valued vector (e.g. a bit vector) which does not modify the environment state but can be received by other agents within their own observation, if they are within a certain range of the sender. Unreliable communication can be modelled by specifying a probability with which messages are lost (i.e. not received by an agent even if within sending range), or

by randomly changing parts of the message. One may even specify combined actions $a_i = (a_{i,s}, a_{i,c})$ where $a_{i,s}$ affects the state s (such as moving around in a physical world) while $a_{i,c}$ is a communication message which may be received by other agents j within their observation o_j .

In Chapter 11 (Section 11.3), we list several multi-agent environments with partial observability used in MARL research.

3.4.1 Belief States and Filtering

In a POSG, since an agent’s observation only provides partial information about the current state of the environment, it is typically not possible to choose optimal actions based only on the current observation. For example, in the level-based foraging environment shown in Figure 3.4, the optimal action for the level-1 agent may be to move towards the level-1 item to its left. However, this item is not included in the agent’s current observation within its vision field, and so the agent cannot infer the optimal action from its current observation alone. In general, if the environment is only partially observed, the agents must maintain estimates of the possible current states and their relative likelihoods based on the history of observations. Continuing with the example, the level-1 agent may have seen the level-1 item to its left in a previous observation, and may thus remember its location within the observation history.

One way of defining such estimates about environment states from the perspective of agent i is as a *belief state* b_i^t , which is a probability distribution over the possible states $s \in S$ that the environment may be in at time t . Consider for simplicity a POSG with only a single agent i – that is, a POMDP. The initial belief state of the agent is given by the distribution of initial states, i.e. $b_i^0 = \Pr^0$. After taking action a_i^t and observing o_i^{t+1} , the belief state b_i^t is updated to b_i^{t+1} by computing a Bayesian posterior distribution

$$b_i^{t+1}(s') = \eta \sum_{s \in S} b_i^t(s) \mathcal{T}(s' | s, a_i^t) \mathcal{O}_i(o_i^{t+1} | a_i^t, s') \quad (3.2)$$

where η is a normalisation constant. The resulting belief state is exact in the sense that it retains all relevant information from the observation history. This belief state is known as a “sufficient statistic”, because it carries enough information needed to choose optimal actions and to make predictions about the future. The process of updating belief states based on observations is also known as (*belief state*) *filtering*.

Unfortunately, the space complexity of storing such exact belief states and the time complexity of updating them using the Bayesian update from Equation 3.2 are each exponential in the number of variables that define the state, making it intractable for complex environments. Hence, developing algorithms for

efficient *approximate* filtering has been the subject of much research; see, for example, Albrecht and Ramamoorthy (2016) and discussions therein.

In a POSG with more than one agent, the definition of belief states and how to update them becomes significantly more complex. In particular, since agents may not observe the chosen actions of other agents and their resulting observations, the agents now also have to infer probabilities over the possible observations and actions of other agents, which in turn requires knowledge of their observation functions and policies (e.g. Gmytrasiewicz and Doshi 2005; Oliehoek and Amato 2016). However, as we will discuss in Section 3.5, in MARL we typically assume that agents do not possess complete knowledge about the elements of the POSG, such as S , \mathcal{T} , and \mathcal{O}_i (including their own observation function), all of which are required in Equation 3.2. Thus, we will not dwell further on the intricacies of defining exact belief states in multi-agent contexts.

To achieve some form of filtering without knowledge of these elements, deep RL algorithms often use recurrent neural networks (discussed in Section 7.5.2) to process observations sequentially. The output vector of the recurrent network learns to encode information about the current state of the environment, and can be used to condition other functions of the agent such as value and policy networks. Part II of this book will discuss how MARL algorithms can integrate such deep learning models in order to learn and operate in POSGs. See also Section 8.3 for a further discussion on this topic.

3.5 Knowledge Assumptions in Games

What do the agents know about the environment they interact in and how it works? In other words, what do agents know about the game they are playing?

In game theory, the standard assumption is that all agents have knowledge of all components that define the game; this is referred to as a “complete knowledge game” (Owen 2013). For normal-form games, this means that the agents know the action spaces and reward functions of all agents (including their own). For stochastic games and POSGs, agents also know the state space and state transition function, as well as the observation functions of all agents. Knowledge of these components can be utilised in different ways. For example, if agent i knows the reward function \mathcal{R}_j of another agent j , then agent i may be able to estimate the best-response action (Section 4.2) of agent j , which in turn could inform the optimal action for agent i . If an agent knows the transition function \mathcal{T} , it may predict the effects of its actions on the state and plan its actions several steps into the future.

However, in most real-world applications of interest (such as those mentioned in Chapter 1), it is infeasible to obtain accurate and complete specifications of these components. For such applications, often the best we can obtain is a simulator of these components which can generate samples of states, (joint) rewards, and (joint) observations. For example, we may have access to a simulator $\widehat{\mathcal{T}}$ which, given an input state s and joint action a , can produce samples of joint rewards and successor states $(r = (r_1, \dots, r_n), s') \sim \widehat{\mathcal{T}}(s, a)$, such that

$$\Pr\left\{\widehat{\mathcal{T}}(s, a) = (r, s')\right\} \approx \mathcal{T}(s' | s, a) \prod_{i \in I} [\mathcal{R}_i(s, a, s') = r_i]_1 \quad (3.3)$$

where $[x]_1 = 1$ if x is true, else 0.

In MARL, we usually operate at the other end of the knowledge spectrum: agents typically do not know the reward functions of other agents, nor even their own reward function; and agents have no knowledge of the state transition and observation functions (in game theory also known as “incomplete knowledge game” (Harsanyi 1967)). Instead, an agent i only experiences the immediate effects of its own actions, via its own reward r_i^t and (in stochastic games) the joint action a^t and resulting next state s^{t+1} or (in POSG) an observation o_i^{t+1} . Agents may use such experiences to construct models of the unknown components in a game (such as \mathcal{T}) or the policies of other agents (Section 6.3.2).

Additional assumptions may pertain to whether certain knowledge about the game is held mutually by all agents or not (symmetric vs asymmetric knowledge), and whether agents are aware of what other agents know about the game. Are the reward functions in a game known to all agents or only to some of the agents? If all agents have knowledge of the reward functions of each agent, are they also aware of the fact that all agents have this knowledge, and the fact that all agents knowing the reward functions is known by all agents, and so on (common knowledge)? While such questions and their impact on optimal decision making have been the subject of much research in game theory (Shoham and Leyton-Brown 2008) and agent modelling (Albrecht and Stone 2018), in MARL they have played a relatively lesser role since the usual assumption is that agents have no knowledge of most game components. Exceptions can be found in MARL research which focused on either zero-sum or common-reward games, where this fact is exploited in some structural way in the algorithm design. Chapters 6 and 9 will describe several MARL algorithms of this latter category.

Lastly, it is usually assumed that the number of agents in a game is fixed and that this fact is known by all agents. While outside the scope of this book, it is interesting to note that recent research in MARL has begun to tackle *open*

multi-agent environments in which agents may dynamically enter and leave the environment (Jiang et al. 2020; Rahman et al. 2021).

3.6 Dictionary: Reinforcement Learning \leftrightarrow Game Theory

This book lies at the intersection of reinforcement learning and game theory. These distinct fields share a number of concepts but use different terminology, and this book primarily uses the terminology which is common in reinforcement learning. We conclude this chapter by providing a small “dictionary” in Figure 3.5 which shows some synonymous terms between the two fields.

RL (this book)	GT	Description
environment	game	Model specifying the possible actions, observations, and rewards of agents, and the dynamics of how the state evolves over time and in response to actions.
agent	player	An entity which makes decisions. “Player” can also refer to a specific role in the game which is assumed by an agent, e.g.: <ul style="list-style-type: none"> • “row player” in matrix game • “white player” in chess
policy	strategy	Function used by an agent/player to assign probabilities to actions.
reward	payoff	Scalar value received by an agent/player after taking an action.
deterministic X	pure X	X assigns probability 1 to one option, e.g.: <ul style="list-style-type: none"> • deterministic policy • pure strategy • deterministic/pure Nash equilibrium
probabilistic X	mixed X	X assigns probabilities < 1 to options, e.g.: <ul style="list-style-type: none"> • probabilistic policy • mixed strategy • probabilistic/mixed Nash equilibrium
joint X	X profile	X is a tuple, typically with one element for each agent/player, e.g.: <ul style="list-style-type: none"> • joint reward • deterministic joint policy • payoff profile • pure strategy profile

Figure 3.5: Synonymous terms in reinforcement learning (RL) and game theory (GT). This book uses RL terminology.

4 Solution Concepts for Games

What does it mean for agents to interact optimally in a multi-agent system? In other words, what is a *solution* to a game? This is a central question of game theory, and many different *solution concepts* have been proposed which specify when a collection of agent policies constitutes a stable or desirable outcome. While Chapter 3 introduced the basic game models to formalise multi-agent environments and interaction, this chapter will introduce a series of solution concepts. Together, a game model and solution concept define a learning problem in MARL (Figure 4.1).

For common-reward games, in which all agents receive the same reward, a straightforward definition for a solution is to maximise the expected return received by all agents (but finding such a solution may not be simple at all, as we will see in later chapters). However, if the agents have differing rewards, the definition of a solution concept becomes more complicated.

In general, a solution to a game is a *joint policy* which consists of one policy for each agent and satisfies certain properties. These properties are expressed in terms of the expected returns yielded to each agent under the joint policy, and the relations between the agents' returns. Thus, this chapter will begin by giving a universal definition of expected returns which applies to all of the game models introduced in Chapter 3. Using this definition, we will introduce a hierarchy of increasingly general equilibrium solution concepts which include the minimax equilibrium, Nash equilibrium, and correlated equilibrium. We will also discuss solution refinements such as Pareto-optimality and social welfare and fairness, and alternative solution concepts such as no-regret. We finish the chapter with a discussion of the computational complexity of computing Nash equilibria.

Note that our definitions of solution concepts and their stated existence properties assume *finite* game models, as defined in Chapter 3. In particular, we assume finite state, action, and observation spaces, and a finite number of agents. Games with infinite elements, such as continuous actions and observations, can

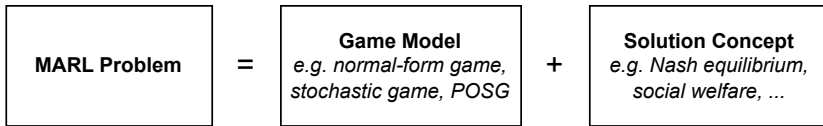


Figure 4.1: A MARL problem is defined by the combination of a game model which defines the mechanics of the multi-agent system and interactions, and a solution concept which specifies the desired properties of the joint policy to be learned. (See also Figure 2.1, page 20.)

use analogous definitions (e.g. by using densities and integrals) but may have different existence properties for solutions.¹⁸

Game theory has produced a wealth of knowledge on foundational questions about solution concepts, including: For a given type of game model and solution concept, is a solution guaranteed to exist in the game? Is the solution unique (i.e. only one solution exists) or are there potentially many solutions, even infinitely many? And is a given learning and decision rule, when used by all agents, guaranteed to converge to a solution? For a more in-depth treatment of such questions, we recommend the books of Fudenberg and Levine (1998), Young (2004), Shoham and Leyton-Brown (2008), and Owen (2013).

4.1 Joint Policy and Expected Return

A solution to a game is a *joint policy*, $\pi = (\pi_1, \dots, \pi_n)$, which satisfies certain requirements defined by the solution concept in terms of the expected return, $U_i(\pi)$, yielded to each agent i under the joint policy. We seek a universal definition of the expected return $U_i(\pi)$ which applies to all of the game models introduced in Chapter 3, so that our definitions of solution concepts also apply to all game models. Thus, we will define the expected return in the context of the POSG model (Section 3.4), which is the most general game model used in this book and includes both stochastic games and normal-form games.

We start by introducing some additional notation. In a POSG, let $\hat{h}^t = \{(s^\tau, o^\tau, a^\tau)_{\tau=0}^{t-1}, s^t, o^t\}$ denote a *full history* up to time t , consisting of the states, joint observations, and joint actions of all agents in each time step before t , and the state s^t and joint observation o^t at time t . The function $\sigma(\hat{h}^t) = (o^0, \dots, o^t)$

18. For example, every two-agent zero-sum normal-form game with finite action spaces has a unique minimax game value, but zero-sum games with continuous actions exist that do not have a minimax game value (Sion and Wolfe 1957).

returns the history of joint observations from the full history \hat{h}^t , and we sometimes abbreviate $\sigma(\hat{h}^t)$ to h^t if it is clear from context. We define the probability of a joint observation, $\mathcal{O}(o^t | a^{t-1}, s^t)$, as the product $\prod_{i \in I} \mathcal{O}_i(o_i^t | a^{t-1}, s^t)$.

To make our definitions valid for both finite and infinite time steps, we use discounted returns¹⁹ and assume the standard convention of absorbing states (Section 2.3) to represent terminal states. That is, once an absorbing (i.e. terminal) state has been reached, the game will subsequently always transition into that same state with probability 1 and return a reward of 0 to all agents; and the observation functions and policies of all agents will become deterministic (i.e. assign probability 1 to a certain observation and action). Thus, our definitions encompass infinite episodes as well as episodes which terminate after a finite number of time steps, including non-repeated normal-form games which always terminate after a single time step.²⁰

In the following, we provide two equivalent definitions of expected returns. The first definition is based on enumerating all full histories in the game, while the second definition is based on a Bellman-style recursion of value computations. These definitions are equivalent, but can provide different perspectives and have been used in different ways. In particular, the first definition resembles a linear sum and may be easier to interpret, while the second definition uses a recursion which can be operationalised such as in value iteration (introduced in Section 6.1).

History-based expected return: Given a joint policy π , we can define the expected return for agent i under π by enumerating all possible full histories and summing the returns for i in each history, weighted by the probability of generating the history under the POSG and joint policy. Formally, define the set \hat{H} to contain all full histories \hat{h}^t for $t \rightarrow \infty$.²¹ Then, the expected return for agent i under joint policy π is given by

$$U_i(\pi) = \mathbb{E}_{\hat{h}^t \sim (\mathbf{P}^0, \mathcal{T}, \mathcal{O}, \pi)} [u_i(\hat{h}^t)] \quad (4.1)$$

$$= \sum_{\hat{h}^t \in \hat{H}} \Pr(\hat{h}^t | \pi) u_i(\hat{h}^t) \quad (4.2)$$

19. Alternative definitions based on average rewards instead of discounted returns are also possible for games (Shoham and Leyton-Brown 2008).

20. In the context of non-repeated normal-form games, the expected return for an agent simply becomes the expected reward of that agent, so we use the term “expected return” in all game models including non-repeated normal-form games.

21. Recall from Chapter 3 that we assume finite game models, and that a^{t-1} before time $t=0$ is the empty set.

where $\Pr(\hat{h}^t \mid \pi)$ is the probability of full history \hat{h}^t under π ,

$$\Pr(\hat{h}^t \mid \pi) = \Pr^0(s^0)\mathcal{O}(o^0 \mid \emptyset, s^0) \prod_{\tau=0}^{t-1} \pi(a^\tau \mid h^\tau) \mathcal{T}(s^{\tau+1} \mid s^\tau, a^\tau) \mathcal{O}(o^{\tau+1} \mid a^\tau, s^{\tau+1}) \quad (4.3)$$

and $u_i(\hat{h}^t)$ is the discounted return for agent i in \hat{h}^t ,

$$u_i(\hat{h}^t) = \sum_{\tau=0}^{t-1} \gamma^\tau \mathcal{R}_i(s^\tau, a^\tau, s^{\tau+1}) \quad (4.4)$$

with discount factor $\gamma \in [0, 1]$.

We use $\pi(a^\tau \mid h^\tau)$ to denote the probability of joint action a^τ under joint policy π after joint observation history h^τ . If we assume that agents act independently, then we can define

$$\pi(a^\tau \mid h^\tau) = \prod_{j \in I} \pi_j(a_j^\tau \mid h_j^\tau). \quad (4.5)$$

If agents do not act independently, such as in correlated equilibrium (Section 4.6) and methods such as central learning (Section 5.3.1), then $\pi(a^\tau \mid h^\tau)$ can be defined accordingly.

Recursive expected return: Analogous to the Bellman recursion used in MDP theory (Section 2.4), we can define the expected return to agent i under joint policy π via two interlocked functions V_i^π and Q_i^π , defined below. In the following, we use $s(\hat{h})$ to denote the last state in \hat{h} (i.e. $s(\hat{h}^t) = s^t$), and we use $\langle \rangle$ to denote the concatenation operation.

$$V_i^\pi(\hat{h}) = \sum_{a \in A} \pi(a \mid \sigma(\hat{h})) Q_i^\pi(\hat{h}, a) \quad (4.6)$$

$$Q_i^\pi(\hat{h}, a) = \sum_{s' \in S} \mathcal{T}(s' \mid s(\hat{h}), a) \left[\mathcal{R}_i(s(\hat{h}), a, s') + \gamma \sum_{o' \in O} \mathcal{O}(o' \mid a, s') V_i^\pi(\langle \hat{h}, a, s', o' \rangle) \right] \quad (4.7)$$

We can understand $V_i^\pi(\hat{h})$ as the expected return, also called *value*, for agent i when following π after the full history \hat{h} . Similarly, $Q_i^\pi(\hat{h}, a)$ is the expected return for i when executing joint action a after \hat{h} and then following π subsequently. (The value iteration procedure for stochastic games (Section 6.1) uses simplified versions of V_i^π , Q_i^π which are conditioned on states rather than histories.) Given Equations 4.6 and 4.7, we can define the expected return as

$$U_i(\pi) = \mathbb{E}_{s^0 \sim \Pr^0, o^0 \sim \mathcal{O}(\cdot \mid \emptyset, s^0)} [V_i^\pi(\langle s^0, o^0 \rangle)]. \quad (4.8)$$

4.2 Best Response

Many existing solution concepts, including most of the solution concepts introduced in this chapter, can be expressed compactly based on *best responses*. Given a set of policies for all agents other than agent i , denoted by $\pi_{-i} = (\pi_1, \dots, \pi_{i-1}, \pi_{i+1}, \dots, \pi_n)$, a best response for agent i to π_{-i} is a policy π_i that maximises the expected return for i when played against π_{-i} . Formally, the set of best-response policies for agent i is defined as

$$\text{BR}_i(\pi_{-i}) = \arg \max_{\pi_i} U_i(\langle \pi_i, \pi_{-i} \rangle) \quad (4.9)$$

where $\langle \pi_i, \pi_{-i} \rangle$ denotes the complete joint policy consisting of π_i and π_{-i} . For convenience and to keep our notation lean, we will sometimes drop the $\langle \rangle$ (e.g. $U_i(\pi_i, \pi_{-i})$).

Note that the best response to a given π_{-i} may not be unique, meaning that $\text{BR}_i(\pi_{-i})$ may contain more than one best-response policy. For example, for a given π_{-i} there may be multiple actions for agent i which achieve equal maximum expected return against π_{-i} , in which case any probability assignment to these actions is also a best response.

Besides being useful for compact definitions of solution concepts, best response operators have also been used in game theory and MARL to iteratively compute solutions. Two example methods we will see include fictitious play (Section 6.3.1) and joint action learning with agent modelling (Section 6.3.2), among others.

4.3 Minimax

Minimax is a solution concept defined for two-agent zero-sum games, in which one agent's reward is the negative of the other agent's reward (Section 3.1). A classical example of such games is the matrix game Rock-Paper-Scissors, whose reward matrix is shown in Figure 3.2a (page 44). More complex examples with sequential moves include games such as chess and Go. The existence of minimax solutions was first proven in the foundational game theory work of von Neumann and Morgenstern (1944).

Definition 5 (Minimax solution) In a zero-sum game with two agents, a joint policy $\pi = (\pi_i, \pi_j)$ is a minimax solution if

$$U_i(\pi) = \max_{\pi'_i} \min_{\pi'_j} U_i(\pi'_i, \pi'_j) \quad (4.10)$$

$$= \min_{\pi'_j} \max_{\pi'_i} U_i(\pi'_i, \pi'_j) \quad (4.11)$$

$$= -U_j(\pi).$$

Every two-agent zero-sum normal-form game has a minimax solution (von Neumann and Morgenstern 1944). Minimax solutions also exist in every two-agent zero-sum stochastic game with finite episode length, and two-agent zero-sum stochastic games with infinite episode length using discounted returns (Shapley 1953). Moreover, while more than one minimax solution π may exist in a game, all minimax solutions yield the same unique value $U_i(\pi)$ for agent i (and, thus, agent j). This value is also referred to as the (*minimax*) *value of the game*.²²

In a minimax solution, each agent uses a policy which is optimised against a worst-case opponent that seeks to minimise the agent's return. Formally, there are two parts in the definition of minimax. Equation 4.10 is the minimum expected return that agent i can guarantee against *any* opponent. Here, π_i is agent i 's *maxmin* policy and $U_i(\pi)$ is i 's maxmin value. Conversely, Equation 4.11 is the minimum expected return that agent j can force on agent i . Here, we refer to j 's policy π_j as the *minmax* policy against agent i , and $U_i(\pi)$ is agent i 's minmax value. In a minimax solution, agent i 's maxmin value is equal to its minmax value. Another way to interpret this is that the order of the min/max operators does not matter: agent i first announcing its policy followed by agent j selecting its policy, is equivalent to agent j first announcing its policy followed by agent i selecting its policy. Neither agent gains from such policy announcements.

A minimax solution can be understood more intuitively as each agent using a best-response policy against the other agent's policy. That is, (π_i, π_j) is a minimax solution if $\pi_i \in \text{BR}_i(\pi_j)$ and $\pi_j \in \text{BR}_j(\pi_i)$. In the non-repeated Rock-Paper-Scissors game, there exists a unique minimax solution which is for both agents to choose actions uniformly randomly (i.e. assign equal probability to all actions). This solution gives an expected return of 0 to both agents. In fact,

22. Recall that we assume finite game models. For zero-sum games with continuous actions, examples exist that do not have a minimax game value (Sion and Wolfe 1957).

it can be verified²³ in this game that if any agent i uses a uniform policy π_i , then *any* policy π_j for the other agent j is a best-response policy to π_i , and all best-response policies $\pi_j \in \text{BR}_j(\pi_i)$ yield an expected reward of 0 to agent j (and to agent i). This example shows that best responses need not be unique, and there can be many (even infinitely many) possible best-response policies. However, the joint policy (π_i, π_j) in which both policies are uniform policies is the *only* joint policy in which *both* policies are best responses to each other, making it a minimax solution. In Section 4.4, we will see that this best-response relation can also be applied to the more general class of general-sum games.

4.3.1 Minimax Solution via Linear Programming

For non-repeated zero-sum normal-form games, we can obtain a minimax solution by solving two linear programmes, one for each agent. Each linear programme computes a policy for one agent by minimising the expected return of the other agent. Thus, agent i minimises the expected return of agent j , and vice versa. We provide the linear programme to compute the policy π_i for agent i ; agent j 's policy is obtained by constructing a similar linear programme in which the indices i and j are swapped. The linear programme contains variables x_{a_i} for each action $a_i \in A_i$ to represent the probability of selecting action a_i (thus, $\pi_i(a_i) = x_{a_i}$ defines the policy of agent i), as well as a variable U_j^* to represent the expected return of agent j , which is minimised as follows:

$$\text{minimise} \quad U_j^* \quad (4.12)$$

$$\text{subject to} \quad \sum_{a_i \in A_i} \mathcal{R}_j(a_i, a_j) x_{a_i} \leq U_j^* \quad \forall a_j \in A_j \quad (4.13)$$

$$x_{a_i} \geq 0 \quad \forall a_i \in A_i \quad (4.14)$$

$$\sum_{a_i \in A_i} x_{a_i} = 1 \quad (4.15)$$

In this linear programme, the constraint in Equation 4.13 means that no single action of agent j will achieve an expected return greater than U_j^* against the policy specified by $\pi_i(a_i) = x_{a_i}$. This implies that no probability distribution over agent j 's actions will achieve a higher expected return than U_j^* . Finally, the constraints in Equations 4.14 and 4.15 ensure that the values of x_{a_i} form a valid

23. Assume agent 1's policy assigns probability $\frac{1}{3}$ to each action R, P, S . The expected reward to agent 2 for any policy π_2 is: $\pi_2(R)(\frac{1}{3}\mathcal{R}_2(R, R) + \frac{1}{3}\mathcal{R}_2(P, R) + \frac{1}{3}\mathcal{R}_2(S, R)) + \pi_2(P)(\frac{1}{3}\mathcal{R}_2(R, P) + \frac{1}{3}\mathcal{R}_2(P, P) + \frac{1}{3}\mathcal{R}_2(S, P)) + \pi_2(S)(\frac{1}{3}\mathcal{R}_2(R, S) + \frac{1}{3}\mathcal{R}_2(P, S) + \frac{1}{3}\mathcal{R}_2(S, S)) = \pi_2(R)(-\frac{1}{3} + \frac{1}{3}) + \pi_2(P)(+\frac{1}{3} - \frac{1}{3}) + \pi_2(S)(-\frac{1}{3} + \frac{1}{3}) = 0$. The same can be shown when switching the roles of agents 1 and 2.

probability distribution. Note that in a minimax solution, we will have $U_i^* = U_j^*$ (see Definition 5).

Linear programmes can be solved using well-known algorithms, such as the simplex algorithm which runs in exponential time in the worst case but often is very efficient in practice; or interior-point algorithms which are provably polynomial-time.

4.4 Nash Equilibrium

The Nash equilibrium applies the idea of a mutual best response to games with general rewards and more than two agents. That such a solution exists in any general-sum non-repeated normal-form game was first proven in the celebrated work of Nash (1950).

Definition 6 (Nash equilibrium) *In a general-sum game with n agents, a joint policy $\pi = (\pi_1, \dots, \pi_n)$ is a Nash equilibrium if*

$$\forall i, \pi'_i : U_i(\pi'_i, \pi_{-i}) \leq U_i(\pi). \quad (4.16)$$

In a Nash equilibrium, no agent i can improve its expected return by changing its policy π_i specified in the equilibrium joint policy π , assuming the policies of the other agents stay fixed. This means that each agent's policy in the Nash equilibrium is a best response to the policies of the other agents, i.e. $\pi_i \in \text{BR}_i(\pi_{-i})$ for all $i \in I$. Thus, Nash equilibrium generalises minimax in that, in two-agent zero-sum games, the set of minimax solutions coincides with the set of Nash equilibria (Owen 2013). Nash (1950) first proved that every finite normal-form game has at least one Nash equilibrium.

Recall the matrix games shown in Figure 3.2 (page 44). In the non-repeated Prisoner's Dilemma matrix game, the only Nash equilibrium is for both agents to choose D. It can be seen that no agent can unilaterally deviate from this choice to increase its expected return. In the non-repeated coordination game, there exist three different Nash equilibria: 1) both agents choose action A, 2) both agents choose action B, and 3) both agents assign probability 0.5 to each action. Again it can be checked that no agent can increase its expected return by deviating from their policies in each equilibrium. Lastly, in the non-repeated Rock-Paper-Scissors game, the only Nash equilibrium is for both agents to choose actions uniform-randomly, which is the minimax solution of the game.

The above examples illustrate two important aspects of Nash equilibrium as a solution concept. First, a Nash equilibrium can be deterministic in that each policy π_i in the equilibrium π is deterministic (i.e. $\pi_i(a_i) = 1$ for some $a_i \in A_i$), such

as in Prisoner’s Dilemma. However, in general, a Nash equilibrium may be probabilistic in that the policies in the equilibrium use randomisation (i.e. $\pi_i(a_i) < 1$ for some $a_i \in A_i$), such as in the coordination game and Rock-Paper-Scissors. In the game theory literature, deterministic and probabilistic equilibria are also called “pure equilibria” and “mixed equilibria”, respectively (see also Figure 3.5 for terminology). As we will see in Chapter 6, this distinction is important in MARL because some algorithms are unable to represent probabilistic policies, and hence cannot learn probabilistic equilibria.

Second, a game may have multiple Nash equilibria, and each equilibrium may entail different expected returns to the agents. In the coordination game, the two deterministic equilibria give an expected return of 10 to each agent; while the probabilistic equilibrium gives an expected return of 5 to each agent. The Chicken game (Section 4.6) and Stag Hunt game (Section 5.4.3) are other examples of games with multiple Nash equilibria that each give different expected returns to the agents. This leads to the important question of which equilibrium the agents should converge to during learning and how this may be achieved. We will discuss this *equilibrium selection* problem in more depth in Section 5.4.3.

The existence of Nash equilibria has also been shown for stochastic games (Fink 1964; Filar and Vrieze 2012). In fact, for games with (infinite) sequential moves, there are various “folk theorems”²⁴ which essentially state that any set of feasible and enforceable expected returns $\hat{U} = (\hat{U}_1, \dots, \hat{U}_n)$ can be achieved by an equilibrium solution if agents are sufficiently far-sighted (i.e. the discount factor γ is close to 1). The assumptions and details in different folk theorems vary and can be quite involved, and here we only provide a rudimentary description for intuition (for more specific definitions, see e.g. Fudenberg and Levine 1998; Shoham and Leyton-Brown 2008). Broadly speaking, \hat{U} is *feasible* if it can be realised within the game by some joint policy π ; i.e. there exists a π such that $U_i(\pi) = \hat{U}_i$ for all $i \in I$. And \hat{U} is *enforceable* if each \hat{U}_i is at least as large as agent i ’s minmax value²⁵

$$v_i = \min_{\pi_{-i}^{\text{mm}}} \max_{\pi_i^{\text{mm}}} U_i(\pi_i^{\text{mm}}, \pi_{-i}^{\text{mm}}). \quad (4.17)$$

(Think: other agents $-i$ minimise the maximum achievable return for agent i .) Under these two conditions, we can construct an equilibrium solution which uses π to achieve \hat{U} , and if at any time t any agent i deviates from its policy π_i in π , the other agents will limit i ’s return to v_i by using their corresponding

24. The name “folk theorem” apparently came about because the general concept was known before it was formalised.

25. We first encountered minmax in Section 4.3 where it was defined for two agents, while in Equation 4.17 we define it for n agents (the *mm* superscript stands for minmax).

minmax policy π_{-i}^{mm} from Equation 4.17 indefinitely after t . Thus, since $v_i \leq \hat{U}_i$, i has no incentive to deviate from π_i , making π an equilibrium.

Given a joint policy π , how can one check whether it is a Nash equilibrium? Equation 4.16 suggests the following procedure, which reduces the multi-agent problem to n single-agent problems. For each agent i , keep the other policies π_{-i} fixed and compute an optimal best-response policy π'_i for i . If the optimal policy π'_i achieves a higher expected return than agent i 's policy π_i in the joint policy π , i.e. $U_i(\pi'_i, \pi_{-i}) > U_i(\pi_i, \pi_{-i})$, then we know that π is not a Nash equilibrium. For non-repeated normal-form games, π'_i can be computed efficiently using a linear programme (e.g. Albrecht and Ramamoorthy 2012). For games with sequential moves, π'_i may be computed using a suitable single-agent RL algorithm.

4.5 ϵ -Nash Equilibrium

The strict requirement of Nash equilibrium – that no agent can gain *anything* by unilaterally deviating from the equilibrium – can lead to practical issues when used in a computational system. It is known that for games with more than two agents, the action probabilities specified by the policies in the equilibrium may be irrational numbers (i.e. cannot be represented as a fraction of two integers). Nash himself pointed this out in his original work (Nash 1950). However, computer systems cannot fully represent irrational numbers using finite-precision floating-point approximations. Moreover, in many applications, reaching a strict equilibrium may be too computationally costly. Instead, it may be good enough to compute a solution which is sufficiently close to a strict equilibrium, meaning that agents could technically deviate to improve their returns but any such gains are sufficiently small.

The ϵ -Nash equilibrium relaxes the strict Nash equilibrium, by requiring that no agent can improve its expected returns by more than some small amount $\epsilon > 0$ when deviating from its policy in the equilibrium. Formally:

Definition 7 (ϵ -Nash equilibrium) *In a general-sum game with n agents, a joint policy $\pi = (\pi_1, \dots, \pi_n)$ is an ϵ -Nash equilibrium for $\epsilon > 0$ if*

$$\forall i, \pi'_i : U_i(\pi'_i, \pi_{-i}) - \epsilon \leq U_i(\pi). \quad (4.18)$$

Since the action probabilities specified by π'_i are continuous and we consider expected returns U_i , we know that every Nash equilibrium is surrounded by a region of ϵ -Nash equilibria for any $\epsilon > 0$. The exact Nash equilibrium corresponds to $\epsilon = 0$. However, although it may be tempting to view ϵ -Nash equilibrium as an approximation of Nash equilibrium, it is important to note that an ϵ -Nash equilibrium may not be close to any real Nash equilibrium, in

	C	D
A	100,100	0,0
B	1,2	1,1

Figure 4.2: Matrix game to illustrate that an ϵ -Nash equilibrium (B,D) ($\epsilon = 1$) may not be close to a real Nash equilibrium (A,C).

terms of the expected returns produced by the equilibrium. In fact, the expected returns under an ϵ -Nash equilibrium may be arbitrarily far away from those of any Nash equilibrium, even if the Nash equilibrium is unique.

Consider the example game shown in Figure 4.2. This game has a unique Nash equilibrium at (A,C).²⁶ It also has an ϵ -Nash equilibrium at (B,D) for $\epsilon = 1$, in which agent 2 could deviate to action C to increase its reward by ϵ . First, note that neither agent's return under the ϵ -Nash equilibrium is within ϵ of its return under the Nash equilibrium. Second, we can arbitrarily increase the rewards for (A,C) in the game without affecting the ϵ -Nash equilibrium (B,D). Thus, in this example, the ϵ -Nash equilibrium does not meaningfully approximate the unique Nash equilibrium of the game.

To check that a joint policy π constitutes an ϵ -Nash equilibrium for some given ϵ , we can use essentially the same procedure for checking Nash equilibrium described at the end of Section 4.4, except that we check for $U_i(\pi'_i, \pi_{-i}) - \epsilon > U_i(\pi_i, \pi_{-i})$ to determine that π is not a ϵ -Nash equilibrium.

4.6 Correlated Equilibrium

A restriction of Nash equilibrium is that the agent policies must be probabilistically independent (as per Equation 4.5), which can limit the expected returns that can be achieved by the agents. Correlated equilibrium (Aumann 1974) generalises Nash equilibrium by allowing for correlation between policies. In the general definition of correlated equilibrium, each agent i 's policy is additionally conditioned on the outcomes of a private random variable d_i for the agent, which are governed by a joint probability distribution over (d_1, \dots, d_n) that is commonly known by all agents. Here, we will present a common version of correlated equilibrium for non-repeated normal-form games, in which d_i corresponds to an action recommendation for agent i given by a joint policy π_c . At the end of this section, we will mention possible extensions to sequential-move games.

26. Saying that a joint action a is an X-equilibrium (e.g. Nash) is a shorthand for saying that a deterministic joint policy π that assigns probability 1 to this joint action is an X-equilibrium.

Definition 8 (Correlated equilibrium) In a general-sum normal-form game with n agents, let $\pi_c(a)$ be a joint policy which assigns probabilities to joint actions $a \in A$. Let $\phi_i : A_i \mapsto A_i$ be an action modifier for agent i . Then π_c is a correlated equilibrium if

$$\forall i, \phi_i : \sum_{a \in A} \pi_c(a) (\mathcal{R}_i(\langle \phi_i(a_i), a_{-i} \rangle) - \mathcal{R}_i(a)) \leq 0. \quad (4.19)$$

Equation 4.19 states that in a correlated equilibrium, in which agents know the probability distribution $\pi_c(a)$ and their own recommended action a_i (but not the recommended actions for other agents), no agent can unilaterally deviate from its recommended action in order to increase its expected return. An example of a correlated joint policy π_c can be seen in the central learning approach discussed in Section 5.3.1, which is a single policy trained directly over the joint action space $A_1 \times \dots \times A_n$ and used to dictate actions to each agent. It can be shown that the set of correlated equilibria contains the set of Nash equilibria (e.g. Osborne and Rubinstein 1994).

To see an example of how correlated equilibrium can achieve greater returns than Nash equilibrium, consider the Chicken matrix game shown in Figure 4.3. This game represents a situation in which two vehicles (agents) are on a collision course and can choose to either stay on course (S) or leave (L). In the non-repeated game, there are the following three Nash equilibria with associated expected returns for the two agents shown as pairs (U_i, U_j) :

- $\pi_i(S) = 1, \pi_j(S) = 0 \rightarrow (7, 2)$
- $\pi_i(S) = 0, \pi_j(S) = 1 \rightarrow (2, 7)$
- $\pi_i(S) = \frac{1}{3}, \pi_j(S) = \frac{1}{3} \rightarrow \approx (4.66, 4.66)$

Now, consider the following joint policy π_c which uses correlated actions:

- $\pi_c(L, L) = \pi_c(S, L) = \pi_c(L, S) = \frac{1}{3}$
- $\pi_c(S, S) = 0$

The expected return under π_c to both agents is: $7 * \frac{1}{3} + 2 * \frac{1}{3} + 6 * \frac{1}{3} = 5$. It can be verified that no agent has an incentive to unilaterally deviate from the action recommended to it by π_c , assuming knowledge of π_c (and without knowing the actions recommended to other agents). For example, say agent i received action recommendation L. Then, given π_c , i knows that agent j will choose S with probability 0.5 and L with probability 0.5. Thus, the expected return to i when choosing L is $2 * \frac{1}{2} + 6 * \frac{1}{2} = 4$, which is higher than the expected return for choosing S ($0 * \frac{1}{2} + 7 * \frac{1}{2} = 3.5$). Thus, i has no incentive to deviate from L.

The above example also illustrates how a correlated equilibrium can again be described as mutual best responses between the agents. In this case, however, we cannot directly use our specific definition from Equation 4.9 of best-response

	S	L
S	0,0	7,2
L	2,7	6,6

Figure 4.3: Chicken matrix game.

policies, since the joint policy π_c cannot in general be factored into individual policies π_1, \dots, π_n . Rather, in correlated equilibrium, we may consider each action recommendation a_i given by π_c to be a best response to the actions a_{-i} recommended to the other agents.

For sequential-move games, various definitions of correlated equilibrium exist (e.g. Forges 1986; Solan and Vieille 2002; Von Stengel and Forges 2008; Farina, Bianchi, and Sandholm 2020). These definitions vary in a number of design choices and can be relatively complex. For instance, the private signals d_i may specify actions a_i that are revealed at each decision point, or they may specify entire policies π_i revealed once at the start of the game. The joint distribution over d_1, \dots, d_n and the action/policy modifier ϕ_i may be conditioned on different types of information, such as the current game state, agent observation histories, or previous values of d_i (Solan and Vieille 2002). The sampled outcomes of d_i may or may not be revealed to agents, such as in “coarse” correlated equilibria which assume that the outcomes of d_i are only revealed to agents if they “commit” to the equilibrium (e.g. Farina, Bianchi, and Sandholm 2020). Definitions of correlated equilibrium can also vary in how agents are treated if they deviate from the equilibrium; for example, no further action recommendations may be issued to an agent after it deviates from the action recommended by the equilibrium (Von Stengel and Forges 2008).

4.6.1 Correlated Equilibrium via Linear Programming

For non-repeated normal-form games, we can obtain a correlated equilibrium by solving a linear programme. The linear programme computes a joint policy π such that no agent can improve its expected return by deviating from the joint actions sampled from π . Thus, the linear programme contains variables x_a for each joint action $a \in A$ to represent the probability of selecting joint action a under the joint policy π (i.e. $\pi(a) = x_a$). To select between different possible equilibria, we here use an objective which maximises the sum of the agents’ expected returns (i.e. social welfare, see Section 4.9), but other objectives could be used such as maximising expected returns for individual agents. This gives

rise to the following linear programme:

$$\text{maximise} \quad \sum_{a \in A} \sum_{i \in I} x_a \mathcal{R}_i(a) \quad (4.20)$$

$$\text{subject to} \quad \sum_{a \in A: a_i = a'_i} x_a \mathcal{R}_i(a) \geq \sum_{a \in A: a_i = \hat{a}'_i} x_a \mathcal{R}_i(\hat{a}_i, a_{-i}) \quad \forall i \in I, a'_i, \hat{a}_i \in A_i \quad (4.21)$$

$$x_a \geq 0 \quad \forall a \in A \quad (4.22)$$

$$\sum_{a \in A} x_a = 1 \quad (4.23)$$

The constraint in Equation 4.21 ensures the property that no agent can gain by deviating from the action a_i sampled under the joint policy $\pi(a) = x_a$ to a different action \hat{a}_i . The constraints in Equations 4.22 and 4.23 ensure that the values of x_a form a valid probability distribution. Note that this linear programme can contain many more variables and constraints than the linear programmes for minimax solutions (Section 4.3.1), since we now have variables and constraints for each joint action $a \in A$, which can grow exponentially in the number of agents (see also the discussion in Section 5.4.4).

4.7 Conceptual Limitations of Equilibrium Solutions

While equilibrium solutions, in particular the Nash equilibrium, have been adopted as the standard solution concept in MARL, they are not without certain shortcomings. Besides the practical issues noted in Section 4.5, we list some of the most important conceptual limitations in equilibrium solutions:

Sub-optimality In general, finding equilibrium solutions is not synonymous with maximising expected returns. The only thing we know about a given equilibrium solution is that each agent's policy is a best response to the policies of the other agents, but this does not mean that the agent's expected returns are the best they could be. A simple example of this can be seen in the Prisoner's Dilemma game (Figure 3.2c), in which the only Nash equilibrium is the joint action (D,D) giving each agent an expected return of -3 , while the joint action (C,C) gives a higher expected return of -1 to each agent but is not a Nash equilibrium (each agent can deviate to improve its return). Similarly, in the Chicken game (Section 4.6), the shown correlated equilibrium π_c achieves an expected return of 5 for each agent, while the joint action (L,L) achieves expected returns of 6 for each agent but is neither a Nash equilibrium nor a correlated equilibrium.

Non-uniqueness Equilibrium solutions may not be unique, which means that there may exist multiple, even infinitely many, equilibria. Each of these equilibria may entail different expected returns to different agents, as can be seen in the three Nash equilibria in the Chicken game. This leads to a difficult challenge: which of these different equilibria should the agents adopt, and how can they agree on a specific equilibrium? This challenge is known as the “equilibrium selection” problem and has been widely studied in game theory and economics. In MARL, equilibrium selection can pose a serious challenge for agents that learn concurrently from local observations, which we will discuss further in Section 5.4.3. One approach to tackle equilibrium selection is to use additional criteria such as Pareto optimality (Section 4.8) and social welfare and fairness (Section 4.9) to differentiate between different equilibria.

Incompleteness An equilibrium solution π is incomplete in that it does not specify equilibrium behaviours for *off-equilibrium paths*. An off-equilibrium path is any full history \hat{h} which has probability $\Pr(\hat{h} \mid \pi) = 0$ (Equation 4.3) under the equilibrium π . For example, this could occur due to some temporary disturbance in the agents’ executed policies leading to actions not normally prescribed by the policies. In such cases, π does not prescribe actions to bring the interaction back to an on-equilibrium path, i.e. a full history \hat{h} with $\Pr(\hat{h} \mid \pi) > 0$ under the equilibrium π . To address such incompleteness, game-theorists have developed refinement concepts such as subgame perfect equilibrium and trembling-hand perfect equilibrium (Selten 1988; Owen 2013).

4.8 Pareto Optimality

As we saw in the previous sections, an equilibrium solution in which agents use mutual best-response policies may be of limited value, because different equilibria may entail very different expected returns to different agents (e.g. Chicken game; see Section 4.7). Moreover, under certain conditions, any feasible and enforceable expected returns can be realised by an equilibrium solution, making the space of equilibrium solutions very large or infinite (folk theorems; see Section 4.4). Therefore, we may want to narrow down the space of equilibrium solutions by requiring additional criteria that a solution must achieve. One such criterion is *Pareto optimality*,²⁷ defined below:

27. Named after Italian economist Vilfredo Pareto (1848–1923).

Definition 9 (Pareto domination and optimality) A joint policy π is Pareto-dominated by another joint policy π' if

$$\forall i : U_i(\pi') \geq U_i(\pi) \text{ and } \exists i : U_i(\pi') > U_i(\pi). \quad (4.24)$$

A joint policy π is Pareto-optimal²⁸ if it is not Pareto-dominated by any other joint policy. We then also refer to the expected returns entailed by π as Pareto-optimal.

Intuitively, a joint policy is Pareto-optimal if there is no other joint policy which improves the expected return for at least one agent, without reducing the expected return for any other agent. In other words, no agent can be better off without making other agents worse off. Note that in zero-sum games, all joint policies are Pareto-optimal; thus, Pareto optimality is not of interest in zero-sum games. In common-reward games, all Pareto-optimal joint policies achieve the same expected return, which by definition is the maximum possible expected return that any joint policy can achieve in the game.

We illustrate Pareto optimality using the non-repeated Chicken matrix game from Figure 4.3 (page 69). Figure 4.4 shows the convex hull of feasible expected joint returns in this game. Each dot corresponds to the expected joint return achieved by some joint policy (π_1, π_2) . In this figure, we discretised the space of joint policies by stepping each policy from $\pi_i(S) = 0$ to $\pi_i(S) = 1$ using a step size of $\frac{1}{30}$, resulting in 30 different policies for each agent, and a total of $30 * 30 = 900$ joint policies. The corners of the convex hull correspond to the four possible deterministic joint policies (i.e. the four joint actions from the game matrix). The expected joint returns obtained by Pareto-optimal joint policies are marked by the red squares. The figure also shows the expected joint returns corresponding to the deterministic and probabilistic Nash equilibria of the game (see Section 4.6 for details).

For the infinitely-repeated game and under the average reward,²⁹ there exists a folk theorem which shows that any expected joint return in this hull that is equal to or larger than the minmax value of the agents (Equation 4.17) can be realised by an equilibrium. In the Chicken game, it can be seen from the game's reward matrix that the minmax value for both agents is 2, and so any expected joint return (U_1, U_2) with $U_1, U_2 \geq 2$ in the hull can be realised by means of an equilibrium. This is where Pareto optimality comes in: we can narrow down the space of desirable equilibria by requiring that the expected joint return achieved by an equilibrium also be Pareto-optimal. The Pareto-optimal joint returns are

28. Some of the game theory literature uses the terms “Pareto-efficient/inefficient” for Pareto-optimal/dominated, respectively.

29. Average returns (or average rewards) correspond to discounted returns with $\gamma \rightarrow 1$.

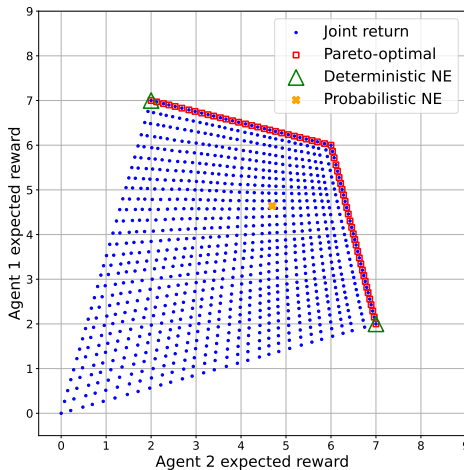


Figure 4.4: Feasible expected joint rewards and Pareto frontier in the Chicken matrix game. Each blue dot shows the expected joint reward obtained by a joint policy. Red squares show joint rewards of Pareto-optimal joint policies. Also shown are the joint rewards corresponding to the deterministic and probabilistic Nash equilibria of the game.

those which reside on the *Pareto frontier* shown by the red squares. Thus, for any given joint policy π , we can project its corresponding expected joint return into the convex hull and detect Pareto optimality if the joint return lies on the Pareto frontier.

We have presented Pareto optimality as a concept to refine equilibrium solutions, but note that a joint policy can be Pareto-optimal without being an equilibrium solution. However, Pareto optimality alone may not be a very useful solution concept. Many joint policies can be Pareto-optimal without being a desirable solution. For example, imagine a game in which the agents divide a finite resource among them. Any solution (division) between the agents would be Pareto-optimal, because any other solution would make some agents worse off. And if the solution is not an equilibrium, then the worse-off agents (relative to the other solutions) are immediately incentivised to deviate from the solution, making it an unstable solution.

4.9 Social Welfare and Fairness

Pareto optimality states that there is no other solution in which at least one agent is better off without making other agents worse off. However, it does not make any statements about the total amount of rewards and their distribution among the agents. For example, the Pareto frontier in Figure 4.4 contains solutions with expected joint returns ranging from $(7, 2)$ to $(6, 6)$ to $(2, 7)$. Thus, we may consider concepts of *social welfare and fairness* to further constrain the space of desirable solutions.

The study of social welfare and fairness has a long history in economics and game theory, and many criteria and social welfare functions have been proposed (Moulin 2004; Fleurbaey and Maniquet 2011; Sen 2018; Amanatidis et al. 2023). The term welfare usually refers to some notion of totality of the agents' returns, while the term fairness relates to the distribution of returns among agents. In this section, we consider two basic definitions of welfare and fairness:

Definition 10 (Welfare and Welfare Optimality) *The (social) welfare of a joint policy π is defined as*

$$W(\pi) = \sum_{i \in I} U_i(\pi). \quad (4.25)$$

A joint policy π is welfare-optimal if $\pi \in \arg \max_{\pi'} W(\pi')$.

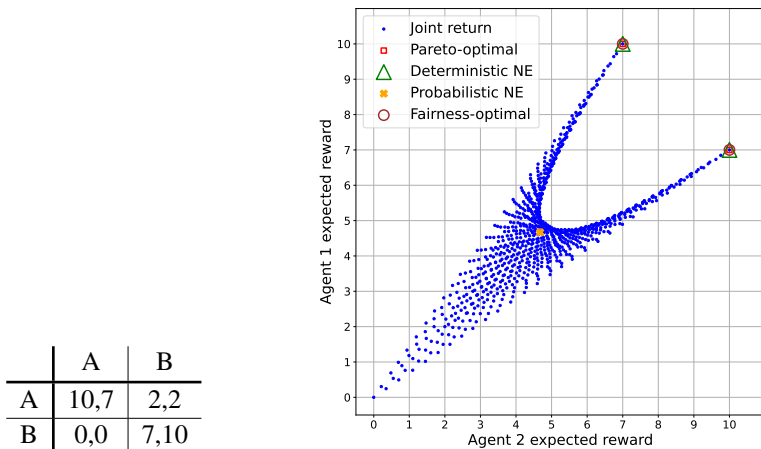
Definition 11 (Fairness and Fairness Optimality) *The (social) fairness of a joint policy π is defined as³⁰*

$$F(\pi) = \prod_{i \in I} U_i(\pi). \quad (4.26)$$

A joint policy π is fairness-optimal if $\pi \in \arg \max_{\pi'} F(\pi')$.

A welfare-optimal joint policy maximises the sum of the agents' expected returns, while a fairness-optimal joint policy maximises the product of the agents' expected returns. This definition of fairness promotes a type of equity between the agents, in the following sense: if we consider a set of joint policies $\pi \in \Pi$ that achieve equal welfare (sum of returns) $W(\pi)$, then the joint policy $\pi \in \Pi$ with the greatest fairness according to $F(\pi)$ is that which gives equal expected return to each agent, i.e. $U_i(\pi) = U_j(\pi)$ for all i, j . For example, three different joint policies in a two-agent game that yield expected joint returns

30. This type of fairness is also known as *Nash social welfare*, defined as the geometric mean $(\prod_{i \in I} U_i(\pi))^{\frac{1}{n}}$ (Caragiannis et al. 2019; Fan et al. 2023).



(a) Battle of the Sexes matrix game. (b) Joint rewards and fairness-optimal outcomes.

Figure 4.5: Feasible joint rewards and fairness-optimal outcomes in the Battle of the Sexes matrix game. This game models a situation where two people (classically, a man and a woman, hence the name of the game) want to meet in one of two places (A or B) but have different preferences about the best place. Agent 1 prefers place A and agent 2 prefers place B. Joint action (A,B) is preferred over (B,A), since in the latter case both agents end up at their respective least-preferred place. The two deterministic joint policies corresponding to joint actions (A,A) and (B,B) are the only joint policies (in the non-repeated game) that are both Pareto-optimal and fairness-optimal. The figure also shows the only probabilistic Nash equilibrium, which is neither Pareto-optimal nor fairness-optimal.

of (1, 5), (2, 4), (3, 3) achieve equal welfare of 6 and a respective fairness of 5, 8, and 9.³¹ When applying these definitions of welfare and fairness to the example game in Figure 4.4, it can be seen that the only solution which is both welfare-optimal and fairness-optimal is the joint policy that achieves expected joint return of (6, 6). Thus, in this example, we have narrowed the space of desirable solutions to a single solution. Another example of fairness-optimality, in the Battle of the Sexes matrix game, is shown in Figure 4.5.

31. The careful reader may have noticed some limitations of the simple definition of fairness in Definition 11. For example, if $U_i(\pi) = 0$ for any agent, then it does not matter what returns the other agents obtain under π . And if we allow $U_i(\pi) < 0$, then the expected joint return $(-0.1, 1, 1)$ would be less fair than the expected joint return $(-0.1, -100, 100)$, which is counter-intuitive.

Social welfare and fairness, such as defined here, can be useful in general-sum games, but are not so useful in common-reward games and zero-sum games. In common-reward games, in which all agents receive identical reward, welfare and fairness are maximised if and only if the expected return of each agent is maximised; hence, welfare and fairness do not add any useful criteria in this class of games. In two-agent zero-sum games, in which one agent's reward is the negative of the other agent's reward, we know that all minimax solutions π have the same unique value $U_i(\pi) = -U_j(\pi)$ for agents i, j . Therefore, all minimax solutions achieve equal welfare and fairness, respectively (besides, the welfare as defined here will always be zero).

It is easy to prove that welfare optimality implies Pareto optimality. To see this, suppose a joint policy π is welfare-optimal but not Pareto-optimal. Since π is not Pareto-optimal, there exists a joint policy π' such that $\forall i : U_i(\pi') \geq U_i(\pi)$ and $\exists i : U_i(\pi') > U_i(\pi)$. However, it follows that $\sum_i U_i(\pi') > \sum_i U(\pi)$ and, therefore, π cannot be welfare-optimal (contradiction). Thus, π must also be Pareto-optimal if it is welfare-optimal. Note that Pareto optimality does not in general imply welfare optimality, hence welfare optimality is a stronger requirement. Moreover, fairness optimality does not imply Pareto optimality, and vice versa.

4.10 No-Regret

The equilibrium solution concepts discussed in the previous sections are based on mutual best responses between agents, and are thus a function of the agents' policies. Another category of solution concepts is based on the notion of *regret*, which measures the difference between the rewards an agent received and the rewards it could have received if it had chosen a different action (or policy) in past episodes against the observed actions (or policies) of the other agents in these episodes. An agent is said to have *no-regret* if, in the limit of infinitely many episodes, the agent's average regret across the episodes is at most zero. Therefore, no-regret considers the performance of learning agents across multiple episodes, which is in contrast to the other solution concepts introduced in this chapter that only consider a single joint policy (and not *how* this joint policy was learned). In this sense, no-regret can be viewed as an example of the prescriptive agenda discussed in Section 1.5, which is concerned with the performance of agents during learning.

There are multiple ways in which regret can be defined. We will first give a standard definition of regret for non-repeated normal-form games, which is based on comparing the rewards of different actions in the episodes. This

Episode e	1	2	3	4	5	6	7	8	9	10
Action a_1^e	C	C	D	C	D	D	C	D	D	D
Action a_2^e	C	D	C	D	D	D	C	C	D	C
Reward $\mathcal{R}_1(a^e)$	-1	-5	0	-5	-3	-3	-1	0	-3	0
Reward $\mathcal{R}_1(\langle C, a_2^e \rangle)$	-1	-5	-1	-5	-5	-5	-1	-1	-5	-1
Reward $\mathcal{R}_1(\langle D, a_2^e \rangle)$	0	-3	0	-3	-3	-3	0	0	-3	0

Figure 4.6: Ten episodes between two agents in the non-repeated Prisoner’s Dilemma matrix game. The bottom two rows show agent 1’s rewards for always choosing actions C/D against agent 2’s observed actions in the episodes.

definition will then be extended to sequential-move games. Let a^e denote the joint action from episodes $e = 1, \dots, z$. Agent i ’s regret for not having chosen the best single action across these episodes is defined as

$$\text{Regret}_i^z = \max_{a_i \in A_i} \sum_{e=1}^z [\mathcal{R}_i(\langle a_i, a_{-i}^e \rangle) - \mathcal{R}_i(a^e)]. \quad (4.27)$$

An agent is said to have no-regret if its average regret in the limit of $z \rightarrow \infty$ is at most zero. As a solution concept, no-regret requires that all agents in the game have no-regret.

Definition 12 (No-regret) *In a general-sum game with n agents, the agents have no-regret if*

$$\forall i: \lim_{z \rightarrow \infty} \frac{1}{z} \text{Regret}_i^z \leq 0. \quad (4.28)$$

Similar to ϵ -Nash equilibrium (Section 4.5), we may replace the ≤ 0 in Equation 4.28 with $\leq \epsilon$, for $\epsilon > 0$, to obtain an ϵ -no-regret.

As a concrete example, Figure 4.6 shows ten episodes of two agents in the non-repeated Prisoner’s Dilemma matrix game. After the episodes, agent 1 has received a total reward of -21 . Always choosing C in the episodes (against the observed actions of agent 2) would have resulted in a total reward of -30 ; while always choosing D would have resulted in a total reward of -15 . Thus, action D was the “best” action against the observed actions of agent 2, and so agent 1’s regret is $\text{Regret}_1^{10} = -15 + 21 = 6$ (with an average regret, dividing by 10, of 0.6). In order for agent 1 to achieve no-regret, the episodes would need to continue in such a way that agent 1’s average regret across the episodes goes to zero.

We can generalise the definition of no-regret to stochastic games and POSGs by re-defining the regret over policies rather than actions. Let π^e denote the joint

policy from episodes $e = 1, \dots, z$. Then, agent i 's regret for not having chosen the best policy across these episodes is defined as

$$\text{Regret}_i^z = \max_{\pi_i \in \Pi_i} \sum_{e=1}^z [U_i(\langle \pi_i, \pi_{-i}^e \rangle) - U_i(\pi^e)] . \quad (4.29)$$

With this definition of regret, Definition 12 applies to all of the game models introduced in Chapter 3.

Our example above in Prisoner's Dilemma illustrates an important conceptual limitation of regret, which is that it assumes that the actions or policies of other agents $-i$ remain fixed in the episodes. This assumption is sensible if the other agents use constant policies that do not change between episodes. However, if the other agents adapt their policies based on past episodes, then this assumption is of course violated. Therefore, regret does not actually quantify what would have happened under counterfactual situations. The second limitation, which is a result of the first limitation, is that minimising regret is not necessarily equivalent to maximising returns (Crandall 2014). For example, in non-repeated and finitely-repeated Prisoner's Dilemma, the only joint policy that has no-regret is for both agents to always choose D. This is analogous to the fact that mutual best-response policies do not necessarily entail maximum returns for the agents, as we discussed in Section 4.7.

Various alternative definitions of regret exist (e.g. Farias and Megiddo 2003; Chang 2007; Arora, Dekel, and Tewari 2012; Crandall 2014). For example, for normal-form games in which agents can choose from more than two actions, rather than replacing all of agent i 's past actions as in Equation 4.27, we may replace only the specific occurrences of a given action a'_i in the history with a counterfactual action a_i . This latter definition of regret is also known as *internal* (or conditional) regret, while the definition in Equation 4.27 is known as *external* (or unconditional) regret. Furthermore, no-regret solutions have connections to equilibrium solutions. In particular, in two-agent zero-sum normal-form games, the empirical distribution of joint actions produced by agents that have no external regret converges to a minimax solution; and in general-sum normal-form games, the empirical distribution of joint actions produced by agents that have no internal regret converges to a correlated equilibrium (Greenwald and Jafari 2003; Young 2004).

4.11 The Complexity of Computing Equilibria

Before we turn to MARL algorithms in Chapters 5 and 6 as a method to compute solutions for games, it is instructive to ask: How difficult is it, in

terms of computational complexity, to compute an equilibrium solution for a game? Do algorithms exist that can compute equilibria *efficiently*, meaning in polynomial time in the size of the game?

These and related questions are studied in algorithmic game theory, which is a research area at the interface of computer science and game theory. Many complexity results exist for various special types of games, and we recommend the books of Nisan et al. (2007) and Roughgarden (2016) for a broad discussion. Here, we focus on non-repeated normal-form games which are the building block of the more complex (partially observable) stochastic game models. Thus, we can expect any complexity results for normal-form games to be a lower bound on the complexity for the more complex game models.

Most computer scientists will have some familiarity with the complexity classes P and NP. P includes all decision problems whose solutions (if they exist) can be computed efficiently in polynomial time in the size of the problem instance. NP includes all decision problems whose solutions (if they exist) can be computed in polynomial time by a non-deterministic Turing machine, and in the worst case require exponential time by a deterministic Turing machine. Unfortunately, these familiar complexity classes are not a good fit for the problem of computing an equilibrium, since P/NP characterise *decision problems* which may or may not have solutions, while we know that games *always* have at least one equilibrium solution. On the other hand, computing an equilibrium which satisfies additional properties *is* a decision problem, since such solutions may or may not exist. Such problems include computing equilibria which:

- are Pareto optimal
- achieve a certain minimum expected return for each agent
- achieve a certain minimum social welfare (sum of returns)
- assign zero or positive probability to certain actions of certain agents.

All of these problems are known to be NP-hard (Gilboa and Zemel 1989; Conitzer and Sandholm 2008).

Some types of games and equilibria do admit polynomial-time algorithms. Computing a minimax solution in a two-agent zero-sum non-repeated normal-form game can be formulated via a linear programme (Section 4.3.1), which can be solved in polynomial time. Similarly, computing a correlated equilibrium in a general-sum non-repeated normal-form game can be done in polynomial time via a linear programme (Section 4.6.1). However, computing a Nash equilibrium in general-sum non-repeated normal-form games—henceforth simply called NASH—cannot be solved via linear programming, due to the independence assumption between policies in a Nash equilibrium.

Problems which always have solutions, such as NASH, are known as *total search problems*. Section 4.11.1 will present one subclass of total search problems, called PPAD. It turns out that NASH is a *complete* problem in PPAD, meaning that any other problem in PPAD can be reduced to NASH. We will discuss the implications for MARL in Section 4.11.2.

