Abstract

Quantum reinforcement learning is an emerging field at the intersection of quantum computing and machine learning. While we intend to provide a broad overview of the literature on quantum reinforcement learning (our interpretation of this term will be clarified below), we put particular emphasis on recent developments. With a focus on already available noisy intermediate-scale quantum devices, these include variational quantum circuits acting as function approximators in an otherwise classical reinforcement learning setting. In addition, we survey quantum reinforcement learning algorithms based on future fault-tolerant hardware, some of which come with a provable quantum advantage. We provide both a birds-eye-view of the field, as well as summaries and reviews for selected parts of the literature.

Contents

1 Introduction and Overview
2 Classical Reinforcement Learning
3 The Quantum Computing Paradigm
4 Quantum Reinforcement Learning Algorithms
   4.1 Quantum-Inspired Reinforcement Learning based on Amplitude Amplification
   4.2 Quantum Reinforcement Learning with Variational Quantum Circuits
   4.3 Projective Simulation for Quantum Reinforcement Learning
   4.4 Boltzmann Machines for Quantum Reinforcement Learning
   4.5 Quantum Policy and Value Iteration
   4.6 Quantum Reinforcement Learning with Oracularized Environments
5 Outlook
1 Introduction and Overview

With recent advances in the fabrication and control of hardware for quantum information processing, the possibilities of merging quantum computing (QC) with machine learning (ML) have received a huge amount of attention within the growing research community. Hereby, reinforcement learning (RL) is the third paradigm besides supervised and unsupervised learning. In this survey article, we provide an overview over so-called quantum reinforcement learning (QRL) algorithms. We understand these as quantum-assisted approaches, that solve a particular task (be they classical or quantum in nature) by employing quantum resources (either in simulation and/or in experiment).

In order to keep this contribution as self-contained as possible, we provide the necessary backgrounds before venturing into the QRL literature. We start out with a brief recap of the essentials of the RL paradigm in the fully classical setting in Sec. 2. Further, in Sec. 3 we provide a quick introduction to QC and variational quantum circuits (VQCs). Readers familiar with either of the topics may safely skip these sections.

In Sec. 4 we turn our attention to the emerging field of QRL, starting out with a quick overview of the literature. Then we delve into summaries of the most prominent contributions. This selection is necessarily subjective and reflects our own research interests. We organized our summaries into several blocks, that are ordered by what one could call an increasing degree of ‘quantization’. The first of these blocks in Subsec. 4.1 covers what we refer to as ‘quantum-inspired’ RL algorithms. The second block in Subsec. 4.2 takes a rather detailed look at QRL algorithms that employ so-called VQCs as function approximators. In many cases, the corresponding algorithms are obtained by simply replacing a standard neural network function approximator (or any other sort) by an appropriate VQC. We provide detailed summaries for most papers in this category, as variational quantum algorithms are believed to offer the potential to obtain quantum advantage despite the

Figure 1: A possible classification matrix for QRL algorithms, where we took into account only those variants of QRL which we focus on in Sec. 4. The algorithm classes are ordered according to their degree of quantum-classical hybridization, ranging from purely classical to purely quantum. A more detailed review of quantum-inspired reinforcement learning (QiRL)–algorithms can be found in Subsec. 4.1. VQC-based approaches are summarized in quite some detail in Subsec. 4.2. QRL-algorithms employing post-noisy intermediate-scale quantum (NISQ) quantum algorithms as subroutines or even fully quantum approaches to QRL are described in Subsec. 4.3, Subsec. 4.4, Subsec. 4.5 and Subsec. 4.6. The dashed vertical line between classical and NISQ compute resources indicates that presently it is unclear whether QRL with NISQ-compatible algorithms offers robust quantum advantage on a broad range of learning problems. The solid vertical line distinguishes post-NISQ algorithms from both classical and NISQ-compatible algorithms, as they typically come with guaranteed quantum advantage (at least relative to their classical counterparts).
limitations of present day NISQ hardware. In Subsecs. 4.3 and 4.4, we take a look at realizations of QRL based on so-called projective simulation and the use of Boltzmann machines as function approximators, respectively. In Subsec. 4.5 we move to a class of approaches that employ quantum algorithms as subroutines. The corresponding hardware requirements will likely be compatible only with universal, fault-tolerant and error-corrected quantum processing units (QPUs). Finally, Subsec. 4.6 provides a summary for a formal approach to QRL, which treats all components of RL `quantumly'. From our point of view, the highest degree of quantization can thus be found in these approaches. Fig. 1 gives an overview of the QRL literature as understood in this survey.

Finally, in Sec. 5 we state our concluding thoughts on the current state-of-the-art of QRL. Before moving to more technical content, we would like to express our hope that this literature survey on QRL will be of use to colleagues and collaborators and the wider QC research community. It represents our effort to familiarize ourselves with QRL and its main research directions.

2 Classical Reinforcement Learning

Compared to the methods of supervised and unsupervised learning, which are typically implemented as passive learning, RL falls into the class of interaction-based learning [SB18]. On an abstract level, the learner interacts with its environment, the state of which it can either fully or only partially observe through a corresponding observation obtained after executing an action according to an underlying policy. In the RL paradigm, the learner is therefore appropriately referred to as an agent: it can - be it in simulation or in the real world - interact with its environment according to its abilities. The aim of RL is to learn a policy through the interaction of the agent with the environment, which is optimal with regard to a reward adapted to the problem. In other words, the agent should find an optimal policy during the learning process in the abstract space of all policies, which maximizes the expected cumulative reward. The theoretical basis for RL is formed by so-called Markov decision processes (MDPs) and the associated Bellman equation, which represents a consistency equation for the so-called value function. In turn, an optimal policy can be extracted from the optimal value function. Alternatively, the optimal policy can also be learned directly. Under certain conditions, the elements of RL can be mapped to their respective equivalents in control theory, where typically a dynamic optimization problem is solved by gradient-based methods with simulation of the corresponding model dynamics. On the RL side, there are both model-based and model-free approaches. The model-free approach in particular is one of the strengths of the RL method, since in many cases state and action spaces are too high-dimensional to design realistic dynamical models and simulate them to efficiently find optimal control strategies. The large dimensions of the spaces that occur in realistic problems make the use of approximation methods for the value function necessary. Driven by the breakthroughs in deep learning (DL), artificial neural networks (NNs) have established themselves as function approximators for both value function and policy (understood as a deterministic or probabilistic mapping of states to actions), thus establishing the field of deep reinforcement learning (DRL).

In the following, we will introduce the various notions pertaining to RL in a more formal way and provide the background necessary to understand the basic RL terminology. In an RL scenario, the algorithm, also referred to as agent, generates its own data by interacting with an environment. This interaction happens over some discrete timesteps \( t \), which are accumulated to episodes with either finite or infinite horizon. In each timestep, the agent is able to make an observation \( s_t \in S \) of the environment. Based on this state information, an action \( a_t \in A \) acting on the environment is selected according to a policy. Based on the (usually unknown) environment dynamics, the next state \( s_{t+1} \in S \) is observed from the environment and the agent receives a reward \( r_t \in R \subseteq \mathbb{R} \) for its choice. The agent should select the actions in such a way that some objective is optimized, usually related to the long term reward. A sketch of this pipeline can be found in Fig. 2. In this survey article, we follow the formalism and notation of Sutton et al. [SB18], with small adaptations wherever we feel that it eases comprehension.
Reinforcement Learning as a Markov Decision Process

More formally, this setup is usually described as an MDP. A finite MDP is a 5-tuple $(S, A, R, T, \gamma)$, where the sets $S$, $A$, and $R$ are finite. It is defined by the following components:

- A set of states $S$ the agent can observe from the environment
- A set of actions $A$ the agent can execute in the environment
- The reward function $R: S \times A \to \mathbb{R}$; The value $R(s, a) := E[r_t | s_t = s, a_t = a]$ gives the reward for the case, that action $a_t$ is executed in state $s_t$.
- The environment dynamics $T: S \times A \times S \to [0, 1]$; The value $T(s' | s, a) := \Pr\{s_{t+1} = s' | s_t = s, a_t = a\}$ gives the probability that the environment transitions to state $s_{t+1}$, if the agents executes action $a_t$ in state $s_t$ at time $t$.
- The discount factor $0 \leq \gamma \leq 1$, more on this below;

The dynamics of the environment are usually not accessible to the agent, otherwise the task would collapse to straightforward dynamic programming. The function $T$ does satisfy the properties of a probability density function (PDF), i.e., it holds $\sum_{s' \in S} T(s' | s, a) = 1$, for all choices of $s \in S$ and $a \in A$. According to the Markov property, the dynamics are completely described by $T$, i.e., the consecutive state $s_{t+1}$ depends solely on the directly preceding state $s_t$ and action $a_t$.

With this framework in mind, the interaction between agent and environment can be described as a trajectory $\tau$. For a finite or infinite horizon $H$, one episode is therefore given by the sequence

$$\tau = [s_0, a_0, r_0, s_1, a_1, r_1, s_2, \cdots, s_{H-1}, a_{H-1}, r_{H-1}],$$

with $s_t \in S$, $a_t \in A$, and $r_t$ a sample from the reward distribution for each timestep $t$.

Long Term Reward as Objective

The agent gets feedback from the environment through the rewards $r_t$. However, instead of maximizing these short-term rewards, it is much more appropriate to use some long term measure as objective. A natural choice is to go for the cumulative reward, also referred to as the expected return

$$G_t := r_t + r_{t+1} + r_{t+2} + \cdots + r_{H-1}. \quad (2)$$

For episodic tasks ($H < \infty$) it is often desirable and for continuous tasks ($H = \infty$) it is necessary to use a discount factor $\gamma$. This leads to the discounted return

$$G_t := \sum_{t'=t}^{H-1} \gamma^{t' - t} \cdot r_{t'}, \quad (3)$$

where each choice of $\gamma$ defines a different MDP. For $\gamma < 1$ the value of $G_t$ is guaranteed to be finite and emphasis on individual rewards decreases with distance from the current time-step. For $\gamma = 0$ the sum reduces to just the immediate reward, so an appropriate choice of this hyperparameter is crucial for the potential success of the RL agent.
Policy, Value Functions and Optimality

In order to describe a meaningful RL setup, there are still some concepts missing. As described above, the agent needs to decide for an action in every timestep, depending on the state information that is observed. This decision making process can be understood as a (stochastic) policy

$$\pi(a|s) := \Pr\{a_t = a|s_t = s\},$$  \hspace{1cm} (4)

where \(\sum_{a \in A} \pi(a|s) = 1\) holds for all \(s \in A\). The overall task of RL is to derive an optimal policy \(\pi^*\) w.r.t. some metric.

A suitable tool to define optimality and also to simplify updates is the notion of value functions. The state value function of state \(s\) under the current policy \(\pi\) is defined as

$$V_\pi(s) := \mathbb{E}_\pi[G_t|s_t = s].$$  \hspace{1cm} (5)

It describes the expected returns when starting in state \(s\) and following policy \(\pi\) from there on, with the value for a terminal state always zero by definition. It can be interpreted as a measure of how good it is to be in a certain state, where quality is measured w.r.t. expected return. Explicitly separating the first step in the definition above gives rise to the Bellman (expectation) equation

$$V_\pi(s) = \sum_{a \in A} \pi(a|s) \sum_{s' \in S} T(s'|s,a) \left[ R(s,a) + \gamma \cdot V_\pi(s') \right],$$  \hspace{1cm} (6)

for all \(s \in S\). Consequently, the value function \(V_\pi\) can be viewed as the unique solution to this Bellman equation. Alternatively, one can define the state-action value function as the expected return when starting in state \(s\), executing action \(a\), and following policy \(\pi\) from there on. It is defined as

$$Q_\pi(s,a) := \mathbb{E}_\pi[G_t|s_t = s, a_t = a],$$  \hspace{1cm} (7)

for all \(s \in S\) and \(a \in A\). It is straightforward to see that it holds \(V_\pi(s) = \sum_{a \in A} \pi(a|s) Q_\pi(s,a)\) for all \(s \in S\). This identity can be used to give the Bellman equation for the state-action value function as

$$Q_\pi(s,a) = \sum_{s' \in S} T(s'|s,a) \left[ R(s,a) + \gamma \cdot \sum_{a' \in A} \pi(a'|s') Q_\pi(s',a') \right].$$

The value function allows to explicitly define and evaluate the quality of policies, i.e., the policy \(\pi\) is better or equal to another policy \(\pi'\), iff \(V_\pi(s) \geq V_{\pi'}(s)\) for all \(s \in S\). If a policy is better or equal to all others, it is considered an optimal policy \(\pi^*\). All optimal policies share the same optimal state-value function

$$V_{\pi^*}(s) := V^*(s) := \max_\pi V_\pi(s),$$  \hspace{1cm} (8)

for all \(s \in S\). A similar notion of optimality for the action-value function is given by

$$Q^*(s,a) := \max_\pi Q_\pi(s,a),$$  \hspace{1cm} (9)

for all \(s \in S\) and \(a \in A\). It is straightforward to formulate the connection of both quantities as \(V^*(s) = \max_\pi \left( \sum_{a \in A} \pi(a|s) Q_\pi(s,a) \right) = \max_\pi Q^*(s,a)\). With this it is possible to derive the Bellman optimality equation for the value function as

$$V^*(s) = \max_{a \in A} \sum_{s' \in S} T(s'|s,a) \left[ R(s,a) + \gamma \cdot V^*(s') \right],$$  \hspace{1cm} (10)

for all \(s \in S\). Using the stated connection this can be reformulated to extend to the state-action value function as \(Q^*(s,a) = \sum_{s' \in S} T(s'|s,a) \left[ R(s,a) + \gamma \cdot \max_{a' \in A} Q^*(s',a') \right]\) for all \(s \in S\) and \(a \in A\).
Solving and Approximating the Bellman Equation

One topic that has to be addressed is the actual representation of the policy and value functions. The most intuitive approach is to just store the values for all state-action pairs in a table, also referred to as the tabular approach. While this formulation offers nice convergence and optimality guarantees for several scenarios, it has some serious drawbacks. Most prominently, it is intractable once the state-action space gets too large, which is the case for most real-world problems. A workaround is to use parametric function approximators, which results in the parameterized functions $\pi_\theta$, $V_\theta$, or $Q_\theta$, respectively. The typical choice is a NN [HSW89], in Subsec. 4.2 the usage of VQCs for this task is considered from several angles. As there now is an approximation in the defining quantities, also convergence guarantees are much less straightforward than for the tabular case. The remaining parts of this section can be understood both for the tabular and parameterized case, although details might vary a bit.

The Bellman optimality equation offers a tool to derive an optimal policy. It has to be noted that the given formulation makes use of the environment dynamics $T$. Therefore, solution methods solving the equation with dynamic programming are referred to as model-based. The two most prominent examples include value iteration [Bel57] and policy iteration [RN94; PRD96].

There is also a whole range of model-free approaches, where the agent does not make use of any model that represents the environment dynamics. Instead, all information is directly acquired by interaction with the environment. One prominent representative is the $Q$-learning approach [WD92], which basically is an approximation of $Q$-value iteration using samples. Starting with a random initialization, the update rule

$$Q(s, a) \leftarrow Q(s, a) + \alpha \left( r_t + \gamma \cdot \max_{a' \in A} Q(s', a') - Q(s, a) \right) \quad (11)$$

directly derives from the Bellman equation, where $\alpha$ is a learning rate hyperparameter. The policy is usually defined to act epsilon-greedily w.r.t. the current action-value function, i.e.

$$\pi(s) := \begin{cases} \arg \max_{a \in A} Q(s, a) & \text{with probability } 1 - \varepsilon, \\ \text{uniformly at random from } A & \text{with probability } \varepsilon. \end{cases} \quad (12)$$

An alternative approach is the policy gradient idea [Sut+99], which directly aims to learn the policy. Based on a parameterized policy $\pi_\theta$, it performs updates

$$\theta \leftarrow \theta + \alpha \nabla_\theta J(\theta) \quad (13)$$

via gradient ascent, where $J(\theta)$ is a performance measure, usually $J(\theta) = V_{\pi_\theta}(s_0)$. Unfortunately the desired gradient likely depends on some environment dynamics, which are not known. The policy gradient theorem [Sut+99] describes a quantity proportional to $\nabla_\theta V_{\pi_\theta}$, which is easier to obtain. It is given by

$$\nabla_\theta V_{\pi_\theta}(s_0) \propto \sum_{s \in S} \mu(s) \sum_{a \in A} Q_{\pi_\theta}(s, a) \nabla_\theta \pi_\theta(a|s), \quad (14)$$

where $\mu(s)$ is a function that expresses the fraction of time that is spend in state $s$. An concrete instance of this idea is the REINFORCE algorithm [Wil92], where a Monte Carlo method is used to estimate the quantity described in the equation above. Furthermore, the training procedure can be stabilized by introducing a suitable baseline function that reduces the variance of the expected return [Zha+11].

Overall, there are several extensions and modifications of the described concepts. One method worth mentioning is the actor-critic approach [KT03], which combines ideas form policy gradient and value functions. As for smaller modifications, there is double $Q$-learning, which introduces an additional target action-value function to reduce some bias caused by the standard $Q$-learning
Similarly, the introduction of an experience replay buffer [Lin92] should improve stability and sample efficiency. This finally leads to offline or batch RL [EGW05], where the agent is not allowed to directly interact with the environment. Instead, it only has access to a set of previously collected experiences. This formulation is especially relevant in practice, as generating data is sometimes quite expensive. There is still a wide range of topics this summary did not touch. Where necessary, additional details will also be introduced in the upcoming chapters. For a more broad introduction to the topic one can refer to Ref. [SB18], more recent developments are e.g. reviewed in Refs. [Aru+17; NLH20].

3 The Quantum Computing Paradigm

The foundations of QC were established at the beginning 20th century when the modern theory of quantum physics was developed. Benioff and Feynman proposed the idea of taking advantage of quantum mechanical systems for computing in the early 1980s [Ben80; Fey82]. QC challenges the strong Church-Turing hypothesis, as it potentially provides efficient solutions to classically intractable problems [NL16]. This section gives a pragmatic introduction to the basics of QC, and also provides an extension to quantum machine learning (QML) (here understood as ML with VQCs as a new class of models) with a focus on QRL.

Single and Multi-Qubit Systems

Similar to RL, notation and conventions regarding quantum computing vary quite a bit throughout the literature. Regarding notation, we closely follow the textbook by Nielsen and Chuang [NL16]. For the moment, let us consider the basic unit of information for classical information processing. A single bit is either in state $0$ or state $1$, consequently, a sequence of $n$ bits can represent $2^n$ unique values. Obviously, the bit register can only be in one of these $2^n$ states at any point in time.

