DE-NOVO DETERMINATION OF PROTEIN TERTIARY STRUCTURE IN A HHPNX 3D -FACE CENTERED CUBIC LATTICE WITH MEAN FIELD MULTI-AGENT REINFORCEMENT LEARNING

by

ADRIAN COUTSOFTIDES URN: 6481554

A dissertation submitted in partial fulfilment of the requirements for the award of

BACHELOR OF SCIENCE IN COMPUTER SCIENCE

May 2020

Department of Computing University of Surrey Guildford GU2 7XH

Supervised by: Sotiris Moschoyanis

I declare that this dissertation is my own work and that the work of others is acknowledged and
indicated by explicit references. Adrian Coutsoftides
May 2020



Abstract

The protein folding problem is the search for a function that maps a proteins primary structure, composed of a string of discrete amino acid residues to their respective native conformation in 3D space denoted as the protein's tertiary structure. Recent breakthroughs in the field have utilize techniques such as multiple sequence alignment coupled with residual convolutional neural networks to derive candidate posteriors over the distribution of inter-residue distances from which multiple energetically favourable tertiary structures can be generated; these results are typically annealed to produce a structure of lowest conformational energy. In this work I propose an alternative deep reinforcement learning system that does not rely on the pre-existing data of other sequences and instead approximates the pair-wise local interactions of the residues on a 3D Face-Centered Cubic Lattice model. I show that mean-field approximations effectively model the the free energy landscape of the system with respect to the expectation of the distribution of rewards in the local neighbourhood for each agent. I propose that the architecture of this system effectively incorporates inductive bias into the problem formulation and thus provides a richer training signal. Additional techniques are also employed in combination to reduce the sample complexity of the search space.

Acknowledgements

Write any personal words of thanks here. Typically, this space is used to thank your supervisor for their guidance, as well as anyone else who has supported the completion of this dissertation, for example by discussing results and their interpretation or reviewing write ups. It is also usual to acknowledge any financial support received in relation to this work.

Contents

1	Intr	oducti	ion											-	12
	1.1	Proble	em Backgr	ound			 	 		 		 			12
	1.2	Projec	et Aims ar	nd Object	ives		 	 	٠	 	 	 			13
	1.3	Succes	ss Criteria				 	 	•	 	 	 			14
	1.4	Struct	ure of Re	port			 	 	•	 		 			14
2	$\operatorname{Lit}_{\epsilon}$	erature	Review											-	16
	2.1	Protei	ns				 	 	•	 		 	•		16
		2.1.1	Amino A	Acids & P	oly-Pept	ides .	 	 	•	 	 	 			16
		2.1.2	Protein S	Structure	s		 	 		 	 	 			17
		2.1.3	The Pro	tein's Ene	ergy Lan	dscape		 		 	 	 			19
		2.1.4	Bioinform	matics .			 	 		 	 	 			22
			2.1.4.1	Homolog	gy Model	lling .	 	 	•	 	 	 	•	 •	22
			2.1.4.2	Lattice I	Models .		 	 	•	 	 	 			23
			2.1.4.3	Bravais	Lattices		 	 		 	 	 			23
			2.1.4.4	HP Mod	ıel		 	 		 	 	 			25
			2.1.4.5	hHPNX	Model .		 	 		 	 	 			27
	2.2	Reinfo	orcement I	Learning			 	 	•	 		 			28
		2.2.1	Markov	Decision 1	Processe	s	 	 		 	 	 			28

		2.2.1.1	Bellman Equation	32
		2.2.1.2	Optimality	32
		2.2.1.3	Generalised Policy Iteration	33
		2.2.1.4	Exploration vs Exploitation	34
		2.2.1.5	Monte Carlo Estimation	34
		2.2.1.6	Model Free vs Model Based	36
	2.2.2	Deep Q	Learning	37
		2.2.2.1	Temporal Difference Error	37
		2.2.2.2	Q Learning	37
		2.2.2.3	Neural Networks	38
		2.2.2.4	Deep Q-Networks	40
	2.2.3	Improve	ements to Vanilla Deep Q-Networks	43
		2.2.3.1	Memory	43
		2.2.3.2	Exploration	43
		2.2.3.3	Exploitation	43
		2.2.3.4	Rainbow DQN	43
2.3	Multi-	Agent Re	einforcement Learning	43
	2.3.1	Stochast	tic Games	43
		2.3.1.1	Game Theory	43
		2.3.1.2	Expected Pay-offs as Expected Rewards	43
		2.3.1.3	Multi-Agent Games	43
		2.3.1.4	Mean Field Games	43
		2.3.1.5	Mean Field multi-agent reinforcement learning	43
2.4	Relate	ed Work		43
	2.4.1	MCMC	methods for lattice models	43

2.4.2	Deep Learning methods for lattice models							
	2.4.2.1 Alpha-Fold	43						
2.4.3	Reinforcement Learning methods for lattice models	43						

List of Figures

2.1	Peptide Bond	17
2.2	Structure of a peptide unit	17
2.3	Ramachandran plot	18
2.4	Secondary and tertiary structures	19
2.5	Gibbs free energy funnel	21
2.6	3D Cubic Bravais Lattices	24
2.7	Markov chain with state space and transition propabilities	29

List of Tables

2.1	 	 •		 •								•		•					16
2.2	 								•			•					•		26
2.3	 																		 27

Glossary

 ΔG Change in Gibbs free energy, measure as the amount of energy in the system than can be turned into work ΔH Change in enthalpy as the change in total heat content of the system ΔS Change in entropy as the measure of disorder in a system T Temperature in Kelvin

 ε Interaction potential of a bond

Abbreviations

MDP Markov Decision Process

DQN Deep-Q-Learning

PSP Protein Structure Prediction

PDB Protein Data Bank

CASP Critical Assessment of Structure Prediction

KL Kullback Liebler

RL Reinforcement Learning

MSA Multiple Sequence Alignment

Chapter 1

Introduction

1.1 Problem Background

Proteins are biological molecules that carry out specific functions within the body, they are assembled out of chemical bonds between smaller sub units called amino acid residues to form a poly-peptide chain. Proteins carry out their functions by conforming into specific shapes and fitting into substrates, this is referred to as the "lock and key" model of proteins. A protein's three-dimensional structure has been proven [reference] to been entirely determined from the ordering of a discrete set of 22 residues in a sequence of arbitrary length. The ability to infer a protein's precise three dimensional structure directly from the sequence of residues would unlock the potential for designer drugs that carry out specific actions within the body; it would also open a path for the treatment of illnesses that arise from malformed proteins such as huntington's disease [reference]. This has however been an open problem, as the search space of possible conformations for a given protein is vast and subject to numerous local minima. Traditional methods of protein structure determination such as X-Ray Crystallography [reference] are extremely costly² and time consuming, sometimes taking up to four years to determine the structure of a protein. This has given rise to multiple computational approaches reference as means to drive down cost and increase the productivity of researchers. Naturally, many approaches using deep learning have been proposed to solve the problem, recently a breakthrough by DeepMind [reference] propelled them to success at the bi-annual Critical Assessment of Structure Prediction (CASP) competition. Though their results were state of

¹Within 1Å

²In the order of millions ^{reference}

the art, their methods still relied on building a predictive model of the properties of available proteins in the Protein Data Bank (PDB)^[reference]; as opposed to inferring the tertiary structure from the primary sequence alone.