4.11.1 PPAD Complexity Class

PPAD (short for “polynomial parity argument for directed graphs”) describes a special class of total search problems. We define PPAD by giving one of its *complete* problems to which all other problems in PPAD can be reduced.³² This PPAD-complete problem is called END-OF-LINE, and is defined as follows:

Definition 13 (END-OF-LINE) *Let $G(k) = \{(V, E)\}$ be a set of directed graphs which consist of*

- *a finite set V containing 2^k nodes (each node is represented as a bit-string of length k)*
- *a finite set $E = \{(a, b) \mid a, b, \in V\}$ of directed edges from a to b such that if $(a, b) \in E$ then $\exists a' \neq a : (a', b) \in E$ and $\exists b' \neq b : (a, b') \in E$.*

Assume functions $Parent(v)$ and $Child(v)$ which, respectively, return the parent node (if any) and child node (if any) of node $v \in V$. These functions are represented as boolean circuits with k input bits and k output bits, and run in time polynomial in k . Given functions $Parent$ and $Child$, and a node $s \in V$ such that $Parent(s) = \emptyset$, find a node $e \neq s$ such that either $Child(e) = \emptyset$ or $Parent(e) = \emptyset$.³³

Figure 4.7 shows an illustration of a END-OF-LINE problem instance. The restriction on E means that any node in the graph can have at most one parent node and at most one child node. A node with no parent is also called a source node, while a node with no child is called a sink node. The “parity argument” in PPAD refers to the fact that any source node in this graph always has a corresponding sink node. Therefore, if a source node s is given, then we know that a node e must exist. The node e can be found by following the directed path starting at the given source node s . However, since we are only given the functions $Parent$ and $Child$ (and not E), the only obvious way to follow this path is by repeatedly calling the function $Child$ starting at the source node s .

32. Problem A can be “reduced” to problem B if there exist polynomial-time algorithms to transform any instance of A into an equivalent instance of B, and any solution of the instance of B to a solution of the original instance of A.

33. To indicate that a node has no parent/child, the respective circuit function simply outputs the same input node.

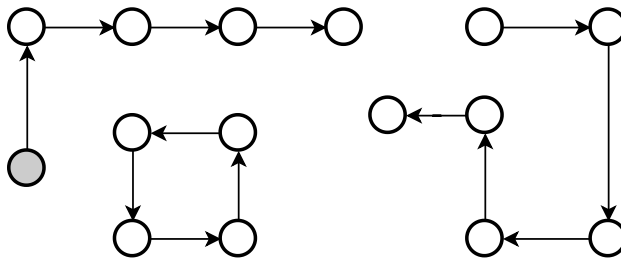


Figure 4.7: Any instance of END-OF-LINE consists of a set of paths and cycles, such as the instance shown here. Given a source node (shaded in grey), is there an efficient polynomial-time algorithm to find a sink node (or a different source node), without following the path starting at the given source node?

Therefore, finding a sink node may require exponential time in the worst case, since there are 2^k nodes.

Why should we care about PPAD? Just as it is unknown whether efficient polynomial-time algorithms exist to solve NP-complete problems (the big “P = NP?” question), it is also unknown whether efficient algorithms exist for PPAD-complete problems (“P = PPAD?”). Indeed, PPAD contains problems for which researchers have tried for decades to find efficient algorithms, including the classic Brouwer fixed-point problem and finding Arrow-Debreu equilibria in markets (see Papadimitriou (1994) for a more complete list). There are currently no known algorithms which can efficiently solve END-OF-LINE (in time polynomial in k) and, thus, any other problem in PPAD. Therefore, establishing that a problem is PPAD-complete is a good indicator that no efficient algorithms exist to solve the problem.

4.11.2 Computing ϵ -Nash Equilibrium is PPAD-Complete

We return to our initial question, “Do algorithms exist that can compute equilibria *efficiently*, in polynomial time in the size of the game?” Unfortunately, the answer is very likely negative. It has been proven that NASH is PPAD-complete, at first for games with three or more agents (Daskalakis, Goldberg, and Papadimitriou 2006, 2009), and shortly after even for games with two agents (Chen and Deng 2006). This means that finding a Nash equilibrium in a non-repeated normal-form game can be described as finding an ϵ -node in an equivalent END-OF-LINE instance. The completeness also means that any other problem in PPAD, including Brouwer fixed-points, can be reduced to NASH.

More precisely, the PPAD-completeness of NASH was proven for approximate ϵ -Nash equilibrium (Section 4.5) for certain bounds on $\epsilon > 0$; and for

exact equilibria (when $\epsilon=0$) in games with two agents.³⁴ The use of ϵ -Nash equilibrium is to account for the fact that a Nash equilibrium may involve irrational-valued probabilities in games with more than two agents. Therefore, the PPAD-completeness of NASH also includes approximation schemes for computing Nash equilibria, such as MARL algorithms which may only learn approximate solutions given a finite number of interactions in the game.

The implication for us is that MARL is unlikely to be a magic bullet for solving games: since no efficient algorithms are known to exist for PPAD-complete problems, it is unlikely that efficient MARL algorithms exist to compute Nash equilibria in polynomial time. Much of the research in MARL has focused on identifying and exploiting structures (or assumptions) in certain game types which lead to improved performance. However, the PPAD-completeness of NASH tells us that, in general and without such assumed structures, it is likely that *any* MARL algorithm still requires exponential time in the worst case.

34. As discussed in Section 4.5, an ϵ -Nash equilibrium may not be close to any actual Nash equilibrium, even if the Nash equilibrium is unique. The problem of approximating an actual Nash equilibrium within a desired distance, as measured in the policy space via norms such as L1 and L2, is in fact much harder than computing an ϵ -Nash equilibrium. This latter problem, for general-sum normal-form games with three or more agents, is complete for a different complexity class called FIXP (Etessami and Yannakakis 2010).

5 Multi-Agent Reinforcement Learning in Games: First Steps and Challenges

The preceding chapters introduced game models as a formalism of multi-agent interaction, and solution concepts to define what it means for the agents to act optimally in a game. In this chapter, we will begin to explore methods to *compute* solutions for games. The principal method by which we seek to compute solutions is via reinforcement learning (RL), in which the agents repeatedly try actions, make observations, and receive rewards. Analogous to the standard RL terminology introduced in Chapter 2, we use the term *episode* to refer to each independent run of a game starting in some initial state. The agents learn their policies based on data (i.e. observations, actions, and rewards) obtained from multiple episodes in a game.

To set the context for the algorithms presented in this book, this chapter will begin by outlining a general learning framework for MARL, as well as different types of convergence definitions used in the analysis and evaluation of MARL algorithms. We will then introduce two basic approaches of applying RL in games, called central learning and independent learning, both of which reduce the multi-agent problem to a single-agent problem. Central learning applies single-agent RL directly to the space of joint actions to learn a central policy that chooses actions for each agent, while independent learning applies single-agent RL to each agent independently to learn agent policies, essentially ignoring the presence of other agents.

Central and independent learning serve as a useful starting point to discuss several important challenges faced by MARL algorithms. One characteristic challenge of MARL is environment non-stationarity due to multiple learning agents, which can lead to unstable learning. Another challenge is multi-agent credit assignment, in which agents must infer *whose* actions contributed to a received reward. Equilibrium selection is the problem of what equilibrium solution the agents should agree on and how they may achieve agreement. Finally, MARL algorithms are typically faced with an exponential growth of

the joint action space as the number of agents is increased, leading to scalability problems. We will discuss each of these challenges and provide examples.

The idea that agents can use their own algorithms to learn policies, such as in independent learning, leads to the possibility that agents may use the same learning algorithm or different algorithms. This chapter will conclude with a discussion of such self-play and mixed-play settings in MARL.

5.1 General Learning Process

We begin by defining *learning*³⁵ in games and the intended learning outcome. In machine learning, learning is a process which optimises a model or function based on *data*. In our setting, the model is a joint policy usually consisting of policies for each agent, and the data (or “experiences”) consist of one or more histories in the game. The learning goal is a solution of the game, defined by a chosen solution concept. Thus, this learning process involves several elements:

Game model: The game model defines the multi-agent environment and how agents may interact. Game models introduced in Chapter 3 include non-repeated normal-form games, repeated normal-form games, stochastic games, and partially observable stochastic games (POSG).

Data: The data used for learning consist of a set of z histories,

$$\mathcal{D}^z = \{h^{t_e} \mid e = 1, \dots, z\}, z \geq 0. \quad (5.1)$$

Each history h^{t_e} was produced by a joint policy π^e used during episode e . These histories may or may not be “complete” in the sense of ending in a terminal state of the game, and different histories may have different lengths t_e . Often, \mathcal{D}^z contains the history so far from the current ongoing episode z and the histories from previous episodes $e < z$.

Learning algorithm: The learning algorithm \mathbb{L} takes the collected data \mathcal{D}^z and current joint policy π^z , and produces a new joint policy,

$$\pi^{z+1} = \mathbb{L}(\mathcal{D}^z, \pi^z). \quad (5.2)$$

The initial joint policy π^0 is typically random.

Learning goal: The goal of learning is a joint policy π^* which satisfies the properties of a chosen solution concept. Chapter 4 introduced a range of possible solution concepts, such as Nash equilibrium.

35. Another commonly used term is “training”. We use the terms learning and training interchangeably, e.g. as in “learning/training a policy”.

We note several nuances in the above elements:

The chosen game model determines the conditioning of the learned joint policy. In a non-repeated normal-form game (where episodes terminate after one time step), policies π_i are not conditioned on histories, i.e. they are simple probability distributions over actions. In a repeated normal-form game, policies are conditioned on action histories $h^t = (a^0, \dots, a^{t-1})$. In a stochastic game, policies are conditioned on state-action histories $h^t = (s^0, a^0, s^1, a^1, \dots, s^t)$. In a POSG, policies are conditioned on observation histories $h_i^t = (o_i^0, \dots, o_i^t)$. These conditionings are general and may be constrained depending the desired form of policies. For example, in a stochastic game we may condition policies only on the current state of the game; and in a POSG we may condition policies using only the most recent k observations.

The histories in \mathcal{D}^z may in general be *full* histories (i.e. contain all states and joint observations/actions; see Section 4.1) or one of the other types of histories listed above. Therefore, the histories in \mathcal{D}^z may contain more information than the histories used to condition the agents' policies. For example, this can be the case in centralised training with decentralised execution regimes (discussed in Section 9.1), in which a learning algorithm has access to the observations of all agents during learning, while the agents' policies only have access to the local observations of agents.

The learning algorithm \mathbb{L} may itself consist of multiple learning algorithms that learn individual agent policies, such as one algorithm \mathbb{L}_i for each agent i . Each of these algorithms may use different parts of the data in \mathcal{D}^z , or use its own data \mathcal{D}_i^z such as in independent learning (Section 5.3.2). Furthermore, an important characteristic of RL is that the learning algorithm is actively involved in the generation of the data by exploring actions, rather than just passively consuming the data. Thus, the policies produced by the learning algorithm may actively randomise over actions to generate useful data for learning.

5.2 Convergence Types

Many different criteria have been developed to evaluate the learning performance of MARL algorithms. In this book, the main theoretical evaluation criterion for learning we use is convergence of the joint policy π^z to a solution π^* of the game in the limit of infinite data,

$$\lim_{z \rightarrow \infty} \pi^z = \pi^*. \quad (5.3)$$

In this book, when we make statements about theoretical convergence properties of MARL algorithms, we mean convergence as per Equation 5.3 unless

specified otherwise. Of course, in practice we cannot collect infinite data, and learning typically stops after a predefined budget is reached (such as a total allowed number of episodes or time steps), or once the changes in policies are below some predetermined threshold. The equality $\pi^z = \pi^*$ may be tested using procedures such as described in Sections 4.4 and 4.5.

Several other theoretical evaluation criteria have been studied in the literature, including both weaker and stronger types of convergence. Weaker types of convergence include:

- Convergence of expected return:

$$\lim_{z \rightarrow \infty} U_i(\pi^z) = U_i(\pi^*), \forall i \in I \quad (5.4)$$

This convergence type means that, in the limit of infinite data $z \rightarrow \infty$, the expected joint return under the learned joint policy π^z will converge to the expected joint return of a Nash equilibrium.

- Convergence of empirical action distribution:

$$\lim_{z \rightarrow \infty} \bar{\pi}^z = \pi^* \quad (5.5)$$

where $\bar{\pi}^z(a|h) = \frac{1}{z} \sum_{e=1}^z \pi^e(a|h)$ is the averaged joint policy across learning. Intuitively, this convergence type can be interpreted as saying that if we generate a new episode for each updated joint policy π^x , then the averaged joint policy $\bar{\pi}^z$ that produces the actions from these episodes in expectation will converge to a Nash equilibrium.

- Convergence of average return:

$$\lim_{z \rightarrow \infty} \bar{U}_i^z = U_i(\pi^*), \forall i \in I \quad (5.6)$$

where $\bar{U}_i^z = \frac{1}{z} \sum_{e=1}^z U_i(\pi^e)$ is the averaged expected return across learning. Intuitively, this convergence type can be interpreted as saying that if we generate a new episode for each updated joint policy π^e , then the average of the joint returns from these episodes will converge to the expected joint return of a Nash equilibrium.

These weaker convergence types are often used when a particular learning algorithm is not technically able to achieve the convergence in Equation 5.3. For example, fictitious play (discussed in Section 6.3.1) produces only deterministic policies, meaning that it cannot represent probabilistic Nash equilibria, such as the uniform-random Nash equilibrium in Rock-Paper-Scissors. However, in certain cases, it can be proven that the empirical action distribution produced by fictitious play converges to a Nash equilibrium, as per Equation 5.5 (Fudenberg

and Levine 1998). Another example is infinitesimal gradient ascent (discussed in Section 6.4.1), which *can* learn probabilistic policies but may still not converge to a probabilistic Nash equilibrium, while it can be shown in non-repeated normal-form games that the average rewards produced by the algorithm do converge to the average rewards of a Nash equilibrium, as per Equation 5.6 (Singh, Kearns, and Mansour 2000).

Note that Equation 5.3 implies all of the above weaker types of convergence. However, it makes no claims about the performance of any individual joint policy π^z for a finite z . In other words, the above convergence types leave open how the agents perform *during* learning. To address this, a stronger evaluation criterion could require additional bounds, such as on the difference between π^z and π^* for finite z . (See also the discussion and references in Section 5.5.2.)

In complex games, it may not be computationally practical to check for these convergence properties. Instead, a common approach is to monitor the expected returns $U_i(\pi^z)$ achieved by the joint policy as z increases, usually by visualising learning curves that show the progress of expected returns during learning, as shown in Section 2.7. A number of such learning curves will be shown for various MARL algorithms presented in this book. However, this evaluation approach may not establish any relationship to the solution π^* of the game. For instance, even if the expected returns $U_i(\pi^z)$ for all $i \in I$ converge after some finite z , the joint policy π^z might not satisfy any of the convergence properties of Equations 5.3 to 5.6.

To reduce notation in the remainder of this chapter (and book), we will omit the explicit z -superscript (e.g. π^z , \mathcal{D}^z) and we usually omit \mathcal{D}^z altogether.

5.3 Single-Agent RL Reductions

The most basic approach to using RL to learn agent policies in multi-agent systems is to essentially reduce the multi-agent learning problem to a single-agent learning problem. In this section, we will introduce two such approaches: *central learning* applies single-agent RL directly to the space of joint actions to learn a central policy that chooses actions for each agent; and *independent learning* applies single-agent RL independently to each agent to learn independent policies, essentially ignoring the presence of other agents.

5.3.1 Central Learning

Central learning trains a single central policy π_c which receives the local observations of all agents and selects an action for each agent, by selecting joint actions from $A = A_1 \times \dots \times A_n$. This essentially reduces the multi-agent problem

Algorithm 4 Central Q-Learning (CQL) for Stochastic Games

- 1: Initialise: $Q(s, a) = 0$ for all $s \in S$ and $a \in A = A_1 \times \dots \times A_n$
- 2: Repeat for every episode:
- 3: **for** $t = 0, 1, 2, \dots$ **do**
- 4: Observe current state s^t
- 5: With probability ϵ : choose random joint action $a^t \in A$
- 6: Else: choose joint action $a^t \in \arg \max_a Q(s^t, a)$
- 7: Apply joint action a^t , observe rewards r_1^t, \dots, r_n^t and next state s^{t+1}
- 8: Transform r_1^t, \dots, r_n^t into scalar reward r^t
- 9: $Q(s^t, a^t) \leftarrow Q(s^t, a^t) + \alpha [r^t + \gamma \max_{a'} Q(s^{t+1}, a') - Q(s^t, a^t)]$

to a single-agent problem, and we can apply existing single-agent RL algorithms to train π_c . An example of central learning based on Q-learning, called *central Q-learning* (CQL), is shown in Algorithm 4. This algorithm maintains *joint action values* $Q(s, a)$ for joint actions $a \in A$, which is a basic concept used by many MARL algorithms presented in this book. Central learning can be useful because it circumvents the multi-agent aspects of the non-stationarity and credit assignment problems (discussed in Sections 5.4.1 and 5.4.2, respectively). However, in practice, this approach has a number of limitations.

The first limitation to note is that, in order to apply single-agent RL, central learning requires transforming the joint reward (r_1, \dots, r_n) into a single scalar reward r . For the case of common-reward games, in which all agents receive identical rewards, we can use $r = r_i$ for any i . In this case, if we use a single-agent RL algorithm that is guaranteed to learn an optimal policy in an MDP (such as the temporal-difference algorithms discussed in Section 2.6), then it is guaranteed to learn a central policy π_c for the common-reward stochastic game such that π_c is a Pareto-optimal correlated equilibrium. The optimality of the single-agent RL algorithm means that π_c achieves maximum expected returns in each state $s \in S$ (as discussed in Section 2.4). Therefore, since the reward is defined as $r = r_i$ for all i , we know that π_c is Pareto-optimal because there can be no other policy that achieves a higher expected return for any agent. This also means that no agent can unilaterally deviate from its action given by π_c to improve its returns, making π_c a correlated equilibrium.

Unfortunately, for zero-sum and general-sum stochastic games, it is less clear how the scalarisation should be done. If one is interested in maximising social welfare (Section 4.9) in general-sum games, one option is to use $r = \sum_i r_i$. However, if the desired solution is an equilibrium type solution, then no scalar transformation may exist which leads to equilibrium policies.

The second limitation is that by training a policy over the joint action space, we now have to solve a decision problem with an action space that grows exponentially in the number of agents.³⁶ In the level-based foraging example shown in Figure 1.2 (page 4), there are three agents which choose from among six actions (up, down, left, right, collect, noop), leading to a joint action space with $6^3 = 216$ actions. Even in this toy example, most standard single-agent RL algorithms do not scale easily to action spaces this large.

Finally, a fundamental limitation of central learning is due to the inherent structure of multi-agent systems. Agents are often localised entities which are physically or virtually distributed. In such settings, communication from a central policy π_c to the agents and vice versa may not be possible or desirable, for various reasons. Therefore, such multi-agent systems require local agent policies π_i for each agent i , which act on agent i 's local observations and independently from other agents.

In stochastic games, in which we assume that agents can observe the full state, it is possible to decompose the central policy π_c into individual agent policies π_1, \dots, π_n . This is because an MDP always has a deterministic optimal policy which assigns probability 1 to some action in every state (see Equation 2.25 in Section 2.4). Thus, we can decompose π_c as

$$\pi_c(s) = (\pi_1(s)=a_1, \dots, \pi_n(s)=a_n) \quad (5.7)$$

where $\pi(s)=a$ is a shorthand for $\pi(a|s)=1$, and analogously for $\pi_i(s)=a_i$. A basic way to achieve this decomposition is simply for every agent to use a copy of π_c : in any given state s , each agent $i \in I$ computes the joint action $(a_1, \dots, a_n) = \pi_c(s)$ and executes its own action a_i from the joint action. However, if the environment is only partially observable by the agents (as in POSGs), then it may not be possible to decompose π_c into individual agent policies π_i , since each policy π_i now only has access to the agent's own observations o_i .

The next section will introduce the independent learning approach that eliminates these limitations, at the expense of introducing other challenges.

5.3.2 Independent Learning

In *independent learning* (often abbreviated as IL), each agent i learns its own policy π_i using only its local history of own observations, actions, and rewards, while ignoring the existence of other agents (Tan 1993; Claus and Boutilier

36. This exponential growth assumes that each additional agent comes with additional decision variables. For example, in level-based foraging (Section 1.1), each agent comes with its own action set. The reverse direction is when we have a fixed set of actions which are partitioned and assigned to agents. In the latter case, the total number of actions remains constant, regardless of the number of agents. We discuss this further in Section 5.4.4.

Algorithm 5 Independent Q-Learning (IQL) for Stochastic Games

```

// Algorithm controls agent i
1: Initialise:  $Q_i(s, a_i) = 0$  for all  $s \in S, a_i \in A_i$ 
2: Repeat for every episode:
3:   for  $t = 0, 1, 2, \dots$  do
4:     Observe current state  $s^t$ 
5:     With probability  $\epsilon$ : choose random action  $a_i^t \in A_i$ 
6:     Else: choose action  $a_i^t \in \arg \max_{a_i} Q_i(s^t, a_i)$ 
7:     (meanwhile, other agents  $j \neq i$  choose their actions  $a_j^t$ )
8:     Observe own reward  $r_i^t$  and next state  $s^{t+1}$ 
9:      $Q_i(s^t, a_i^t) \leftarrow Q_i(s^t, a_i^t) + \alpha [r_i^t + \gamma \max_{a_i'} Q_i(s^{t+1}, a_i')] - Q_i(s^t, a_i^t)]$ 

```

1998). Agents do not observe or use information about other agents, and the effects of other agents' actions are simply part of the environment dynamics from the perspective of each learning agent. Thus, similar to central learning, independent learning reduces the multi-agent problem to a single-agent problem from the perspective of each agent, and existing single-agent RL algorithms can be used to learn the agent policies. An example of independent learning based on Q-learning, called *independent Q-learning* (IQL), is shown in Algorithm 5. Here, each agent uses its own copy of the same algorithm.

Independent learning naturally avoids the exponential growth in action spaces that plagues central learning, and it can be used when the structure of the multi-agent system requires local agent policies. It also does not require a scalar transformation of the joint reward, as is the case in central learning. The downside of independent learning is that it can be significantly affected by non-stationarity caused by the concurrent learning of all agents. In an independent learning algorithm such as IQL, from the perspective of each agent i the policies π_j of other agents $j \neq i$ become part of the environment's state transition function via³⁷

$$\mathcal{T}_i(s^{t+1} | s^t, a_i^t) = \eta \sum_{a_{-i} \in A_{-i}} \mathcal{T}(s^{t+1} | s^t, \langle a_i^t, a_{-i} \rangle) \prod_{j \neq i} \pi_j(a_j | s^t) \quad (5.8)$$

where \mathcal{T} is the game's original state transition function defined over the joint action space (Section 3.3), and η is a normalisation constant to ensure that $\mathcal{T}_i(\cdot | s^t, a_i^t)$ is a valid probability distribution.

As each agent j continues to learn and update its policy π_j , the action probabilities of π_j in each state s may change. Thus, from the perspective of agent i ,

37. Recall that we use $-i$ in the subscript to mean "all agents other than agent i " (e.g. $A_{-i} = \times_{j \neq i} A_j$).

Subclass	1a	1b	2a	2b	3a	3b
# deterministic NE	0	0	2	2	1	1
# probabilistic NE	1	1	1	1	0	0
Dominant action?	No	No	No	No	Yes	Yes
Det. joint act. > NE?	No	Yes	No	Yes	No	Yes
IQL converges?	Yes	No	Yes	Y/N	Yes	Y/N

Figure 5.1: Convergence of “infinitesimal” independent Q-learning (IQL) in repeated general-sum normal-form games with two agents and two actions (Wunder, Littman, and Babes 2010). Games are characterised by: 1) number of deterministic Nash equilibria; 2) number of probabilistic Nash equilibria; 3) whether at least one of the agents has a dominant action in the game; 4) whether there exists a joint action which leads to higher rewards for both agents than the Nash equilibrium of the game (or equilibrium with lowest rewards in class 2a/b). Rewards are real-valued, hence each game class contains an infinite number of games. Class 2b includes Chicken, and class 3b includes Prisoner’s Dilemma. In the bottom row, “Yes” means that IQL will converge to a Nash equilibrium, “No” means that IQL will not converge, and “Y/N” means that IQL converges under certain conditions.

the transition function \mathcal{T}_i appears to be non-stationary, when really the only parts that change over time are the policies π_j of the other agents. As a result, independent learning approaches may produce unstable learning and may not converge to any solution of the game. Section 5.4.1 discusses non-stationarity in MARL in further detail.

The learning dynamics of independent learning have been studied in various idealised models (Rodrigues Gomes and Kowalczyk 2009; Wunder, Littman, and Babes 2010; Kianercy and Galstyan 2012; Barfuss, Donges, and Kurths 2019; Hu, Leung, and Leung 2019; Leonardos and Piliouras 2022). For example, Wunder, Littman, and Babes (2010) study an idealised model of IQL with epsilon-greedy exploration (as shown in Algorithm 5) that uses infinitely small learning steps $\alpha \rightarrow 0$, which makes it possible to apply methods from linear dynamical systems theory to analyse the dynamics of the model. Based on this idealised model, predictions about the learning outcomes of IQL can be made for different classes of repeated general-sum normal-form games with two agents and two actions, which are summarised in Figure 5.1. These classes are primarily characterised by the number of deterministic and probabilistic Nash equilibria the games possess. As can be seen, this idealised version of IQL is predicted to converge to a Nash equilibrium in some of the game classes,

while in others it may not converge at all or only under certain conditions. An interesting finding of this analysis is that in games such as Prisoner’s Dilemma, which is a member of class 3b, IQL can have a chaotic non-convergent behaviour which results in rewards that average above the expected reward under the unique Nash equilibrium of the game.

Despite their relative simplicity, independent learning algorithms still serve as important baselines in MARL research. In fact, they can often produce results which are competitive with state-of-the-art MARL algorithms, as shown by the study of Papoudakis et al. (2021). In Chapter 9, we will see some of the independent learning algorithms used in current MARL research.

5.3.3 Example: Level-Based Foraging

We compare the performance of central Q-learning (CQL), given in Algorithm 4, and independent Q-learning (IQL), given in Algorithm 5, in an instance of the level-based foraging environment shown in Figure 5.2. In this level-based foraging task, two agents must collaborate in order to collect two items in a 11 by 11 grid world. In each time step, each agent can move in one of the four directions, attempt to collect an item, or do nothing. Each agent and item has a skill level, and one or more agents can collect an item if they are positioned next to the item, they simultaneously attempt to collect, and the sum of their levels is equal or higher than the item’s level. Every episode begins with the same start state shown in Figure 5.2, in which both agents have level 1 and are initially located in the top corners of the grid, and there are two items located in the centre with levels 1 and 2, respectively. Therefore, while the level-1 item can be collected by any individual agent, the level-2 item requires that the two agents collaborate in order to collect the item.

We model this learning problem as a stochastic game in which the agents observe the full environment state, using a discounted return objective with discount factor $\gamma = 0.99$. An agent receives a reward of $\frac{1}{3}$ when collecting the level-1 item, and both agents receive a reward of $\frac{1}{3}$ when collecting the level-2 item; thus, the total (undiscounted) reward between the agents after collecting all items is 1.³⁸ Due to the reward discounting, the agents will need to collect the items as quickly as possible in order to maximise their returns. Episodes terminate when all items have been collected or after a maximum of 50 time steps. In this example, both algorithms use a constant learning rate $\alpha = 0.01$, and an exploration rate which is linearly decayed from $\epsilon = 1$ to $\epsilon = 0.05$ over the first

38. See Section 11.3.1 for a more general definition of the reward function in level-based foraging.

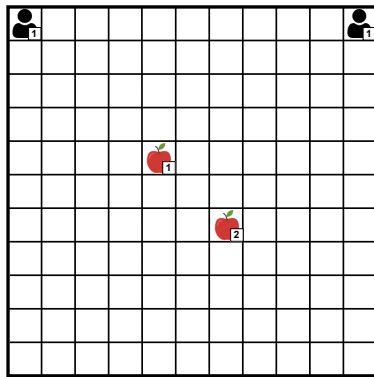


Figure 5.2: Level-based foraging task used to compare central Q-learning and independent Q-learning. This example task consists of a 11 by 11 grid world with two agents and two items. Both agents have level 1, while one item has level 1 and another item has level 2. All episodes start in this shown state with the shown positions and levels.

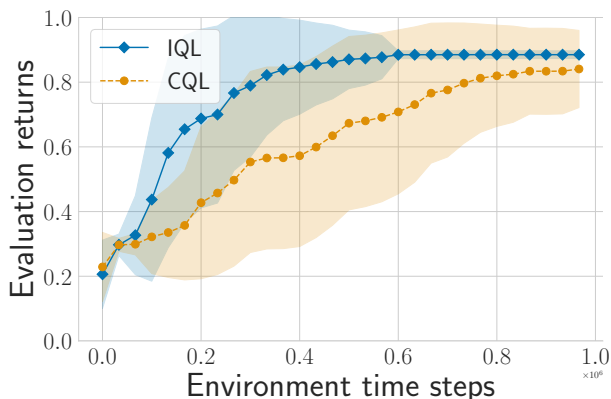


Figure 5.3: Average discounted evaluation returns of central Q-learning (CQL) and independent Q-learning (IQL) in the level-based foraging task shown in Figure 5.2, with discount factor $\gamma = 0.99$. Results are averaged over 50 independent training runs. The shaded area shows the standard deviation over the averaged returns from each training run.

80,000 time steps. For CQL, we obtain a scalar reward (Line 8 in Algorithm 4) by summing the individual rewards, i.e. $r^t = r_1^t + r_2^t$.

Figure 5.3 shows the evaluation returns³⁹ achieved by CQL and IQL in the level-based foraging task. Note that Figure 5.3 shows discounted returns with discount factor $\gamma = 0.99$, hence the maximum achievable return is also less than 1. IQL learns to solve this task faster than CQL, which is a result of the fact that the IQL agents only explore 6 actions in each state while the CQL agent has to explore $6^2 = 36$ actions in each state. This allows the IQL agents to learn more quickly to collect the level-1 item, which can be seen in the early jump in evaluation returns of IQL. Eventually, both IQL and CQL converge to the optimal joint policy in this task, in which the agent in the right corner goes directly to the level-2 item and waits for the other agent to arrive; meanwhile, the agent in the left corner first goes to the level-1 item and collects it, and then goes to the level-2 item and collects it together with the other agent. This optimal joint policy requires 13 time steps to solve the task.

5.4 Challenges of MARL

MARL algorithms, including central Q-learning and independent Q-learning, inherit certain challenges from single-agent RL such as non-stationarity from bootstrapping and temporal credit assignment, and face additional conceptual and algorithmic challenges as a result of learning in a dynamic multi-agent system. This section describes several of the core challenges which together characterise MARL.

5.4.1 Non-Stationarity

A central problem in MARL is the non-stationarity resulting from the continual co-adaptation of multiple agents as they learn from interactions with one another. This non-stationarity can lead to cyclic dynamics, whereby a learning agent adapts to the changing policies of other agents, which in turn may adapt their policies to the learning agent’s policy, and so on.

An illustrative example of such learning dynamics can be seen in Figure 5.4, in which two agents play the repeated Rock-Paper-Scissors matrix game using the WoLF-PHC algorithm (which will be introduced in Section 6.4.4) to update their policies in each time step. The lines show how the agent’s policies co-adapt over time until converging to the unique Nash equilibrium of the game, in which both agents choose each action with equal probability.

³⁹ For a reminder of the term “evaluation return” and how these learning curves are produced, see Section 2.7.

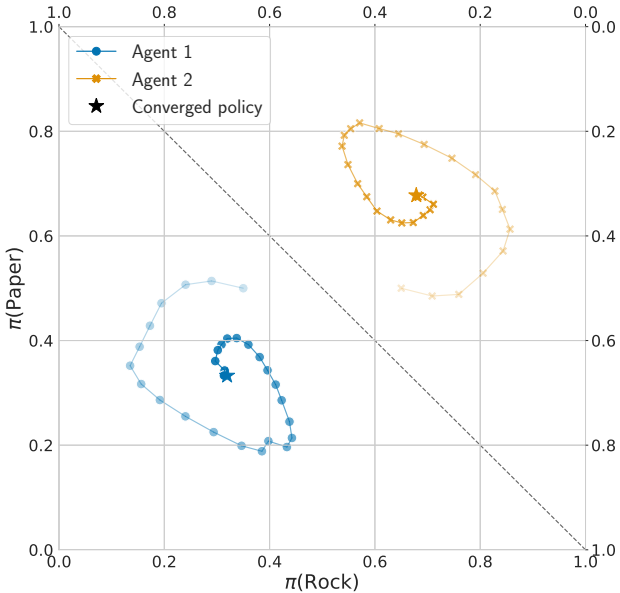


Figure 5.4: This plot shows the evolving policies of two agents in the Rock-Paper-Scissors game where both agents use the WOLF-PHC algorithm (Section 6.4.4) to update their policies. Each point in the simplex for an agent corresponds to a probability distribution over the three available actions. Each line shows the current policy at time steps $t = 0, 5, 10, 15, \dots, 145$ as well as the converged policy at $t = 100,000$.

While non-stationarity caused by the learning of multiple agents is a defining characteristic of MARL, it is important to note that non-stationarity already exists in the single-agent RL setting. To gain clarity on the problem, it is useful to define the notion of stationarity and see how stationarity and non-stationarity appear in single-agent RL, before moving to the multi-agent RL setting.

A stochastic process $\{X^t\}_{t \in \mathbb{N}^0}$ is said to be stationary if the probability distribution of $X^{t+\tau}$ does not depend on $\tau \in \mathbb{N}^0$, where t and $t + \tau$ are time indices. Intuitively, this means that the dynamics of the process do not change over time.

Now, consider a stochastic process X^t which samples the state s^t at each time step t . In a Markov decision process, the process X^t is completely defined by the state transition function, $\mathcal{T}(s^t | s^{t-1}, a^{t-1})$, and the agent's policy, π , which selects the actions $a \sim \pi(\cdot | s)$. If this policy does not change over time (meaning that no learning takes place), then it can be seen that the process X^t is indeed stationary, since s^t depend only on the state s^{t-1} and action a^{t-1} from the previous time step

(known as the Markov property), and a^{t-1} depends only on s^{t-1} via $\pi(\cdot | s^{t-1})$. Therefore, the process dynamics are independent of the time t .

However, in reinforcement learning, the policy π changes over time as a result of the learning process. Using the definition of a learning algorithm from Section 5.1, the policy π^z at time t is updated via $\pi^{z+1} = \mathbb{L}(\mathcal{D}^z, \pi^z)$, where \mathcal{D}^z contains all data collected up to time t in the current episode z as well as data from the previous episodes. Therefore, in single-agent RL, the process X^t is non-stationary since the policy depends on t .

This non-stationarity is a problem when learning the values of states or actions, since the values depend on subsequent actions which can change as the policy π changes over time. For example, when using temporal-difference learning to learn action values as described in Section 2.6, the value estimate $Q(s^t, a^t)$ is updated via an update target that depends on value estimates for a different state, such as in the target $r^t + \gamma Q(s^{t+1}, a^{t+1})$ used in the Sarsa algorithm. As the policy π changes during learning, the update target becomes non-stationary since the value estimate used in the target is also changing. For this reason, the non-stationarity problem is also referred to as the *moving target problem*.

In MARL, the non-stationarity is exacerbated by the fact that *all* agents change their policies over time; here, $\pi^{z+1} = \mathbb{L}(\mathcal{D}^z, \pi^z)$ updates an entire joint policy $\pi^z = (\pi_1^z, \dots, \pi_n^z)$. This adds another difficulty for the learning agents in that not only do the value estimates face non-stationarity (as in single-agent RL), but the entire environment appears to be non-stationary from the perspective of each agent. As we discussed in Section 5.3.2, this is encountered in independent learning algorithms such as IQL: from the perspective of agent i , the policies of other agents $j \neq i$ become part of the environment's state transition function, as shown in Equation 5.8 (page 90).

Because of these non-stationarity issues, the usual stochastic approximation conditions required for temporal-difference learning in single-agent RL (defined in Equation 2.53, page 33) are usually not sufficient in MARL to ensure convergence. Indeed, all known theoretical results in MARL for convergent learning are limited to restricted game settings and mostly only work for specific algorithms. For example, IGA (Section 6.4.1) provably converges to the average reward of a Nash equilibrium as per Equation 5.6, and WoLF-IGA (Section 6.4.3) provably converges to a Nash equilibrium as per Equation 5.3; however, both results are limited to normal-form games with only two agents and two actions. Designing general, scalable MARL algorithms that have useful learning guarantees is very difficult and the subject of ongoing research.

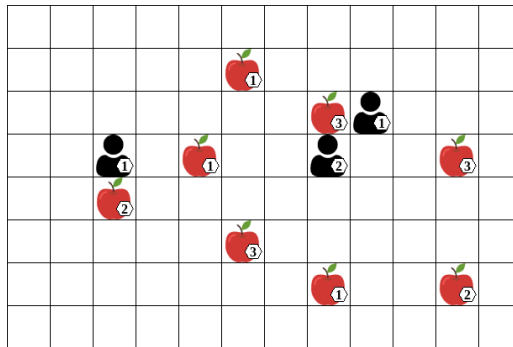


Figure 5.5: A level-based foraging task in which a group of three robots, or agents, must collect all items (shown as apples). Each agent and item has an associated skill level, shown inset. A group of one or more agents can collect an item if they are located next to the item, and the sum of the agents' levels is greater than or equal to the item's level.

5.4.2 Multi-Agent Credit Assignment

Credit assignment in RL is the problem of determining which past actions have contributed to received rewards. In multi-agent RL, there is the additional question of *whose* actions among multiple agents contributed to the rewards. To distinguish the two problems, the first type is usually referred to as temporal credit assignment while the second type is referred to as multi-agent credit assignment.

To illustrate multi-agent credit assignment and how it complicates learning, consider the level-based foraging example introduced in Chapter 1, and shown again in Figure 5.5 for convenience. In this situation, assume that all three agents attempt the “collect” action, and that as a result they all receive a reward of +1. Whose actions led to this reward? To us, it is clear that the action of the agent on the left did not contribute to this reward since it failed to collect the item (the agent’s level was not large enough). Moreover, we know that the *combined* actions of the two agents on the right led to the reward, rather than any individual agent’s action. However, for a learning agent which only observes the environment state (before and after) the agents’ chosen actions, and the received collective reward of +1, disentangling the action contributions in such detail can be highly non-trivial. It requires a detailed understanding of the world dynamics, specifically the relationship between agent/item locations and levels and how the collect action depends on these.

Note that the above example considers multi-agent credit assignment in one specific time instance. Over time, this is further compounded by temporal credit assignment. Agents must also learn to give appropriate credit to their own past actions and those of other agents. For example, not only did the collect action of the agent on the left not contribute to the +1 reward, but neither did its previous move actions that brought it to its current location.

To resolve multi-agent credit assignment, an agent needs to understand the effect of its own actions on the received reward versus the effect of the other agents' actions. Ascribing values to *joint actions*, as is done in central learning, is a useful way to disentangle each agent's contribution to a received reward. As a simple example, consider the Rock-Paper-Scissors matrix game from Section 3.1. During training, suppose the two agents choose actions $(a_1, a_2) = (R, S)$ and agent 1 receives a reward of +1 (because rock beats scissors), then the agents choose actions $(a_1, a_2) = (R, P)$ and agent 1 receives a reward of -1 (because paper beats rock). If agent 1 uses an action value model $Q(s, a_1)$ which ascribes values to its own actions (as in independent Q-learning), then the average value of taking action R may appear to be 0 since Q does not explicitly model the impact of agent 2's action. In contrast, a joint action value model $Q_1(s, a_1, a_2)$ (as in central Q-learning) can correctly represent the impact of agent 2's action, by ascribing different values to joint actions (R, S) and (R, P) . The class of *joint action learning* algorithms, introduced in Chapter 6, uses such joint action values to learn solutions for games.

Joint action value models also enable an agent to consider counterfactual questions such as "What reward would I have received if agent j had done action X instead of Y?". *Difference rewards* (Wolpert and Tumer 2002; Tumer and Agogino 2007) formulate this approach, where X corresponds to doing nothing or a "default action". Unfortunately, it is generally unclear whether such a default action exists for a given environment and what it should be. Other approaches attempt to learn a decomposition of the value function corresponding to the agents' individual contributions to a collective reward (e.g. Sunehag et al. 2017; Rashid et al. 2018; Son et al. 2019; Zhou, Liu, et al. 2020). Some of these methods will be introduced in Chapter 9.

5.4.3 Equilibrium Selection

As was highlighted in Section 4.7, a game may have multiple equilibrium solutions which may yield different expected returns to the agents in the game. The (non-repeated) Chicken game discussed in Section 4.6, shown again in Figure 5.6a, has three different equilibria which, respectively, yield expected returns of $(7, 2)$, $(2, 7)$, and $\approx(4.66, 4.66)$ to the two agents. *Equilibrium*

	S	L
S	0,0	7,2
L	2,7	6,6

(a) Chicken

	S	H
S	4,4	0,3
H	3,0	2,2

(b) Stag Hunt

Figure 5.6: Matrix games with multiple equilibria.

selection is the problem of which equilibrium the agents should agree on and how they can achieve agreement (Harsanyi and Selten 1988b).

Equilibrium selection can be a significant complicating factor for reinforcement learning in games, in which agents typically do not have prior knowledge about the game. Indeed, even if a game has a Nash equilibrium which yields maximum returns to all agents, there can be forces which make the agents prone to converging to a sub-optimal equilibrium.

Consider the Stag Hunt matrix game shown in Figure 5.6b. This game models a situation in which two hunters can either hunt a stag (S) or a hare (H). Hunting a stag requires cooperation and yields higher rewards, while hunting a hare can be done alone but gives lower rewards. It can be seen that the joint actions (S,S) and (H,H) are two Nash equilibria in the game (there is also a third probabilistic equilibrium, which we omit for clarity). While (S,S) yields the maximum reward to both agents and is Pareto-optimal, (H,H) has a relatively lower risk in that each agent can guarantee a reward of at least 2 by choosing H. Thus, (S,S) is the *reward-dominant* equilibrium meaning that it yields higher rewards than the other equilibria, and (H,H) is the *risk-dominant* equilibrium in that it has lower risk (agents can guarantee higher minimum reward) than the other equilibria. Algorithms such as independent Q-learning can be prone to converging to a risk-dominant equilibrium if they are uncertain about the actions of other agents. In the Stag Hunt example, in the early stages of learning when agents choose actions more randomly, each agent will quickly learn that action S can give a reward of 0 while action H gives a reward of 2 or higher. This can steer the agents to assign greater probability to action H, which reinforces the risk-dominant equilibrium in a feedback loop since deviating from H is penalised if the other agent chooses H.

Various approaches can be considered to tackle equilibrium selection. One approach is to refine the space of solutions by requiring additional criteria, such as Pareto optimality and welfare/fairness optimality. As we saw in the example discussed in Section 4.9, in some cases this may reduce an infinite space of solutions to a single unique solution. In some types of games, the structure of the game can be exploited for equilibrium selection. For instance,

minimax Q-learning (Section 6.2.1) benefits from the uniqueness of equilibrium values in zero-sum games, meaning that all minimax solutions give the same expected joint returns to agents. In no-conflict games such as Stag Hunt, in which there always exists a Pareto-optimal reward-dominant equilibrium, the Pareto Actor-Critic algorithm (Chapter 9) uses the fact that all agents know that the reward-dominant equilibrium is also preferred by all the other agents. Agent modelling (Section 6.3) can also help with equilibrium selection by making predictions about the actions of other agents. In the Stag Hunt game, if agent 1 expects that agent 2 is likely to choose action S if agent 1 chose S in past time steps, then agent 1 may learn to choose S more frequently and the process could converge to both agents choosing (S,S). However, whether such an outcome occurs depends on several factors, such as the details of the agents' exploration strategies and learning rates, and how the agent model is used by the agent.

If the agents are able to communicate with each other, then they may send messages about their preferred outcomes or future actions to each other, which could be used to agree on a particular equilibrium. However, communication can bring its own challenges if the agents are not bound to the information they communicate (such as deviating from their announced actions) and other agents cannot verify the received information. Moreover, if different equilibria yield different returns to the agents, then there may still be inherent conflicts about which equilibrium is most preferred.

5.4.4 Scaling to Many Agents

The ability to scale efficiently to many agents is an important goal in MARL research. This goal is significantly complicated by the fact that the number of joint actions can grow exponentially in the number of agents, since we have

$$|A| = |A_1| * \dots * |A_n|. \quad (5.9)$$

In the level-based foraging example discussed in Section 5.4.2, if we change the number of agents from 3 to 5, we also increase the number of joint actions from 216 to 7776. Moreover, if the agents have their own associated features in the state s , as they do in level-based foraging (agent positions), then the number of states $|S|$ also increases exponentially in the number of agents.

MARL algorithms can be affected in various ways by this exponential growth. For example, algorithms that use joint action values $Q(s, a)$, $a \in A$, such as central Q-learning and joint action learning (Section 6.2), are faced with an exponential growth of the space required to represent Q as well as the number of observations required to fill Q . Algorithms which do not use joint action values, such as independent learning (Section 5.3.2), can also be significantly affected by additional agents. In particular, a larger number of agents can increase

the degree of non-stationarity caused by multiple learning agents, since each additional agent adds another moving part that the other agents must adapt to. Multi-agent credit assignment can also become more difficult with more agents, since each additional agent adds a potential cause for an observed reward.

We stated at the beginning that the number of joint actions *can* grow exponentially. In fact, it is important to note that this exponential growth may not always exist, in particular when using the decomposition approach of multi-agent systems discussed in Section 1.2. As an example, suppose we want to control a power plant which has 1000 control variables, and each variable can take one of k possible values. Thus, an action is a vector of length 1000, and there are k^{1000} possible actions (value assignments). To make this problem more tractable, we could factor the action vector into n smaller vectors and use n agents, one for each of the smaller action vectors. Each agent now deals with a smaller action space, for example $|A_i| = k^{\frac{1000}{n}}$ if the factored action vectors have the same length. However, note that the total number of joint actions between agents, $|A| = |A_1| * \dots * |A_n| = k^{1000}$, is independent of the number of agents n .

Indeed, while exponential growth due to the number of agents is an important challenge in MARL, it is not unique to MARL. Single-agent reductions of the problem, such as central learning, are still faced with an exponential growth in the number of actions. Moreover, other approaches to optimal decision making in multi-agent systems, such as model-based multi-agent planning (e.g. Oliehoek and Amato 2016), also have to handle the exponential growth. Still, scaling efficiently with the number of agents is an important goal in MARL research. Part II of this book will introduce deep learning techniques as one way to improve the scalability of MARL algorithms.

5.5 What Algorithms Do Agents Use?

MARL approaches such as independent learning open up the possibility that agents may use different learning algorithms. Two basic modes of operation in MARL are self-play and mixed-play. In self-play, all agents use the same learning algorithm, or even the same policy. In mixed-play, agents use different learning algorithms. We will discuss these approaches in turn in this section.

5.5.1 Self-Play

The term *self-play* has been used to describe two related but distinct modes of operation in MARL. The first definition of self-play simply refers to the assumption that all agents use the same learning algorithm and the same algorithm parameters (Bowling and Veloso 2002; Banerjee and Peng 2004; Powers and

Shoham 2005; Conitzer and Sandholm 2007; Shoham, Powers, and Grenager 2007; Wunder, Littman, and Babes 2010; Chakraborty and Stone 2014). Developing algorithms that converge to some type of equilibrium solution in self-play is at the core of much of the literature in MARL. Essentially all of the MARL algorithms introduced in this book operate in this way. For independent learning, self-play is usually implicitly assumed (such as in independent Q-learning), but note that this is not strictly a requirement since the agents may use different algorithms within the independent learning approach.

This definition of self-play is rooted in game theory, specifically the literature on “interactive learning” (Fudenberg and Levine 1998; Young 2004), which studies basic learning rules for players (i.e. agents) and their ability to converge to equilibrium solutions in game models. The principal focus is the theoretical analysis of learning outcomes in the limit, under the standard assumption that all players use the same learning rule. This assumption serves as an important simplification, since the non-stationarity caused by multiple learning agents (discussed in Section 5.4.1) can be further exacerbated if the agents use different learning approaches. From a practical perspective, it is appealing to have one algorithm that can be used by all agents regardless of differences in their action and observation spaces.

Another definition of self-play, which was primarily developed in the context of zero-sum sequential games, uses a more literal interpretation of the term by training an agent’s policy directly against itself (or against previous versions of the policy). In this process, the agent learns to exploit weaknesses in its own play as well as how to eliminate such weaknesses. One of the earliest applications of this self-play approach in combination with temporal-difference learning was the algorithm TD-Gammon by Tesauro (1994), which was able to achieve champion-level performance in the game of backgammon. More recently, the AlphaZero algorithm (Silver et al. 2018) used self-play combined with deep RL techniques and was able to reach champion-level performance in diverse zero-sum sequential games, such as chess, shogi and Go. We will describe this type of self-play and AlphaZero in Section 9.7.

To distinguish these two types of self-play, the terms “algorithm self-play” and “policy self-play” could be used, respectively. Note that policy self-play implies algorithm self-play. An important benefit of policy self-play is that it may learn significantly faster than algorithm self-play, since the experiences of all agents can be combined to train a single policy. However, policy self-play is also more restricted in that it requires that the agents in the game have symmetrical roles and ego-centric observations (as we will describe in Section 9.7.2), such that the same policy can be used from the perspective of each agent. In contrast, algorithm self-play has no such restriction.

5.5.2 Mixed-Play

Mixed-play describes the case in which the agents use different learning algorithms. One example of mixed-play can be seen in trading markets, in which the agents may use different learning algorithms developed by the different users or organisations which control the agents. Ad hoc teamwork (Stone et al. 2010; Mirsky et al. 2022) is another example, in which agents must collaborate with previously unknown other agents whose behaviours may be initially unknown.

The study of Albrecht and Ramamoorthy (2012) considered such mixed-play settings and empirically compared different learning algorithms, including JAL-AM (Section 6.3.2), Nash-Q (Section 6.2), WoLF-PHC (Section 6.4.4) and others, in many different normal-form games using a range of metrics, including several of the solution concepts discussed in Chapter 4. The study concluded that there was no clear winner among the tested algorithms, each having relative strengths and limitations in mixed-play settings. While Papoudakis et al. (2021) provide a benchmark and comparison of contemporary deep learning-based MARL algorithms (such as those discussed in Chapter 9) for self-play in common-reward games, there is currently no such study for deep learning-based MARL algorithms in mixed-play settings.

MARL research has also developed algorithms which aim to bridge (algorithm) self-play and mixed-play. For example, the agenda of converging to an equilibrium solution in self-play was extended with an additional agenda to converge to a best-response policy if other agents use a stationary policy (Bowling and Veloso 2002; Banerjee and Peng 2004; Conitzer and Sandholm 2007). Algorithms based on *targeted optimality and safety* (Powers and Shoham 2004) assume that other agents are from a particular class of agents and aim to achieve best-response returns if the other agents are indeed from that class, and otherwise achieve at least maxmin (“security”) returns that can be guaranteed against any other agents. For example, we could assume that other agents use a particular policy representation, such as finite state automata or decision trees, or that their policies are conditioned on the previous x observations from the history (Powers and Shoham 2005; Vu, Powers, and Shoham 2006; Chakraborty and Stone 2014). In self-play, these algorithms aim to produce Pareto-optimal outcomes (Shoham and Leyton-Brown 2008).

6 Multi-Agent Reinforcement Learning: Foundational Algorithms

In Chapter 5, we took some first steps towards applying RL to compute solutions in games: we defined a general learning process in games and different convergence types for MARL algorithms, and we introduced the basic concepts of central learning and independent learning which apply single-agent RL in games. We then discussed several core challenges faced in MARL, including non-stationarity, multi-agent credit assignment, and equilibrium selection.

Continuing in our exploration of RL methods to compute game solutions, the present chapter will introduce several classes of foundational algorithms in MARL. These algorithms go beyond the basic central/independent learning approaches by explicitly modelling and using aspects of the multi-agent interaction. We call them *foundational* algorithms because of their basic nature, and because each algorithm type can be instantiated in different ways. As we will see, depending on the specific instantiation used, these MARL algorithms can successfully learn or approximate (as per the different convergence types given in Section 5.2) different types of solutions in games.

Specifically, we will introduce three classes of MARL algorithms. *Joint action learning* is a class of MARL algorithms which use temporal-difference learning to learn estimates of joint action values. These algorithms can make use of game-theoretic solution concepts to compute policies and update targets for temporal-difference learning. Next we will discuss *agent modelling*, which is the task of learning explicit models of other agents to predict their actions based on their past chosen actions. We will show how joint action learning can use such agent models in combination with best-response actions to learn optimal joint policies. The third class of MARL algorithms covered in this chapter are *policy-based learning* methods, which directly learn policy parameters using gradient-ascent techniques.

To simplify the descriptions in this chapter, we will focus on normal-form games and stochastic games in which we assume full observability of environment states and actions. Part II of this book will introduce MARL algorithms

that use deep learning techniques and can be applied to the more general POSG model. Many of the basic concepts and algorithm categories discussed in this chapter still feature in these deep learning-based MARL algorithms.

6.1 Dynamic Programming for Games: Value Iteration

In his seminal work on stochastic games, Shapley (1953) described an iterative procedure to compute the optimal expected returns (or “values”) $V_i^*(s)$ for each agent i and state s , in zero-sum stochastic games with two agents. This method is analogous to the classical value iteration algorithm for MDPs (Section 2.5), and forms the foundation for a family of temporal-difference learning algorithms discussed in Sections 6.2 and 6.3.

Algorithm 6 shows the pseudocode of the value iteration algorithm for stochastic games. The algorithm requires access to the reward functions \mathcal{R}_i and state transition function \mathcal{T} of the game, as is the case in MDP value iteration. The algorithm starts by initialising functions $V_i(s)$, for each agent i , which associate a value to each possible state of the game. In the pseudocode we initialise V_i to 0, but arbitrary initial values may be used. The algorithm then makes two sweeps over the entire state space S :

1. The first sweep computes for each agent $i \in I$ and state $s \in S$ a matrix⁴⁰ $M_{i,s}$ which contains entries for each joint action $a \in A$. This matrix can be viewed as a reward function for agent i in a normal-form game associated with the state s , i.e. $\mathcal{R}_i(a) = M_{i,s}(a)$.
2. The second sweep updates each agent’s value function $V_i(s)$ for each state s , by using the expected return to agent i under the minimax solution (defined in Equation 4.10 on page 62) of the non-repeated normal-form game given by $M_{1,s}, \dots, M_{n,s}$. We denote the minimax value for agent i by $Value_i(M_{1,s}, \dots, M_{n,s})$. This minimax value is unique and can be computed efficiently as a linear programme, as shown in Section 4.3.1.

The sweeps are repeated and the process converges to the optimal value functions V_i^* for each agent i , which satisfy

$$V_i^*(s) = Value_i(M_{1,s}^*, \dots, M_{n,s}^*) \quad (6.1)$$

$$M_{i,s}^* = \sum_{s' \in S} \mathcal{T}(s' | s, a) [\mathcal{R}_i(s, a, s') + \gamma V_i^*(s')] . \quad (6.2)$$

Given value functions V_i^* for each $i \in I$, the corresponding minimax policies of the stochastic game for each agent, $\pi_i(a_i | s)$, are obtained by computing the

40. Technically, $M_{i,s}$ is a *hypermatrix* if the game has more than two agents.

Algorithm 6 Value Iteration for Stochastic Games

- 1: Initialise: $V_i(s) = 0$ for all $s \in S$ and $i \in I$
- 2: Repeat until all V_i have converged:
- 3: **for all** agents $i \in I$, states $s \in S$, joint actions $a \in A$ **do**
- 4: Compute matrix:

$$M_{i,s}(a) = \sum_{s' \in S} \mathcal{T}(s' | s, a) [\mathcal{R}_i(s, a, s') + \gamma V_i(s')] \quad (6.3)$$

- 5: **for all** agents $i \in I$, states $s \in S$ **do**
- 6: Update V_i :

$$V_i(s) \leftarrow \text{Value}_i(M_{1,s}, \dots, M_{n,s}) \quad (6.4)$$

minimax solution in the non-repeated normal-form game given by $M_{1,s}^*, \dots, M_{n,s}^*$, for each state $s \in S$. Notice that, analogous to optimal policies in MDPs, the minimax policies in the stochastic game are conditioned only on the state rather than state-action histories.

Notice that Equation 6.4 resembles the value update in MDP value iteration defined in Equation 2.46 and repeated below,

$$V(s) \leftarrow \max_{a \in A} \sum_{s' \in S} \mathcal{T}(s' | s, a) [\mathcal{R}(s, a, s') + \gamma V(s')], \quad (6.5)$$

but replaces the \max_a -operator with the Value_i -operator. In fact, in stochastic games with a single agent (i.e. MDPs), the value iteration algorithm discussed here reduces to MDP value iteration.⁴¹ This can be seen as follows: First, in stochastic games with a single agent i , the value update becomes $V_i(s) \leftarrow \text{Value}_i(M_{i,s})$ where $M_{i,s}$ is a vector of action values for agent i in state s . Next, recall the definition of the maxmin value from Equation 4.10 (page 62), which simply becomes $\max_{\pi_i} U_i(\pi_i)$ in the single-agent case. In the single-agent case (i.e. MDPs), we know that there is always a deterministic optimal policy which in each state chooses an optimal action with probability 1. Using all of the aforementioned, the definition of Value_i becomes

$$\text{Value}_i(M_{i,s}) = \max_{a_i \in A_i} M_{i,s}(a_i) \quad (6.6)$$

$$= \max_{a_i \in A_i} \sum_{s' \in S} \mathcal{T}(s' | s, a_i) [\mathcal{R}(s, a_i, s') + \gamma V(s')], \quad (6.7)$$

41. Hence, despite the focus on stochastic games, Shapley (1953) can also be regarded as an early inventor of value iteration for MDPs.

and, thus, Equation 6.4 reduces to MDP value iteration.

To see why the value iteration process in stochastic games converges to the optimal values V_i^* , one can show that the update operator defined in Equation 6.4 is a *contraction mapping*. We described the proof for MDP value iteration in Section 2.5, and the proof for value iteration in stochastic games follows an analogous argument which we summarise here. A mapping $f: \mathcal{X} \mapsto \mathcal{X}$ on a $\|\cdot\|$ -normed complete vector space \mathcal{X} is a γ -contraction, for $\gamma \in [0, 1)$, if for all $x, y \in \mathcal{X}$:

$$\|f(x) - f(y)\| \leq \gamma \|x - y\| \quad (6.8)$$

By the Banach fixed-point theorem, if f is a contraction mapping, then for any initial vector $x \in \mathcal{X}$ the sequence $f(x), f(f(x)), f(f(f(x))), \dots$ converges to a unique fixed point $x^* \in \mathcal{X}$ such that $f(x^*) = x^*$. Using the max-norm $\|x\|_\infty = \max_i |x_i|$, it can be shown that Equation 6.4 satisfies Equation 6.8 (Shapley 1953). Thus, repeated application of the update operator in Equation 6.4 converges to a unique fixed point which, by definition, is given by V_i^* for all agents $i \in I$.

6.2 Temporal-Difference Learning for Games: Joint Action Learning

While the value iteration algorithm introduced in the previous section has useful learning guarantees, it requires access to the game model (in particular, the reward functions R_i and state transition function T), which may not be available. Thus, the question arises whether it is possible to learn solutions to games via a process of repeated interaction between the agents, using ideas based on temporal-difference learning in RL (Section 2.6).

Independent learning algorithms (such as Independent Q-Learning) can use temporal-differencing, but because they ignore the special structure of games – in particular, that the state is affected by actions of multiple agents – they suffer from non-stationarity and multi-agent credit assignment problems. On the other hand, central learning algorithms (such as Central Q-Learning) address these issues by learning values of joint actions, but suffer from other limitations such as the need for reward scalarisation.

Joint action learning (JAL) refers to a family of MARL algorithms based on temporal-difference learning which seek to address the above problems. As the name suggests, JAL algorithms learn *joint action value models* that estimate the expected returns of joint actions in any given state. Analogous to the Bellman equation for MDPs (Section 2.4), in a stochastic game the expected return to agent i when the agents select joint action $a = (a_1, \dots, a_n)$ in state s and

subsequently follow joint policy π is given by

$$Q_i^\pi(s, a) = \sum_{s' \in S} \mathcal{T}(s' | s, a) \left[\mathcal{R}_i(s, a, s') + \gamma \sum_{a' \in A} \pi(a' | s') Q_i^\pi(s', a') \right]. \quad (6.9)$$

Specifically, the JAL algorithms we present in this section are all off-policy algorithms which aim to learn *equilibrium* Q-values, $Q_i^{\pi^*}$, where π^* is an equilibrium joint policy for the stochastic game. To reduce notation, we will drop the π from Q_i^π unless required.

In contrast to using single-agent Q-values $Q(s, a)$, using joint action values $Q_i(s, a_1, \dots, a_n)$ alone is no longer enough to select the best action for agent i in a given state, i.e. finding $\max_{a_i} Q_i(s, a_1, \dots, a_n)$, since it depends on the actions of the other agents in that state. Moreover, since a game can have multiple equilibria which may yield different expected returns to the agents, learning $Q_i^{\pi^*}$ requires some way to agree on a particular equilibrium (we discussed this *equilibrium selection* problem in Section 5.4.3). Therefore, using joint action values requires additional information or assumptions about the actions of other agents in order to select optimal actions and to compute target values for temporal-difference learning. In this section, we will discuss a class of JAL algorithms that use solution concepts from game theory to derive this additional information; hence, we will refer to them as JAL-GT.