A qubit is the quantum version of a bit. We use the Dirac notation [NL16] to define $|0\rangle$ and $|1\rangle$ as two distinct, orthogonal states of the qubit system. These basis states span a 2-dimensional Hilbert space $\mathcal{H} \cong \mathbb{C}^2$, which contains all 1-qubit (pure) quantum states. The qubits are subject to the laws of quantum mechanics and can be realized with, e.g., spin systems of subatomic particles [PMV02], ion traps [BBA14], neutral atoms [SWM10], or superconducting circuits [YN06]. This gives rise to some interesting properties. In fact, a qubit can not only be in either state $|0\rangle$ or $|1\rangle$, but in a superposition of both. An arbitrary 1-qubit state is given as

$$|\psi\rangle = \alpha |0\rangle + \beta |1\rangle.$$  \hspace{1cm} (15)

The amplitudes $\alpha$ and $\beta$ are complex numbers, which must satisfy $|\alpha|^2 + |\beta|^2 = 1$. To get a nice visual representation, Eq. (15) can be reformulated as

$$|\psi\rangle = e^{i\gamma} \left( \cos \frac{\theta}{2} |0\rangle + e^{i\phi} \sin \frac{\theta}{2} |1\rangle \right),$$  \hspace{1cm} (16)

with $\gamma, \theta, \phi \in \mathbb{R}$. As any global phase has no observable effect [NL16], the prefactor $e^{i\gamma}$ in Eq. (16) can be omitted. This representation makes it possible to visualize the state of a 1-qubit system on the surface of the Bloch sphere, see Fig. 3. The north and south poles w.r.t. the $z$-axis correspond to the basis states $|0\rangle$ and $|1\rangle$, which are also referred to as computational basis states of a single qubit. Another, less commonly used basis is given by the poles w.r.t. the $x$-axis, the elements are related by $|+\rangle = \frac{|0\rangle + |1\rangle}{\sqrt{2}}$ and $|-\rangle = \frac{|0\rangle - |1\rangle}{\sqrt{2}}$. Similarly, one could also use $|R\rangle = \frac{|0\rangle + i|1\rangle}{\sqrt{2}}$ and $|L\rangle = \frac{|0\rangle - i|1\rangle}{\sqrt{2}}$. An alternative representation associates quantum states with amplitude vectors:

$$|0\rangle \rightarrow \begin{bmatrix} 1 \\ 0 \end{bmatrix} \text{ and } |1\rangle \rightarrow \begin{bmatrix} 0 \\ 1 \end{bmatrix}$$  \hspace{1cm} (17)
Figure 3: Bloch sphere representation of a 1-qubit state.

Multiple-qubit systems are the point where things get interesting. An \( n \)-qubit system gives access to the \( 2^n \)-dimensional Hilbert space, in which an arbitrary pure quantum state is defined as
\[
|\psi\rangle = c_0 |00\cdots00\rangle + c_1 |00\cdots01\rangle + \ldots + c_{2^n-1} |11\cdots11\rangle,
\]
with \( c_i \in \mathbb{C} \) and \( \sum_{i=0}^{2^n-1} |c_i|^2 = 1 \). The basis states, e.g. \( |00\cdots01\rangle = |0\rangle \otimes |0\rangle \otimes \cdots \otimes |0\rangle \otimes |1\rangle \), consist of tensor products of the individual qubits. The state \( |\psi\rangle \to [c_0, c_1, \ldots, c_{N-1}] \) possesses \( N = 2^n \) complex amplitudes, whose absolute squared values must sum up to one. Due to the principle of superposition, an \( n \)-qubit system is able to encode and process information scaling in \( \mathcal{O}(2^n) \), while for a classical setting, it is limited to \( \mathcal{O}(n) \).

**Evolution of Closed Quantum Systems**

In order for computation to be possible, there must be some method to manipulate quantum states. Exactly this is achieved by *operators* acting on the Hilbert space \( \mathcal{H} \). By definition, all operators, which describe the time evolution of a closed quantum system are reversible. Hence, they can be represented as unitary matrices, i.e., for an operator \( U \) it must hold that \( U^\dagger U = I \). This constraint also conveys length preserving properties, i.e., applying a unitary operator to a quantum state will again yield a valid quantum state satisfying Eq. (18).

In the following, explicit matrix representations of operators are specified in the computational basis. Starting simple, consider the bit-flip operator \( \sigma_x \). This operator just flips the amplitudes of the \( |0\rangle \) and \( |1\rangle \) basis state, on the Bloch sphere this is equivalent to a rotation by \( \pi \) about the \( x \)-axis. The corresponding operators also exist for \( y \)-axis and \( z \)-axis, in matrix notation those are given as
\[
X := \sigma_x = \begin{bmatrix} 0 & 1 \\ 1 & 0 \end{bmatrix}, \quad Y := \sigma_y = \begin{bmatrix} 0 & -i \\ i & 0 \end{bmatrix}, \quad Z := \sigma_z = \begin{bmatrix} 1 & 0 \\ 0 & -1 \end{bmatrix}.
\]

Allowing an additional degree of freedom, one can define an operator for arbitrary rotation with \( \theta \) about axis \( i \) as
\[
R_i(\theta) = e^{-i \frac{\theta}{2} \sigma_i}, \quad \text{for } i \in \{x, y, z\}.
\]

The last 1-qubit operator we introduce is the Hadamard matrix:
\[
H = \frac{1}{\sqrt{2}} \begin{bmatrix} 1 & 1 \\ 1 & -1 \end{bmatrix},
\]

\[8\]
which basically performs a change of basis with $H |0⟩ = |+⟩$ and $H |1⟩ = |−⟩$. By employing the tensor product for operators, we can extend 1-qubit operators to act on single qubits comprising a multi-qubit system. We now move to genuine multi-qubit operators, acting non-trivially on two or more qubits. For our purposes, the most relevant 2-qubit operators are the controlled $X$ ($CX$) and controlled $Z$ ($CZ$), where one qubit acts as the control and the other as the target. More concretely, the $CX$-gate flips the amplitudes of the target qubit, iff the control is in state $|1⟩$. Similar to this, the $CZ$ operator performs a conditional phase flip. The matrix notations are given by

$$CX = \begin{bmatrix} 1 & 0 & 0 & 0 \\ 0 & 1 & 0 & 0 \\ 0 & 0 & 0 & 1 \\ 0 & 0 & 1 & 0 \end{bmatrix} \quad \text{and} \quad CZ = \begin{bmatrix} 1 & 0 & 0 & 0 \\ 0 & 1 & 0 & 0 \\ 0 & 0 & 1 & 0 \\ 0 & 0 & 0 & -1 \end{bmatrix}. \quad (22)$$

Quantum circuit diagrams are a nice way to visualize what is going on in a quantum algorithm. The individual qubits are represented as wires, where the order of operators, also called gates, is defined by their relative position. To be more precise, the top wire gets associated with the leftmost qubit. A few common circuit symbols for the operators introduced so far are depicted in Fig. 4.

Extracting Classical Information via Measurements

In classical computing, it is trivial to observe the exact states of all bits. For quantum systems, in order to extract information, an observable quantity has to be measured. To build the bridge to quantum computing, for each physical observable there exists a Hermitian operator $O$ [NL16], i.e., it holds $O^\dagger = O$. The eigenstates of $O$ define a basis of the quantum system’s Hilbert space.

Once an observable $O$ is measured, the corresponding measurement device outputs an eigenvalue of $O$. The post-measurement state of the system is given by the eigenstate corresponding to the eigenvalue that is measured. The most commonly used observable might be Pauli-$Z$, which corresponds to a measurement in the computational basis for a single qubit, see also Eq. (19). It has eigenvalues $\lambda_1 = +1$, $\lambda_2 = -1$ and corresponding eigenstates $v_1 = [1 \ 0]^t$, $v_2 = [0 \ 1]^t$.

The consequences for quantum computing are quite sobering, as observing superpositions w.r.t. the basis defined by the observable is impossible. Rather, one of the postulates of quantum mechanics states the Born rule, which defines a probabilistic relationship between quantum state and measurement output. Let $|0⟩$, $|1⟩$, ..., $|N − 1⟩$ be the basis defined by observable $O$ and $c_0$, $c_1$, ..., $c_N$ the corresponding amplitudes of state $|ψ⟩$ expressed in this basis. It holds, that measuring $O$ will result in the measurement outcome $\lambda_i$ with probability $|c_i|^2$. Consequently, having obtained $\lambda_i$, the post-measurement state of the system is $|i⟩$.

The first algorithm claiming provable quantum advantage, i.e., an improvement w.r.t. some complexity metric compared to any classical approach, was published in 1992 by Deutsch and Josza [DJ92] for a specially constructed problem. Most famous might be Shor’s algorithm [Sho97],
which provides an exponential speedup for tasks like prime factorization. Unfortunately, it requires large-scale, fault-tolerant and error-corrected quantum computers. All current hardware can be considered NISQ devices, which makes the execution of these algorithms infeasible. Despite this, the first claim of experimental quantum advantage was published just two years ago [Aru+19]. Yet, the considered problem was quite far from general practical applicability. A demonstration for achievable quantum supremacy on a practically relevant problem has still to be given. There are some promising candidates like quantum chemistry and material science. Recently, ideas have been put forward on combining quantum computing and machine learning [Ben+20; SP18]. These algorithms are expected to bypass at least some of the problems with execution on presently available NISQ hardware.

Quantum Machine Learning with Variational Quantum Circuits

The research on QML just really took off in the last two decades, yet there exists already a variety of approaches. As a rough clue, the hoped-for benefit of QML relies, to a large extent, on the access to the high dimensional Hilbert space granted by quantum systems. Here, we want to briefly collect the background for the summaries of VQC-based QRL approaches in Subsec. 4.2.

QML frequently deals with expectation values of quantum measurements. The expectation value of an observable $O$ w.r.t. the quantum state $|\psi\rangle$ is denoted as

$$\langle O \rangle_{\psi} := \langle \psi | O | \psi \rangle.$$  \hfill (23)

While VQCs define a new class of ML models, one can make the case for the loose analogy to NNs, where the relation of in- and output depends on a set of weights. An example for a parameterized quantum operator is given in Eq. (20). The corresponding gate applies a rotation about a specific axis by some angle $\theta_0$. Multiple rotation gates form a quantum circuit, where $\theta$ summarizes all free parameters. Varying these values gives the possibility to determine the evolution of the quantum system. Let $U_\theta$ denote the corresponding unitary. An schematic example of a VQC is displayed in Fig. 5. Most RL tasks use the concept of states, based on which an informed decision should be taken. This state information is encoded into the quantum system with an appropriate feature map. In general, the inputs $s$ are pre-processed with some mapping function $\Phi$. The results $\Phi(s)$ can be neatly integrated into the quantum circuit via the unitary $U_{\Phi(s)}$. To enhance the expressive power of the VQC, one can use more sophisticated data encoding routines like data re-uploading [Pér+20] or incremental data-uploading [Per+22]. Eventually, some observable has to be measured. A common choice is the computational basis with $O = Z^\otimes n$. Overall, the output of the VQC-model can be described as

$$\langle O \rangle_{s,\theta} := \langle 0 | \left( U_\theta U_{\Phi(s)} \right)^\dagger OU_\theta U_{\Phi(s)} | 0 \rangle.$$  \hfill (24)

For most tasks, this value is post-processed using some function $f$. Keeping things as general as possible, one can define a loss function $L$ on $f\left( \langle O \rangle_{s,\theta} \right)$ (based on the concrete problem at hand). The update of the parameters can be performed using, e.g., gradient-based techniques:

$$\theta \leftarrow \theta + \alpha \cdot \nabla_\theta L \left( f\left( \langle O \rangle_{s,\theta} \right) \right)$$  \hfill (25)

The required gradient can be obtained using the parameter-shift rule [Cro19; Wie+22b].
4 Quantum Reinforcement Learning Algorithms

In QML, there are approaches that either aim to stabilize the coherent function of the QPU using ML methods, or use the structure of a hybrid variational algorithm for ML purposes. Very often, RL is used to generate a solution for a quantum control problem, e.g., to learn quantum error correction strategies [Fös+18] or to generate control policies at a lower level [Zha+19; Dal+20]. Other work considers RL as the optimizer of a variational quantum algorithm (VQA) [Kha+19; Kha+20]. While this represents a fascinating research topic in itself, here we will focus on the application of QRL algorithms for solving specific tasks, be they classical or quantum. Research in the field of QML has so far mostly focused on supervised and unsupervised learning. However, the literature already proposes quite a few theoretical concepts and even some small-scale experimental realizations for QRL. Recent developments mostly focus on employing VQCs as function approximators. When transferring from RL to QRL, i.e., the ‘quantization’ of the RL paradigm, there are various possibilities of how quantum computing enters the game. This has led to the development of different QRL variants. A few works exist, that review current progress in QRL [KSG21; ML22]. There is also recent work towards a fair comparison of RL and QRL in restricted settings [MK21; Fra+22].

Quantum-Inspired Approaches. The earliest idea for combining RL with a quantum routine relies on the method of amplitude amplification, as it is used in Grover-type search algorithms [CDC06; Don+08; Don+06; CD08; Li+20a; Nir+21; LAD21; Cho+22]. Several qubit registers embed the states and actions relevant for the RL system in a suitable Hilbert space. Starting from a uniform superposition, amplitudes favored by the reward or the value function are selectively amplified. The action selection is based on Born’s rule, i.e., a measurement is carried out on the qubit register with regard to the ‘action-basis’. The algorithm was also investigated independently of QPUs [Don+12] and recently further developed [GH19]. As it turns out, these early variants should rather be considered a set of QiRL algorithms, that do not offer an intrinsic potential for quantum advantage. Recently, the technique was transferred to sampling from the experience replay buffer in Q-learning [Wei+22]. A summary and review of this type of QiRL can be found in Subsec. 4.1.

VQC-Based Function Approximation. In DRL, deep neural networks (DNNs) are employed as powerful function approximators. Typically, the approximation either happens in policy space (actor), in value space (critic), or both, resulting in so-called actor-critic approaches. Recently, VQCs were proposed and analyzed in their role as function approximators in the RL setting. On the one hand, this approach basically replaces a more or less well understood heuristic with a poorly understood heuristic. For the quantum heuristic many open questions regarding computational power, scalability and trainability remain. On the other hand, VQCs nonetheless have spurred the hope for quantum advantage already with NISQ devices. The earliest work in this direction introduces VQC-based Q-learning in Ref. [Che+20], followed by extensions in Refs. [LS20; LS21; SJD22]. Adoptions to policy approximation were presented in Refs. [Jer+21a; SSB22]. Analogous actor-critic approaches were discussed in Refs. [Wu+20; Kwa+21], a soft actor-critic version is considered in Ref. [Lan21]. The paradigm is extended to quantum multi-agent RL in Ref. [Yun+22], with a follow-up work incorporating meta multi-agent RL in Ref. [YPK22]. Other authors follow similar ideas, but instead consider quantum components in the purely photonic paradigm [HH19a; HH19b; HH19c; SH20]. More recent work by Ref. [Kim+21] proposes an extension to partially observable environments. The usual gradient-based parameter update is replaced with evolutionary optimization in Ref. [Che+22]. Ref. [Hsi+22] proposes the usage of VQCs with no entanglement structure. In Ref. [San+22a], multiple VQCs were employed as subroutines of a big classical model with a so-called attention mechanism. More detailed summaries and reviews are conducted in Subsec. 4.2.

Projective Simulation. Another QRL method is based on projective simulation (PS), which in the broadest sense is a particular learning paradigm and similar in spirit to RL [BD12]. Based on experiences made through interaction with the environment, a memory network is created by the
agent. The network has a directed structure with adaptive weights between the nodes of the network. The learning process and action selection are based on a random process (more precisely, a random walk) on the graph of the network, with the transition probabilities between nodes being given by the respective adaptive weights. PS can be ‘quantized’ by replacing the random walk with a so-called quantum random walk [Pap+14; Mel+17]. A formal analysis of convergence properties was given in Ref. [Boy+20]. In fact, there is already work on a proof-of-principle implementation in the laboratory [DFB15; Sri+18]. Possible quantum advantages over classical PS lie in the acceleration of the process of action selection, also referred to as deliberation in the literature. A more detailed summary is provided in Subsec. 4.3.

Quantum Boltzmann Machines. Another line of research proposes to use Boltzmann machines as function approximators. These models are assumed to be advantageous compared to typical NNs in environments with large action spaces. Ref. [Jer+21b] demonstrates, that Boltzmann machines are closely related to energy-based models. For specific instances, those allow for a quantum representation, which enables potential quantum speed-up for post-NISQ devices. A similar concept is also proposed for the annealing-based QC paradigm [Cra+18]. A summary of these ideas can be found in Subsec. 4.4.

Quantum Subroutines. Another approach to go from RL to QRL replaces certain subroutines in existing RL approaches. One idea is to replace policy or value iteration with some quantum-enhanced analogues. While this approach is limited to universal, fault-tolerant and error-corrected quantum hardware, several such algorithms have been proposed and analyzed [Wie21; Wie+22a; Wan+21a; CKP22]. Most importantly, these algorithms come with guarantees regarding speed-up, compared to their classical counterparts. QRL in these settings is often limited to the tabular case and assumes a quantum version of the RL environment, i.e., oracle access. Our summaries and reviews can be found in Subsec. 4.5.

Full-Quantum Formulation. An approach which not only ‘quantizes’ certain subroutines, but all components of the pipeline, is considered in Refs. [DTB15; DTB16; DTB17]. Extensions [HDW21; HW22], applied to specific problems [Wan+21b; Wan+22], and small-scale experimental realizations [Sag+21a] were presented. An alternative route to fully quantized QRL was taken in [Cor18]. For our review of this line of research, see Subsec. 4.6.

Various Concepts. For the sake of completeness, we mention different approaches found in the literature. We note, however, that we did not pursue a detailed review for those works, typically because we focused on what we identified as the most considered lines of research. While some of the works listed in the following simply do not fit directly with the learning-based QRL approach, for others it might not seem obvious how to generalize their particular setting to a broader class of problems. While quantum algorithms for dynamic programming have been discussed [Ron19; Amb+19], it currently remains unclear how to move from dynamic programming to a learning-based approach such as RL. Similarly, quantum algorithms have been employed to solve planning tasks [NW05], but again the transfer to a learning-based approach is far from obvious. A series of papers discussed QRL in the setting of photonic circuits, see Refs. [Fia+20; Lam21; Sag+21b]. The corresponding variant for superconducting qubits has been discussed in Ref. [Lam17]. Another approach, which we did not review in detail, is given by combining RL with the paradigm of quantum annealing [Neu+18; AHF20; Neu+20]. Hybrid quantum-classical strategies have been considered for a series of specialized tasks [PPR20]. Similarly, various interpretations of QRL for specialized tasks in the quantum domain exist [Alv+16; Alv+18; Cár+18; Alb+20; Oli+20; Bha+19]. Different approaches have been proposed for combining RL with quantum walks [Dal+22; Che+19]. There has also been work on extending VQC-based algorithms with post-NISQ amplitude amplification [San+22b]. From a purely theoretical perspective, an analysis of partially observable Markov decision problems in the quantum setting was given in Ref. [BBA14]. Further work on optimization tasks rather than RL, such as Ref. [Jaš+19], have also not been reviewed in detail.
4.1 Quantum-Inspired Reinforcement Learning based on Amplitude Amplification

| Citation     | First Author | Title                                                                 |
|--------------|--------------|----------------------------------------------------------------------|
| [Don+08]     | D. Dong      | Quantum reinforcement learning                                         |
| [Don+06]     | D. Dong      | Quantum mechanics helps in learning for more intelligent robots       |
| [CD08]       | C.-L. Chen   | Superposition-Inspired Reinforcement Learning and Quantum Reinforcement Learning |
| [CDC06]      | C.-L. Chen   | Quantum computation for action selection using reinforcement learning  |
| [Don+12]     | D. Dong      | Robust Quantum-Inspired Reinforcement Learning for Robot Navigation   |
| [GH19]       | M. Ganger    | Quantum Multiple Q-Learning                                           |
| [Li+20a]     | J.-A. Li     | Quantum reinforcement learning during human decision-making           |
| [LAD21]      | J.-A. Li     | Intelligent Trajectory Planning in UAV-Mounted Wireless Networks: A Quantum-Inspired Reinforcement Learning Perspective |
| [Cho+22]     | B. Cho       | Quantum bandit with amplitude amplification exploration in an adversarial environment |
| [Nir+21]     | D. Niraula   | Quantum deep reinforcement learning for clinical decision support in oncology: application to adaptive radiotherapy |
| [Wei+22]     | Q. Wei       | Deep Reinforcement Learning With Quantum-Inspired Experience Replay   |

Table 1: Work considered for “QiRL based on amplitude Amplification” (Subsec. 4.1)

Quantum reinforcement learning, Dong et al. (2008) and related work

Summary. Ref. [Don+08] discusses a new RL algorithm that is inspired by the superposition principle of quantum mechanics. The authors propose an algorithm that modifies the action-selection procedure and balances exploration and exploitation in a novel way. The authors present their ideas in modified form in a sequence of papers, see Refs. [Don+08; Don+06; CD08; CDC06; Don+12; GH19; Li+20a; LAD21]. We note that this work should be viewed as a QiRL approach.