1.2 Project Aims and Objectives

Over the course of this dissertation I aim to explore and contrast modern deep learning approaches for approximating the native conformations of proteins on a discrete lattice structure. I will then go on to propose a novel algorithm based on mean-field approximations that addresses some of the drawbacks and biases inherent in the approaches I have reviewed. The following is a list of the project's aims overall:

- 1. Provide a succinct introduction to the molecular mechanics that govern the conformations of proteins
- 2. Introduce and compare different approaches to modelling the problem computationally
- 3. Provide an extended analysis of lattice models for proteins and their ability to encode correct conformations
- 4. Introduce reinforcement learning and progress to modern Deep Q-Learning
- 5. Building on Deep Q-Learning, introduce improvements to the algorithm since it's inception
- 6. Compare and contrast deep learning approaches to the lattice model with an emphasis on reinforcement learning methods
- 7. Introduce multi-agent learning within the framework of stochastic games
- 8. Introduce mean-field approximations in particular reference to mean field games
- 9. Formulate the conformation of residues on a lattice as a cooperative game of incomplete information
- 10. Provide a novel approach to de-novo structure determination using mean-field multi-agent learning
- 11. Benchmark my approach against the results of other groups on the same proteins to evaluate the effectiveness of the novel algorithm

1.3 Success Criteria

In order to evaluate the utility of both my findings and subsequent algorithm, I have defined the following high-level requirements that must be satisfied to mark this project as a success.

- 1. Provide comprehensive overview of the underlying problem of protein folding
- 2. Describe markov decision processes (MDPs) and its ties to reinforcement learning
- 3. Highlight drawbacks to the default DQN algorithms and notable improvements to address those
- 4. Demonstrate multi-agent learning as a generalisation of single MDPs into markov games
- 5. Successfully benchmark my multi-agent approach against similar lattice-based approaches to protein folding

1.4 Structure of Report

1. Introduction

In this section I provided an overview the protein folding problem and introduced the components that I will be synthesising into novel approach.

2. Literature Review

In this section I will provide an overview as-well as an introduction to select topics and that comprise the necessary background knowledge. This section ends with a comparative analysis of related work on the problem.

3. System Requirements and Specification

In this section I will analyse the drawbacks of methods utilized in related work, and from this evaluation derive the requirements of a systems required to address these limitations.

4. System Design

This section seeks to unify the selected systems and concepts into an integrated learning algorithm that coherently reflects the underlying problem's structure.

5. Testing & Validation

In order to verify the efficacy of the learning agents, a set of lattice structure of known

proteins will be calculated and evaluated against the results for the same set of proteins in related work.

6. Discussions

Limitations of my implementation are discussed here and possible solutions and research directions are proposed.

7. Conclusion

Here I will present my concluding thoughts and results of my final analyses of the architecture.

Chapter 2

Literature Review

2.1 Proteins

2.1.1 Amino Acids & Poly-Peptides

Proteins are the mechanism by which executive function takes place in the cell. This is includes the implementation of activities such as "metabolism, growth, architecture and regulation of cell and organism" -(Lesk 2018). Proteins themselves are composed of discrete molecular units termed amino acids, the precise way in which they amino acids arrange themselves is dependent of the genetic sequence of the parent organism itself. There exist 20 unique amino acids (examples given below) that can be combined that can be sequenced into strings of arbitrary length of any order following chemical laws. These molecules come together to form peptide

Table 2.1

Examples of proteins and their codes										
Glycine	G									
Alanine	A									
Serine	S									
Cysteine	C									
Threorine	Т									
Proline	Р									
Valine	V									
Leucine	K									

Figure 2.1: Peptide Bond

†Image Source: https://en.wikipedia.org/wiki/Peptide_bond

bonds, this occurs when the carboxyl group (^-COOH) bonds to the amino group ($^+NH_3$) of another amino acid producing water as a by-product of the reaction (Lesk 2018). Despite all amino acids possessing an amino and carboxyl group, every amino acid has a unique side chain, an additional molecular group attached to the main chain. The unique chemical properties of these side chains are responsible for the interactions in that molecule's neighbourhood; a term that is expanded on in the next few sections.

2.1.2 Protein Structures

Figure 2.2: Structure of a peptide unit

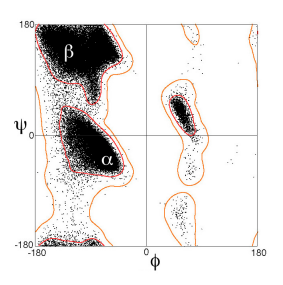
$$\begin{array}{c|c} \mathbf{C}_{\beta_{\mathbf{i}}} \\ \mathbf{O} & & \\ & \mathbf{C}_{\alpha_{\mathbf{i}}} \\ \mathbf{C}_{\beta_{\mathbf{i}-1}} - \mathbf{N}_{\mathbf{i}} & \mathbf{C}_{\beta_{\mathbf{i}+1}} \\ & & \\ & \mathbf{O} \end{array}$$

The carbon atom that joins the side chain to the main chain is denoted as the α carbon atom and the carbon atoms in immediate contact with the α carbons are known as β carbons¹. The bond angles between the $C_{\alpha_i} - C_{\beta_i+1}$ and $C_{\alpha_i} - N_i$ groups are commonly denoted as ψ and ϕ respectively. Stereochemically feasible angles can be described by a Ramachandran plot as in figure 2.3.

Poly-peptide conformations can be described by three structures:

 $^{^{1}}$ In figure 2.2 subscript *i* denotes membership to a unique unit

Figure 2.3: Ramachandran plot



†Source: https://proteopedia.org/wiki/index.php/Ramachandran_Plots

1. Primary Structure

This is the encoding of a protein as a 1-D sequence of its constituent amino acids.

I.e:

DTYGYWEPYT

2. Secondary Structure

This encoding represents local, repeating structures with respect to the entire conformations. These structures satisfy chemical restraints common to most proteins². In particular, the formation of structures known as α^3 helices and β^4 sheets largely facilitate the energetic and conformational constraints imposed by the interactions amongst the side-chains orthogonal to the backbone. These structures form the dense regions in the Ramachandran plot.

3. Tertiary Structure

A protein's tertiary structure are its 3D atomic coordinates in space. The protein's torsion backbone (main-chain) traces out a curve through space parameterised by all pairs (ϕ, ψ) for each residue pair. Hydrogen bonds, which arise when neighbouring residues are in close proximity although not directly connected by the backbone, hold together the various

²The hydrogen bonding potential of main-chain N-H and C=O groups (Lesk 2018)

 $^{^3}B$ in fig 2.4

 $^{^4}E$ in fig 2.4

secondary structures in specific conformations such that the global energy of the system is minimized (Yang, Ji & Liu 2013).

Figure 2.4: Secondary and tertiary structures

†Source: (Boyle 2018)

The primary goal of protein structure prediction is the *ab initio* determination of a protein's tertiary structure from its primary structure (Yang et al. 2013) which can be described as a mapping:

$$\mathbf{X}^n := \text{Set of all sequences of length } n$$
 (2.1)

$$\mathbf{F}: \mathbf{X}^n \to ((\phi, \psi)_1 \dots (\phi, \psi)_n) \tag{2.2}$$

(Anfinsen 1972) et al showed that the protein's tertiary structure is entirely encoded by its primary structure; the implications of this are explored next.

2.1.3 The Protein's Energy Landscape

Due in part to the continuous spectrum of stereochemically plausible values of (ϕ, ψ) , the possible number of conformations for a given sequence are extraordinarily large. Given that each possible conformation has an associated free energy, Levinthal's paradox states that if all conformations were equally energetically favourable, then protein would essentially have to undergo a random walk along the Gibbs free energy surface until it has found its native conformation. The free energy surface in this respect refers to the manifold that is formed by taking the distribution over conformations $((\phi, \psi)_1 \dots (\phi, \psi)_n)$ at every time-step and ΔG at every point to form a surface in 2n + 1 dimensional space. Given that common proteins have in the order of 100-1000+

constituent residues, the combinatorial size of the state space would prohibit any efforts to find a specific conformation by random walk with vanishing probability.