The underlying idea in JAL-GT algorithms is that the set of joint action values $Q_1(s, \cdot), \dots, Q_n(s, \cdot)$ can be viewed as a non-repeated normal-form game Γ_s for state s , in which the reward function for agent i is given by

$$\mathcal{R}_i(a_1, \dots, a_n) = Q_i(s, a_1, \dots, a_n). \quad (6.10)$$

As convenient notation, we may also write this as $\Gamma_{s,i}(a)$ for joint actions a . Visually, for a stochastic game with two agents (i and j) and three possible actions for each agent, the normal-form game $\Gamma_s = \{\Gamma_{s,i}, \Gamma_{s,j}\}$ in state s can be written as

$$\Gamma_{s,i} = \begin{array}{|ccc|} \hline & Q_i(s, a_{i,1}, a_{j,1}) & Q_i(s, a_{i,1}, a_{j,2}) & Q_i(s, a_{i,1}, a_{j,3}) \\ \hline Q_i(s, a_{i,2}, a_{j,1}) & Q_i(s, a_{i,2}, a_{j,2}) & Q_i(s, a_{i,2}, a_{j,3}) \\ Q_i(s, a_{i,3}, a_{j,1}) & Q_i(s, a_{i,3}, a_{j,2}) & Q_i(s, a_{i,3}, a_{j,3}) \\ \hline \end{array}$$

with an analogous matrix $\Gamma_{s,j}$ for agent j using Q_j . The notation $a_{i,k}$ means that agent i uses action number k , and similarly for $a_{j,k}$ and agent j .

We can solve the normal-form game Γ_s using existing game-theoretic solution concepts, such as minimax or Nash equilibrium, to obtain an equilibrium joint policy π_s^* . Note that π_s^* is a joint policy for a *non-repeated normal-form game*

Algorithm 7 Joint Action Learning with Game Theory (JAL-GT)

```

// Algorithm controls agent  $i$ 
1: Initialise:  $Q_j(s, a) = 0$  for all  $j \in I$  and  $s \in S, a \in A$ 
2: Repeat for every episode:
3: for  $t = 0, 1, 2, \dots$  do
4:   Observe current state  $s^t$ 
5:   With probability  $\epsilon$ : choose random action  $a_i^t$ 
6:   Else: solve  $\Gamma_{s^t}$  to get policies  $(\pi_1, \dots, \pi_n)$ , then sample action  $a_i^t \sim \pi_i$ 
7:   Observe joint action  $a^t = (a_1^t, \dots, a_n^t)$ , rewards  $r_1^t, \dots, r_n^t$ , next state  $s^{t+1}$ 
8:   for all  $j \in I$  do
9:      $Q_j(s^t, a^t) \leftarrow Q_j(s^t, a^t) + \alpha [r_j^t + \gamma Value_j(\Gamma_{s^{t+1}}) - Q_j(s^t, a^t)]$ 

```

(Γ_s) , while π^* is the equilibrium joint policy we seek to learn for the stochastic game. Given π_s^* , action selection in state s of the stochastic game is simply done by sampling $a \sim \pi_s^*$ with some random exploration (e.g. ϵ -greedy). Update targets for temporal-difference learning of $Q_i^{\pi^*}$ are derived by using agent i 's expected return (or value) of the solution to the game $\Gamma_{s'}$ for the next state s' , given by

$$Value_i(\Gamma_{s'}) = \sum_{a \in A} \Gamma_{s', i}(a) \pi_{s'}^*(a) \quad (6.11)$$

where $\pi_{s'}^*$ is the equilibrium joint policy for the normal-form game $\Gamma_{s'}$.

The pseudocode for a general JAL-GT algorithm for stochastic games is shown in Algorithm 7. The algorithm observes the actions and rewards of all agents in each time step, and maintains joint action value models Q_j for every agent $j \in I$, in order to produce the games Γ_s . Several well-known MARL algorithms can be instantiated from Algorithm 7, which differ in the specific solution concept used to solve Γ_s . These algorithms can learn equilibrium values for the stochastic game under certain conditions, as we will see in the following subsections.

6.2.1 Minimax Q-Learning

Minimax Q-learning (Littman 1994) is based on Algorithm 7 and solves Γ_s by computing a minimax solution, for example via linear programming (Section 4.3.1). This algorithm can be applied to two-agent zero-sum stochastic games. Minimax Q-learning is guaranteed to learn the unique minimax value of the stochastic game under the assumption that all combinations of states and joint actions are tried infinitely often, as well as the usual conditions on learning rates used in single-agent RL (Littman and Szepesvári 1996). This algorithm

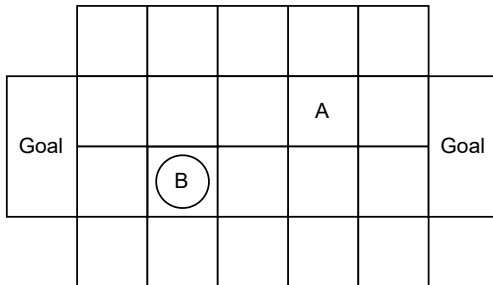


Figure 6.1: Simplified grid-world soccer game with two agents (A and B). The circle marks ball possession. Based on Littman (1994).

is essentially the temporal-difference version of the value iteration algorithm discussed in Section 6.1.

To see the differences in the policies learned by minimax Q-learning compared to independent Q-learning (Algorithm 5), it is instructive to look at the experiments of the original paper which proposed minimax Q-learning (Littman 1994). Both algorithms were evaluated in a simplified soccer game represented as a zero-sum stochastic game with two agents. The game is played on a 4 by 5 grid in which each episode (i.e. match) starts in the initial state shown in Figure 6.1, with the ball randomly assigned to one agent. Each agent can move up, down, left, right, or stand still, and the agents' selected actions are executed in random order. If the agent with the ball attempts to move into the location of the other agent, it loses the ball to the other agent. An agent scores a reward of +1 if it moves the ball into the opponent goal (and the opponent gets a reward of -1), after which a new episode starts. Each episode terminates as a draw (reward of 0 for both agents) with probability 0.1 in each time step, hence the discount factor was set to $\gamma = 0.9$ ($1 - \gamma$ is the probability of terminating in each time step). Both algorithms learned over one million time steps, used exploration rate $\epsilon = 0.2$, and an initial learning rate $\alpha = 1.0$ which was reduced in each time step by multiplying with 0.9999954. After training, one of the learned policies from each algorithm was tested over 100,000 time steps against 1) a random opponent which picks actions uniformly randomly; 2) a deterministic hand-built opponent which uses heuristic rules for attacking and defending; and 3) an optimal (i.e worst-case) opponent which used Q-learning to train an optimal policy against the other agent's fixed policy.

	minimax Q		independent Q	
	% won	ep. len.	% won	ep. len.
vs. random	99.3	13.89	99.5	11.63
vs. hand-built	53.7	18.87	76.3	30.30
vs. optimal	37.5	22.73	0	83.33

Figure 6.2: Percentage of won episodes and average episode length (time steps) in the simplified soccer game, when policies learned by minimax Q-learning and independent Q-learning are tested against random, hand-built, and optimal opponents. Based on Littman (1994).

Figure 6.2 shows the percentage of won episodes and average episode length⁴² for policies learned by minimax Q-learning and independent Q-learning when tested against the random, hand-built, and optimal opponents. Against the random opponent, the policies of both algorithms won in almost all games and achieved similar episode lengths. Against the hand-built opponent, minimax Q-learning achieved a win rate of 53.7% which is close to the theoretical 50% of the exact minimax solution. Independent Q-learning achieved a higher win rate of 76.3% because its learned policy was able to exploit certain weaknesses in the hand-built opponent. On the other hand, against the optimal opponent, minimax Q-learning achieved a win rate of 37.5%. The fact that this win rate was a little further from the theoretical 50% suggests that the algorithm had not fully converged during learning, resulting in weaknesses in the policy which the optimal opponent was able to exploit in some situations. The policy learned by independent Q-learning lost in all episodes against its optimal opponent. This is because any fully-deterministic policy can be exploited by an optimal opponent. These results show that the policies learned by minimax Q-learning do not necessarily exploit weaknesses in opponents when they exist, but at the same time these policies are robust to exploitation by assuming a worst-case opponent during training.

6.2.2 Nash Q-Learning

Nash Q-learning (Hu and Wellman 2003) is based on Algorithm 7 and solves Γ_s by computing a Nash equilibrium solution. This algorithm can be applied to the general class of general-sum stochastic games with any finite number of agents.

42. The original work (Littman 1994) showed the number of completed episodes within the 100,000 time steps in testing. We find the average episode length (100,000 divided by number of completed games) more intuitive to interpret.

Nash Q-learning is guaranteed to learn a Nash equilibrium of the stochastic game, albeit under highly restrictive assumptions. In addition to trying all combinations of states and joint actions infinitely often, the algorithm requires that all of the encountered normal-form games Γ_s either a) all have a global optimum or b) all have a saddle point. A joint policy π is a global optimum in Γ_s if each agent individually achieves its maximum possible expected return, i.e. $\forall i, \pi' : U_i(\pi) \geq U_i(\pi')$. Thus, a global optimum is also an equilibrium since no agent can deviate to obtain higher returns. A joint policy π is a saddle point if it is an equilibrium *and* if any agent deviates from π , then all other agents receive a higher expected return.

Either of these conditions is unlikely to exist in any encountered normal-form game, let alone in *all* of the encountered games. For example, global optimality is significantly stronger than Pareto optimality (Section 4.8), which merely requires that there is no other joint policy that increases the expected return of at least one agent without making other agents worse off. As an example, in the Prisoner’s Dilemma matrix game (Figure 3.2c), agent 1 defecting and agent 2 cooperating is Pareto optimal since any other joint policy will reduce the expected return for agent 1. However, there is no global optimum since no single joint policy can achieve the maximum possible reward for both agents at the same time.

To see why these assumptions are required, notice that all global optima are equivalent to each agent, meaning that they give the same expected return to each agent. Thus, choosing a global optimum to compute $Value_i$ circumvents the equilibrium selection problem mentioned earlier. Similarly, it can be shown that all saddle points of a game are equivalent in the expected returns they give to agents, and so choosing any saddle point to compute $Value_i$ circumvents the equilibrium selection problem. However, an implicit additional requirement here is that all agents consistently choose either global optima or saddle points when computing target values.

6.2.3 Correlated Q-Learning

Correlated Q-learning (Greenwald and Hall 2003) is based on Algorithm 7 and solves Γ_s by computing a correlated equilibrium. Like Nash Q-learning, correlated Q-learning can be applied to general-sum stochastic games with a finite number of agents.

This algorithm has two important benefits:

1. Correlated equilibrium spans a wider space of solutions than Nash equilibrium, including solutions with potentially higher expected returns to agents (see discussion in Section 4.6).

2. Correlated equilibria can be computed efficiently for normal-form games via linear programming (Section 4.6.1), while computing Nash equilibria requires quadratic programming.

Since correlated Q-learning solves Γ_s via correlated equilibrium, we require a small modification to Algorithm 7: in Line 6 of the algorithm, it may not be possible to factorise a correlated equilibrium π into individual agent policies π_1, \dots, π_n , and hence there would be no policy π_i to sample an action a_i^t from. Instead, agent i can sample a joint action $a^t \sim \pi$ from the correlated equilibrium, and then simply take its own action a_i^t from this joint action. To maintain correlations between the agents' actions, a further modification would be to include a central mechanism which samples a joint action from the equilibrium and sends to each agent its own action from the joint action.

Since the space of correlated equilibria can be larger than the space of Nash equilibria in a game, the problem of equilibrium selection – how the agents should select a common equilibrium in Γ_s – becomes even more pronounced. Greenwald and Hall (2003) define different mechanisms to select equilibria, which ensure that there is a unique equilibrium value $Value_j(\Gamma_s)$. One such mechanism is to choose an equilibrium which maximises the sum of expected rewards of the agents, and this is what we used in the linear programme presented in Section 4.6.1. Other mechanisms include selecting an equilibrium which maximises the minimum or maximum of the agents' expected rewards. In general, however, no formal conditions are known under which correlated Q-learning converges to a correlated equilibrium of the stochastic game.

6.2.4 Limitations of Joint Action Learning

We saw that Nash Q-learning requires some very restrictive assumptions to ensure convergence to a Nash equilibrium in general-sum stochastic games, while correlated Q-learning has no known convergence guarantees. This begs the question: is it possible to construct a joint action learning algorithm based on Algorithm 7 which converges to an equilibrium solution in *any* general-sum stochastic game? As it turns out, there exist stochastic games for which the information contained in the joint action value models $Q_j(s, a)$ is insufficient to reconstruct equilibrium policies.

Consider the following two properties:

- Since Q_j is conditioned on the state s (rather than the history of states and joint actions), any equilibrium joint policy π for the stochastic game which is derived from Q_j is also only conditioned on s to choose actions. Such an equilibrium is called *stationary*.

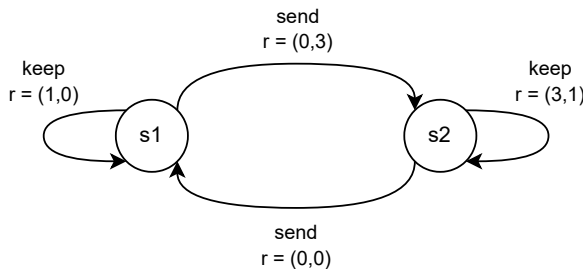


Figure 6.3: NoSDE (“No Stationary Deterministic Equilibrium”) game with two agents, two states $\{s_1, s_2\}$, and two actions $\{\text{send}, \text{keep}\}$. Agent 1 chooses an action in state s_1 and agent 2 chooses an action in state s_2 . State transitions are deterministic and shown via the arrows, with the resulting joint reward for the agents. The discount factor is $\gamma = \frac{3}{4}$.

- If in a given state s only one agent has a choice to make, i.e. $|A_{i,s}| > 1$ for some agent i and $|A_{j,s}| = 1$ for all other agents $j \neq i$, where $A_{i,s}$ is the set of available actions to agent i in state s ; then any equilibrium concept (such as Nash equilibrium and correlated equilibrium) will reduce to a max-operator when applied to Q_j in s , i.e. $\max_a Q_i(s, a)$. This is by virtue of the fact that the best response of the other agents $j \neq i$ is trivially their only available action, and hence the best response for agent i is to deterministically choose the action with the highest reward to agent i . (For simplicity, assume that any ties between actions are resolved by choosing one action with probability 1.)

In a “turn-taking” stochastic game, where in every state only one agent has a choice, the above two properties mean that any JAL-GT algorithm would attempt to learn a stationary deterministic equilibrium (joint policy). Unfortunately, there exist turn-taking stochastic games which have a unique stationary *probabilistic* equilibrium, but no stationary deterministic equilibrium.

Zinkevich, Greenwald, and Littman (2005) provide an example of such a game, shown in Figure 6.3. In this game, there are two agents, two states, and two actions available to whichever agent’s turn it is. All state transitions in this game are deterministic. It can be seen that none of the four possible combinations of deterministic policies between the agents constitute an equilibrium, because each agent always has an incentive to deviate to increase its returns. For example, given the deterministic joint policy $\pi_1(\text{send} \mid s_1) = 1, \pi_2(\text{send} \mid s_2) = 1$, agent 1 would be better off choosing the keep action in state s_1 . Similarly, given joint policy $\pi_1(\text{keep} \mid s_1) = 1, \pi_2(\text{send} \mid s_2) = 1$, agent 2 would be better off

choosing keep. In fact, this game has a unique probabilistic stationary equilibrium which is $\pi_1(\text{send} \mid s1) = \frac{2}{3}$, $\pi_2(\text{send} \mid s2) = \frac{5}{12}$. Zinkevich, Greenwald, and Littman (2005) refer to this class of games as NoSDE games (which stands for “No Stationary Deterministic Equilibrium”) and proved the following theorem:

Theorem *Let $Q_i^{\pi, \Gamma}/V_i^{\pi, \Gamma}$ denote the Q_i^π/V_i^π functions in game Γ . For any NoSDE game Γ with a unique equilibrium joint policy π , there exists another NoSDE game Γ' which differs from Γ only in the reward functions and has its own unique equilibrium joint policy $\pi' \neq \pi$, such that:*

$$\forall i : Q_i^{\pi, \Gamma} = Q_i^{\pi', \Gamma'} \quad \text{and} \quad \exists i : V_i^{\pi, \Gamma} \neq V_i^{\pi', \Gamma'}. \quad (6.12)$$

This result establishes that the joint action value models learned by JAL-GT algorithms may not carry sufficient information to derive the correct equilibrium joint policy in a game. However, while JAL-GT cannot learn the unique probabilistic stationary equilibrium in a NoSDE game, algorithms based on value iteration for games (such as JAL-GT) can converge to a cyclic sequence of actions which constitutes a “cyclic equilibrium” in NoSDE games (Zinkevich, Greenwald, and Littman 2005).

6.3 Agent Modelling

The game-theoretic solution concepts used in JAL-GT algorithms are *normative*, in that they prescribe how agents *should* behave in equilibrium. But what if some of the agents deviate from this norm? For example, by relying on the minimax solution concept, minimax Q-learning (Section 6.2.1) assumes that the other agent is an optimal worst-case opponent, and so it learns to play against such an opponent – independently of the actual chosen actions of the opponent. We saw in the soccer example in Figure 6.2 that this hard-wired assumption can limit the algorithm’s achievable performance: minimax-Q learning learned a generally robust minimax policy, but would fail to exploit the weaknesses of the hand-built opponent if it was trained against such an opponent, whereas independent Q-learning was able to outperform minimax Q-learning against the hand-built opponent.

An alternative to making implicit normative assumptions about the behaviour of other agents is to directly model the actions of other agents based on their observed behaviours. *Agent modelling* (also known as opponent modelling⁴³)

43. The term “opponent modelling” was originally used because much of the early research in this area focused on competitive games, such as chess. We prefer the more neutral term “agent modelling” since other agents may not necessarily be opponents.



Figure 6.4: An agent model makes predictions about another agent (such as the action probabilities, goals, and beliefs of the agent) based on past observations about the agent.

is concerned with constructing models of other agents which can make useful predictions about their behaviours. A general agent model is shown in Figure 6.4. For example, an agent model may make predictions about the actions of the modelled agent, or about the long-term goal of the agent such as wanting to reach a certain goal location. In a partially observable environment, an agent model may attempt to infer the beliefs of the modelled agent about the state of the environment. To make such predictions, an agent model may use various sources of information as input, such as the history of states and joint actions, or any subset thereof. Agent modelling has a long history in artificial intelligence research and, accordingly, many different methodologies have been developed (Albrecht and Stone 2018).⁴⁴

The most common type of agent modelling used in MARL is called *policy reconstruction*. Policy reconstruction aims to learn models $\hat{\pi}_j$ of the policies π_j of other agents based on their observed past actions. In general, learning the parameters of $\hat{\pi}_j$ can be framed as a supervised learning problem using the past observed state-action pairs $\{(s^\tau, a_j^\tau)\}_{\tau=1}^t$ of the modelled agent. Thus, we may choose a parameterised representation for $\hat{\pi}_j$ and fit its parameters using the (s^τ, a_j^τ) -data. Various possible representations for $\hat{\pi}_j$ could be used, such as look-up tables, finite state automata, and neural networks. The chosen model representation should ideally allow for iterative updating to be compatible with the iterative nature of RL algorithms.

Given a set of models for the other agents' policies, $\hat{\pi}_{-i} = \{\hat{\pi}_j\}_{j \neq i}$, the modelling agent i can select a best-response policy (Section 4.2) with respect to these models,

$$\pi_i \in \text{BR}_i(\hat{\pi}_{-i}). \quad (6.13)$$

Thus, while in Chapter 4 we used the concept of best-response policies as a compact characterisation of solution concepts, in this section we will see how the best-response concept is operationalised in reinforcement learning.

44. The survey of Albrecht and Stone (2018) was followed by a dedicated special journal issue on agent modelling which contains further articles on the topic (Albrecht, Stone, and Wellman 2020).

In the following subsections, we will see different methods that use policy reconstruction and best responses to learn optimal policies.

6.3.1 Fictitious Play

One of the first and most basic learning algorithms defined for non-repeated normal-form games is called *fictitious play* (Brown 1951). In fictitious play, each agent i models the policy of each other agent j as a stationary probability distribution $\hat{\pi}_j$. This distribution is estimated by taking the empirical distribution of agent j 's past actions. Let $C(a_j)$ denote the number of times that agent j selected action a_j prior to the current episode. Then, $\hat{\pi}_j$ is defined as

$$\hat{\pi}_j(a_j) = \frac{C(a_j)}{\sum_{a'_j} C(a'_j)}. \quad (6.14)$$

At the start of the first episode, before any actions have been observed from agent j (i.e. $C(a_j) = 0$ for all $a_j \in A_j$), the initial model $\hat{\pi}_j$ can be the uniform distribution $\hat{\pi}_j(a_j) = \frac{1}{|A_j|}$.

In each episode, each agent i chooses a best-response action against the models $\hat{\pi}_{-i} = \{\hat{\pi}_j\}_{j \neq i}$, given by

$$\text{BR}_i(\hat{\pi}_{-i}) = \arg \max_{a_i \in A_i} \sum_{a_{-i} \in A_{-i}} \mathcal{R}_i(\langle a_i, a_{-i} \rangle) \prod_{j \neq i} \hat{\pi}_j(a_j) \quad (6.15)$$

where $A_{-i} = \times_{j \neq i} A_j$, and a_j refers to the action of agent j in a_{-i} .

It is important to note that this definition of best response gives an optimal *action*, in contrast to the more general definition (see Equation 4.9, page 61) which gives best-response *policies*. Therefore, fictitious play cannot learn equilibrium solutions which require randomisation, such as the uniform-random equilibrium for Rock-Paper-Scissors which we saw in Section 4.3. However, the empirical distribution of actions (defined in Equation 5.5) in fictitious play *can* converge to such randomised equilibria. Indeed, fictitious play has several interesting convergence properties (Fudenberg and Levine 1998):

- If the agents' actions converge, then the converged actions form a Nash equilibrium of the game.
- If in any episode the agents' actions form a Nash equilibrium, then they will remain in the equilibrium in all subsequent episodes.
- If the empirical distribution of each agent's actions converges, then the distributions converge to a Nash equilibrium of the game.
- The empirical distributions converge in several game classes, including in two-agent zero-sum games with finite action sets (Robinson 1951).

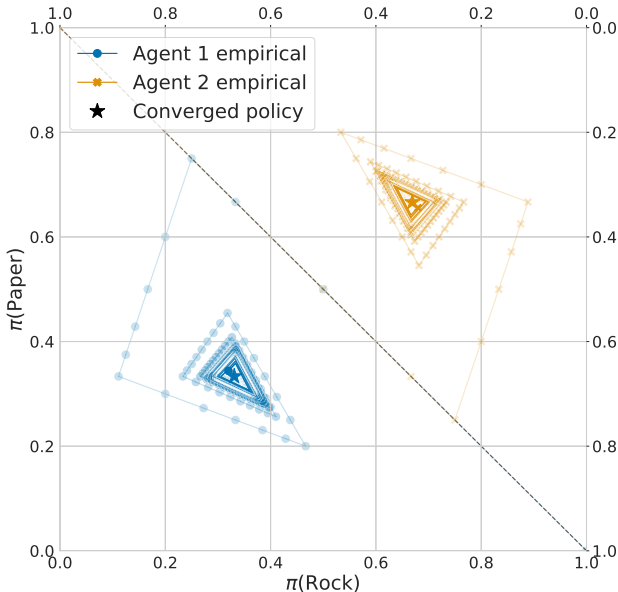


Figure 6.5: This plot shows the evolution of empirical action distributions of two agents in the Rock-Paper-Scissors game, in which both agents use fictitious play to choose actions. Each point in the simplex for an agent corresponds to a probability distribution over the three available actions. The lines show empirical action distributions over the first 500 episodes of the game. The empirical action distributions of the agents converge to the unique Nash equilibrium of the game, in which both agents choose actions uniform-randomly.

Figure 6.5 shows the evolution of the agents’ empirical action distributions defined in Equation 6.14 in the non-repeated Rock-Paper-Scissors matrix game (see Section 3.1 for a description of the game), in which both agents use fictitious play to choose actions. The first 10 episodes are shown in Figure 6.6. Both agents start with the R action and then follow identical trajectories.⁴⁵ In the next step, both agents choose the P action and the empirical action distribution gives 0.5 to R and P each, and 0 to S. The agents continue to choose the P action until episode 3, after which the optimal action for both agents with respect to their current agent models becomes S. The agents continue to choose S until the next “corner” of the trajectory, where both agents switch to choosing R,

⁴⁵. The agents could also start with different initial actions and then follow non-identical trajectories; but they would still follow a similar spiral pattern and converge to the Nash equilibrium.

Episode e	Joint action (a_1^e, a_2^e)	Agent model $\hat{\pi}_2$	Agent 1 action values
1	R,R	(0.33, 0.33, 0.33)	(0.00, 0.00, 0.00)
2	P,P	(1.00, 0.00, 0.00)	(0.00, 1.00, -1.00)
3	P,P	(0.50, 0.50, 0.00)	(-0.50, 0.50, 0.00)
4	P,P	(0.33, 0.67, 0.00)	(-0.67, 0.33, 0.33)
5	S,S	(0.25, 0.75, 0.00)	(-0.75, 0.25, 0.50)
6	S,S	(0.20, 0.60, 0.20)	(-0.40, 0.00, 0.40)
7	S,S	(0.17, 0.50, 0.33)	(-0.17, -0.17, 0.33)
8	S,S	(0.14, 0.43, 0.43)	(0.00, -0.29, 0.29)
9	S,S	(0.13, 0.38, 0.50)	(0.12, -0.38, 0.25)
10	R,R	(0.11, 0.33, 0.56)	(0.22, -0.44, 0.22)

Figure 6.6: First 10 episodes in the non-repeated Rock-Paper-Scissors game when both agents use fictitious play. The columns show the episode number e , the joint action a^e in episode e , agent 1’s model $\hat{\pi}_2$ of agent 2 (probabilities assigned to R/P/S), and agent 1’s action values for R/P/S (expected reward for each action with respect to the current model of agent 2). If multiple actions have equal maximum value, then the first action is deterministically chosen.

and so forth. As the history length increases, the changes in the empirical action distributions become smaller. The lines in Figure 6.5 show how the empirical action distributions change and co-adapt over time, converging to the only Nash equilibrium of the game which is for both agents to choose actions uniform-randomly.

A number of modifications of fictitious play have been proposed in the literature to obtain improved convergence properties and generalise the method (e.g. Fudenberg and Levine 1995; Hofbauer and Sandholm 2002; Young 2004; Leslie and Collins 2006; Heinrich, Lanctot, and Silver 2015). Note that in *repeated* normal-form games, the best response defined in Equation 6.15 is myopic, in the sense that this best response only considers a single interaction (recall that fictitious play is defined for non-repeated normal-form games). However, in repeated normal-form games, the future actions of agents can depend on the history of past actions of other agents. A more complex definition of best responses in sequential game models, as defined in Equation 4.9, also accounts for the long-term effects of actions. In the following sections, we will see how MARL can combine a best-response operator with temporal-difference learning to learn non-myopic best-response actions.

6.3.2 Joint Action Learning with Agent Modelling

Fictitious play combines policy reconstruction and best-response actions to learn optimal policies for non-repeated normal-form games. We can extend this idea to the more general case of stochastic games by using the joint action learning framework introduced in Section 6.2. This class of JAL algorithms, which we will call JAL-AM, uses the joint action values in conjunction with agent models and best responses to select actions for the learning agents and to derive update targets for the joint action values.

Analogous to fictitious play, the agent models learn empirical distributions based on the past actions of the modelled agent, albeit this time conditioned on the state in which the actions took place. Let $C(s, a_j)$ denote the number of times that agent j selected action a_j in state s , then the agent model $\hat{\pi}_j$ is defined as

$$\hat{\pi}_j(a_j | s) = \frac{C(s, a_j)}{\sum_{a'_j} C(s, a'_j)}. \quad (6.16)$$

Note that the action counts $C(s, a_j)$ include actions observed in the current episode and previous episodes. If a state s is visited for the first time, then $\hat{\pi}_j$ can assign uniform probabilities to actions. Conditioning the model $\hat{\pi}_j$ on the state s is a sensible choice, since we know that the optimal equilibrium joint policy in a stochastic game requires only the current state rather than the history of states and actions.

If the true policy π_j is fixed or converges, and is indeed conditioned only on the state s , then in the limit of observing each (s^τ, a_j^τ) -pair infinitely often, the model $\hat{\pi}_j$ will converge to π_j . Of course, during learning, π_j will not be fixed. To track changes in π_j more quickly, one option is to modify Equation 6.16 to give greater weight to more recent observed actions in state s . For example, the empirical probability $\hat{\pi}_j(a_j | s)$ may only use up to the most recent 10 observed actions of agent j in state s . Another option is to maintain a probability distribution over a space of possible agent models for agent j , and to update this distribution based on new observations via a Bayesian method (we will discuss this approach in Section 6.3.3). In general, many different approaches can be considered for constructing an agent model $\hat{\pi}_j$ from observations.

JAL-AM algorithms use the learned agent models to select actions for the learning agent and to derive update targets for the joint action values. Given agent models $\{\hat{\pi}_j\}_{j \neq i}$, the value (expected return) to agent i for taking action a_i in state s is given by

$$AV_i(s, a_i) = \sum_{a_{-i} \in A_{-i}} Q_i(s, \langle a_i, a_{-i} \rangle) \prod_{j \neq i} \hat{\pi}_j(a_j | s) \quad (6.17)$$

Algorithm 8 Joint Action Learning with Agent Modelling (JAL-AM)

```

// Algorithm controls agent i

1: Initialise:
2:    $Q_i(s, a) = 0$  for all  $s \in S, a \in A$ 
3:   Agent models  $\hat{\pi}_j(a_j | s) = \frac{1}{|A_j|}$  for all  $j \neq i, a_j \in A_j, s \in S$ 
4: Repeat for every episode:
5:   for  $t = 0, 1, 2, \dots$  do
6:     Observe current state  $s^t$ 
7:     With probability  $\epsilon$ : choose random action  $a_i^t$ 
8:     Else: choose best-response action  $a_i^t \in \arg \max_{a_i} AV_i(s^t, a_i)$ 
9:     Observe joint action  $a^t = (a_1^t, \dots, a_n^t)$ , reward  $r_i^t$ , next state  $s^{t+1}$ 
10:    for all  $j \neq i$  do
11:      Update agent model  $\hat{\pi}_j$  with new observations (e.g.  $(s^t, a_j^t)$ )
12:     $Q_i(s^t, a^t) \leftarrow Q_i(s^t, a^t) + \alpha [r_i^t + \gamma \max_{a'_i} AV_i(s^{t+1}, a'_i) - Q_i(s^t, a^t)]$ 

```

where Q_i is the joint action value model for agent i learned by the algorithm. Using the action values AV_i , the best-response action for agent i in state s is given by $\arg \max_{a_i} AV_i(s, a_i)$; and the update target in the next state s' uses $\max_{a_i} AV_i(s', a_i)$.

Algorithm 8 provides pseudocode for a general JAL-AM algorithm for stochastic games. Similar to fictitious play, this algorithm learns best-response *actions* rather than probabilistic best-response policies. Unlike JAL-GT, which requires observing the rewards of other agents and maintaining joint action value models Q_j for every agent, JAL-AM does not require observing the rewards of other agents and only maintains a single joint action value model Q_i for the learning agent. Both JAL-AM and JAL-GT are off-policy temporal-difference learning algorithms.

Figure 6.7 shows the evaluation returns of JAL-AM using agent models defined in Equation 6.16, compared to CQL and IQL in the level-based foraging task from Figure 5.2 (page 93). The learning curves for CQL and IQL are copied over from Figure 5.3. In this specific task, JAL-AM converged to the optimal joint policy faster than IQL and CQL. The agent models learned by the JAL-AM agents reduced the variance in their update targets, as is reflected by the smaller standard deviation in evaluation returns compared to IQL and CQL. This allowed JAL-AM to converge (on average) to the optimal joint policy after about 500,000 training time steps, while IQL required about 600,000 training time steps to converge to the optimal joint policy.

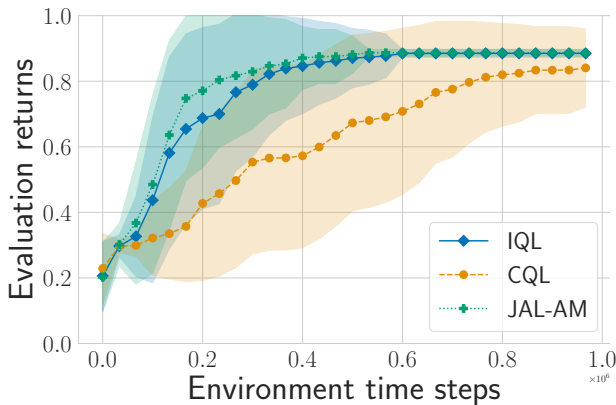


Figure 6.7: Joint action learning with agent modelling (JAL-AM) compared to central Q-learning (CQL) and independent Q-learning (IQL) in the level-based foraging task from Figure 5.2. Results are averaged over 50 independent training runs. The shaded area shows the standard deviation over the averaged returns from each training run. All algorithms use a constant learning rate $\alpha = 0.01$ and an exploration rate which is linearly decayed from $\epsilon = 1$ to $\epsilon = 0.05$ over the first 80,000 time steps.

6.3.3 Bayesian Learning and Value of Information

Fictitious play (using Equation 6.14) and JAL-AM (using Equation 6.16) learn a *single* model for each other agent, and compute best-response actions with respect to the models. What these methods are lacking is an explicit representation of *uncertainty* about the models; that is, there may be different possible models for the other agents, and the modelling agent may have beliefs about the relative likelihood of each model given the past actions of the modelled agents. Maintaining such beliefs enables a learning agent to compute best-response actions with respect to the different models and their associated likelihoods. Moreover, as we will show in this section, it is possible to compute best-response actions that maximise the *value of information* (VI) (Chalkiadakis and Boutilier 2003; Albrecht, Crandall, and Ramamoorthy 2016). VI evaluates how the outcomes of an action may influence the learning agent’s beliefs about the other agents, and how the changed beliefs will in turn influence the future actions of the learning agent. VI also accounts for how the action may influence the future behaviour of the modelled agents. Thus, best responses based on VI can optimally trade-off between exploring actions to obtain more accurate beliefs about agent models on one hand, and the potential benefits and costs involved

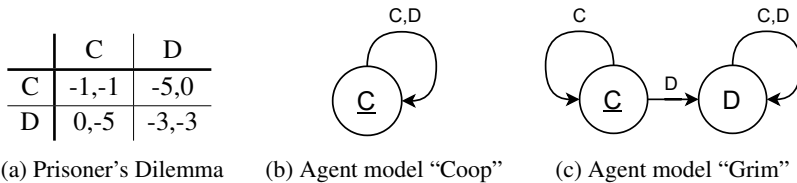


Figure 6.8: Two agent models for Prisoner's Dilemma. The models are shown as simple finite state automata, where the state (circle) specifies the model's action (C for cooperating, D for defecting) and the state transition arrows are labelled by the previous action of the other agent. Starting states are shown underlined. The model Coop always cooperates. The model Grim cooperates initially until the other agent defects, after which Grim defects indefinitely.

in the exploration on the other hand. Before we define the notions of beliefs and VI in more detail, we will give an illustrative example.

Suppose two agents play the repeated Prisoner's Dilemma matrix game, first discussed in Section 3.1 and shown again in Figure 6.8a, where every episode lasts 10 time steps. In each time step, each agent can choose to either cooperate or defect, where mutual cooperation gives a reward of -1 to each agent, but each agent has an incentive to defect in order to achieve higher rewards. Suppose agent 1 believes that agent 2 can have one of two possible models, which are shown in Figures 6.8b and 6.8c. The model Coop always cooperates, while the model Grim cooperates initially until the other agent defects after which Grim defects indefinitely. Assume agent 1 has a uniform prior belief which assigns probability 0.5 to each model. Given this uncertainty about agent 2's policy, how should agent 1 choose its first action? Consider the following cases (summarised in Figure 6.9):

- If agent 1 cooperates initially and agent 2 uses Coop, then agent 1 has not learned anything new about agent 2's model (meaning the probabilities assigned by agent 1's belief to the models will not change), since both Coop and Grim cooperate in response to agent 1 cooperating. If agent 1 cooperates throughout the episode, it will obtain a return of $10 * (-1) = -10$. Therefore, agent 1 will get a lower return than it could have achieved by using the defect action against Coop.
- On the other hand, if agent 1 defects initially and agent 2 uses Coop, then agent 2 will continue to cooperate and agent 1 will learn that agent 2 uses Coop (meaning that agent 1's belief will assign probability 1 to Coop), since Grim would not cooperate in this case. With this new knowledge about

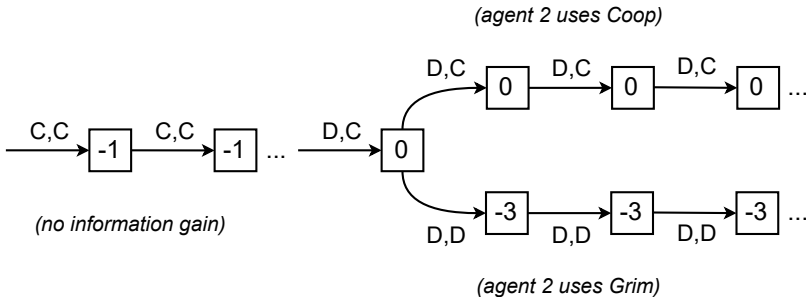


Figure 6.9: Value of information in the Prisoner’s Dilemma example. Arrows show joint action of agents 1 and 2, boxes show reward for agent 1. Action C provides no information gain to agent 1 since both Coop and Grim cooperate in response to agent 1 cooperating. Action D will cause agent 2 to either cooperate (Coop) or defect (Grim) in response, after which agent 1 will know which model agent 2 uses and can select the appropriate best-response action.

agent 2, agent 1 can achieve maximum returns by defecting throughout the episode, resulting in a return of $10 * 0 = 0$.

- However, if agent 1 defects initially but agent 2 uses Grim, then agent 2 will defect subsequently and agent 1 will learn that agent 2 uses Grim, which will lead agent 1 to continue defecting as its best response. Therefore, this new knowledge about agent 2 comes at the cost of a lower return, which is $0 + 9 * (-3) = -27$ (one reward of 0, followed by -3 rewards until the end of the episode).

This example illustrates that some actions may reveal information about the policies of other agents, and the resulting more accurate beliefs can be used to maximise the returns; however, the same actions may carry a risk in that they may inadvertently change the other agents’ behaviours and result in lower achievable returns.

In the above example, if agent 1 uses randomised exploration methods such as ϵ -greedy exploration, then eventually it will defect, which will cause agent 2 to go into the “defect mode” if it uses Grim. VI instead evaluates the different constellations of agent 1’s models and actions, and the impact on agent 1’s future beliefs and actions under the different constellations, essentially as we did in the above example. Depending on how far VI looks into the future when evaluating an action, it may decide that exploring the defect action is not worthwhile considering the potential cost (i.e. lower achievable return) if agent

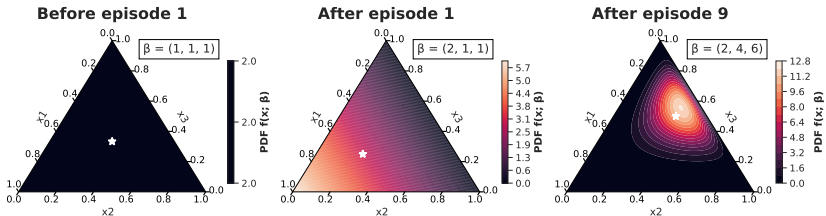


Figure 6.10: Agent 1’s beliefs about agent 2’s model (policy) after varying episodes from the Rock-Paper-Scissors non-repeated game example in Figure 6.6 (page 120): before episode 1 (left), after episode 1 (middle), and after episode 9 (right). Beliefs are represented as Dirichlet distributions over probability distributions $x = (x_1, x_2, x_3)$ for actions R,P,S. Each point in the simplex triangle shows the Dirichlet probability density (defined in Footnote 46) for a distribution x that is mapped into the simplex coordinate space via barycentric coordinates. Pseudocount parameters (β_k) are shown next to each simplex. Means are shown as white stars.

2 uses Grim. In this case, if both agents maintain beliefs over Coop/Grim and use VI, they may learn a joint policy in which both agents cooperate.

We now describe these ideas more formally, starting with beliefs over agent models. Assume we control agent i . Let $\hat{\Pi}_j$ be the space of possible agent models for agent j , where each model $\hat{\pi}_j \in \hat{\Pi}_j$ can choose actions based on the interaction history h^t . (Recall that we assume stochastic games, in which agents fully observe the state and actions.) In our example above, $\hat{\Pi}_j$ contains two models $\hat{\pi}_j^{Coop}$ and $\hat{\pi}_j^{Grim}$. Agent i starts with a prior belief $\Pr(\hat{\pi}_j | h^0)$ which assigns probabilities to each model $\hat{\pi}_j \in \hat{\Pi}_j$ before observing any actions from agent j . If no prior information is available which makes one agent model more likely than others, then the prior belief could simply be a uniform probability distribution, as used in our example. After observing agent j ’s action a_j^t in state s^t , agent i updates its belief by computing a Bayesian posterior distribution:

$$\Pr(\hat{\pi}_j | h^{t+1}) = \frac{\hat{\pi}_j(a_j^t | h^t) \Pr(\hat{\pi}_j | h^t)}{\sum_{\hat{\pi}'_j \in \hat{\Pi}_j} \hat{\pi}'_j(a_j^t | h^t) \Pr(\hat{\pi}'_j | h^t)} \quad (6.18)$$

Note that this Bayesian belief update is computed *across* episodes, meaning that the posterior $\Pr(\hat{\pi}_j | h^T)$ at the end of episode e is used as the prior $\Pr(\hat{\pi}_j | h^0)$ at the start of episode $e + 1$.

The above definition of beliefs assumes that $\hat{\Pi}_j$ contains a finite number of models. We can also define $\hat{\Pi}_j$ to be a continuous space of agent models, in which case the beliefs are defined as probability densities over $\hat{\Pi}_j$ rather than

discrete probability distributions. For example, we may define $\hat{\Pi}_j$ to contain all possible policies $\hat{\pi}_j(a_j \mid s)$ that are conditioned on states, as we did in JAL-AM in Equation 6.16. We can then define the belief over $\hat{\Pi}_j$ based on a set of Dirichlet distributions,⁴⁶ with one Dirichlet δ_s corresponding to each state $s \in S$ to represent a belief over possible policies $\hat{\pi}_j(\cdot \mid s)$ in state s , where δ_s is parameterised by pseudocounts $(\beta_1, \dots, \beta_{|A_j|})$ (one for each action). An initial setting of $\beta_k = 1$ for all k produces a uniform prior belief. After observing action a_j^t in state s^t , the belief update is simply done by incrementing (by 1) the pseudocount β_k in δ_s that is associated with a_j^t . The resulting posterior distribution is again a Dirichlet distribution. Note that the mean of a Dirichlet distribution δ_s is given by the probability distribution $\hat{\pi}_j(a_{j,k} \mid s) = \frac{\beta_k}{\sum_l \beta_l}$ where β_k is the pseudocount associated with action $a_{j,k}$. Therefore, the mean of the Dirichlet corresponds to the definition of the empirical distribution used in fictitious play (Equation 6.14) and JAL-AM (Equation 6.16).⁴⁷ Example Dirichlet distributions after varying episodes from the Rock-Paper-Scissors non-repeated game (i.e. stochastic game with a single state) example in Figure 6.6 (page 120) can be seen in Figure 6.10.

Given finite model spaces $\hat{\Pi}_j$ and beliefs $\Pr(\hat{\pi}_j \mid h^t)$ for each other agent $j \neq i$, define $\hat{\Pi}_{-i} = \times_{j \neq i} \hat{\Pi}_j$ and $\Pr(\hat{\pi}_{-i} \mid h) = \prod_{j \neq i} \Pr(\hat{\pi}_j \mid h)$ for $\hat{\pi}_{-i} \in \hat{\Pi}_{-i}$. Let $s(h)$ denote the last state in history h (i.e. $s(h^t) = s^t$), and $\langle \rangle$ denote the concatenation operation. The VI of action $a_i \in A_i$ after history h is defined via a recursive combination of two functions:

$$VI_i(a_i \mid h) = \sum_{\hat{\pi}_{-i} \in \hat{\Pi}_{-i}} \Pr(\hat{\pi}_{-i} \mid h) \sum_{a_{-i} \in A_{-i}} Q_i(h, \langle a_i, a_{-i} \rangle) \prod_{j \neq i} \hat{\pi}_j(a_j \mid h) \quad (6.19)$$

$$Q_i(h, a) = \sum_{s' \in S} \mathcal{T}(s' \mid s(h), a) \left[\mathcal{R}_i(s(h), a, s') + \gamma \max_{a'_i \in A_i} VI_i(a'_i \mid \langle h, a, s' \rangle) \right] \quad (6.20)$$

46. A Dirichlet distribution of order $K \geq 2$ with parameters $\beta = (\beta_1, \dots, \beta_K)$ has a probability density function given by $f(x_1, \dots, x_K; \beta) = B(\beta)^{-1} \prod_{k=1}^K x_k^{\beta_k-1}$, where $\{x_k\}_{k=1,\dots,K}$ belong to the standard K-1 simplex (i.e. the x_k define a probability distribution), and $B(\beta)$ is the multivariate beta function. A Dirichlet distribution can be interpreted as specifying a likelihood over different probability values for $\{x_k\}$. It is the conjugate prior of the categorical and multinomial distributions.

47. However, note the difference that the Dirichlet prior is initialised with pseudocounts $\beta_k = 1$ for all actions (prior to observing any actions), while the empirical distribution used in fictitious play and JAL-AM is defined over the observed history of actions. Therefore, after one observed action, the empirical distribution assigns probability 1 to that action while the Dirichlet mean assigns probability less than 1. However, in practice, this difference is “washed away” as the number of observed actions increases, and in the limit of observing infinitely many actions (or state-action pairs), the Dirichlet mean and the empirical distribution used by fictitious play and JAL-AM converge to the same probability distribution. Moreover, by using initial action counts of $C(a_j) = 1$ (or $C(s, a_j) = 1$) for all actions, the empirical distribution is identical to the Dirichlet mean.

(The case of continuous $\hat{\Pi}_j$ can be defined analogously, by using integrals and densities instead of sums and probabilities.) $VI_i(a_i | h)$ computes the VI of action a_i by summing the action values $Q_i(h, \langle a_i, a_{-i} \rangle)$ for all possible action combinations a_{-i} of the other agents, weighted by the probabilities assigned by agent i 's belief to models $\hat{\pi}_{-i}$ and the probabilities with which $\hat{\pi}_{-i}$ would take actions a_{-i} . The action value $Q_i(h, a)$ is defined analogously to the Bellman optimality equation (defined in Equation 2.24, page 28), except that the max-operator uses VI_i rather than Q_i . Importantly, Q_i extends the history h to $\langle h, a, s' \rangle$ and recurses back to VI_i with the extended history. Since VI_i uses the posterior belief $\Pr(\hat{\pi}_{-i} | h)$, this means that the computation of VI_i accounts for how the beliefs will change in the future for different possible histories h . The depth of the recursion⁴⁸ between Q_i and VI_i determines how far VI evaluates the impact of an action into the future of the interaction. Using this definition of VI, at time t after history h^t , agent i selects an action a_i^t with maximum VI, i.e. $a_i^t \in \arg \max_{a_i} VI_i(a_i | h^t)$.

When applied in the Prisoner's Dilemma example given above, using a uniform prior belief over the two models $\Pr(\hat{\pi}_j^{Coop} | h^0) = \Pr(\hat{\pi}_j^{Grim} | h^0) = 0.5$, no discounting (i.e. $\gamma = 1$), and a recursion depth equal to the length of the episode (i.e. 10), we obtain the following VI values for agent 1's actions in the initial time step $t=0$:

$$VI_1(C) = -9 \quad VI_1(D) = -13.5 \quad (6.21)$$

Therefore, in this example when using a uniform prior belief, cooperating has a higher value of information than defecting. Essentially, while VI realises that the defect action can give absolute certainty about the model used by the other agent, if that agent uses the Grim model then the cost of causing the agent to defect until the end of the episode is not worth the absolute certainty in beliefs. If both agents use this VI approach to select actions, then they will both cooperate initially, and will continue to cooperate until the last time step. In the last time step $t=9$, the agents obtain the following VI values (for $i \in \{1, 2\}$):

$$VI_i(C) = -1 \quad VI_i(D) = 0 \quad (6.22)$$

Thus, both agents will defect in the last time step. In this case, as this is the last time step, there is no more risk in defecting since both Coop and Grim

48. To make the recursion anchor explicit in the definition of VI, we can define VI_i^d and Q_i^d for recursion depth $d \geq 0$, where VI_i^d calls Q_i^d and Q_i^d calls VI_i^{d-1} , and we define $VI_i^d(a_i | h) = 0$ if $d = 0$ or if $s(h)$ is a terminal state (i.e. end of episode). For the case of $d = 0$, instead of assigning value $VI_i^d(a_i | h) = 0$, Chalkiadakis and Boutilier (2003) describe an alternative "myopic" approach by fixing the belief $\Pr(\hat{\pi}_{-i} | h)$ at the end of the recursion, sampling agent models $\hat{\pi}_{-i}$ from the fixed belief, solving the corresponding MDPs in which agents $-i$ use those sampled models, and averaging over the resulting Q-values from the MDPs.

are expected to cooperate (hence, in either case, defecting will give reward 0) and the episode will finish. This is an example of the end-game effects first mentioned in Section 3.2.

The idea of maintaining Bayesian beliefs over a space of agent models and computing best-responses with respect to the beliefs has been studied in game theory under the name “rational learning” (Jordan 1991; Kalai and Lehrer 1993; Nachbar 1997; Foster and Young 2001; Nachbar 2005). A central result in rational learning is that, under certain strict assumptions, the agents’ predictions of future play will converge to the true distribution of play induced by the agents’ actual policies, and the agents’ policies will converge to a Nash equilibrium (Kalai and Lehrer 1993). One important condition of the convergence result is that any history that has positive probability under the agents’ actual policies must have positive probability under the agents’ beliefs (also known as “absolute continuity”). This assumption can be violated if the best-responses of agent j are not predicted by any model in $\hat{\Pi}_j$. In our Prisoner’s Dilemma example, if the agents start with a prior belief that assigns probability 0.8 to Coop and 0.2 to Grim, then the agents will obtain VI values of $VI_i(C) = -5.8$, $VI_i(D) = -5.4$ and both agents will initially defect, which is not predicted by the Coop nor Grim model.⁴⁹ On the other hand, when using the Dirichlet beliefs discussed earlier, then any finite history will always have non-zero probability under the agents’ beliefs. Note that an important difference between the Bayesian learning presented here and rational learning is that the latter does not use value of information as defined here. Instead, rational learning assumes that agents compute best-responses with respect to their current belief over models, without considering how different actions may affect their beliefs in the future.

6.4 Policy-Based Learning

The algorithms presented so far in this chapter all have in common that they estimate or learn the *values* of actions or joint actions. The agents’ policies are then derived using the value functions. We saw that this approach has some important limitations. In particular, the joint action values learned by JAL-GT algorithms may not carry sufficient information to derive the correct equilibrium joint policy (Section 6.2.4); and algorithms such as fictitious play and JAL-AM are unable to represent probabilistic policies due to their use of best-response actions, which means that they cannot learn probabilistic equilibrium policies.

49. In fact, the choice of prior belief can have a substantial effect on the achievable returns and convergence to equilibrium (Nyarko 1998; Dekel, Fudenberg, and Levine 2004; Albrecht, Crandall, and Ramamurthy 2015).

Another major category of algorithms in MARL instead uses the learning data to directly optimise a parameterised joint policy, based on gradient-ascent techniques. These policy-based learning algorithms have the important advantage that they can directly learn the action probabilities in policies, which means they can represent probabilistic equilibria. As we will see, by gradually varying the action probabilities in learned policies, these algorithms achieve some interesting convergence properties. Thus, as is the case in single-agent RL, there are value-based and policy-based methods in MARL. This section will present several of the original methods developed in this latter category.

6.4.1 Gradient Ascent in Expected Reward

We begin by examining gradient-ascent learning in general-sum non-repeated normal-form games with two agents and two actions. It will be useful to introduce some additional notation. First, we will write the reward matrices of the two agents as follows:

$$\mathcal{R}_i = \begin{bmatrix} r_{1,1} & r_{1,2} \\ r_{2,1} & r_{2,2} \end{bmatrix} \quad \mathcal{R}_j = \begin{bmatrix} c_{1,1} & c_{1,2} \\ c_{2,1} & c_{2,2} \end{bmatrix} \quad (6.23)$$

\mathcal{R}_i is the reward matrix of agent i and \mathcal{R}_j is the reward matrix of agent j . Thus, if agent i chooses action x and agent j chooses action y , then they will receive rewards $r_{x,y}$ and $c_{x,y}$, respectively.

Since the agents are learning policies for a non-repeated normal-form game (where each episode consists of a single time step), we can represent their policies as simple probability distributions:

$$\pi_i = (\alpha, 1 - \alpha) \quad \pi_j = (\beta, 1 - \beta), \quad \alpha, \beta \in [0, 1] \quad (6.24)$$

where α and β are the probabilities with which agent 1 and agent 2 choose action 1, respectively. Note that this means that a joint policy $\pi = (\pi_i, \pi_j)$ is a point in the unit square $[0, 1]^2$. We will write (α, β) as a shorthand for the joint policy.

Given a joint policy (α, β) , we can write the expected reward for each agent as follows:

$$\begin{aligned} U_i(\alpha, \beta) &= \alpha\beta r_{1,1} + \alpha(1-\beta)r_{1,2} + (1-\alpha)\beta r_{2,1} + (1-\alpha)(1-\beta)r_{2,2} \\ &= \alpha\beta u + \alpha(r_{1,2} - r_{2,2}) + \beta(r_{2,1} - r_{2,2}) + r_{2,2} \end{aligned} \quad (6.25)$$

$$\begin{aligned} U_j(\alpha, \beta) &= \alpha\beta c_{1,1} + \alpha(1-\beta)c_{1,2} + (1-\alpha)\beta c_{2,1} + (1-\alpha)(1-\beta)c_{2,2} \\ &= \alpha\beta u' + \alpha(c_{1,2} - c_{2,2}) + \beta(c_{2,1} - c_{2,2}) + c_{2,2} \end{aligned} \quad (6.26)$$

where

$$u = r_{1,1} - r_{1,2} - r_{2,1} + r_{2,2} \quad (6.27)$$

$$u' = c_{1,1} - c_{1,2} - c_{2,1} + c_{2,2}. \quad (6.28)$$

The gradient-ascent learning method we consider in this section updates the agents' policies to maximise their expected rewards defined above. Let (α^k, β^k) be the joint policy at episode k . Each agent updates its policy in the direction of the gradient in expected reward using some step size $\kappa > 0$:

$$\alpha^{k+1} = \alpha^k + \kappa \frac{\partial U_i(\alpha^k, \beta^k)}{\partial \alpha^k} \quad (6.29)$$

$$\beta^{k+1} = \beta^k + \kappa \frac{\partial U_j(\alpha^k, \beta^k)}{\partial \beta^k} \quad (6.30)$$

where the partial derivative of an agent's expected reward with respect to its policy takes the simple form of

$$\frac{\partial U_i(\alpha, \beta)}{\partial \alpha} = \beta u + (r_{1,2} - r_{2,2}) \quad (6.31)$$

$$\frac{\partial U_j(\alpha, \beta)}{\partial \beta} = \alpha u' + (c_{2,1} - c_{1,2}). \quad (6.32)$$

An important special case in this procedure is when the updated joint policy moves outside the valid probability space, i.e. the unit square. This can happen when (α^k, β^k) is on the boundary of the unit square, meaning at least one of α^k and β^k has value 0 or 1, and the gradient points outside the unit square. In this case, the gradient is redefined to project back onto the boundary of the unit square, ensuring that $(\alpha^{k+1}, \beta^{k+1})$ will remain valid probabilities.

From the update rule in Equations 6.29 and 6.30, we can see that this method of learning implies some strong knowledge assumptions. In particular, each agent must know its own reward matrix and the policy of the other agent in the current episode k . In Section 6.4.5, we will see a generalisation of this learning method which does not require this knowledge.

6.4.2 Learning Dynamics of Infinitesimal Gradient Ascent

What kind of joint policy will the two agents learn if they follow the learning rule given in Equations 6.29 and 6.30? Will the joint policy converge? And if yes, to what type of solution?

We can study these questions using dynamical systems theory (Singh, Kearns, and Mansour 2000). If we consider "infinitely small" step sizes $\kappa \rightarrow 0$, then the joint policy will follow a continuous trajectory $(\alpha(t), \beta(t))$ in continuous time t

which evolves according to the following differential equation (we will write (α, β) to refer to $(\alpha(t), \beta(t))$ in the rest of this section):

$$\begin{bmatrix} \frac{\partial \alpha}{\partial t} \\ \frac{\partial \beta}{\partial t} \end{bmatrix} = \underbrace{\begin{bmatrix} 0 & u \\ u' & 0 \end{bmatrix}}_F \begin{bmatrix} \alpha \\ \beta \end{bmatrix} + \begin{bmatrix} (r_{1,2} - r_{2,2}) \\ (c_{2,1} - c_{2,2}) \end{bmatrix} \quad (6.33)$$

F denotes the off-diagonal matrix which contains the terms u and u' . This learning algorithm using an infinitesimal step size is referred to as *Infinitesimal Gradient Ascent*, or IGA for short.

The centre point (α^*, β^*) which has a zero-gradient can be found by setting the left-hand side of Equation 6.33 to zero and solving to obtain:

$$(\alpha^*, \beta^*) = \left(\frac{c_{2,2} - c_{2,1}}{u'}, \frac{r_{2,2} - r_{1,2}}{u} \right) \quad (6.34)$$

It can be shown that the dynamical system described in Equation 6.33 is an affine dynamical system, which means that (α, β) will follow one of three possible types of trajectories. The type of trajectory depends on the specific values of u and u' . First, recall that in order to compute the eigenvalue λ of matrix F , we have to find λ and a vector $x \neq 0$ such that $Fx = \lambda x$. Solving for λ gives $\lambda = \sqrt{uu'}$. Then, the three possible types of trajectories are the following:

1. If F is not invertible, then (α, β) will follow a divergent trajectory, as shown in Figure 6.11a. This happens if u or u' (or both) are zero, which can occur in common-reward, zero-sum, and general-sum games.
2. If F is invertible and has purely real eigenvalues, then (α, β) will follow a divergent trajectory to and away from the centre point, as shown in Figure 6.11b. This happens if $uu' > 0$, which can occur in common-reward and general-sum games, but not in zero-sum games (since $u = -u'$, thus $uu' \leq 0$).
3. If F is invertible and has purely imaginary eigenvalues, then (α, β) will follow an ellipse trajectory around the centre point, as shown in Figure 6.11c. This happens if $uu' < 0$, which can occur in zero-sum and general-sum games, but not in common-reward games (since $u = u'$, thus $uu' \geq 0$).

Note that this dynamical system does not take into account the constraint that (α, β) must lie in the unit square. Thus, the centre point in general may not lie in the unit square. In the unconstrained system, there exists at most one point with zero-gradient (there is no such point if F is not invertible). The constrained system, which projects gradients on the boundary of the unit square back into the unit square, may include additional points with zero-gradient on

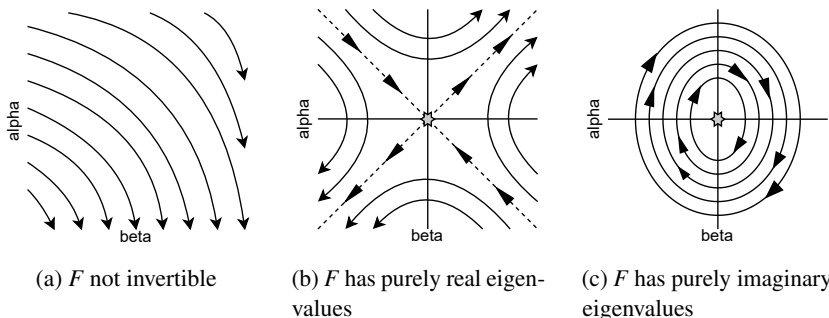


Figure 6.11: The joint policy (α, β) learned by Infinitesimal Gradient Ascent (IGA) in the unconstrained space (i.e. (α, β) may be outside the unit square) will follow one of three possible types of trajectories, depending on properties of the matrix F from Equation 6.33. Shown here are the general schematic forms of the trajectory types; the exact trajectory will depend on the values of u and u' . The plot axes correspond to values of α, β and arrows indicate the direction of the (α, β) -trajectories. The centre point which has zero-gradient (if one exists) is marked by the star in the centre. For non-invertible F , (a) shows one possible set of trajectories, but others are possible depending on u and u' .

the boundary of the unit square. In any case, IGA converges if and only if it reaches a point where the projected gradient is zero.

Based on the dynamical system described above, several properties of gradient ascent learning can be established:

- (α, β) does not converge in all cases.
- If (α, β) does not converge, then the average rewards received during learning converge to the expected rewards of some Nash equilibrium.
- If (α, β) converges, then the converged joint policy is a Nash equilibrium.

While if F is non-invertible or has real eigenvalues it can be shown that (α, β) converges to a Nash equilibrium in the constrained system, this does not hold if F has imaginary eigenvalues where ellipses can be wholly contained inside the unit square, in which case (α, β) will cycle indefinitely if it follows such an ellipse. Notably, however, Singh, Kearns, and Mansour (2000) showed that the average rewards obtained by following such an ellipse converge to the expected rewards of a Nash equilibrium of the game. Thus, this is an example of the convergence type defined in Equation 5.6 (Section 5.2). An implication of this result is that when (α, β) converges, then (α, β) must be a Nash equilibrium. This can also be seen by the fact that (α, β) converges if and only if reaching a

point where the (projected) gradient is zero, and such points can be shown to be Nash equilibria in the constrained system (since otherwise the gradient cannot be zero). Finally, Singh, Kearns, and Mansour (2000) showed that these results also hold for finite step sizes κ if κ is appropriately reduced during learning (e.g. $\kappa^k = \frac{1}{k^{2/3}}$).

6.4.3 Win or Learn Fast

The IGA learning method presented in the preceding sections guarantees that the average rewards received by the agents converge in the limit to the expected rewards of a Nash equilibrium. In practice, this type of convergence is relatively weak since the reward received at any time may be arbitrarily low, as long as this is compensated by an arbitrarily high reward in the past or future. We would prefer that the actual policies of the agents converge to a Nash equilibrium, as per Equation 5.3.