Algorithmic Concepts and Extensions. Initially, the algorithm is formulated as merely quantum inspired in Ref. [CD08] (i.e., it is developed for a classical computer that simulates a quantum superposition). The motivation is to design an algorithm with better exploration-exploitation trade-off compared to e.g. $\epsilon$-greedy action selection. The underlying routine is a modification of temporal difference (TD), more concretely TD(0) in the following way: For each state the set of possible actions is in a ‘superposition’ and the agent (in state $s$) now selects an action with a given probability. The action is taken and the new state $s'$ and reward $r$ is observed. Afterwards, the probability of the taken action is increased by $k(r + V(s'))$, where $V(s')$ is the value function of state $s'$, and $k$ is a hyperparameter. The term $r + V(s')$ samples a quantity similar to $Q(s,a)$. Consequently, the update creates a probability distribution, where for a given state the probability to select an action
increases as the value of $Q(s, a)$ increases. Therefore, this action selection process corresponds to sampling from a stochastic policy dependent on the value of the state-action pairs.

Now the algorithm is translated to be run on a quantum computer. The stochastic policy is replaced by a quantum superposition. That is, for each state $s$ the possible actions are represented by the eigenstates of some observable and a superposition of these states is created. If the observable is measured, the state will collapse to an eigenstate associated with an action which will be taken by the agent and therefore constitutes the selection process. After receiving the reward and the new state, the Grover operator is applied $L = \min\{k(r + V(s')), L_{\text{max}}\}$ times to a copy of the superposition state to enhance the amplitude corresponding to the previous selected action. The variable $L_{\text{max}}$ guarantees that the Grover operator is not applied too many times. Note that repeated application of the procedure requires a new copy of the state after each measurement. Due to the no cloning theorem, this could be realized by many different independent copies of the initial memory, or by a purely classical representation of the states. The latter realization reduces the algorithm to the initial proposal of a quantum-inspired action selection process.

In Ref. [Don+12] the QiRL algorithm is applied to robot navigation. It is stated explicitly that QiRL is a classical action-selection method that differs from the ideas of QRL, which in principle could benefit from a quantum computer. In Ref. [GH19] the algorithms are generalized to $Q$-learning and double- and multiple $Q$-learning. Also these approaches should be understood in the context of QiRL. Finally, Refs. [Li+20a; Nir+21] apply QiRL to human decision making behavior. Recently, the quantum-inspired approach to action selection in RL was transferred to experience replay buffer sampling in $Q$-learning [Wei+22].

Remarks. Although it is mentioned in Ref. [Don+08; Don+06; CD08; CDC06; Don+12; GH19; Li+20a; LAD21; Cho+22] that the whole algorithm could be run in quantum superposition on a quantum device, no details of such kind of genuine QRL algorithm are given. Overall, it is unclear if such an algorithm might exist. Indeed, subsequent work focuses on the QiRL paradigm.

The claims made in Refs. [Don+08; Don+06; CD08; CDC06; Don+12; GH19; Li+20a; LAD21; Cho+22] can be summarized as follows: speed-up in learning by better balancing exploration-exploitation; less GPU power needed on classical computer compared to algorithms like classical $Q$-learning; more robust against changes of learning rate. More experiments on larger environments for deeper insights into the scaling of the algorithm and a rigorous complexity analysis would be an interesting topic for future work.

4.2 Quantum Reinforcement Learning with Variational Quantum Circuits

This section summarizes the state-of-the-art on VQC-based RL. Several ideas have been proposed in this field, with extensions in different directions. Their common ground is the usage of a VQC as parameterized function approximator.

The typical hybrid pipeline is summarized in Fig. 6. It was originally proposed for $Q$-function approximation by Chen et al. [Che+20] and extended to policy approximation by Jerbi et al. [Jer+21a]. Other work proposes several modifications to this pipeline, which we will describe in the respective summaries. The algorithm must be understood as hybrid, as a lot of the work, especially the optimization, is executed on classical hardware. The agent observes the current state of the environment $s_t$, and applies some pre-processing $\phi$. The result is encoded using the feature map $U_{\phi(s)}$. With the current variational parameters $\theta_t$, a quantum state is prepared and a (potentially action-dependent) observable $O_a$ is measured. The expectation value $\langle O_a \rangle_{s, \theta}$ can be post-processed to represent, e.g., a state-action value function $Q_\theta(s, a)$, or the policy $\pi_\theta(a|s)$. Depending on the instance, the agent employs this function to sample an action $a_t$, and executes it in the environment. The reward $r_t$ (and potentially also the consecutive state $s_{t+1}$) is observed by the classical optimizer. To enable gradient-based parameter updates, an additional hybrid module uses the parameter-shift rule [Cro19; Wie+22b] to compute the gradients of the VQC outputs w.r.t. the variational parameters $\theta_t$. The classical optimizer determines the new parameter set $\theta_{t+1}$ and instantiates the VQC.
with these updated parameters. This overall iterative procedure of environment interaction, function approximation, and parameter update is repeated for several episodes, in the same way as for, e.g., DRL.

Unfortunately, thus far there is no guaranteed quantum advantage for this approach, apart from some cryptography inspired artificial datasets [Jer+21a; SJD22]. However, several of the papers and preprints summarized below demonstrate promising experimental results.

| Citation | First Author | Title |
|----------|--------------|-------|
| [Che+20] | S. Y.-C. Chen | Variational Quantum Circuits for Deep Reinforcement Learning |
| [LS20]   | O. Lockwood  | Reinforcement Learning with Quantum Variational Circuits |
| [LS21]   | O. Lockwood  | Playing Atari with Hybrid Quantum-Classical Reinforcement Learning |
| [SJD22]  | A. Skolik     | Quantum agents in the Gym: a variational quantum algorithm for deep Q-learning |
| [Jer+21a]| S. Jerbi      | Parameterized Quantum Policies for Reinforcement Learning |
| [LS22]   | A. Sequeira   | Variational Quantum Policy Gradients with an Application to Quantum Control |
| [Wu+20]  | S. Wu         | Quantum reinforcement learning in continuous action space |
| [Kwa+21] | Y. Kwak       | Introduction to Quantum Reinforcement Learning: Theory and PennyLane-based Implementation |
| [Lan21]  | Q. Lan        | Variational Quantum Soft Actor-Critic |
| [Yun+22] | W. J. Yun     | Quantum Multi-Agent Reinforcement Learning via Variational Quantum Circuit Design |
| [YPK22]  | W. J. Yun     | Quantum Multi-Agent Meta Reinforcement Learning |
| [HH19a]  | W. Hu         | Q Learning with Quantum Neural Networks |
| [HH19c]  | W. Hu         | Reinforcement Learning with Deep Quantum Neural Networks |
| [HH19b]  | W. Hu         | Distributional Reinforcement Learning with Quantum Neural Networks |
| [SH20]   | E. Sorensen   | Practical Meta-Reinforcement Learning of Evolutionary Strategy with Quantum Neural Networks for Stock Trading |
| [Kim+21] | T. Kimura     | Variational Quantum Circuit-Based Reinforcement Learning for POMDP and Experimental Implementation |
| [Che22]  | S. Y.-C. Chen | Variational quantum reinforcement learning via evolutionary optimization |
| [Hsi+22] | J.-Y. Hsiao   | Unentangled quantum reinforcement learning agents in the OpenAI Gym |
| [San+22a]| F. Sanches    | Short quantum circuits in reinforcement learning policies for the vehicle routing problem |

Table 2: Work considered for “QRL with Variational Quantum Circuits” (Subsec. 4.2)
Variational Quantum Circuits for Deep Reinforcement Learning, Chen et al. (2020)

Summary. This paper by Chen et al. [Che+20] represents the first attempt to utilize VQCs for RL. This is done in the context of using VQCs as function approximators for the state-action value function. The authors perform simulations on simple benchmark environments and report.

Hybrid Algorithm. The algorithm is inspired by deep Q-learning (DQL) [Mni+15], where a DNN represents the Q-function. The authors replace the DNN by a VQC. The update is performed w.r.t. the mean square error (MSE) loss function $L(\theta) = \mathbb{E}[(r_t + \gamma \cdot \max_{a'} Q_\theta'(s_{t+1}, a') - Q_\theta(s_t, a_t))^2]$ using, e.g., gradient descent. Additionally, experience replay and target networks (second set of parameters $\theta'$) are employed to address the instabilities stemming from bootstrapping the value function, forming a double deep Q-learning (DDQL) algorithm. Fig. 7 gives the complete algorithm.

VQC Architecture. The feature map uses simple computational basis encoding on individual qubits. More concretely, the RL state is interpreted as bitstring, which can be encoded using the identity $R_z(\pi) R_x(\pi) |0\rangle = |1\rangle$. The entanglement structure connects nearest neighbors with CZ gates. The variational parameters are incorporated in single qubit rotations about the x, y, and z axis. The state-action value is decoded by measuring Pauli-Z observables on a number of qubits, that corresponds to the number of actions in the environment. The full VQC is visualized in Fig. 8.

Experimental Results and Discussion. The proposed VQC-DQL algorithm is simulated for two environments. The first one is FrozenLake, with 16 states and 4 actions. The second one is CognitiveRadio, which is adapted to VQCs sizes of 2 to 5 qubits. The authors report that their VQC-based agent performs at least equally well as a NN. Moreover, they claim that this requires fewer parameters (about one order of magnitude compared to DNNs), which points towards potential quantum advantage. The model is tested on actual quantum hardware with competitive results.

Remarks. The employed encoding scheme (computational basis encoding) could be simplified by
Algorithm 1: Variational Quantum Deep Q Learning

Initialize replay memory \( \mathcal{D} \) to capacity \( N \)
Initialize action-value function quantum circuit \( Q \) with random parameters

for episode = 1, 2, …, \( M \) do
    Initialize state \( s_1 \) and encode into the quantum state
    for \( t = 1, 2, \ldots, T \) do
        With probability \( \epsilon \) select a random action \( a_t \)
        otherwise select \( a_t = \text{argmax}_a Q(s_t, a_t; \theta) \) from the output of the quantum circuit
        Execute action \( a_t \) in emulator and observe reward \( r_t \) and next state \( s_{t+1} \)
        Store transition \( (s_t, a_t, r_t, s_{t+1}) \) in \( \mathcal{D} \)
        Sample random minibatch of transitions \( (s_j, a_j, r_j, s_{j+1}) \) from \( \mathcal{D} \)
        Set \( y_j = \begin{cases} r_j & \text{for terminal } s_{j+1} \\ r_j + \gamma \text{max}_a Q(s_{j+1}, a'; \theta) & \text{for non-terminal } s_{j+1} \end{cases} \)
        Perform a gradient descent step on \( (y_j - Q(s_j, a_j; \theta))^2 \)
    end for
end for

Figure 7: Hybrid algorithm proposed by and taken from Chen et al. \cite{Che+20}; This algorithm uses a VQC to approximate the state-action value function and follows the typical steps of DQL. Note, that the authors notation for the \( Q \)-function slightly deviates from our conventions.

Figure 8: VQC proposed by and taken from Chen et al. \cite{Che+20}; The \( R_x \) and \( R_z \) gates are used for state encoding. Several parameterized layers (dashed box) are repeated to form the \( Q \)-function approximator. The values of the function are decoded using 1-qubit Pauli-Z observables.

omitting the \( R_Z \) rotations, as these only introduce a global phase. The CognitiveRadio environment might be oversimplified. We also note that the claim on reduced parameter count should be substantiated by experiments with environments of different scale.

Algorithmic Characteristics - Chen et al. \cite{Che+20}

| Environment                          | Algorithm Type | Quantum Component | State Space | Action Space | Qubits | Parameterized Gates\(^a\) |
|--------------------------------------|----------------|-------------------|-------------|--------------|--------|--------------------------|
| FrozenLake (OpenAI Gym)             | DDQL           | \( Q \)-function  | discrete 16 | discrete 4   | 4      | 4 × 2 (encoding) 4 × 4 × 3 (weights) |
| CognitiveRadio (see \cite{Che+20})  | DDQL           | \( Q \)-function  | \( n^2 \)   | \( n \)       | \( n \) | \( n \) × 2 (encoding) \( n \) × 4 × 3 (weights) |

\(^a\) encoding gates: \( \text{qubits} \times \text{per qubit} \); variational gates: \( \text{qubits} \times \text{layers} \times \text{per qubit per layer} \);
Reinforcement Learning with Quantum Variational Circuits, Lockwood and Si (2020)

Summary. The work by Lockwood and Si [LS20] modifies several aspects of the routine proposed by Chen et al. [Che+20]. Most importantly, they introduce two new encoding schemes to deal with a continuous state space.

Modification of Architecture. The first proposed encoding is denoted as scaled encoding. It scales the RL state values to the range $[0, 2\pi)$, which are then encoded using some 1-qubit parameterized rotations. The second one (so-called directional encoding) only encodes the sign of the value. More concretely, if a state variable is positive, $R_x$ and $R_z$ rotations by $\pi$ are applied to the encoding qubit (following a similar idea as the computational state encoding [Che+20]).

The architecture for the variational layer consists of an entangling block (nearest-neighbor CX gates) and parameterized 1-qubit rotations about $x$, $y$, and $z$ axis. This block is repeated three times. For decoding the state-action value, the authors employ two different strategies. The first one feeds the measurement result into a classical fully-connected layer where the number of outputs corresponds to the number of possible actions. In the other case, a so-called quantum pooling operation, condenses the information of the quantum state into a subset of the qubits [CCL19]. This allows for a more flexible architecture, independent of the number of actions in the environment.

Experimental Results. The proposed algorithm and the encoding schemes are benchmarked on the CartPole and Blackjack environment. While the former one uses a combination of scaled and directional encoding, the second one only employs scaled encoding. Their findings agree with those reported previously in the literature, namely that VQC-based models achieve similar performance to NN-based function approximators. As also stated by Chen et al. [Che+20], the usage of VQCs reduces the required parameter complexity.

Remarks. While the scaled encoding should be a sound choice, the directional encoding could be inappropriate for most environments. Usually, not only the sign of a specific state is relevant, but the concrete state contains relevant information. With this encoding, this information is lost, which should lead to a drop in performance for more complex environments. As stated previously, the reduced parameter complexity should be investigated for larger problem instances.

Algorithmic Characteristics - Lockwood and Si [LS20]

| Environment       | Algorithm Type | Quantum Component | State Space | Action Space | Qubits | Parameterized Gates $^a$ |
|-------------------|----------------|-------------------|-------------|--------------|--------|---------------------------|
| CartPole (OpenAI Gym) | DDQL           | $Q$-function      | continuous 4-dim | discrete 2   | 4      | $4 \times 2$ (encoding) $4 \times 3 \times 3$ (weights) |
| Blackjack (OpenAI Gym) | DDQL           | $Q$-function      | discrete 31x11x2 | discrete 2   | 3      | $3 \times 2$ (encoding) $3 \times 3 \times 3$ (weights) |

$^a$ encoding gates: qubits $\times$ per_qubit; variational gates: qubits $\times$ layers $\times$ per_qubit $\times$ per_layer;

Playing Atari with Hybrid Quantum-Classical Reinforcement Learning, Lockwood and Si (2021)

Summary. This work by Lockwood and Si [LS21] extends their previous paper [LS20], which, in turn, was based on Chen et al. [Che+20], where $Q$-learning with VQC function approximation has been introduced. The paper considers the Atari environments Pong and Breakout, with continuous state space of dimensionality 28,224 (the observations are cropped and converted to images with 84x84x4 pixels). This environment complexity is not tractable with previously introduced encoding schemes, which require one qubit for each dimension. The proposed workaround uses a classical NN to reduce the state dimensionality before encoding it into the VQC.
**Underlying Algorithm and Simulation.** Similar to Refs. [Che+20; LS20], the concept of DDQL is used. The pipeline is modified by replacing the pure VQC function approximator with a hybrid model. Several different choices are considered, the most important details are highlighted below. The training is performed in an end-to-end manner, i.e., the gradients w.r.t. the VQC parameters are propagated back through the classical encoding network.

**Model Architecture.** The VQC architecture is, as usually, composed of three parts (i.e. state encoding, variational layers, and action decoding). To encode the state, the raw data is first fed through a classical NN. This outputs a number of values equal to the number of parameters in the feature map, which itself consists of 1-qubit parameterized rotations. The authors compare the performance of a densely connected and a convolutional neural network (CNN) for this task (the concrete architecture of these networks is not specified). Apart from that, encoding layers of different sizes (and therefore different number of parameters) from 5 to 15 qubits are compared.

The variational layers itself consists of two parts, where the first one is a quantum convolutional neural network (QCNN) [CCL19]. The authors state two motivations for this choice: First, it should help capture the spatial structure of the input images (but it is unclear, whether the encoding part retains the spatial structure). Second, QCNNs help to avoid barren plateaus [Pes+21] (while the experiments show no sign of barren plateaus, it is not clear if this is due to this choice, or the limited size of the employed circuits). After this QCNN there are three repetitions of entanglement gates and parameterized rotations, similar to those also used for state encoding.

The paper proposes two methods to deal with the problem of measurement for unequal number of qubits and actions. The first method performs Pauli-Z measurements on all qubits and uses an appended dense NN. Alternatively, quantum pooling operations [CCL19] are used, which subsequently compress the measurement of two qubits into one.

**Experimental Results and Discussion.** To demonstrate the basic functionality of the model, initial experiments are conducted on the CartPole environment. The results demonstrate a similar performance to Lockwood and Si [LS20]. On the two Atari environments, the paper considers 12 different hybrid architectures (dense vs. convolutional encoding, 5 vs. 10 vs. 15 qubits, dense vs. pooling decoding), which are compared to a well-established classical architecture.

It turns out, that the hybrid models are not able to learn at all. The authors state, that this is down to the lack of expressibility of the hybrid models, which only make use of about $10^4$ parameters, while the classical model uses about $10^6$. It is expected, that for more expressive models the performance improves, as learning on the much simpler CartPole environment was successful.