(Yang et al. 2013) address both the paradox and subsequent critiscisms; part of their work can be summarised by the following lemmas:

Lemma 1. Not all conformations are equally energetically favourable.

Proof. Proteins fold spontaneously in the order of nanoseconds into their native conformation in a solvent at constant temperature, thus they cannot be traversing the whole state space. \Box

Lemma 2. There must therefore be a driving reaction that narrows down the search space.

Proof. Folding appears to undergo two consecutive stages, tier 1 occurs of a timescale of nanosec-onds and tier 2 appears to occur over a scale of picoseconds, a reduction in 3 orders of magnitude. Thus the peptide must undergo a slower interaction that constrains the state space followed by a final, faster "relaxation" into the native state.

Lemma 3. By undergoing hydrophobic collapse, the remaining residues have restricted degrees of freedom, thus a reduced subspace consisting of only energetically feasible conformations is explored, within this subspace there exists one unique solution whose energy is minimized.

Proof. The relative timescales of the tier 1 and tier 2 interactions indicate a smooth slope in conformational space that greatly constrains the possible conformations, followed by a second tier of faster interactions that enables the system to quickly overcome the local maxima and minima which gives way to the global optimum at the bottom of the subset. \Box

Lemma 4. If there exists only one unique conformation whose Gibbs free energy is minimal amongst all possible conformations, the manifold must be "funnel shaped".

Proof. The nature of the hydrophobic collapse "drops" the full state space into a smaller subspace, the unique global minimum lies within this subspace. If there existed more than one unique solution, then a given sequence could form into multiple possible native conformations, this would violate the marginal stability property of the native state so there cannot exist more than one unique solution.

$$\Delta G = \Delta H - T\Delta S$$

By the second law of thermodynamics, the entropy (amount of disorder) in a closed system can never decrease over time and can remain constant only if the system is in equilibrium, thus a state of maximum entropy (Jaffe 2018). The work of (Yue & Dill 1993, Yang et al. 2013) show that the overall increase in entropy of the peptide-solvent system is primarily motivated by a phenomenon known as the *hydrophobic collapse* whereby all the water produced as by-product of the peptide bond ⁵ "squeezes" the hydrophobic residues into globular core. Following **Lemma 4**, the process appears to be guided by a heuristic search in this subspace, enabling it to swiftly reach its native state.

Path
Path
Path
Hydrophobically
collapsed
molten globule

Global free energy minimum

Figure 2.5: Gibbs free energy funnel

†Source: (Yang et al. 2013)

Critically, the notion of ΔG is with respect to the distribution over conformations, indicating that at any point the protein exists as an ensemble of possible conformations, and the native state consists of a collection of conformations whose joint distribution minimises the value of ΔG . I reformulate this property as maximum likelihood objective:

Theorem 1.

- 1. Let $T \in {\{\mathbf{X}\}}^n$ denote the native state as a tuple of n torsion angles (follows from eq. 2.1).
- 2. Let $P(\mathbf{F}(\mathbf{X}^n; \theta))$ denote the distribution over all functional mappings from a primary sequence to a tuple of n torsion angles with parameters θ (follows from eq. 2.2).

The search for a function that maps the protein's primary structure to its tertiary structure can be formulated as the maximization of the posterior probability that function \mathbf{F} with parameters θ

 $^{^5}$ See fig 2.1

maps sequence \mathbf{X}^n to state T.

$$P(T|\mathbf{F}(\mathbf{X}^n;\theta)) = \frac{P(\mathbf{F}(\mathbf{X}^n;\theta)|T) \cdot P(T)}{P(\mathbf{F}(\mathbf{X}^n;\theta))}$$
(2.3)

This formulation is further explored in the **System Requirements and Specification** section, however an interesting detail to note is that this is equivalent to sampling the maximum likelihood estimate from the posterior of a *Gaussian process* ⁶.

2.1.4 Bioinformatics

There exists experimental techniques to determine the protein's native (or *crystalline*) structure⁷, however these approaches are out of the scope of this project and are not given a detailed analysis. Otherwise, these techniques can often require years of trial and error and can be very costly (Alberts 2002).

In recent years, advancements in both hardware and processing power have enabled computational methods to play increasingly significant role in the PSP problem. I will focus briefly what's known as homology modelling as this is relevant to understanding related supervised learning methods, and then I will then provide extended review of another class of models used for *ab initio* structure prediction, lattice models.

2.1.4.1 Homology Modelling

All proteins are members of an evolutionary tree and so related proteins tend to demonstrate high structural similarity with their relatives. Although older studies suggest that proteins are in fact random strings only slightly edited by evolution (Weiss, Jiménez-Montaño & Herzel 2000); more recent studies have shown that this is not the case, as certain methods were able to distinguish between random strings and real proteins with an accuracy of up to 88 – 94.36% (Lucrezia, Slanzi, Poli, Polticelli & Minervini 2012, Tsygvintsev 2019). Given the apparent specificity of the sequences, Homology modelling, focuses on trying to predict the tertiary structure of an unknown protein from the structures of closely related proteins (homologues). This involves measuring Hamming distances⁸ between closely related proteins and deriving an *inverse scoring* matrix with each pair-wise comparison. When this is done for more than one possible pairing,

⁶See appendix A.1

⁷Such as X-Ray Crystallography (Chayen 2005)

⁸See appendix A.2

this is known as Multiple Sequence Alignment (MSA). Highly correlated proteins with relatively low MSA scores tend to exhibit very similar tertiary structures (Lesk 2018); this property is used as the basis for many machine learning methods which is explored in **Related Work**.

2.1.4.2 Lattice Models

Although approaches that utilise homology modelling in some form have been very successful in recent years⁹, they rely on the pre-determined structures of other proteins, this poses a barrier to *de novo* protein design and research into undetermined structures for which we do not have a readily available dataset of relatives.

An alternative model of computation was proposed by (Yue & Dill 1993) that seeks to address the combinatorial conformational space. Instead, they proposed that the space should be discretised to restrict the possible number of conformations a given sequence can take on, a *Bravais* lattice was the perfect tool for this.

2.1.4.3 Bravais Lattices

A Bravais lattice is a simple mathematical description of a lattice structure, whereby each point in space is some linear combination of unit vectors with integer multiples (Kittel 2005).

$$\mathbf{r}, \mathbf{a} \in \mathbb{R}^n, u \in \mathbb{Z}$$

$$\mathbf{r} = u_1 \mathbf{a}_1 + u_2 \mathbf{a}_2 + u_3 \mathbf{a}_3$$
(2.4)

In matrix form:

$$\mathbf{r} \in \mathbb{R}^n, \mathbf{u} \in \mathbb{Z}^n$$

$$\mathbf{a} \in \mathbb{R}^n \times \mathbb{R}^n$$

$$\mathbf{r} = \mathbf{a} \cdot \mathbf{u}^\top$$
(2.5)

Different values for the unit vectors give rise to different classes of lattices, the number of neighbours for a given vertez is denoted as z. In the interest of brevity 3 lattices are discussed as they are pertinent to the related work and system design.