As it turns out, the problem in IGA which prevents convergence of the policies in all cases is when using a constant step size (or learning rate) κ . However, if we allow the step size to vary over time, we can construct a sequence of step sizes such that (α^k, β^k) is always guaranteed to converge to a Nash equilibrium of the game. Specifically, we modify the learning rule to

$$\alpha^{k+1} = \alpha^k + l_i^k \kappa \frac{\partial U_i(\alpha^k, \beta^k)}{\partial \alpha^k} \quad (6.35)$$

$$\beta^{k+1} = \beta^k + l_j^k \kappa \frac{\partial U_j(\alpha^k, \beta^k)}{\partial \beta^k} \quad (6.36)$$

where $l_i^k, l_j^k \in [l_{\min}, l_{\max}] > 0$, and we still use $\kappa \rightarrow 0$. Thus, l_i^k, l_j^k may vary in each update, and the overall step size $l_{ij}^k \kappa$ is bounded.

The principle by which we vary the learning rates l_i^k, l_j^k is to learn quickly when “losing” (i.e. use l_{\max}) and to learn slowly when “winning” (i.e. use l_{\min}). This principle is known as *Win or Learn Fast*, or WoLF (Bowling and Veloso 2002). The idea here is that if an agent is losing, it should try to adapt quickly to catch up with the other agent. If the agent is winning, it should adapt slowly since the other agent will likely change its policy. The determination of losing/winning is based on comparing the actual expected rewards with the expected rewards achieved by a Nash equilibrium policy. Formally, let α^e be an equilibrium policy chosen by agent i , and β^e be an equilibrium policy chosen by agent j . Then, the agents will use the following variable learning rates:

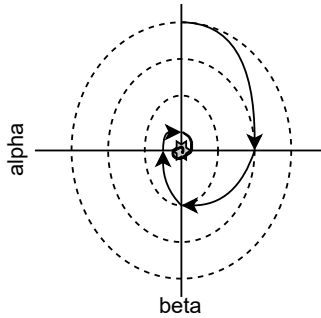


Figure 6.12: General form of joint policy (α, β) trajectory when using WoLF-IGA, for the case when $F(t)$ has purely imaginary eigenvalues and the unique centre point is contained in the unit square. The trajectory (solid line) tightens in each quadrant and converges to the centre point (grey star), which is a Nash equilibrium. The plot axes correspond to values for α, β .

$$l_i^k = \begin{cases} l_{\min} & \text{if } U_i(\alpha^k, \beta^k) > U_i(\alpha^e, \beta^k) \\ l_{\max} & \text{otherwise} \end{cases} \quad (\text{winning}) \quad (6.37)$$

$$l_j^k = \begin{cases} l_{\min} & \text{if } U_j(\alpha^k, \beta^k) > U_j(\alpha^k, \beta^e) \\ l_{\max} & \text{otherwise} \end{cases} \quad (\text{winning}) \quad (6.38)$$

This modified IGA learning rule using a variable learning rate is called *WoLF-IGA*. Note that α^e and β^e need not be from the same equilibrium, meaning that (α^e, β^e) may not form a Nash equilibrium.

Using this variable learning rate, it can be proven that WoLF-IGA is guaranteed to converge to a Nash equilibrium in general-sum games with two agents and two actions. The analysis of WoLF-IGA is near-identical to IGA: using infinitesimal step sizes, the joint policy will follow a continuous trajectory $(\alpha(t), \beta(t))$ in continuous time t which evolves according to the following differential equation:

$$\begin{bmatrix} \frac{\partial \alpha}{\partial t} \\ \frac{\partial \beta}{\partial t} \end{bmatrix} = \underbrace{\begin{bmatrix} 0 & l_i(t)u \\ l_j(t)u' & 0 \end{bmatrix}}_{F(t)} \begin{bmatrix} \alpha \\ \beta \end{bmatrix} + \begin{bmatrix} l_i(t)(r_{1,2} - r_{2,2}) \\ l_j(t)(c_{2,1} - c_{2,2}) \end{bmatrix}. \quad (6.39)$$

As was the case for IGA, we again examine three qualitatively distinct trajectory types based on properties of the matrix $F(t)$, which now depends on t due to the variable learning rates. These cases are when 1) $F(t)$ is non-invertible,

2) $F(t)$ is invertible and has purely real eigenvalues, and 3) $F(t)$ is invertible and has purely imaginary eigenvalues. The crucial part in the analysis is case 3), and in particular the sub-case where the centre point is contained within the unit square. This is the problematic case where IGA will not converge to a Nash equilibrium. The WoLF variable learning rate has an important effect in this sub-case: Bowling and Veloso (2002) showed that in WoLF-IGA, the trajectories of (α, β) are in fact piecewise elliptical, with four pieces given by the four quadrants around the centre point (as shown in Figure 6.12), such that the trajectories spiral toward the centre point. Note that in this case, there exists only one centre point and this is by definition a Nash equilibrium. In each quadrant, the ellipse will “tighten” by a factor of $\sqrt{\frac{l_{\min}}{l_{\max}}} < 1$, and thus the trajectory will converge to the centre point.

6.4.4 Win or Learn Fast with Policy Hill Climbing

So far, both IGA and WoLF-IGA have been defined only for normal-form games with two agents and two actions. Moreover, these methods require complete knowledge about the learning agent’s reward function and the policy of the other agent, which can be quite restrictive assumptions in an RL setting. *Win or Learn Fast with Policy Hill Climbing* (WoLF-PHC) (Bowling and Veloso 2002) is a MARL algorithm which can be used in general-sum stochastic games with any finite number agents and actions; and it does not require knowledge about the agents’ reward functions nor the policies of other agents.

Algorithm 9 shows the pseudocode for WoLF-PHC. The algorithm learns action values $Q(s, a_i)$ as in standard Q-learning (Line 11; here α denotes the learning rate for Q-updates), and updates the policy π_i using the WoLF learning principle. When determining winning/losing to compute the learning rate, instead of comparing the expected rewards to Nash equilibrium policies as is done in WoLF-IGA, WoLF-PHC compares the expected reward of π_i to the expected reward of an “average” policy $\bar{\pi}_i$ which averages over past policies (Line 12). The idea is that this average policy replaces the (unknown) Nash equilibrium policy for agent i . A similar idea is used in fictitious play (Section 6.3.1), where it is shown that the average policy does in fact converge to a Nash equilibrium policy in many types of games.

WoLF-PHC uses two parameters $l_l, l_w \in (0, 1]$ with $l_l > l_w$, corresponding to the learning rates for losing and winning, respectively. The action probabilities $\pi_i(s^t, a_i)$ are updated by adding a term $\Delta(s^t, a_i)$, defined as

$$\Delta(s, a_i) = \begin{cases} -\delta_{s,a_i}, & a_i \notin \arg \max_{a'_i} Q(s, a'_i) \\ \sum_{a'_i \neq a_i} \delta_{s,a'_i}, & \text{else} \end{cases} \quad (6.43)$$

Algorithm 9 Win or Learn Fast with Policy Hill Climbing (WoLF-PHC)

-
- // Algorithm controls agent i
- 1: Initialise:
 - 2: Learning rates $\alpha, l_l, l_w \in (0, 1]$ with $l_l > l_w$
 - 3: $Q(s, a_i) \leftarrow 0$ and $\pi_i(s, a_i) \leftarrow \frac{1}{|A_i|}$, for all $s \in S, a_i \in A_i$
 - 4: $\bar{\pi}_i \leftarrow \pi_i$
 - 5: Repeat for every episode:
 - 6: **for** $t = 0, 1, 2, \dots$ **do**
 - 7: Observe current state s^t
 - 8: With probability ϵ : choose random action $a_i^t \in A_i$
 - 9: Else: sample action a_i^t using probabilities $\pi_i(s^t, \cdot)$
 - 10: Observe reward r_i^t and next state s^{t+1}
 - 11: Update Q-value:
- $$Q(s^t, a_i^t) \leftarrow Q(s^t, a_i^t) + \alpha \left[r_i^t + \gamma \max_{a'_i} Q_i(s^{t+1}, a'_i) - Q_i(s^t, a_i^t) \right] \quad (6.40)$$
- 12: Update average policy $\bar{\pi}_i$ for all $a_i \in A_i$:
- $$\bar{\pi}_i(s^t, a_i) \leftarrow \bar{\pi}_i(s^t, a_i) + \frac{1}{t+1} (\pi_i(s^t, a_i) - \bar{\pi}_i(s^t, a_i)) \quad (6.41)$$
- 13: Update policy π_i for all $a_i \in A_i$:
- $$\pi_i(s^t, a_i) \leftarrow \pi_i(s^t, a_i) + \Delta(s^t, a_i) \quad (6.42)$$
- with $\Delta(s^t, a_i)$ computed as per Equation 6.43.
-

where

$$\delta_{s, a_i} = \min \left(\pi_i(s, a_i), \frac{\delta}{|A_i| - 1} \right) \quad (6.44)$$

$$\delta = \begin{cases} l_w, & \sum_{a'_i} \pi_i(s, a'_i) Q(s, a'_i) > \sum_{a'_i} \bar{\pi}_i(s, a'_i) Q(s, a'_i) \\ l_l, & \text{else} \end{cases} . \quad (6.45)$$

Therefore, the lower learning rate l_w is used if the expected reward of π_i is greater than the expected reward of $\bar{\pi}_i$ (winning), and the larger learning rate l_l is used if the expected reward of π_i is lower than or equal to the expected reward of $\bar{\pi}_i$ (losing).

Figure 6.13 shows WoLF-PHC applied in the Rock-Paper-Scissors matrix game, where it can be seen that the agents' policies gradually co-adapt and converge to the unique Nash equilibrium of the game, in which both agents choose actions uniform-randomly. The learning trajectories produced by WoLF-PHC

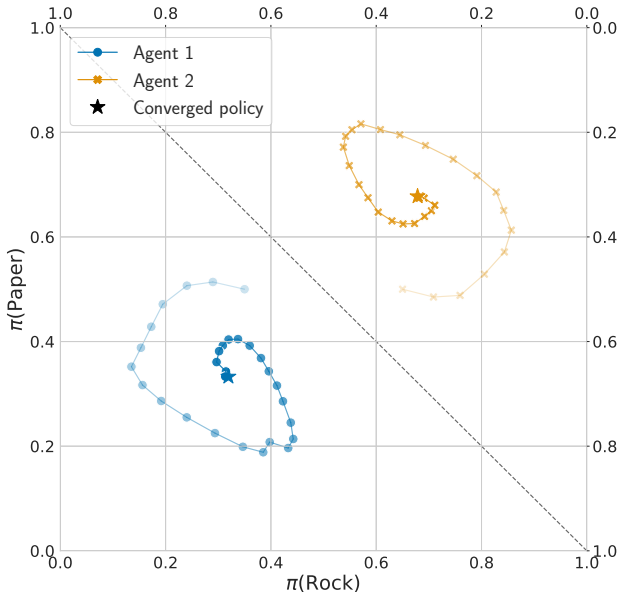


Figure 6.13: This plot shows the evolving policies of two agents in the Rock-Paper-Scissors game where both agents use the WoLF-PHC algorithm to update their policies in each time step. Each point in the simplex for an agent corresponds to a probability distribution over the three available actions. Each line shows the current policy at time steps $t=0, 5, 10, 15, \dots, 145$ as well as the converged policy at $t=100,000$.

are similar to the trajectories produced by fictitious play shown in Figure 6.5 (page 119), but are “smoother” and circular rather than triangular. This is because fictitious play uses deterministic best-response actions, which can change abruptly over time, whereas WoLF-PHC learns action probabilities which can vary gradually over time.

6.4.5 Generalised Infinitesimal Gradient Ascent

We have so far examined IGA based on its ability to learn Nash equilibria or Nash equilibrium rewards. Another major solution concept is no-regret, which was presented in Section 4.10. In this section, we will present a gradient-based learning algorithm which generalises IGA to normal-form games with more than two agents and actions. As we will see, this *Generalised Infinitesimal Gradient Ascent* (GIGA) algorithm achieves no-regret (Zinkevich 2003), which implies that IGA also achieves no-regret.

GIGA does not require knowledge of the other agents' policies, but assumes that it can observe the past actions of the other agents. Similarly to IGA, GIGA updates policies using unconstrained gradients which are projected back into the space of valid probability distributions. However, while IGA uses a gradient in *expected* reward with respect to the agents' policies, GIGA uses a gradient in *actual* rewards after observing the past actions of the other agents. In the following, we will present GIGA for games with two agents i and j , and we will later return to the case of $n > 2$ agents.

Given a policy π_i for agent i and an action a_j for agent j , the expected reward for agent i against action a_j is

$$U_i(\pi_i, a_j) = \sum_{a_i \in A_i} \pi_i(a_i) \mathcal{R}_i(a_i, a_j). \quad (6.46)$$

Therefore, the gradient of this expected reward with respect to policy π_i is simply the vector of rewards for each of agent i 's available actions 1, 2, 3...,

$$\nabla_{\pi_i} U_i(\pi_i, a_j) = \left[\frac{\partial U_i(\pi_i, a_j)}{\partial \pi_i(1)}, \frac{\partial U_i(\pi_i, a_j)}{\partial \pi_i(2)}, \frac{\partial U_i(\pi_i, a_j)}{\partial \pi_i(3)}, \dots \right] \quad (6.47)$$

$$= [\mathcal{R}_i(1, a_j), \mathcal{R}_i(2, a_j), \mathcal{R}_i(3, a_j), \dots]. \quad (6.48)$$

Given policy π_i^k and observed action a_j^k in episode k , GIGA updates π_i^k via two steps:

$$\begin{aligned} (1) \quad & \tilde{\pi}_i^{k+1} \leftarrow \pi_i^k + \kappa^k \nabla_{\pi_i^k} U_i(\pi_i^k, a_j^k) \\ (2) \quad & \pi_i^{k+1} \leftarrow P(\tilde{\pi}_i^{k+1}) \end{aligned} \quad (6.49)$$

where κ^k is the step size. Step (1) updates the policy in the direction of the unconstrained gradient $\nabla_{\pi_i^k} U_i(\pi_i^k, a_j^k)$, while Step (2) projects the result back into a valid probability space via the projection operator

$$P(x) = \arg \min_{x' \in \Delta(A_i)} \|x - x'\| \quad (6.50)$$

where $\|\cdot\|$ is the standard L2-norm.⁵⁰ $P(x)$ projects a vector x back into the space of probability distributions $\Delta(A_i)$ defined over the action set A_i .

Zinkevich (2003) showed that if all agents use GIGA to learn policies with a step size of $\kappa^k = \frac{1}{\sqrt{k}}$, then in the limit of $k \rightarrow \infty$ the policies will achieve no-regret. Specifically, recall the definition of regret in Equation 4.27 (page 77), then it can be shown that the regret for agent i is bounded by

$$\text{Regret}_i^k \leq \sqrt{k} + \left(\sqrt{k} - \frac{1}{2} \right) |A_i|r_{\max}^2 \quad (6.51)$$

50. The L2-norm is defined as $\|x\| = \sqrt{x \cdot x}$, where \cdot is the dot product.

where r_{\max} denotes the maximum possible reward for agent i . Thus, the average regret $\frac{1}{k} \text{Regret}_i^k$ will go to zero for $k \rightarrow \infty$ (since k grows faster than \sqrt{k}), satisfying the no-regret criterion given in Definition 12.

Finally, note that the above definitions also work for games with more than two agents, since we can replace j by $-i$ to represent a collection of other agents which accordingly choose joint actions a_{-i} . All of the above definitions and results still hold for this case.

II MULTI-AGENT DEEP REINFORCEMENT LEARNING: ALGORITHMS AND PRACTICE

Part II of this book will build upon the foundations introduced in Part I and present MARL algorithms which use deep learning to represent value functions and agent policies. As we will see, deep learning is a powerful tool which enables MARL to scale to more complex problems than is possible with tabular methods. The chapters in this part will introduce the basic concepts of deep learning, and show how deep learning techniques can be integrated into RL and MARL to produce powerful learning algorithms.

Chapter 7 provides an introduction to deep learning, including the building blocks of neural networks, foundational architectures, and the components of gradient-based optimisation used to train neural networks. This chapter primarily serves as a concise introduction to deep learning for readers unfamiliar with the field, and explains all foundational concepts required to understand the following chapters. Chapter 8 then introduces deep RL, explaining how to use neural networks to learn value functions and policies for RL algorithms.

Chapter 9 builds on the previous chapters and introduces multi-agent deep RL algorithms. The chapter begins by discussing different modes of training and execution which determine the information available to agents during training and execution, and how this information affects the learning process. The chapter then revisits the class of independent learning MARL algorithms with deep learning, before presenting advanced topics including multi-agent policy gradient algorithms and value decomposition. The chapter also shows how agents can share parameters and experiences in multi-agent deep RL algorithms to further improve the learning efficiency. Finally, the chapter considers the setting of competitive zero-sum games and how self-play can be used with deep learning to solve such games. Chapters 10 and 11 conclude this part of the book by discussing practical considerations for the implementation of MARL algorithms, and presenting environments which can serve as benchmarks and playground to study these algorithms.

7 Deep Learning

In this chapter, we provide a concise introduction to *deep learning*, a learning framework for function approximation. The chapter begins by motivating why we need function approximation in RL and MARL to solve complex environments. We will introduce feedforward neural networks as the foundational architecture of neural networks. The chapter then introduces gradient-based optimisation techniques as the main approach to train neural networks, before introducing advanced architectures for high-dimensional and sequential inputs. This chapter only covers the fundamental concepts of deep learning and is not meant to be a comprehensive or complete summary of this vast field. We refer the interested reader to the textbook by Goodfellow, Bengio, and Courville (2016) for a more comprehensive overview of deep learning. Chapters 8 and 9 will build on the present chapter and show how deep learning can be used in RL and MARL.

7.1 Function Approximation for Reinforcement Learning

Before discussing *what* deep learning is and *how* it works, it is useful to see *why* deep learning is omnipresent in RL research nowadays. What does deep learning offer over other techniques used to learn value functions, policies, and other models in RL?

Part I introduced MARL algorithms in their classical form, using tabular methods to represent value functions and policies of agents. These methods are referred to as “tabular” because their value function can be thought of as a large table with each state-action pair corresponding to a single table entry containing the value estimate for that particular input to the value function. This representation of tabular MARL has two important limitations. First, the table grows linearly with the number of possible inputs to the represented value function, rendering tabular MARL infeasible for complex problems like Go,

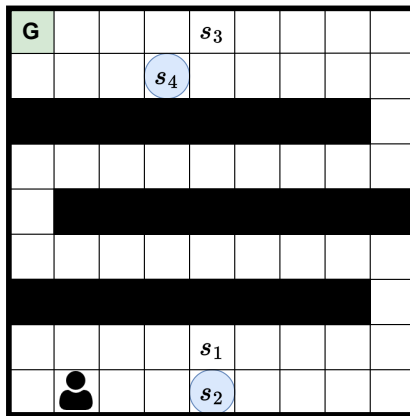


Figure 7.1: A maze environment in which a single agent must reach a goal location (G). The agent has seen several states, including s_1 and s_3 during its trajectories, and tries to estimate values for the circled states s_2 and s_4 . An agent with a tabular value function cannot generalise in this manner and would need to explicitly experience trajectories with s_2 and s_4 to be able to learn accurate value estimates. In contrast, a value function using function approximation techniques (such as linear value functions or deep learning) may be able to generalise and estimate the values of s_2 and s_4 to be similar to s_1 and s_3 , respectively.

video games, and most real-world applications. For example, the state space of the board game of Go is estimated to contain approximately 10^{170} possible states. Storing, managing and updating a table with this number of entries is completely infeasible for modern computers. Second, tabular value functions update each value estimate in isolation. For tabular MARL, an agent will only update its value estimate for the state s after applying an action in state s , and leave its value function for any other inputs unchanged. This consistency with prior value estimates comes with theoretical guarantees for tabular MARL algorithms, but means that an agent has to encounter many states before being able to learn their values and hence identify a desirable policy. For learning in complex tasks, it is therefore essential that agents have the ability to *generalise* to novel states. Encountering a specific state should ideally allow an agent to update not just its value estimate for this particular state, but also update its estimates for different states which are in some sense similar to the encountered state.

To illustrate the benefits of generalisation, consider the following example depicted in Figure 7.1. In this single-agent environment, the agent needs to

navigate a maze and reach the goal location (G) to receive positive reward. Given this reward function, the expected discounted returns given an optimal policy in a state depend on the distance of a state to the goal. Hence, two states which are similar in their distance to the goal should have similar value estimates. Considering the circled states in Figure 7.1, we would expect the value estimate for state s_2 to be similar to the value of s_1 . Likewise, the value of s_4 should be similar to the value of s_3 . Tabular value functions would need to be trained on each of these states in isolation, but function approximation offers value functions which can generalise and, thereby, pick-up on the relationships of states to their respective values to enable reasonable value estimates for s_2 and s_4 *before* ever encountering these states.

7.2 Linear Function Approximation

Given these limitations of tabular MARL, there is a clear need for value functions which generalise across states. Function approximation addresses both of these limitations and can be applied to represent value functions and policies of MARL agents. One possibility is to represent these functions as linear function approximators defined over pre-defined features of states. For instance, a linear state-value function can be written as

$$\hat{v}(s; \theta) = \theta^\top x(s) = \sum_{i=1}^d \theta_i x_i(s) \quad (7.1)$$

with $\theta \in \mathbb{R}^d$ and $x(s) \in \mathbb{R}^d$ denoting the vector of parameters and state feature vector, respectively. The state feature vector represents a predetermined encoding of states into a d -dimensional vector. An example for such an encoding of states would be a vector of polynomials where each entry represents a combination of values that make up the full state up to a fixed degree. During training, the value function can be continually updated by updating its parameters θ , typically following gradient-based optimisation techniques. Section 7.4 will provide a more detailed description of gradient-based optimisation (for the more involved case of deep learning, which includes linear function approximation), but in a nutshell we search for parameters θ which minimise an objective function. Consider the hypothetical example in which we aim to learn a linear state-value function and already knew the true values $v_\pi(s)$ for several states. We might minimise the mean squared error between the approximate value function $\hat{v}(s; \theta)$ and true values $v_\pi(s)$:

$$\theta^* = \arg \min_{\theta} \mathbb{E}_{s \in S} \left[(v_\pi(s) - \hat{v}(s; \theta))^2 \right] \quad (7.2)$$

In order to optimise the parameters θ and get closer to the optimal parameters θ^* which minimise the mean squared error, we can compute the gradient of the error with respect to θ and follow this gradient “downwards”.

The main benefit of linear value functions is their simplicity and ability to generalise. However, linear function approximation heavily relies on the selection of state features represented in $x(s)$, as $\hat{v}(s; \theta)$ is constrained to be a linear function with respect to these state features. Finding such state features can be non-trivial, often requires domain knowledge and features need to be carefully chosen depending on the task. In particular for environments with high-dimensional state representations, e.g. including images or language which might occur in MARL applications such as robotics and autonomous cars, finding appropriate features such that the desired value function can be represented as a linear function is very difficult.

In contrast to linear function approximation, deep learning provides a universal method for function approximation which is able to automatically learn feature representations of states, generalises to novel states, and is able to represent non-linear, complex functions. In the following sections, we will introduce the fundamental building blocks of deep learning and their optimisation process before discussing their concrete application to RL and MARL in Chapter 8 and Chapter 9 respectively.

7.3 Feedforward Neural Networks

Deep learning encompasses a family of machine learning techniques which apply *neural networks* as function approximators. These networks consist of many units organised in sequential layers with each unit computing comparably simple but non-linear calculations (visualised in Figure 7.2), thereby allowing neural networks to approximate complex functions which cannot be represented with linear function approximation. Neural networks are very flexible and have emerged as the most prominent type of function approximators in machine learning. In the following sections of this chapter, we will explain how neural networks work and how we can optimise these highly complex function approximators.

Feedforward neural networks – also called deep feedforward networks, fully-connected neural networks, or multilayer perceptrons (MLPs) – are the most prominent architecture of neural networks and can be considered the building block of most neural networks. As a general-purpose function approximator, a neural network learns a function $f(x; \theta)$ for some input $x \in \mathbb{R}^{d_x}$. Similar to linear function approximation, we denote the parameters of the function $\theta \in \mathbb{R}^d$ with

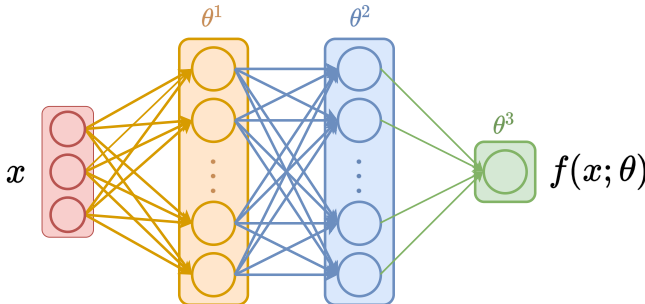


Figure 7.2: Illustration of a feedforward neural network with three layers. The input $x \in \mathbb{R}^3$ is processed by two hidden layers (orange and blue) with parameters θ^1 and θ^2 , respectively, and a single scalar output is computed in the final output layer (green) with parameters θ^3 .

d corresponding to the total number of learnable parameters. *Training* a neural network involves an optimisation process to find a set of parameter values θ such that f accurately approximates a target function f^* :

$$\forall x : f(x; \theta) \approx f^*(x) \quad (7.3)$$

For example, in RL f^* might be a value function representing the expected returns for a given state s as input.

Feedforward neural networks are structured into multiple sequential *layers* with the first layer processing the given input x and any subsequent layer processing the output of the previous layer. Each layer is defined as a parameterised function and is composed of many units with the output of the overall layer consisting of the concatenation of its units' outputs. An entire feedforward neural network is then defined by the composition of the functions of all layers by sequentially feeding the output of the previous layer into the next layer. For example, a feedforward neural network with three layers, visualised in Figure 7.2, can be written as

$$f(x; \theta) = f_3(f_2(f_1(x; \theta^1); \theta^2); \theta^3) \quad (7.4)$$

where θ^i corresponds to the parameters of layer i and $\theta = \bigcup_i \theta^i$ denotes the parameters of the entire network. The number of layers is referred to as the *depth* of the neural network and the number of units in a layer is sometimes referred to as the *width* of the layer. The generality and representational capacity, i.e. the complexity of functions the network can represent, largely depend on their depth and the width of each layer. We further denote the number of units in the i^{th} layer with d_i . To understand more deeply how these networks

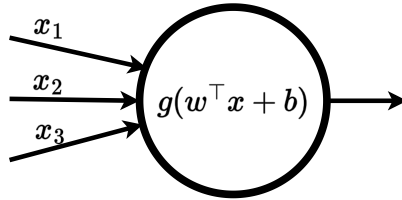


Figure 7.3: Illustration of a single neural unit computing a scalar output given an input $x \in \mathbb{R}^3$. First, a weighted sum over the input features is computed using the vector multiplication of $w \in \mathbb{R}^3$ and the input x and a scalar bias $b \in \mathbb{R}$ is added. Finally, a non-linear activation function $g : \mathbb{R} \mapsto \mathbb{R}$ is applied to obtain the scalar output.

approximate the overall function $f(x; \theta)$, we will first explain individual units within layers before explaining how to compose all these pieces together to form the entire network.

7.3.1 Neural Unit

An individual unit of a neural network in layer i represents a parameterised function $f_{i,u} : \mathbb{R}^{d_{i-1}} \mapsto \mathbb{R}$ which processes the output of the previous layer given by the output of each of its d_{i-1} th units. Note, for a unit in the first layer the input $x \in \mathbb{R}^{d_x}$ is used instead. First, a unit computes a weighted sum over its input features using a weight vector $w \in \mathbb{R}^{d_{i-1}}$ and adding a scalar bias $b \in \mathbb{R}$ to the weighted sum. This first computation represents a linear transformation and is followed by a non-linear *activation function* g_i . The entire computation of a neural unit is visualised in Figure 7.3 and can be formalised as

$$f_{i,u}(x; \theta_u^i) = g_i(w^\top x + b) \quad (7.5)$$

with parameters $\theta_u^i \in \mathbb{R}^{d_{i-1}+1}$ containing the weight vector w as well as the scalar bias b . Applying non-linear activation functions to the output of each unit is essential, because the composition of two linear functions, f and g , can only represent a linear function itself and hence non-linear functions could not be approximated by composing an arbitrary number of neural units without non-linear activation functions. During the optimisation of a feedforward neural network, the parameters θ_u^i of each unit in the network are optimised.

7.3.2 Activation Functions

There are many choices of activation functions g to apply in neural units. The most common activation function is the *rectified linear unit*, short ReLU (Jarrett

Name	Equation	Hyperparameters
Rectified linear unit (ReLU)	$\max(0, x)$	None
Leaky ReLU	$\max(cx, x)$	$0 < c < 1$
Exponential linear unit (ELU)	$\begin{cases} x & \text{if } x > 0 \\ \alpha(e^x - 1) & \text{if } x \leq 0 \end{cases}$	$\alpha > 0$
Hyperbolic tangent	$\tanh(x) = \frac{e^{2x}-1}{e^{2x}+1}$	None
Sigmoid	$\frac{1}{1+e^{-x}}$	None

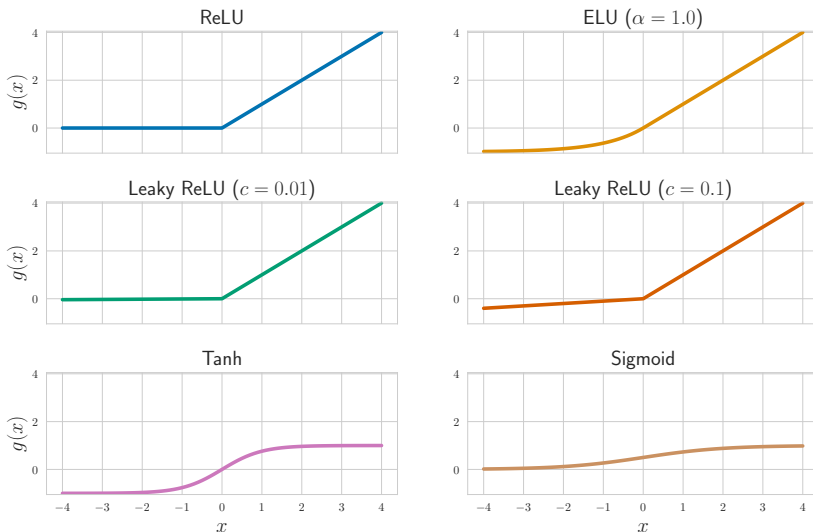
Figure 7.4: Summary of commonly-used activation functions for input $x \in \mathbb{R}$.

Figure 7.5: Common non-linear activation functions defined in Figure 7.4.

et al. 2009; Nair and Hinton 2010). ReLU applies a non-linear transformation but remains “close to linear”. This has useful implications for gradient-based optimisation used to update neural network parameters. Also, ReLU is able to output zero values instead of outputting values close to zero, which can be desirable for representational sparsity and computational efficiency. Other activation functions which are still commonly applied in feedforward neural networks, are \tanh , sigmoid, and several variations of the ReLU function such as leaky ReLU and exponential linear units (ELU). The previously mentioned activation functions are defined in Figure 7.4. For a visualisation and comparison of these activation functions, see Figure 7.5. The \tanh and sigmoid activation

functions are most commonly applied to restrict the output of a neural network to be within the ranges $(-1, 1)$ or $(0, 1)$, respectively. If no such constraints are required, ReLU and variations of the ReLU function are most commonly applied as default activation functions.

7.3.3 Composing a Network from Layers and Units

A feedforward neural network is composed of several layers (Figure 7.2) and each layer itself is composed of many neural units, all of which represent a parameterised function (Section 7.3.1). The i^{th} layer of a feedforward neural network receives the output of the previous layer $x_{i-1} \in \mathbb{R}^{d_{i-1}}$ as input and itself computes an output $x_i \in \mathbb{R}^{d_i}$. By aggregating all neural units of the i^{th} layer, we can write its computation as

$$f_i(x_{i-1}; \theta^i) = g_i(W_i^\top x_{i-1} + b_i) \quad (7.6)$$

with activation function g_i , weight matrix $W_i \in \mathbb{R}^{d_{i-1} \times d_i}$, and bias vector $b_i \in \mathbb{R}^{d_i}$. The parameters of the layer i consist of the weight matrix as well as the bias vector $\theta^i = W_i \cup b_i$. Note that the computation of the layer can be seen as the parallel computation of its d_i neural units and aggregating their outputs to a vector x_i . The weight matrix and bias vector contain the weight vectors and bias of each of the neural units within the layer, respectively, as their column entries. Considering the calculation of a single layer in this vectorised way naturally gives rise to an efficient computation and alternative interpretation of the operation of a single layer: The high-dimensional linear transformation of a single layer can also be described as a a single matrix multiplication with the weight matrix, followed by a vector addition with the bias vector and lastly applying the activation function element-wise to the resulting vector.

7.4 Gradient-Based Optimisation

The parameters θ of a neural network, sometimes referred to as *weights* of the network, have to be optimised to obtain a function f which accurately represents a target function f^* . A neural network may have many parameters. For example, recent advances in large language models like the GPT family of models (Brown et al. 2020) trained neural networks with many billions of parameters, so an efficient and automated process for the optimisation of their parameters is needed. Neural networks are complex non-linear functions and there exists no general closed-form solution to find their optimal parameters with respect to an optimisation objective. Instead, non-convex gradient-based optimisation methods can be used. These methods randomly initialise and

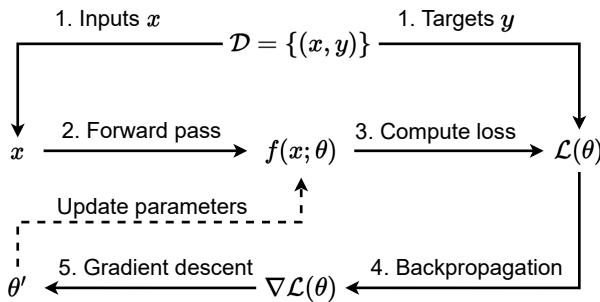


Figure 7.6: The training loop for gradient-based optimisation of a neural network parameterised by θ : 1. From a dataset D of input-output pairs (x, y) , a batch is sampled. 2. The prediction $f(x; \theta)$ is computed for each input x in the batch. 3. The loss function \mathcal{L} is computed between the predictions $f(x; \theta)$ and the target values y . 4. The gradients of the loss function with respect to the network parameters θ are computed using backpropagation. 5. The parameters θ are updated using a gradient-based optimiser. Then the loop starts again with the updated parameters θ .

sequentially update the parameters of a neural network to find parameters which improve the approximation of the neural network. In the following sections, we will explain the three key components for gradient-based optimisation:

1. *Loss function*: An optimisation objective which needs to be minimised and is defined over the network parameters θ .
2. *Gradient-based optimiser*: The choice of gradient-based optimisation technique.
3. *Backpropagation*: A technique to efficiently compute gradients of the loss function with respect to the network parameters θ , including the weights and bias of each unit.

The training loop of a neural network is illustrated in Figure 7.6 and the following sections will explain each of the three components in detail.

7.4.1 Loss Function

Our objective during the optimisation is to obtain parameters θ^* such that the loss function, denoted with \mathcal{L} , is minimised:

$$\theta^* \in \arg \min_{\theta} \mathcal{L}(\theta) \quad (7.7)$$

It is essential for this loss to be differentiable, because we need to compute gradients of the loss function to enable gradient-based optimisation of the parameters θ . The choice of loss function is dependent on the type of function the neural network approximates and the available data used for the optimisation.

To go back to our example of linear function approximation in Section 7.2, the network may be optimised to approximate a state-value function $\hat{v}(s; \theta)$ for RL. This value function should accurately approximate the true state-value function $v_\pi(s)$ for any state s under the current policy π . For this optimisation objective, we can define the loss as the mean squared error (MSE) of our approximated state-value function and the true state-value function for a batch of M states:

$$\mathcal{L}(\theta) = \frac{1}{M} \sum_{i=1}^M (v_\pi(s_i) - \hat{v}(s_i; \theta))^2 \quad (7.8)$$

The data used for the optimisation in this case contains M states $\{s_i\}_{i=1}^M$ with their corresponding true state values $\{v_\pi(s_i)\}_{i=1}^M$. Minimising this loss will gradually update the parameters θ of our neural network, representing \hat{v} , such that it approximates the true state-value function more closely for all states in the training data. The MSE is a commonly used loss function which can be applied to any setting in which pairs of inputs and ground truth outputs for the approximated function, here states with their true state values $\{(s_i, v_\pi(s_i))\}_{i=1}^M$, are available for training. This setting is commonly referred to as *supervised learning*. However, note that in RL we typically do not know the true state-value function a priori. Fortunately, temporal difference learning gives us a framework with which we can formulate a loss for our value function to approximate discounted state-value estimates using bootstrapped value estimates (Section 2.6):

$$\mathcal{L}(\theta) = \frac{1}{M} \sum_{i=1}^M (r_i + \gamma \hat{v}(s'_i; \theta) - \hat{v}(s_i; \theta))^2 \quad (7.9)$$

In this loss, we make use of a batch of experience of the agent consisting of state s , reward r , and next state s' , respectively. Minimising this loss will optimise our network parameters θ , such that \hat{v} will gradually provide accurate state-value estimates. Later in Chapter 8, we will discuss details on RL using neural networks with concrete algorithms, examples, and performance comparisons.

7.4.2 Gradient Descent

A common optimisation technique to update parameters in neural networks is *gradient descent*. Gradient descent sequentially updates parameters θ by following the negative gradients of the loss function with respect to the parameters for given data. This technique is similar to the gradient ascent in expected returns for policy learning we already saw in Section 6.4. The gradient $\nabla_\theta \mathcal{L}(\theta)$

is defined as the vector of partial derivatives for each parameter $\theta_i \in \theta$

$$\nabla_{\theta} \mathcal{L}(\theta) = \left(\frac{\partial \mathcal{L}(\theta)}{\partial \theta_1}, \dots, \frac{\partial \mathcal{L}(\theta)}{\partial \theta_d} \right) \quad (7.10)$$

and can be interpreted as the vector in the space of all parameters which points in the direction where our loss function increases the fastest. As we want to minimise the loss function, we can follow the negative gradient to update our parameters in the direction of the steepest descent. In the simplest case of *vanilla gradient descent*, network parameters are updated as

$$\theta \leftarrow \theta - \alpha \nabla_{\theta} \mathcal{L}(\theta | \mathcal{D}) \quad (7.11)$$

with *learning rate* $\alpha > 0$ typically taking on small values between $1e^{-5}$ and $1e^{-2}$. Vanilla gradient descent, also called batch gradient descent, computes a single gradient for the entire training data \mathcal{D} to update the parameters. This application of gradient descent comes with two main downsides. First, the training data will often not fit into memory in its entirety which makes the computation of the gradient for the entire training data difficult. Second, computing the gradient for the entire dataset for a single update of the parameters is costly and, hence, vanilla gradient descent is slow to converge to a local optima.

Stochastic gradient descent (SGD) addresses these difficulties by following the gradient for any individual sample d drawn from the training data:

$$\theta \leftarrow \theta - \alpha \nabla_{\theta} \mathcal{L}(\theta | d) |_{d \sim \mathcal{D}} \quad (7.12)$$

SGD is significantly faster to compute as it only requires computing the gradient for a single sample from the training data, but its updates exhibit high variance due to the dependency on the sample drawn.

Mini-batch gradient descent lies in between these two extremes of vanilla and stochastic gradient descent. Instead of using the entire dataset or individual samples to compute gradients, mini-batch gradient descent uses, as the name suggests, batches of samples from the training data to compute gradients:

$$\theta \leftarrow \theta - \alpha \nabla_{\theta} \mathcal{L}(\theta | \mathcal{B}) |_{\mathcal{B}=\{d_i \sim \mathcal{D}\}_{i=1}^B} \quad (7.13)$$

The number of samples in a batch \mathcal{B} used for each gradient computation, called the *batch size* B , is chosen as a hyperparameter and offers a trade-off between variance of gradients and computational cost. For small batch sizes, mini-batch gradient descent approaches SGD with fast computation of gradients but high variance. For larger batch sizes, mini-batch gradient descent approaches vanilla gradient descent with slower computation of gradients but lower variance of gradients and hence stable convergence. Common batch sizes for optimisation of neural networks lie between 32 and 1028, but the batch size should always

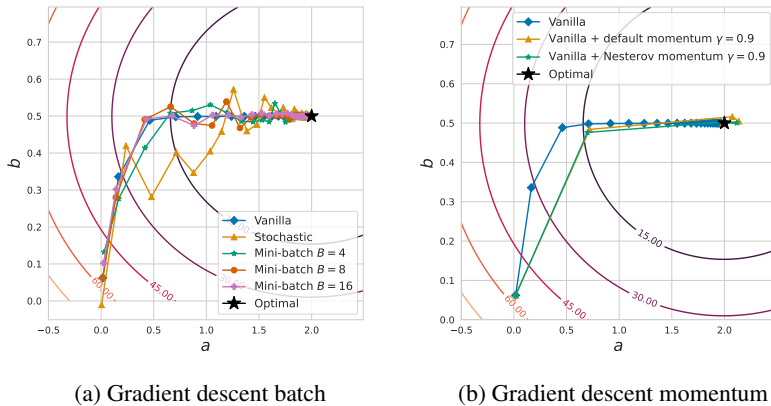


Figure 7.7: Contour plots showing the optimisation of a function approximation to fit a polynomial function with gradient-based optimisation. The concentric circles represent loss values given by the mean-squared error. Figure (a) compares vanilla, stochastic, and mini-batch gradient descent with various batch sizes, and (b) compares vanilla gradient descent with and without momentum.

be carefully chosen depending on the available data, computational resources, neural network architecture, and loss function.

We show a comparison of vanilla gradient descent, SGD, and mini-batch gradient descent with varying batch sizes for the optimisation of a polynomial function in Figure 7.7a. In this example, we train a function approximator $f(x; a, b) = ax + bx^2$ to approximate a target function $f^*(x; a^* = 2, b^* = 0.5)$. For training, we generate a dataset $\mathcal{D} = \{(x, f^*(x))\}$ for 1000 input values $x \in [-5, 5]$. For all optimisation, we use $\alpha = 5e^{-4}$ and initial parameters $a = b = 0$. The results indicate that the optimisation with vanilla gradient descent is very stable but it is important to remember that each update with vanilla gradient descent is computationally expensive. In contrast, stochastic gradient descent is computationally cheap but its optimisation is less stable due to the high variance of its gradients. Mini-batch gradient descent provides an appealing trade-off of both these approaches. Even with a small batch size of $B = 4$, we see that mini-batch gradient descent approaches the stability of vanilla gradient descent for a fraction of its computational cost. We note that in the deep learning literature, mini-batch gradient descent is sometimes also referred to as just SGD due to its approach of approximating the expected gradient of the entire training data using samples drawn from the data.

Many gradient-based optimisation techniques have been proposed to extend mini-batch gradient descent. One common concept is the idea of *momentum* (Polyak 1964; Nesterov 1983) in which a moving average over past gradients is computed and added to the gradients to “accelerate” optimisation. Momentum significantly speeds up gradient-based optimisation as long as it continues in a similar direction, but also increases the risk of “overshooting” local minima of the loss function, as seen by default momentum in Figure 7.7b. For a comparison of vanilla gradient descent with and without two types of momentum, we show contour plots of the optimisation of a polynomial function, as discussed before, in Figure 7.7b. We can see significant improvements in the efficiency of gradient descent when using momentum, with significantly fewer updates being needed to obtain parameters close to the ground truth parameters. Also, we observe that Nesterov momentum is considerably more stable than traditional momentum. Several more recent approaches follow the idea of a dynamically adapting the learning rate during optimisation which can be considered similar to momentum. Learning rate adaptation has the main benefits of simplifying the process of choosing the initial learning rate as a hyperparameter, making the optimisation less sensitive to the choice of the (initial) learning rate, and to speed-up the optimisation. All gradient-based optimisers, which are commonly used to optimise the parameters of neural networks, apply different learning rate adaptation schemes (Duchi, Hazan, and Singer 2011; Hinton, Srivastava, and Swersky 2012; Zeiler 2012; Kingma and Ba 2015). None of the optimisers is consistently observed to perform better than the others, but Adam (Kingma and Ba 2015) has emerged as the most prominently used optimiser and is often applied as the default choice to optimise neural networks.⁵¹

7.4.3 Backpropagation

Gradient-based optimisers require the gradients $\nabla_{\theta}\mathcal{L}(\theta)$ of the loss with respect to all parameters of the network. These gradients are needed to understand how we should change each parameter to minimise the loss. The computation of the output of the neural network for some given input is often referred to as a *forward pass* because it passes outputs of layers sequentially forward through the network. To compute gradients of the loss function with respect to all parameters in the network, i.e. including the parameters of every layer within the network, the *backpropagation* algorithm (Rumelhart, Hinton, and Williams 1986) is used. This algorithm considers the fact that a neural network computes

51. For a more detailed overview of gradient-based optimisation techniques, we refer the interested reader to Ruder (2016).

a compositional function for which the chain rule of derivatives can be applied. As a reminder, the chain rule states for $y = g(x)$ and $z = f(y) = f(g(x))$

$$\nabla_x z = \left(\frac{\partial y}{\partial x} \right)^\top \nabla_y z \quad (7.14)$$

with $\frac{\partial y}{\partial x}$ being the Jacobian matrix of function g . In words, the chain rule allows us to compute the gradients of a compositional function with respect to the inputs of the inner function as a multiplication of the Jacobian of the inner function g and the gradient of the outer function with respect to its inputs.

As discussed in Section 7.3, feedforward neural networks are compositional functions with each layer defining its own parameterised function consisting of a non-linear activation function, matrix multiplication and vector addition as defined in Equation 7.6. By computing the inner gradients, $\nabla_y z$ for each inner operation computed throughout the network, the gradients of the parameters can be efficiently computed with respect to its parameters for some given input by traversing the network from its output layer to the input layer. Throughout the network, the chain rule is applied to propagate gradients backwards until the inputs are reached. Computing gradients with respect to every parameter of the network in a single process backwards from the last to the first layer is also referred to as a *backwards pass*. The backpropagation algorithm has been implemented using computational techniques of automatic differentiation in any major deep learning framework. Thanks to these techniques, the details of the backpropagation algorithm are hidden away and computing gradients of neural networks is as simple as calling a function in the respective framework.

7.5 Convolutional and Recurrent Neural Networks

Feedforward neural networks can be considered the backbone of deep learning and are universally applicable to any type of data, but many specialised architectures exist which build upon the idea of feedforward neural networks. The most common specialised architectures also found in RL and MARL algorithms are *convolutional neural networks* and *recurrent neural networks*. Both of these architectures are designed for specific types of inputs and, hence, suitable for particular types of problems. Convolutional neural networks are specifically constructed to process spatially structured data, in particular images. Recurrent neural networks are designed to process sequences, in RL most commonly histories of observations in partially observable environments.

7.5.1 Learning from Images – Exploiting Spatial Relationships in Data

Feedforward neural networks can be used to process any inputs, but are not well suited for processing spatial data such as images for two primary reasons. First, to process an image with a feedforward neural network, the image representation would need to be flattened from a multi-dimensional matrix $x \in \mathbb{R}^{c \times h \times w}$ where c, h and w correspond to the number of colour channels (many images are represented with three colour channels corresponding to the colours red, blue and green), height, and width of the image. The dimensionality of this flattened image vector $\tilde{x} \in \mathbb{R}^{c \cdot h \cdot w}$ corresponds to the number of pixels in the image, multiplied by the number of colour channels. Processing such large input vectors with feedforward neural networks would require the first layer to contain many parameters, which makes the optimisation difficult and is hence undesirable. For example, consider a feedforward neural network which processes images of 128×128 pixels with RGB colours for three colour channels. Representing images of this size, which is significantly smaller than photos taken with any modern smartphone camera, corresponds to vectors of dimensionality $128 \times 128 \times 3 = 49,152$. With a small number of neural units of 128 in the first layer, the weight matrix would be a $49,152 \times 128$ dimensional matrix with a total of 6,291,456 parameters which need to be learned! While six million parameters is not a large neural network in comparison to large language models, it appears excessive and overly computationally expensive to require so many parameters to process even small images.⁵²

Secondly, images contain spatial relationships with pixels close to each other, often corresponding to similar objects shown in the image. Feedforward neural networks do not consider such spatial relationships and process every input value individually.

Convolutional neural networks (CNNs) (Fukushima and Miyake 1982; LeCun et al. 1989) directly make use of the spatial relationships within input data such as images by processing patches of nearby pixels at a time. Small groups of parameters called *filters* or *kernels* are “slid” over the image in a *convolution* operation. For the convolution, each filter moves through the pixels in each row of the image and encodes all pixel values it covers at a time. For each patch of pixels the filter moves over, its parameters are multiplied with the corresponding patch of pixels of the same size in a matrix multiplication to obtain a single output value. The patch of input values involved in a single convolution are also referred to as the *receptive field* of the corresponding

52. To process 4K RGP images with a resolution of 3840×2160 instead of 128×128 pixels, the weight matrix in the first layer of a feedforward neural network alone would contain more than 3 billion parameters!

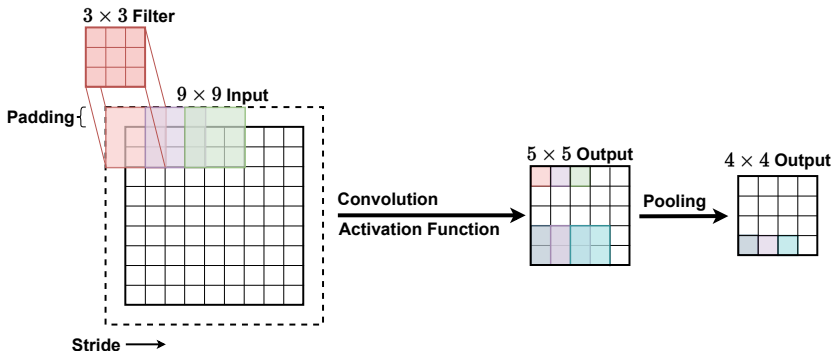


Figure 7.8: Illustration of a convolutional neural network with a single 3×3 kernel processing a 9×9 input. First, the kernel is applied to the image with padding one and a stride of two (kernel moving two pixels at a time) for a 5×5 output. Second, a pooling operation is applied to 2×2 groups of pixels with no padding and a stride of one for a final output of 4×4 . Note that the image is processed with only 10 learned parameters (9 weights and a scalar bias).

output value or neuron. Formally, the convolution of a filter with parameters $W \in \mathbb{R}^{d_w \times d_w}$ over input $x \in \mathbb{R}^{d_x \times d_x}$ for output $y_{i,j} \in \mathbb{R}$ is defined as follows:

$$y_{i,j} = \sum_{a=1}^{d_w} \sum_{b=1}^{d_w} W_{a,b} x_{i+a-1, j+b-1} \quad (7.15)$$

This operation is repeated until the filter has been moved over the entire image. Note that each filter has one parameter for each cell, which are re-used in every convolution operation. This sharing of parameters across multiple computations leads to significantly less parameters which need to be optimised, in contrast to a fully-connected network layer, and makes use of the spatial relationship of images with patches of nearby pixels being highly correlated. Filters move across the input depending on the *stride* and *padding*. The stride determines the number of pixels the filter moves at each slide, and padding refers to an increase in the image size, typically by adding zero values around its borders. These hyperparameters can be used to influence the dimensionality of the convolution output. Following the convolution, a non-linear activation function is applied element-wise to each cell of the output matrix.

CNNs often apply multiple such convolution operations in a sequence using varying sizes of filters. In between each convolution, it is common to additionally apply *pooling* operations. In these operations, patches of pixels are aggregated together using operations such as taking the maximum value within

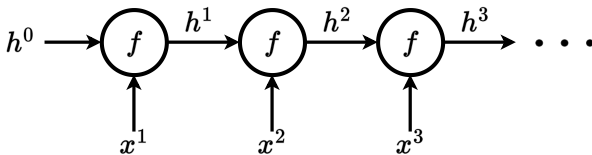


Figure 7.9: Illustration of a recurrent neural network which represents a function f used to process a sequence of inputs x^1, x^2, \dots . At time step t , the network takes both the current input x^t and the previous hidden state h^{t-1} as inputs to compute the new hidden state h^t . An initial hidden state h^0 is given to the model at the first step, and then the hidden state is continually updated by the recurrent neural network to process the input sequence.

the patch, referred to as max-pooling, or taking the average. These pooling operations in between convolutions further reduce the output dimensionality throughout the convolutional neural networks and make the output of the network insensitive to small local changes in the image. This effect is desirable because the learned locally-insensitive features might generalise better than features which are associated with a particular location in the image.

Figure 7.8 shows a convolutional neural network with a single kernel processing an input and applying pooling for aggregation. After processing highly spatial inputs, such as images, with several layers of CNNs, it is common to further process the obtained representations using highly parameterised feedforward neural networks.

7.5.2 Learning from Sequences with Memory

Similar to images, feedforward neural networks are ill-equipped to process sequential inputs. For example, in partially observable tasks an RL agent is conditioned on its history of observations (see Section 3.4). Conditioning a feedforward neural network on such a history requires the entire sequence as input. Similar to image inputs, processing a long sequence of observations with a feedforward neural network requires many parameters. Additionally, observations within a history will likely be correlated, thus sharing parameters to process each observation appears desirable, similar to filters being used many times in a convolutional neural network.

Recurrent neural networks (RNNs) (Rumelhart, Hinton, and Williams 1986) are neural networks specifically designed to process sequential data. Instead of using long concatenated sequences of inputs, recurrent neural networks

sequentially process inputs and additionally condition the computation at each step on a compact representation of the history of previous inputs. This compact representation, also called the *hidden state* of the recurrent neural network, is continually updated to encode more information as the sequence is processed and serves as a form of memory. In this way, the same neural network and computation can be applied at each time step, but its function continually adapts as the sequence is processed and the hidden state changes. Formally, the hidden state is given by the output of the network

$$h^t = f(x^t, h^{t-1}; \theta) \quad (7.16)$$

with the initial hidden state h^0 usually being initialised as a zero-valued vector. In line with common notation in deep learning, we denote the hidden state of a recurrent neural network with h , but highlight that elsewhere in the book h refers to the history of observations. Figure 7.9 illustrates the computational graph for a recurrent neural network processing a sequence of inputs. Some recurrent neural network architectures provide separate outputs for the updated hidden state and the main output of the network.

Optimising recurrent neural networks over long sequences is difficult, with gradients often vanishing (becoming close to zero) or exploding (becoming very large) due to the backpropagation repeatedly multiplying gradients and Jacobians of the network parameters throughout the sequence. Various approaches have been proposed to address this challenge, including skip connections (Lin et al. 1996), which add connections between computations across multiple time steps, and leaky units (Mozer 1991; El Hihi and Bengio 1995), which allow tuning the accumulation of previous time steps through linear self-connections. However, the most commonly applied and effective recurrent neural networks architectures are *long short-term memory cells* (LSTMs) (Hochreiter and Schmidhuber 1997) and *gated recurrent units* (GRUs) (Cho et al. 2014). Both approaches are based on the idea of allowing the recurrent neural network to decide when to try to remember or discard accumulated information within the hidden state.

8 Deep Reinforcement Learning

In Part I, we represented value functions with large tables. After encountering a particular state, only its corresponding value estimate, as given by its entry in the table, is updated and all other value estimates remain unchanged. This inability of tabular value functions to *generalise*, i.e. to update its value estimation for states similar but not identical to encountered states, severely limits the possible applications of tabular MARL. In problems with large or continuous state spaces, encountering any state multiple times is unlikely, so reaching accurate value estimates is infeasible. Chapter 7 introduced deep learning and neural networks with their universal optimisation techniques. RL and MARL can leverage these powerful function approximators to represent value functions which, unlike tabular value functions, can generalise to previously unseen states.

Before we discuss how MARL algorithms can make use of neural networks (Chapter 9), this chapter introduces the underlying techniques in the context of single-agent RL. It naturally builds upon the content of Chapters 2 and 7 and aims to answer the questions of how to effectively use neural networks to approximate value functions and policies. We first introduce how to use neural networks to approximate value functions. As we will see, the integration of deep learning into value functions for RL also introduces several challenges. In particular, the previously encountered moving target problem (Section 5.4.1) is further exacerbated when neural networks come into play, and neural networks tend to specialise to the most recent experience. We will discuss these difficulties, and how we can mitigate them. Then, we will discuss a new family of RL algorithms – policy gradient algorithms – which directly learn a policy represented as a parameterised function. We will introduce the theorem underlying these algorithms, introduce fundamental policy gradient algorithms, and discuss how to efficiently train these algorithms.

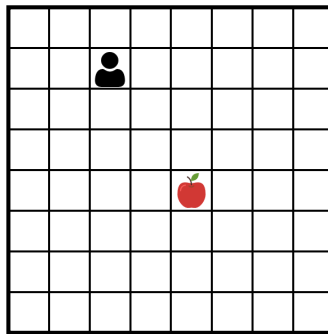


Figure 8.1: A simplified single-agent variant of the level-based foraging environment (Figure 1.2). The agent moves within the gridworld and has to pick-up a single, randomly located item.

8.1 Deep Value Function Approximation

Value functions are an essential component of most RL algorithms. This section presents fundamental ideas on how to leverage neural networks to approximate value functions. Perhaps surprisingly, introducing neural networks to represent value functions in RL introduces several challenges which did not previously occur with tabular value functions. In particular, extending common off-policy RL algorithms such as Q-learning (Section 2.6) with neural networks requires careful consideration. In this section, we piece-by-piece extend Q-learning, as introduced for tabular RL, to the commonly used deep RL algorithm of deep Q-networks (DQN) (Mnih et al. 2015). DQN is the first and one of the most frequently applied deep RL algorithms. Moreover, DQN serves as a common building block for many single- and multi-agent RL algorithms, so we will explain all its components in detail. We start from the Q-learning algorithm we are already familiar with and present each extension with its formalisation, pseudocode, and compare the resulting algorithms in a single-agent variant of the level-based foraging environment to study the impact of these extensions.

In this environment, visualised in Figure 8.1, the agent moves within an 8×8 gridworld to pick-up a single item.⁵³ The agent and the item are randomly placed at the beginning of each episode. To collect the item, and receive a reward of +1, the agent has to move next to the item and select its picking action.

⁵³ In contrast to the multi-agent level-based foraging environment introduced in Figure 1.2, the agent and item have no levels and pick-up will always be successful as long as the agent is located next to the item.

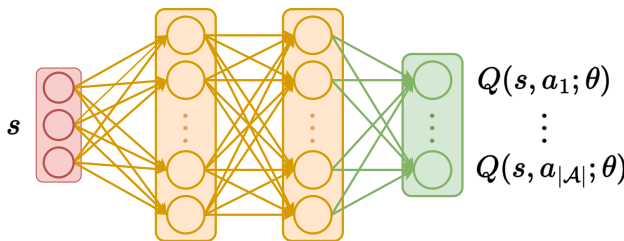


Figure 8.2: Neural network architecture for action-value functions: The network receives a state as its input and outputs action-value estimates for every possible (discrete) action.

For any other interaction, the agent receives a reward of 0. To formalise the algorithms for this chapter, we assume the environment to be represented by a fully-observable MDP (Section 2.2). We will briefly discuss the extension to partially-observable games in Section 8.3.

8.1.1 Deep Q-Learning – What Can Go Wrong?

In tabular Q-learning the agent maintains an action-value function Q in the form of a table. After taking action a^t in state s^t at time step t , the agent receives reward r^t and observes a new state s^{t+1} . Using this experience, the agent can use the Q-learning update rule to update its action-value function as follows

$$Q(s^t, a^t) \leftarrow Q(s^t, a^t) + \alpha \left(r^t + \gamma \max_{a'} Q(s^{t+1}, a') - Q(s^t, a^t) \right) \quad (8.1)$$

with α denoting the learning rate.

How can the tabular value function of Q-learning be substituted with a neural network? In Chapter 7, we saw that in order to train a neural network, we need to define the network architecture as well as the loss function, and choose a gradient-based optimiser. The choice of an optimiser is important and may significantly impact the learning process, as we discussed in Section 7.4, but neural networks can be optimised with any gradient-based optimiser. Therefore, we focus on defining the architecture and loss function. To this end, we define a neural network Q to represent the action-value function for a deep version of Q-learning, visualised in Figure 8.2. The network receives a state as its input and outputs its estimated action-value for each possible action. This architecture is computationally efficient by computing the action-value estimates for all actions in one forward pass through the network but also limits the deep Q-learning algorithm, just as tabular Q-learning, to environments with finite (discrete) action spaces. For the loss function, which is to be minimised by updating

Algorithm 10 Deep Q-Learning

- 1: Initialise value network Q with random parameters θ
 - 2: Repeat for every episode:
 - 3: **for** time step $t = 0, 1, 2, \dots$ **do**
 - 4: Observe current state s^t
 - 5: With probability ϵ : choose random joint action $a^t \in A$
 - 6: Else: choose $a^t \in \arg \max_a Q(s^t, a; \theta)$
 - 7: Apply action a^t ; observe rewards r^t and next state s^{t+1}
 - 8: Update parameters θ by minimising the loss from Equation 8.2
-

parameters θ , we define the squared error between the value function estimate and the target value using the same experience of a single transition as for tabular Q-learning:

$$\mathcal{L}(\theta) = \left(\underbrace{r^t + \gamma \max_{a'} Q(s^{t+1}, a'; \theta)}_{\text{target value}} - Q(s^t, a^t; \theta) \right)^2 \quad (8.2)$$

It is important to note that the value estimate which we would like to update using this loss function is the current action-value estimate for (s^t, a^t) , given by $Q(s^t, a^t; \theta)$. In contrast, the remaining term within the loss, $r^t + \gamma \max_{a'} Q(s^{t+1}, a'; \theta)$, constitutes the bootstrapped target value we aim to approximate. Hence, when computing the gradients of the loss with respect to θ , the backpropagation should only propagate through the value estimate for (s^t, a^t) and ignore the gradients of the bootstrapped target value.⁵⁴ Pseudocode for this algorithm with an ϵ -greedy exploration policy, which we will refer to as deep Q-learning, is shown in Algorithm 10. In the following sections and chapters, we will denote the parameters of neural networks which represent value functions with θ .

This simple extension of Q-learning suffers from two important issues. First, the moving target problem is further exacerbated by the application of an approximated value function, and the strong correlation of consecutive samples used to update the value function leads to an undesirable specialising of the network to the most recent experiences. The following two sections will focus on these challenges and how to address them for deep reinforcement learning.

⁵⁴ Deep learning libraries such as PyTorch, Tensorflow and Jax all provide functionality to restrict the gradient computation to specific components within the loss. This is an implementation detail, even though an important one, and not essential to understand the core components within the following algorithms.