### Algorithmic Characteristics - Lockwood and Si [LS21]

| Environment          | Algorithm Type | Quantum Component | State Space | Action Space | Qubits | Parameterized Gates |
|----------------------|----------------|-------------------|-------------|--------------|--------|---------------------|
| CartPole (OpenAI Gym)| DDQL           | $Q$-function      | continuous 4-dim | discrete 2 | 5      | $N/A$ (classical)<sup>a</sup> $O(10^1)$ (encoding) $O(10^2)$ (weights) |
| Pong-v0 (OpenAI Gym) | DDQL           | $Q$-function      | 28224-dim<sup>b</sup> | discrete 6 | 5 to 15 | $O(10^6)$ (classical) $O(10^2)$ (encoding) $O(10^4)$ (weights) |
| Breakout-v0 (OpenAI Gym) | DDQL          | $Q$-function      | 28224-dim<sup>b</sup> | discrete 4 | 5 to 15 | $O(10^6)$ (classical) $O(10^2)$ (encoding) $O(10^4)$ (weights) |

<sup>a</sup> potentially also uses a classical NN for pre-processing, details are not stated;  
<sup>b</sup> dimensionality of feature space is reduced with a NN to fit size of feature map;
Remarks. The experiments are conducted with a restricted set of hybrid models. Consequently, the claim that these results do not demonstrate the inapplicability of QRL to more complex environments like Atari is reasonable. The assumption that this approach could be made to work on complex environments, as it succeeds on e.g. CartPole, should be sustained with additional experiments. For a modified architecture succeeding on the Atari environments, it is not completely clear, which part of the work is done by the classical and quantum part of the model. This is a typical caveat, whenever quantum and classical architectures are combined.

Quantum agents in the Gym: a variational quantum algorithm for deep $Q$-learning, Skolik et al. (2022)

Summary. This work by Skolik et al. [SJD22] proposes another instance of $Q$-learning with VQCs as function approximators. Being aware of preceding literature, the authors set out to analyze the role of architecture design, RL state encoding schemes, and observables for action decoding. With regard to the previous work, the authors remark that the CartPole environment cannot be considered solved.

Importance of Architecture Design. In terms of architecture choices, the problem of barren plateaus is emphasized: Architectures with many qubits and layers (which naively is required for high expressivity) are hard to train. Contrarily, over-parameterized architectures are easier to train, but probably less expressive and therefore less effective on a given task.

The authors chose a hardware-efficient ansatz, despite being known to run into the barren plateau problem for large circuits. For the small circuit sizes considered in the present work, the barren-plateau problem does not appear to be relevant.

Encoding Schemes. As for encoding schemes, discrete RL states are encoded in the computational basis. Continuous states are scaled to the finite interval $[-\pi/2, +\pi/2]$ by applying arctan to the raw observations. The result serves as the rotation angle for an $R_x$ rotation, which is very similar to the scaled encoding proposed by Lockwood and Si [LS20]. In order to increase expressivity w.r.t. to the input, the encoding layer can be repeated through the circuit, forming a data re-uploading structure [Pér+20]. Effectively, this allows to learn and approximate a Fourier sum of a certain order, where the order is tied to the number of repetitions of the encoding layer [SSM21]. The encoding is further modified by introducing learnable re-scaling parameters, that are multiplied with the raw states before computing the arctan.

Experimental Results and Discussion. The authors benchmark their architecture choices on the FrozenLake and CartPole environment. The performance on CartPole is compared to a small NN with the same number of parameters, which seems to be inferior.

Further, the range of $Q$-values that can be encountered in the two benchmark environments is investigated. For FrozenLake, representing the $Q$-value with the expectation values of 1-qubit $Z$-operators is sufficient. For the CartPole environment, this strategy is found not to be adaptable enough. Instead, they chose the expectation values of the parities (of 2 non-overlapping pairs of qubits) and allow for additional trainable classical weights that set the scale for the $Q$-value approximation.

Remarks. The authors emphasize the critical role of architectural choices at the outset of their manuscript. While they offer valuable insights into this topic, also open questions remain for future work in this direction. For the CartPole environment, several trainable classical weights are incorporated in the algorithm. Therefore, it is not completely clear, what part of the training is achieved by which part of the hybrid model.
Parameterized Quantum Policies for Reinforcement Learning, Jerbi et al. (2021)

Summary. The paper by Jerbi et al. [Jer+21a] starts out with a small summary of VQC-based ML models. They cite several reports of quantum advantage in the supervised and unsupervised QML. This motivates their approach to go beyond the scope of \( Q \)-function approximation [Che+20; LS20; LS21; SJD22], and use the VQC to directly approximated the policy.

Quantum Policy Gradient. After a brief recap of policy gradient methods for solving RL problems, the authors extend those ideas to a quantum policy gradient (QPG) approach. More concretely, they quantize the REINFORCE algorithm [Wil92] with value-function baselines by using VQCs as function approximators for the (stochastic) policy. The define two families of VQC-based policies: (1) A RAW-VQC policy, where the action selection follows Born’s rule. It is defined as \( \pi_\theta(a|s) = \langle P_a \rangle_{s,\theta} \), where \( P_a \) are the projectors on the elements of the computational basis. This allows action selection with only one evaluation of the quantum circuit; (2) A SOFTMAX-VQC policy, defined as \( \pi_\theta(a|s) = \frac{e^{\beta O_a}}{\sum_{a'} e^{\beta O_{a'} s,\theta}} \). The measurement result of an action-dependent observable \( O_a \) is fed into a single-parameter softmax-function, to form a PDF. The inverse-temperature parameter \( \beta \) allows to adjust the peak-width of the distribution, i.e., the greediness of the policy.

Circuit Architecture. The ansatz for the VQC is chosen to be hardware-efficient, i.e., only single and two-qubit gates. The RL state is encoded with 1-qubit rotations. To increase the expressivity of the model, the authors introduce additional learnable state-scaling parameters \( \lambda \). Those are multiplied to the rotational parameter denoting the state value, i.e., \( \lambda_i \cdot s_i \) is the value of a 1-qubit rotation. This also helps circumvent the problem of being restricted to a finite set of frequencies in such an encoding scheme [SSM21]. The feature map is repeated several times, alternating with the variational layer, which forms a data re-uploading structure [Pér+20]. A variational layer consists of \( CZ \)-gates for creating entanglement in a circular structure. The learnable parameters are used in 1-qubit parameterized rotation gates. Depending on the policy type, measurements are either conducted in the computational basis, or more complex observables are measured.

Experimental Results. Overall, all agents are able to learn meaningful behavior in the OpenAI Gym environments CartPole, MountainCar, and Acrobat. Further experiments are reported, which serve the purpose of assessing the importance of the various design choices: (1) Circuit depth increases performance and learning speed, where SOFTMAX-VQC policies outperform RAW-VQC policies in all instances; (2) Incorporating learnable state scaling parameters increases learning performance, trainable classical weights (in case of SOFTMAX-VQC) multiplied to expectation values leads to increase in performance; (3) The performance gap between RAW-VQC and softmax-VQC policies seems to stem from the ability to adjust greediness.

Provable and Empirical Quantum Advantage. To the best of our knowledge, this work is the first to corroborate the idea quantum advantage with VQCs in the RL setting. Therefore, the authors devise RL environments (based on the discrete logarithm problem (DLP)), which are supposed to

### Table: Algorithmic Characteristics - Skolik et al. [SJD22]

| Environment               | Algorithm Type | Quantum Component | State Space | Action Space | Qubits | Parameterized Gates
g | Encoding Gates: \( qubits \times \text{per}_\text{qubit} \); Variational Gates: \( qubits \times \text{layers} \times \text{per}_\text{qubit}_\text{per}_\text{layer} \); b Model incorporates classical weights after measurement, details are not stated. |
|---------------------------|----------------|-------------------|-------------|--------------|--------|--------------------------|
| CartPole (OpenAI Gym)     | DDQL           | \( Q \)-function  | continuous  | discrete     | 4      | 4 \times 1 (encoding)    |
|                           |                |                   | 4-dim       | 2            |        | 4 \times 15 \times 2 (weights) |
|                           |                |                   | 4 \times 1   |              |        | N/A (classical)          |
| FrozenLake (OpenAI Gym)   | DDQL           | \( Q \)-function  | discrete     | discrete     | 4      | 4 \times 1 (encoding)    |
|                           |                |                   | 16           | 4            |        | 4 \times 15 \times 2 (weights) |

Parameterized Gates:

- a
- b
be classically intractable. Any classical algorithm would need a number of samples that scales exponential in the problem size to achieve a low generalization error. A VQC-based algorithm with a very specific architecture only requires a polynomial amount of data. This implies an exponential advantage w.r.t. sample complexity, assuming it is infeasible to efficiently simulate the VQC on classical hardware for large problem instances. The construction of the environment is inspired by previous results from QML, where similar learning separations between classical and quantum models have been demonstrated [LAT21].

Further, the authors report numerical evidence of potential quantum advantage for environments based on expectation values sampled from VQCs. The motivation lies in the (potential) intractability of simulating the given VQC classically for large systems. More concretely, one uses a VQC to define a labeling function (in the sense of a classification task) over the domain $[0, 2\pi]^2$ (so-called SL-VQC). This synthetic classification dataset is then rephrased as a RL environment by incorporating some temporal structure (denoted as Cliffwalk-VQC). Numerically, the authors observe a performance separation of models with classical DNNs and VQC-based policies. They claim, that this is likely due to the oscillatory structure in the labeling function.

**Remarks.** While the proposal of provable quantum advantage is obviously quite encouraging, the practical realization is probably out of reach for the NISQ-era. The idea of solving the task efficiently on quantum hardware is based on Shor’s algorithm. Formulated as a VQC-based RL problem, this would require circuits of complexity far beyond current scope. We think it requires also some more large-scale experiments, to support the empirical learning separation on the SL-VQC and Cliffwalk-VQC environments. A comparison to other hybrid models [Che+20; LS20] shows, that the proposed QPG approach is superior in terms of RL performance on various environments.

### Algorithmic Characteristics - Jerbi et al. [Jer+21a]

| Environment                  | Algorithm Type | Quantum Component | State Space   | Action Space | Qubits | Parameterized Gates$^a$ |
|------------------------------|----------------|-------------------|---------------|--------------|--------|-------------------------|
| CartPole (OpenAI Gym)        | REINFORCE      | Policy            | continuous 4-dim | discrete     | 4      | 30                      |
| MountainCar (OpenAI Gym)     | REINFORCE      | Policy            | continuous 2-dim | discrete     | 2      | 36                      |
| Acrobot (OpenAI Gym)         | REINFORCE      | Policy            | continuous 2-dim | discrete     | 2      | 37                      |
| SL-VQC Cliffwalk-VQC (see [Jer+21a]) | REINFORCE      | Policy            | continuous 2-dim | discrete     | 2      | 37                      |
| CognitiveRadio (see [Che+20]) | REINFORCE      | Policy            | discrete $n^2$  | discrete $n$ | $n$    | 30 to 75 for $n = 2$ to $5$ |

$^a$ this entails encoding, scaling, and variational parameters; the SOFTMAX-VQC also uses classical parameters;

### Variational Quantum Policy Gradients with an Application to Quantum Control, Sequeira et al. (2022)

**Summary.** The article by Sequeira et al. [SSB22] proposes a quantum version of the REINFORCE algorithm with a VQC-based function approximator, very similar to Jerbi et al. [Jer+21a]. The methods are applied to the classical environments CartPole and Acrobot but also to a simple quantum control problem. It proposes an initialization technique for the variational parameters of a VQC. Following the experimental results, a quantum advantage w.r.t. the number of required parameters and trainability of the models is claimed.
Underlying Reinforcement Learning Algorithm. As in Jerbi et al. [Jer+21a], the policy is defined as \( \pi_\theta(a|s) = e^{\beta \langle O_a \rangle}/\sum_{a'} e^{\beta \langle O_{a'} \rangle} \), and REINFORCE updates are performed. Hereby, the expectation values \( \langle O_a \rangle \) for action \( a \) is defined as the expectation \( \langle \sigma_a^z \rangle \), i.e., the expectation value of 1-qubit Pauli-\( Z \) observable measured on the \( a \)-th qubit.

VQC Architecture. The architecture follows the typical three-part structure. In the beginning, the states are encoded with \( R_x \) rotations, with the state values normalized to the range \([-\pi, \pi)\). Consequently, the number of qubits has to correspond to \( \max\{|A|, |S|\} \). There are several parameterized layers (see Fig. 9) which incorporate variational parameters in \( 1 \)-qubit \( R_y \) and \( R_z \) rotations. The entanglement structure can be described as \( CX[i,(i+l) \mod n] \), where \( n \) is the number of qubits, and \( l \) the index of the layer. The measurement of \( 1 \)-qubit Pauli-\( Z \) observables is a deviation to the procedure proposed by Jerbi et al. [Jer+21a], where multi-qubit observables were used.

![VQC architecture](image)

Figure 9: VQC architecture proposed by and taken from Sequeira et al. [SSB22]; It deviates from the typical circular entanglement structure.

Complexity of Gradient Estimation. The paper gives an estimation of the required number of samples to get an \( \epsilon \)-approximation of the log-policy gradient. According to this consideration, for a success probability of \( 1 - \delta \), the number of required measurements is bounded by \( c \cdot \frac{(1-\epsilon)^2}{\epsilon^2} \cdot \log\left(\frac{k}{\delta}\right) \). Hereby, \( c \) is a constant depending on algorithmic hyperparameters and \( k \) is the number of variational parameters. Although this theoretical result is quite promising, it is not stated how to achieve such an efficient estimation using the parameter-shift rule on an actual quantum device.

Initialization Technique. There is some work proposing a technique for parameter initialization to avoid barren plateaus [Gra+19]. However, a technique to boost the overall performance has not yet been proposed. Inspired by classical ML, the authors aim to break symmetries between different neurons (as usually initialization with constant values is a bad choice). A typical strategy is to select values uniformly at random from \([-\pi, \pi)\), or drawn them following a Gaussian distribution. Inspired by the classical Glorot initialization scheme [GB10], the paper proposed to use a normal distribution \( \mathcal{N}(0, \text{std}^2) \) with \( \text{std} = g \cdot \sqrt{2/(\text{fan}_{\text{in}} + \text{fan}_{\text{out}})} \). Here, \( g \) is a constant multiplicative factor, \( \text{fan}_{\text{in}} \) is the number of embedded features, and \( \text{fan}_{\text{out}} \) is the number of computational basis measurements. This technique demonstrates some promising experimental results, but no theoretical justification is given.

Analysis of Fisher Information Spectrum. The paper analyzes the spectrum of the Fisher information matrix (FIM), which serves as a tool to quantify the trainability of a model. The empirical FIM is computed as \( F(\theta) = \frac{1}{T} \sum_{t=1}^{T} \nabla_{\theta} \log \pi(a_t|s_t, \theta) \nabla_{\theta} \log \pi(a_t|s_t, \theta)^t \). A similar analysis has also been proposed for QML [Abb+21].

The results show, that the spectrum of the FIM associated with the quantum model exhibits significantly larger averaged eigenvalues. The compared NN was optimized over several architectures, but not many details are provided in the paper. The authors conclude, that the quantum models are beneficial in terms of trainability, and might be resilient to barren plateaus.
Experimental Results and Discussion of Potential Quantum Advantage. The proposed algorithm is tested on the classical benchmark environments CartPole and Acrobot. The performance is compared to the best classical NN (it is not clear, what best means in this case, and to what extend this holds). The authors claim a significant advantage in terms of convergence speed.

Additional experiments are conducted with the proposed Quantum-Glorot initialization technique. In the two environments CartPole and Acrobot, this technique demonstrates to be beneficial in terms of convergence speed and training stability.

Finally, the experiments are extended to a QuantumControl environment. It requires to learn the mapping $|0\rangle \rightarrow |1\rangle$ via the time dependent Hamiltonian $H(t) = 4J(t)\sigma_z + h\sigma_x$. This is converted to a set of unitary gates $U(t)$, such that $|\psi_{t+1}\rangle = U(t)|\psi_t\rangle$. The reward is defined as the overlap between the prepared state and $|1\rangle$, i.e. $r_t = |\langle \psi_t|1\rangle|^2$. The agent has to decide between the two actions $0 \doteq$ no pulse and $1 \doteq$ apply pulse. The usage of a quantum environment removes the necessity of encoding classical states. Unfortunately, it is not described, how $|\psi_t\rangle$ is incorporated in the VQC (a 1-qubit parameterized circuit is apparently used to solve the task). The results on this environment suggest, that the agent is able to learn the optimal pulses in a low number of epochs.

Summarizing the experiments, the authors claim an advantage in convergence speed compared to classical approaches (questionable, as there should be NNs which perform much better). Additionally, there seems to be a clear advantage in terms of parameter complexity.

Remarks. The authors claim, that it is possible to estimate the log-policy gradient with only an logarithmic amount of samples (in the number of variational parameters). While this certainly holds for simulation, it is not clear, if such a technique can be applied on quantum hardware (e.g., some kind of sparse or perturbed gradients). The introduced initialization strategy gives some good experimental results, although some additional experiments and theoretical justifications would be desirable. The formulation of the empirical FIM drops the dependency on the prior state distribution, which potentially renders the considered spectrum less representative of the model than for a generic supervised learning problem. The claim of quantum advantage w.r.t. parameter complexity and absence of barren plateaus should be supported with experiments on larger-scale environments.

Algorithmic Characteristics - Sequeira et al. [SSB22]

| Environment       | Algorithm Type | Quantum Component | State Space | Action Space | Qubits | Parameterized Gates |
|-------------------|----------------|-------------------|-------------|--------------|--------|---------------------|
| CartPole (OpenAI Gym) | REINFORCE      | Policy            | continuous 4-dim | discrete     | 2      | 4 (encoding)        |
| Acrobot (OpenAI Gym)   | REINFORCE      | Policy            | continuous 6-dim | discrete     | 3      | 6 (encoding)        |
| QuantumControl (see [SSB22]) | REINFORCE  | Policy, Environment | quantum     | discrete     | N/A   | 0 (encoding)$^a$    |

$^a$ the RL state is a quantum state, i.e. no classical information has to be encoded;

Quantum reinforcement learning in continuous action space, Wu et al. (2020)

Summary. This paper by Wu et al. [Wu+20] extends the concept of VQC-based RL to continuous action spaces. The authors choose a quantum control environment, more concretely one that encodes an eigenvalue problem. This allows to interpret the action as a (parameterized) unitary. The experimental results suggest an exponential reduction in model complexity compared to classical approaches.
Eigenvalue Problem as RL Environment. The RL agent has to solve an eigenvalue problem, i.e., find the eigenvalue of a given Hamiltonian. This should be done in the following iterative procedure: Let \( H \) be the Hamiltonian of an \( n \)-qubit quantum system \( E \) and \( |s_0\rangle \) an initial state from \( E \). The system should be driven towards the eigenstate of \( H \), denoted as \( |u_0\rangle \). Also the corresponding eigenvalue \( \lambda_0 \) should be returned. Although not explicitly stated in the paper, we assume the agent should search for the eigenstate with the associated smallest eigenvalue, as this corresponds to the ground state.

The observation for this environment is the current quantum state \( |s_t\rangle \), which is provided to the agent via some quantum channel. The actions the agent can execute correspond to parameterized unitaries \( U(\theta_t) \), where \( \theta_t \) are classical parameters sampled from the VQC via measurements. Once instantiated, this unitary is applied to \( |s_t\rangle \) to evolve the state \( U(\theta_t) |s_t\rangle = |s_{t+1}\rangle \). The agent receives a classical reward, which describes the closeness of the current state to the searched eigenstate of the Hamiltonian.

The authors state, that their proposed technique has some parallels to Grover’s search. More concretely, the trained agent provides an alternative to the amplitude amplification procedure, which could alternatively be used to solve the task at hand.