⁹See AlphaFold 2.4.2.1

1. 2D Square Lattice

A simple lattice whose unit vectors are two dimensional:

$$\mathbf{a} = \begin{bmatrix} 1 & 0 \\ 0 & 1 \end{bmatrix}$$

$$z = 4$$
(2.6)

2. 3D Primitive Cubic Lattice

A 3D lattice with unit vectors in each direction of space:

$$\mathbf{a} = \begin{bmatrix} 1 & 0 & 0 \\ 0 & 1 & 0 \\ 0 & 0 & 1 \end{bmatrix}$$

$$z = 6$$
(2.7)

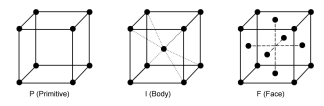
3. 3D Face-Centered Cubic Lattice

A 3D cubic lattice with additional vertex points at the center of each square face; notably, this lattice has the highest sphere packing density ¹⁰:

$$\mathbf{a} = \begin{bmatrix} 0 & 0.5 & 0.5 \\ 0.5 & 0 & 0.5 \\ 0.5 & 0.5 & 0 \end{bmatrix}$$

$$z = 12$$
(2.8)

Figure 2.6: 3D Cubic Bravais Lattices



†Source: https://biochem.co/2008/08/crystal-structure-studies/

For clarity, any point within the vector space defined by the lattice structure is an integer multiple of any linear combination of the unit vectors. Nomenclature on the subject dictates that a vertex on the lattice is referred to as a "site".

 $^{^{10}}$ Highest amount of rigid spheres that can be packed per unit space ≈ 0.74 (Hoque, Chetty & Sattar 2009)

2.1.4.4 HP Model

(Yue & Dill 1993) initially proposed the use of a 3D primitive cubic lattice to model the conformations of proteins. Each amino acid was divided into two categories according to their properties: Hydrophobic and Polar, and a conformation was formulated as a self avoiding walk (SAW) on the lattice.

Some definitions:

- A contact on the lattice is define as the adjacency of two immediate neighbours on the lattice not connected by the torsion backbone, if two residues are connected through the torsion backbone directly this is referred to as a bond.
- A segment is a run of monomers of a particular category (a)¹¹, a singlet occurs when a monomer is situated between two monomers of another category (b)¹².

$$\begin{array}{ccc} HPPPPH & PHP \\ ^{(a)} & ^{(b)} \end{array}$$

- $x \in \{H, P\}$
- n_x is the number of x monomers in the sequence. I.e. n_H is the number of H monomers in the entire sequence.
- b_{xx} is number of bonds between xx.
- t_{xx} is the number of bonds + contacts between xx.
- h_{term} is the number of segements that terminate in H monomers

These quantities are related by the following equations:

$$2b_{HH} + b_{HP} + h_{term} + (z - 2)n_H = 2(t_{HH} + b_{HH}) + b_{HP} + t_{HP} + t_{H-Solvent}$$
(2.9)

$$G = \frac{b_{HP} + h_{term}}{2}$$

$$S = b_{HP} + t_{HP} + t_{H-Solvent}$$
(2.10)

• G is the total number of H segments in a sequence

¹¹P segment of length 4

¹²H singlet

• S is the total surface area of the hydrophobic core, assuming that the monomers as unconnected. This acts as an upper bound of the surface area.

Substituting (2.10) into (2.9) produces the "folding equation":

$$t_{HH} + \frac{S}{2} = G + \frac{(z-2)n_H}{2} \tag{2.11}$$

Where the left side is dependent on the *conformation*, and the right side is a constant that solely depends on the lattice structure and given sequence. Motivated by the role of the hydrophobic collapse, the folding problem within this framework is a function that maximises t_{HH} ; this is equivalent to minimizing S. The logic follows that the most tightly packed core, that with the minimal surface area should also have the densest packing of H - H contacts.

(Dill, Bromberg, Yue, Chan, Ftebig, Yee & Thomas 2008, Lau & Dill 1989) show that an unguided exhaustive search through the coformational landscape is unlikely to find a single candidate native state but instead produce a set of states with high degeneracy; indicating that many conformations shared the same free energy value due to the numerous local minima. However by introducing a heuristic that guided the conformation through as hydrophobic collapse, a much smaller set of candidate conformations with much less degeneracy was produced. Genetic algorithms were developed to overcome the rugged landscape, their fitness function was determined by the favourability of any contact pair derived for experimental data. This resulted in the interaction potential (ε) matrix, which is a measure of that interaction's contribution to the total ΔG of the system (Hoque et al. 2009):

Table 2.2

	Н	Р
Н	-1	0
Р	0	0

When (Lau & Dill 1989) first proposed the model, only 2D lattices were explored at the time with exhaustive search techniques and the simulations were evaluated against *mean-field* approximations of the system. (Yue & Dill 1993) then introduced linear integer programming methods that conducted volumetric optimization such as those described by (2.10-2.11) on a 3D primitive cubic lattice. There are some key limitations imposed on the mean-field approximations which are explored in **Related Work** and **System Design and Specification**; subsequent

improvements to the original HP model were also introduced and are explored next.

2.1.4.5 hHPNX Model

(Hoque et al. 2009) showed that previously explored variants of the HP model on 2D and 3D lattices were still prone to large degeneracy in the solutions they produced. This was a result of the limitations imposed by having only two possible categories; stochastic methods end up traversing many redundant solutions for each non-native conformation visited in the search-space¹³. The original HP categories were further subdivided according the relative polarities of the side chains:

- $P \mapsto \{P, N, X\}$
- $H \mapsto \{h, H\}$

They then correcte errors in the calculated interaction potentials of the previously proposed HPNX and YhHX models (Bornberg-Bauer 1997) producing the hHPNX interaction matrix:

Table 2.3

	h	Н	Р	N	X
h	2	-4	0	0	0
Н	-4	-3	0	0	0
Р	0	0	1	-1	0
N	0	0	-1	1	0
X	0	0	0	0	0

The extended categories were also motivated by experimatal data that emphasises the importance of treating Alanine and $Valine^{14}$ separately from other hydrophobic residues as they consistently observed that these groups underwent different reactions, resulting in different potentials; this was attributed to their geometrical positions in the folded protein (Crippen 1991).

More detailed work has been conducted in the work I has cited thus far, including but not limited to:

 $^{^{13}}$ This can be proved using Group Theory axioms as lattices form geometric groups - needs citation

 $^{^{14}}A, V \in H \text{ in HPNX}$

- Investigating geometric and volumentric constraints of various lattice structures¹⁵.
- Optimizing linear integer programming constraints with tighter bounds on H-H contacts¹⁶.
- Analysis of conformational lanscapes with respect to free energy landscapes¹⁷.
- Probabilistic thermodynamic analysis of folding pathways¹⁸.

Thus far I have summarised the most important results from these authors within the context of this project. In **Related Work** I will explore how these models have been re-contextualised within the reinforcement learning framework and then propose a novel method that incorporates results covered in this section that have yet to be exploited in recent work.

2.2 Reinforcement Learning

In this section I will introduce the Reinforcement Learning (RL) paradigm. Then, its integration with deep learning is explored, and subsequent improvements in the algorithm are also elaborated upon. The end result, the Rainbow DQN agent, is integrated as part of the **System Design** and **Specification**.

2.2.1 Markov Decision Processes

A finite markov chain is a process that consists of a set of states $\mathbf{S}^{\mathbf{n}} \coloneqq \{S_1, S_2, \dots, S_n\}$ and a transition function $F(\mathbf{S})$ that takes the state at the current timestep t and outputs a new state at timestep t+1.

$$F(S_t) \mapsto S_{t+1} \quad \forall S \in \mathbf{S}, \forall t \in \mathbf{T}$$
 (2.12)

For a given process, the states are linked by a transition probability $P(S_{t+1}, S_t)$. A process is markov only if the markov property holds:

$$P(S_{t+1}, S_t) = P(S_{t+1}|S_t)$$
(2.13)

 $^{^{15}}$ (Yue & Dill 1993, Dill et al. 2008)

¹⁶(Yue & Dill 1993)

¹⁷(Dill et al. 2008)

¹⁸(Yang et al. 2013, Dill et al. 2008)

The next state in a process that obeys the markov property is determined solely by the value of the current state, and for any given sequence, the transition probability between any two states remains the same. Thus, markov chains can begin characterised by a transition matrix $\mathbf{P} := |\mathbf{S}^n| \times |\mathbf{S}^n|$ where each row j is a distribution $P(i, \cdot)$ such that:

$$i \in \mathbf{S}^n, \sum_{j \in \mathbf{S}^n} P(i, j) = 1 \tag{2.14}$$

The product along the column space $j \in \mathbf{S}^n$, $\prod_{i \in \mathbf{S}^n} P(i,j)$ of the transition matrix reveals the stationary probability of being in any particular state $P(S_j)$.