8.1.2 Moving Target Problem

In Section 5.4.1, we have already seen that the learning of value functions in RL is challenging due to non-stationarity. This non-stationarity is caused by two factors. First, the policy of the agent is changing throughout training. Second, the target estimates are computed with bootstrapped value estimates of the next state which change as the value function is trained. This challenge, which we also refer to as the *moving target problem*, is further exacerbated in deep RL. In contrast to tabular value functions, value functions with function approximation, such as neural networks, generalise their value estimates across inputs. This generalisation is essential in order to apply value functions to complex environments but also introduces an additional challenge. Updating the value estimate for a single state also changes the value estimate in all other states. This can cause the bootstrapped target estimate to change significantly more rapidly than for tabular value functions, which can render optimisation unstable.

To reduce the instability of training caused by the moving target problem, we can equip the agent with an additional network. This so-called *target network* with parameters $\bar{\theta}$ is of the same architecture as the primary network representing the value function and is initialised with the same parameters. The target network can then be used instead of the primary network to compute bootstrapped target values in the following loss function:

$$\mathcal{L}(\theta) = \left(r^t + \gamma \max_{a'} \underbrace{Q(s^{t+1}, a'; \bar{\theta})}_{\text{target network value}} - \underbrace{Q(s^t, a^t; \theta)}_{\text{main network value}} \right)^2 \quad (8.3)$$

Instead of optimising the target network with gradient descent, the parameters of the target network are periodically updated by copying the current parameters of the value function network to the target network. This process ensures that the bootstrapped target values are not too far from the estimates of the main value function, but remain unchanged for a fixed number of updates and thereby increase the stability of target values.

8.1.3 Breaking Correlations

The second problem of naive deep Q-learning is the correlation of consecutive experiences. In many paradigms of machine learning, it is generally assumed that the data used to train the function approximation are *independent of each other and identically distributed* or “i.i.d. data” in short. This assumption guarantees that, firstly, there are no correlations of individual samples within the training data, and secondly, that all data points are sampled from the same

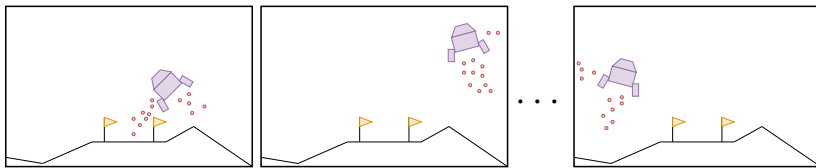


Figure 8.3: Illustration of the correlations of consecutive experiences in an environment where the agent controls a spaceship to land. In the first two episodes, the agent approaches the goal location from the right side. In the third episode, the agent has to approach the goal location from the left side and, thus, experiences very different states than in the previous episodes.

training distribution. Both of these components of the i.i.d. assumption are typically violated in RL. For the former, individual samples of experiences, given by tuples of state, action, reward, and next state, are clearly not independent and highly correlated. This correlation is obvious given the formalisation of an environment as an MDP. The experience at time step t is directly dependent on and, thus, not independent from the experience at time step $t - 1$, with s^t and r^t being determined by the transition function conditioned on s^t and a^t . Regarding the assumption of data points being sampled from the same training distribution, the distribution of encountered experiences in RL depends on the currently executed policy. Therefore, changes in the policy also cause changes in the distribution of experiences.

But why are we concerned about these correlations? What impact do these correlations have on the training of the agent using deep value functions? Consider the example illustrated in Figure 8.3, in which an agent controls a spaceship. For several timesteps, the agent receives similar sequences of experiences in which it approaches the goal location from the right direction (Section 8.1.3). The agent sequentially updates its value function parameters using these experiences, which may lead to the agent becoming specialised for these particular, recent experiences of approaching the goal location from the right side. Suppose that, after several such episodes, the agent is forced to approach the goal from the left side (Section 8.1.3), but throughout the previous updates has learned a value function which is specialised to states where the spaceship is located to the right of the goal location. This specialised value function may provide inaccurate value estimates for states where the spaceship is located to the left of the goal location and, thus, may fail to successfully land from this side. Moreover, updating the value function with the experience samples from this most recent episode might lead to the agent forgetting how

to approach the goal from the right side. This phenomenon is referred to as *catastrophic forgetting* and is a fundamental challenge of deep learning. This challenge is further exacerbated in RL since changes in the policy will lead to changes in the data distribution encountered by the agent, and changes in the data distribution might change what policy is optimal. This dependence can lead to oscillating or even diverging policies.

To address these issues, we can randomise the experience samples used to train the agent. Instead of using the sequential experiences as the agent receives them to update the value function, experience samples are collected in a so-called *replay buffer* \mathcal{D} . For training of the value function, a batch of experiences \mathcal{B} is sampled uniformly at random from the replay buffer, $\mathcal{B} \sim \mathcal{D}$. This sampling has two additional benefits for the optimisation of the value function: (1) experiences can be reused multiple times for training, which can improve sample efficiency, and (2) by computing the value loss over batches of experiences rather than for an individual experience sample, we can obtain a more stable gradient for the network optimisation with lower variance. During training, the mean squared error loss is computed over the batch and minimised to update the parameters of the value function:

$$\mathcal{L}(\theta) = \frac{1}{B} \sum_{(s^t, a^t, r^t, s^{t+1}) \in \mathcal{B}} \left(r^t + \gamma \max_{a'} Q(s^{t+1}, a'; \theta) - Q(s^t, a^t; \theta) \right)^2 \quad (8.4)$$

Formally, the replay buffer can be denoted as a set $\mathcal{D} = \{(s^t, a^t, r^t, s^{t+1})\}$ of experience samples. Typically, the replay buffer is implemented with a fixed capacity as a first-in-first-out queue, i.e. once the buffer is filled to its capacity with experience samples, the oldest experiences are continually replaced as new experience samples are added to the buffer. It is important to note that experiences within the buffer have been generated using the policy of the agent at earlier time steps during training. Therefore, these experiences are off-policy and, thus, a replay buffer can only be used to train off-policy RL algorithms such as algorithms based on Q-learning.

8.1.4 Putting It All Together: Deep Q-Networks

These ideas bring us to the first and one of the most influential deep RL algorithms: Deep Q-networks (DQN) (Mnih et al. 2015). DQN extends tabular Q-learning by introducing a neural network to approximate the action-value function, as shown in Figure 8.2. To address the challenges of moving target values and correlation of consecutive samples, DQN uses a target network and replay buffer, as discussed in Sections 8.1.2 and 8.1.3. All these ideas together define the DQN algorithm, as shown in Algorithm 11. The loss function is

Algorithm 11 Deep Q-Networks (DQN)

- 1: Initialise value network Q with random parameters θ
 - 2: Initialise target network with parameters $\bar{\theta} = \theta$
 - 3: Initialise an empty replay buffer $\mathcal{D} = \{ \}$
 - 4: Repeat for every episode:
 - 5: **for** time step $t = 0, 1, 2, \dots$ **do**
 - 6: Observe current state s^t
 - 7: With probability ϵ : choose random joint action $a^t \in A$
 - 8: Else: choose $a^t \in \arg \max_a Q(s^t, a; \theta)$
 - 9: Apply action a^t ; observe rewards r^t and next state s^{t+1}
 - 10: Store transition in replay buffer \mathcal{D}
 - 11: Sample random mini-batch of transitions from replay buffer \mathcal{D}
 - 12: Update parameters θ by minimising the loss from Equation 8.5
 - 13: In a set interval, update target network parameters $\bar{\theta}$
-

given as follows:

$$\mathcal{L}(\theta) = \frac{1}{B} \sum_{(s^t, a^t, r^t, s^{t+1}) \in \mathcal{B}} \left(r^t + \gamma \max_{a'} Q(s^{t+1}, a'; \bar{\theta}) - Q(s^t, a^t; \theta) \right)^2 \quad (8.5)$$

To see the impact of both the target network and replay buffer on the learning of the agent, we show the learning curves of four algorithms in the single-agent level-based foraging environment (Figure 8.1) in Figure 8.4: Deep Q-learning (Algorithm 10), deep Q-learning with target networks, deep Q-learning with a replay buffer, and the full DQN algorithm with a replay buffer and target networks (Algorithm 11). We see that training the agent with deep Q-learning leads to a slow and unstable increase in evaluation returns. Adding target networks leads to no notable improvement in performance. Training the agent with deep Q-learning and batches sampled from a replay buffer slightly increases evaluation returns in some runs, but performance remains noisy across runs. Finally, training the agent with the full DQN algorithm, i.e. using target networks and a replay buffer, leads to a stable and quick increase in performance and convergence to near-optimal discounted returns.

This experiment demonstrates that, in isolation, neither the addition of target networks nor of a replay buffer are sufficient to train the agent with deep Q-learning in this environment. Adding a target network reduces stability issues caused by the moving target problem, but the agent still receives highly correlated samples and therefore is unable to train its value function to generalise over all initial positions of the agent and item which are randomised at the

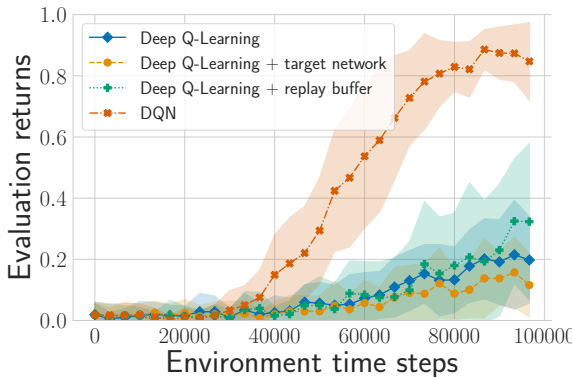


Figure 8.4: Learning curves for deep Q-learning, deep Q-learning with target networks, deep Q-learning with a replay buffer, and the full DQN algorithm in the single-agent level-based foraging environment shown in Figure 8.1. We train all algorithms for 100,000 time steps and in frequent intervals compute the average evaluation returns of the agent over 10 episodes using a near-greedy policy ($\epsilon = 0.05$). Visualised learning curves and shading correspond to the mean and standard deviation across discounted evaluation returns across five runs with different random seeds. To ensure consistency, we use identical hyperparameters: discount factor $\gamma = 0.99$, learning rate $\alpha = 3e^{-4}$, ϵ is decayed from 1.0 to 0.05 over half of training (50,000 time steps) and then kept constant, batch size $B = 512$ and buffer capacity is set to 10,000 experience tuples for algorithms with a replay buffer, and target networks are updated every 100 time steps where applied.

beginning of each episode. Training the agent with a replay buffer addresses the issue of correlated experiences, but without target networks suffers from unstable optimisation due to the moving target problem. Only the combination of both of these ideas in the DQN algorithm leads to a stable and fast learning process.

8.2 Policy Gradient Algorithms

So far in this chapter, we have discussed value-based RL algorithms. These algorithms learn a parameterised value function, represented by a neural network, and the agent follows a policy which is directly derived from this value function. As we will see, it can be desirable to directly learn a policy as a separate parameterised function. Such a parameterised policy can be represented by

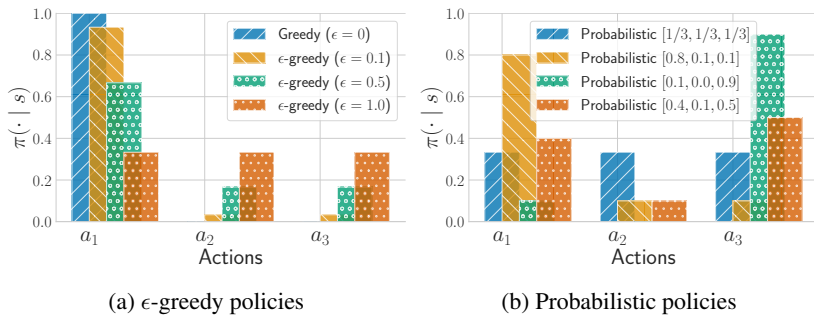


Figure 8.5: Illustration of the varying flexibility of (a) ϵ -greedy policies and (b) softmax policies (Equation 8.7) for a task with three discrete actions, i.e. $A = \{a_1, a_2, a_3\}$. (a) ϵ -greedy policies are limited to selecting the greedy action a_1 given the underlying action-value function with probability $1 - \epsilon$ and a uniform policy over all actions otherwise. In contrast, the softmax policies of policy gradient algorithms, shown in (b), can represent arbitrary distributions over all actions. For example in the multi-agent game of Rock-Paper-Scissors, the Nash equilibrium is for all agents to follow a uniform policy. The ϵ -greedy policy for any action-value function, however, is unable to represent such a uniform policy for $\epsilon < 1$ which may prevent the ability of agents to learn optimal policies for the environment.

any function approximation technique, most commonly using linear function approximation (see Section 7.2) or deep learning. In the RL literature, these algorithms are referred to as *policy gradient algorithms* because they compute gradients with respect to the parameters of their policy to update and continually improve the learned policy. In Section 6.4, we have already seen simple parameterised policies for MARL which are updated using gradient ascent. In this section, we will discuss more advanced policy gradient algorithms for single-agent RL which make use of neural networks to represent the policy, and in Chapter 9 we will extend the algorithms of this section for MARL.

8.2.1 Advantages of Learning a Policy

Directly representing the policy of an RL agent has two key advantages. Firstly, in environments with discrete actions, a parameterised policy can naturally represent any arbitrary probabilistic policy which leads to significantly more flexibility in its action selection compared to value-based RL algorithms. A

value-based RL agent following an ϵ -greedy policy with respect to its action-value function⁵⁵ is more restricted in its policy representation depending on the current value of ϵ and its greedy action. For example, assume an action-value function where the first action has the largest value estimate in a given state. The ϵ -greedy policies derived from this greedy action for varying values of ϵ are visualised in Figure 8.5a. These policies are restricted in their expressiveness due to the ϵ -greedy policy representation. In contrast, a policy gradient algorithm learns a parameterised policy which can represent arbitrary policies (Figure 8.5b). This expressiveness of a probabilistic policy can be important in partially observable and multi-agent games, where the only optimal policy might be probabilistic. Consider for example the multi-agent game of Rock-Paper-Scissors. The Nash equilibrium (and the minimax solution) of this game is for both agents to choose each of its three actions with uniform probability (see Sections 4.3 and 4.4 for a refresher on these solution concepts). Therefore, a deterministic policy would not be able to represent the equilibrium, whereas a probabilistic policy can be used to represent this solution.

Moreover, by representing a policy as a separate learnable function, we can represent policies for continuous action spaces. In environments with continuous actions, an agent selects a single or several continuous values (typically within a certain interval) as its actions. For example, the agent might control (real or simulated) robots and vehicles, with continuous actions corresponding to the amount of torque applied by an actuator, rotational angles of a steering wheel, or the force applied to a brake pedal. All these actions are most naturally represented by continuous actions within a predefined interval of possible values. The value-based RL algorithms introduced in Section 8.1 are not applicable to such settings, because their neural network architecture has one output value for every possible (discrete) action corresponding to the action-value estimate of that particular action. However, there are an infinite number of continuous actions and thus the same architecture cannot be applied. In contrast, we can learn parameterised policies over continuous actions based on the policy gradient theorem, which will be introduced in Section 8.2.2. For example, we can represent a parameterised policy over continuous actions with a Gaussian distribution, in which the mean μ and standard deviation σ are computed by

⁵⁵. We note that there are other policy representations for value-based RL algorithms than ϵ -greedy policies. For example, agents can follow a Boltzmann policy which is similar to a softmax policy over the action-value estimates with a decaying temperature parameter, or agents can follow a deterministic exploration policy given with upper confidence bounds (UCB) from the bandit literature. However, all these policies of value-based algorithms are applied for exploration and converge to a deterministic policy for evaluation.

parameterised functions:

$$\pi(\cdot | s; \theta) = \mathcal{N}(\mu(s; \phi), \sigma(s; \phi)^2) \quad (8.6)$$

We note that a Gaussian policy is only one example of a parameterised policy over continuous actions and alternative approaches exist. Continuous action spaces can also be discretised by splitting the space of continuous actions into a fixed number of discrete actions. Discretisation of continuous action spaces allows to use any RL algorithm for discrete action spaces, but also restricts the action selection of the agent and may limit the achievable returns. This book focuses on policy gradient algorithms for discrete action spaces, which are commonly used in the MARL literature.

To represent a probabilistic policy for discrete action spaces using neural networks, we can use an identical architecture as for action-value functions (Figure 8.2). The policy network receives a state as input and outputs a scalar output for every action, $l(s, a)$, to represent the preference of the policy to select action a in state s . These preferences are transformed into a probability distribution using the *softmax function*, defined as the exponential of the preferences divided by the sum of the exponentials of all preferences:

$$\pi(a | s; \phi) = \frac{e^{l(s, a; \phi)}}{\sum_{a' \in A} e^{l(s, a'; \phi)}} \quad (8.7)$$

In the remainder of this book, we will always denote the parameters of a parameterised value function with θ and denote the parameters of a parameterised policy with ϕ .

8.2.2 Policy Gradient Theorem

In order to train a parameterised policy for policy gradient RL algorithms, we want to be able to use the same gradient-based optimisation techniques introduced in Section 7.4. For gradient-based optimisation, we require the function approximation of the policy to be differentiable. Besides a differentiable function, we need to specify the loss function to compute gradients to update the parameters of the policy. Generally, minimising the loss function should correspond to increasing the “quality” of the policy. But what constitutes a “good” policy? One sensible metric for the quality of a policy in episodic environments is expected episodic returns (Section 2.3). These expected returns in any given state under a given policy can also be represented by the value of the policy for that particular state, $V^\pi(s)$, or the action value $Q^\pi(s, a)$. However, optimising the policy to maximise such values can be challenging since changes of the policy directly affect not just the action selection, and thus the rewards received by the agent, but also the distribution of states encountered by the

agent. The action selection and changes in expected returns can be captured by the returns of experienced episodes, but the state distribution directly depends on the transition dynamics of the environment which is generally assumed to be unknown.

The *policy gradient theorem* (Sutton and Barto 2018) formulates a solution to these challenges and provides a theoretically founded expression for the gradient of the performance of a parameterised policy with respect to the parameters of the policy. The policy gradient theorem for episodic environments is given by

$$\nabla_{\phi} J(\phi) \propto \sum_{s \in S} \Pr(s \mid \pi) \sum_{a \in A} Q^{\pi}(s, a) \nabla_{\phi} \pi(a \mid s; \phi) \quad (8.8)$$

where J represents a function measuring the quality of a policy π with parameters ϕ , $\Pr(\cdot \mid \pi)$ represents the state-visitation distribution of a given policy π in the environment, and Q^{π} represents the value for a given action and state under the policy. The function J is similar to a loss function (Section 7.4.1) but with the key difference that we aim to *maximise* rather than minimise it. As we can see, the policy gradient does not depend on unknown information of the environment, such as the transition function and reward function, and hence can be instantiated with various techniques to approximate the value of the policy to guide the optimisation of a parameterised policy.⁵⁶ Note that the policy gradient theorem assumes the state-visitation distribution and action values to be given under the currently optimised policy π . This assumption becomes more apparent by rewriting the policy gradient as an expectation under the current policy. For this derivation, we make use of the fact that the policy gradient theorem defines a sum over states of the environment weighted by their

56. We note that in this section we provide the policy gradient for episodic environments. For continual environments, in which the experience of the agent is not split into finite episodes, we need to substitute the expected returns as a measure of quality of the policy with a suitable metric for the continual setting. One candidate for a metric of the quality of the policy for continual environments would be average rewards which capture the expected reward received by the agent under the state-visitation distribution and action selection of the current policy at any time step.

respective probability of occurring under the current policy, $\Pr(s \mid \pi)$:

$$\nabla_\phi J(\phi) \propto \sum_{s \in S} \Pr(s \mid \pi) \sum_{a \in A} Q^\pi(s, a) \nabla_\phi \pi(a \mid s; \phi) \quad (8.9)$$

$$= \mathbb{E}_{a \sim \pi(\cdot \mid s; \phi)} \left[\sum_{a \in A} Q^\pi(s, a) \nabla_\phi \pi(a \mid s; \phi) \right] \quad (8.10)$$

$$= \mathbb{E}_{a \sim \pi(\cdot \mid s; \phi)} \left[\sum_{a \in A} \pi(a \mid s; \phi) Q^\pi(s, a) \frac{\nabla_\phi \pi(a \mid s; \phi)}{\pi(a \mid s; \phi)} \right] \quad (8.11)$$

$$= \mathbb{E}_{a \sim \pi(\cdot \mid s; \phi)} \left[Q^\pi(s, a) \frac{\nabla_\phi \pi(a \mid s; \phi)}{\pi(a \mid s; \phi)} \right] \quad (8.12)$$

$$= \mathbb{E}_{a \sim \pi(\cdot \mid s; \phi)} [Q^\pi(s, a) \nabla_\phi \log \pi(a \mid s; \phi)] \quad (8.13)$$

We denote the natural logarithm with \log unless stated otherwise. We can see that the state distribution and action selection are both induced by the policy and, thus, fall within the expectation. In the end, we obtain a simple expression within the expectation under the current policy to express the gradients of the policy parameters towards policy improvement. This expectation also clearly illustrates the restriction that follows from the policy gradient theorem: the optimisation of the parameterised policy is limited to *on-policy data*, i.e. the data used to optimise π is generated by the policy π itself (Section 2.6). Therefore, data collected by interacting with the environment using any different policy π' , in particular also including previous policies obtained during the training of π , can *not* be used to update π following the policy gradient theorem. The application of a replay buffer, as seen in Section 8.1.3, would constitute such a violation of the assumption, because it contains experiences generated by “older and outdated” versions of our currently optimised policy. Therefore, a replay buffer cannot be used to update a policy following the policy gradient theorem. Furthermore, we have to approximate the expected returns *under our current policy* denoted with Q^π . Most algorithms which train an action-value function, such as DQN and other algorithms based on the classical Q-learning algorithm, do not satisfy this requirement. Instead, they directly approximate the optimal value function, i.e. the expected returns under the optimal policy, by following the Bellman optimality equation (Section 2.6).

Looking at the terms within the obtained expression (Equations 8.9 to 8.13) of the policy gradient theorem further provides an intuitive interpretation for policy improvement. This interpretation is most obvious using the following expression from our previous derivation:

$$\nabla_\phi J(\phi) = \mathbb{E}_\pi \left[Q^\pi(s, a) \frac{\nabla_\phi \pi(a \mid s; \phi)}{\pi(a \mid s; \phi)} \right] \quad (8.14)$$

The numerator of the fraction within the expression, $\nabla_\phi \pi(a|s; \phi)$, represents the gradient of the policy with respect to its parameters. This gradient can be used to modify the parameters of the policy and is further weighted by the quality of any particular action a in state s as given by the value or expected returns of the policy $Q^\pi(s, a)$. This weighting ensures that the policy optimises its parameters such that actions with higher expected returns become more probable than actions with lower expected returns. Lastly, the denominator of the fraction, $\pi(a|s; \phi)$, can be thought of as a normalising factor to correct for the data distribution induced by the policy. The policy π might take some actions with significantly higher probabilities than others and hence more updates might be done to the policy parameters to increase the probability of more likely actions. To account for this factor, the policy gradient needs to be normalised by the inverse of the action probability under the policy.

8.2.3 REINFORCE: Monte Carlo Policy Gradient

The policy gradient theorem defines the gradient to update a parameterised policy to gradually increase its expected returns. In order to use the theorem to compute gradients and update the parameters of the policy, we need to either approximate the derived expectation (Equation 8.13) or obtain samples from it. Monte Carlo estimation is one possible sampling method, which uses on-policy samples of episodic returns to approximate the expected returns of the policy. Using the policy gradient theorem with Monte Carlo samples gives rise to REINFORCE (Williams 1992) which minimises the following loss for the full history of an episode, $\hat{h} = \{s^0, a^0, r^0, \dots, s^{T-1}, a^{T-1}, r^{T-1}, s^T\}$:

$$\mathcal{L}(\phi) = -\frac{1}{T} \sum_{t=0}^{T-1} \left(\sum_{\tau=t}^{T-1} \gamma^{\tau-t} r^\tau \right) \log \pi(a^t | s^t; \phi) \quad (8.15)$$

$$= -\frac{1}{T} \sum_{t=0}^{T-1} \left(\sum_{\tau=t}^{T-1} \gamma^{\tau-t} \mathcal{R}(s^\tau, a^\tau, s^{\tau+1}) \right) \log \pi(a^t | s^t; \phi) \quad (8.16)$$

Given the policy gradient theorem provides a gradient in the direction of policies with *higher* expected returns and we want to define a *to-be-minimised* loss, this loss corresponds to the *negative* policy gradient (Equation 8.13) with Monte Carlo estimates of the expected returns under the current policy π . During training, the REINFORCE algorithm first collects a full episodic trajectory with history \hat{h} by using its current policy π . After an episode has terminated, the return estimate is computed to estimate the policy gradient that minimises the loss given in Equation 8.16. Pseudocode for the REINFORCE algorithm is given in Algorithm 12.

Algorithm 12 REINFORCE

- 1: Randomly initialise policy network π with parameters ϕ
 - 2: Repeat for every episode:
 - 3: **for** time step $t = 0, 1, 2, \dots, T - 1$ **do**
 - 4: Observe current state s^t
 - 5: Sample action $a^t \sim \pi(\cdot | s^t; \phi)$
 - 6: Apply action a^t ; observe rewards r^t and next state s^{t+1}
 - 7: Collect history $\hat{h} \leftarrow \{s^0, a^0, r^0, \dots, s^{T-1}, a^{T-1}, r^{T-1}\}$
 - 8: Update parameters ϕ by minimising the loss from Equation 8.16
-

Unfortunately, Monte Carlo return estimates have high variance, which leads to high variance of computed gradient estimates and unstable training. This high variance arises due to the returns of each episode depending on all states and actions encountered within the episode. Both states and actions are samples of a probabilistic transition function and policy, respectively. To reduce the variance of return estimates, we can subtract a *baseline* from the return estimates. For any baseline $b(s)$ defined over the state s , the gradient derived from the policy gradient theorem (Equation 8.13) remains unchanged in expectation. Therefore, even when subtracting a baseline we still optimise the parameters of the policy to maximise its quality but we reduce the variance of the gradient estimates. To see that the policy gradient remains unchanged, we can rewrite the policy gradient theorem as follows:

$$\nabla_\phi J(\phi) \propto \sum_{s \in S} \Pr(s | \pi) \sum_{a \in A} (Q^\pi(s, a) - b(s)) \nabla_\phi \pi(a | s; \phi) \quad (8.17)$$

$$= \mathbb{E}_\pi \left[\sum_{a \in A} (Q^\pi(s, a) - b(s)) \nabla_\phi \pi(a | s; \phi) \right] \quad (8.18)$$

$$= \mathbb{E}_\pi \left[\sum_{a \in A} \pi(a | s; \phi) (Q^\pi(s, a) - b(s)) \frac{\nabla_\phi \pi(a | s; \phi)}{\pi(a | s; \phi)} \right] \quad (8.19)$$

$$= \mathbb{E}_\pi \left[(Q^\pi(s, a) - b(s)) \frac{\nabla_\phi \pi(a | s; \phi)}{\pi(a | s; \phi)} \right] \quad (8.20)$$

$$= \mathbb{E}_\pi [(Q^\pi(s, a) - b(s)) \nabla_\phi \log \pi(a | s; \phi)] \quad (8.21)$$

$$= \mathbb{E}_\pi [Q^\pi(s, a) \nabla_\phi \log \pi(a | s; \phi)] - \mathbb{E}_\pi [b(s) \nabla_\phi \log \pi(a | s; \phi)] \quad (8.22)$$

$$= \mathbb{E}_\pi [Q^\pi(s, a) \nabla_\phi \log \pi(a | s; \phi)] - \sum_{s \in S} \Pr(s | \pi) \sum_{a \in A} b(s) \nabla_\phi \pi(a | s; \phi) \quad (8.23)$$

$$= \mathbb{E}_\pi [Q^\pi(s, a) \nabla_\phi \log \pi(a | s; \phi)] - \sum_{s \in S} \Pr(s | \pi) b(s) \nabla_\phi \sum_{a \in A} \pi(a | s; \phi) \quad (8.24)$$

$$= \mathbb{E}_\pi [Q^\pi(s, a) \nabla_\phi \log \pi(a | s; \phi)] - \sum_{s \in S} \Pr(s | \pi) b(s) \nabla_\phi 1 \quad (8.25)$$

$$= \mathbb{E}_\pi [Q^\pi(s, a) \nabla_\phi \log \pi(a | s; \phi)] - \sum_{s \in S} \Pr(s | \pi) b(s) 0 \quad (8.26)$$

$$= \mathbb{E}_\pi [Q^\pi(s, a) \nabla_\phi \log \pi(a | s; \phi)] \quad (8.27)$$

A state-value function $V(s)$ is a common choice for a baseline, which can be trained to minimise the following loss given history \hat{h} :

$$\mathcal{L}(\theta) = \frac{1}{T} \sum_{t=0}^{T-1} \left(u(\hat{h}^t) - V(s^t; \theta) \right)^2 \quad (8.28)$$

Similarly, the REINFORCE policy loss with a state-value function as a baseline can be written as:

$$\mathcal{L}(\phi) = -\frac{1}{T} \sum_{t=0}^{T-1} \left(u(\hat{h}^t) - V(s^t; \theta) \right) \log \pi(a^t | s^t; \phi) \quad (8.29)$$

8.2.4 Actor-Critic Algorithms

Actor-critic algorithms are a family of policy gradient algorithms which train a parameterised policy, the *actor*, and a value function, the *critic*, alongside each other. As seen before, the actor is optimised using gradient estimates derived from the policy gradient theorem. However, in contrast to REINFORCE (with or without a baseline), actor-critic algorithms use the critic to compute bootstrapped return estimates. Using bootstrapped value estimates to optimise policy gradient algorithms has two primary benefits.

Firstly, bootstrapped return estimates allow us, as seen with temporal-difference algorithms (Section 2.6), to estimate episodic returns just from the experience of a single step. Using bootstrapped return estimates of a state-value function V , we can estimate episodic returns as follows:

$$\mathbb{E}_\pi [u(\hat{h}^t) | s^t] = \mathbb{E}_\pi [\mathcal{R}(s^t, a^t, s^{t+1}) + \gamma u(\hat{h}^{t+1:T}) | s^t, a^t \sim \pi(s^t)] \quad (8.30)$$

$$= \mathbb{E}_\pi [\mathcal{R}(s^t, a^t, s^{t+1}) + \gamma V(s^{t+1}) | s^t, a^t \sim \pi(s^t)] \quad (8.31)$$

By using bootstrapped return estimates, actor-critic algorithms are able to update the policy (and critic) from any immediate experience, irrespective of the following history of the episode. In particular, in non-terminating environments

or environments with long episodes, this allows significantly more frequent updates than with REINFORCE, which often improves the training efficiency.

Secondly, bootstrapped return estimates exhibit lower variance compared to the Monte Carlo estimates of episodic returns used in REINFORCE. Variance is reduced because these return estimates only depend on the current state, received reward, and next state, and unlike episodic returns do not depend on the entire future history of the episode. However, this reduction in variance comes at a price of introduced bias, because the used value function might not (yet) approximate the true expected returns. In practice, we find that the trade-off of bias for lower variance often improves training stability. Moreover, N -step return estimates can be used. Instead of directly computing a bootstrapped value estimate of the next state, these estimates aggregate the received rewards of N consecutive steps before computing a bootstrapped value estimate of the following state:

$$\mathbb{E}_\pi[u(\hat{h}^t) | s^t] = \mathbb{E}_\pi \left[\left(\sum_{\tau=0}^{N-1} \gamma^\tau \mathcal{R}(s^{t+\tau}, a^{t+\tau}, s^{t+\tau+1}) \right) + \gamma^N V(s^{t+N}) \mid s^t, a^\tau \sim \pi(s^\tau) \right] \quad (8.32)$$

For $N = T$, with T being the episode length, the computed return estimate corresponds to the Monte Carlo episodic returns with no bootstrapped value estimates, as used in REINFORCE. These return estimates have high variance but are unbiased. For $N = 1$, we obtain one-step bootstrapped return estimates, as given in Equation 8.31, with low variance and high bias. Using the hyperparameter of N allows us to arbitrarily choose between bias and variance of return estimates. We will illustrate this trade-off of N -step return estimates at the example of an actor-critic algorithm in Figure 8.6. N -step returns are commonly applied for small N , such as $N = 5$ or $N = 10$, to obtain return estimates with fairly low bias and variance.

For notational brevity, we will write pseudocode and equations using one-step bootstrapped return estimates, but note that N -step return estimates can be applied to substitute any of these value estimates. In the following, we will introduce two actor-critic algorithms: advantage actor-critic (A2C) and proximal policy optimisation (PPO).

8.2.5 A2C: Advantage Actor-Critic

Advantage actor-critic (A2C)⁵⁷ (Mnih et al. 2016) is a foundational actor-critic algorithm which, as the name suggests, computes estimates of the *advantage* of a policy to guide the policy gradients. The advantage for a state s and action a is given by

$$Adv^\pi(s, a) = Q^\pi(s, a) - V^\pi(s) \quad (8.33)$$

with Q^π and V^π representing action-value and state-value functions with respect to a policy π , respectively. For the policy π , the action value $Q^\pi(s, a)$ represents the expected returns of π when first applying action a in state s and afterwards following π . In contrast, the state value $V^\pi(s)$ represents the expected returns when following the policy π already in state s rather than taking a specific predetermined action. The advantage can therefore be understood as quantifying how much higher the expected returns are when applying the specific action a compared to following the policy π in state s . The advantage takes on positive values whenever the chosen action a achieves higher expected returns than the current policy π is expected to achieve in state s . Similarly, the advantage is negative when the chosen action a achieves lower expected returns than the current policy π . This interpretation of the advantage can directly be used to guide the optimisation of the policy. For a positive advantage, we should increase the probability of the policy π to select action a in state s , and we should decrease the probability of the policy π to select action a in state s whenever the advantage is negative. In the following sections, we will omit the superscript π for brevity and assume that the advantage and value functions are computed with respect to the current policy π .

As defined above, estimating the advantage would require both an action-value function and a state-value function. Fortunately, we can estimate an action-value function using the immediate rewards and the state-value estimate of the following state:

$$Q(s^t, a^t) = \mathcal{R}(s^t, a^t, s^{t+1}) + \gamma V(s^{t+1}) = r^t + \gamma V(s^{t+1}) \quad (8.34)$$

With this estimation of the action-value function, we only rely on a state-value function to approximate the advantage:

$$Adv(s^t, a^t) = Q(s^t, a^t) - V(s^t) = r^t + \gamma V(s^{t+1}) - V(s^t) \quad (8.35)$$

57. Mnih et al. (2016) originally proposed the Asynchronous Advantage Actor Critic (A3C) algorithm which uses asynchronous threads to collect experience from the environments. For simplicity, and because in practice it often does not make a difference, we avoid the asynchronous aspect of this algorithm and introduce a simplified version of its synchronous implementation (A2C). We will further discuss asynchronous and synchronous parallelisation of training in Section 8.2.7.

Algorithm 13 Simplified Advantage Actor-Critic (A2C)

- 1: Randomly initialise policy network π with parameters ϕ
 - 2: Randomly initialise value function network V with parameters θ
 - 3: Repeat for every episode:
 - 4: **for** time step $t = 0, 1, 2, \dots$ **do**
 - 5: Observe current state s^t
 - 6: Sample action $a^t \sim \pi(\cdot | s^t; \phi)$
 - 7: Apply action a^t ; observe rewards r^t and next state s^{t+1}
 - 8: Update parameters ϕ by minimising the loss from Equation 8.37
 - 9: Update parameters θ by minimising the loss from Equation 8.38
-

Similar to return estimates (Equation 8.32), we can estimate the advantage using multiple steps of rewards before bootstrapping the value estimate of the final state to reduce the variance of the obtained estimate:

$$Adv(s^t, a^t) = \sum_{\tau=0}^{N-1} \gamma^\tau \mathcal{R}(s^{t+\tau}, a^{t+\tau}, s^{t+\tau+1}) + \gamma^N V(s^{t+N}) - V(s^t) \quad (8.36)$$

In A2C, we optimise the parameters ϕ of the actor to maximise the advantage by minimising the following loss:

$$\mathcal{L}(\phi) = -Adv(s^t, a^t) \log \pi(a^t | s^t; \phi) \quad (8.37)$$

For the optimisation of the parameters of the critic θ , we compute the squared error of the value estimate of the current state and the bootstrapped target estimate. Below, we show the respective loss with an one-step bootstrapped target estimate.

$$\mathcal{L}(\theta) = (r^t + \gamma V(s^{t+1}; \theta) - V(s^t; \theta))^2 \quad (8.38)$$

Commonly, multi-step target estimates are used to reduce the variance of the critic's loss. In this case, the target can be computed as shown in Equation 8.32.

Full pseudocode for the A2C algorithm is given in Algorithm 13. We denote this algorithm as “Simplified A2C” because the algorithm was originally proposed with two further techniques: asynchronous or synchronous parallelisation of training, and entropy regularisation to incentivise exploration. We will discuss parallelisation of training in Section 8.2.7. Entropy regularisation adds an additional term to the actor loss given by the negative entropy of the policy in the current state:

$$-\mathcal{H}(\pi(\cdot | s; \phi)) = \sum_{a \in A} \pi(a | s; \phi) \log \pi(a | s; \phi) \quad (8.39)$$

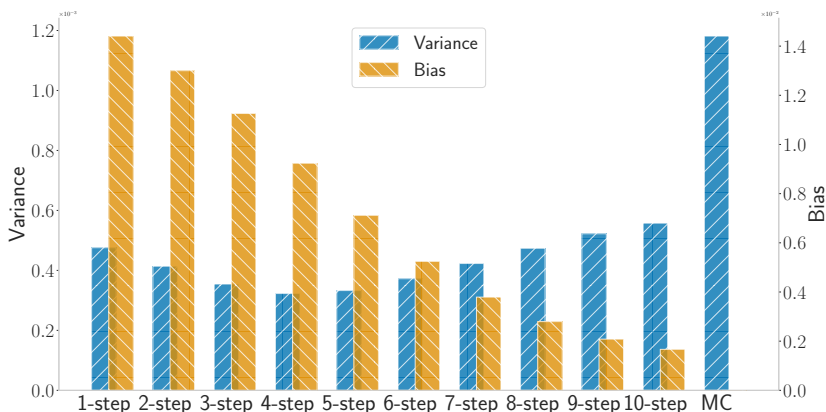


Figure 8.6: Variance and bias of N -step return estimates for $N \in \{1, \dots, 10\}$ and Monte Carlo returns for a state-value function trained with A2C for 100,000 time steps in the single-agent level-based foraging environment (Figure 8.1). We use N -step return estimates with $N = 5$ during training.

The entropy of the policy π is a measure of the policy’s uncertainty. The entropy is maximised for a uniform distribution, i.e. when the policy selects all actions with equal probability. Minimising the negative entropy, i.e. maximising the entropy, as part of the actor loss penalises the policy for assigning a very high probability to any action. This regularisation term discourages premature convergence to a suboptimal close-to-deterministic policy and, thus, incentivises exploration.

To illustrate the trade-off of variance and bias when using N -step return estimates (Equation 8.32), we train a policy and state-value function with A2C in the single-agent level-based foraging environment (Figure 8.1). After training with $N = 5$, we collect 10,000 episodes with the policy at the end of training. Figure 8.6 visualises the variance and bias of N -step return estimate with $N \in [1, 10]$ and Monte Carlo return estimates using the trained critic on the entire dataset of 10,000 episodes. As expected, the variance of the N -step return estimates increases with increasing N and Monte Carlo returns exhibit the highest variance. In contrast, the bias gradually decreases with increasing N until Monte Carlo returns which are fully unbiased. In practice, we often choose $N = 5$ or $N = 10$ for comparably low variance and bias.

8.2.6 PPO: Proximal Policy Optimisation

In all algorithms discussed in the previous sections, the policy parameters are continually updated using gradients derived with the policy gradient theorem. These gradients aim to move the policy parameters towards a policy with higher expected returns. However, any individual gradient update step, even for small learning rates, might lead to significant changes of the policy and could reduce the expected performance of the policy. The risk of such significant changes of the policy as a consequence of a single gradient optimisation step can be reduced using *trust regions*. Intuitively, trust regions define an area within the space of policy parameters in which the policy would not change significantly and, thus, we would “trust” that the resulting policy with such parameters would not lead to a significant reduction in performance. *Trust region policy optimisation* (TRPO) (Schulman et al. 2015) constrains each optimisation step of the policy in policy gradient RL algorithms to a small trust region. In this way, TRPO aims to reduce any drops in quality of the policy and thereby gradually and safely improve the quality of the policy. However, each update with TRPO requires either solving a constrained optimisation problem or computing a penalty term which are both computationally expensive.

Proximal policy optimisation (PPO)⁵⁸ (Schulman et al. 2017) builds on the idea of trust regions for policy optimisation and computes a computationally efficient surrogate objective to avoid large jumps in policy in a single optimisation step. This surrogate objective makes use of importance sampling weights $\rho(s, a)$ which are defined as the fraction of probabilities of selecting a given action a in state s for two policies:

$$\rho(s, a) = \frac{\pi(a|s; \phi)}{\pi_\beta(a|s)} \quad (8.40)$$

For the importance sampling weight ρ , the policy π , parameterised by ϕ , represents the policy we want to optimise and π_β represents a behaviour policy which was followed to generate the data. More specifically, the behaviour policy π_β was used to select the action a in s . The importance sampling weight can be thought of as a factor to shift from the distribution of data encountered under policy π_β to the data distribution of policy π . This factor adjusts the data distribution to make the data generated by π_β “appear” on-policy to the policy π . Using these weights, PPO is able to update the policy multiple times using the same data. Typical policy gradient algorithms rely on the policy gradient

58. Schulman et al. (2017) propose two versions of the PPO algorithm in their work. In this section, we describe the PPO algorithm with a clipped surrogate objective. This algorithm is simpler and more common than the alternative PPO algorithm with a KL divergence penalty term, and often just referred to as PPO.

Algorithm 14 Simplified Proximal Policy Optimisation (PPO)

-
- 1: Randomly initialise policy network π with parameters ϕ
 - 2: Randomly initialise value function network V with parameters θ
 - 3: Repeat for every episode:
 - 4: **for** time step $t = 0, 1, 2, \dots$ **do**
 - 5: Observe current state s^t
 - 6: Sample action $a^t \sim \pi(\cdot | s^t; \phi)$
 - 7: Apply action a^t ; observe rewards r^t and next state s^{t+1}
 - 8: **for** epoch $e = 1, \dots, N_e$ **do**
 - 9: Update parameters ϕ by minimising the loss from Equation 8.41
 - 10: Update parameters θ by minimising the loss from Equation 8.38
-

theorem and, therefore, assume data to be on-policy. However, after a single update of the policy, the policy changes and any previously collected data is not on-policy anymore. Additionally, the importance sampling weight can be seen as a measure of divergence of the policies, with an importance weight of 1 corresponding to both policies having equal probabilities of selecting an action a in state s . PPO makes use of these properties to update the policy multiple times using the same data, and restrict the change of the policy by restricting the importance sampling weight. This is achieved using a policy loss with clipped importance sampling weights

$$\mathcal{L}(\phi) = -\min \left(\frac{\rho(s^t, a^t) \text{Adv}(s^t, a^t)}{\text{clip}(\rho(s^t, a^t), 1 - \epsilon, 1 + \epsilon) \text{Adv}(s^t, a^t)} \right) \quad (8.41)$$

where ρ represents the importance sampling weight as defined in Equation 8.40, the advantage $\text{Adv}(s^t, a^t)$ is computed using a state-value function as given in Equation 8.35, and ϵ represents a hyperparameter which determines how much the policy is allowed to deviate from the previous policy π_β .

Pseudocode for PPO is given in Algorithm 14 with N_e denoting the number of epochs, i.e. number of updates, for a given batch of data. Similar to A2C, we denote the presented algorithm as “Simplified PPO” because it is typically applied in combination with parallel training (Section 8.2.7), entropy regularisation (Section 8.2.5), and N -step returns (Section 8.2.4) to obtain larger batches of data for more stable optimisation and improve the exploration of the algorithm.

We compare the policy gradient algorithms REINFORCE, A2C, and PPO in the single-agent level-based foraging environment introduced in Figure 8.1. We see that REINFORCE agents learn to solve the task in most runs at the

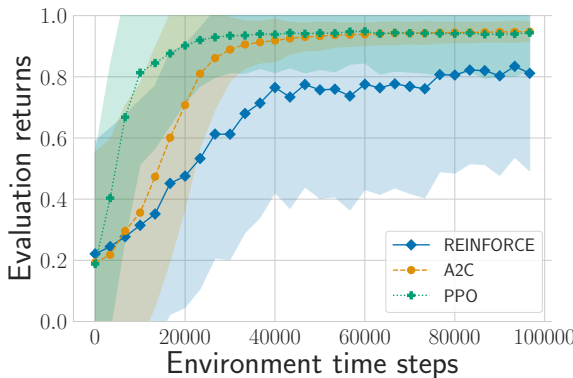


Figure 8.7: Learning curves for REINFORCE, A2C, and PPO in the single-agent level-based foraging environment shown in Figure 8.1. We train all algorithms for 100,000 time steps. Visualised learning curves and shading correspond to the mean and standard deviation across discounted episodic returns across five runs with different random seeds. Across all algorithms, we use a discount factor $\gamma = 0.99$ during training, small critic and actor networks with two layers of 32 hidden units, ReLU activation function, and conduct a small grid search to identify suitable hyperparameters. REINFORCE is trained without a baseline and with a learning rate of $\alpha = 1e^{-3}$. For A2C and PPO we use N -step returns with $N = 5$ and a learning rate of $3e^{-4}$. Lastly, PPO uses a clipping parameter $\epsilon = 0.2$ and optimises its networks for $N_e = 4$ epochs using the same batch of experience.

end of training, but the episodic returns exhibit high variance all throughout training. This variance can be explained by the high variance of Monte Carlo returns (Figure 8.6) and, thus, highly variant policy gradients during training. In contrast, A2C and PPO with N -step returns reach the optimal performance across all runs within 20,000 time steps. This experiment demonstrates the improved stability and sample efficiency of actor-critic algorithms such as A2C and PPO. In particular, with N -step returns, training is significantly more stable than REINFORCE due to less variant return estimates, and the agent robustly obtains optimal returns across all episodes thereafter. Lastly, we see that PPO is able to learn slightly faster than A2C, which can be explained by its optimisation being able to use each batch of experiences for multiple updates.

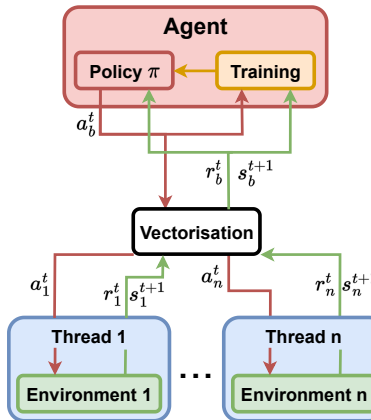


Figure 8.8: Visualisation of synchronous data collections to parallelise interactions of the agent across multiple environment instances running in parallel. Environment instances are executed across different threads. At each time step, the agent selects a vector of actions a_b^t with one action for each environment conditioned on the last batch of states s_b^t . Each environment instance receives its action and transitions to a new state, returning both the reward and new state. The batch of rewards r_b^t and new states s_b^{t+1} from all environment instances is then passed to the agent as vectors for training and its next action selection. The parallelisation of this technique is synchronous because the agent has to wait with its next action selection until all environment instances finished their current transition.

8.2.7 Concurrent Training of Policies

On-policy policy gradient algorithms cannot make use of a replay buffer, as applied in off-policy value-based RL algorithms such as DQN (Section 8.1.3). However, the replay buffer is a key component of off-policy RL algorithms in order to break correlations between consecutive experiences and provides larger batches of data to compute the loss. This raises the question of how to break correlations and obtain batches of data for sample efficient optimisation of on-policy policy gradient algorithms. In this section, we introduce two approaches to target this problem by parallelising the interaction of the agent with the environment using multi-threading capabilities of modern hardware: synchronous data collection and asynchronous training.

Synchronous data collection, visualised in Figure 8.8, initiates separate instances of the environment for every thread. At every time step, the agent

receives a batch of states and rewards from all environment instances and independently decides on its action selection for every environment. A batch of selected actions is then sent to each of the environments in its respective thread to transition to a new state and receive a new reward. This interaction is repeated throughout all of training, and is synchronous because the agent has to wait for its next action selection until all environment instances have transitioned to their new state. Synchronous data collection is simple to deploy with minimal changes required for the training of the RL algorithm, and significantly increases the amount of data samples available for each update. Similarly to the batches sampled from a replay buffer, averaging gradients across such batches of experience makes gradients more stable and optimisation more efficient. Moreover, the forward pass over batches of inputs through a neural network can be parallelised using efficient vector and matrix operations, so the computation required for synchronous data collection is highly efficient. The benefits of vectorised computation are particularly significant for modern hardware such as GPUs, which are able to perform many operations in parallel. Lastly, correlations of consecutive experiences are partly broken because experiences across different environment instances may vary significantly due to different initial states and probabilistic transitions.

We show the pseudocode for the simplified A2C algorithm with synchronous data collection in Algorithm 15. The algorithm is identical to Algorithm 13 except that the agent computes its loss over batches of experience from all environment instances and independently interacts with every environment. The actor loss for A2C over a batch of experience is given by

$$\mathcal{L}(\phi) = \frac{1}{K} \sum_{k=1}^K - \underbrace{(r_k^t + \gamma V(s_k^{t+1}; \theta) - V(s_k^t; \theta))}_{\text{Adv}(s_k^t, a_k^t)} \log \pi(a_k^t | s_k^t; \phi) \quad (8.42)$$

and the critic loss is computed as follows:

$$\mathcal{L}(\theta) = \frac{1}{K} \sum_{k=1}^K (r_k^t + \gamma V(s_k^{t+1}; \theta) - V(s_k^t; \theta))^2 \quad (8.43)$$

Note that simplified A2C optimises its networks once each of its K environments has completed a single time step⁵⁹. Therefore, the agent collects more experiences within the same wall-clock time⁶⁰ for larger values of K but also

59. We commonly use N -step returns to obtain value estimates with reduced bias. In this case, the batch of $K * N$ experiences across N time steps and K environments can be used to optimise the networks.

60. Wall-clock time is the elapsed time that a clock or stopwatch would measure between the start and end of the training, irrespective of the required resources and number of parallel threads or processes.

Algorithm 15 Simplified Advantage Actor-Critic (A2C) with Synchronous Environments

- 1: Randomly initialise policy network π with parameters ϕ
 - 2: Randomly initialise value function network V with parameters θ
 - 3: Initialise K parallel environments
 - 4: Repeat for every episode:
 - 5: **for** time step $t = 0, 1, 2, \dots$ **do**
 - 6: Observe a batch of current states for all environments $[s_1^t \dots s_K^t]^T$
 - 7: Sample actions $a_k^t \sim \pi(\cdot | s_k^t; \phi)$ for $k = 1, \dots, K$
 - 8: Execute action a_k^t in k th environment for $k = 1, \dots, K$; observe rewards $[r_1^t \dots r_K^t]^T$ and next states $[s_1^{t+1} \dots s_K^{t+1}]^T$
 - 9: Update parameters ϕ by minimising the loss from Equation 8.42
 - 10: Update parameters θ by minimising the loss from Equation 8.43
-

uses more experience for each optimisation of its networks as it makes use of the experience across all K environments.

To illustrate the impact of synchronous parallel environments on the training of the agent, we train the simplified A2C algorithm with synchronous environments (Algorithm 15) with varying numbers of synchronous environments in a single-agent level-based foraging environment. The environment has a larger 12×12 grid and the agent has to collect two items to receive all possible episodic rewards. We train the agent with $K \in \{1, 4, 16, 64\}$ for five minutes and present (discounted) evaluation returns across time steps trained and wall-clock training time in Figure 8.9. On one hand, the experiment illustrates that training for smaller values of K can be comparably sample efficient because of the frequent optimisations of the agent’s networks (Figure 8.9a). On the other hand, the optimisation is less stable because each optimisation is computed over a smaller batch of experiences, so the agent trained with $K=1$ environment does not converge to the optimal policy. Inspecting the wall-clock efficiency in Figure 8.9b, we see that training with larger values of K can be considerably more efficient (whilst making use of larger amounts of computational resources). However, these benefits diminish with growing number of synchronous environments. For large values of K , individual threads have to wait for other threads to finish their transition to receive the next action from the agent and continue their interaction, so idle time of threads increases the more parallel environments are deployed.

Asynchronous training, visualised in Figure 8.10, instead parallelises the optimisation of the agent. In addition to an instance of the environment, each

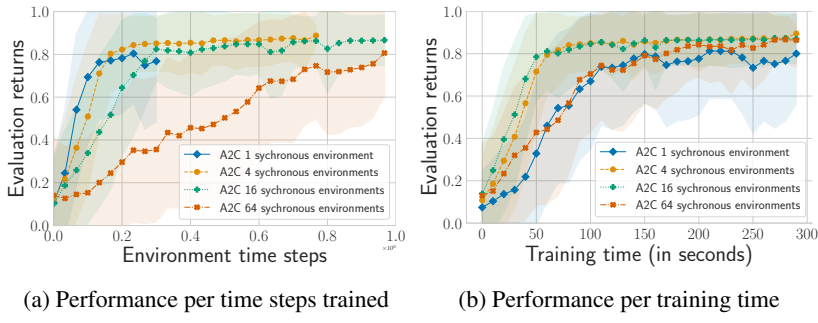


Figure 8.9: Learning curves for A2C in a single-agent level-based foraging environment similar to the one shown in Figure 8.1 but with a 12×12 grid and a total of two items the agent has to collect each episode for optimal performance. We train simplified A2C for 5 minutes with $K \in \{1, 4, 16, 64\}$ synchronous environments and compare the (a) sample efficiency as given by the episodic returns per time steps trained and (b) wall-clock efficiency as given by the episodic returns per time trained. Visualised learning curves and shading correspond to the mean and standard deviation across discounted episodic returns across five runs with different random seeds. For all algorithms, we use a discount factor $\gamma = 0.99$ during training, small critic and actor networks with two layers of 32 hidden units, ReLU activation function, a learning rate of $\alpha = 1e^{-3}$, and N -step returns with $N = 10$.

thread keeps a copy of the agent to interact with its environment instance. Each thread separately computes the loss and gradients to optimise the parameters of the agent’s networks based only on data collected within the environment instance of the thread. Once gradients are computed, the networks of the central agent are updated and the newly obtained parameters are sent to all threads to update their agent copies. Therefore, the agent’s networks are separately updated by all threads, with each thread’s optimisation only using data collected within that particular thread. Implementing memory-safe asynchronous parallelisation and optimisation is more involved and requires careful engineering considerations, but has the major benefit of not relying on information of all threads to proceed for every interaction. Threads can independently complete transitions and optimisations, which minimises potential idle time.

Due to the parallelised computation, both synchronous environments and asynchronous training can efficiently leverage multiple threads ranging from a few threads supported by CPUs found in most consumer laptops and desktop

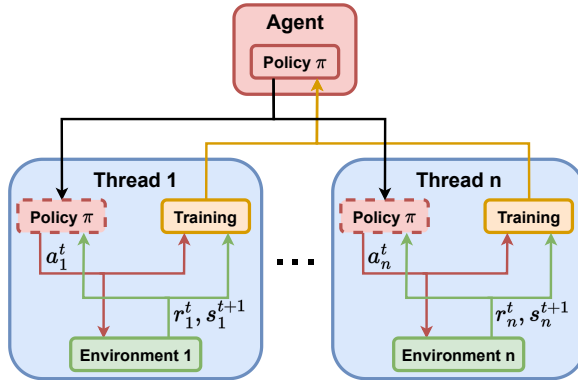


Figure 8.10: Asynchronous parallelisation

Figure 8.11: Visualisation of asynchronous training to parallelise the optimisation of the agent across multiple threads. Each thread keeps a copy of the agent to interact with its separate environment instance. Training is done within each thread and only uses the data collected from the environment instance in that particular thread. Whenever gradients are computed to update the agent's networks within the training of any thread, the networks of the central agent are updated and the updated parameters are shared with all threads. This ensures that all threads use the most up-to-date networks at all times and updates of any thread affect the policy executed by other threads.

computers, up to thousands of threads executed across large distributed computing clusters. Whenever multiple CPU threads are available, synchronous data collection is a comparably simple approach to significantly improve the efficiency of policy gradient algorithms. If multiple machines with dedicated accelerators for deep learning models are available, asynchronous training may be preferred due to its ability to independently optimise network parameters within each thread. However, it is worth noting that both approaches assume that multiple instances of environment can be executed in parallel. This is not always the case, for example, when the environment is a physical system with a single instance such as a robot. In these cases, parallelisation of data collection with these techniques is not possible and other techniques have to be used to improve the efficiency of policy gradient algorithms.

We further note that these techniques are most commonly applied with on-policy policy gradient because these algorithms cannot use replay buffers, but parallel training and data collection are also applicable to off-policy algorithms. Also, in this section we focused on two conceptually simple ideas to improve

the efficiency of policy gradient algorithms. More complex ideas have been proposed in the literature which largely focus on parallelisation across large computing infrastructures (e.g. Espeholt et al. (2018) and Espeholt et al. (2020)).

8.3 Observations, States, and Histories in Practice

In this chapter, we defined deep RL algorithms conditioned on states of the environment. However, as discussed in Section 3.4, the agent might not observe the full state of the environment but only receive a partial view of the current state. Consider for example an environment in which the agent has to control a robot. Using its sensors, the agent perceives its environments but some objects might be out of its sensory view or be occluded by other objects. In such partially observable environments, learned value functions and policies should be conditioned on the episodic history of observations $h^t = (o^0, \dots, o^t)$ to make use of all information perceived within an episode until time step t .

In order to condition value functions and policies on the history of observations, we could concatenate all observations into a single vector representing h^t and use this vector as the input to the policy and value networks. However, this approach is not practical because the dimensionality of the input vector grows as past observations are accumulated. Most neural network architectures, including the most commonly used feedforward neural networks, require a constant input dimensionality. To represent such a concatenated observation vector as an input of constant dimensionality for a deep value functions or policy networks, we could represent the history as a zero-padded vector of sufficient dimensionality to represent a history of maximum episode length⁶¹. However, such an input vector would be of high dimensionality and very sparse, i.e. it would contain mostly zero values for most histories. These properties would make it difficult to learn a good policy and value function conditioned on such episodic histories.

We have already seen a deep learning technique to address these challenges. Recurrent neural networks, as introduced in Section 7.5.2, are designed to process sequences of inputs. By treating the history of observations as such a sequence, a recurrent neural network is able to process the history one observation at a time. At each time step, the network only receives the most recent observation as its input and continually updates its hidden state to represent information about the full episodic history of observations. At the beginning of each episode, this hidden state is initialised to a zero-valued vector. By

61. For tasks with potentially infinite episodes, this approach already falls flat.

using recurrent neural networks within the architecture of policies and value functions in partially observable environments, RL algorithms can receive one observation at a time to internally represent the full episodic history. Using this architectural change for all networks, the deep RL algorithms introduced in this chapter can be applied to partially observable environments.

9 Multi-Agent Deep Reinforcement Learning

In Chapter 8, we saw that tabular MARL algorithms, introduced in Part I, are limited because their value functions are only updated for visited states. This inability to generalise to previously unseen inputs makes tabular MARL algorithms ineffective in environments with many states because the agent might not encounter states a sufficient number of times to obtain accurate value estimates. Fortunately, deep learning, introduced in Chapter 7, provides us with the tools to train neural networks as flexible function approximators that generalise over large input spaces. Chapter 8 already demonstrated how deep learning can be used to train parameterised value functions and policies for RL. This chapter will extend these ideas to multi-agent RL and introduce fundamental algorithms for training multiple agents to solve complex tasks.

To set the context for algorithms presented in this chapter, we will begin by discussing different paradigms of MARL training which differ in the information available for training and execution of agent policies. Then, we will discuss deep independent learning algorithms for MARL which naively apply deep single-agent RL by training each agent’s policy while ignoring the presence of other agents. After that, we will introduce more sophisticated algorithms that make use of joint information from multiple agents available during training to improve the learning process for policy gradient and value-based MARL algorithms. Concurrently training multiple agents often requires large numbers of samples to learn effective policies. Therefore, we will also discuss how multiple agents can share networks and experiences to make the training more sample efficient. Lastly, we will cover policy self-play for zero-sum games. Under this paradigm, a single agent is trained to play a zero-sum game by playing against copies of its own policy. Self-play has been highly impactful and is a core component of several MARL breakthroughs in competitive board-and video-game playing.

9.1 Training and Execution Modes

MARL algorithms can be categorised based on the information available during the learning and execution of policies. During training, each agent may be restricted to only use local information observed by itself (“decentralised training”), or might be able to leverage information about all agents in the multi-agent system (“centralised training”). After the training of agent policies, the question of available information remains: What information can agents use to make their action selection, i.e. to condition their policy on? Most commonly, policies of agents are only conditioned on their local history of observations (“decentralised execution”), but under some circumstances it might be reasonable to assume availability of information from all agents (“centralised execution”). This section will give a brief description of the three main categories of MARL algorithms based on their modes of training and execution.

9.1.1 Centralised Training and Execution

In Centralised Training and Execution, the learning of agent policies as well as the policies themselves use some type of information or mechanism that is centrally shared between the agents. Centrally shared information may include the agents’ local observation histories, learned world and agent models, value functions, or even the agents’ policies themselves. In the case of centralised training and execution, we knowingly depart from the typical setting defined by a POSG (Section 3.4), since agents are no longer limited to only receiving local observations of the environment. Therefore, centrally shared information can be considered privileged information which may benefit the training or execution of agents if the application scenario allows for it.