Model Architecture and Underlying RL Algorithm. The overall approach can be considered hybrid, as the optimization of the VQC parameters is still conducted on classical hardware. A schematic description of the approach is given in Fig. 10. The agent observes a quantum state from the environment, which is used as the initial state \( |s_t\rangle \) of the VQC function approximator. Measurements on the prepared quantum state determine the parameters \( |\theta_t\rangle \). Those are then fed into the unitary operator \( U(\theta_t) \) and applied to the environment state. The new state \( |s_{t+1}\rangle \), combined with an ancilla reward qubit initialized to \( |0\rangle \), is then evolved using some user-defined reward unitary \( U_r \). Measurements are performed on this state to determine the reward produced by the executed action. This procedure repeats several timesteps, with the objective to approximate the eigenstate \( |u_0\rangle \).

Figure 10: Hybrid model for quantum environment proposed by and taken from Wu et al. [Wu+20] (including subcaptions); We note an ambiguity in notation, as the parameters \( \theta_{t,i} \) must not be confused with the parameters of the action unitary \( U(\theta) \). The first set are the ones updated by the RL algorithm, the other ones are extracted via measurements from the quantum state prepared by the VQC.

The VQC architecture does not incorporate a feature map, as the observation \( |s_t\rangle \) is used as the initial state \( |\Phi\rangle \). Each parameterized layer consists of 1-qubit rotations and a circular entanglement structure. For every element of the action parameters \( \theta_j \), there is an associated observable \( B_j \), which
is measured on the prepared quantum state. (The paper does not mention, how the action unitary $U(\theta)$ is explicitly constructed.) Following this step, a phase estimation circuit implements the reward unitary $U_r = U_{PE}$. This transforms the state to the basis of eigenstates, i.e., $U_{PE} |0 \rangle |s_{t+1} \rangle = \sum_{k=1}^{n} \alpha_{t+1,k} |\lambda_k \rangle |u_k \rangle$. With a measurement of the eigenvalue phase register, the desired eigenvalue $\lambda_0$ is observed with a probability of $p_{t+1} = |\alpha_{t+1,0}|^2 = |\langle s_{t+1} |u_0 \rangle|^2$. The reward can then be defined as e.g. $r_{t+1} = p_{t+1} - p_t$. Obviously, for $p_{t+1} \rightarrow 1$, the state $|s_{t+1} \rangle$ converges to $|u_0 \rangle$.

The underlying RL routine is an actor-critic method. Therefore, the paper combines a policy-VQC as actor and a Q-function-VQC as critic to a so-called quantum deep deterministic policy gradient (QDDPG) algorithm. The experience of the agent, i.e., tuples $(|s_t \rangle, \theta_t, r_t, |s_{t+1} \rangle)$, are stored in a replay buffer to prevent overfitting. Additionally, target networks are employed for both, the actor and the critic.

**Experimental Results and Model Complexity.** All experimental results in the paper are based on classical simulations. For training, the Hamiltonian $H = \frac{1}{4}(s_x \sigma_x + s_y \sigma_y + s_z \sigma_z + I)$ is instantiated with the coefficients $(s_x, s_y, s_z) = (0.13, 0.28, 0.95)$. Concrete details on the training procedure, e.g., the number of episodes, are not stated. The trained model is applied to 1000 random initial states. The overlap with the respective $|u_0 \rangle$ is approaching one, consequently the agent is able to get quite close to the desired eigenstate in all cases. The trained model shows good generalization capabilities, i.e., it can be applied to various initial states. This is in contrast to e.g. a variational quantum eigensolver (VQE), where the control pulse for one initial state is meaningless for other ones.

The overall gate complexity for one RL episode is stated as $O(m \cdot \text{polylog}(N))$. Here, $m$ is the number of shots for sampling expectation values and $N$ denotes the number of qubits. This statement assumes that $H$ can be efficiently simulated as otherwise the complexity of $U_{PE}$ would exceed $O(\text{polylog}(N))$. Additionally, all VQCs in the method are also assumed to have a gate complexity of at most $O(\text{polylog}(N))$. With this perquisites, the authors claims an exponential advantage in model complexity compared to classical approaches.

**Generalization to Discrete Action Spaces.** The paper also generalizes the presented concept to discrete action spaces, with the FrozenLake environment as an example. The observations are encoded as basis states into the VQC via computational encoding, similar to Chen et al. [Che+20]. The movements applied by the actions are formulated as unitaries acting on the VQC state. A slight generalization of Chen et al. [Che+20] is used for this, which allows to perform the transforms $|0 \rangle \rightarrow |1 \rangle$ and $|1 \rangle \rightarrow |0 \rangle$ in a parameterized manner. The reward unitary is formulated in a similar fashion. It is stated that experiments with this configuration were successful, but no concrete results are provided.

**Remarks.** There are some caveats and ambiguities we identified regarding the proposed approach. First, the algorithm requires knowledge of and ability to prepare the desired eigenstate $|u_0 \rangle$ for the training procedure. With this state already known, the whole procedure of reproducing it is a somewhat circular task. However, as the learned model seems to generalize to different input states, the technique offers clear advantage over approaches like quantum phase estimation. Second, the model requires repeated preparation of the environment state $|s_t \rangle$, as it is disturbed by measurements to extract the reward information. This should be doable, as one knows the state preparation routine $|s_t \rangle = U(\theta_{t-1}) \cdots U(\theta_0) |s_0 \rangle$. The influence of this additional overhead is unfortunately not considered in the complexity considerations discussed above. Third, the claim of exponential quantum advantage w.r.t. model complexity (i.e. $O(\text{polylog}(N))$ for all VQCs) should be supported by larger-scale experiments.
Algorithmic Characteristics - Wu et al. [Wu+20]

| Environment   | Algorithm Type | Quantum Component | State Space | Action Space | Qubits | Parameterized Gates |
|---------------|----------------|-------------------|-------------|--------------|--------|---------------------|
| Quantum       | Actor-Critic   | Q-function,       | quantum     | continuous^a | n      | 0 (encoding)^b       |
| Eigenvalues   | “QDDPG”        | Policy,           |             |              |        | n × d × 3 (weights)^c|
| (see [Wu+20])|                | Environment       |             |              |        |                     |
| FrozenLake    | Actor-Critic   | Q-function,       | discrete^d  | discrete^d   | n      | 0 (encoding)^b       |
| (OpenAI Gym)  | “QDDPG”        | Policy            | 16          | 4            |        | N/A (weights)        |

^a output is interpreted as parameters of a unitary, i.e. a quantum operation applied to the environment;
^b the RL state is a quantum state, i.e. no classical information has to be encoded;
^c variational gates: qubits × layers × per_qubit_per_layer; details are not specified;
^d state and action space are encoded into the quantum realm for a neat integration into the pipeline;

Introduction to Quantum Reinforcement Learning: Theory and PennyLane-based Implementation, Kwak et al. (2021)

Summary. The paper by Kwak et al. [Kwa+21] gives a short introduction to both RL and (variational) QC. This is followed up by a tutorial on how to implement a VQC-enhanced RL algorithm with PennyLane to solve the CartPole environment.

Hybrid RL Agent. The paper employs the typical hybrid structure, where the VQC is used as a function approximator. The optimization of the parameters and the interaction with the CartPole environment is executed on classical hardware. The underlying algorithm uses an actor-critic approach, where the actor is quantum and the critic is classical.

The architecture of the VQC is split up into three parts. A set of 1-qubit rotations is used to encode the state of the CartPole environment into the four-qubit system. This encoding layer is followed by 4 layers with learnable 1-qubit rotations and an unspecified entanglement structure. The result is extracted from the measurement of 2 qubits in the computational basis and the respective expectation values are interpreted as the action-value function.

Experimental Results. The agent is able to surpass random behavior, but still lacks behind other hybrid approaches proposed by e.g. [LS20; Jer+21a].

Remarks. To the best of our understanding, the implemented quantum actor-critic approach deviates in some details from previously considered approaches. Most importantly, a hybrid approach is used, where the actor is represented with a VQC and the critic employs a classical DNN.

Algorithmic Characteristics - Kwak et al. [Kwa+21]

| Environment   | Algorithm Type | Quantum Component | State Space | Action Space | Qubits | Parameterized Gates |
|---------------|----------------|-------------------|-------------|--------------|--------|---------------------|
| CartPole      | Actor-Critic   | Q-function         | 4-dim       | discrete     | 4      | 4 × 1 (encoding)    |
| (OpenAI Gym)  |                |                   |             |              |        | 4 × 4 × 3 (weights) |

^a encoding gates: qubits × per_qubit; variational gates: qubits × layers × per_qubit_per_layer;
^b only the actor employs a VQC, the critic uses a classical DNN;

Variational Quantum Soft Actor-Critic, Q. Lan (2021)

Summary. The paper by Q. Lan [Lan21] introduces a quantum version of a soft actor-critic (SAC) approach. The advantage of this algorithm, compared to previous suggestions, is the possibility to work with a continuous action space. The algorithm is tested on the Pendulum environment.
Soft Actor-Critic for Continuous Control. The term continuous control refers to a setup, in which the agent acts in a continuous action space. Most publications in the context of QRL deal with discrete action spaces, while a few others discuss continuous control for quantum environments [Wu+20; SSB22]. This work focuses on classical environments, which requires some kind of action decoding based on measurements of the quantum state. Instead of directly selecting the actions based on measurement results, the parameterized hybrid model learns the parameters of a distribution, from which the action is sampled. The VQC, and a downstream NN, are used to represent mean $\mu$ and variance $\sigma$ of a Gaussian distribution. This allows the agent to act in a continuous action space in a straightforward manner.

In contrast to the standard RL setup, SAC [Haa+18] not only aims to optimize the expected return, but also the policy entropy [Zie+08; Haa+17]. Therefore, the expected return is defined as $G_t = \sum_{i=t}^{\infty} \gamma^{i-t}(r(s_i, a_i) + \alpha H[\pi_\theta(s_i)])$, where $H[p] = -\int p(x) \log p(x) dx$ is the differential entropy for the probability density function $p(x)$. Among other advantages, this entropy normalization potentially enhances exploration by encouraging more stochastic policies [Haa+17].

VQC Architecture. The paper considers two different VQC architectures. The first one uses the typical three-part structure of rotational encoding, variational layers, and measurements. The second architecture is more complex, as it uses data re-uploading [Pér+20], and a more complex encoding structure [SJD22; Jer+21a]. It can be expected, that the second choice gives rise to more expressive models, which usually correlates with RL performance.

Experimental Results. The experimental section of the paper compares the performance of the two resulting quantum SAC approaches to a classical NN on the Pendulum environment. On the one hand, the quantum model with the simple VQC architecture is inferior to the other two approaches. On the other hand, the quantum model with data re-uploading performs similar to the classical model, and both are able to learn near-optimal behavior. The quantum model incorporates only 41 parameters, while the classical one uses 1250. This is interpreted as an quantum advantage w.r.t. parameter complexity.

Some additional architecture experiments are conducted, mainly focusing one the depth of the underlying VQCs. It is observed, that a certain number of variational layers is required to enable training. Overall, the performance is strongly correlated with the concrete architecture choice, which is in line with the results known from literature [Fra+22].

Remarks. To substantiate the claim of quantum advances w.r.t. parameter complexity, more experiments with increasing environment size should be performed. By using NNs in combination with VQCs, it is not completely clear, which part of the learning is actually conducted by the quantum part. The differing performance of the two architecture choices highlight the importance of designing a sophisticated data encoding scheme.

Algorithmic Characteristics - Q. Lan [Lan21]

| Environment       | Algorithm Type | Quantum Component | State Space | Action Space | Qubits | Parameterized Gates$^a$ |
|-------------------|----------------|-------------------|-------------|--------------|--------|------------------------|
| Pendulum (OpenAI Gym) | Quantum-SAC    | Q-function        | continuous 3-dim | continuous | 3      | 3 to 12 (encoding)     |
|                   |                |                   |              |              |        | 36 (weights)           |

$^a$ the hybrid model also incorporates additional classical parameters in an appended NN;

Quantum Multi-Agent Reinforcement Learning via Variational Quantum Circuit Design, Yun et al. (2022)

Summary. This paper by Yun et al. [Yun+22] introduces a quantum multi-agent reinforcement learning (QMARL) approach. It is applied to an environment inspired by wireless communica-
tion. The authors achieve results that are competitive with classical NNs with higher parameter complexity.

**QMARL Framework and VQC Architecture.** The approach is inspired by the classical method of centralized training with decentralized execution (CTDE). This approach deals with the problems introduced by a non-stationary reward structure, caused by the interaction of multiple agents [Low+17].

The actor-critic structure employs only a single critic (i.e. represented by a single VQC), which receives the rewards. A naive implementation would increase the qubit count with the number of agents. To resolve this problem, the state encoding routine is modified, such that only one qubit is required for each agent.

The general VQC architecture follows the typical three-part structure. The states are encoded using a feature map with 1-qubit rotations. The state space of the environment is four-dimensional. Consequently, four qubits are used to represent the actor associated to each of the four agents. For the critic, all rotations for the state of one agent are applied to a single qubit. This implies a qubit count equal to the number of agents (i.e. implemented for 4 qubits in the article). The following learnable layer(s) consist of 1-qubit rotations and some unspecified entanglement structure. The choice of the measured observables $M$ are not explicitly stated.

**Experimental Results and Discussion.** The QMARL algorithm is applied to a communication task referred to as **Single-Hop Offloading** environment. It simulates two clouds, between which packages have to be distributed along four edges. Each cloud and edge has a queue with a certain capacity. One agent is used to learn the actions of its associated edge. The objective is to minimize the overflow and underflow of queues.

The paper compares four different multi-agent reinforcement learning (MARL) and QMARL frameworks: (1) The described version, where actor and critic are represented with a VQC; (2) A modified pipeline, where the critic is represented with a classical NN; (3) A small-scale classical MARL approach; All three setups contain 50 trainable parameters each. (4) A large-scale classical MARL algorithm with over 40000 trainable parameters.

The results demonstrate, that the QMARL approach (1) is competitive with the large-scale MARL algorithm (4). In contrast, the hybrid QMARL method (2) and also the small-scale classical MARL seem to lack expressivity to solve this task. The authors conclude, that QMARL yields some quantum advantage, as the parameter complexity is drastically reduced.

**Remarks.** Potentially, compressing all observations of one agent into one qubit is not sufficient to represent the information in a lossless manner. Therefore, larger-scale experiments should be conducted to get more insights into the proposed quantum multi-agent architecture. The same holds for the reduced parameter complexity compared to classical models.

**Algorithmic Characteristics - Yun et al. [Yun+22]**

| Environment          | Algorithm Type   | Quantum Component | State Space | Action Space | Qubits | Parameterized Gates |
|----------------------|------------------|-------------------|-------------|--------------|--------|---------------------|
| Single-Hop Offloading | Multi-Agent      | Q-function Policy | continuous  | discrete     | 4      | 4 or 16 (encoding)$^a$ |
| (see [Yun+22])      | Actor-Critic     | Policy            | 4-dim       |              |        | N/A (weights)       |
| “QMARL”              |                   |                   |             |              |        |                     |

$^a$ the 4 quantum actors use 4 encoding parameters each; the quantum centralized critic contains 16;

**Quantum Multi-Agent Meta Reinforcement Learning, Yun et al. (2022)**

**Summary.** The second paper by Yun et al. [YPK22] extends their previous approach [Yun+22] with various new techniques for QMARL. It proposes to use meta-learning by pre-training only one
individual agent. This is followed by a fine-tuning the multi-agent scenario. Therefore, two different types of trainable parameters are used, i.e. trainable measurements are introduced to complement the typical variational parameters. The approach is also extended to continual learning, where meta-learning is performed on multiple environments at once.

VQC Architecture and meta-QMARL. The underlying RL algorithm employs an SAC approach with the VQC as function approximator for the action-value function. The quantum circuit uses the three-layer structure of 1-qubit rotation data encoding, variational layers with entanglement gates, and measurement. The paper applies QRL to multi-agent problems and extends the original proposal on quantum CTDE by Yun et al. [Yun+22]. An additional step is introduced for the training procedure, resulting in a meta-learning approach.

In order to realize these concepts, the authors define two different sets of parameters. First, there are the typical variational parameters $\phi$, usually parameterizing 1-qubit rotations. Second, it is also possible to parameterize and train the measurement observables. The paper proposes to use $M_{\theta_{12}^{(m)}} = R_z(\theta_1) \cdot R_y(\theta_2) \cdot Z \cdot R_y(\theta_2) \cdot R_z(\theta_1)$ as observable on the $m$-th qubit, i.e. two trainable parameters for each 1-qubit observable. Basically, this trainable observable introduces a change of basis, as final measurements are always performed in the computational basis. The instantiated observable can be visualized on the Bloch sphere as the angle w.r.t. which the measurement is performed.

Both parameter sets are trained in alternating steps, where the first one is referred to as meta quantum neural network (QNN) angle training, and focuses exclusively on the variational parameters $\phi$. This step trains only a single quantum agent, which interacts with several other agents in the multi-agent environment. Unfortunately, the authors do not state how this interaction is actually realized. We assume, that the quantum agent interacts with other classical agents in this initial training phase. During training, the pole parameters $\theta$ are not updated, but they can be varied with some randomly selected value to form a kind of angle-to-pole regularization. The second phase, the local QNN pole training, focuses on the parameterized observables. Those are fine-tuned individually for each copy of the meta-trained QNN, corresponding to the all-quantum agents interacting in the multi-agent environment. The authors propose, that by meta-training the network, it is more efficient to fine-tune the individual agents. This is justified with the lower parameter complexity, as the variational parameters remain constant in the second training phase. The loss function is the sum of all $Q$-learning losses of the individual agents.

Additionally, the paper introduces the concept of pole memory, which refers to storing the trained pole parameters for the individual agents. As these sets are much smaller than the set of variational parameters, it is more efficient to store the full configuration.

Experimental Results. The introduced training routing is executed on a two-step two-agent environment. It is observed, that the meta-training convergence is slower than direct training of a QMARL agent. However, once this training has converged, finetuning is much more efficient. Overall, the authors conclude, that the additional step of meta-training enhances convergence in a multi-agent environment.

Extension to Continual Learning. The above setting is also extended to continual learning, i.e. training in more than one environment (or typically the same environment with slightly altered dynamics).

The investigation focuses on the difference in performance with and without the use of pole memory. The results suggest that resetting the poles to the initial state (i.e. the parameter setting with which meta training was conducted) benefits convergence speed and stability in an environment with alternating dynamics. Meta training with a higher degree of angle-to-pole regularization seems to enhance the generalization performance of the meta-QNN.

Remarks. The paper does not state explicitly how exactly the initial meta training is conducted. Considering the results, the VQCs seem to have some capability w.r.t. transfer learning, as which
the meta-learning and continual training can be interpreted. The idea of employing trainable observables has also potential for other approaches, as it partially avoids the necessity to explicitly pre-select an action decoding scheme. Practically, these trainable observables are introduced by adding an additional layer to the VQC which learns a specific measurement. A significant difference to pre-existing procedures is that these parameters are not trained simultaneously with the typical variational parameters. It is not completely clear, whether this two-step training procedure is beneficial in a general setup.