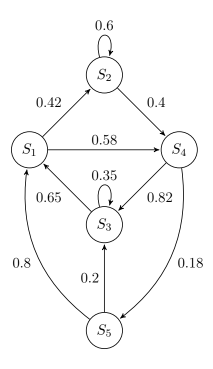


Figure 2.7: Markov chain with state space and transition propabilities

A Markov Decision Process (MDP) is a generalisation of this framework to sequential decision making (Sutton 2018). An MDP consists of an agent-environment interface, where the *Agent* is the learner and decision maker and the *Environment* consists of everything outside of the agent. The agent interacts with the environment bny taking action $a \in \mathbf{A}$ in the current state S_t and receives a reward R_{t+1}^{19} , the outcome of the agents actions transitions the environment from state S_t to S_{t+1} . The agent's long term reward is maximised by taking actions that move the agents into favourable states that yield higher rewards, the **expected** reward in any given state

¹⁹Reward received at t+1 to indicate the fact that an action must be taken first to move to a new state in order to obtain a reward

is equal to the probability of entering state S_{t+1} multiplied by the value of the reward R_{t+1} in that state.

$$\mathbb{E}_{S \in \mathbf{S}^n} \left[R_{t+1} \mid S_t, A_t \right] = P(R_{t+1} \mid S_t, A_t) \cdot R_{t+1}$$
(2.15)

The value of a given state $\mathcal{V}(S)$ is the total expected reward for that state for any action taken in that state. This is taken to be the long run return of state S if the process was repeated in the infinite limit under a stationary distribution of rewards²⁰:

$$\mathcal{V}(S) = \sum_{\forall a \in \mathbf{A}} \mathbb{E}_{S \in \mathbf{S}^n} \left[R_{t+1} \mid S, a \right]$$
 (2.16)

An agent in the environment seeks to take actions that maximise its long term reward at each timestep:

$$\sum_{t \in \mathbf{T}} \max_{a} \left(\mathbb{E}_{S \in \mathbf{S}^n} \left[R_{t+1} \mid S_t, a \right] \right) \tag{2.17}$$

In episodic settings the set of tasks underlying the MDP contains a terminal state that is reached after a finite number of timesteps, whereas continuous processes may not necessarily have a finite number of time steps. These settings can be unified under the notion of an absorbing state τ which is one that transitions only to itself with a reward of $R_{\tau} = 0$. For a given MDP, successive runs from S_0 to S_{τ} are termed episodes. In applied settings, an agent will repeatedly play through episodes with the Goal of maximising the cumulative reward in each episode:

$$G_t := R_{t+1} + R_{t+2} + \dots + R_{\tau-1} = \sum_{t=0}^{t=\tau-1} R_{t+1}$$
 (2.18)

Equations (2.15-18) assume a stationary distribution of rewards, and (2.18) is incompatible with continuous domains where the cumulative reward that the agent tries to maximise can be infinite if $\tau = \infty$. Many real world tasks such as playing video games (Mnih, Kavukcuoglu, Silver, Rusu, Veness, Bellemare, Graves, Riedmiller, Fidjeland, Ostrovski, Petersen, Beattie, Sadik, Antonoglou, King, Kumaran, Wierstra, Legg & Hassabis 2015) and dialogue generation (Weisz, Budzianowski, Su & Gasic 2018) fall into the category of continuous, non-stationary MDPs. By instead considering the discounted cumulative reward, the analysis of both episodic and continuous tasks with non-stationary distributions of rewards becomes computationally tractable:

$$0 < \gamma \leqslant 1$$

$$R_{t+1} + \gamma R_{t+2} + \gamma^2 R_{t+3} \dots + \gamma^{\tau - t - 1} R_{\tau - 1}$$
(2.19)

 $^{^{20}}$ "Stationary" indicating that the probability of receiving a reward in any state does not change as the process evolves

Where γ is referred to as the *discount factor*. The use of a discount factor contracts rewards further into the future asymptotically towards zero as (2.19) forms a geometric series²¹ with ratio r < 1:

$$\lim_{n \to \infty} \left(\sum_{k=0}^{n} \alpha \cdot r^k \right) = \frac{\alpha}{1-r}, \quad \forall \ 0 < r < 1, \quad \alpha \in \mathbb{R}^+$$
 (2.20)

Therefore by (2.20):

$$\lim_{n \to \infty} \left(\sum_{k=0}^{n} 1 \cdot \gamma^k \right) = \frac{1}{1 - \gamma} \tag{2.21}$$

Intuitively, this ensures that the agents values immediate rewards more so than delayed rewards as the rewards in the current timestep can be more accurately estimated than rewards further into the future due to the non-stationarity of the process. And so we can rewrite (2.19) in terms of cumulative reward (2.18):

$$G_{t} = R_{t+1} + \gamma R_{t+2} + \gamma^{2} R_{t+3} \dots + \gamma^{\tau-t-1} R_{\tau-1}$$

$$= R_{t+1} + \gamma \left[R_{t+2} + \gamma R_{t+3} + \dots + \gamma^{\tau-t-2} R_{\tau-1} \right]$$

$$= R_{t+1} + \gamma G_{t+1}$$
(2.22)

From (2.22) we can see that a stationary process (2.18) is a special case of the cumulative discounted reward where $\gamma = 1$. The closer the discount factor is to 1, the more the agent values future rewards.

In order for an agent to choose actions that maximise the discounted future rewards, it must adhere to a policy π . A policy is a mapping of states to a distribution of possible actions:

$$\pi : \mathbf{S} \to P(\mathbf{A})$$

$$\sum_{a \in \mathbf{A}} \pi(a \mid s) = 1, \quad \forall s \in \mathbf{S}$$
(2.23)

Under this policy, the *value* of a state (2.16) can be rewritten as:²²

$$v_{\pi}(s) = \sum_{a \in \mathbf{A}} \pi(a \mid s) \cdot \mathbb{E} \left[R_{t+1} + \gamma G_{t+1} \middle| S_t = s \right]$$

$$= \mathbb{E} \left[R_{t+1} + \gamma G_{t+1} \middle| S_t = s \right]$$
(2.24)

Equation (2.24) describes an agent's state-value function, which measures the expected return of a given state if the process was repeated infinitely. Similarly, we can define the action value function $q_{\pi}(s, a)$ to express the average return of taking a particular action in a given state:

$$q_{\pi}(s, a) = \mathbb{E}_{\pi} \left[R_{t+1} + \gamma G_{t+1} \, \middle| \, S_t = s, A_t = a \right]$$
 (2.25)

²¹See Appendix A.3

²²Needs further derivation

Formally, the action value function describes the expected return of being in state s and taking action a and following π thereafter. It is expected reward described in (2.24) conditioned additionally upon a fixed action being taken in the current state.

2.2.1.1 Bellman Equation

The state value function and the action value function for a given MDP can be estimated from experience, the Bellman Equation combines the equations described so far and codifys the relationship between the value of a state and the value of successor states:

$$\upsilon_{\pi}(s) = \mathbb{E}_{\pi} \left[R_{t+1} + \gamma G_{t+1} \mid S_{t} = s \right]$$

$$= \sum_{a \in \mathbf{A}} \pi(a \mid s) \sum_{s' \in \mathbf{S}} \sum_{r \in \mathbf{R}} P(s', r \mid s, a) \left(r + \gamma \mathbb{E}_{\pi} \left[r + G_{t+1} \mid S_{t+1} = s' \right] \right)$$

$$= \sum_{a \in \mathbf{A}} \pi(a \mid s) \sum_{s' \in \mathbf{S}} \sum_{r \in \mathbf{R}} P(s', r \mid s, a) \left(r + \gamma \upsilon_{\pi}(s') \right)$$

$$(2.26)$$

This represents a weighted sum of the expectations over all possible next states given the current state s and action taken a (Sutton 2018).