An example of this category is central learning (Section 5.3.1), which reduces a multi-agent game to a single-agent problem by using the joint observation history, i.e. the history of observations of all agents, to train a single central policy, which then sends actions to all agents. This approach has the primary benefit of being able to leverage the joint observation space of the environment, which can be useful in environments with partial observability or where complex coordination is required by the agents. For instance, a value function can be conditioned on the history of joint observations to better estimate the expected returns. However, central learning is often not feasible or applicable for multiple reasons: (1) The joint reward across all agents has to be transformed into a single reward for training, which might be difficult or impossible in general-sum games, (2) the central policy has to learn over the joint action space

which typically⁶² grows exponentially in the number of agents, and (3) agents might be physically or virtually distributed entities which might not allow for communication from and to a central policy for centralised control. Autonomous vehicles, for example, cannot be realistically expected to transmit and receive the sensor and camera information of all surrounding vehicles in real time. Furthermore, even if information sharing across vehicles was possible and instantaneous, learning a centralised control policy to control all vehicles would be very difficult due to the scale and complexity of the problem. In this case, decentralised control is a more reasonable approach to implement individual agents for each vehicle and to decompose the larger single-agent problem into multiple smaller multi-agent problems.

9.1.2 Decentralised Training and Execution

In Decentralised Training and Execution, the training of agent policies and the policies themselves are fully decentralised between the agents, meaning that they do not rely on centrally shared information or mechanisms. Decentralised training and execution is a natural choice for MARL training in scenarios in which agents lack the information or ability to be trained or executed in a central manner. Financial markets are an example of such a scenario. Trading individuals and companies cannot possibly know how other agents might act or how they affect the markets, and any such influence can only be partially observed.

An example of this category is independent learning (Section 5.3.2), in which each agent ignores the existence of other agents and trains its policy in a completely local way using single-agent RL techniques. Independent learning has the benefit of scalability by avoiding the exponential growth in action spaces of central learning, and is naturally applicable in scenarios where agents are physically or virtually distributed entities that cannot communicate with each other. However, independent learning has two downsides: (1) The agents' policies are not able to leverage information about other agents (neither during training of their policies nor for their execution), and (2) training can be significantly affected by non-stationarity caused by the concurrent training of all agents, as we discussed in Section 5.4.1. Effectively, with changes in the policies of other agents, the transition, observation and reward functions, as perceived by each individual agent, change as well. These changes can lead to unstable learning and poor convergence of independent learning. Despite these challenges, independent learning can often be found to perform well

62. In Section 5.4.4 we discussed an example in which the growth is not considered exponential.

in practice and serves as a first step to build more complex algorithms with decentralised execution. Building on independent learning algorithms with tabular value functions (Section 5.3.2), we will introduce deep independent learning algorithms in Section 9.3. We note that independent learning is not the only way decentralised training can be implemented. Agent modelling, for example, covers a variety of methods that can be used to model the changing behaviour of other agents in the environment (Section 6.3).

9.1.3 Centralised Training with Decentralised Execution

Centralised training and decentralised execution (CTDE) represents the third paradigm of MARL. These algorithms use centralised training to train agent policies, while the policies themselves are designed to allow for decentralised execution. For example, during training the algorithm may utilise the shared local information of all agents to update the agent policies, while each agent's policy itself only requires the agent's local observation to select actions, and can thus be deployed in a fully decentral fashion. In this way, CTDE algorithms aim to combine the benefits of both centralised training and decentralised execution.

CTDE algorithms are particularly common in deep MARL because they enable conditioning approximate value functions on privileged information in a computationally tractable manner. A multi-agent actor-critic algorithm, for example, may train a policy with a centralised critic which can be conditioned on the joint observation history and, therefore, be informed to provide more accurate estimation of the value of a state compared to a critic which only receives a local observation history. During execution, the value function is no longer needed since the action selection is done by the policy. To preserve decentralised execution, the policies of agents are only conditioned on the local observation history. We will further discuss deep MARL algorithms which make use of this paradigm in Section 9.4.

Using centralised training in competitive games requires further considerations, since it makes several restrictive assumptions. For example, the assumption that the training will include the same opponent policies as the evaluation is harder to justify. Even if they are the same opponents, it is unlikely that they will willingly provide information that will help each other during training. Training agents under the CTDE paradigm, however, can still be effective approach to train robust policies for competitive games. In this case, one could train a set of agents with the intention of creating a variety of opponent policies – and still share information during training to improve their robustness. After training, one or more policies can be selected from the pool of agents that were trained to compete against others. These policies can be executed

decentrally, and as such, the assumption that opponents do not share information with each other is maintained.

9.2 Notation for Multi-Agent Deep Reinforcement Learning

In line with the notation used in Chapter 8, we will use θ to denote the parameters of learned value functions, and ϕ to denote the parameters of policies. Similarly, the parameters of the value function and policy of agent i will be denoted with θ_i and ϕ_i , respectively. We will denote the policy, state-value function, and action-value function of agent i with $\pi(\cdot; \phi_i)$, $V(\cdot; \theta_i)$, and $Q(\cdot; \theta_i)$, respectively. To keep notation slim, we do not explicitly denote the policy and value functions with a subscript of the respective agent when it is clear from the parameterisation which agent is considered. For example, we will write $\pi(\cdot; \phi_i)$ instead of $\pi_i(\cdot; \phi_i)$. We note that this notation simplifies the distinction of networks of agents, since the networks of two agents might differ beyond the parameterisation, e.g. due to varying input and output dimensions for different observation and action spaces across agents.

In partially observable multi-agent games, agents only receive local observations about the environment which might be different across agents (Section 3.4). During centralised training, most commonly in the CTDE paradigm, agents might make use of joint information across all agents during training but condition their policies only on their local observation history. To represent this discrepancy of information available during training and execution, we will introduce all following MARL algorithms using notation for partially observable environments. For this purpose, we will use h to denote histories of observations. However, we note that some centralised training algorithms make use of the full state s of the environment during training. In these cases, we will specifically highlight where the local observation history $h_i^t = (o_i^0, o_i^1, \dots, o_i^t)$ of agent i , the joint observation history $h^t = (o^0, o^1, \dots, o^t)$, or the state s^t at time step t should be used. In environments where the full state is not available and only observations are practically accessible, the state of the environment can be approximated by the joint observation history $s^t \approx h^t$. In fully observable environments, agents should use the state s of the environment instead of individual or joint observation histories to make use of the full information available to agents.

Section 8.3 discussed the application of recurrent neural networks to efficiently condition deep value functions and policies on the history of observations. These networks can receive one observation at a time and internally represent

the observation history as a hidden state. Due to this practice and for notational brevity, many publications define the policy and value functions of deep RL algorithms as a function conditioned only on the most recent observation. Instead, we will explicitly condition the policy and value function networks at time step t on the history of local or joint observations, denoted with h_i^t and h^t , respectively.

9.3 Independent Learning

In MARL, multiple agents act and learn concurrently in a shared environment. When agents perceive other agents as part of the environment and learn using (single-agent) RL algorithms, we consider them to learn independently. Despite its simplicity, *independent learning* has been shown to perform competitively to more sophisticated algorithms in multi-agent deep RL (Palmer 2020; Papoudakis et al. 2021). In this section, we will show how to use existing deep RL algorithms (Chapter 8) to train multiple agents.

9.3.1 Independent Value-based Learning

Independent value-based learning consists of algorithms that aim to learn value functions that are conditioned on the observations and actions of individual agents. A representative example is the Independent DQN (IDQN) algorithm, where each agent trains its own action-value function $Q(\cdot; \theta_i)$, maintains a replay buffer \mathcal{D}_i and only learns from its own observation history, actions and rewards using DQN (Section 8.1.4). The DQN loss function for each agent i is

$$\mathcal{L}(\theta_i) = \frac{1}{B} \sum_{(h_i^t, a_i^t, r_i^t, h_i^{t+1}) \in \mathcal{B}} \left(r_i^t + \gamma \max_{a_i} Q(h_i^{t+1}, a_i; \bar{\theta}_i) - Q(h_i^t, a_i^t; \theta_i) \right)^2 \quad (9.1)$$

with $\bar{\theta}_i$ denoting the parameters of agent i 's target network. The final loss $\mathcal{L}(\theta_1) + \mathcal{L}(\theta_2) + \dots + \mathcal{L}(\theta_N)$ should be minimised simultaneously for each agent. We also present pseudocode of the IDQN algorithm in Algorithm 16.

It is worth noting that the replay buffer has specific disadvantages in IDQN. In multi-agent environments, the behaviour of an agent is not only determined by its own actions but also influenced by the actions of other agents in the environment. Therefore, the observation and action of an agent can lead to significantly different results depending on the policies of other agents. This creates a challenge when using a replay buffer, as it assumes that the stored experiences will remain relevant over time. However, in MARL, the policies of other agents are constantly changing as they learn, and this can make the experiences stored in the replay buffer quickly become outdated.

Algorithm 16 Independent Deep Q-Networks

-
- 1: Initialise n value networks with random parameters $\theta_1 \dots \theta_n$
 - 2: Initialise n target networks with parameters $\bar{\theta}_1 = \theta_1 \dots \bar{\theta}_n = \theta_n$
 - 3: Initialise a replay buffer for each agent D_1, D_2, \dots, D_n
 - 4: Collect environment observations $o_1^0 \dots o_n^0$
 - 5: **for** time step $t = 0, 1, 2, \dots$ **do**
 - 6: **for** agent $i = 1 \dots n$ **do**
 - 7: With probability ϵ : choose random action a_i^t
 - 8: Else: choose $a_i^t \in \arg \max_{a_i} Q(h_i^t, a_i; \theta_i)$
 - 9: Apply actions and collect observations o_i^{t+1} and rewards r_i^t .
 - 10: Store transitions in replay buffers D_1, D_2, \dots, D_n
 - 11: **for** agent $i = 1 \dots n$ **do**
 - 12: Sample random mini-batch of transitions from replay buffer D_i
 - 13: Update parameters θ_i by minimising the loss from Equation 9.1
 - 14: In a set interval, update target network parameters $\bar{\theta}_i$ for each agent i
-

To understand the problem that can occur with off-policy algorithms like IDQN that use a replay buffer to store experiences in multi-agent settings consider an example of two agents learning to play chess. Say that Agent 1 is using a specific opening that is initially effective, but is actually a weak strategy in the long run. Agent 2 has not yet learned to counter this opening and so is not penalizing agent 1 for using it. As agent 2 learns to counter the opening, the old opening sequences where agent 1 had success will still be stored in the replay buffer. Agent 1 will keep learning from these old examples, even though they are no longer relevant to the current state of the learning process, since the improved policy of agent 2 can counter those actions. This can lead to a situation where agent 1 continues to use the weak opening even after it has been countered by the agent 2.

To address this issue in multi-agent settings, smaller replay buffers may be used. As a result, the buffer will quickly reach its maximum capacity, and older experiences will be removed, which diminishes the likelihood of stored experiences becoming outdated and allows agents to learn from recent data. However, there are also more elaborate methods to address this challenge of non-stationarity with using a replay buffer. Foerster et al. (2017) proposed two methods to address the challenge of non-stationarity when using a replay buffer for MARL, based on importance sampling weights and fingerprints of agent policies. In the former approach, the replay buffer is extended by

storing policies alongside experience such that importance sampling weight correction can be applied to account for the changing probability of selecting an action. The second approach extends the observation of an agent with a fingerprint of the policy such that agents can consider the changing policies of other agents. Hysteretic Q-learning (Matignon, Laurent, and Le Fort-Piat 2007) uses smaller learning rates for updates of action values which would decrease these value estimates. This approach is motivated by the observation that these decreasing estimates might be the result of the stochasticity of other agents' policies. Similarly, the concept of leniency (Panait, Tuyls, and Luke 2008) ignores decreasing updates of action-value estimates with a given probability which decreases throughout training to especially consider the stochasticity of agent policies early in training. Both concepts of hysteretic and lenient learning have been applied to deep multi-agent RL algorithms (Omidshafiei et al. 2017; Palmer et al. 2018) and extended by distinguishing between negative updates as a consequence of miscoordination or stochasticity (Palmer, Savani, and Tuyls 2019).

9.3.2 Independent Policy Gradient Methods

Similarly to independent learning with value-based methods, policy gradient methods can independently be applied in MARL. To independently train each agent with the REINFORCE algorithm (Section 8.2.3) in multi-agent settings, each agent maintains its own policy and learns independently from its own experiences. The policy gradient is computed based on the agent's own actions and rewards, without taking into account the actions or policies of other agents.

Each agent can follow the policy gradient by computing the gradient of the expected return with respect to its own policy parameters. At the end of every episode, each agent updates its policy with the following policy gradient:

$$\begin{aligned}\nabla_{\phi_i} J(\phi_i) &= \mathbb{E}_{\pi} \left[u_i^t \frac{\nabla_{\phi_i} \pi(a_i^t | h_i^t; \phi_i)}{\pi(a_i^t | h_i^t; \phi_i)} \right] \\ &= \mathbb{E}_{\pi} [u_i^t \nabla_{\phi_i} \log \pi(a_i^t | h_i^t; \phi_i)]\end{aligned}\quad (9.2)$$

This gradient updates the policy parameters in the direction in which the probability of selecting an action increases ($\nabla_{\phi_i} \pi(a_i^t | h_i^t; \phi_i)$) proportional to the returns (u_i^t), with gradients being normalised by the inverse of the current probability of selecting the action under the policy ($\pi(a_i^t | h_i^t; \phi_i)$). The Independent REINFORCE algorithm is also shown in Algorithm 17.

In multi-agent settings, on-policy algorithms like REINFORCE have an advantage over off-policy algorithms in that they always learn from the most up-to-date policies of the other agents. This is because the policy gradient is computed based on the most recent experiences, which are generated by the

Algorithm 17 Independent REINFORCE

-
- 1: Initialise n actor networks with random parameters $\phi_1 \dots \phi_n$
 - 2: Collect environment observations $o_1^0 \dots o_n^0$
 - 3: **for** time step $t = 0, 1, 2, \dots$ **do**
 - 4: **for** agent $i = 1 \dots n$ **do**
 - 5: Sample actions a_i^t from $\pi(a_i^t | o_i^t; \phi_i)$
 - 6: Execute actions and collect observations o_i^{t+1} and rewards r_i^t .
 - 7: **for** agent $i = 1 \dots n$ **do**
 - 8: Update parameters ϕ_i by minimising the loss from Equation 9.2
-

agents' current policies. As the policies of the agents evolve over time, the experiences collected by each agent reflect the most up-to-date policies of the other agents in the environment. This feature of on-policy algorithms is important in multi-agent settings because the policies of the agents are constantly evolving. Learning from the most up-to-date policies of other agents enables each agent to adapt to changes in the environment or the policies of the other agents, leading to more stable learning.

Consider again the chess example from Section 9.3.1, which discussed how algorithms that use replay buffers may be unable to learn that an opening has been countered by the other agent. On-policy algorithms like REINFORCE are less susceptible to this problem because they always learn from the most up-to-date policies of the other agents. In the chess example, as agent 2 learns to counter the opening, the trajectories collected by agent 1 will immediately reflect that change. In this way, on-policy algorithms can adapt more quickly to changes in the policies of the other agents. This is particularly important in multi-agent settings where the policies of the agents are constantly evolving.

The A2C (Section 8.2.5) and PPO (Section 8.2.6) algorithms can also be extended similarly to REINFORCE and be applied independently in multi-agent settings. Let us now discuss the independent learning algorithm for A2C with multiple environments (Section 8.2.7). In A2C with parallel environments, the trajectories from the parallel environments along with the multiple agents form batches of higher dimensionality. For example, the observations collected from K environments on a timestep t form a two dimensional matrix:

$$\begin{bmatrix} o_1^{t,1} \dots o_n^{t,1} \\ \vdots \\ o_1^{t,K} \dots o_n^{t,K} \end{bmatrix}$$

Algorithm 18 Independent Learning with A2C

- 1: Initialise n actor networks with random parameters $\phi_1 \dots \phi_n$
 - 2: Initialise n critic networks with random parameters $\theta_1 \dots \theta_n$
 - 3: Initialise K parallel environments
 - 4: Build a batch of initial observations for each agent
 - 5: and environment:
$$\begin{bmatrix} o_1^{0,1} \dots o_n^{0,1} \\ \vdots \\ o_1^{0,K} \dots o_n^{0,K} \end{bmatrix}$$
 - 6: **for** timestep $t = 0 \dots$ **do**
 - 7: Sample actions
$$\begin{bmatrix} a_1^{t,1} \dots a_n^{t,1} \\ \vdots \\ a_1^{t,K} \dots a_n^{t,K} \end{bmatrix}$$
 from $\pi(a_i^t | o_i^t; \phi_i)$
 - 8: Execute actions and observe
$$\begin{bmatrix} o_1^{t+1,1} \dots o_n^{t+1,1} \\ \vdots \\ o_1^{t+1,K} \dots o_n^{t+1,K} \end{bmatrix}$$
 and rewards
$$\begin{bmatrix} r_1^{t,1} \dots r_n^{t,1} \\ \vdots \\ r_1^{t,K} \dots r_n^{t,K} \end{bmatrix}$$
.
 - 9: **for** agent $i = 1 \dots n$ **do**
 - 10: Update θ_i by minimising the loss from Equation 9.5 over the batch.
 - 11: Update ϕ_i by minimising the loss from Equation 9.3 over the batch.
-

The actions and the reward would also form such matrices. Calculating the final A2C loss requires iterating and summing over the individual losses. The policy loss of a single agent over the data collected from an environment k becomes

$$\mathcal{L}(\phi_i | k) = -\log \pi(a_i^{t,k} | h_i^{t,k}; \phi_i) \left(r_i^{t,k} + \gamma V(h_i^{t+1,k}; \theta_i) - V(h_i^{t,k}; \theta_i) \right) \quad (9.3)$$

and the final policy loss sums and averages over the batch

$$\mathcal{L}(\phi) = \frac{1}{K} \sum_{i \in I} \sum_{k=1}^K \mathcal{L}(\phi_i | k) \quad (9.4)$$

where i iterates over the agents and k over the environments.

The value loss also makes use of the batch by iterating over all its elements similarly to the policy loss:

$$\mathcal{L}(\theta_i | k) = \left(V(h_i^{t,k}; \theta_i) - y_i \right)^2 \text{ with } y_i = r_i^{t,k} + \gamma V(h_i^{t+1,k}; \bar{\theta}_i) \quad (9.5)$$

Pseudocode for Independent Learning with A2C (IA2C) is shown in Algorithm 18. Independently applying PPO does not require any other considerations and is very similar to IA2C.

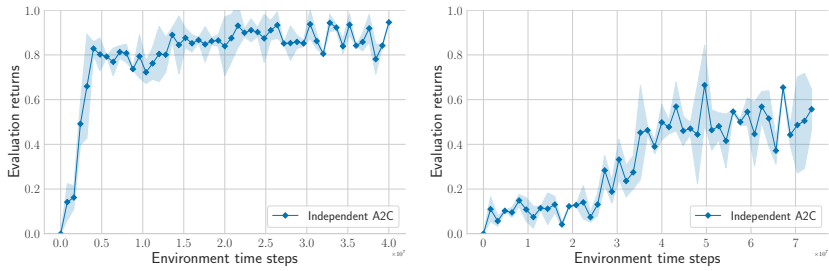
9.3.3 Example: Deep Independent Learning in a Large Task

Section 5.3.2 showed how independent learning can, with tabular MARL algorithms, learn on the level-based foraging environment. The level-based foraging environment used in the experiments of that section used the same initial state in each episode, so that the two agents and items started in the same locations and had the same levels in all episodes. With the 11×11 grid size, two agents and two items, the size of the state space can be calculated to be 42,602, which means that as many Q values for each agent’s action will need to be stored. While such a state space can be manageable for a tabular algorithm, any significant increases to the space will be limited by the algorithm’s need for maintaining large Q tables.

Independent learning algorithms such as IA2C make use of neural networks to learn policies and action-value functions. The ability of neural networks to generalise to similar states allows IA2C to handle environments with much larger state spaces. To demonstrate this, we train IA2C on level-based foraging with a grid size of 15×15 , and random initial locations and levels for the agents and items, which adheres to the open-sourced level-based foraging environment discussed in Chapter 11. This learning problem has a state space that is many orders of magnitude larger than the tasks we explored in Part I. Indicatively, two agents and two items in a 15×15 grid result in approximately 5 billion (5×10^9) combinations.

Our experiments with IA2C on the larger level-based foraging environment demonstrate the power of deep RL algorithms in tackling tasks with larger state spaces. As shown in Figure 9.1a, IA2C learned a joint policy that collects all the available items, which is indicated by the evaluation returns reaching values close to 1 (as detailed in Section 11.3.1). This result was reached within 40,000,000 environment timesteps and required approximately 3 hours on typical hardware (running on an Intel i7-2700K CPU), indicating its scalability to environments with much larger state spaces.

We extended our experiments to the case of three agents and three items in the level-based foraging environment, which results in a state space of approximately three hundred trillion possible states (3.6×10^{14}). As shown in Figure 9.1b, IA2C still learned to navigate the environment and collect some of the items (a return of 0.5 signifies that on average half of the items are collected in each episode) in less than 6 hours on the same hardware (Intel i7-2700K CPU), even in the presence of multiple agents and a much larger state space. These results highlight the potential of deep MARL algorithms, such as IA2C, for solving complex multi-agent environments.



(a) Task with two agents and two items. (b) Task with three agents and three items.

Figure 9.1: Independent A2C (IA2C) algorithm in the level-based foraging environment with a 15×15 grid, and (a) two agents and two items, (b) three agents and three items. Each episode starts with random initial locations and levels for agents and items. IA2C used 8 parallel environments, N -step returns with $N = 10$, a learning rate of $3e-4$ and two neural networks with 2 hidden layers of 64 units for the actor and critic networks. The discount factor for the environment was set to $\gamma = 0.99$.

9.4 Multi-Agent Policy Gradient Algorithms

So far in this chapter, we discussed deep independent learning algorithms for MARL. These algorithms extend single-agent RL algorithms with deep neural networks for value function and policy approximation to multi-agent RL. In Section 5.4.1, we discussed the problem of non-stationarity in RL and explained how multi-agent learning and partial observability exacerbate it. Independent learning suffers from this problem in particular, as each agent perceives other agents as part of the environment, rendering the environment non-stationary from each agent’s perspective. We can, however, use the CTDE paradigm (Section 9.1.3) to mitigate the effects of non-stationarity. Under this paradigm, agents can share information during training to stabilise learning, as long as they are still able to execute their policies in a decentralised manner. In this section, we will focus on how to apply CTDE to policy gradient algorithms, where centralised training allows us to train value functions conditioned on information of all agents. First, we extend the policy gradient theorem to multi-agent RL, then discuss how to train centralised critics as state-value functions and, finally, how to train centralised critics as action-value functions conditioned on the state and actions of all agents.

9.4.1 Multi-Agent Policy Gradient Theorem

The policy gradient theorem (Section 8.2.2) is the foundation of all single-agent policy gradient algorithms which define various update rules for the parameters of a parameterised policy. As a reminder, the policy gradient theorem states that the gradients of the quality of a parameterised policy, as given by its expected returns, with respect to the policy parameters can be written as follows:

$$\nabla_{\phi} J(\phi) \propto \sum_{s \in S} \Pr(s | \pi) \sum_{a \in A} Q^{\pi}(s, a) \nabla_{\phi} \pi(a | s; \phi) \quad (9.6)$$

$$= \mathbb{E}_{\pi} [Q^{\pi}(s, a) \nabla_{\phi} \log \pi(a | s; \phi)] \quad (9.7)$$

But can the policy gradient theorem be extended to the setting of MARL? Fortunately, we can define the *multi-agent policy gradient theorem* by considering that the expected returns received by any agent's policy in multi-agent RL are dependent on the policies of all agents. Using this insight, we can write the multi-agent policy gradient theorem for the policy of agent i with an expectation over the policies of all agents:

$$\nabla_{\phi} J(\phi_i) \propto \mathbb{E}_{a_i \sim \pi_i, a_{-i} \sim \pi_{-i}} [Q_i^{\pi}(s, \langle a_i, a_{-i} \rangle) \nabla_{\phi_i} \log \pi_i(a_i | s; \phi_i)] \quad (9.8)$$

Similar to the single-agent policy gradient theorem, the multi-agent policy gradient theorem can be used to derive various policy gradient update rules by estimating the expected returns. We have already seen several instantiations of multi-agent policy gradient algorithms in the form of independent learning policy gradient algorithms (Section 9.3.2). For these algorithms, the expected return estimate of agent i , given by $Q_i(s, \langle a_i, a_{-i} \rangle)$, is estimated using a value function which is only conditioned on the history of observations and action of agent i itself: $Q_i(h_i, a_i) \approx Q_i(s, \langle a_i, a_{-i} \rangle)$. In the following, we will focus on the CTDE paradigm and derive estimates of the expected return which are conditioned on centralised information. In particular, we will see that we can obtain more precise estimates of expected returns when using the state of the environment and the actions of all agents. We will then use these value functions to derive multi-agent policy gradient algorithms under the CTDE paradigm.

9.4.2 Centralised State-Value Critics

To define an actor-critic algorithm under the CTDE paradigm, we have to consider both the actor and critic networks. The actor network was previously defined as $\pi(h_i^t; \phi_i)$. With this definition, the actor network requires only the local observation history of agent i to select its actions. This ability is critical to ensure decentralised execution and, therefore, must stay as-is.

Algorithm 19 Centralised A2C

```

1: Initialise  $n$  actor networks with random parameters  $\phi_1 \dots \phi_n$ 
2: Initialise  $n$  critic networks with random parameters  $\theta_1 \dots \theta_n$ 
3: Collect environment observations  $o_1^0 \dots o_n^0$ 
4: for time step  $t = 0 \dots$  do
5:   for agent  $i = 1 \dots n$  do
6:     Sample actions  $a_i^t$  from  $\pi(a_i^t | h_i^{1:t}; \phi_i)$ 
7:     Execute actions and collect state  $s^t$ , observations  $o_i^{t+1}$  and rewards  $r_i^t$ .
8:     for agent  $i = 1 \dots n$  do
9:       Update parameters  $\theta_i$  by minimising the loss from Equation 9.9
10:      Update parameters  $\phi_i$  by minimising the loss from Equation 9.3

```

However, the critic network does not have this restriction *during training*. Indeed, the network is discarded after training, and only the actor is used to generate the agent actions. Therefore, a decentralised version of a critic network is not required and can be replaced with a centralised one. We can redefine the critic as $V(s^t; \theta_i)$, a network that is conditioned on the full state of the environment but still approximates the value of the state for agent i . The advantage is clear: the network has access to the full state, allowing it to make use of more information of the environment and other agents. Access to such centralised information is particularly valuable in partially observable environments, where the critic would otherwise lack potentially relevant information to estimate values of states, and to reduce the impact of non-stationarity. By having access to information about all other agents, a centralised critic facilitates the adaptation to varying behaviour of other agents. As stated in Section 9.2, the full state is often unavailable in multi-agent environments. In this case, we can use the joint observation history h^t as an approximation to the full state of the environment. The value loss of this *centralised state-value critic*, shown in Figure 9.2 becomes:

$$\mathcal{L}(\theta_i) = (V(s^t; \theta_i) - y_i)^2 \quad \text{with } y_i = r_i^t + \gamma V(s^{t+1}; \theta_i). \quad (9.9)$$

Any independent actor-critic RL algorithm which learns a state-value function (Section 9.3.2) can be instantiated with a centralised state-value critic to learn a value function in MARL. Algorithm 19 presents pseudocode for the centralised A2C algorithm which can be seen as multi-agent A2C with a centralised critic. For notational clarity, we present the pseudocode of multi-agent policy gradient

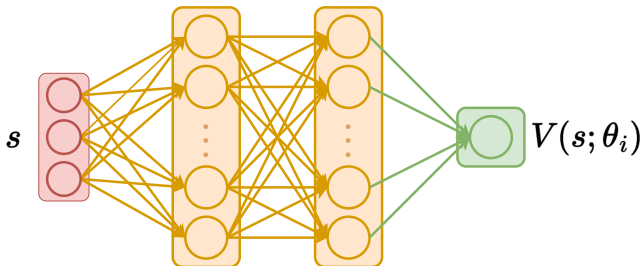
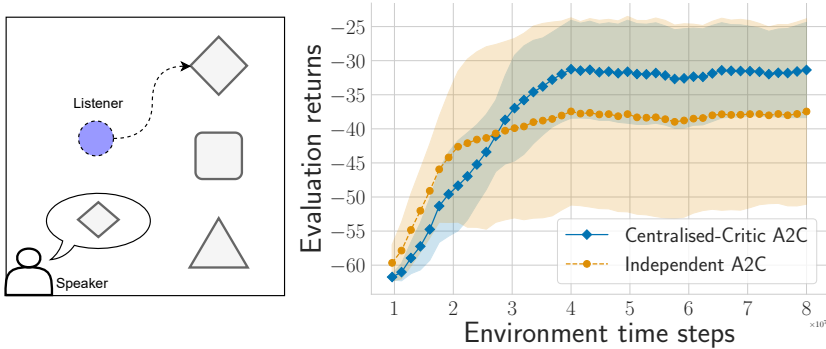


Figure 9.2: Architecture of a centralised state-value critic for agent i . The critic is conditioned on the full state of the environment and outputs a single scalar to represent the approximated value of the state.



(a) Speaker-listener game.

(b) Training curves.

Figure 9.3: Using a centralised state-value critic with A2C for the Speaker-Listener game allows the agents to learn the task by addressing the partial observability of the environment. With the centralised critic, the algorithm converges to higher returns than IA2C.

algorithms in this section for training in a single environment. To train multi-agent policy gradient algorithms with multiple parallel environments, we refer to the pseudocode of IA2C (Algorithm 18).

In practice, such a centralised critic can lead to more robust learning. Consider the speaker-listener game shown in Figure 9.3a (see also Section 11.3.2 for more information). This is a common-reward game, in which two agents need to cooperate to reach their goals. One agent, the listener, is placed in the environment and can observe its location and that of three distinct landmarks (shapes in Figure 9.3a). The other agent, the speaker, can only observe the

shape of the landmark that will maximize the common reward and can transmit an integer from 1 to 3 to the listener. The goal of the game is for the two agents to learn to cooperate such that the listener always moves to the goal landmark. Given the partial observability, this game is quite challenging: the speaker has to learn to recognise the different shapes and transmit a distinct message long enough for the listener to learn to move to the correct landmark. Conditioning the critic of each agent on the full state (the landmark goal and the positions of the agent and landmarks) allows them to learn despite the partial observability, and leads to the performance advantages seen in Figure 9.3b.

9.4.3 Centralised Action-Value Critics

As we have seen, a centralised state-value critic can stabilise the training of multi-agent actor-critic algorithms, in particular in partially observable multi-agent environments. However, it can be desirable to learn action-value functions as critics instead. These value functions condition their value estimation not just on the current state, but also on the actions of agents. To train a centralised action-value critic for multi-agent actor-critic algorithms, similar to the setting described in Section 9.4.2, each agent i trains a policy π_i which is conditioned on agent i 's local observation history. For the critic, agent i trains an action-value function Q_i which is conditioned on the full state and actions of all agents. If the full state is not available, we use the joint observation history instead. We can instantiate this idea by training the centralised critic to minimise the following value loss:

$$\mathcal{L}(\theta_i) = (Q(s^t, a^t; \theta_i) - y_i)^2 \quad \text{with } y_i = r_i^t + \gamma Q(s^{t+1}, a^{t+1}; \theta_i) \quad (9.10)$$

For this loss, we compute the target value y_i for agent i using the next state and the next actions applied by all agents, similar to the on-policy Sarsa algorithm (Equation 2.52, page 33). Using the critic, we can define the policy loss for agent i as:

$$\mathcal{L}(\phi_i) = -\log \pi(a_i^t | h_i^t; \phi_i) Q(s^t, a^t; \theta_i) \quad (9.11)$$

Using the centralised critic, the multi-agent policy gradient for agent i is given by:

$$\nabla_{\phi_i} J(\phi_i) = \mathbb{E}_{\pi} [\nabla_{\phi_i} \log \pi_i(a_i^t | h_i^t; \phi_i) Q_i(s^t, a^t; \theta_i)] \quad (9.12)$$

We have previously seen action-value functions in value-based RL algorithms like DQN. These algorithms optimise their value function with the max operator over the next action, and use off-policy batches of experiences sampled from a replay buffer. To understand why we do not use these techniques to train the action-value critic for multi-agent actor-critic algorithms, we have to remind ourselves that the multi-agent policy gradient theorem requires the estimation

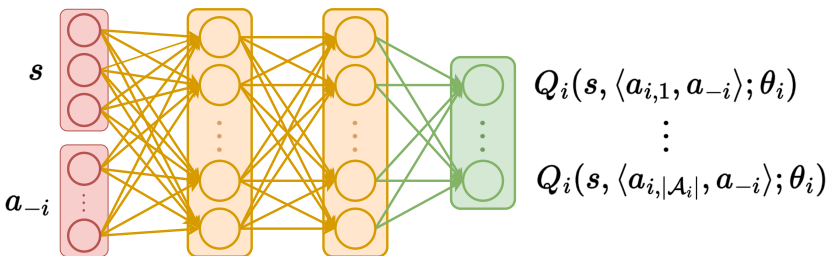


Figure 9.4: Architecture of a centralised action-value critic for agent i . The critic is conditioned on the full state of the environment and the actions of all other agents. The network outputs a single scalar for every action of agent i to represent the approximated value of the state and respective joint actions.

of the expected returns *under the current policies* of all agents. A replay buffer contains off-policy data which does not represent the distribution of experiences under the current policies. Likewise, DQN updates directly train the critic to approximate the optimal returns instead of the expected returns under the current policies. Therefore, we cannot use these techniques, despite optimising an actor-value function for the critic.

Naively modelling the centralised action-value critic as a neural network with the state as input and one action-value for each joint action as output is difficult, because the dimensionality of the output would scale with the joint action space which grows exponentially with the number of agents. To avoid this, we can instead model the action-value critic of agent i to receive the actions of all other agents a_{-i} as input. Then, the network only computes a single output for each action of agent i which corresponds to the action-value for the input state and the joint action given by the concatenation of the particular action of agent i and the joint action of all other agents a_{-i} . This architecture is illustrated in Figure 9.4.

9.4.4 Counterfactual Action-Value Estimation

So far, we have defined an algorithm which replaces the centralised state-value critic with a centralised action-value critic. The motivation for training an action-value function instead of a state-value function was its ability to directly estimate the impact of the action selection on the expected returns. However, a state-value function can be used to approximate the advantage estimate (Equation 8.33) which also provides preferences over particular actions. Additionally, training an action-value function with one output for every possible action may be more difficult than training a state-value function with only a single output since each

individual experience only updates the action-value output for a single action. Given these considerations, why would we want to train action-value critics for multi-agent actor-critic algorithms?

One idea leverages action-value critics to address the multi-agent credit assignment problem (Section 5.4.2) based on the concept of *difference rewards* (Wolpert and Tumer 2002; Tumer and Agogino 2007). Difference rewards approximate the difference between the received reward and the reward agent i would have received if they had chosen a different action \tilde{a}_i :

$$d_i = \mathcal{R}(s, \langle a_i, a_{-i} \rangle) - \mathcal{R}(s, \langle \tilde{a}_i, a_{-i} \rangle) \quad (9.13)$$

The action \tilde{a}_i is also referred to as the *default action*. Difference rewards aim to consider the counterfactual question of “Which reward would agent i have received if they instead had selected their default action?”. Answering this question is valuable in settings where all agents receive a common reward, because it provides information about the concrete contribution of agent i to the received reward. However, computing difference rewards in practice is often difficult because (1) it is not clear how to select the default action for agent i , and (2) computing $\mathcal{R}(s, \langle \tilde{a}_i, a_{-i} \rangle)$ requires access to the reward function. The need to determine a default action for each agent can be avoided by using the definition of the *aristocrat utility* (Wolpert and Tumer 2002):

$$d_i = \mathcal{R}(s, \langle a_i, a_{-i} \rangle) - \mathbb{E}_{a'_i \sim \pi_i(\cdot | h_i)} [\mathcal{R}(s, \langle a'_i, a_{-i} \rangle)] \quad (9.14)$$

Given access to the reward function, **castellini2020difference** derive return estimates over difference rewards determined by the aristocrat utility and incorporate them into the REINFORCE algorithm for MARL. If the reward function is not available, they propose to learn a model of the reward function from experience in the environment, and use this model to estimate the difference rewards.

Counterfactual multi-agent policy gradient (COMA) (Foerster et al. 2018) uses the same concepts to derive a centralised action-value critic to compute a counterfactual baseline, which marginalises out the action of agent i to estimate the advantage for selecting action a_i over following the current policy π_i :

$$Adv_i(s, h_i, a) = Q(s, a) - \underbrace{\sum_{a'_i \in A_i} \pi_i(a'_i | h_i) Q(s, \langle a'_i, a_{-i} \rangle)}_{\text{counterfactual baseline}} \quad (9.15)$$

This advantage estimate looks similar to the one discussed in Section 8.2.5, but its baseline is following the aristocrat utility computing the expected centralised value estimate with agent i following its own policy and actions of other actions being fixed. This baseline is shown to not change the multi-agent policy gradient (Equation 9.8) in expectation, and can efficiently be computed

	A	B
A	4,4‡	0,3
B	3,0	2,2†

Figure 9.5: The stag hunt matrix game, also seen in Section 5.4.3. The Pareto-dominated equilibrium is denoted with \dagger and the Pareto-optimal one with \ddagger .

using the previously introduced architecture for centralised action-value critics (Figure 9.4). To train the policy of agent i in COMA, the action-value estimate in Equation 9.11 is replaced by the advantage estimate in Equation 9.15. Despite its clear motivation, COMA empirically suffers from high variance in its baseline (Kuba et al. 2021) and inconsistent value estimates (Vasilev et al. 2021) which result in unstable training that can lead to poor performance (Papoudakis et al. 2021).

9.4.5 Equilibrium Selection with Centralised Action-Value Critics

A centralised action-value critic offers flexibility when computing the advantage. Christianos, Papoudakis, and Albrecht (2023) alter the advantage to guide the learning agents to a Pareto-optimal equilibrium in no-conflict games. No-conflict games are a class of games where all agents agree on the most-preferred outcome. A list of 2×2 no-conflict matrix games can be found Section 11.2. Formally, a game is no-conflict if:

$$\arg \max_{\pi} U_i(\pi) = \arg \max_{\pi} U_j(\pi) \quad \forall i, j \in I \quad (9.16)$$

An example is the stag hunt game (Figure 9.5), which is a no-conflict game with two agents and was previously discussed in Section 5.4.3. In the stag hunt game, both agents prefer the outcome (A, A), making it no-conflict, although they disagree on the second-best outcome: (B, A) for agent 1 and (A, B) for agent 2. These two joint actions do not represent Nash equilibria, however, since agents can unilaterally improve their rewards by changing their actions. Finally, action (B, B) is a Nash equilibrium since an agent changing its action (with the other agent not doing so) decreases its reward.

When such games have multiple Nash equilibria, learning agents tend to converge to less risky equilibria by preferring less risky actions (Papoudakis et al. 2021). In the stag hunt example, agents selecting action B are guaranteed a reward of 2, while action A might lead to a reward of 0. Therefore, even if the highest reward can only be achieved by choosing action A, the agents tend to learn the suboptimal (B, B) solution. This preference is easy to illustrate

	A	B	C
A	11‡	-30	0
B	-30	7†	0
C	0	6	5

Figure 9.6: The climbing matrix game. The climbing game is a common-reward (which is always no-conflict) game that shares similar characteristics to the stag-hunt game: an optimal Nash equilibrium and joint actions which are less rewarding but are easier to reach. The Pareto-dominated equilibrium is denoted with † and the Pareto-optimal with ‡.

for agents that do not model the actions of the other agents. Suppose the two agents are initialised with a uniform random policy, $\pi(A)=\pi(B)=0.5$, before learning starts. At that moment, the expected reward of choosing action A for agent 1 is $0.5 * 4 + 0.5 * 0 = 2.0$ and for action B the expected reward is $0.5 * 3 + 0.5 * 2 = 2.5$ (where 0.5 is the probability of the other agent selecting A or B). Learning the resulting Q values or following the policy gradient results in both agents gradually preferring B even more, making action A even less attractive for the other agent.

Pareto Actor-Critic (ParetoAC) (Christianos, Papoudakis, and Albrecht 2023) addresses this problem in no-conflict games by incorporating into the policy gradient the fact that other agents have the same most preferred outcome. The algorithm assumes that the policy of the other agents is

$$\pi_{-i}^{\text{pareto}} = \arg \max_{a_{-i}^t} Q(a_{-i}^t | o^t, a_i^t; \theta_i) \quad (9.17)$$

and modifies the original Nash equilibrium objective (Sections 4.2 and 4.4) to

$$\pi_i \in \text{BR}_i(\pi_{-i}^{\text{pareto}}) \quad \forall i \in I. \quad (9.18)$$

This objective leads to a policy that can be optimised with the gradient:

$$\nabla_{\phi_i} J(\phi_i) = \mathbb{E}_{\pi_i, \pi_{-i}^{\text{pareto}}} [\nabla_{\phi_i} \log \pi_i(a_i^t | h_i^t; \phi_i) Q(s^t, a_i^t, a_{-i}^t; \theta_i)] \quad (9.19)$$

With ParetoAC, agents tend to converge to more rewarding equilibria in no-conflict games even if they are riskier. We show experiments in two environments to illustrate the difference of ParetoAC to Centralised A2C. First, in Figure 9.7a, ParetoAC is shown to converge to the Pareto-optimal equilibrium (A, A) in the climbing game (Figure 9.6) while the A2C algorithm with a centralised state-value only converges to action (B,B) which is a suboptimal solution for both agents. However, matrix games are not the only situation in which this problem appears. Take the example of level-based foraging (Section 5.4.2) with two agents and an item that always needs both agents to

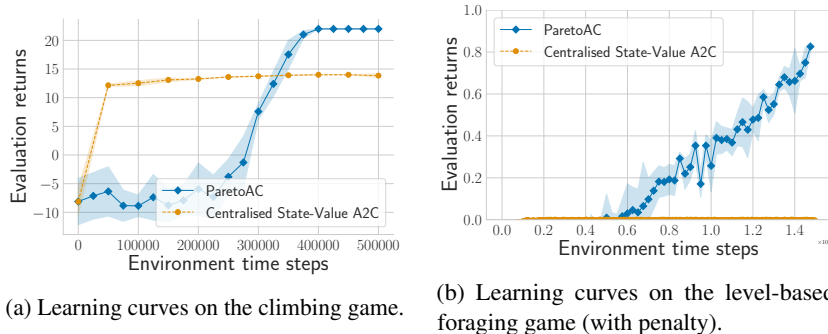


Figure 9.7: Training curves comparing centralised state-value A2C with ParetoAC in the climbing and level-based foraging games.

cooperate to collect it. In addition, apply a penalty (-0.6 in this example) if an agent attempts to gather an item alone and fails. Such a game has a suboptimal equilibrium of never trying to gather items in order to avoid getting penalised. In Figure 9.7b, we can see the results of ParetoAC and a centralised state-value algorithm in level-based foraging with two agents that always need to cooperate to gather an item that has been placed on a 5×5 grid. A2C quickly learns to avoid collecting the item so as to not receive any penalties. In contrast, ParetoAC is optimistic by using the centralised action-value critic and eventually learns to solve the task.

ParetoAC is an example of how a centralised action-value function can be used to improve learning in multi-agent problems. The centralised action-value function is not only used to learn a joint-action value but also to guide the policy gradient to the most promising equilibria. With its decentralised actors which are only conditioned on the observations, the algorithm adheres to the CTDE paradigm enabling the individual agents to execute their actions independently during execution.

9.5 Value Decomposition in Common-Reward Games

As we have seen in Section 9.4, centralised value functions can be used to address or mitigate several challenges in MARL, such as non-stationarity, partial observability (Section 9.4.2), multi-agent credit assignment (Section 9.4.4), and equilibrium selection (Section 9.4.5). One challenge of centralised value functions is that they are difficult to learn directly. In particular, centralised action-value functions are difficult to learn due to the exponential growth of

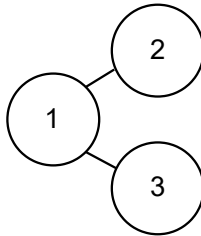


Figure 9.8: Example of a simple coordination graph. Each node represents an agent, and each edge connects agents that interact with each other. In this example, agent 1 interacts with agent 2 and agent 3. However, agent 2 and agent 3 do not interact with each other.

the joint action space with the number of agents. Motivated by this challenge, we discuss the following question in this section: how can agents effectively learn an action-value function that maximises the common reward and, equally important, understand their contribution to the common reward?

There is a long history of research on how value functions can be factorised to facilitate learning them. At first, Crites and Barto (1998) employed independent learners that attempt to learn $Q_i(s, a_i) \approx Q(s, a)$. However, it soon became evident that the underlying joint action-value function can be difficult to learn and cannot be approximated as easily. A key observation that helped simplify the problem was that not all agents interact with each other, and the interacting agents could be represented as a graph, known as a *coordination graph* (Guestrin, Koller, and Parr 2001; Guestrin, Lagoudakis, and Parr 2002; Kok and Vlassis 2005). The sparsity of the coordination graph can be exploited to approximate the joint action-value as the sum of the values of interacting agents, which can be easier to estimate. In the coordination graph example shown in Figure 9.8, in which two agents interact with a third but not with each other, the joint action value $Q(s, a_1, a_2, a_3)$ can be approximated as the sum $Q(s, a_1, a_2) + Q(s, a_1, a_3)$. Numerous studies have since explored approaches to learning near-exact (Oliehoek, Witwicki, and Kaelbling 2012) or approximate (Oliehoek 2010; Oliehoek, Whiteson, Spaan, et al. 2013) versions of these value functions, and applied these methods to the deep RL setting (Pol 2016; Böhmer, Kurin, and Whiteson 2020).

In this section, we will discuss more recent methods that have been successful at learning action-value functions using deep learning. These value decomposition algorithms decompose centralised value functions into simpler functions which can be learned more efficiently in common-reward games, in which all agents have the same reward functions, i.e. $\mathcal{R}_i = \mathcal{R}_j$ for all $i, j \in I$. In this setting,

all agents have the same objective and can benefit from a centralised value function which accurately estimates the expected returns over the common rewards. The centralised action-value function $Q(s, a; \theta)$, conditioned on the state and joint action, can be written as

$$Q(s^t, a^t; \theta) = \mathbb{E} \left[\sum_{\tau=t}^{\infty} \gamma^{\tau-t} r^{\tau} \mid s^t, a^t \right] \quad (9.20)$$

where r^{τ} denotes the common reward at time step τ .

9.5.1 Individual-Global-Max Property

One natural way to decompose the centralised action-value function is to learn individual utility functions for each agent. The utility function of agent i , written as $Q(h_i, a_i; \theta_i)$, is conditioned only on the individual observation history and action of the agent. These functions can use the same architecture as action-value functions (Figure 8.2), but we refer to them as utility functions because they are not optimised to approximate the expected returns of their respective agents. Instead, the utility functions of all agents are jointly optimised to approximate the centralised action-value function while satisfying the *individual-global-max* (IGM) property. To define the IGM property, we first define the sets of greedy actions with respect to a decomposed centralised action-value function and the individual utility function of agent i , respectively,

$$A^*(s; \theta) = \arg \max_{a \in A} Q(s, a; \theta) \quad (9.21)$$

$$A_i^*(h_i; \theta_i) = \arg \max_{a_i \in A_i} Q(h_i, a_i; \theta_i) \quad (9.22)$$

with $Q(s, a; \theta)$ and $Q(h_i, a_i; \theta_i)$ denoting the centralised action-value function and individual utility function of agent i , respectively.

Formally, the IGM property is satisfied if the following holds for all full histories \hat{h} with corresponding observation histories of each agent, $h_i = \sigma_i(\hat{h})$, and last state $s = s(\hat{h})$:⁶³

$$\forall a = (a_1, \dots, a_n) \in A : a \in A^*(s; \theta) \iff \forall i \in I : a_i \in A_i^*(h_i; \theta_i) \quad (9.23)$$

Intuitively, the IGM property states that if a joint action is greedy with respect to the centralised action-value function, i.e. $(a_1, \dots, a_n) \in A^*(s; \theta)$, then each agent i 's action within this joint action is also a greedy action with respect to its individual utility function, i.e. $a_i \in A_i^*(h_i; \theta_i)$. Likewise, if each agent i selects

63. Recall from Section 4.1 that a full history \hat{h} contains the history of states, joint observations, and joint rewards; $\sigma(\hat{h})$ returns the history of joint observations within full history \hat{h} ; and $s(\hat{h})$ denotes the last state in the full history. We use $\sigma_i(\hat{h})$ to denote the observation history of agent i .

a greedy action $a_i^* \in A_i^*(h_i; \theta_i)$ with respect to its individual utility function, then the joint action of all individually greedy actions itself is a greedy joint action with respect to the decomposed centralised action-value function, i.e. $(a_1^*, \dots, a_n^*) \in A^*(s; \theta)$.

If a decomposition upholds the IGM property, then each agent can decentrally follow the greedy policy with respect to its individual utility function, and all agents together will select the greedy *joint* action with respect to the decomposed centralised action-value function. During the optimisation of the decomposed centralised action-value function, the target computation with the greedy joint action of the decomposed centralised action-value function can be efficiently computed by computing the greedy individual actions of all agents with respect to their individual utility. If the individual utility functions satisfy the IGM property for the centralised action-value function, we also say that the utility functions *factorise* the centralised value function (Son et al. 2019). This property in the literature is also known as *consistency* (Rashid et al. 2018).⁶⁴

Besides providing an easier-to-learn decomposition of the centralised action-value function and enabling decentralised execution, the individual utilities learned by value decomposition algorithms can provide an estimate for the contribution of each agent to the common reward. This is because the individual utility functions are jointly optimised to approximate the centralised action-value function in aggregation, and individually are only conditioned on the local observation history and action of their corresponding agent. Hence, if an agent contributed to the common reward with its action, then its utility should approximate such contribution. In this way, value decomposition can address the multi-agent credit assignment problem (Section 5.4.2), similarly to the counterfactual action-value estimation seen in Section 9.4.4.

9.5.2 Linear Value Decomposition

A simple method of decomposing $Q(s, a; \theta)$ that satisfies the IGM property is to assume a linear decomposition of common rewards, i.e. the sum of the individual utilities of agents equals the common reward

$$\bar{r}^t = \bar{r}_1^t + \dots + \bar{r}_n^t \quad (9.24)$$

where \bar{r}_i^t denotes the utility of agent i at time step t . The bar over the reward symbol denotes that these utilities are obtained by the decomposition, and do

64. Prior work typically defines the IGM property as the equivalence of the greedy joint action with respect to $Q(s, a; \theta)$ and the joint action of the individually greedy actions. However, these definitions disregard the possibility that there could be *multiple* greedy actions and, thus, we define the IGM property as the equivalence of the sets of individually greedy actions and the set of greedy joint actions.

not represent true rewards received by the environment. Using this assumption, the centralised action-value function of agents can be decomposed as follows:

$$Q(s^t, a^t; \theta) = \mathbb{E}_\pi \left[\sum_{\tau=t}^{\infty} \gamma^{\tau-t} r^\tau \mid s^t, a^t \right] \quad (9.25)$$

$$= \mathbb{E}_\pi \left[\sum_{\tau=t}^{\infty} \gamma^{\tau-t} \left(\sum_{i \in I} \bar{r}_i^\tau \right) \mid s^t, a^t \right] \quad (9.26)$$

$$= \sum_{i \in I} \mathbb{E}_\pi \left[\sum_{\tau=t}^{\infty} \gamma^{\tau-t} \bar{r}_i^\tau \mid s^t, a^t \right] \quad (9.27)$$

$$= \sum_{i \in I} Q(h_i^t, a_i^t; \theta_i) \quad (9.28)$$

Any such linear decomposition satisfies the IGM property (Equation 9.23), as we will show below:

Proof. Let \hat{h} be a full history, $h_i = \sigma_i(\hat{h})$ be the observation history of agent i , and $s = s(\hat{h})$ denote the last state within \hat{h} . We will first prove that for any greedy joint action $a^* \in A^*(s; \theta)$, the individual actions in a^* of all agents are also greedy with respect to their individual utility functions. Then, we will show that any greedy individual actions of all agents together represent a greedy joint action with respect to $Q(s, a; \theta)$.

" \Rightarrow " Let $a^* = (a_1^*, \dots, a_n^*) \in A^*(s; \theta)$ be a greedy joint action with respect to $Q(s, \cdot; \theta)$. To prove that the individual actions of all agents a_1^*, \dots, a_n^* are also greedy with respect to agents' individual utility functions, i.e.

$$\forall i \in I : a_i^* \in A_i^*(h_i; \theta_i), \quad (9.29)$$

we need to show that for any agent i and any action $a_i \in A_i$, we have

$$\forall i \in I : Q(h_i, a_i^*; \theta_i) \geq Q(h_i, a_i; \theta_i). \quad (9.30)$$

We will prove this by contradiction. Assume that there exists an agent i and an action $a_i \in A_i$ such that

$$Q(h_i, a_i^*; \theta_i) < Q(h_i, a_i; \theta_i). \quad (9.31)$$

Given the linear decomposition of $Q(s, a; \theta)$, we have

$$Q(s, a^*; \theta) = \sum_{i \in I} Q(h_i, a_i^*; \theta_i) \quad (9.32)$$

$$= Q(h_i, a_i^*; \theta_i) + \sum_{j \in I \setminus \{i\}} Q(h_j, a_j^*; \theta_j) \quad (9.33)$$

$$< Q(h_i, a_i; \theta_i) + \sum_{j \in I \setminus \{i\}} Q(h_j, a_j^*; \theta_j) \quad (9.34)$$

$$(9.35)$$

which contradicts the assumption that a^* is a greedy joint action with respect to $Q(s, a; \theta)$. Therefore, Equation 9.31 cannot hold, thus, Equation 9.30 must be true and the individual actions of all agents within the greedy joint action a^* are also greedy with respect to their individual utility functions.

" \Leftarrow " Let $a_1^* \in A_1^*(h_1; \theta_1), \dots, a_n^* \in A_n^*(h_n; \theta_n)$ be greedy individual actions of all agents with respect to their individual utility functions. Let $a^* = (a_1^*, \dots, a_n^*)$ be the joint action composed of such greedy individual actions of all agents. To prove that a^* is a greedy joint action with respect to $Q(s, a; \theta)$, we need to show that for any joint action $a' \in A(s; \theta)$, we have

$$Q(s, a^*; \theta) \geq Q(s, a'; \theta). \quad (9.36)$$

Given the linear decomposition of the centralised action-value function, we have

$$Q(s, a^*; \theta) = \sum_{i \in I} Q(h_i, a_i^*; \theta_i) \quad (9.37)$$

$$= \sum_{i \in I} \max_{a_i} Q(h_i, a_i; \theta_i) \quad (9.38)$$

$$\geq \sum_{i \in I} Q(h_i, a'_i; \theta_i) \quad (9.39)$$

$$= Q(s, a'; \theta) \quad (9.40)$$

where $a' = (a'_1, \dots, a'_n)$ is any joint action composed of arbitrary individual actions $a'_1 \in A_1, \dots, a'_n \in A_n$. Therefore, a^* is a greedy joint action with respect to $Q(s, a; \theta)$ (Equation 9.36).

This concludes the proof. \square

The introduced linear decomposition defines the approach of *Value Decomposition Networks* (VDN) (Sunehag et al. 2017). VDN maintains a replay buffer \mathcal{D} containing the experience of all agents and jointly optimises the following loss over the approximate centralised value function for all agents. The loss

is computed over a batch \mathcal{B} sampled from the replay buffer and propagates its optimisation objective through the individual utilities of all agents

$$\mathcal{L}(\theta) = \frac{1}{B} \sum_{(h^t, a^t, r^t, h^{t+1}) \in \mathcal{B}} \left(r^t + \gamma \max_a Q(h^{t+1}, a; \bar{\theta}) - Q(h^t, a^t; \theta) \right)^2 \quad (9.41)$$

with

$$Q(h^t, a^t; \theta) = \sum_{i \in I} Q(h_i^t, a_i^t; \theta_i) \text{ and} \\ \max_a Q(h^{t+1}, a; \bar{\theta}) = \sum_{i \in I} \max_{a_i} Q(h_i^{t+1}, a_i; \bar{\theta}_i).$$

Using this optimisation objective, all agents implicitly learn their individual utility functions. These functions are computationally tractable and naturally gives rise to policies with agent i choosing the greedy action with respect to its individual utility function for any given history. Note that no constraints are imposed on the individual utility functions and only the shared rewards are used during the optimisation.

Figure 9.9a illustrates the architecture of VDN. The algorithm follows the same pseudocode as IDQN (Algorithm 16) but optimises the loss given in Equation 9.41. It is worth noting that VDN can benefit from any of the optimisations which can be applied to IDQN mentioned in Section 9.3.1.

9.5.3 Monotonic Value Decomposition

While the linear decomposition assumed by VDN is natural and simple, it is not necessarily realistic. In many cases, the contribution of agents might be better represented by a non-linear relation, which VDN is unable to capture. Various approaches have been proposed to represent non-linear decompositions of rewards and value functions, with QMIX (Rashid et al. 2018) being a widely adopted approach. QMIX builds on the observation that the IGM property can be ensured if (strict) *monotonicity* of the centralised action-value function with respect to individual utilities holds, i.e. the derivative of the centralised action-value function with respect to agent utilities is positive:

$$\forall i \in I, \forall a \in A : \frac{\partial Q(s, a; \theta)}{\partial Q(h_i, a_i; \theta_i)} > 0 \quad (9.42)$$

Intuitively, this means that an increase in the utility of any agent i for its action a_i must lead to an increase in the decomposed centralised action-value function for joint actions containing a_i .

Similar to VDN, QMIX builds on top of IDQN and represents each agent's individual utility function as a deep Q-network. In order to be able to represent any monotonic decomposition of the centralised action-value function into

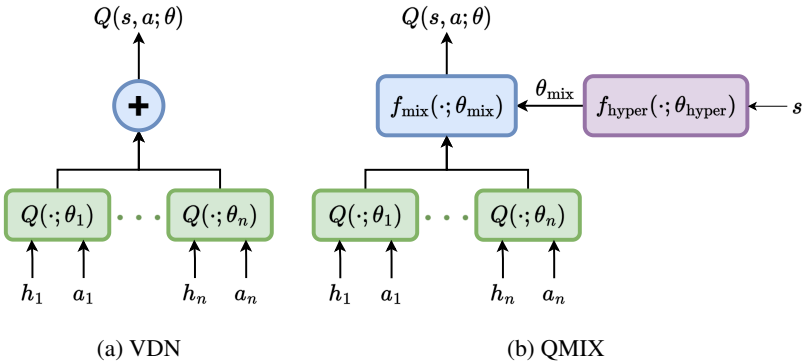


Figure 9.9: Network architecture of VDN and QMIX. The input to the individual utility networks are the observation histories and actions of the respective agents. For VDN, the centralised action-value function is approximated by the sum of the individual utilities. For QMIX, the centralised action-value function is approximated by a monotonic aggregation computed by the mixing network, which is parameterised by the output of a trained hypernetwork.

these individual utilities, QMIX defines a mixing network f_{mix} given by a feedforward neural network which combines individual utilities to approximate the centralised action-value function:

$$Q(s^t, a^t, \theta) = f_{\text{mix}}(Q(h_1^t, a_1^t; \theta_1), \dots, Q(h_n^t, a_n^t; \theta_n); \theta_{\text{mix}}) \quad (9.43)$$

This decomposition ensures the monotonicity property from Equation 9.42 if the mixing function is monotonic with respect to the utilities of all agents.

The monotonic decomposition of QMIX is a sufficient condition to ensure the IGM property. We can prove this similarly to the previous proof for a linear decomposition.

Proof. Let \hat{h} be a full history, $h_i = \sigma_i(\hat{h})$ be the observation history of agent i , and $s = s(\hat{h})$ denote the last state within \hat{h} . We will first prove that for any greedy joint action $a^* \in A^*(s; \theta)$, the individual actions in a^* of all agents are also greedy with respect to their individual utility functions. Then, we will show that any greedy individual actions of all agents together represent a greedy joint action with respect to $Q(s, a; \theta)$.

" \Rightarrow " Let $a^* = (a_1^*, \dots, a_n^*) \in A^*(s; \theta)$ be a greedy joint action with respect to $Q(s, \cdot; \theta)$. To prove that the individual actions of all agents a_1^*, \dots, a_n^* are also greedy with respect to agents' individual utility functions, i.e.

$$\forall i \in I : a_i^* \in A_i^*(h_i; \theta_i), \quad (9.44)$$

we need to show that for any agent i and any action $a_i \in A_i$, we have

$$\forall i \in I : Q(h_i, a_i^*; \theta_i) \geq Q(h_i, a_i; \theta_i). \quad (9.45)$$

We will prove this by contradiction. Assume that there exists an agent i and an action $a_i \in A_i$ such that

$$Q(h_i, a_i^*; \theta_i) < Q(h_i, a_i; \theta_i). \quad (9.46)$$

Given the monotonic decomposition of $Q(s, a; \theta)$, we have

$$Q(s, a^*; \theta) \quad (9.47)$$

$$= f_{\text{mix}}(Q(h_1, a_1^*; \theta_1), \dots, Q(h_i, a_i^*; \theta_i), \dots, Q(h_n, a_n^*; \theta_n); \theta_{\text{mix}}) \quad (9.48)$$

$$< f_{\text{mix}}(Q(h_1, a_1^*; \theta_1), \dots, Q(h_i, a_i; \theta_i), \dots, Q(h_n, a_n^*; \theta_n); \theta_{\text{mix}}) \quad (9.49)$$

$$= Q(s, \langle a_{-i}^*, a_i \rangle; \theta) \quad (9.50)$$

where the inequality follows from Equation 9.46 and the monotonicity of f_{mix} with respect to its inputs (Equation 9.42). This contradicts the assumption that a^* is a greedy joint action with respect to $Q(s, a; \theta)$. Therefore, Equation 9.46 cannot hold, and we have Equation 9.45, so the individual actions of all agents within the greedy joint action a^* are also greedy with respect to their individual utility functions.

" \Leftarrow " Let $a_1^* \in A_1^*(h_1; \theta_1), \dots, a_n^* \in A_n^*(h_n; \theta_n)$ be greedy individual actions of all agents with respect to their individual utility functions.