Algorithmic Characteristics - Yun et al. [YPK22]

| Environment                | Algorithm Type        | Quantum Component | State Space | Action Space | Qubits | Parameterized Gates a |
|----------------------------|-----------------------|-------------------|-------------|--------------|--------|-----------------------|
| Single-Hop Offloading      | Meta-Multi-Agent SAC  | Q-function        | continuous  | discrete     | 4      | 4 (encoding)          |
|                            | “Meta-QMARL”          |                   | 4-dim       |              |        | N/A (weights)         |
| Two-Step Meta-Multi-Agent  | Meta-Multi-Agent SAC  | Q-function        | continuous  | discrete     | 2      | 2 (encoding)          |
| Game Offloading            | “Meta-QMARL”          |                   | 4-dim       |              |        | N/A (weights)         |

a the parameter counts are denoted for a single agent;

Q-Learning with Quantum Neural Networks, Hu and Hu (2019)

Summary. The paper by Hu and Hu [HH19a] uses a QNN to approximate the action-value function in Q-learning-based RL. The proposed algorithm is tested in the discrete state-action space FrozenLake environment (of size $2 \times 3$) from OpenAI Gym.

Big parts of the literature [Che+20; LS20; SJD22; Jer+21a] follow the gate-based QC paradigm (with simulators such as Qiskit, PennyLane or TensorflowQuantum). The authors of this paper use a photonic simulator (StrawberryField) to simulate a QNN in the continuous-variable (CV)-QC paradigm. The following photonic gates are used in sequence (compare e.g. Ref. [Kil+19]): (1) interferometer; (2) displacement; (3) rotation squeeze; (4) Kerr gates; Non-linearity is introduced into the model by the non-Gaussian Kerr gate. According to the paper, a measurement in the Fock basis (i.e. the photon number) represents the output of the QNN.

In the present study, the authors employ a single-layer CV-QNN, where the states of the RL environment are encoded as integers. Details on the architecture of the network can be found in the authors’ previous work [HH19d] (which we did not explicitly summarize, as it applies similar techniques to a more restricted set of ContextualBandits environments).

The authors claim that using a QNN to approximate the $Q$-value in a Q-learning-based RL algorithm is possible and stable. They also state that the quantum learner demonstrates advantage over a purely classical learner. However, these experimental results are not explicitly stated in the paper.

Remarks. We are not sure if the proposed QNN uses techniques that are beyond the scope of classical computing. Whether it is possible to realize a QNN that cannot be simulated efficiently on classical devices is an interesting topic for future work. We could not completely identify, which measurements are actually employed, or whether averaging is involved in producing the estimates for the action-value function.

Reinforcement Learning with Deep Quantum Neural Networks, Hu and Hu (2019)

Summary. This paper by Hu and Hu [HH19c] is an extension of the authors’ work [HH19d]. Similar to the previous work, a CV-QNN approximates the action-value function in Q-learning and actor-
critic based RL. The agent is trained on a $2 \times 3$ FrozenLake gridworld environment.

The proposed architecture is similar to the previous works [HH19d; HH19a], but employs a higher number of layers repeated in a sequential manner.

The authors of the paper claim that using a QNN to approximate the $Q$-value in both $Q$-learning and actor-critic based RL algorithms is possible and stable. They compare the performance of a three-layer network to the performance of a single-layer network. A larger number of layers (i.e. also parameters) in the function approximator improves the performance.

Remarks. The paper does not explicitly state some practical details, e.g. the encoding scheme and the post-processing of the measurement results. The possibility of quantum advantage over a classical counterpart is interesting future work.

Distributional Reinforcement Learning with Quantum Neural Networks, Hu and Hu (2019)

Summary. This article by Hu and Hu [HH19b] proposes distributional reinforcement learning (DistRL) [Dab+17] for a photonic quantum computer. It is an extension of previous work of the authors [HH19a; HH19c].

Distributional RL. Standard RL techniques, such as e.g. $Q$-learning, estimate the action-value function over the expectation value of the return distribution. In contrast, DistRL employs the full return distribution. The motivation is that a probability distribution generally contains more information than its expectation value. By moving from the expectation value to the entire distribution in the Bellman equation, one finds the so-called distributional Bellman equation. A parameterization of the distribution is chosen in terms of $N$ quantiles (that is the distribution is represented by an equally weighted sum over Dirac delta functions located at the position of the quantiles). The derivation of a DistRL algorithm is inspired by the construction of TD learning. Here, one first shows that the Bellman operator $T$ is a contraction with respect to the $L_2$ norm. Now TD learning iteratively minimizes the distance between $TQ$ and $Q$, which in turn guarantees convergence of the algorithm. The parameterization of the distributional Bellman operator in terms of quantiles as described above again is a contraction, but now with respect to the Wasserstein metric. Consequently, a learning algorithm adapts the reward distributions for each state-action pair (in terms of the quantile locations), such that the difference between $TQ$ and $Q$ is again minimized with respect to the Wasserstein metric. Unfortunately, it turns out that this minimization is biased. This problem is solved by using quantile regression, a method to determine the quantiles of a distribution by minimization of a loss function. Note that in this algorithm the next action is still selected greedily from the expected value of the $Q$-function. This algorithm is known as quantile regression (QR) $Q$-learning. Using function approximation, the input to the NN is the current state. The output is the location of the $N$ quantiles for each action. Consequently, the output space is of dimensionality $N \cdot |A|$, where $|A|$ is the number of possible actions.

Proposed Method. Hu and Hu [HH19b] implement the DistRL algorithm using a QNN for function approximation. The network architecture is adapted to a photonic device and run on a simulator. Unfortunately, it is not explicitly stated, how the measurement is decoded into actions and values for the quantiles. The overall architecture is similar to the one used in previous work [HH19a; HH19c].

Experimental Results. Experiments are conducted for a simple Markov-chain environment. It consists of a start state, from which the agent can reach a terminal state in two directions after two intermediate states. One final state produces a reward of $r = 1$, the other $r = 0.001$, such that there exists an optimal and a suboptimal policy. Compared to quantum $Q$-learning (one quantile) [HH19a; HH19c], the agent seems to explore better using the DistRL version (the former learns the suboptimal policy, the latter the optimal one). Apart from that, the average reward increases as the number of quantiles is increased.
Remarks. The paper compares the performance of a standard QRL approach [HH19a; HH19c] to the quantum DistRL approach with encouraging results. Over and beyond, also a comparison to a classical agent might be insightful.

Practical Meta-Reinforcement Learning of Evolutionary Strategy with Quantum Neural Networks for Stock Trading, Sorensen and Hu (2020)

Summary. The work by Sorensen and Hu [SH20] introduces a meta-RL system for stock trading on a photonic quantum computer. The term meta learning describes training a model on one task and finetuning it with a small number of samples on another one. The agent is trained on m portfolios containing n stocks. Subsequently, the agent learns to trade portfolios of different composition by meta-RL. The authors apply MAML [FAL17] and its extension CAVIA [Zin+19], which are few-shot meta-RL algorithms. Both classical versions use NNs for function approximation. The optimization method employed in the paper is based on evolutionary strategies.

For the quantum version of the agent, the classical NN is replaced by a parameterized circuit run on a photonic simulator. Since only 4 qumodes (the basic quantum-mechanical system for computation on photonic quantum computers) are used to limit the computation time, also the portfolio size is decreased to 2 stocks.

Two different architectures (with and without entanglement gates) are considered. The former yields better performance. The learning curves of the quantum agents show learning process for both architectures. As the quantum agent still is able to learn, the authors conjecture that its superiority might be observed as the number of qubits increases.

Remarks. The stated potential for quantum advantage should be supported with larger-scale experiments.

Variational Quantum Circuit-Based Reinforcement Learning for POMDP and Experimental Implementation, Kimura et al. (2021)

Summary. The paper by Kimura et al. [Kim+21] extends the concept of VQC-based RL to partially observable environments. The approach is inspired by classical model-free, complex-valued RL [HSS06]. Additionally, a novel VQC architecture (novel with regard to measurement procedure) is proposed. A detailed description of the gradient computation with backpropagation techniques is provided (it is not quite clear how this method generalizes to quantum hardware).

Partially Observable MDP. A partially observable Markov decision process (POMDP) is described as a tuple $\langle S, A, T, R, \Omega, O \rangle$ and is a generalization of a MDP. The variable $S$ denotes a discrete state space, $A$ is a discrete set of actions, $T(s'|s,a)$ describes the state transition probabilities and $R(s,a)$ is a reward function. Extending the fully-observable case, $\Omega$ is a discrete set of observations and $O(o|s,a)$ is an observation probability matrix with $o \in \Omega$.

One caveat of partially observable environments is the perceptual aliasing problem. This refers to the property, that the agent cannot distinguish two different states due to the limited observation ability. An example of such an environment is the partially observable maze used in Kimura et al. [Kim+21]. Similar to most gridworld environments, the task is to navigate from the start state to the goal state on the shortest path possible. However, the observations provided to the agent are ambiguous as several cells return the same state indicator.

Solving POMDPs with Complex Valued RL. One way to bypass this state ambiguity is to introduce a belief distribution over possible states. Unfortunately, this is computationally expensive. An alternative approach is complex-valued RL [HSS06; MNM17]. It incorporates time series information into the action-value function, which represented as complex numbers. More concretely, the complex $\hat{Q}$-function ($\hat{Q}$ denotes complex values) encodes the history of the agent, i.e. the previously visited states. The cumulative reward value is expressed by the absolute value of $\hat{Q}$-function, while the path length of the propagated reward is represented by the phase of the $\hat{Q}$-function on the complex...
plane. Therefore, \( \dot{Q} \)-function-Learning keeps continuity w.r.t. the described internal reference value. This helps distinguish states which are affected by the perceptual aliasing problem. Formally, this is achieved by updating the complex values in the opposite phase direction. The complex-valued \( \dot{Q} \)-function can be represented with tabular methods [HSS06], or with complex-valued NNs [MNM17].

The update mechanism represents a generalized \( Q \)-learning approach, i.e. the objective is to optimize the loss function \( L_\theta = \frac{1}{2} |\dot{Q}(\alpha_{t-k}, \alpha_{t-k}) - (r_{t+1} + \gamma \cdot \dot{Q}_{\text{max}}(t))\dot{u}_t(k)| \). Here, \( \dot{u}_t(k) = \beta^{k+1} \) is a complex hyperparameter and \( k \) is the trace length. The NN is replaced with a VQC as action-value function approximator in the following.

**VQC Architecture and Gradient Computation.** The paper deviates in several design choices from the standard method. Most importantly, the \( \dot{Q} \)-values for the different actions are not extracted from the same circuit (e.g. measurement on different qubits corresponding to different actions). Instead, the actions are encoded into the VQC with a feature map similar to the one used for state encoding. Consequently, different circuits have to be evaluated for each action. This encoding can happen either directly in the feature map, or alternatively into the decoding unitary. However, the three-part structure of the circuit is preserved.

The encoding unitary \( U_{\text{encoder}} \) consists of simple parameterized 1-qubit rotations, where three different concrete encodings are considered as shown in Fig. 11. The **Type 1** feature map encodes the observations directly with an \( \text{arcsin} \) function. **Type 2** uses a computational encoding for the observations, basically equivalent to the one proposed by Chen et al. [Che+20]. **Type 3** also uses an \( \text{arcsin} \) transform, but directly encodes the action information into the feature map.

The variational part repeats several layers of parameterized 1-qubit rotations, followed by a circular entanglement structure. In the experimental part, the authors consider different circuit depths.

The output of the circuit is evaluated with the Hadamard Test, which measures the prepared state against an output unitary \( U_{\text{out}} \). This introduces an overhead since a controlled version of the unitary \( U_{\text{out}} \) needs to be implemented (details in Fig. 11). Additionally, an ancilla qubit, and three 1-qubit gates are required. The output unitary itself consists also of learnable 1-qubit rotations. If encoding unitaries of **Type 1** or **Type 2** are used, it additionally encodes the action information. The real and imaginary output of the Hadamard test are used to construct the complex-valued \( \dot{Q} \)-function.

The evaluation of the circuits is straightforward on quantum hardware. However, this does not apply to evaluating gradients w.r.t. the parameters, which is necessary for training. The paper gives a detailed derivation on how to compute the gradients via simulation on classical hardware. The
idea is inspired by classical backpropagation and somewhat looks like the adjoint method [Luo+20]. This makes it infeasible, at least in the given form, for actual quantum hardware.

**Experimental Results and Discussion.** The paper compares the training results (on the described maze environment) for the three types of quantum agents to different classical agents. The classical tabular approach outperforms all other methods, as the underlying algorithm guarantees an optimal solution. The authors argue, that there seems to be some intrinsic advantage of the Type 2 quantum circuits, as these perform better then the other approximate algorithms.

**Remarks.** We think there needs to be some further investigation regarding the applicability of the algorithm to actual quantum hardware. Currently, we propose to consider the approach as QiRL. We agree, that QC offers great potential for complex-valued RL, as QC itself deals with complex numbers. However, there are still open questions regarding the most promising way to exploit this connection.

### Algorithmic Characteristics - Kimura et al. [Kim+21]

| Environment   | Algorithm Type | Quantum Component | State Space (Qubits) | Action Space (Qubits) | Parameterized Gates |
|---------------|----------------|-------------------|----------------------|-----------------------|--------------------|
| PO Maze       | complex-valued DQL | Q-function discrete | 18                   | 4                     | 6 × 2 (encoding) |
|               |                 |                   |                      |                       | 6 × 4 × 2 (weights) |
| (see [Kim+21])|                 |                   |                      |                       | 6 × 3 (decoding)  |

*a encoding and decoding gates: qubits × per qubit; variational gates: qubits × layers × per qubit_per_layer;*

**Variational quantum reinforcement learning via evolutionary optimization, Chen et al. (2022)**

**Summary.** The main focus of the paper by Chen et al. [Che+22] is the investigation of gradient-free evolutionary optimization for Q-learning with VQCs. This routine is tested in two different scenarios, for each of which also a state encoding scheme is proposed. More concretely, amplitude encoding is applied to the CartPole environment. For the gridworld environment MiniGrid with larger state space (147 dimensional), the paper proposes a hybrid model with an encoding mechanism based on tensor network (TN) techniques.

**Amplitude Encoding.** The observation space of the CartPole environment is 4-dimensional. The state values are continuous. This allows the use of amplitude encoding, i.e. two qubits can be used to encode the (re-scaled) values into the four amplitudes of the system. The authors follow the method described in Schuld and Petruccione [SP18]. This works fine for small systems, but requires not NISQ-compatible operators for bigger instances.

**TN-based Encoding.** The MiniGrid environment is similar to FrozenLake, as the goal in both environments is to navigate from a start to a goal state on the shortest way possible. The paper uses simple environment configurations, with state spaces of size 5×5, 6×6, and 8×8. The observation space is of dimensionality 7×7×3. The agent has to decide between 6 actions, of which only 4 are relevant in the simplified scenario. The reward is defined as $1 - 0.9 \cdot \text{number\_steps} / \text{max\_number\_steps}$. Apart from the larger observation space, we assume this environment to be about the same complexity as FrozenLake.

The paper addresses the problem of encoding the 147-dimensional state into a quantum feature map with just 8 variational parameters. Other work uses e.g. CNNs to reduce the dimensionality of the feature space [LS21]. As the encoding networks have to be pre-trained, it is not quite clear, what part of the work is really done by the VQC. The authors suggest to use a hybrid encoding scheme based on TNs, similar to Chen et al. [Che+21]. The proposed TN technique encodes the
observation \([v_1, \ldots, v_{147}]^t\) into the product state \([1 - v_1, v_1]^t \otimes [1 - v_2, v_2]^t \otimes \cdots \otimes [1 - v_N, v_N]^t\), where the individual elements are normalized. Those encoded states represented by the red nodes in Fig. 12a. The trainable part of the matrix product state (MPS) outputs an 8-dimensional compressed feature vector. This is represented by the 147 + 1 blue nodes and the open leg (i.e. outgoing edge) in Fig. 12a. The bond dimension is a hyperparameter of the MPS, which correlates with the number of trainable parameters [Per+06].

VQC Architecture. The model follows the typical three-part architecture, i.e. first the feature map, then the variational part, and finally some measurements. For the CartPole environment, a simple 2-qubit circuit with amplitude encoding and 4 variational layers is used. Both qubits are measured in the Pauli-Z basis and the action corresponding to the higher expectation value is selected. For the MiniGrid environment, the 8-qubit circuit with just one repetition of the variational layer is used. The encoding is done with the TN-compressed state, i.e. the output from the TN is encoded into the circuit as shown in Fig. 12b. As the environment has 6 actions, the top 6 qubits are measured, and the action corresponding to the highest expectation value is executed.

RL with Evolutionary Optimization. The underlying algorithm is a Q-learning RL approach. The updates of the QNN representing the action-value function are conducted via evolutionary optimization. This implies, that no gradients have to be computed. Usually, this is one major bottleneck of VQC-based RL, which might be circumvented by this approach.

The paper uses a simplistic instance of an evolutionary algorithm, where mutation, but no recombination operations are employed. An initial population of \(M\) individuals is generated, which are used to simulate some episodes on the environment. The best \(T\) agents (the ones producing the highest reward averaged over several runs) are selected as parents for the next generation. Random Gaussian noise is applied to this parents (mutation), until \(M - 1\) children are generated. Additionally, the best individual from the previous generation is kept, i.e. again \(M\) individuals. This procedure is repeated until a certain convergence criteria is met, e.g. a high enough reward.

Experimental Findings and Discussion. The paper applies the two different encoding methods, combined with the evolutionary optimization idea, to the respective environments. All experiments are conducted as noiseless simulations. On the CartPole environment, the 2-qubit architecture achieves an near-optimal performance with only 26 parameters, which is significantly less than in most state-of-the-art NNs. The authors claim, that with their method the number of parameters can be reduced to \(O(\text{polylog}(n))\). In contrast, classical ML requires \(O(\text{poly}(n))\) parameters.

The experiments on the MiniGrid environment employ the described hybrid TN-based architecture. Results are compared to an encoding based on a simple NN, presumably similar to Lockwood and Si [LS21]. All approaches achieve a near-optimal performance. Overall, the TN-model (with large enough bond dimension) slightly outperforms the classical approach. The authors consider
this as a proof-of-principle for effectiveness of the MPS encoding for RL learning.

Remarks. The amplitude encoding is currently not feasible for more complex problems, due to the lack of an NISQ-compatible state-preparation routine. The evolutionary optimization approach could circumvent some of the problems typically associated with gradient based techniques. Experiments on larger-scale environments might be an interesting direction for future work, to investigate how the evolutionary algorithm deals with more complex optimization landscapes. We suggest to incorporate some recombination procedures into the evolutionary algorithm, to enhance its performance.

Algorithmic Characteristics - Chen et al. [Che+22]

| Environment       | Algorithm Type | Quantum Component | State Space | Action Space | Qubits | Parameterized Gates |
|-------------------|----------------|-------------------|-------------|--------------|--------|---------------------|
| CartPole (OpenAI Gym) | DQL\(^a\) | Q-function | continuous 4-dim | discrete | 2 | 3 (encoding) 26 (weights) |
| MiniGrid (OpenAI Gym) | DQL\(^a\) | Q-function | discrete 147 | discrete | 8 | 8 (encoding)\(^b\) 24 (weights) |

\(^a\) the standard QRL pipeline is reformulated to allow training via evolutionary optimization;
\(^b\) this does not include parameters of the TN for dimensionality reduction; details are not stated;

Unentangled quantum reinforcement learning agents in the OpenAI Gym, Hsiao et al. (2022)

Summary. The paper by Hsiao et al. [Hsi+22] uses an hybrid proximal policy optimization (PPO) algorithm, with a combination of VQC and NN as policy function approximator. The quantum circuit architecture is untypical, as it only uses 1-qubit rotations. Consequently, no entanglement is created, and all qubits can be considered as independent systems. Still, the resulting RL agent is able to learn good policies on some standard environments (CartPole, Acrobot, and LunarLander). The learned parameters are ported to quantum hardware and tested with sophisticated results.