2.2.1.2 Optimality

Optimal state and value functions (v_*, q_*) are those that maximise the expected return under policy π . A policy π is defined to be better or equal to policy π' if its expected return is greater or equal to π' for all states.

$$v_*(s) = \max_{\pi} v_{\pi}(s)$$

$$q_*(s) = \max_{\pi} q_{\pi}(s, a)$$

$$\Rightarrow q_*(s) = \mathbb{E} \left[R_{t+1} + \gamma v_*(S_{t+1}) \mid S_t = s, A_t = a \right]$$

$$(2.27)$$

As v_* is the optimal value function, its expected return must equal the expected return for greedily selecting the *best action* in that state everytime, otherwise there would exist $v_{\pi*} > v_*$:

$$v_{*}(s) = \max_{a \in \mathbf{A}} q_{\pi*}(s, a)$$

$$= \max_{a} \mathbb{E} \left[R_{t+1} + \gamma v_{*}(S_{t+1}) \mid S_{t} = s, A_{t} = a \right] \quad \text{by (2.27)}$$

$$= \max_{a} \sum_{s' \in \mathbf{S}} \sum_{r \in \mathbf{R}} P(s', r \mid s, a) [r + \gamma v_{*}(s')]$$
(2.28)

From (2.28) we can see that the optimal value function is defined independently of any policy, indicating that it can be reached starting from any arbitary policy, and improving on that, this is formalised under the framework of *Generalised Policy Iteration* (GPI).

2.2.1.3 Generalised Policy Iteration

GPI consists of an alternating process of making the value function more accurate with respect to the policy and making the policy greedy with respect to the current value function. This process is decomposed into two separate steps:

1. Policy Evaluation

An initial value function v_0 is parameterised with arbitary values for each $s \in \mathbf{S}$. Given a sequence of value functions v_0, v_1, v_2, \ldots each succesive value can be calculated by:

$$v_{k+1}(s) = \mathbb{E}_{\pi} \left[R_{t+1} + \gamma v_k(S_{t+1}) \, \middle| \, S_t = s \right]$$
 (2.29)

The value of each state $s \in \mathbf{S}$ is updated "in place" with the revised estimate garnered from the current reward (which may not have been achieved previously) and the expected immediate rewards approximated by the previous value function. A pass over all \mathbf{S} produces a new state value function.

2. Policy Iteration

For a given value function, (2.28) shows that the optimal policy is that which greedily selections actions that maximise the expected future returns at each state. It follows that:

$$\pi \in \{\Pi\} := \text{ Set of all policies}$$

$$v \in \{\Upsilon\} := \text{ Set of all value functions}$$
 (2.30)
$$q \in \{Q\} := \text{ Set of all action value functions}$$

$$\forall \pi \exists q_* \mid q_* \ge (q' \ne q_*) \Longleftrightarrow \forall \pi \exists v_* \mid v_* \ge (v' \ne v_*) \tag{2.31}$$

$$\pi' \ge \pi \to \upsilon_{\pi'}(s) \ge \upsilon_{\pi}(s) \quad \forall s \in \mathbf{S}$$

$$\therefore q_*^{\pi'}(s, a) \ge q_*^{\pi}(s, a)$$
(2.32)

$$\Rightarrow q_{\pi}(s, \pi'(s)) \ge \upsilon_{\pi}(s) \tag{2.33}$$

This inductive line of logic reasons that if $\exists \pi' \geq \pi$ then the expected returns of a given state calculated under policy π' will be higher, thus if the less optimal policy had instead chosen the action according to the optimal policy, then the expected returns would be greater (2.33). The optimal policy is obtained by making the current policy greedy with respect to the action value function:

$$\pi'(s) = \underset{a}{\operatorname{argmax}} \ q_{\pi}(s, a)$$

$$= \underset{a}{\operatorname{argmax}} \sum_{s' \in \mathbf{S}} \sum_{r \in \mathbf{R}} P(s', r, | s, a) \left[r + \upsilon_{\pi}(s') \right]$$
(2.34)

By succesively cycling through policy evaluation and improvement, we continuously update our estimate of the value function and then greedily select actions in each state according to our updated estimates. This cycle is referred to as *Generalised Policy Iteration* and can be proven to monotonically converge to an optimal value function and an optimal policy. Upon convergence, the process reaches an equilibrium where the policy does not change because the value of the states has not changed (Sutton 2018).

2.2.1.4 Exploration vs Exploitation

The full reinforcement learning problem can be characterised by the models we have described so far with the addition of the exploration - exploitation trade off. So far we have considered agents with full knowledge of the environment, but for many MDPs, the agent may never access all available state action pairs, or indeed the number of state action pairs may be unenumerable. In order to maximise returns, the agent must strike a balance between exploring new, unseen states and exploiting the greedy strategy it has already learned. A naive exploration strategy²³ is to choose the greedy action with fixed probability ε and otherwise choose a random action with probability $1 - \varepsilon$; this is called an ε greedy strategy. However the agent would still require an infinite amount of episodes to form the optimal policy and value functions, they must instead be approximated in such a way that preserves the gurantees of GPI.

2.2.1.5 Monte Carlo Estimation

Monte carlo estimation enables an agent to *learn* optimal policies and value functions from experience alone. In order to take advanatge of Monte Carlo methods, the MDP is modelled in

²³More recent and general exploration strategies are discussed in **2.2.3.2** and **Related Work**.

the following way:

- Each state has its own stationary distribution of rewards.
- Each state is inter-related, i.e taking an action in one state depends on actions taken in later states.
- As all action selections are undergoing learning, the distribution of rewards becomes *non-stationary* from the point of view of later states.

Monte Carlo methods sample average returns for a given state-action pair for a given episode and use those samples to update estimates of action values. Two distinct approaches enable us to do so:

1. On Policy Methods

In "On Policy Control" we aim to evaluate or improve the policy that is used to make decisions, which in turn produces data in the form of the outcomes of these decisions.

2. Off Policy Methods

In "Off-Policy Control" attempt to improve a policy different than the one used to generate the data.

Within the scope of this project we will focus on Off Policy methods. First we consider two policies $\pi \neq b$ and we refer to the former as the target policy and the latter as the behaviour policy. The goal is to estimate v_{π} or q_{π} . We also assume that $\pi(a|s) > 0 \rightarrow b(a|s) > 0$ implying that every action taken under π is also taken at least once under b; this is the assumption of coverage. From coverage it follows that b must be stochastic in states where it is not identical to π as π_* , being the optimal policy, is greedy and deterministic.