Let $a^* = (a_1^*, \dots, a_n^*)$ be the joint action composed of such greedy individual actions of all agents. To prove that a^* is a greedy joint action with respect to $Q(s, a; \theta)$, we need to show that for any joint action $a' \in A(s; \theta)$, we have

$$Q(s, a^*; \theta) \geq Q(s, a'; \theta). \quad (9.51)$$

Given the monotonic decomposition of $Q(s, a; \theta)$, we have

$$Q(s, a^*; \theta) = f_{\text{mix}}(Q(h_1, a_1^*; \theta_1), \dots, Q(h_n, a_n^*; \theta_n); \theta_{\text{mix}}) \quad (9.52)$$

$$\geq f_{\text{mix}}(Q(h_1, a'_1; \theta_1), \dots, Q(h_n, a'_n; \theta_n); \theta_{\text{mix}}) \quad (9.53)$$

$$= Q(s, a'; \theta) \quad (9.54)$$

where $a' = (a'_1, \dots, a'_n)$ is any joint action composed of arbitrary individual actions $a'_1 \in A_1, \dots, a'_n \in A_n$. The inequality follows from the monotonicity of f_{mix} with respect to its inputs (Equation 9.42) and the fact that a_1^*, \dots, a_n^* are greedy individual actions of all agents with respect to their individual utility functions. Therefore, Equation 9.51 holds, so a^* is a greedy joint action with respect to $Q(s, a; \theta)$.

This concludes the proof.⁶⁵ \square

In practise, the monotonicity assumption is satisfied if the mixing network f_{mix} is a network with only positive weights for the utility inputs. Note, the same constraint does not need to be imposed on the bias vectors in θ_{mix} . The parameters θ_{mix} of the mixing function are obtained through a separate hypernetwork f_{hyper} parameterised by θ_{hyper} which receives the full state s as its input and outputs the parameters θ_{mix} of the mixing network (hence the name "hyper"). To ensure positive weights, the hypernetwork f_{hyper} applies an absolute value function as activation function to the outputs corresponding to the weight matrix of the mixing network f_{mix} and, thus, ensures monotonicity⁶⁶. Whenever $Q(s, a; \theta)$ is needed for optimisation, the individual utilities $Q(h_1, a_1; \theta_1), \dots, Q(h_n, a_n; \theta_n)$ are computed and the mixing network parameters θ_{mix} are obtained by feeding the state into the hypernetwork. The utilities are then aggregated to $Q(s, a; \theta)$ using the mixing network with the parameters received by the hypernetwork.

The entire architecture of QMIX is illustrated in Figure 9.9b. During optimisation, all parameters θ of the decomposed centralised action-value function, including the parameters of individual utility networks $\theta_1, \dots, \theta_n$ and parameters

65. We note that in their original paper, Rashid et al. (2018) do not assume strict monotonicity of the mixing network f_{mix} with respect to its inputs. Instead, they assume that f_{mix} is monotonic with respect to its inputs, i.e.

$$\forall i \in I, \forall a \in A : \frac{\partial Q(s, a; \theta)}{\partial Q(h_i, a_i; \theta_i)} \geq 0 \quad (9.55)$$

In this case, the proof of the implication " \Rightarrow " that the joint action composed of greedy individual actions with respect to all agents' utility functions is a greedy joint action with respect to $Q(s, a; \theta)$ still holds. However, the proof of the implication " \Leftarrow " does not. To see why the strict monotonicity assumption is necessary, consider the following counterexample for two agents with actions $A_1 = \{a_{1,1}, a_{1,2}\}$ and $A_2 = \{a_{2,1}, a_{2,2}\}$, respectively. Let $a_{1,1}$ and $a_{2,1}$ be the greedy actions with respect to the agents' individual utility functions

$$Q(h_1, a_{1,1}; \theta_1) > Q(h_1, a_{1,2}; \theta_1) \quad \text{and} \quad Q(h_2, a_{2,1}; \theta_2) > Q(h_2, a_{2,2}; \theta_2), \quad (9.56)$$

and let the decomposed centralised action-value function be

$$Q(s, (a_1, a_2); \theta) = f_{\text{mix}}(Q(h_1, a_1; \theta_1), Q(h_2, a_2; \theta_2); \theta_{\text{mix}}) = Q(h_1, a_1; \theta_1) \quad (9.57)$$

for some actions $a_1 \in A_1$ and $a_2 \in A_2$, meaning that the mixing function only considers the first agent's utility and ignores the second agent's utility. We can see that

$$\frac{\partial Q(s, a; \theta)}{\partial Q(h_1, a_1; \theta_1)} = 1 \geq 0 \quad \text{and} \quad \frac{\partial Q(s, a; \theta)}{\partial Q(h_2, a_2; \theta_2)} = 0 \geq 0 \quad (9.58)$$

so the monotonicity assumption (Equation 9.55) holds. However, the joint action $a = (a_{1,1}, a_{2,2})$ is a greedy joint action with respect to $Q(s, a; \theta)$, since $a_{1,1}$ maximises the individual utility of agent 1 and the mixing function is maximised whenever the individual utility of agent 1 is maximised, but $a_{2,2}$ is not a greedy action with respect to the individual utility of agent 2. Therefore, the implication " \Leftarrow " does not hold without the strict monotonicity assumption (Equation 9.42).

66. We note that the absolute value function could allow zero weights which violate the necessary strict monotonicity assumption. However, for the strict monotonicity to be violated, all weights corresponding to a single agent's utility would need to be zero which does not occur in practise.

of the hypernetwork θ_{hyper} , are jointly optimised by minimising the value loss given by

$$\mathcal{L}(\theta) = \frac{1}{B} \sum_{(s^t, a^t, r^t, s^{t+1}) \in \mathcal{B}} \left(r^t + \gamma \max_a Q(s^{t+1}, a; \bar{\theta}) - Q(s^t, a^t; \theta) \right)^2 \quad (9.59)$$

over a batch \mathcal{B} sampled from the replay buffer \mathcal{D} with the centralised value function and its target network being given by Equation 9.43 with parameters θ and $\bar{\theta}$, respectively. The parameters of the mixing network are not optimised by gradient-based optimisation but instead are always obtained as an output of the optimised hypernetwork. The training in QMIX follows the same pseudocode as for IDQN (Algorithm 16), but additionally initialises the mixing network and hypernetwork and optimises the value loss given in Equation 9.59. It is also worth noting that the replay buffer in QMIX stores the full state s^t as a superset of the individual observation histories h_1^t, \dots, h_n^t of each agent used in VDN and IDQN. The full state is needed in QMIX to compute the monotonic mixing, because the hypernetwork function f_{hyper} is conditioned on the full state of the environment.

Furthermore, it is straightforward to see that any linear decomposition of the centralised action-value function as seen in Equation 9.28 also upholds the monotonicity property, but there exist monotonic decompositions of the centralised action-value function which are not linear. A simple example is the following linear decomposition

$$Q(s, a; \theta) = \sum_{i \in I} \alpha_i Q(h_i, a_i; \theta_i) \quad (9.60)$$

where $\alpha_i \geq 0$ are positive weights. The weights can be interpreted as the relative importance of each agent's contribution to the centralised action-value function. Any such weighting constitutes a monotonic decomposition, but VDN is only able to represent equal weighting for all agents with $\alpha_i = 1$ for $i \in I$. This example illustrates that the monotonicity assumed by QMIX subsumes the linearity assumption of VDN, so the set of centralised action-value functions which can be represented with QMIX is a superset of the centralised action-value functions which can be decomposed with VDN. In Section 9.5.4, we will see concrete examples of games where QMIX is able to decompose the centralised action-value function but VDN is unable to accurately represent the value function.

QMIX has been shown to outperform VDN and many other value-based MARL algorithms in a diverse set of common-reward environments (Rashid et al. 2018; Papoudakis et al. 2021). However, it is worth noting that the original implementation of QMIX (Rashid et al. 2018) includes several implementation details, some of which may significantly contribute to its performance. Among

<table border="1"> <tr><td></td><td>A</td><td>B</td></tr> <tr><td>A</td><td>1</td><td>5</td></tr> <tr><td>B</td><td>5</td><td>9</td></tr> </table>		A	B	A	1	5	B	5	9	<table border="1"> <tr><td></td><td>A</td><td>B</td></tr> <tr><td>A</td><td>0</td><td>0</td></tr> <tr><td>B</td><td>0</td><td>10</td></tr> </table>		A	B	A	0	0	B	0	10	<table border="1"> <tr><td></td><td>$Q_2(A)$</td><td>$Q_2(B)$</td></tr> <tr><td>$Q_1(A)$</td><td>$Q(A, A)$</td><td>$Q(A, B)$</td></tr> <tr><td>$Q_1(B)$</td><td>$Q(B, A)$</td><td>$Q(B, B)$</td></tr> </table>		$Q_2(A)$	$Q_2(B)$	$Q_1(A)$	$Q(A, A)$	$Q(A, B)$	$Q_1(B)$	$Q(B, A)$	$Q(B, B)$
	A	B																											
A	1	5																											
B	5	9																											
	A	B																											
A	0	0																											
B	0	10																											
	$Q_2(A)$	$Q_2(B)$																											
$Q_1(A)$	$Q(A, A)$	$Q(A, B)$																											
$Q_1(B)$	$Q(B, A)$	$Q(B, B)$																											
(a) Linear game	(b) Monotonic game	(c) Decomposition format																											

Figure 9.10: Common-reward matrix games and decomposition format. The centralised action-value function of game (a) can be represented using a linear decomposition, but the game (b) requires a more complex decomposition. (c) shows the decomposition format used in this section.

others, individual agent utility networks are shared across agents, i.e. $\theta_i = \theta_j$ for all $i, j \in I$, and these utility networks receive an additional onehot-encoded agent ID to allow for different utility functions across agents. We will discuss such parameter sharing with its benefits and disadvantages for deep MARL in more detail in Section 9.6. Also, agent utility networks are modelled as recurrent neural networks, and the observation of agent i additionally includes its last action to allow the utility function to consider the concrete action applied while stochastic exploration policies are followed. Lastly, an episodic replay buffer is used to store and sample batches of entire episodes to update all networks after the completion of each episode. All these details are not specific to QMIX and could equally be applied in other deep MARL algorithms.

9.5.4 Value Decomposition in Practice

To better understand value decomposition algorithms in practice, and to what extent they are able to represent the centralised action-value function, we will analyse VDN and QMIX in two simple 2×2 matrix games, shown in Figure 9.10. The first game can be decomposed using a linear aggregation of utilities, and the second game requires a more complex monotonic decomposition. To compactly represent the learned individual utility functions and the centralised action-value function for any of these matrix games, we present tables as shown in Figure 9.10c and in the following denote the individual utility function of agent i by Q_i instead of $Q(\cdot; \theta_i)$.

The first matrix game (Figure 9.10a) is linearly decomposable, i.e. the centralised action-value function can be represented by the sum of individual utility functions. To see this, we can write the centralised action-value function as

$$Q(a_1, a_2) = Q_1(a_1) + Q_2(a_2) \quad (9.61)$$

	0.12	4.12
0.88	1.00	5.00
4.88	5.00	9.00

(a) VDN – linear game

	-0.21	0.68
0.19	1.00	5.00
0.96	5.00	9.00

(b) QMIX – linear game

Figure 9.11: Visualisation of the learned value decomposition of (a) VDN and (b) QMIX in the linearly decomposable common-reward matrix game (Figure 9.10a).

and assign concrete values to the individual utility functions:

$$Q_1(A) = 1, \quad Q_1(B) = 5, \quad Q_2(A) = 0, \quad Q_2(B) = 4 \quad (9.62)$$

It is easy to verify that these utilities indeed lead to the desired centralised action-value function, but many similar assignments are possible. Figure 9.11 shows the individual utility functions and the centralised action-value functions learned by VDN and QMIX in the linear game. As expected, both VDN and QMIX are able to learn an accurate centralised action-value function⁶⁷. The learned individual utility functions of VDN are different from the ones given in Equation 9.62, but also accurately represent the centralised action-value function. However, these results also emphasise the fact that these algorithms are optimised to learn accurate centralised action-value functions, which may result in hard-to-interpret individual action-value functions. In particular for QMIX, which aggregates individual utilities through its monotonic mixing network, the individual utilities are difficult to interpret besides their ability to represent the centralised action-value function and policies of both agents.

The second matrix game (Figure 9.10b) is monotonic but not linear, i.e. the centralised action-value function can be represented by a monotonic decomposition of the individual action-value functions. The learned individual utilities and centralised action-value function for VDN and QMIX in the monotonic game are shown in Figure 9.12. As expected, VDN is unable to learn an accurate centralised action-value function due to its inability to represent non-linear value functions. In contrast, using its monotonic mixing function, QMIX learns the

67. For all experiments in this section, we train a feedforward neural network with two layers of 64 hidden units and ReLU activation function for each agent’s utility function in both VDN and QMIX. All networks are optimised with the Adam optimiser using a learning rate of $3e^{-4}$ and batches of 128 experiences sampled from the replay buffer. Target networks are updated every 200 time steps. To ensure sufficient exploration of all joint actions, agents follow a uniform policy throughout training. For QMIX, the hypernetwork is represented by a two-layer feedforward neural network with 32 hidden units each and the mixing network is represented by a single-layer feedforward neural network with 64 units. Both algorithms were trained until convergence (at most 20,000 time steps).

	-1.45	3.45
-0.94	-2.43	2.51
4.08	2.60	7.53

(a) VDN – monotonic game

	-4.91	0.82
-4.66	0.00	0.00
1.81	0.00	10.00

(b) QMIX – monotonic game

Figure 9.12: Visualisation of the learned value decomposition of (a) VDN and (b) QMIX in the monotonically decomposable common-reward matrix game (Figure 9.10b).

optimal centralised action-value function. To further illustrate the monotonic aggregation of individual utilities of both agents through the mixing network f_{mix} in QMIX, we visualise the mixing function over the individual agents' utilities in Figure 9.13. Due to the monotonicity constraint, the estimated joint action values increase with increasing individual utilities. The optimal joint action has a value of +10, which is significantly larger than the value of all the other joint actions, which is 0. QMIX's monotonic aggregation is able to accurately represent all these values by learning a mixing function with a steep incline from the value estimates of suboptimal actions to the optimal joint action value, as seen by the sharp change in shading in the top-right corner of the visualisation.

However, it is worth noting that despite inaccurate value estimates, VDN learns the optimal policy of (B,B) by greedily following its individual value function. We often evaluate MARL algorithms based on the evaluation returns (or rewards in non-repeated matrix games) and here VDN and QMIX would both obtain the optimal reward of +10 by greedily following the learned individual utility functions. To see that the inaccurate value estimates of VDN can be problematic, we evaluate both VDN and QMIX in a two-step common-reward stochastic game shown in Figure 9.14. This game has three states. In the initial state, the agents receive 0 reward for any action but the action of agent 1 determines the next and final state. With action $a_1 = A$, the agents will play the previously introduced linearly decomposable matrix game. With action $a_1 = B$, the agents will play the monotonically decomposable matrix game. The optimal policy in this two-step game is for agent 1 to use action B in the initial state and for both agents to use action B in the monotonic game for a reward of +10. QMIX accurately learns the centralised action-value function in both games and is able to learn this optimal policy (Figure 9.15b). However, VDN is only able to learn accurate value estimates in the linearly decomposable game and underestimates the value of the optimal policy in the other game (Figure 9.15a). This underestimation leads to VDN preferring to play the linearly decomposable

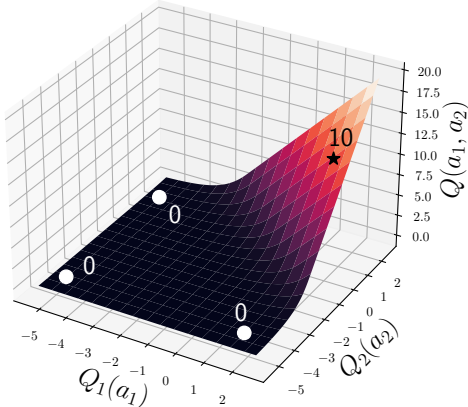


Figure 9.13: Visualisation of the QMIX mixing function f_{mix} in the monotonic matrix game. The mixing function is shown after convergence over the individual utilities of both agents, with shading corresponding to the estimated centralised action-value function. The mixed utilities of all four joint actions are highlighted together with their centralised action-value estimate, with the optimal joint action shown as a star.

game and converging to the suboptimal policy of playing $a_1 = A$ in the initial state and then playing the policy (B,B) for a reward of +9.⁶⁸

Lastly, we evaluate both QMIX and VDN in the climbing matrix game shown in Figure 9.16a. As we have seen before (Section 9.4.5), solving this game is challenging due to the coordination required to obtain the optimal rewards of +11, with deviations by any individual agent leading to significantly lower rewards. By inspecting the decomposition learned by VDN (Figure 9.16b) and QMIX (Figure 9.16c), we see that the inaccurate decomposition results in both algorithms converging to suboptimal policies. VDN converges to (C,C) with a reward of +5 and QMIX converges to (C, B) with a reward of +6. This illustrates that both VDN and QMIX are unable to learn accurate action-value estimates and recover the optimal policy for challenging matrix games.

68. The value and utility functions learned by VDN and QMIX at both steps at $t = 1$ in the two-step game are different from the ones seen in Figures 9.11 and 9.12 due to different initialisation of the networks and stochasticity introduced by the ϵ -greedy exploration policy. We train both algorithms with $\gamma = 1.0$.

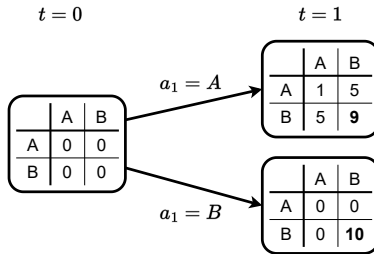


Figure 9.14: Two-step common-reward stochastic game with three states. The transition function across the three states and the reward function within each state is shown here. In the initial state the agents receive no rewards for any action and agent 1 decides which matrix game to play in the next time step: either the linearly decomposable game, as represented by the state in the top right, or the monotonically decomposable game, as represented by the state in the bottom right.

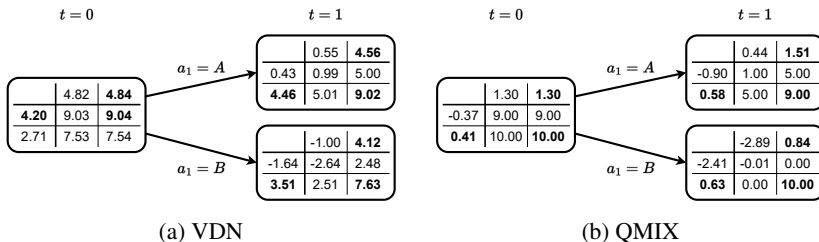


Figure 9.15: Learned value decomposition of (a) VDN and (b) QMIX in the two-step common-reward stochastic game. Both algorithms were trained to estimate undiscounted returns ($\gamma = 1$).

9.5.5 Beyond Monotonic Value Decomposition

As we see in the climbing game, even the monotonicity constraint of QMIX on the value decomposition might be too restrictive to be able to represent an accurate centralised action-value function. As we will see below, the monotonicity constraint is a sufficient but not necessary condition to ensure the IGM property (Equation 9.23). In that sense, a less restrictive constraint can be formalised to factorise the centralised action-value function into individual utility functions.

Son et al. (2019) formulate the following conditions sufficient to ensure the IGM property of the value decomposition, i.e. the utility functions factorise the

	A	B	C
A	11	-30	0
B	-30	7	0
C	0	6	5

(a) Climbing game

	-4.56	-4.15	3.28		-16.60	-0.24	-4.68
-4.28	-8.84	-8.43	-1.00	-7.44	-11.16	-11.16	-11.16
-6.10	-10.66	-10.25	-2.82	7.65	-11.15	2.34	-1.37
5.31	0.75	1.16	8.59	11.27	-4.95	8.72	5.01

(b) VDN – climbing game

	-16.60	-0.24	-4.68
-16.60	-11.16	-11.16	-11.16
7.65	-11.15	2.34	-1.37
11.27	-4.95	8.72	5.01

(c) QMIX – climbing game

Figure 9.16: (a) Reward table of the climbing matrix game with the learned value decomposition of (b) VDN and (c) QMIX.

centralised action-value function if

$$\sum_{i \in I} Q(h_i, a_i; \theta_i) - Q(s, a; \theta_q) + V(s; \theta_v) = \begin{cases} 0 & \text{if } a = a^* \\ \geq 0 & \text{if } a \neq a^* \end{cases} \quad (9.63)$$

where $a^* = (a_1^*, \dots, a_n^*)$ is the greedy joint action with $a_i^* = \arg \max_{a_i} Q(h_i, a_i; \theta_i)$. $V(s; \theta_v)$ denotes a state-utility function defined as follows

$$V(s; \theta_v) = \max_a Q(s, a; \theta_q) - \sum_{i \in I} Q(h_i, a_i^*; \theta_i) \quad (9.64)$$

and a centralised action-value function Q over the state and joint action is parameterised by θ_q .

It can be further shown that these conditions are also necessary to ensure the IGM property under an affine transformation g of the individual utilities, where

$$g(Q(h_i, a_i; \theta_i)) = \alpha_i Q(h_i, a_i; \theta_i) + \beta_i \quad (9.65)$$

with $\alpha_i \in \mathbb{R}_+$ and $\beta_i \in \mathbb{R}$ for all $i \in I$. Given such a necessary and sufficient condition, we know that for any centralised action-value function which can be factorised into individual utility functions, there exists a decomposition which satisfies the above conditions. Likewise, if a decomposition of this form is found, we know that it satisfies the IGM property and, therefore, factorises the centralised action-value function.

Based on these conditions, the QTRAN value decomposition algorithm is defined (Son et al. 2019). From Equation 9.63, we can identify three components required for this condition: (1) individual utility functions $\{Q(\cdot; \theta_i)\}_{i \in I}$, (2) a (non-decomposed) centralised action-value function $Q(s, a; \theta_q)$, and (3) a

state-utility function $V(s; \theta_v)$. QTRAN optimises neural networks for each of these components. For each agent, an individual utility function is trained, and QTRAN trains a single network to approximate the global state-utility function V and centralised action-value function $Q(s, a; \theta_q)$, respectively. Therefore, in contrast to the previously discussed value decomposition algorithms, QTRAN directly optimises a centralised action-value function for the purpose of optimising the individual utility functions used for action selection. As previously discussed in Section 9.4.3, naively training a centralised action-value function with the state as input and one output for the value estimate of each joint action is infeasible for larger numbers of agents. Therefore, the network for the centralised action-value function receives the state and joint action as input and computes a single scalar output for the corresponding value estimate. To train the centralised action-value function, the following TD-error is minimised over a batch \mathcal{B} sampled from a replay buffer \mathcal{D}

$$\mathcal{L}_{\text{td}}(\theta_q) = \frac{1}{B} \sum_{(s^t, a^t, r^t, s^{t+1}) \in \mathcal{B}} (r^t + \gamma Q(s^{t+1}, a^{*t+1}; \bar{\theta}_q) - Q(s^t, a^t; \theta_q))^2 \quad (9.66)$$

where $\bar{\theta}_q$ denotes the parameters of an architecturally identical target network, and $a^{*t} = (\arg \max_{a_i} Q(h_i^t, a_i; \theta_i))_{i \in I}$ is the greedy joint action at time step t . To train the individual utility networks of all agents and ensure that the conditions of Equation 9.63 are satisfied, QTRAN computes soft regularisation terms in the overall loss function. By minimising the first regularisation term given by

$$\mathcal{L}_{\text{opt}}(\{\theta_i\}_{i \in I}, \theta_v) = \frac{1}{B} \sum_{(s^t, a^t, r^t, s^{t+1}) \in \mathcal{B}} \left(\sum_{i \in I} Q(h_i^t, a_i^{*t}; \theta_i) - Q(s^t, a^{*t}; \theta_q) + V(s^t; \theta_v) \right)^2 \quad (9.67)$$

QTRAN optimises for the property stated in Equation 9.63 for the greedy joint action. The second regularisation term is computed as follows

$$m = \sum_{i \in I} Q(h_i^t, a_i^t; \theta_i) - Q(s^t, a^t; \theta_q) + V(s^t; \theta_v) \quad (9.68)$$

$$\mathcal{L}_{\text{nopt}}(\{\theta_i\}_{i \in I}, \theta_v) = \frac{1}{B} \sum_{(s^t, a^t, r^t, s^{t+1}) \in \mathcal{B}} \min(0, m)^2 \quad (9.69)$$

and optimises for the property stated for non-greedy joint actions a .⁶⁹ We note that the optimisation of QTRAN does not directly enforce the formulated properties Equations 9.63 and 9.64 but minimises the additional loss terms

69. The QTRAN algorithm presented in this section is the QTRAN-base algorithm proposed by Son et al. (2019). In their work, they also propose an alternative algorithm QTRAN-alt which uses a counterfactual centralised action-value function, similar to COMA (Section 9.4.4), to compute an alternative condition for non-greedy actions.

Equations 9.67 and 9.69, which reach their minimum when the properties from Equation 9.63 are satisfied. Therefore, these properties are only satisfied asymptotically and not throughout the entire training process.

To get a sense for the centralised action-value functions QTRAN is able to represent, we train the algorithm in the linear game, the monotonic game, and the climbing game seen in Section 9.5.4. We can see that the unconstrained centralised action value function $Q(s, a; \theta_q)$ of QTRAN almost exactly converges to the true reward table for all three games (Figures 9.17a, 9.18a and 9.19a). The learned linear decomposition does not converge to the correct centralised action values (Figures 9.17b, 9.18b and 9.19b), but together with the learned state utilities is converging to values which satisfy the formulated constraints in Equation 9.63 for the linear and climbing game. For the monotonically decomposable matrix game, the constraint for the optimal action $a^* = (B, B)$ is not satisfied exactly with

$$\sum_{i \in I} Q(h_i, a_i^*; \theta_i) - Q(s, a^*; \theta_q) + V(s; \theta_v) = 7.58 - 10.00 + 2.45 = 0.03 \neq 0 \quad (9.70)$$

but is still close to zero. Given the constraints of Equation 9.63 are a sufficient and necessary condition to satisfy the IGM property, we can see that QTRAN is able to learn a decomposition of the centralised action-value function which satisfies the IGM property for the linear and climbing game, and is close to satisfying it for the monotonic game. For all three games, the learned decomposition of QTRAN gives rise to the optimal policy, indicating its ability to express more complex decompositions than VDN and QMIX which were unable to learn the optimal policy in the climbing game. However, it is worth noting that training the non-decomposed centralised action-value function becomes challenging in tasks with large number of agents or actions, since the joint action space grows exponentially to the number of agents. In contrast, the number of parameters of the decomposed centralised action-value function of QTRAN grows linearly with the number of agents and actions, making it more scalable to large multi-agent systems.

9.6 Environments with Homogeneous Agents

As the number of agents increases, the parameter space of multi-agent deep RL methods can increase significantly. For example, IDQN has an action value network containing a large number of parameters for each agent in the environment, which we have denoted with θ_i so far. Having more agents in IDQN requires more sets of such parameters, each defining the weights of an

	A	B			
A	0.99	5.00	2.01	4.13	$V(s; \theta_v) = 1.10$
B	5.00	9.00	1.82	3.83	5.95
(a) $Q(s, a; \theta_q)$	(b) Linear decomposition	(c) State utility			

Figure 9.17: Visualisation of the learned (a) centralised action-value function, (b) value decomposition, and (c) state utility of QTRAN in the linearly decomposable common-reward matrix game (Figure 9.10a).

	A	B			
A	0.01	0.00	0.75	3.83	$V(s; \theta_v) = 2.45$
B	0.00	10.00	0.72	1.47	4.55
(a) $Q(s, a; \theta_q)$	(b) Linear decomposition	(c) State utility			

Figure 9.18: Visualisation of the learned (a) centralised action-value function, (b) value decomposition, and (c) state utility of QTRAN in the monotonically decomposable common-reward matrix game (Figure 9.10b).

	A	B	C			
A	11.01	-30.25	0.00	4.98	4.29	2.63
B	-30.27	5.38	5.73	4.67	9.65	8.96
C	0.00	6.18	4.31	3.98	8.96	8.27
				2.45	7.43	6.74
(a) $Q(s, a; \theta_q)$	(b) Linear decomposition	(c) State utility				

Figure 9.19: Visualisation of the learned (a) centralised action-value function, (b) value decomposition, and (c) state utility of QTRAN in the climbing game (Figure 9.16a).

action value neural network. Updating and training multiple neural networks can be a slow process and require a lot of computational power. In this section, we discuss methods to improve the scalability and sample efficiency of MARL algorithms in environments where agents are *homogeneous*.

A game where the agents are homogeneous intuitively means that all agents share the same or similar capabilities and characteristics, such as the same set of actions, observations, and rewards. Homogeneous agents can be thought of as identical or nearly identical copies of each other, and they interact with the environment in similar ways. Intuitively, this allows agents to interchange their policies while still preserving the overall result in solving a task, regardless

of which agent is executing which policy. The level-based foraging example (Section 11.3.1) could be such a game: the agents are all randomly assigned a level in the beginning of an episode, they move in the environment using the same set of actions, and receive similar rewards when successfully gathering the items. We define such a game as an environment with *weakly homogeneous agents*.

Definition 14 (Environment with weakly homogeneous agents) *Formally, an environment has weakly homogeneous agents if for any joint policy $\pi = (\pi_1, \pi_2, \dots, \pi_n)$ and permutation between agents $\sigma : I \mapsto I$, the following holds:*

$$U_i(\pi) = U_{\sigma(i)}((\pi_{\sigma(1)}, \pi_{\sigma(2)}, \dots, \pi_{\sigma(n)})), \forall i \in I \quad (9.71)$$

At a first glance, one could assume that in such an environment, the optimal policy of the agents would be to behave identically. This is not necessarily true as we will discuss in the rest of the section, especially given the numerous solution concepts in MARL (Chapter 4). However, to capture environments in which all agents act identically under the optimal policy, we also define an environment with *strongly homogeneous agents*.

Definition 15 (Environment with strongly homogeneous agents) *An environment has strongly homogeneous agents if the optimal joint policy consists of identical individual policies, formally:*

- *The environment has weakly homogeneous agents.*
- *The optimal joint policy $\pi^* = (\pi_1, \pi_2, \dots, \pi_n)$ consists of identical policies $\pi_1 = \pi_2 = \dots = \pi_n$.*

Figure 9.20 shows two environments with weakly homogeneous (left) and strongly homogeneous (right) agents. In both examples, the agents start at a random location and must reach a location to receive a positive reward and end the game. The agents can only observe their own position relative to the landmark(s) and move in one of the four cardinal directions.

In the environment shown in Figure 9.20a, there are two landmarks that the agents must reach. The agents are weakly homogeneous in this example because their policies can be switched without affecting the outcome of the game; if they have learned to move to opposite landmarks, they will still move to them even if their policies are switched. However, the optimal policy (e.g. Agent 1 moving to the left landmark and Agent 2 moving to the right landmark) requires the agents to behave differently by moving to different landmarks, and as such, the environment does not have strongly homogeneous agents.

In contrast, the environment shown in Figure 9.20b has a single landmark for both agents to move to. The optimal policy in this environment requires both

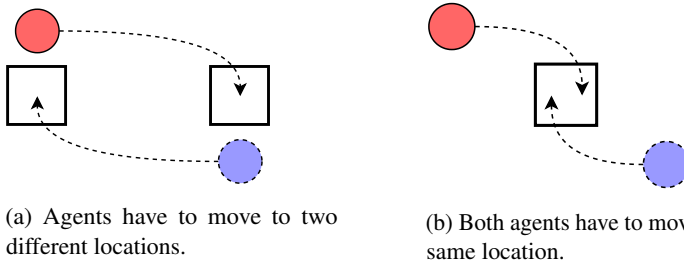


Figure 9.20: Illustrations of environments with weakly (left) and strongly (right) homogeneous agents. Agents (circles) have to move to certain locations (squares) to receive a positive reward. In the beginning of the episode, the agents begin in a random location. For simplicity, agents can only observe the squares, they cannot collide, and each agent chooses to move in one of the four cardinal directions. The agents receive a common reward of 1 only when they both arrive at their goal.

agents to move to the same landmark as soon as possible, making the agents *strongly homogeneous*.

In the next two subsections, we will further discuss environments with weakly and strongly homogeneous agents, and how the properties of such environments can be exploited for improved sample and computational efficiency.

9.6.1 Parameter Sharing

Sharing parameters across agents can offer immediate improvements on an algorithm's sample efficiency in environments with strongly homogeneous agents. Parameter sharing consists of agents using the same set of parameter values in their neural networks, i.e.

$$\begin{aligned}\theta_{\text{shared}} &\equiv \theta_1 \equiv \theta_2 \equiv \dots \equiv \theta_n \text{ and/or} \\ \phi_{\text{shared}} &\equiv \phi_1 \equiv \phi_2 \equiv \dots \equiv \phi_n.\end{aligned}\tag{9.72}$$

Equation 9.72, which constrains the policy space to identical policies, aligns with the definition of strongly homogeneous agents introduced earlier (Definition 15). Any experience generated by those agents, in the form of observations, actions and rewards, is then used to update shared parameters simultaneously. Sharing parameters among agents has two main benefits. Firstly, it keeps the number of parameters constant, regardless of the number of agents, whereas not sharing would result in a linear increase in the number of parameters with the number of agents. Secondly, the shared parameters are updated using the

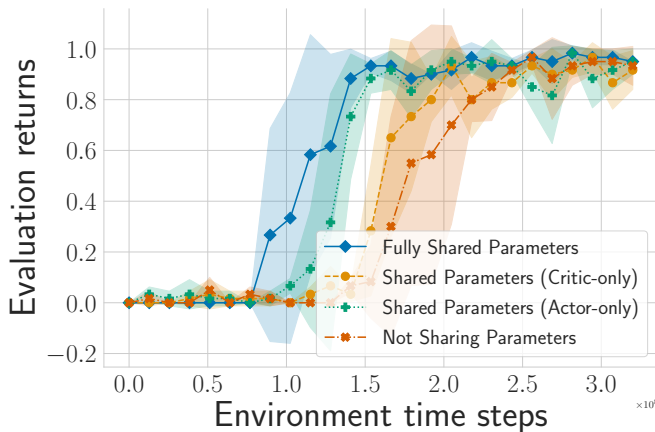


Figure 9.21: Learning curves for the Independent Actor-Critic algorithm on the level-based foraging environment.

experiences generated by all agents, resulting in a more diverse and larger set of trajectories for training.

The caveat is that strongly homogeneous agents is a strong assumption, and one that might not even be known beforehand. If agents share parameters, the policies of the agents are identical. An environment with weakly homogeneous agents will not see the same benefit from parameter sharing. An example was already shown in Figure 9.20a, in which the agents are weakly homogeneous but the policies are different and cannot be learned if the constraint of Equation 9.72 is enforced. So, when approaching a problem one should consider the properties of the environment and decide whether techniques such as parameter sharing could be useful.

Theoretically, we might be able to train diverse policies and still retain the benefits of shared parameters in environments with *weakly* homogeneous agents. To achieve that, the observations of the agents can include their index i to create a new observation \bar{o}_i^t . Since each agent will always receive an observation which contains their own index, they will theoretically be able to develop distinct behaviours. But in practice, learning distinct policies by relying on the index is not the case, as the representational capacity of neural networks might not be enough to represent several distinct strategies. The work of Christianos et al. (2021) discusses the limitations of parameter sharing and parameter sharing with an observation that contains the index.

To further illustrate the advantages of parameter sharing in an environment with strongly homogeneous agents, we have executed four variations of the

Independent A2C algorithm (Algorithm 18) in a level-based foraging environment. The four variations are i) using parameter sharing in both actor and critic ii) parameter sharing only on the critic iii) parameter sharing only on the actor iv) no parameter sharing. Figure 9.21 presents the average returns during training for the four variations in an level-based foraging environment with a grid-world of size 6×6 , two level-1 agents, and one level-2 item. In the figure, we observe that sharing parameters leads to the algorithm converging in less timesteps. However, it does not necessarily increase the final converged returns, as all algorithms eventually reach a similar policy in this example.

9.6.2 Experience Sharing

Parameter sharing offers computational advantages by retaining only a single set of policy (and/or value) network parameters. Learning only one set of parameters is also very beneficial in environments with strongly homogeneous agents since it constrains the policy search space to identical individual policies. A different approach would be to train a different set of parameters for each agent, but share the trajectories generated between the agents which relaxes the strongly homogeneous agent assumption and allows for different policies to be learned. For example, consider the IDQN algorithm, where each agent collects experience tuples and stores them in an experience replay. In an environment with weakly homogeneous agents, the experience replay could be shared across all agents. Now, an agent could sample and learn from a transition collected by a different agent.

In Algorithm 20 we show how DQN can be implemented with a shared replay. In the pseudocode shown, the experience replays of IDQN ($D_{1\dots n}$) are replaced with a single shared one D_{shared} . This simple change can improve learning in environments with weakly homogeneous agents as the variety of experiences found in the replay increases. Also, any successful policy by an agent that achieves higher rewards will further populate the replay with those experiences from which the other agents can also learn from.

Implementing DQN with shared experiences is relatively simple because DQN is an off-policy algorithm. Off-policy algorithms can make use of transitions collected by other agents. That is not the case for algorithms such as Independent Actor-Critic (Algorithm 18). *Shared Experience Actor-Critic (SEAC)* (Christianos, Schäfer, and Albrecht 2020) uses importance sampling to correct for transitions collected by other algorithms and apply the idea of experience sharing to on-policy settings.

In on-policy based algorithms, such as A2C or PPO, each agent generates one on-policy trajectory in every episode. The algorithms we have seen so

Algorithm 20 Deep Q-Networks with Shared Experience Replay

-
- 1: Initialise n value networks with random parameters $\theta_1 \dots \theta_n$
 - 2: Initialise a single replay D_{shared} buffer for all agents
 - 3: Collect environment observations $o_1^0 \dots o_n^0$
 - 4: **for** time step $t = 0 \dots$ **do**
 - 5: **for** agent $i = 1 \dots n$ **do**
 - 6: With probability ϵ : choose random action a_i^t
 - 7: Else: choose $a_i^t \in \arg \max_{a_i} Q(h_i^t, a_i; \theta_i)$
 - 8: Execute actions and collect observations o_i^{t+1} and rewards r_i^t .
 - 9: Store all transitions in replay buffer D_{shared}
 - 10: **for** agent $i = 1 \dots n$ **do**
 - 11: Sample random mini-batch of transitions from replay buffer D_{shared}
 - 12: Update parameters θ_i by minimising the loss from Equation 9.1
 - 13: In a set interval, update target network parameters $\bar{\theta}_i$ for each agent i
-

far, such as Independent A2C or A2C with centralised state-value critics, used the experience of each agent's own sampled trajectory to update the agent's networks with respect to their policy loss (e.g. Equation 9.3). The SEAC algorithm reuses trajectories of other agents while considering that they have been collected as off-policy data, i.e. the trajectories were generated by agents executing different policies than the one optimised. Correcting for off-policy samples can be achieved with importance sampling. The loss for such off-policy policy gradient optimisation from a behavioural policy π_β can be written as

$$\mathcal{L}(\phi) = -\frac{\pi(a^t | o^t; \phi)}{\pi_\beta(a^t | o^t)} (r^t + \gamma V(s^{t+1}; \theta) - V(s^t; \theta)) \log \pi(a^t | s^t; \phi),$$

in a similar fashion to Equation 8.40 in Section 8.2.6. In the independent A2C framework of Section 9.3.2, we can extend the policy loss to use the agent's own trajectories (denoted with i) along with the experience of other agents (denoted with k), shown below:

$$\begin{aligned} \mathcal{L}(\phi_i) = & - (r_i^t + \gamma V(s_i^{t+1}; \theta_i) - V(s_i^t; \theta_i)) \log \pi(a_i^t | s_i^t; \phi_i) \\ & - \lambda \sum_{k \neq i} \frac{\pi(a_k^t | o_k^t; \phi_i)}{\pi(a_k^t | o_k^t; \phi_k)} (r_k^t + \gamma V(s_k^{t+1}; \theta_i) - V(s_k^t; \theta_i)) \log \pi(a_k^t | s_k^t; \phi_i) \end{aligned} \quad (9.73)$$

Using this loss function, each agent is trained on both on-policy data while also using the off-policy data collected by all other agents at each training step. The

hyperparameter λ controls how much the experience of others is weighted, with $\lambda = 1$ indicating that the experience of others has the same weight to the agent and $\lambda = 0$ collapsing to no-sharing. The value function can also make use of experience sharing if desired. The final algorithm is identical to Independent A2C (Algorithm 18), but uses Equation 9.73 for learning the parameters.

Why would one consider using experience sharing over parameter sharing? Experience sharing is certainly more expensive in terms of computational power per environment step as it increases the size of the batch and has a number of neural network parameters that scale with the number of agents. However, it has been shown (Christianos, Schäfer, and Albrecht 2020) that initialising agent-specific neural networks leads to learning better policies: not making the assumption that an environment contains strongly homogeneous agents and not restricting policies to be identical leads to higher converged returns.

In contrast to parameter sharing, experience sharing does not assume that the optimal policy $\pi^* = (\pi_1, \pi_2, \dots, \pi_n)$ consists of identical policies $\pi_1 = \pi_2 = \dots = \pi_n$. In addition, the benefit of experience sharing over not sharing any experience or parameters is that it increases the sample efficiency of algorithms by using all the available trajectories for learning. But another less apparent benefit is that it ensures a uniform learning progression between agents. Agents learning from the experience of others can quickly pick up on policies that achieve higher returns, since those trajectories are treated similarly to expert data. In turn, when agents have a similar learning progression, they get more opportunities to explore actions that require coordination, resulting in better efficiency in data collection.

In the end, sharing experience or parameters rests on the choices of the MARL practitioner and is highly sensitive to the environment.

9.7 Policy Self-Play in Zero-Sum Games

In this section, we will turn our attention to zero-sum games with two agents and fully observable states and actions, in particular turn-taking board games such as chess, shogi, backgammon, and Go. Such games are characterised by three primary aspects, which in combination can make for a very challenging decision problem for an agent:

1. The game terminates after a finite number of moves, at which point the winning player receives a +1 reward and the losing player receives a -1 reward; or both players receive a 0 reward in the case of a draw outcome. In all other cases there is no reward signal, meaning the reward is always 0 in non-terminal states, until a terminal state is reached.

2. Reaching a terminal state (i.e. win/lose/draw) can require tens or even hundreds of moves in some games. Therefore, the agents may have to explore long sequences of actions before they receive any reward.
3. The agents can typically choose from a large number of actions, such as moving the many available pieces on the board. This leads to a large branching factor in the search space, which can be infeasible to fully explore and navigate with a limited compute budget.

A standard approach to tackle such games is to use heuristic search algorithms such as alpha-beta minimax search, which expands a search tree from each encountered state of the game to compute an optimal action for that state. Such algorithms rely heavily on specialised evaluation functions which are used to guide the search, as well as many other domain-specific adaptations (Levy and Newborn 1982; Campbell, Hoane Jr, and Hsu 2002). This makes such algorithms highly specialised for a specific game, and difficult to adapt to other games. Moreover, the heuristic design choices in such algorithms can limit their achievable performance, such as due to inaccuracies in heuristic evaluations of game states.

Monte Carlo Tree Search (MCTS) is a sampling-based method which, similar to alpha-beta search, expands a search tree from the game state, but does not require knowledge of specialised evaluation functions (however, if available, such knowledge can also be used in MCTS to further improve its performance). MCTS algorithms grow the search tree by generating a number of simulations from the game state, where each simulation is produced by sampling actions based on information contained in the current search tree. MCTS is essentially a reinforcement learning method, in that it can use the same action selection mechanisms and temporal-difference learning operators. However, while reinforcement learning algorithms aim to learn complete policies that choose optimal actions in each state, the focus in MCTS is to compute optimal actions for the *current* state of the game rather than complete policies.

Algorithms that use MCTS in combination with policy self-play and deep learning have achieved “super-human” performance in several games, including chess, shogi, and Go (Silver et al. 2016; Silver et al. 2017; Silver et al. 2018; Schrittwieser et al. 2020). These algorithms use a self-play approach (specifically, “policy self-play” as described in Section 5.5.1) whereby an agent’s policy is trained against itself. In this section, we will present one such algorithm called AlphaZero (Silver et al. 2018). The section begins by describing a general MCTS algorithm for Markov decision processes, based on which we will introduce policy self-play in MCTS for zero-sum turn-taking games with

fully observable states and actions. The AlphaZero algorithm uses this self-play MCTS algorithm in combination with deep learning to learn an effective evaluation function for the game.

9.7.1 Monte Carlo Tree Search

The pseudocode for a general MCTS algorithm in a Markov decision process is shown in Algorithm 21. In each state s^t , the algorithm performs k simulations⁷⁰ $\hat{s}^\tau, \hat{a}^\tau, \hat{r}^\tau, \hat{s}^{\tau+1}, \hat{a}^{\tau+1}, \hat{r}^{\tau+1}, \hat{s}^{\tau+2}, \hat{a}^{\tau+2}, \hat{r}^{\tau+2}, \dots$ (where k is a parameter of the MCTS algorithm) by sampling actions and growing the search tree. We use the notation $\hat{s}^\tau, \hat{a}^\tau$, and \hat{r}^τ to refer to the states, actions, and rewards in a simulation, respectively, starting with state $\hat{s}^\tau = s^t$ and time $\tau = t$. To simplify the pseudocode, we implicitly assume that \hat{s}^τ refers to both a state and a corresponding node in the search tree; hence, we do not introduce explicit notation to represent trees and nodes. The algorithm can either use the state transition function \mathcal{T} if this is known, or it can use a simulation model $\widehat{\mathcal{T}}$ as described in Section 3.5. Note that, rather than building a new tree in each state s^t , MCTS continually uses and grows the tree across all episodes and time steps.

Each node in the search tree carries certain statistics used by the algorithm. In particular, each node \hat{s}^τ contains a counter $N(\hat{s}^\tau, \hat{a})$ which counts the number of times that action \hat{a} was tried in state \hat{s}^τ , and an action value function $Q(\hat{s}^\tau, \hat{a})$ to estimate the value (i.e. expected return) of each action in the state. When a new node \hat{s}^τ is created and added to the tree, the function $InitialiseNode(\hat{s}^\tau)$ sets the counter and action value function to zero, i.e. $N(\hat{s}^\tau, \hat{a}) = 0$ and $Q(\hat{s}^\tau, \hat{a}) = 0$ for all $\hat{a} \in A$.

To generate a simulation, the function $ExploreAction(\hat{s}^\tau)$ returns an action to be tried in each visited state \hat{s}^τ in the simulation. A basic way to sample actions is ϵ -greedy action selection, such as used in Algorithm 3 (page 35). Another, often used action selection method in MCTS is to first try each action once, and then deterministically choose an action which has the highest *upper confidence bound* (UCB),⁷¹ formally

$$\hat{a}^\tau = \begin{cases} \hat{a} & \text{if } N(\hat{s}^\tau, \hat{a}) = 0, \text{ else} \\ \arg \max_{\hat{a} \in A} \left(Q(\hat{s}^\tau, \hat{a}) + \sqrt{\frac{2 \ln N(\hat{s}^\tau)}{N(\hat{s}^\tau, \hat{a})}} \right) \end{cases} \quad (9.74)$$

70. Other terms used to refer to simulations in MCTS include “rollouts” and “playouts”. These terms come from the original research focus of using MCTS in competitive board games. We prefer the neutral term “simulation” since MCTS can be applied in a broad range of decision problems.

71. This version of UCB was originally called “UCB1” in the work of Auer, Cesa-Bianchi, and Fischer (2002).

Algorithm 21 Monte Carlo Tree Search (MCTS) for MDPs

```

1: Repeat for every episode:
2: for  $t=0, 1, 2, 3, \dots$  do
3:   Observe current state  $s^t$ 
4:   for  $k$  simulations do
5:      $\tau \leftarrow t$ 
6:      $\hat{s}^\tau \leftarrow s^t$                                  $\triangleright$  Perform simulation
7:     while  $\hat{s}^\tau$  is non-terminal and  $\hat{s}^\tau$ -node exists in tree do
8:        $\hat{a}^\tau \leftarrow ExploreAction(\hat{s}^\tau)$ 
9:        $\hat{s}^{\tau+1} \sim \mathcal{T}(\cdot | \hat{s}^\tau, \hat{a}^\tau)$ 
10:       $\hat{r}^\tau \leftarrow \mathcal{R}(\hat{s}^\tau, \hat{a}^\tau, \hat{s}^{\tau+1})$ 
11:       $\tau \leftarrow \tau + 1$ 
12:    if  $\hat{s}^\tau$ -node does not exist in tree then
13:       $InitialiseNode(\hat{s}^\tau)$                        $\triangleright$  Expand tree
14:      while  $\tau > t$  do                           $\triangleright$  Backpropagate
15:         $\tau \leftarrow \tau - 1$ 
16:         $Update(Q, \hat{s}^\tau, \hat{a}^\tau)$ 
17:    Select action  $a^t$  for state  $s^t$ :
18:     $\pi^t \leftarrow BestAction(s^t)$ 
19:     $a^t \sim \pi^t$ 

```

where $N(\hat{s}) = \sum_{\hat{a}} N(\hat{s}, \hat{a})$ is the number of times state \hat{s} has been visited. (If multiple actions \hat{a} have $N(\hat{s}^\tau, \hat{a}) = 0$, then we can choose them in any order.) UCB assumes that the rewards \hat{r}^τ lie in the normalised range [0, 1] (Auer, Cesa-Bianchi, and Fischer 2002). UCB action selection tends to be more effective than ϵ -greedy action selection if only a relatively small number of actions can be tried in a state. Moreover, the finite-sample estimation errors of UCB are bounded when used in MCTS (Kocsis and Szepesvári 2006).

The simulation stops once a state \hat{s}^l has been reached which is a leaf node in the search tree, meaning the state \hat{s}^l has never been visited and no corresponding node exists in the search tree. The search tree is then expanded by using $InitialiseNode(\hat{s}^l)$ to initialise a new node corresponding to \hat{s}^l . $InitialiseNode(\hat{s}^l)$ can use an evaluation function $f(\hat{s}^l)$ to get an initial estimate of the value of the state \hat{s}^l . Such a value estimate could be obtained in different ways. If domain knowledge is available, then a heuristic function f could be manually created to compute value estimates based on the domain knowledge. For instance, sophisticated evaluation functions have been created for the game of chess based

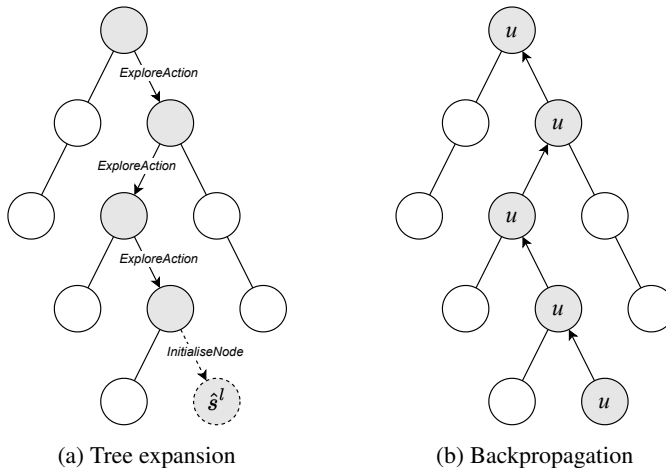


Figure 9.22: Tree expansion and backpropagation in MCTS. Each node in the tree corresponds to a state. The tree is expanded by sampling actions (edges) until a previously unvisited state \hat{s}' is reached, and a new node for the state is initialised and added to the tree (dashed circle). The new state \hat{s}' is evaluated via $u = f(\hat{s}')$, and u is propagated back through the predecessor nodes in the tree until reaching the root node (e.g. using Equation 9.75).

on vast amounts of expert knowledge (Levy and Newborn 1982; Campbell, Hoane Jr, and Hsu 2002). More generally, a domain-agnostic approach could be to sample actions uniform-randomly until a terminal state is reached, but such an approach can be highly inefficient.

Once the new state \hat{s}^l has been initialised in the search tree and a value estimate $u = f(\hat{s}^l)$ is obtained, MCTS backpropagates u and the rewards \hat{r}^τ through the nodes that have been visited in the simulation, starting with the predecessor node \hat{s}^{l-1} and following its predecessors until reaching the root node. For each node \hat{s}^τ to be updated, the function $Update(Q, \hat{s}^\tau, \hat{a}^\tau)$ increases the counter $N(\hat{s}^\tau, \hat{a}^\tau) = N(\hat{s}^\tau, \hat{a}^\tau) + 1$, and the action value function Q is updated. In general, Q may be updated using any of the known temporal-difference learning rules used in RL (Section 2.6). For example, we may update $Q(\hat{s}^\tau, \hat{a}^\tau)$ using the off-policy Q-learning update rule with learning target $\hat{r}^\tau + \gamma u$ if $\tau = l - 1$, and learning target $\hat{r}^\tau + \gamma \max_{a' \in A} Q(\hat{s}^{\tau+1}, a')$ if $\tau < l - 1$. However, if the MDP is terminating (i.e. each episode will terminate at some point) and all rewards are zero until a terminal state is reached, such as in the zero-sum board games

considered in this section, then u can be backpropagated directly via⁷²

$$Q(\hat{s}^\tau, \hat{a}^\tau) \leftarrow Q(\hat{s}^\tau, \hat{a}^\tau) + \frac{1}{N(\hat{s}^\tau, \hat{a}^\tau)} [u - Q(\hat{s}^\tau, \hat{a}^\tau)] \quad (9.75)$$

which computes the average value of u for state-action pair $\hat{s}^\tau, \hat{a}^\tau$. Note that, in this case, if \hat{s} is a terminal state giving reward \hat{r} (such as +1 for win, -1 for lose), then the evaluation $u = f(\hat{s})$ should be equal or close to the reward \hat{r} . Figure 9.22 depicts the process of expanding the search tree and backpropagating u through the predecessor nodes.

After the k simulations have been completed, MCTS uses $BestAction(s^t)$ to select an action a^t for the current state s^t , based on the information stored in the node corresponding to s^t . In general, $BestAction(s^t)$ returns a probability distribution π^t over actions $a \in A$. A common choice is to select a most-tried action, $a^t \in \arg \max_a N(s^t, a)$, or an action with the highest value, $a^t \in \arg \max_a Q(s^t, a)$. After executing a^t and observing the new state s^{t+1} , the MCTS procedure repeats and k simulations are performed in s^{t+1} , and so forth.

9.7.2 Self-Play MCTS

The idea of policy self-play is to train an agent's policy against itself, meaning the same policy is used to choose actions for each agent. This requires that the agents in the game have perfectly symmetrical roles and ego-centric observations, such that the same policy can be used from the perspective of each agent. In zero-sum board games, the focus of this section, the agents have symmetric roles since they are direct opponents (each agent is trying to defeat the other agent) and they generally have access to the same types of actions. In contrast, if the agents have non-symmetrical roles in the game, such as offence and defence players in a football team, then using the same policy for each agent does not make sense since the different roles require different actions.

An agent's observation is ego-centric if the information contained in the observation is relative to the agent. In the chess example, in which agents observe the full state of the game, the state may be represented as a vector $s = (i, x, y)$ where i is the number of the agent whose turn it is, x is a vector containing the locations of agent i 's pieces (set to -1 if the piece was removed), and y is a similar vector for the opponent's pieces. Suppose agent 1 controls the white pieces and its policy π_1 is conditioned on this state vector. Therefore, π_1 will assume that the information about white pieces is located in the x -part of the state vector. In order to use agent 1's policy π_1 to control the black pieces for agent 2, we can transform the state vector $s = (2, x, y)$ (in which x stores the

72. Equation 9.75 additionally assumes an undiscounted return objective.

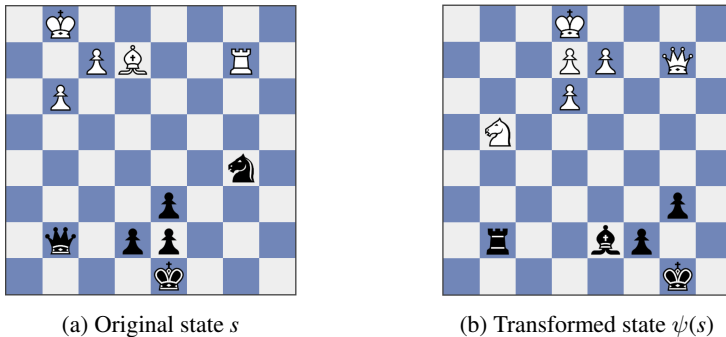


Figure 9.23: State transformation in chess. The state s is transformed to $\psi(s)$ by rotating the board by 180 degrees and swapping the colour of the pieces.

locations of the black pieces, and y for white pieces) by changing the agent number and swapping the order of the x/y vectors, $\psi(s) = (1, y, x)$. If we visualise the state s , as shown in Figure 9.23, where white pieces are located at the top of the board and black pieces are at the bottom, then the transformation $\psi(s)$ corresponds to rotating the board by 180 degrees and swapping the colours of the pieces from black to white and vice versa. The transformed state $\psi(s)$ can now be used by agent 1's policy to choose an action for agent 2, i.e. $a_2 \sim \pi_1(\cdot | \psi(s))$.

Given such a state transformation method, we can adapt the general MCTS algorithm shown in Algorithm 21 to implement a policy self-play approach for zero-sum games in which the agents take turns. Essentially, from the perspective of agent 1, in self-play the MCTS simulations become a Markov decision process in which agent 1 chooses the action in each time step. Using \hat{s}_i^τ to denote the state and agent i whose turn it is to choose an action in that state, the self-play simulation process for agent 1 becomes:

$$\pi_1(\hat{s}_1^\tau) \rightarrow \hat{a}^\tau \rightarrow \pi_1(\psi(\hat{s}_2^{\tau+1})) \rightarrow \hat{a}^{\tau+1} \rightarrow \pi_1(\hat{s}_1^{\tau+2}) \rightarrow \hat{a}^{\tau+2} \rightarrow \pi_1(\psi(\hat{s}_2^{\tau+3})) \rightarrow \hat{a}^{\tau+3} \dots \quad (9.76)$$

To implement this process in the MCTS simulations, we modify the algorithm to replace \hat{s}^τ with \hat{s}_i^τ to keep track of which agent is choosing the action in the state, and in the function calls of *ExploreAction* (Line 8), *InitialiseNode* (Line 13), and *Update* (Line 16) we input \hat{s}_i^τ if $i = 1$, and $\psi(\hat{s}_i^\tau)$ if $i \neq 1$. Additionally, since the evaluation $u = f(s^l)$ is always from the perspective of agent 1 (e.g. +1 reward if agent 1 wins), in function *Update* when using Equation 9.75 to update $Q(\hat{s}_i^\tau, \hat{a}^\tau)$, we have to flip the sign of u if $i \neq 1$. Thus, these functions always operate on states as if it was agent 1's turn in each state. Note that

MCTS controls agent 1, hence the time steps t in Line 2 include only the times in which it is agent 1's turn.

The above version of self-play MCTS uses the *current* version of the policy to select actions for each agent. This algorithm can be further modified by training the policy not just against the current version of itself, but also against *past* versions of the policy. For example, we may maintain a set Π which contains copies of agent 1's policies (or the Q functions from which they are derived) from different points of the training process. Using Π , the self-play MCTS algorithm may sample any policy from this set when selecting actions for agent 2. In this way, we may obtain a potentially more robust policy for agent 1 by forcing it to perform well against not just one specific policy (itself) but also against past versions of the policy. Thus, Π serves a similar purpose to the replay buffer \mathcal{D} (Section 8.1.3), which is to reduce overfitting.

9.7.3 Self-Play MCTS with Deep Neural Networks: AlphaZero

AlphaZero (Silver et al. 2018) is based on MCTS with self-play against the current policy, as described in the previous sections. Additionally, it uses a deep convolutional neural network parameterised by θ to learn an evaluation function

$$(u, p) = f(s; \theta) \quad (9.77)$$

which, for any input state s , predicts two elements:

- u : The expected outcome of the game from state s , $u \approx \mathbb{E}[z|s]$, where z is either +1 for a win, -1 for a loss, and 0 for a draw. Thus, u estimates the (undiscounted) value of state s .
- p : A vector of action selection probabilities in state s , with entries $p_a = \Pr(a|s)$ for each action $a \in A$, to predict the action probabilities computed by MCTS in the *BestAction* function for state s .

AlphaZero learns these value estimates and action probabilities entirely from self-play, starting with randomly initialised parameters θ and using stochastic gradient descent to minimise a combined loss function

$$\mathcal{L}(\theta) = (z - u)^2 - \pi^\top \log p + c \|\theta\|^2 \quad (9.78)$$

with parameter c to control the weight of the squared L2-norm, $\|\theta\|^2 = \theta^\top \theta$, which acts as a regulariser to reduce overfitting. This loss is computed based on data $D = \{(s^t, \pi^t, z^T)\}$, where for each episode in the MCTS algorithm (Algorithm 21), s^t is the state at time t (Line 3), π^t is the action probability distribution for state s^t computed in *BestAction* (Line 18), and z^T is the game outcome in the last time step T of the episode. As usual, a mini-batch of this data is sampled when computing the loss in stochastic gradient descent. Thus, $f(s)$ learns value

AlphaZero (playing white) vs.	Win	Draw	Loss
Stockfish (chess)	29.0%	70.6%	0.4%
Elmo (shogi)	84.2%	2.2%	13.6%
AlphaGo Zero (Go)	68.9%	–	31.1%

Figure 9.24: AlphaZero match results reported by Silver et al. (2018). Results show percentage of won/drawn/lost games for AlphaZero when playing white.

estimates u to predict the outcome z , and action probabilities p to predict the MCTS action probabilities π , for any given state s .

The action probability vector p is used in AlphaZero to guide the action selection in the MCTS simulations. When expanding the search tree for new node \hat{s}^l (Line 13), the function *InitialiseNode*(\hat{s}^l) computes $(u, p) = f(\hat{s}^l; \theta)$ with the current parameters θ and, in addition to setting $N(\hat{s}^l, \hat{a}) = 0$ and $Q(\hat{s}^l, \hat{a}) = 0$, initialises action prior probabilities $P(\hat{s}^l, \hat{a}) = p_{\hat{a}}$ for all $\hat{a} \in A$. The function *ExploreAction* (Line 8) uses N, Q, P in a formula similar to the UCB (Equation 9.74) method

$$\hat{a}^\tau = \begin{cases} \hat{a} & \text{if } N(\hat{s}^\tau, \hat{a}) = 0, \text{ else} \\ \arg \max_{\hat{a} \in A} \left(Q(\hat{s}^\tau, \hat{a}) + C(\hat{s}^\tau)P(\hat{s}^\tau, \hat{a}) \frac{\sqrt{N(\hat{s}^\tau)}}{1+N(\hat{s}^\tau, \hat{a})} \right) \end{cases} \quad (9.79)$$

where $C(\hat{s})$ is an additional exploration rate. Therefore, AlphaZero explores actions similarly to UCB, but biases the action selection based on the predicted probabilities p . After completing the MCTS simulations from the root node s^t , AlphaZero uses a function *BestAction*(s^t) that selects an action a^t either proportionally (for exploration/training) with respect to the root visit counts $N(s^t, \cdot)$, i.e. $a^t \sim \frac{N(s^t, \cdot)}{\sum_{a \in A} N(s^t, a)}$, or greedily $a^t \in \arg \max_a N(s^t, a)$.