Underlying RL Algorithm and Model Architecture. The classical RL algorithm is PPO, i.e. a policy-based approach. It follows the typical hybrid setup, as the VQC is used as function approximator, and parameter updates are computed on classical hardware. To enhance the expressivity of the model, a classical NN is appended. It uses the measured expectation values as inputs. The outputs of the network are post-processed using a softmax function.

The structure of the hybrid model is displayed in Fig. 13. The feature map consists of 1-qubit rotations, which is a common choice in the literature. The variational (‘parameter’ in Fig. 13) layer incorporates 1-qubit parameterized rotations. It is important to highlight that the circuit does not contain any multi-qubit gates. Consequently, no entanglement between the qubits is created. As efficient classical simulation of the circuit is possible, the approach should be counted towards QiRL. Despite this, the authors demonstrate, that a good RL training performance can be achieved with this model.

Experimental Results. The described hybrid agent is trained on three tasks from the OpenAI Gym, i.e. CartPole, Acrobot, and LunarLander. The quantum agents outperform several classical architectures. As this is achieved with much fewer parameters, the authors claim that the approach points towards potential advantage.

The results on LunarLander are remarkable in that regard, that it might be the most complex environment solved with VQC-based RL thus far. While the classical simulability prohibits any intrinsic quantum advantage, the models still are able to achieve a good performance. This gives
rise to the questions, whether one can draw inspiration from quantum mechanics for purely classical approaches.

Testing on Quantum Hardware. Once the models are trained, they are tested with the learned parameters on IBMQ hardware (with up to 8 qubits, depending on the environment). The models are able to replicate the learned near-optimal behavior.

Remarks. As without entanglement the VQCs can be simulated classically, we agree with the authors that the proposed algorithm should be considered as a QiRL approach. As the proposed model incorporates also a classical network, it is not clear, what part of the learning is conducted with the VQC. The simple circuit structure might also explain, that the results for testing on hardware are stable. Usually, a big portion of the noise is caused by two-qubit gates, which are not present in the used VQC.

Algorithmic Characteristics - Hsiao et al. [Hsi+22]

| Environment          | Algorithm Type | Quantum Component | State Space     | Action Space | Qubits | Parameterized Gates$^a$ |
|----------------------|----------------|-------------------|-----------------|--------------|--------|------------------------|
| CartPole (OpenAI Gym)| PPO            | Policy            | continuous 4-dim| discrete 2   | 4      | 12                     |
| Acrobot (OpenAI Gym) | PPO            | Policy            | continuous 6-dim| discrete 3   | 6      | 18                     |
| LunarLander (OpenAI Gym) | PPO        | Policy            | continuous 8-dim| discrete 4   | 8      | 72                     |

$^a$ this entails encoding and variational parameters; an appended NN adds classical parameters;

Short quantum circuits in reinforcement learning policies for the vehicle routing problem, Sanches et al. (2022)

Summary. This work by Sanches et al. [San+22a] uses small-scale VQCs to replace shallow classical NNs within a model with attention mechanism. The algorithm is applied to find solutions to instances of the vehicle routing problem (VRP), which is an NP-hard combinatorial optimization task. The authors explicitly states that they do not look for any kind of quantum advantage. Instead, they want to demonstrate, that VQCs are competitive with NNs in a certain scenario. Additionally, some technical tricks for running experiments on quantum hardware are presented.
The Vehicle Routing Problem. The VRP is a generalization of the more well-known traveling salesman problem (TSP), which is also an NP-hard combinatorial optimization task. An instance of the VRP can be represented as a graph with $n$ nodes, where one node is the depot, and the other $n-1$ ones are places we have to make deliveries to. For each of these nodes there is a positive real number which states the demand of the respective destination. The edges of the graph are labeled with positive real values, which denote the cost of driving from one node to the adjacent one.

RL with Attention Mechanism. As the VRP is NP-hard, most (RL) approaches focus on learning heuristics, that provide a good approximation to the optimal solution. The paper makes use of an encoder-decoder model with an attention mechanism [KWW18]. The concrete structure and functionality of the model is of not too much importance for this summary. In short words, the attention mechanism allows to put more emphasis on the important parts of the data. The authors focus on modifications to the encoder part, more concretely the attention heads. These are usually implemented with shallow NNs, which are replaced with 4-qubit VQCs. In the RL setup, the authors make the choice to interpret the solution as only one action. This means, that the action $a_0$ consists of the complete route taken by the truck, and the reward $r_1$ represents the negation of the traveled distance. It turns out, that this formulation allows the agent to train successfully.

Quantum Architecture of The Attention Heads. As explained above, the VQCs are used to replace the attention heads. These are given a pair of nodes and output values which are trained to describe some compatibility between them. With this concept in mind, the authors define a 4-qubit quantum circuit, where one node (the key) is encoded into the upper two qubits, and the other one (the query) into the bottom two. For practical reasons, this is extended with two ancilla qubits. It is not explicitly described, how this encoding is performed. The parameters are trained with an instance of REINFORCE. The observables $\frac{1}{4}(X_1X_3 + Y_1Y_3 + Z_1Z_3)$ and $\frac{1}{4}(X_2X_4 + Y_2Y_4 + Z_2Z_4)$ are measured on the quantum state and the results are post-processed.

Experiments and Discussion. The described models are simulated on a VRP instance with 14 nodes randomly located on the unit circle (depot in the middle, random demands). Both, the classical and the hybrid model, are able to learn near-optimal behavior. It seems, that the quantum model performs slightly better, but this is considered to be statistically insignificant. The authors also execute a reduced test with 5 nodes on the hardware (with parameters trained via simulation). In this experiment, the agent is able to find the optimal route. As this is only a single instance of a quite simple problem, the implications of this success are weakened a bit by the authors. It is concluded, that components of a classical ML model can be replaced with a VQCs without losing performance.

Remarks. The attention heads are usually quite small networks in the classical case. Therefore, it is not clear, how the proposed idea could be extended to scenarios where quantum advantage is possible. However, the integration into a complex classical pipeline demonstrates, that considering hybrid models is an interesting research direction. This would allow to combine the benefits of classical hardware (e.g. error-free, scale-able) with quantum hardware (e.g. potentially more expressive model).

Algorithmic Characteristics - Sanches et al. [San+22a]

| Environment | Algorithm Type | Quantum Component | State Space | Action Space | Qubits | Parameterized Gates |
|-------------|----------------|-------------------|-------------|--------------|--------|---------------------|
| VRP$^a$     | REINFORCE      | Attention-Head    | continuous  |              | 4      | 4 (encoding)        |
|             |                |                   | 2-dim       |              |        | 2 (weights)         |

$^a$ several instances of this component are used as subroutine of a complex classical model;
4.3 Projective Simulation for Quantum Reinforcement Learning

| Citation | First Author | Title |
|----------|--------------|-------|
| [BD12]   | H. J. Briegel | Projective simulation for artificial intelligence |
| [Mel+17] | A. A. Melnikov | Projective simulation with generalization |
| [Boy+20] | W. L. Boyajian | On the convergence of projective-simulation–based reinforcement learning in Markov decision processes |
| [Pap+14] | G. D. Paparo | Quantum Speedup for Active Learning Agents |
| [DFB15]  | V. Dunjko | Quantum-enhanced deliberation of learning agents using trapped ions |
| [Sri+18] | T. Sriarunothai | Speeding-up the decision making of a learning agent using an ion trap quantum processor |

Table 3: Work considered for “Projective Simulation for QRL” (Subsec. 4.3)

Projective simulation for artificial intelligence, Briegel et al. (2012) and related work

Summary. Projective simulation for artificial intelligence by Briegel et al. [BD12] is the first in a series of articles, which propose a learning scheme for creative behavior. This is understood in the sense that the agent can deal with unseen experiences by relating to other conceivable situations. The method is developed for classical agents. There is only a brief final paragraph, outlining a quantum-mechanical implementation. Since subsequent papers ‘quantize’ the original idea heavily, a brief summary is in order: The approach is based on a random walk on a previous-experience network (memory), simulating an agent pondering its next action. More specifically, previous experiences compose a network of clips, which is dynamically modified by new experiences. It is important to note that clips, in contrast to actual experiences, are e.g. remembered observations, states or actions. To select the next action, an observation of the agent activates a clip, followed by a random walk through the network (projective simulation). This is repeated until an action is ‘excited’ and coupled out from the network and the action is selected. It is worthwhile noting that the term projective as used here is not related to its use in quantum physics, such as in projective measurement.

Action Selection. The process of action selection is slightly more sophisticated than described above. If a percept $s$ is observed, a random walk through the network starts from the corresponding percept clip. After some deliberation time the random walk reaches an action clip, which is only out-coupled and taken in reality if the percept-action pair $(s,a)$ was rewarded in the past (i.e. tagged positively). If not, a new simulation is started. This process repeats until an action clip with positive tag, or a predefined reflection time is reached; in the latter case the action is out-coupled irrespective of the tag.

Learning Procedure. The actual learning process can be summarized as follows:

1. If a transition $(s,a)$ is rewarded, increase the network weight of the direct transition $s \rightarrow a$. (Note that the agent might have chosen $s \rightarrow a$ after many steps of PS; by reinforcing the direct transition, it might be exploited directly next time);

2. Increase the weights of the indirect transition (all weights of the network that led to the transition $s \rightarrow a$ in the random walk through the network). Thus, the agent discovers useful actions after deliberation of fictitious clips;

3. Introduce damping of all weights to let the agent forget, in order to be able to adapt to new situations (as appearing for example in a time-dependent environment);
4. If a new situation is discovered, a corresponding clip is added to the network and directed edges from all the other clips to the new one are added;

5. Additional extensions can be implemented, such as modifications of clips and creation of completely fictitious compositions of episodes;

This line of research has been continued in Ref. [Mel+17] (generalization) and [Boy+20] (convergence). In the last paragraph of Ref. [BD12] a quantum version of the algorithm is briefly discussed. The idea is to replace the random walk on the network by a quantum walk. A number of subsequent papers investigate the quantum approach more rigorously:

In order to define a quantum walk algorithm as done in Ref. [Pap+14], the PS approach is viewed slightly different. The given clip network with the percept set $S$ is separated in $|S|$ disjoint networks. Thus one obtains a directed weighted graph (a Markov chain) for each percept with action clips as absorber states. Each of the actions is initially flagged (corresponding to the emotion tags of the initial projected simulation proposal). If an actual out-coupled action did not lead to a reward, this particular flag is removed. Now the action selection proceeds in the following way: If the agent observes a percept $s$, a random walk starts through the graph (deliberation) until an action is reached, which is out-coupled only if the action is flagged (reflection). Thus, action selection corresponds to sampling from the conditional probability distribution over the flagged action space. Given the transition matrix $P$ of the Markov chain, subsequent applications of $P$ to the initial state (probability one for the percept clip) realizes the approximate stationary distribution (subsequently referred to as diffusion). Sampling from this distribution and disregarding un-flagged actions produces the correct samples. As $p_s$ is the probability to sample a flagged action from the equilibrium distribution obtained by diffusion, one needs to repeat the sampling process $O(1/p_s)$ times until a flagged action is sampled. The quantum random walk search algorithm is closely related to Grover’s algorithm. By elevating the transition matrix to a diffusion operator and introducing an oracle that marks flagged actions, the quantum algorithm only needs $O(1/\sqrt{p_s})$ oracle calls. Consequently, a quadratic speed-up for the deliberation process can be achieved. Therefore, this quantum algorithm speeds up the agent’s internal computation time for action selection.

Experimental Implementation. In Ref. [DFB15] the authors investigate the implementation of the algorithm proposed by Ref. [Pap+14] on an ion trap quantum computer. Results are also backed up by numerical simulations. The actual proof-of-principle experiment with two qubits is discussed in Ref. [Sri+18], where signatures of the quadratic speed up are observed.

4.4 Boltzmann Machines for Quantum Reinforcement Learning

| Citation | First Author | Title |
|----------|--------------|-------|
| [Jer+21b] | S. Jerbi     | Quantum Enhancements for Deep Reinforcement Learning in Large Spaces |
| [Cra+18]  | D. Crawford   | Reinforcement Learning Using Quantum Boltzmann Machines |

Table 4: Work considered for “Boltzmann Machines for QRL” (Subsec. 4.4)

Quantum Enhancements for Deep Reinforcement Learning in Large Spaces, Jerbi et al. (2021) and related work

Summary. The work presented in Ref. [Jer+21b] investigates an alternative NN architecture to those often used for learning the $Q$-function (or more generally the merit function) in RL tasks. The authors argue that these alternative models perform advantageously in large action spaces. This is
due to their capability to represent multimodal functions better than standard network architectures, while using a similar number of parameters. It is further found that these alternative architectures are closely related to energy-based models, some of which admit quantum representations. In turn, this allows quantum evaluations, enabling a provable quantum speed-up for fault-tolerant quantum computing.

**Motivation.** The standard architecture for $Q$-learning with NNs is depicted in Fig. 14 (upper part). The representation of a state is fed into a NN, which outputs the values of the so-called merit function (the $Q$-value in case of $Q$-learning) for each possible action (given the state). The policy can be derived from this function by with softmax post-processing. The effective-temperature parameter is decreased over time to reduce exploration and enhance exploitation.

The authors argue that this NN architecture is not suited for large action spaces. It has to output a high dimensional function, i.e. the merit functions for all actions simultaneously for a given state. The authors argue that this network is unable to approximate a multimodal merit function in case of complex state-action correlations. Instead, the authors discuss the NN structure shown in the lower part of Fig. 14. Here, the state and action is fed to the NN, which outputs the corresponding merit function. Action selection is done by sampling from the probability distribution, given by a softmax function on the values of the merit function. Therefore, sampling requires $|A|$ forward passes, where $A$ is the action set, making action selection a computationally expensive task for large action spaces.

Experiments are conducted on a generalized GridWorld environment with a large set of actions. The associated complex transition function gives rise to one optimal and many sub-optimal policies. The authors find that the NN architecture shown in the lower part of Fig. 14 indeed performs better, but at the cost of the expensive sampling described before.

**Energy-based Models.** The potential for quantum speed up comes from the observation that the second architecture in Fig. 14 is equivalent to a certain kind of energy-based model. Energy-based function approximators are used for generative modeling of probability distributions based on the Boltzmann-Gibbs distribution with respect to an energy functional. Boltzmann machines are one instance of such energy-based models where the energy functional is given by a spin-spin interaction model. However, Boltzmann machines are hard to train which led to the development of restricted Boltzmann machines where a special interaction structure with a hidden layer enables more efficient training. In Ref. [Jer+21b] the authors observe that the lower architecture in Fig. 14 is equivalent to a generalized form of restricted Boltzmann machines.

**Quantum Speed-Up.** Inspired by this insight, the authors next investigate quantum energy-based models. Here, the classical spin-spin interaction energy is promoted to a spin Hamiltonian, known as quantum Boltzmann machines and restricted quantum Boltzmann machines. Some of these models allow efficient training, while the hardness of sampling remains. To speed up sampling in the classical and quantum setting, the following quantum subroutines for a RL algorithm are discussed:

1. Quantum Gibbs sampling: The Gibbs-Boltzmann distribution is prepared as a qsample, from which expectations values can be sampled with quadratic speed-up, compared to classical Monte-Carlo sampling methods. (2) Gibbs-state preparation by Hamilton simulation: Using Hamilton-simulation techniques, an approximation to the Gibbs qsample can be prepared, leading to quadratic speed-up compared to exact sampling (calculating all energies and explicitly normalizing the probability distribution). (3) Quantum simulated annealing: This method uses a quantum method for the approximate Monte-Carlo sampling of the Gibbs state itself by leveraging quantum random walks on graphs.

All methods discussed so far need oracularized access to the Hamiltonian and it is unlikely that they could be realized on current hardware. A realization on near-term hardware might be achieved by (4) Variational Gibbs-state preparation: Here, a variational circuit can be employed to approximate a Gibbs qsample, using the free energy as an objective. Any quantum speed up,
however, for this method is heuristic and has not been made rigorous so far.

Remarks. Crawford et al. [Cra+18] propose models based on quantum Boltzmann machines for quantum annealing hardware. Since this literature survey focuses on algorithms proposed for gate-based QC, we do not include a detailed summary here.

4.5 Quantum Policy and Value Iteration

So far, we have considered QRL algorithms that employ QC for function approximation or propose quantum approaches to alternative learning frameworks such as PS. We now turn to proposals that replace subroutines of existing RL frameworks by quantum algorithms such as amplitude estimation, quantum maximum finding and, respectively, quantum matrix inversion. As a result, the proposed QRL algorithms guarantee improved sample or computational complexity. As these methods need oracular access to the environment, they should be categorized as post-NISQ algorithms.

| Citation | First Author | Title |
|----------|--------------|-------|
| [Wan+21a] | D. Wang | Quantum algorithms for reinforcement learning with a generative model |
| [CKP22] | E. A. Cherrat | Quantum Reinforcement Learning via Policy Iteration |
| [Wie+22a] | S. Wiedemann | Quantum Policy Iteration via Amplitude Estimation and Grover Search - Towards Quantum Advantage for Reinforcement Learning |

Table 5: Work considered for “Quantum Policy and Value Iteration” (Subsec. 4.5)

Quantum algorithms for reinforcement learning with a generative model, Wang et al. (2021)

Summary. The work in Ref. [Wan+21a] proposes two algorithms for RL with a generative model and rigorously derives bounds for their sample complexity.
Classical Generative Models. Classically, the term generative model describes a simulator, which queried with a state-action pair \((s, a)\), produces a sample \(s' \sim P(\cdot|s, a)\). Thus, by repeated sampling for each state-action pair, one can estimate the transition matrix of the underlying MDP. This allows to subsequently obtain an approximation of the optimal policy by means of value iteration. Over the years there has been tremendous effort devoted to improving sample efficiency (defined as the number of times the simulator has to be queried). This performance metric is meaningful if one assumes that every query of the simulator is costly. The best classical algorithm \([Li+20b]\) requires a total number of \(O(|S||A|\Gamma^3/\epsilon^2)\) samples, where \(|S|\) and \(|A|\) are the number of states and actions, \(\Gamma = 1/(1 - \gamma)\) is the effective horizon of the MDP, and \(\epsilon\) is the deviation of the optimal value function from the approximation. The sample complexity is linear in the product \(|S||A|\), since the transition matrix has to be estimated for each \((s, a)\). The factor \(1/\epsilon^2\) originates from Hoeffding’s inequality (indeed bounding the deviation of a sample average from its real value by \(\epsilon\), requires \(O(1/\epsilon^2)\) samples). The origin of the third power of \(\Gamma\), in contrast, is less intuitive. Note that the sample complexity of the classical algorithm is also a lower bound (in the classical case) and therefore optimal.

Incorporating Quantum Subroutines. As shown in Ref. \([Wan+21a]\), the classical sample complexity can be reduced by replacing the classical mean-estimation subroutine in Ref. \([Li+20b]\) by a quantum routine based on the quantum mean-estimation algorithm \([Bra+02]\). Even though the optimal classical algorithm is more sophisticated as outlined above and so is its quantization, the following discussion captures the essential features. The quantum subroutine requires the generative model in oracle form and can then be used to estimate the expectation value \(E(V) = \sum_{s', s} P(s'|s, a) V(s')\) (which appears in the Bellman equation) individually for every pair \((s, a)\) in time \(O(1/\epsilon)\). This quadratic speed-up originates from Grover’s algorithm, on which the quantum-mean estimation algorithm is based upon. As a consequence, the quantum-policy iteration algorithm achieves the sample complexity \(O(|S||A|^{1.5}/\epsilon)\) with a quadratic improvement in \(\Gamma\) and \(\epsilon\).