First I will introduce *importance sampling*, a method for estimating expected values under on distribution given samples from another distribution. This is a general technique used in off-policy methods (Sutton 2018) as well as in later discussed *Prioritised Experience Replay*²⁴. Given a starting state S_t the probability of a subsequent state action trajectory $A_t, S_{t+1}, A_{t+1}, \ldots$ is:

$$\pi(A_t \mid S_t) \cdot P(S_{t+1} \mid S_t, A_t) \cdot \pi(A_{t+1} \mid S_{t+1}) \dots P(S_T \mid S_{T-1}, A_{T-1})$$

$$= \prod_{k=t}^{T-1} \pi(A_k \mid S_k) \cdot P(S_{k+1} \mid S_k, A_k)$$
(2.35)

²⁴See Memory

Therefore, in order to calculate the relative probability (importance sampling ratio) of a trajectory given the target policy and and the behaviour policy:

$$\rho_{t:T-1} = \frac{\prod_{k=t}^{T-1} \pi(A_k \mid S_k) \cdot P(S_{k+1} \mid S_k, A_k)}{\prod_{k=t}^{T-1} b(A_k \mid S_k) \cdot P(S_{k+1} \mid S_k, A_k)}$$

$$= \prod_{k=t}^{T-1} \frac{\pi(A_k \mid S_k)}{b(A_k \mid S_k)}$$
(2.36)

Intuitively, this is a ratio of how much more or less probably the trajectory is under π relative to b, given that $v_b(s) = \mathbb{E}\left[G_t \mid S_t = s\right]$ we can transform the returns under b to have to correct expected value by multiplying them by the importance sampling ratio:

$$\mathbb{E}\left[\rho_{t:T-1}G_t \mid S_t = s\right] = \nu_{\pi}(s) \tag{2.37}$$

In the case of full monte carlo methods, the returns are averaged at the end of each episode, weighted importance sampling applies the importance sampling ratio as weighted average of the returns over the whole episode with terminal state T(t) to form our estimate V:

$$V_{\pi}(s) = \frac{\sum_{t \in T(t)} \rho_{t:T-1} \cdot G_t}{\sum_{t \in T(t)} \rho_{t:T-1}}$$
(2.38)

This can be rewritten as an incremental update that occurs on an episode by episode basis where we start with random weights W_i and for each state maintain the cumulative sum of the weights given to the first n returns (C_n) :

$$W_i = \rho_{t_i:T(t_i)-1}$$

$$C_n = \sum_{k=1}^n W_k$$

$$V_n + 1(s) = V_n(s) + \frac{W_n}{C_n} \left[G_n - v_n(s) \right]$$

$$(2.39)$$

Where the text in red is a target update as measurement of the error between the actual returns for that episode G_n and the previously estimated returns $v_n(s)$. This is then multiplied proportionally by a weighted importance sampling ratio. The original estimate of $v_n(s)$ is the updated proportionally in the direction towards the actual expected returns.

2.2.1.6 Model Free vs Model Based

Sometimes it is possible to provide the agent with access to a model of the environment. If an agent has access to a model of the environment, typically in the form of a simulator, it can simulate states n steps ahead and use that to inform its estimated returns trivially using the

approaches described so far. The trouble with this approach however is that when modelling a real world task, no coarse grained model will ever approximately the task perfectly. A poorly designed model can also introduce unintended biases into the agent (Sutton 2018). On the other hand model free methods, such as monte carlo estimation, can approximate the state value or action value function directly, and have shown better generalization properties with regard to acting in unforseen scenarios.

In this project I will focus on the model-free off-policy methods as I will show that the PSP problem on a lattice is a non-stationary MDP for which a model is not computationally tractable. With the foundations lain, I will introduce the central algorithm to contemporary reinforcement learning.

2.2.2 Deep Q Learning

2.2.2.1 Temporal Difference Error

The temporal difference (TF) error is a special case of monte carlo estimation where the action value functions and the state value functions are updates with each timestep instead of at the end of each episode.

$$V(S_t) \leftarrow V(s_t) + \alpha \left[R_{t+1} + \gamma V(S_{t+1}) - V(S_t) \right]$$
 (2.40)

This is similar to the target update in monte carlo estimation, however instead our we update our existing estimate based on our current estimate, the target in (2.39) is expanded using (2.22). In this particular case for each episode the agent takes an action, observes the outcome of this action at S_{t+1} and evaluates the value function at this timestep; this evaluation is then used as part of a target update that is weighted by some constant step size parametere a.

2.2.2.2 Q Learning

We can go a step further and unpack (2.40) some more by observing (2.28) and (2.32 - 34) to form an estimate of our action value function Q directly forming an off policy TD control update rule:

$$Q(S_t, A_t) \leftarrow Q(S_t, A_t) + \alpha \left[R_{t+1} + \gamma \cdot \max_{a} Q(S_{t+1}, a) - Q(S_t, A_t) \right]$$
 (2.41)

This update target is formally known as the Bellman Error denoted δ_t . I introduce the central algorithm (Sutton 2018): The Q learning algorithm forms the core of methods discussed and

Algorithm 1: Q Learning

```
\alpha \in (0,1];
0 < \varepsilon \ll 1;
Init Q(s,a) \forall s \in \mathbf{S}, a \in \mathbf{A}(s) randomly;
Q(terminal, \cdot) = 0;
forall episodes do
    Init S;
    forall steps in episode do
         r\backsim U(0,1); // Sample random number r
         if r > \varepsilon then
             A \leftarrow \underset{a}{argmax} \ Q(s,a);
         else
              A \leftarrow rand(a \in \mathbf{A}); // Choose a random action
         end
         Take action A, observe R, S';
         Q(S, A) \leftarrow Q(S, A) + \alpha [R + \gamma \cdot \max_{a} Q(S', a) - Q(S, A)] ;
    end
end
```

proposed throughout this project. After a brief review of neural networks, I will show how they can parameterise the Q function.

2.2.2.3 Neural Networks

A neural network is a universal non-linear function approximator. It consists of smaller sub units called perceptrons. Each perceptron acts as linear function with respect to its inputs \mathbf{x} and can be parameterised by a weight vector \mathbf{w} and bias b.

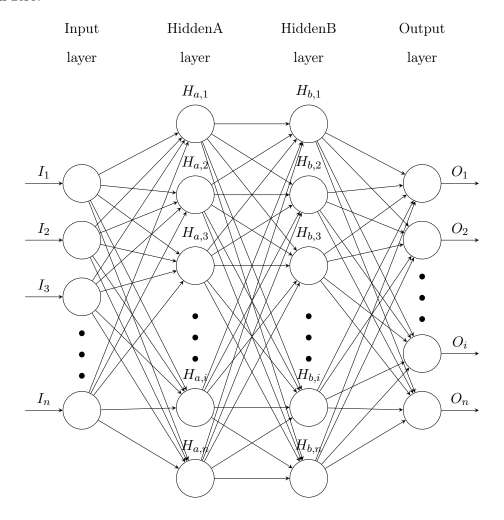
$$\hat{\mathbf{y}} = \mathbf{x} \cdot \mathbf{w}^{\top} + b \tag{2.42}$$

Perceptrons can be organised into layers hence become multi-layer perceptrons MLP. If an MLP contains more than a single intermediate layer between the input and the output layer, it is known as a deep neural net. When organised into layers, the output of each perceptron must

be passed through a non-linearity

$$ReLU(\hat{\mathbf{y}}) = max(\hat{\mathbf{y}}, 0)$$

to remove the correlation between layers, unlocking its potential as a non-linear function approximator. Contemporary architecture use non-linearities such as rectified linear units (ReLU) as they demonstrate faster convergence due to sparser representations induced by zeroing weights less than zero.