The complete specification of AlphaZero includes various additional implementation details about the network representation of f , the exploration rate C , the addition of exploration noise in action probabilities, the sampling of data batches to compute the loss, action masking, and so on. These details can be found in the supplementary material of (Silver et al. 2018). However, the core idea behind AlphaZero is to learn an effective evaluation function f purely from self-play MCTS, and to use this function to guide the tree search in a vast search space in order to find optimal actions.

Silver et al. (2018) reported results for AlphaZero in the games of chess, shogi, and Go, in which it was matched against several strong game-playing programs: “Stockfish” for chess, “Elmo” for shogi, and AlphaGo Zero (Silver et al. 2017) for Go trained for three days. In each game, AlphaZero performed

$k=800$ MCTS simulations in each state s^t . Separate instances of AlphaZero were trained for 9 hours (44 million games) in chess, 12 hours (24 million games) in shogi, and 13 days (140 million games) in Go. Based on Elo rating (Elo 1960), AlphaZero first outperformed Stockfish after 4 hours, Elmo after 2 hours, and AlphaGo Zero after 30 hours. Figure 9.24 shows the reported match results for the final trained instances of AlphaZero (when playing white) in the three games. AlphaZero was able to achieve this performance using the same general self-play MCTS method in each game, without requiring heuristic evaluation functions based on expert knowledge.⁷³ While Stockfish and Elmo evaluated approximately 60 million and 25 million states per second, respectively, AlphaZero evaluated only about 60 thousand states per second in chess and shogi. AlphaZero compensated for the lower number of evaluations by using its deep neural network to focus more selectively on promising actions.

73. However, AlphaZero *did* use domain knowledge in the construction of the input and output features of f (e.g. legality of castling, repetition count for current position, underpromotions for pawn moves, piece drops in shogi, etc).

10

Multi-Agent Deep RL in Practice

In this chapter, we will dive into the implementation of MARL algorithms. The book comes with a codebase, called “FastMARL”, which is built using Python and the PyTorch framework, and it implements some of the key concepts and algorithms we have discussed in earlier chapters. The codebase provides a practical and easy-to-use platform for experimenting with MARL algorithms. The goal of this chapter is to provide an overview of how these algorithms are implemented in the codebase, and provide the reader with an understanding of coding patterns that are repeatedly found in MARL algorithm implementations.

This chapter assumes knowledge of the Python programming language and a basic understanding of the PyTorch framework. The code examples shown in this chapter are meant to be educational and describe some of the ideas that this book explored in earlier chapters, and are not necessarily to be used as-is. For MARL, many codebases exist that efficiently implement many of the algorithms discussed in the deep MARL chapters: EPyMARL for common-reward games (<https://github.com/uoe-agents/epymarl>), Mava (<https://github.com/instadeepai/Mava>), and more.

10.1 The Agent-Environment Interface

Interacting with an environment is critical to the implementation of MARL algorithms. In a MARL setting, multiple agents interact with an environment simultaneously, and each agent’s actions affect the environment state and other agents’ actions. However, unlike the single-agent RL case, MARL does not have a unified interface for environments. Different frameworks use different environment interfaces, making it challenging to implement an algorithm that seamlessly works on all environments.

Nevertheless, the general idea of an agent-environment interface remains the same in MARL as in RL. The environment provides a set of functions that

agents can use to interact with the environment. Typically, the environment has two critical functions: `reset()` and `step()`. The `reset()` function initialises the environment and returns the initial observation, while the `step()` function advances the environment by one timestep, taking an agent's action as input and returning the next observation, reward, and termination flag. In addition to these functions, the environment should be able to describe the observation space and the action space. The observation space defines the possible observations that agents can receive from the environment, while the action space defines the possible actions that agents can take.

In single-agent RL, the most common interface is called Gym (Brockman et al. 2016) and, with some minor modifications, can support many multi-agent environments. An example of this interface can be found in level-based foraging, which can be imported in Python using the code in Code Block 10.1. The `gym.make()` command takes as an argument the name of the environment, which in this case further defines the exact task parameters (grid size, number of agents, and number of items).

```
1 import lbforaging
2 import gym
3
4 env = gym.make("Foraging-8x8-2p-1f-v2")
```

Code Block 10.1: Creating an environment.

The observation and action spaces can be retrieved from the environment using the code in Code Block 10.2. In the case of level-based foraging, the observation space is a 15-dimensional vector for each agent (noted as `Box` in the Gym library when it contains float values) containing information on the placement of agents and items, along with their levels. The action space, which is `Discrete(6)` for each agent, specifies that the expected actions are discrete numbers (0, 1, ..., 5) corresponding to the four movement actions, the loading action, and the noop action which does nothing. Notably, both the observation and action spaces are *tuples* and their n elements correspond to each agent. In Code Block 10.2, the tuples are of size two, which means that two agents exist in the environment.

```
1 env.observation_space
2 >> Tuple(Box(..., 15), Box(..., 15))
3
4 env.action_space
5 >> Tuple(Discrete(6), Discrete(6))
```

Code Block 10.2: Observation and action spaces.

The crucial step of interacting with the environment is shown in Code Block 10.3. The `reset()` function initialises the environment and returns the initial observation for each agent. This can be used to select the actions a_i^0 for each agent i which are then passed to the `step()` function. The `step()` function simulates the transition from a state to the next given the joint action, and returns the n observations (of the next state), a list of n rewards, whether the episode has terminated, and an optional `info` dictionary (which we ignore in the Code Block using Python's `_` notation).

```
1 observations = env.reset()
2 next_observations, rewards, terminal_signal, _ = env.step(
    actions)
```

Code Block 10.3: Observing and acting on an environment.

Some multi-agent environments (which will be discussed in Chapter 11) do not conform to this exact interface. However, the simple interface described above can model POSGs, with other interfaces mostly used to extend it with additional capabilities such as limiting the actions available to agents during the episode, or providing the full environment state. FastMARL uses the above interface and requires wrappers for environments that have a different way to provide the necessary information to the learning algorithms.

10.2 MARL Neural Networks in PyTorch

The first step in implementing a deep learning algorithm is the process of setting up a simple neural network model. In RL, neural networks can be used to represent a policy, a state value, or a state-action value function. The exact specifications of a model depend on the algorithm that will be implemented, but the main structure is similar across algorithms.

The FastMARL codebase provides a flexible architecture for designing a fully connected neural network per agent to be used in a MARL scenario. The code defines a specific number of hidden layers, number of units in each hidden layer, and type of non-linearity applied in each unit, which can be changed to best fit the specific problem being modelled. The input to the module's constructor allows for different networks to be created, by changing the size of the inputs and outputs for each of the agents. In this implementation, the networks defined in Code Block 10.4 below are *independent* of each other.

```
1 import torch
2 from torch import nn
```

```

3 from typing import List
4
5 class MultiAgentFCNetwork(nn.Module):
6     def __init__(self,
7                  in_sizes: List[int],
8                  out_sizes: List[int]
9                  ):
10        super().__init__()
11
12        # We use the ReLU activation function:
13        activ = nn.ReLU
14        # We use two hidden layers of 64 units each:
15        hidden_dims = (64, 64)
16
17        n_agents = len(in_sizes)
18        # The number of agents is the length of the
19        # input and output vector
20        assert n_agents == len(out_sizes)
21
22        # We will create 'n_agents' (independent) networks
23        self.networks = nn.ModuleList()
24
25
26        # For each agent:
27        for in_size, out_size in zip(in_sizes, out_sizes):
28            network = [
29                nn.Linear(in_size, hidden_dims[0]),
30                activ(),
31                nn.Linear(hidden_dims[0], hidden_dims[1]),
32                activ(),
33                nn.Linear(hidden_dims[1], out_size),
34            ]
35            self.networks.append(nn.Sequential(*network))
36
37    def forward(self, inputs: List[torch.Tensor]):
38
39        # The networks can run in parallel:
40        futures = [
41            torch.jit.fork(model, inputs[i])
42            for i, model in enumerate(self.networks)
43        ]
44        results = [torch.jit.wait(fut) for fut in futures]
45        return results

```

Code Block 10.4: Example of neural network for MARL.

Code Block 10.4, Lines 27 to 35 iterate over the input and output sizes of each agent and extend the list of the separate neural networks (Line 35). While all modules are included in the same model, the actual networks are independent and can even be run in parallel. The loop in Line 40 begins a parallel forward pass through these networks, which makes the computation begin in the background. The program can continue while PyTorch computes the forward pass. In Line 44, the program waits until the computation finishes

and the result is ready (similarly to the `async/await` pattern found in many programming languages).

10.2.1 Seamless Parameter Sharing Implementation

Section 9.6.1 discusses parameter sharing, a prevalent paradigm when implementing MARL algorithms, and especially useful in environments with homogeneous agents. Parameter sharing requires agents to have a single network and all inputs and outputs can derive from that. Parameter sharing can be seamlessly interchanged in the model by defining a parameter-sharing variation as shown in Code Block 10.5.

```

1 class MultiAgentFCNetwork_SharedParameters(nn.Module):
2
3     def __init__(
4         self,
5         in_sizes: List[int],
6         out_sizes: List[int]
7     ):
8
9         # ... same as MultiAgentFCNetwork
10
11        # We will create one (shared) network
12        # This assumes that input and output size
13        # is identical across agents. If not, we
14        # should first pad the inputs and outputs
15
16        network = [
17            # ... same as MultiAgentFCNetwork
18        ]
19        self.network = nn.Sequential(*network)
20
21    def forward(self, inputs: List[torch.Tensor]):
22
23        # A forward pass of the same network in parallel
24        futures = [
25            torch.jit.fork(self.network, inp)
26                for inp in inputs)
27        ]
28        results = [torch.jit.wait(fut) for fut in futures]
29        return results

```

Code Block 10.5: Example of shared neural network.

The difference between the networks created in Code Block 10.4 and 10.5 is the lack of a loop that creates many sub-networks in the second (Code Block 10.4, Line 27), and instead, the definition of a single sequential module (Code Block 10.5, Line 19). The parallel execution of the forward operation in Code Block 10.5's Line 24 calls the same network each time instead of iterating over a list.

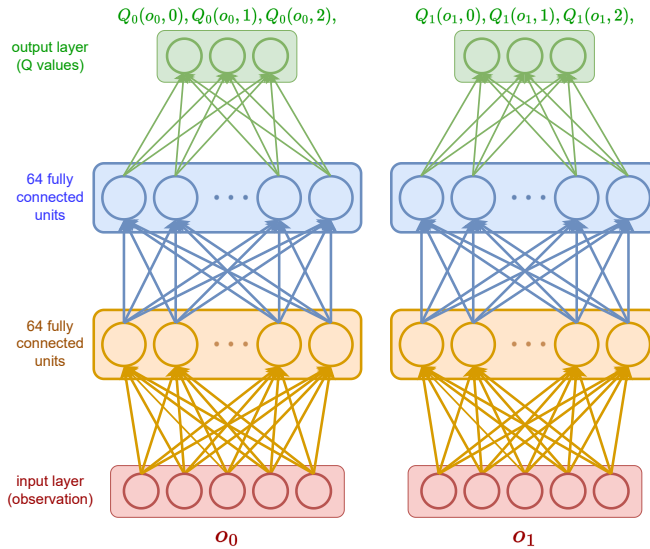


Figure 10.1: The resulting MultiAgentFCNetwork of Code Block 10.6 for the IDQN algorithm with two agents, an observation of size 5, and three possible actions for each agent.

10.2.2 Defining the Models: An Example with IDQN

The models that were discussed in the previous section can be easily initialised for use in a MARL setting. For instance, to use either of them in IDQN (Algorithm 16) we only have to understand and define the sizes of our action and observation spaces. DQN uses a single value network that is structured in a way that receives as an input the individual observation \$o_i\$ and outputs a vector of Q-values for each of the possible actions. Below (Code Block 10.6) is an example of how to create a model to be used with IDQN, with two agents, an observation size of five, and three possible actions per agent.

```

1 # Example of observation of agent 2:
2 # obs1 = torch.tensor([1, 0, 2, 3, 0])
3
4 # Example of observation of agent 1:
5 # obs2 = torch.tensor([0, 0, 0, 3, 0])
6
7 obs_sizes = (5, 5)
8
9 # Example of action of agent 1:
10 # act1 = [0, 0, 1] # one-hot encoded

```

```

11
12 # Example of action of agent 2:
13 # act2 = [1, 0, 0] # one-hot encoded
14
15 action_sizes = (3, 3)
16
17 model = MultiAgentFCNetwork(obs_sizes, action_sizes)
18
19 # Alternatively, the shared parameter model can be used instead:
20 # model = MultiAgentFCNetwork_SharedParameters(
21 #     obs_sizes, action_sizes
22 #)
```

Code Block 10.6: Example of initialising the models.

The resulting networks produced by the Code Block 10.6 above can be seen in Figure 10.1. The IDQN algorithm can be built on top of these networks. To predict (and later learn) state-action value for each agent for the IDQN algorithm, one can use Code Block 10.7, below.

```

1 # obs1, obs2, model as above
2
3 q_values = model([obs1, obs2])
4 >> ([Q11, Q12, Q13], [Q21, Q22, Q23])
5 # where  $Q_{ij}$  is the Q value of agent i doing action j
```

Code Block 10.7: Sampling Q-values.

In PyTorch, executing operations (such as summation, or other functions) in an additional “agent” dimension is a very powerful tool. Say, for example, that the algorithm needs to calculate the actions with the highest Q values per agent (e.g. for use in the Bellman equation). This can be done in parallel in a similar way as in the single agent case, as shown in Code Block 10.8.

```

1 # we are creating a new "agent" dimension
2 q_values_stacked = torch.stack(q_values)
3 print(q_values_stacked.shape)
4 >> [2, 3]
5 # 2: agent dimension, 3: action dimension
6
7 # calculating best actions per agent (index):
8 _, a_prime = q_values_stacked.max(-1)
```

Code Block 10.8: Stacking Q-values.

Implementing the full IDQN algorithm presented in Algorithm 16 requires a framework to interact with the environment, an implementation of a replay buffer, and a target network. However, with the basics covered in this section, implementing IDQN should be a relatively simple exercise.

10.3 Centralised Value Functions

This section will cover another basic idea in MARL, which is to build algorithms which condition value functions or policies on “privileged” information (Section 9.4.2).

The IDQN algorithm uses the observations as input to the action value network. A simple alternative is conditioning the action value on the concatenation of the observations, which may give a better approximation of the state in partially observable environments. The State-Conditioned Actor-Critic algorithm, first presented in Section 9.4.2, uses such a critic. An example in PyTorch for concatenating the observations and creating the critic for this algorithm is shown in Code Block 10.9, below.

```

1 centr_obs = torch.cat([obs1, obs2])
2 print(centr_obs)
3 >> tensor([1, 0, 2, 3, 0, 0, 0, 0, 3, 0])
4
5 # we use an input size of 5+5=10, once for each agent
6 critic = MultiAgentFCNetwork([10, 10], [1, 1])
7
8 values = critic(2*[centr_obs]) # outputs the state value for
      each agent

```

Code Block 10.9: Concatenated observations.

Line 1 uses PyTorch’s concatenation function to merge the observations of two agents. The input size of the critic is increased accordingly to 10 (Line 6), while the dimensionality of the output is 1 given that it returns the value of the joint observation under the policy (but not action). This model is now conditioned on the concatenation of the observations, and produces better estimates in partially observable environments.

To adhere to the CTDE paradigm, the policy network (actor) is just conditioned on the observation of the agent. An example of how to initialise and how to sample actions from that network can be seen in Code Block 10.10.

```

1 actor = MultiAgentFCNetwork([5, 5], [3, 3])
2
3 from torch.distributions import Categorical
4 actions = [Categorical(logits=y).sample() for y in actor([obs1,
      obs2])]

```

Code Block 10.10: Sampling actions from a policy network.

Line 4 creates a categorical distribution for each agent, from the outputs of the network. The categorical distribution can be sampled to produce the actions that the agents will perform.

10.4 Value Decomposition

In common-reward environments, an algorithm could employ a value decomposition method such as VDN or QMIX (Sections 9.5.2 and 9.5.3). In Code Block 10.11 we show how value decomposition with VDN can be implemented in practice.

```

1 # The critic and target are both MultiAgentFCNetwork
2 # The target is "following" the critic using soft or hard
   updates
3
4 # obs and nobs are List[torch.tensor] containing the
5 # observation and observation at t+1 respectively
6 # For each agent
7
8 with torch.no_grad():
9     q_tp1_values = torch.stack(critic(nobs))
10    q_next_states = torch.stack(target(nobs))
11    all_q_states = torch.stack(self.critic(obs))
12
13 _, a_prime = q_tp1_values.max(-1)
14
15
16 target_next_states = q_next_states.gather(
17                         2, a_prime.unsqueeze(-1)
18                         ).sum(0)
19
20 # Notice .sum(0) in the line above.
21 # This command sums the Q values of the next states
22
23 target_states = rewards + gamma*target_next_states*(1-
   terminal_signal)
```

Code Block 10.11: Value decomposition with VDN.

Remember that in the example above, our rewards are one-dimensional, meaning one reward for the joint action. We, therefore, sum the outputs of the state-action networks and attempt to approximate the returns using that sum. More complicated solutions (e.g. QMIX) can be used in place of a simple summation operation to enforce assumptions (e.g. non-linearity, or the monotonicity constraint).

10.5 Practical Tips for MARL Algorithms

Similarly to other areas of machine learning, the implementation of MARL algorithms also demands significant engineering efforts. This section provides a number of useful tips for implementing such algorithms. It should be noted that not all of these tips will be applicable to every MARL problem, given the wide-ranging and diverse assumptions that exist in this field. Nonetheless, gaining a grasp of these concepts can prove to be useful.

10.5.1 Stacking Timesteps vs. Recurrent Network vs. Neither

In Part I of the book, we focused on the theoretically sound approach of conditioning policies on the history of observations (i.e. $o_i^{1:t}$), especially in POSGs. But, as discussed in Section 8.3, neural networks have a predefined structure which does not allow a variable input length. Even if this was circumvented (e.g. by padding the inputs), in large and complicated environments, large inputs with repeated information could even have a detrimental effect on learning. A MARL practitioner implementing an algorithm with neural networks can choose between three options: i) stack a small number of observations (e.g. $o_i^{t-5:t}$), ii) use a recurrent neural network (e.g. LSTM or GRU), or iii) simply ignore the previous observations and assume that o_i^t carries all necessary information for deciding on an action. Using recurrent structures is the closest to the theoretically sound solution, but in practice, recurrent architectures suffer from the vanishing gradient problem which can lead to information far in the past being unused.

When should we use each approach? We cannot have a definite answer to this question before we have experimental results on a specific environment. We could, however, consider the following: How important is the information contained in previous observations to the agents' decisions? If almost all information is contained in the last observation (e.g. a fully observable environment), then it is possible that using previous observations will not lead to improved performance. If all information is contained in a set amount of timesteps, then a starting point would be to concatenate the last timesteps before providing them to the network. Finally, if the information contained far in the past is required (e.g. navigating a long maze), then a recurrent network might prove useful.

10.5.2 Standardising Rewards

Research in single-agent RL has demonstrated empirical improvements when standardising the rewards, and returns. These improvements are also observed in MARL. Many MARL environments have rewards that span many orders of

magnitude (e.g. multi-agent particle environments discussed in Section 11.3.2), hindering the ability of neural networks to approximate them. So, an algorithm could be more efficient by using rewards that are standardised: the mean of rewards should be zero, with a standard deviation of one. In practice, there are many different ways to implement such a mechanism. For example, one could standardise over the batch, or over a running average and standard deviation of the rewards. Or one could standardise the returns instead of the rewards, as shown in Code Block 10.12, attempting to keep the output of the state value or state-action value network close to zero.

```
1 # Standardising the returns requires a running mean and variance
2 returns = (returns - MEAN) / torch.sqrt(VARIANCE)
```

Code Block 10.12: Example of return standardisation.

Notably, reward standardisation is an empirical tip but can distort the underlying assumptions and goals of an algorithm. For example, consider an environment with only negative rewards: a one-state MDP with two actions. The first one rewards the agent with -1 and ends the episode, while the second action rewards the agent with -0.1 but only ends the episode with a 1% chance. Standardisation would lead to the second action being considered to provide positive rewards instead. While the overall preference between actions would remain the same (the first action leads to lower rewards than the second), the nature of the problem has changed. With negative rewards, the agent aims for the episode to end as early as possible, but now, with potentially positive rewards, the agents could instead prefer staying in the environment for longer. This difference in an agent's goals is subtle, but should be well understood before standardisation of rewards is applied in practice.

10.5.3 Centralised Optimisation

Stochastic gradient descent remains one of the slowest parts of training an RL agent. Many independent learning implementations use a different optimiser instance for each agent. Multiple co-existing optimisers can ensure the independence between agents by having a separate list of trainable and internal parameters. However, such an implementation can be extremely time-consuming and does not use parallelisation. Instead, using a single optimiser encompasses all the trainable parameters, even if the agents consist of different neural networks or algorithms, and can be significantly faster. The final losses can just be added before the stochastic gradient descent steps. An example in PyTorch can be seen in Code Block 10.13

```

1 params_list = list(nn_agent1.parameters())
2         + list(nn_agent2.parameters())
3         + list(nn_agent3.parameters())
4         + ...
5 common_optimiser = torch.optim.Adam(params_list)
6 ...
7 loss = loss1 + loss2 + loss3 + ...
8 loss.backward()
9 common_optimizer.step()

```

Code Block 10.13: Example of single optimiser.

Lines 1 to 4 create a list of parameters to be used by the optimiser. (The MultiAgentFCNetwork introduced in Code Block 10.6 already does this automatically). Summing the losses of each agent (Line 7) creates a single loss to be used for the gradient descent step (Lines 8 and 9).

10.6 Presentation of Experimental Results

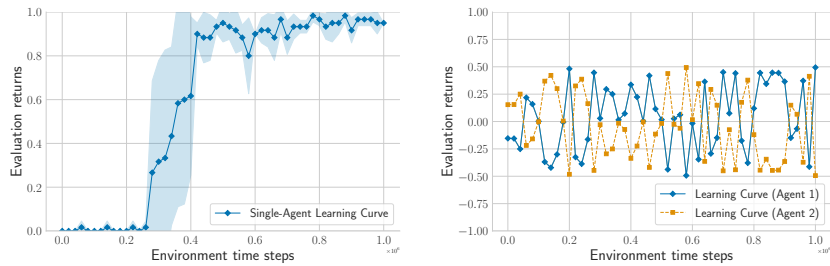
Comparing algorithms and presenting their differences is arguably more difficult in MARL than typical supervised learning or even single-agent RL. There are two main reasons these comparisons are not straightforward: i) sensitivity to hyperparameters or training seeds and ii) solution concepts that go beyond one-dimensional representations (such as the accuracy in supervised learning).

This section discusses how to perform fair comparisons between algorithms in MARL.

10.6.1 Learning Curves

Single-agent RL can often present learning performance in a “learning curve” (see Section 2.7 in page 36), a two-dimensional plot where the x-axis represents the training time, or environment timesteps, and the y-axis shows an estimation of the agent’s episodic returns. An example of such a learning curve can be seen in Figure 10.2a, where the performance of a single agent learning on an environment is displayed. There are two steps to creating the information needed to reproduce such a figure:

- An evaluation procedure for the parameters of a policy ϕ , or action value θ , that outputs an estimation of the mean returns of an agent in an environment. This procedure should run at regular intervals during training.
- The outputs of the evaluations *across different seeds*. Then, the average and the standard error (or deviation) of the mean returns approximated in the previous step are needed for the final plots.



(a) Example of a single-agent learning curve.

(b) Example of a learning curve in a zero-sum game with two agents.

Figure 10.2: Examples of learning curves in single-agent and multi-agent reinforcement learning with independent A2C. The learning curve in a zero-sum multi-agent game is not informative and cannot demonstrate whether agents learn during training.

Learning curves are easy to read and to parse, therefore they are often used in MARL too. In common-reward games, the common reward can be used during evaluation and should be sufficient to provide informative learning curves similar to single-agent RL. However, this might not be the case in other types of games, such as zero-sum games. For example, Figure 10.2b shows the learning curves of two agents in a zero-sum game. It is immediately clear that the learning curves in this example are not informative and cannot inform of the learned abilities of the agents.

There are still options available to make learning curves informative even in zero-sum games. For example, a pre-trained or heuristic agent could be used as a static opponent and every evaluation procedure should match the learning agents with that stationary opponent. This way, the MARL practitioner can assess the abilities of the agents as they learn. The disadvantage of this method is apparent when the game is hard enough to make heuristic agents difficult to program, or when there is a gap between the abilities of the heuristic and learned agents (e.g. a heuristic chess agent with high Elo rating will not offer informative signals before the learned agent also approaches that rating). Another way to monitor the performance in zero-sum games is by saving previous instances of the trainable parameters, creating a pool of opponent agents, and ensuring that newer agents can typically win against them. However, this does not necessarily make comparisons easy, unless the pool of agents is common between different algorithms.

Finally, the information in the learning curves could potentially be condensed into a single number, either by presenting the maximum value of the curve or its average. These values offer different functions, the maximum can be used to signify whether an algorithm at any point *solved* the environment, and the average signifies how fast the algorithm learned in the environment as well as its stability. The maximum can be considered more important since it can inform us whether a desired behaviour has been achieved, and should be preferred unless the algorithms reach the same value (at which point stability and speed become important).

10.6.2 Hyperparameter Search

The fact that multiple agents are learning concurrently, and they each affect the learning of other agents, makes MARL especially sensitive to hyperparameters. In turn, this sensitivity to hyperparameters makes comparisons between algorithms complicated.

Therefore, in most cases, a thorough hyperparameter search is necessary for finding hyperparameters that work in a specific problem. The quantity of hyperparameters to be tested depends on the complexity of the problem and the computational resources available. A simple, highly parallelisable solution is to run a grid search across many combinations of hyperparameters and multiple seeds. Then, the best hyperparameter combination can be found by comparing the runs using the metrics (e.g. maximum) described in Section 10.6.1.

A parallel hyperparameter search can be executed by independently calling the training programs with different input parameters (the hyperparameters). For example, the Bash script shown in Code Block 10.14 iterates over various learning rates by providing them as inputs to a Python script. The ampersand symbol (&) at the end of Line 4 indicates the end of the command, but causes Bash to execute it asynchronously.

```

1 for s in {1..5}
2   for i in $(seq 0.01 0.01 0.1)
3     do
4       python algorithm.py --lr=$i --seed=$s &
5     done
6 done

```

Code Block 10.14: Example of Hyperparameter Search.

Of course, more complicated hyperparameter searches would require more complicated scripts to launch and collect the experiments. However, the procedure should be similar in nature. The size of the hyperparameter search is limited to the available computational power, but still, a larger search leads to

more confidence in the ability of an algorithm to learn in an environment. A rule of thumb is to start the hyperparameter search near values that are known to be sensible and mainly focus on the hyperparameters that control the exploration, such as the entropy coefficient.

11

Multi-Agent Environments

Chapter 3 introduced a hierarchy of game models to formalise interaction processes in multi-agent systems, including normal-form games, stochastic games, and POSG. Based on these game models, a number of multi-agent environments have been implemented in the MARL research community which serve as benchmarks and “playgrounds” for MARL algorithms, and allow us to evaluate and study such algorithms. This chapter presents a selection of existing multi-agent environments.

The purpose of this chapter is two-fold: Firstly, the environments presented here serve as concrete examples of the game models used in this book. They illustrate a range of situations and learning challenges which MARL algorithms have to tackle. Secondly, for the reader interested in experimenting with MARL algorithms, the environments presented in this chapter are a first starting point. The code implementations for the environments are freely available from their respective sources.

In Section 11.1, we will first discuss a set of criteria to consider when selecting multi-agent environments, in particular regarding the mechanics of the environment (e.g. state/action dynamics and observability) and the different types of learning challenges involved. Section 11.2 will then present a taxonomy of 2×2 matrix games (i.e. normal-form games with two agents and two actions each) which are further classified into no-conflict games and conflict games. This taxonomy is complete in the sense that each game is structurally distinct from the other games in the set, and includes ordinal versions of games such as Prisoner’s Dilemma, Chicken, and Stag Hunt which were discussed in previous chapters of this book. Moving on to the more complex game models of stochastic games and POSGs, Section 11.3 will present a selection of multi-agent environments in which agents are faced with challenges such as complex state/action spaces, partial observability, and sparse rewards.

11.1 Criteria for Choosing Environments

There are a number of considerations when choosing environments to test MARL algorithms. Which properties and learning abilities do we want to test in a MARL algorithm? Relevant properties may include the algorithm’s ability to robustly converge to specific solution concepts; how efficiently it scales in the number of agents; and the algorithm’s ability to learn successfully when there are large state and/or action spaces, partial observability, and sparse rewards (meaning rewards are zero most of the time).

Normal-form games can serve as simple benchmarks and are particularly useful when evaluating fundamental properties of MARL algorithms, such as convergence to specific types of solution concepts. For non-repeated normal-form games, methods exist to compute exact solutions for the different solution concepts, such as the linear programmes for minimax and correlated equilibria shown in Sections 4.3.1 and 4.6.1. The joint policies learned by MARL algorithms in a normal-form game can then be compared to the exact solutions to give an indication of learning success. Normal-form games (if they are relatively small) are also useful for manual inspection of learning processes, and can be used as illustrative examples as we have done in many places in this book.

Environments based on stochastic games and POSGs can be used to test an algorithm’s ability to deal with state/action spaces of varying complexity, partial observability, and sparse rewards. Many such environments can be configured to create tasks of increasing complexity, such as by varying the number of agents and world size; and by varying the degree of partial observability (e.g. setting the observation radius). The most difficult learning tasks usually feature a combination of large state/action spaces, limited observability for agents, and sparse rewards. A downside of using such environments is that it is usually not tractable to compute exact solutions, such as Nash equilibria; though we can test whether a learned joint policy is an equilibrium, as outlined in Section 4.4.

In addition to the properties mentioned above, it is important to consider what kinds of skills do the agents have to learn in an environment in order to solve the task. Different environments may require different kinds of skills in agents, such as deciding when and with what other agents to cooperate (as in LBF, Section 11.3.1), what information to share (as in some MPE tasks, Section 11.3.2), how to position oneself in a team and distribute responsibilities (such as in SMAC and GRF, Sections 11.3.3 and 11.3.5), and many others. A MARL algorithm may succeed at learning some types of skills but not others, and it is important to evaluate such learning abilities.

11.2 Structurally Distinct 2×2 Matrix Games

This section contains a listing of all 78 structurally distinct, strictly ordinal 2×2 matrix games (i.e. normal-form games with two agents and two actions each), based on the taxonomy of Rapoport and Guyer (1966).⁷⁴ The games are *structurally distinct* in that no game can be reproduced by any sequence of transformations of any other game, which includes interchanging the rows, columns, agents, and any combinations thereof in the game's reward matrix. The games are *strictly ordinal*, meaning that each agent ranks the four possible outcomes from 1 (least preferred) to 4 (most preferred), and no two outcomes can have the same rank. The games are further categorised into *no-conflict* games and *conflict* games. In a no-conflict game, the agents have the same set of most preferred outcomes. In a conflict game, the agents disagree on the most preferred outcomes.

The games are presented in the following format:

X	(Y)
$\underline{a_{1,1}}, b_{1,1}$	$a_{1,2}, b_{1,2}$
$a_{2,1}, b_{2,1}$	$\underline{a_{2,2}}, b_{2,2}$

X is the number of the game in our listing, and Y is the corresponding number of the game in the original taxonomy (Rapoport and Guyer 1966). The variables $a_{i,j}$ and $b_{i,j}$, where $i,j \in \{1, 2\}$, contain the rewards for agent 1 (row) and agent 2 (column), respectively, if agent 1 chooses action i and agent 2 chooses action j . A reward pair is underlined if the corresponding joint action constitutes a pure Nash equilibrium, as defined in Section 4.4.

11.2.1 No-Conflict Games

$1 \quad (1)$	$2 \quad (2)$	$3 \quad (3)$	$4 \quad (4)$	$5 \quad (5)$
$\underline{4, 4} \quad 3, 3$	$\underline{4, 4} \quad 3, 3$	$\underline{4, 4} \quad 3, 2$	$\underline{4, 4} \quad 3, 2$	$\underline{4, 4} \quad 3, 1$
$2, 2 \quad 1, 1$	$1, 2 \quad 2, 1$	$2, 3 \quad 1, 1$	$1, 3 \quad 2, 1$	$1, 3 \quad 2, 2$
$6 \quad (6)$	$7 \quad (22)$	$8 \quad (23)^{75}$	$9 \quad (24)$	$10 \quad (25)$
$\underline{4, 4} \quad 2, 3$	$\underline{4, 4} \quad 3, 3$	$\underline{4, 4} \quad 3, 3$	$\underline{4, 4} \quad 3, 2$	$\underline{4, 4} \quad 3, 2$
$3, 2 \quad 1, 1$	$2, 1 \quad 1, 2$	$1, 1 \quad 2, 2$	$2, 1 \quad 1, 3$	$1, 1 \quad 2, 3$

74. The matrix games can be downloaded for use with this book's codebase at: <https://github.com/uoe-agents/matrix-games>

75. Game no. 23 in the original listing (Rapoport and Guyer 1966) has a typo: the reward $a_{2,1}$ should be 1.

11	(26)	12	(27)	13	(28)	14	(29)	15	(30)
<u>4, 4</u>	2, 3	<u>4, 4</u>	2, 2	<u>4, 4</u>	3, 1	<u>4, 4</u>	3, 1	<u>4, 4</u>	2, 1
3, 1	1, 2	3, 1	1, 3	2, 2	1, 3	1, 2	2, 3	3, 2	1, 3

16	(58)	17	(59)	18	(60)	19	(61)	20	(62)
<u>4, 4</u>	2, 3	<u>4, 4</u>	2, 2	<u>4, 4</u>	2, 1	<u>4, 4</u>	1, 3	<u>4, 4</u>	1, 2
1, 1	3, 2	1, 1	3, 3	1, 2	3, 3	3, 1	2, 2	3, 1	2, 3

21	(63)
<u>4, 4</u>	1, 2

11.2.2 Conflict Games

22	(7)	23	(8)	24	(9)	25	(10)	26	(11)
<u>3, 3</u>	4, 2	<u>3, 3</u>	4, 2	<u>3, 3</u>	4, 1	<u>2, 3</u>	4, 2	<u>2, 3</u>	4, 1
2, 4	1, 1	1, 4	2, 1	1, 4	2, 2	1, 4	3, 1	1, 4	3, 2

27	(12)	28	(13)	29	(14)	30	(15)	31	(16)
<u>2, 2</u>	4, 1	<u>3, 4</u>	4, 2	<u>3, 4</u>	4, 2	<u>3, 4</u>	4, 1	<u>3, 4</u>	4, 1
1, 4	3, 3	2, 3	1, 1	1, 3	2, 1	2, 3	1, 2	1, 3	2, 2

32	(17)	33	(18)	34	(19)	35	(20)	36	(21)
<u>2, 4</u>	4, 2	<u>2, 4</u>	4, 1	<u>3, 4</u>	4, 3	<u>3, 4</u>	4, 3	<u>2, 4</u>	4, 3
1, 3	3, 1	1, 3	3, 2	1, 2	2, 1	2, 2	1, 1	1, 2	3, 1

37	(31)	38	(32)	39	(33)	40	(34)	41	(35)
<u>3, 4</u>	2, 2	<u>3, 4</u>	2, 1	<u>3, 4</u>	1, 2	<u>3, 4</u>	1, 1	<u>2, 4</u>	3, 2
1, 3	4, 1	1, 3	4, 2	2, 3	4, 1	2, 3	4, 2	1, 3	4, 1

42	(36)	43	(37)	44	(38)	45	(39)	46	(40)
<u>2, 4</u>	3, 1	<u>3, 4</u>	2, 3	<u>3, 4</u>	1, 3	<u>2, 4</u>	3, 3	<u>3, 4</u>	4, 1
1, 3	4, 2	1, 2	4, 1	2, 2	4, 1	1, 2	4, 1	2, 2	1, 3

47	(41)	48	(42)	49	(43)	50	(44)	51	(45)
<u>3, 4</u>	4, 1	<u>3, 3</u>	4, 1	<u>3, 3</u>	4, 1	<u>2, 4</u>	4, 1	<u>3, 2</u>	4, 1
1, 2	2, 3	2, 2	1, 4	1, 2	2, 4	1, 2	3, 3	2, 3	1, 4

52 (46)	53 (47)	54 (48)	55 (49)	56 (50)
<u>3, 2</u> 4, 1	<u>2, 3</u> 4, 1	<u>2, 2</u> 4, 1	<u>3, 4</u> 4, 3	<u>3, 4</u> 4, 3
1, 3 2, 4	1, 2 3, 4	1, 3 3, 4	2, 1 1, 2	1, 1 2, 2
57 (51)	58 (52)	59 (53)	60 (54)	61 (55)
<u>3, 4</u> 4, 2	<u>3, 4</u> 4, 2	<u>3, 3</u> 4, 2	<u>3, 3</u> 4, 2	<u>2, 4</u> 4, 3
2, 1 1, 3	1, 1 2, 3	2, 1 1, 4	1, 1 2, 4	1, 1 3, 2
62 (56)	63 (57)	64 (64)	65 (65)	66 (66)
<u>2, 4</u> 4, 2	<u>2, 3</u> 4, 2	<u>3, 4</u> 2, 1	<u>2, 4</u> 3, 1	<u>3, 3</u> <u>2, 4</u>
1, 1 3, 3	1, 1 3, 4	1, 2 <u>4, 3</u>	1, 2 <u>4, 3</u>	4, 2 1, 1
67 (67)	68 (68)	69 (69)	70 (70)	71 (71)
<u>2, 3</u> <u>3, 4</u>	<u>2, 2</u> <u>3, 4</u>	<u>2, 2</u> <u>4, 3</u>	<u>3, 4</u> 2, 1	<u>3, 3</u> 2, 1
4, 2 1, 1	4, 3 1, 1	3, 4 1, 1	4, 2 1, 3	4, 2 1, 4
72 (72)	73 (73)	74 (74)	75 (75)	76 (76)
3, 2 2, 1	2, 4 4, 1	2, 4 3, 1	2, 3 4, 1	2, 3 3, 1
4, 3 1, 4	3, 2 1, 3	4, 2 1, 3	3, 2 1, 4	4, 2 1, 4
77 (77)	78 (78)			
2, 2 4, 1	2, 2 3, 1			
3, 3 1, 4	4, 3 1, 4			

11.3 Complex Environments

With the emergence of increasingly complex MARL algorithms, a plethora of multi-agent environments have been developed to evaluate and study algorithms. In this section, we present a selection of such multi-agent environments which have seen significant adoption by the MARL research community.⁷⁶ Our selection includes individual environments as well as environment collections. For each environment, we describe the core properties in terms of state/action representation and observability, and the main learning challenges involved.

76. Many more environments exist which are not covered in this chapter. For additional environments, see: <https://agents.inf.ed.ac.uk/blog/multiagent-learning-environments>

Environment (Section)	Observability	Observations	Actions	Rewards
<i>Environments:</i>				
LBF (11.3.1)	full, part	dis	dis	spa
MPE (11.3.2)	full, part	con	dis, con	den
SMAC (11.3.3)	part	mix	dis	den
RWARE (11.3.4)	part	dis	dis	spa
GRF (11.3.5)	full, part	mix	dis	den, spa
Hanabi (11.3.6)	part	dis	dis	spa
<i>Environment collections:</i>				
Melting Pot (11.4.1)	part	con	dis	den, spa
OpenSpiel (11.4.2)	full, part	dis	dis	den, spa
Petting Zoo (11.4.3)	full, part	mix	dis, con	den, spa

Figure 11.1: List of multi-agent environments and environment collections, with core properties. **Observability** indicates full (full) or partial (part) observability. **Observations** indicate discrete (dis), continuous (con), or mixed (mix) observations and states. **Actions** indicates discrete (dis) or continuous (con) actions. **Rewards** indicates dense (den) or sparse (spa) rewards. Multiple values in a column indicate that the environment provides options for each value.

Figure 11.1 provides a summary of the environments along with their core properties. Download URLs can be found in the references for each environment, given in the respective sections below. Many environments use parameters to control the complexity or difficulty of the learning problem (such as by setting the number of agents, word size, number of items, etc.), and we use the term “task” to refer to a specific parameter setting of an environment.

11.3.1 Level-Based Foraging

Throughout this book, we have used a number of examples based on the level-based foraging (LBF) environment. LBF was first introduced in the work of Albrecht and Ramamoorthy (2013), and was adopted as a benchmark for multi-agent deep RL (e.g. Christianos, Schäfer, and Albrecht 2020; Papoudakis et al. 2021; Jafferjee et al. 2022; Yang et al. 2022). In LBF, n agents are placed in a fully observable grid-world environment and tasked with collecting items which are randomly located in the environment. Each agent and item has a numerical skill level. All agents have the same action space, $A = \{up, down, left, right, collect, noop\}$, to navigate in the environment, collect items, or do nothing (noop). A group of one or more agents can collect an item if: they are positioned adjacent to the item, they all select the *collect*

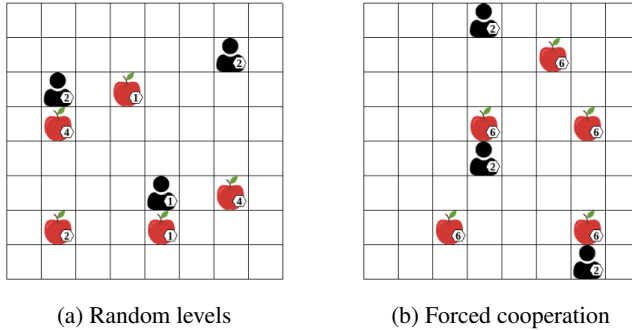


Figure 11.2: Illustrations of two LBF tasks in an 8×8 grid-world with 3 agents, 5 items, and either (a) random levels or (b) forced cooperation. Tasks with forced cooperation assign item level, such that each item requires all agents to cooperate to collect the item.

action, and the sum of the agents' levels is equal or greater than the item's level. Thus, some items require cooperation between (subsets of) agents. Agents receive a normalised reward for the collection of an item which depends on the level of the item as well as the agents' contribution to the collection of the item. Specifically, agent i receives the following reward when collecting item f

$$r_i = \frac{l_f \cdot l_i}{\sum_{f' \in \mathcal{F}} l_{f'} \sum_{j \in I(f')} l_j} \quad (11.1)$$

where l_f is the level of item f , l_i is the level of agent i , \mathcal{F} is the set of all items, and $I(f)$ is the set of agents involved in collecting the item f . Hence, the rewards of agent i are normalised with respect to the level of all items which can be collected in the environment as well as the relative level of agent i compared to the level of all agents contributing to the collection of item f .

An LBF task is specified by the size of the grid-world, the number of agents and items, and their skill levels. Agents and items can be randomly assigned a level at the beginning of each episode and placed in a random location within the grid-world. See Figure 11.2 for an illustration of two LBF tasks.

The environment is flexible with tasks requiring varying degrees of cooperation, observability, and scale. Challenging exploration problems can be defined in LBF by “enforcing cooperation”. In tasks with forced cooperation, item levels are assigned such that all agents in the environment need to cooperate to collect any of the items in the grid-world. See Figure 11.2b for an example environment with forced cooperation. It is also worth noting that many LBF tasks have mixed objectives across agents, with agents competing to receive

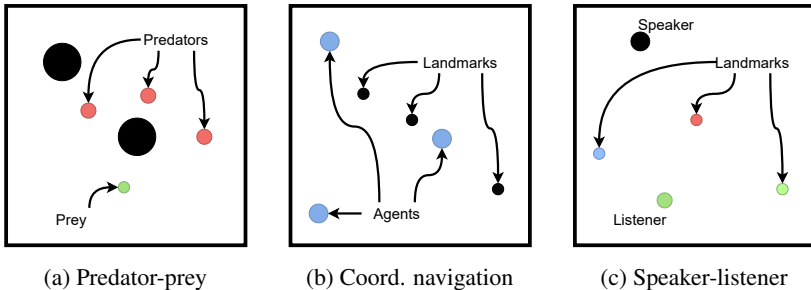


Figure 11.3: Three MPE tasks. (a) Predator-prey: a set of predator agents must catch a faster prey agent and avoid obstacles (large black circles). (b) Coordinated navigation: three agents need to scatter to cover three landmarks in the environment while avoiding collisions with each other. (c) Speaker-listener: a “listener” agent is randomly assigned one of three landmarks and needs to navigate to its assigned landmark (highlighted with the same colour). The listener does not observe its own colour and relies on a “speaker” agent which can see the listener’s colour and needs to learn to communicate the colour through binary communication actions.

individual reward for items they can collect by themselves and needing to cooperate with other agents to collect items of higher levels. Such conflicting objectives make for interesting multi-agent environments.

11.3.2 Multi-Agent Particle Environment

The multi-agent particle environment (MPE) contains several two-dimensional navigations tasks focusing on agent coordination. The environment includes competitive, cooperative, and common-reward tasks with full and partial observability. Agents observe high-level features, such as their velocity and relative positions to landmarks and other agents in the environment. Agents can either choose between discrete actions corresponding to movement in each cardinal direction, or use continuous actions for the velocity to move in any direction. Tasks include the common predator-prey problem in which a team of agents, the predators, must chase and reach an escaping prey; tasks in which agents need to communicate parts of their local observation to other agents; and more coordination problems.

Mordatch and Abbeel (2018) introduced the environment and Lowe et al. (2017) proposed an initial set of tasks commonly used for MPE. As the environment is extendable, more variations and tasks have been proposed (e.g. Iqbal and Sha (2019)). Three common MPE tasks are shown in Figure 11.3.



(a) Symmetric task

(b) Asymmetric task

Figure 11.4: A SMAC task in which two teams fight against each other. One team is controlled by the agents (one agent per unit) while the other team is controlled by a built-in AI. (a) A symmetric task in which each team consists of the same number and type of units (3 “marines” on each team). (b) An asymmetric task in which the teams consist of different unit types.

To further extend MPE, Bettini et al. (2022) propose the vectorized multi-agent simulator (VMAS). It also supports 2D multi-agent tasks with continuous or discrete action spaces. In contrast to the original MPE environment, VMAS can directly be simulated on GPUs to speed up training, offers more tasks, and supports multiple interfaces for compatibility across multiple RL frameworks.

11.3.3 StarCraft Multi-Agent Challenge

The StarCraft Multi-Agent Challenge (SMAC) environment (Samvelyan et al. 2019) contains common-reward tasks based on the real-time strategy game StarCraft II. In SMAC tasks, a team of agents controls units (one per agent) which combat against a team of units controlled by a fixed built-in AI. Tasks vary in the number and types of units, and maps in which the combat scenarios take place. Depending on the terrain, area and units, micromanagement strategies such as “kiting” are required to coordinate the team and successfully defeat the opponent team. SMAC contains symmetric tasks in which both teams consist of the same units (Figure 11.4a), and asymmetric tasks in which the teams have different compositions (Figure 11.4b). All SMAC tasks are partially observable, with agents only observing information about their own and nearby units up to a certain radius. Agents have actions to move within the map and to attack opponent units, and receive a dense common reward based on damage dealt and enemy units defeated, as well as a large bonus reward for winning the combat scenario.

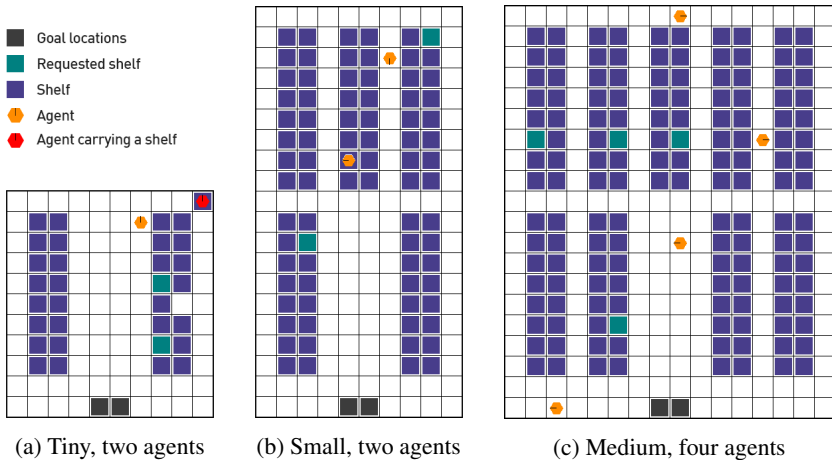


Figure 11.5: Three RWARE tasks with varying sizes and number of agents (from Christianos, Schäfer, and Albrecht (2020)).

The main challenge of SMAC lies in the common rewards across all agents. The credit assignment problem (Section 5.4.2) is particularly prominent in this setting because the actions of agents can have long-term consequences (such as destroying an opponent unit early on in an episode), and the common reward makes it difficult to disentangle each agent’s contribution to the achieved returns. This makes value decomposition methods, introduced in Section 9.5, particularly suitable for SMAC tasks.

11.3.4 Multi-Robot Warehouse

In the multi-robot warehouse (RWARE) environment (Christianos, Schäfer, and Albrecht 2020; Papoudakis et al. 2021), agents control robots navigating a grid-world warehouse and need to find, collect and deliver shelves with requested items. Tasks vary in the layout of the warehouse and number of agents (Figure 11.5). Agents observe information about shelves and agents in their close proximity, defined by an observation range that can be specified as part of the task. Agents select actions to rotate to the left or right, move forward, stay, or pick-up/drop-off load in their current location if possible. Agents can move underneath shelves as long as they carry no shelves. Agents only receive individual positive rewards for successfully delivering shelves with requested items to the goal locations. Upon each delivery, a new shelf without requested items in the warehouse is randomly sampled and then requested.

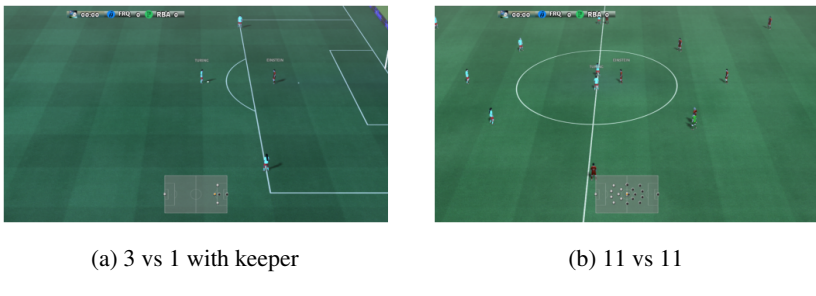


Figure 11.6: Two GRF environment scenarios.

The main challenge in RWARE tasks lies in its very sparse rewards. Delivering shelves with requested items requires agents to execute very specific, long sequences of actions to receive any non-zero reward. This makes the sample-efficient approaches of sharing parameters and experiences, introduced in Section 9.6, particularly well suited in this environments as demonstrated by Christiansos, Schäfer, and Albrecht (2020).

11.3.5 Google Research Football

The Google Research Football (GRF) environment (Kurach et al. 2020) provides a visually complex, physics-based 3D simulation of the game of football (Figure 11.6). The environment supports a multi-agent interface with either two agents controlling each of the teams, or agents controlling individual players as part of a cooperating team competing against a team controlled by a fixed built-in AI. The environment includes the full game of 11 vs. 11 football as well as a set of reference tasks with progressively harder scenarios to evaluate specific situations, such as defending and scoring with a smaller number of players. Agents can choose between 16 discrete actions, including eight movement actions, variations of passes, shots, dribbling, sprinting, and defensive actions. There are two reward functions which can be chosen: (1) providing a reward of +1 and -1 for scoring and conceding a goal, respectively, and (2) additionally providing positive rewards for maintaining the ball and moving forward towards the opponent’s goal region. Similar to reward functions, there are three modes of observations including (1) a pixel-based observation showing a rendered screen with a view of the pitch following the ball, a small global map as well as a scoreboard, (2) a significantly smaller compressed global map showing information about both teams, ball location, and highlighting the location of the active player, and (3) a feature vector of 115 values encoding player locations, ball possession, direction, game mode, active player, and more.

The different modes of rewards, observations, and progressively harder scenarios provided by the environment make it a suitable benchmark to evaluate complex multi-agent interactions for cooperative games. The environment can also be used for two-player competitive games, with each agent controlling one team, to investigate competitive self-play algorithms.

11.3.6 Hanabi

Hanabi is a cooperative turn-based card game for two to five players in which each player (agent) holds a set of cards with numerical rank (1 to 5) and colours (red, green, blue, yellow, white) drawn from a set of 50 cards in total. Players need to build up ordered stacks of cards, with each stack containing cards of the same colour with increasing rank. The twist of Hanabi is that each player does not see their own card but only sees the cards of all other players. Players take actions in turns with the acting player having three possible actions: giving a hint, playing a card from their hand, or discarding a card from their hand. When giving a hint, the acting player selects another player and is allowed to point out all cards of that player's hand which match a chosen rank or colour. Each given hint consumes an information token, with the entire team of players starting the game with eight information tokens. Once all information tokens have been used, players can no longer give hints. Information tokens can be recovered by discarding a card (which also makes the active player draw a new card from the deck) and by completing a stack of a colour by placing the fifth and final card on it. When a card is played, it is either successfully continuing one of the five stacks or is unsuccessful, in which case the team loses a life. The game ends when the team loses all their three lives, all five stacks were successfully completed with cards of all five ranks in increasing order, or when the last card of the deck has been drawn and each player took a final turn. The team receives a reward for each successfully placed card, for final scores between 0 and 25.

Based on the partial observability of Hanabi, in which agents lack significant information to guide their action selection, Bard et al. (2020) proposed Hanabi as a challenging multi-agent environment with agents needing to adopt conventions to successfully use their limited communication to cooperate. Agents need to establish implicit communication beyond the limited resource of hints to succeed. These properties make Hanabi an interesting challenge for cooperative self-play algorithms, ad hoc teamwork (Mirsky et al. 2022), and communicating and acting under imperfect information.

11.4 Environment Collections

The environment collections in this section include many different environments, each representing a different game which can differ in properties such as state/action representation and dynamics, full/partial observability and type of observation, and reward density. Hence, the focus of these environment collections is to provide a unified representation and agent-environment interface for the games, such that a MARL algorithm which is compatible with the interface can be trained in each of its environments. These environment collections typically also provide additional functionality, such as tools for analysis and the creation of new environments, and even implementations of MARL algorithms.

While some of the previously introduced environments also include multiple tasks, these tasks usually use the same state-action dynamics and observation specification. In contrast, the environments contained in the below collections define a set of diverse games.

11.4.1 Melting Pot

Melting Pot (Leibo et al. 2021) is a collection of over 50 different multi-agent tasks based on DeepMind Lab2D (Beattie et al. 2020) (see Figure 11.7 for some examples). It focuses on two aspects of generalisation for MARL: (1) generalisation across different tasks and (2) generalisation across different co-players. The first aspect is achieved by providing a diverse set of tasks, including tasks with different numbers of agents, different objectives, and different dynamics. The second aspect is achieved by providing a set of diverse populations of pre-trained agent policies for each task. During training, a so-called focal population of agents is trained in a task. During evaluation, the focal population is evaluated in the same task but with varying co-players: A set of agents is sampled from the trained focal population, and some so-called background agents are controlled by pre-trained policies. Therefore, Melting Pot evaluates the ability of trained focal agents to be able to zero-shot generalise to the diverse behaviours of the background agents in the task.

Melting pot tasks vary in their number of agents, their objectives ranging from zero-sum competitive, fully cooperative common-reward, and mixed-objective games. Tasks are partially observable with agents observing a partial 88×88 RGB image of the environment. The action space is discrete with agents having six movement actions in all tasks: move forward, backward, strafe left or right, turn left or right, and potentially additional actions depending on the task.

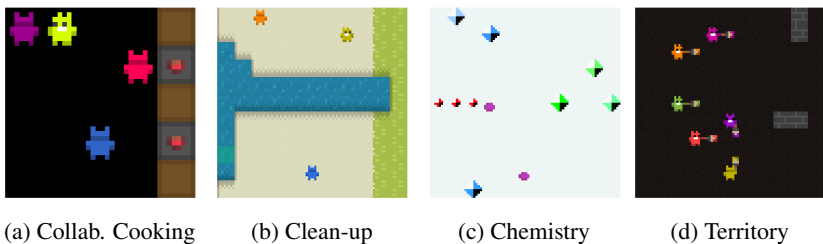


Figure 11.7: Four Melting Pot environments. (a) Collaborative Cooking: Agents need to cook a meal together. There exist several versions of the task which vary in the layout of the kitchen and, thus, required cooperation and specialisation of agents. (b) Clean-up: Seven agents can collect apples in the environment to gain reward. The rate of apples spawning is dependent on agents cleaning the nearby river which leads to a social dilemma with competition between agents for apples, and the need to clean-up for long-term rewards. (c) Chemistry: Agents can carry molecules in the environment. When two molecules are brought close to each other, they may react to synthesise new molecules and generate reward according to a task-specific reaction graph. (d) Territory: Agents need to capture resources and eliminate opponent agents by shooting a beam.

11.4.2 OpenSpiel

OpenSpiel⁷⁷ (Lanctot et al. 2019) is a collection of environments and MARL algorithms, as well as other planning/search algorithms (such as MCTS, see Section 9.7.1), with a focus on turn-based games, also known as “extensive-form” games. A large diversity of classical turn-based games are provided in OpenSpiel, which includes games such as Backgammon, Bridge, Chess, Go, Poker, Hanabi (Section 11.3.6), and many more. The agent-environment interface used in OpenSpiel is designed with a focus on turn-based games, though it also supports simultaneous-move games such as the game models used in this book. The environments in OpenSpiel use a mix of full or partial observability, and all environments specify discrete actions, observations, and states. A challenge in many of the games is that long interaction sequences are often required by the agents before they receive any rewards.

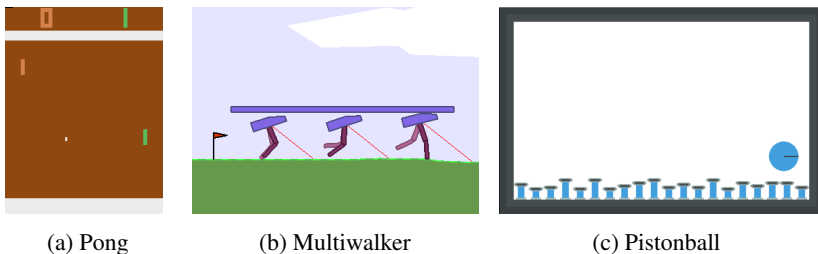


Figure 11.8: Three Petting Zoo environments. (a) Pong: Two agents play against each other in the classic Atari Pong game. (b) Multiwalker: Three agents control bipedal robots and need to learn to walk together without dropping the package placed on their heads. (c) Pistonball: Agents control pistons on the ground and move the ball from the right side of the screen to the left.

11.4.3 Petting Zoo

Petting Zoo (Terry et al. 2020) is a library for MARL research which contains a large number of multi-agent environments, including multi-agent games based on the Atari Learning Environment (Bellemare et al. 2013), various classic games such as Connect Four, Go, and Texas Holdem, and continuous control tasks (see Figure 11.8 for some examples). Petting Zoo also integrates the multi-agent particle environment (Section 11.3.2). Petting Zoo covers a wide range of learning problems by including environments with full and partial observability, discrete and continuous actions, and dense and sparse rewards. Besides providing a large set of tasks, Petting Zoo unifies the interface across all its tasks, offers additional tools to customise the environment interface, and integrates training with various MARL frameworks.

⁷⁷. Spiel is the German word for game, pronounced “shpeel”.

A Surveys on Multi-Agent Reinforcement Learning

At the time of writing this book, MARL is a highly active and fast-moving field of research, as evidenced by the many survey articles which have been published in this area in recent years. This book focuses on the foundations of MARL and does not include many algorithms which are at the forefront of MARL research. To complement the book, this chapter provides a list of survey articles of MARL research in reverse chronological order, dating back to the first survey of the field (to our knowledge) published in 1999. Additional survey articles about particular application domains of MARL exist but have been omitted here.

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K. Zhang, Z. Yang, T. Basar (2021). Multi-Agent Reinforcement Learning: A Selective Overview of Theories and Algorithms. In: *Handbook of Reinforcement Learning and Control*.

S. Gronauer, K. Diepold (2021). Multi-Agent Deep Reinforcement Learning: A Survey. In: *Artificial Intelligence Review*.

C. Moulin-Frier, P. Oudeyer (2020). Multi-Agent Reinforcement Learning as a Computational Tool for Language Evolution Research: Historical Context and Future Challenges. In: *arXiv:2002.08878*.

M. Zaïem, E. Bennequin (2019). Learning to Communicate in Multi-Agent Reinforcement Learning: A Review. In: *arXiv:1911.05438*.

P. Hernandez-Leal, B. Kartal, M.E. Taylor (2019). A Survey and Critique of Multiagent Deep Reinforcement Learning. In: *Autonomous Agents and*

Multi-Agent Systems, vol. 33, no. 6, pp. 1-48.

R. Rădulescu, P.K. Mannion, D.M. Roijers, A. Nowé (2019). Multi-Objective Multi-Agent Decision Making: A Utility-based Analysis and Survey. In: *arXiv:1909.02964*.

G. Papoudakis, F. Christianos, A. Rahman, S.V. Albrecht (2019). Dealing with Non-Stationarity in Multi-Agent Deep Reinforcement Learning. In: *arXiv:1906.04737*.

F. Silva, A. Costa (2019). A Survey on Transfer Learning for Multiagent Reinforcement Learning Systems. In: *Journal of Artificial Intelligence Research*, vol. 64, pp. 645-703.

T.T. Nguyen, N.D. Nguyen, S. Nahavandi (2019). Deep Reinforcement Learning for Multi-Agent Systems: A Review of Challenges, Solutions and Applications. In: *arXiv:1812.11794*.

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