A network can be represented as a function f parameterised by weight matrix \mathbf{W} and bias matrix \mathbf{b} with the outputs of each layer being passed into the next layer in a fully connected network.

$$\hat{\mathbf{y}} = f(\mathbf{x}; \mathbf{W}, \mathbf{b}) \tag{2.43}$$

The network is optimised against a target output Y using stochastic gradient descent.

$$L(Y, \hat{\mathbf{y}}) = \|Y - \hat{\mathbf{y}}\|_2$$

$$= \frac{1}{N} \cdot \sum_{i=1}^{N} (Y_i - (\mathbf{x}_i \cdot \mathbf{w}_i^{\top} + b))^2$$
(2.44)

$$\frac{\partial E}{\partial \mathbf{w}} = -\frac{2}{N} \cdot \sum_{i=1}^{N} \mathbf{x}_{i} (Y_{i} - (\mathbf{x}_{i} \cdot \mathbf{w}_{i}^{\top} + b))$$

$$\frac{\partial E}{\partial b} = -\frac{2}{N} \cdot \sum_{i=1}^{N} (Y_{i} - (\mathbf{x}_{i} \cdot \mathbf{w}_{i}^{\top} + b))$$
(2.45)

$$\nabla_{\mathbf{W}} E = \frac{\partial L(Y, \hat{\mathbf{y}})}{\partial \mathbf{W}}, \quad \nabla_{\mathbf{b}} E = \frac{\partial L(Y, \hat{\mathbf{y}})}{\partial \mathbf{b}}$$
 (2.46)

The error is calculated first for the network's output layers, the weights are then updated in the direction of negative gradient, steepest descent with respect to error E with some constant stepsize η :

$$\mathbf{W}_{k+1} = \mathbf{W}_k - \eta \nabla E_{\mathbf{W}_k} \tag{2.47}$$

The gradients for the biases are updates likewise. This error is then back-propagated using reverse mode differentiation on a computational graph (Margossian 2019) through the remaining nodes, where the weights adjusted after the gradient updates are used as the target for the preceding layer.

Remark. In recent years additional properties of neural networks have been discovered relating to their nature as universal function approximators. Neural networks with a single hidden layer can approximate any function, and as the number of nodes tends to infinity it simplifies to a linear model (Lee, Xiao, Schoenholz, Bahri, Novak, Sohl-Dickstein & Pennington 2019). Alternatively, infinitely wide, deep neural networks are equivalent to a gaussian process (Lee, Bahri, Novak, Schoenholz, Pennington & Sohl-Dickstein 2017). The have also demonstrated the ability to sort lists in practically O(1) time (Zhu, Cheng, Zhang, Liu, He, Yao & Zhou 2019).

2.2.2.4 Deep Q-Networks

Neural networks can be used to approximate the Q function directly using Q learning, however the structure of the gradient update introduces complexity when trying to estimate expected value of rewards. We begin by parameterising our Q fuction with the weights of our network:

$$Q(s, a) = f(\mathbf{s}; \mathbf{W}, \mathbf{b}) \quad s \in \mathbf{S}$$

$$\Rightarrow Q(s, a; \mathbf{W}, \mathbf{b}) \to Q(s, a; \mathbf{W})$$
(2.48)

We can develop a loss function in the off policy setting using the Bellman Error δ_t . This holds because the update target $R_{t+1} + \gamma \max_a Q(S_{t+1}, a)$ acts as a "peek ahead" from the perspective

of the current state. That is we take an action, observe the outcome, and update the action-value of our old state with this new information.

$$L(\delta_{t}, Q(s, a; \mathbf{W})) = \|\delta_{t} - Q(s, a; \mathbf{W})\|_{2}$$

$$\frac{\partial L(\delta_{t}, Q(s, a; \mathbf{W}))}{\partial \mathbf{W}} = \frac{\partial (\delta_{t} - Q(s, a; \mathbf{W}))^{2}}{\partial \mathbf{W}}$$

$$= (\delta_{t} - Q(s, a; \mathbf{W})) \cdot \frac{\partial (\delta_{t} - Q(s, a; \mathbf{W}))}{\partial \mathbf{W}}$$

$$= (\delta_{t} - Q(s, a; \mathbf{W})) \nabla_{\mathbf{W}} Q(s, a; \mathbf{W})$$
(2.49)

The target acts as a constant because the maximal estimated Q value at the next state is taken to be the value of the state under the optimal policy (see 2.32). In their original paper, (Mnih et al. 2015) used a neural network to take in the state as input and then output a vector of action values, one for each action. If we used the interpretation of Q learning developed in (2.41), this structure could lead to unstable gradient updates or could cause the action values to diverge. This can be described in terms of the correlation between various elements of the model, consider two dependent random variables X and Y:

The corelation between two dependent random variables is proportional to the covariance of the

$$Correlation(X,Y) = \frac{Covar(X,Y)}{\sqrt{Var(X)Var(Y)}}$$
 (2.50)

random variables normalised by their joint standard deviatiation $\sqrt{Var(X)} = \sigma$. As correlation is a dimensionless constant on the interval [-1,1], if X and Y are correlated $Corr(X,Y) \approx 1$, then they increasingly co-vary across each dimension, then their standard deviation must increase likewise, and as the co-variance goes to infinity, so does their individual variance. And so under (2.41) we would be using a very similar distribution of weights to calculate $\max_a Q(S_{t+1}, a)$ and $Q(S_t, a)$ which are both dependent according to the definition of our MDP. In addition to that, if we consider successive frames of pixels on a screen as our states, it is trivial note that these observations will also be highly correlated; in the case of the atari game "Pong", the position on the paddle on the next frame is highly informed by position of the paddle in the next frame. Instead, (Mnih et al. 2015) propose two solutions to this problem. A biologically inspired mechanism termed Experience Replay and a separate Target Network. To remove the correlations between successive observations $(S_t, A_t, R_{t+1}, S_{t+1})$, they instead store these tuples in a Target Network. In this case the replay buffer is a FIFO array of constant size that overwrites older memories with new ones. The target network is a separate neural network whose weights θ^- are initialised with the weights of the behaviour network and then frozen. Periodically, this target

network is updated with the current weights of the behaviour network θ . Every timestep, a batch of memories are uniformly sampled, the target network is used to compute the Bellman Error for each memory and the output of the behaviour network is updated according to the loss defined in (2.49), averaged across the whole batch. This algorithm proceeded to lead a breakthrough in contemporary deep reinforcment learning by outperforming human benchmarks at a super human level for the first time (adapted from the original authors):

Algorithm 2: Deep Q Network (DQN)

```
Init replay memory D to capacity N;
Init Q with random weights \theta;
Init \hat{Q} with \theta^- = \theta;
forall episodes do
    Init state s_1;
    Preprocess sequence \phi_1 = \phi(s_1);
    forall timesteps in episode do
        r \backsim U(0,1);
        if r < \varepsilon then
            a_t \backsim A; // Randomly sample action from action space
        else
            a_t = \underset{a}{argmax} \ Q(\phi(s_t), a; \theta);
        end
        Execute action a_t observe R_{t+1}, s_{t+1}, d; // d = 1 if state is terminal else 0
        Store (\phi_1, a_t, r_{t+1}, \phi_{t+1}) in D;
        Sampled random minibacth from D;
        \delta_t = r_t + (1 - d)\gamma \max_{a'} \hat{Q}(\phi_{t+1}, a'; \theta^-);
        Perform gradient update (\delta_t - Q(s, a; \theta)) \nabla_{\theta} Q(s, a; \theta);
        Every C steps reset \hat{Q} = Q;
    end
```

 \mathbf{end}

2.2.3 Improvements to Vanilla Deep Q-Networks

The success of the original DQN algorithm was not without flaws however.

- 2.2.3.1 Memory
- 2.2.3.2 Exploration
- 2.2.3.3 Exploitation
- 2.2.3.4 Rainbow DQN

2.3 Multi-Agent Reinforcement Learning

- 2.3.1 Stochastic Games
- 2.3.1.1 Game Theory
- 2.3.1.2 Expected Pay-offs as Expected Rewards
- 2.3.1.3 Multi-Agent Games
- 2.3.1.4 Mean Field Games
- 2.3.1.5 Mean Field multi-agent reinforcement learning
- 2.4 Related Work
- 2.4.1 MCMC methods for lattice models
- 2.4.2 Deep Learning methods for lattice models
- 2.4.2.1 Alpha-Fold
- 2.4.3 Reinforcement Learning methods for lattice models

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