The dependence on the size of the action space can be further reduced by using quantum maximum finding \([Mon15]\) to calculate the maximum over actions in the Bellman optimality equation. However, using this quantum routine, one cannot fully exploit the power of the classical optimal algorithm. Hence, while the dependence on \(|A|\) is reduced quadratically and the improvement in \(\epsilon\) is kept, the improvement in \(\Gamma\) is lost. As a result, the algorithm based on both quantum-mean estimation and quantum maximum finding achieves a sample complexity \(O(|S|\sqrt{|A|}\Gamma^3/\epsilon)\).

Finally, the lower bound \(O(|S||A|^{1.5}/\epsilon)\) is derived and possible improvements of the algorithm to reach this limit are discussed.

Quantum Reinforcement Learning via Policy Iteration, Cherrat et al. (2022)

Summary. Ref. \([CKP22]\) proposes a quantum algorithm for an iterative scheme of \(Q\)-value evaluation and policy improvement. The algorithm evaluates the \(Q\)-value on a quantum computer, with the state vector representing the \(Q\)-values, being extracted by measurements. The policy afterwards is improved on a classical device. The algorithm can achieve quantum advantage in certain situations.

To set up the general framework, the authors first formulate the Bellman equation for \(Q\)-value evaluation as a matrix equation \([LP03]\)

\[
Q = R + \gamma P \Pi Q.
\]

Denoting the size of the action and state space as \(|A|\) and \(|S|\), the \(|A||S|\) dimensional vectors \(Q\) and \(R\) represent the \(Q\)-values and the reward vector, respectively; the environment transition function is the \(|A||S| \times |S|\) dimensional matrix \(P\); the policy is represented by an \(|S| \times |A||S|\)-dimensional matrix \(\Pi\); \(\gamma\) denotes the usual discount factor; The authors propose to compute \((\mathbb{I} - \gamma P \Pi)^{-1} R\) on a quantum device.

Quantum Subroutine: Block Encodings and Linear Algebra. To perform this task, Ref. \([CKP22]\) relies on so-called block encodings of matrices \([Gil+19]\). This powerful framework gives rise to
various quantum algorithms for encoding general complex (not necessarily rectangular) matrices in the leading principal block of a larger unitary matrix. Once the data has been loaded, the framework further provides linear-algebra routines such as matrix multiplication, addition [Gil+19] and inversion [CKS17]. The encoding algorithms need quantum access to the data, i.e. via oracles. Therefore, the methods can be attributed to the post-NISQ algorithms category. A well-known data-loading scheme is the sparse-input model, viable for sparse matrices. The authors of Ref. [CKP22] apply a more general scheme, the so-called \( \mu_p(A) \) [CGJ19] block encoding of a matrix \( A \). Here, the quality (i.e. the probability to obtain the correct output of the algorithm, e.g. after matrix-vector multiplication and a subsequent measurement) of the encoding depends on the maximum of the column and row norms of the matrix. The aforementioned norm is a function of \( p \) and can be chosen freely to optimize the encoding quality. Based on this formalism, the authors show that policy evaluation requires time
\[
\mathcal{O}(\mu_p \Gamma \text{polylog}(|S||A|/\epsilon)) .
\] (26)

In Eq. (26), the parameter \( \Gamma = (1 - \gamma)^{-1} \), \( \epsilon \) denotes the accuracy of the matrix inversion subroutine. The term \( \mu_p \) describes the quality of the encoding of the environment-transition matrix, which depends on the structure of the environment. In the worst case it scales as \( \sqrt{|S||A|} \). Due to the sparsity of the transition function of many environments, a better scaling is often expected. As discussed below, for the \texttt{frozen-lake} environment one even finds \( \mu_p = \mathcal{O}(1) \). The complexity in Eq. (26) assumes an efficient loading routine for the matrices. To achieve efficient loading also for the policy matrix, a QRAM data structure for the policy needs to be constructed. This needs to happen in time \( \mathcal{O}(|S||A|) \) for each policy-evaluation step. Afterwards, the matrix can be loaded efficiently for each cycle of the measurement protocol.

**Classical Subroutine: Policy Improvement.** The policy improvement step on a classical device requires reading out the \( Q \)-vector from the quantum computer after matrix inversion. Naively, one would expect that the measurement process introduces exponential overhead. However, since convergence results for the Bellman equations are based on the maximum norm \( (L_\infty \text{ norm}) \), the authors employ \( L_\infty \)-norm state tomography [KP18]. This is efficient, i.e. requires \( \mathcal{O}(1/\epsilon^2) \) shots, where \( \epsilon \) now is the target accuracy for the optimal \( Q \)-values (under \( L_\infty \)-norm). Consequently, the overall time complexity (neglecting logarithmic terms) of the algorithm is
\[
\mathcal{O}(|S||A| + \mu_p \Gamma /\epsilon^2) .
\] (27)

In Eq. (27) the factor \( 1/\epsilon^2 \) appears in the second term since the matrix inversion subroutine is called for each of the \( 1/\epsilon^2 \) shots. The first term is the classical complexity of calculating the \( \text{argmax} \) function for policy improvement and construction of the policy oracle prior to each evaluation step.

**Example Environments.** The authors consider the \texttt{FrozenLake} and the \texttt{InvertedPendulum} environments as examples. We will briefly discuss the insights from the former here: The simple form of the environment allows choosing \( \mu_p = 1/2 \), which thus is independent of the size of the action and state space. Note that the gate complexity is still of the order of \( |S||A| \). It only becomes efficient for special structured instances of the environment such as all ‘holes’ on the diagonal of the grid.

**Quantum advantage.** The leading term in Eq. (27) is linear in \( |S| \) and \( |A| \), showing a speed-up with respect to classical linear-system of equations solvers. These exhibit complexity \( \mathcal{O}((|S||A|)^\omega) \), with \( \omega > 1 \), and vanilla \( Q \)-value iteration with complexity \( \mathcal{O}(|S|^2|A|) \). Even though a more detailed characterization of possible quantum advantage is not provided in Ref. [CKP22], it is clear that the speed up can be at most polynomial.

**Least-Squares Policy Iteration.** Finally, the authors generalize the method to least-squares policy iteration [LP03], where the \( Q \)-vector is approximated by a set of basis functions. For details refer to Refs. [LP03; CKP22].
Quantum Policy Iteration via Amplitude Estimation and Grover Search - Towards Quantum Advantage for Reinforcement Learning, Wiedemann et al. (2022)

Summary. In the QRL scheme proposed in Refs. [Wie+22a; Wie21], a policy is evaluated by constructing a superposition of all possible trajectories of an MDP with fixed-horizon and with finite action and state space. Making use of amplitude estimation [Bra+02], the number of calls to a state-transition oracle for estimation of the value function (up to some fixed additive error) can be quadratically reduced. A second algorithm finds the optimal policy in the policy space quadratically faster compared to direct policy search by means of Grover’s algorithm.

First Algorithm. The first algorithm assumes access to a policy oracle $\Pi$ and an environment oracle $E$ which act on an initial state $|s\rangle$ as

$$
\Pi(|s\rangle|0\rangle_A) = \sum_a \sqrt{\pi(a|s\rangle|s\rangle|a\rangle} 
$$

$$
E(|s\rangle|a\rangle|0\rangle_R|0\rangle_S) = \sum_{r,a,s'} \sqrt{p(r,s'|s,a)r\rangle|s\rangle|a\rangle|r\rangle|s'\rangle}. 
$$

Applying these operators sequentially on partially fresh registers as shown in Fig. 15 results in a superposition of all possible trajectories

$$
|t\rangle = |s_0\rangle|a_0\rangle|r_1\rangle|s_1\rangle\ldots|r_H\rangle|s_H\rangle
$$

where $H$ is the horizon of the MDP, such that the quantum state reads

$$
|\psi^\pi\rangle = \sum_t \sqrt{p_t}|t\rangle|G_t\rangle.
$$

Here, $p_t$ is the probability of trajectory $t$. An additional unitary operator has been applied that calculates the return $G_t$ of trajectory $t$ and encodes the value into an additional register entangled with the corresponding trajectory. The superscript $\pi$ on $|\psi^\pi\rangle$ denotes that the state corresponds to the superposition of trajectories for a given policy $\pi$.

![Figure 15: Sequence of policy and environment operator application to an initial state $s$. This constructs a superposition of all possible trajectories for a fixed horizon MDP as shown in Wiedemann et al. [Wie21].](image)

The next step of the algorithm attaches an ancilla qubit. With bit-by-bit rotations of the state the digital encoding of $G_t$ is transformed into amplitude encoding (assuming here for simplicity $G_t \in [0,1]$). A simple calculation reveals that the probability of finding the ancilla qubit in state $|1\rangle$ is given by the average return, that is the value function of the initial state $s$. With this in mind, the authors propose amplitude estimation [Bra+02]. This involves the phase-estimation algorithm, to extract the value function. While classically sampling from the superposition of trajectories would require $O(1/\epsilon^2)$ preparations of the state, the quantum algorithm achieves the same error with $O(1/\epsilon)$, resulting in a quadratic speed-up. Hereby, $\epsilon$ denotes the fixed additive error to which the value function is to be determined.
Second Algorithm. The second algorithm shown in Ref. [Wie+22a] is a quantum version of direct policy search. The authors propose to create a superposition

$$\frac{1}{\sqrt{|P|}} \sum_{\pi} |\pi\rangle |\psi^{\pi}\rangle$$

where $|\pi\rangle$ is a digital representation of the policy, $|\psi^{\pi}\rangle$ the superposition of all trajectories corresponding to policy $\pi$ as before, and $|P|$ the size of the policy space. Quantum minimum finding [DH96] can now be applied to find the optimal policy (the one with maximal expected return starting from initial state $s$), requiring only $O(\sqrt{|P|})$ preparations of the state. This is opposed by $O(|P|)$ in classical direct policy search. Note, however, that the space of all policies scales as $O(|A|^{|S|})$, where $|A|$ and $|S|$ are the sizes of action and state space, respectively. Consequently, the quantum algorithm scales exponentially worse compared to policy iteration where the Bellman optimality equation is iterated with polynomial complexity in $|S|$ and $|A|$. The method proposed in Refs. [Wie+22a; Wie21] therefore should be seen as a quantum version of direct policy search.

4.6 Quantum Reinforcement Learning with Oracularized Environments

In this final section we summarize work that proposes fully quantum-mechanical approaches to QRL. In the articles we survey below, the environment is a quantum system or oracle that can be queried by superpositions of states and actions. Interactions with a quantum-mechanical agent create superpositions of trajectories as input for subroutines like Grover search, quantum-maximum finding, and amplitude estimation. Provable quantum advantage renders some of these proposals interesting candidates for the post-NISQ era.

| Citation   | First Author | Title                                           |
|------------|--------------|-------------------------------------------------|
| [DTB16]    | V. Dunjko    | Quantum-Enhanced Machine Learning                |
| [DTB15]    | V. Dunjko    | Framework for learning agents in quantum environments |
| [DTB17]    | V. Dunjko    | Advances in quantum reinforcement learning       |
| [HDW21]    | A. Hamann    | Quantum-accessible reinforcement learning beyond strictly epochal environments |
| [Wan+21b]  | D. Wang      | Quantum exploration algorithms for multi-armed bandits |
| [Wan+22]   | Z. Wan       | Quantum Multi-Armed Bandits and Stochastic Linear Bandits Enjoy Logarithmic Regrets |
| [Sag+21a]  | V. Saggio    | Experimental quantum speed-up in reinforcement learning agents |
| [HW22]     | A. Hamann    | Performance analysis of a hybrid agent for quantum-accessible reinforcement learning |
| [Cor18]    | A. Cornelissen | Quantum gradient estimation and its application to quantum reinforcement learning |

Table 6: Work considered for “QRL with Oracularized Environments” (Subsec. 4.6)

Quantum-Enhanced Machine Learning, Dunjko et al. (2016) and related works

Summary. In Ref. [DTB16] and in a more detailed preprint [DTB15] a general framework of an agent-environment interaction where both entities are quantum-mechanical systems is developed.
To query the environment by a superposition of action states (intuitively the agent learns in parallel), clearly the environment must be modeled by some form of an oracle. As it turns out, this oracularization is much more involved than one might naively think. The focus of the work is therefore:

- Formalizing a quantum mechanical version of agent-environment interaction
- Investigation of the classical limit
- Properties of the general quantum mechanical set-up
- Treatment of special oracularizable environments
- Identification of quantum advantage for these environments

General Setup. The interaction between agent and environment is modeled as shown in Fig. 2a in Ref. [DTB15]. The register $R_A$ processes the computations of the agent, while the register $R_E$ represents the environment. The communication register stores one action and one state. The interaction is described by completely positive trace preserving (CPTP) maps or, if we wish, unitary maps on a larger system. The first map $M_E^1$ outputs the initial state and stores it into the communication register. The map $M_A^1$ (modeling the agent) reads this state and, after some processing on $R_A$, outputs an action state which is added to the communication register. Now this action processed by $M_E^2$, which outputs a new state. Consecutively, the previous state in the communication register is overwritten, and so on. The particular form of the states of $R_C$ (if in superpositions of action or not) will be discussed later.

While $R_C$ only contains a state-action pair, the agent’s register stores all previous states and actions (because the next action proposed by the learning algorithm depends on all actions and states encountered before, note here the distinction between algorithm and policy). The same is true for the (in general non-Markovian) environment.

Next, as shown in Fig. 16, a tester register $R_T$ is introduced, which is designed to ‘observe’ the elapsed history (all encountered states and actions during a learning sequence). This copying from $R_C$ to $R_T$ is modeled by controlled unitaries (so they do not modify $R_C$). Each of them act on a fresh part of the register $R_T$.

The term copying the register here means that a superposition of computational basis states is concatenated with a second register, on which then each basis state is copied to. This produces in general a highly entangled state, which cannot be factorized into the initial state on the first register and a copy on the second (note the no-cloning theorem only rules out a transformation producing this factorized copy for a general initial state). The most general form of the tester interaction treated in this work allows additional unitary transformations, such that the copying can be described in the form of controlled unitaries. A tester interaction that merely copies the states will be referred to as classical.

After training, the register $R_T$ contains the sequence of actions and states, the so-called history. Any metric measuring performance of learning can be phrased as a function of the histories and their probabilities. Therefore, it can be formulated as the expectation value of an observable on the $R_T$ register.

Classical Limit. For recovering the classical learning set-up, the notion of classical interaction is defined by restricting the form of the maps, such that the state in $R_A - R_C - R_E$ remains separable (note that no entanglement between the registers does not prohibit entangled agent or environment states, thus quantum mechanical environments and agents equipped with a quantum computer are not excluded). Additionally, the tester interaction is supposed to be classical (in the sense as defined above). For this setup it is shown that for every scenario with separable register state there exists a classical environment and a classical agent that produce the same history. Consequently, no quantum improvements are possible. Hence, there can be no improvement in the figure of merit, even when the agent has access to a quantum computer.
Figure 16: Adding a tester as proposed in Dunjko et al. [DTB16].

**General Quantum-Mechanical Set-Up.** What happens when we allow general maps and general states on the registers? The authors prove that the state on $R_T$ is still an incoherent mixture, and therefore no quantum advantage can be expected. The reason for this result lies in the memory of agent and possibly the environment: The agent in general has to remember all previous encountered states and actions, because the learning algorithm run by the agent is a function of that particular elapsed history. The quantum state therefore is a superposition of histories entangled with a state, which describes the agent that has seen this particular history. The states of this agent are orthogonal, since a different agent state translates into a different bit state of the memory. Thus, when tracing out these degrees of freedoms, the resulting reduced density matrix on $R_T$ is an incoherent mixture and no quantum advantage can be achieved in the figure of merit. (Side remark: This does not exclude a quantum advantage in terms of computational complexity in the internal processing of the agent. The result is about exploiting the ‘quantumness’ of the environment-agent interaction)

We note that one has to be careful with the interpretation of density matrices. One might be inclined to think that an incoherent mixture of history states weighted by their probability in some sense corresponds to traversing all of the histories simultaneously but note that the correct expectation value with respect to this density matrix is only obtained in the limit of infinitely many runs corresponding to sampling trajectories one after another.

**Oracularization of Environments.** The next part of the work focuses on a special class of environments and learning setting without memory, which overcome the decoherence problem. These oracularized environments are of the following form:

- episodic with fixed horizon $\rightarrow$ fixed sequence of interactions
- deterministic $\rightarrow$ action sequence fully determines the history, states can be disregarded
- binary rewards issued at final state $\rightarrow$ allows use of Grover search

**Quantum Advantage.** With these assumptions a proper oracle can be constructed, that can be queried with a superposition of actions. This allows to use it as a phase flip oracle, as in the Deutsch-Jozsa or Grover algorithm. The time required for finding a rewarded-action sequence is therefore quadratically reduced. Consequently, this setting is meaningful for learning tasks, where the reward is very sparse. That is, the agent cannot learn until it has first seen a reward. After this initial exploration phase, the agent can now be further trained in simulation. Finally, some of the assumptions are relaxed. The authors also show, how stochastic oracles can be constructed.

**Further Work.** There is further work that builds upon the results of Refs. [DTB15; DTB16]. In Ref. [DTB17], the algorithm is applied to the optimization of parameters describing the properties of the agent (hyperparameter). It also discusses the notion of register hijacking, where the agent has access to hidden memory registers of the environment. This assumption allows the oracularization of more general environments. This class is further generalized in Ref. [HDW21] beyond episodic environments. A closer investigation of amplitude amplification techniques for the special case of multi-armed bandits environments is conducted in Refs. [Wan+21b; Wan+22]. In Ref. [Sag+21a], the learning setting is implemented experimentally for a two-qubit system and an experimental
quantum advantage is observed. Finally, the performance of an agent in this setting is investigated in Ref. [HW22].

**Quantum gradient estimation and its application to quantum reinforcement learning, Cornelissen (2018)**

Summary. The master’s thesis [Cor18] considers model-based RL and develops quantum algorithms for policy evaluation and policy optimization. For the former method a quadratic improvement in sample complexity is found.

Quantum Policy Evaluation. A quantum algorithm for quantum policy evaluation is presented in Sec. 6.2 of the thesis and will be summarized in the following: The algorithm is executed on a register that is capable to store $T$ states and actions of a $T$-step MDP. To generate a sequence, a transition-probability oracle and a policy oracle are defined. They generate a superposition of all possible action-state sequences of the Markov problem, weighted by the square root of the corresponding probabilities. Note that the state is normalized as the probabilities sum up to one. Next, a reward oracle is defined which, when acting on a state-action pair, multiplies the state with a phase factor. The phase is the discounted reward for this state action pair. The discount factors are introduced by making use of fractional phase oracles. This is discussed in detail in Sec. 4 and 5 of the thesis, which are based on Refs. [GAW19; Gil+19]. The fractional reward oracle is applied to every state-action pair in the register, resulting in the product of phase factors containing the individual discounted rewards. Thus, when merging the exponentials to one exponential, the full quantum state is a superposition of all state-action sequences, weighted by the square root of the individual probability and a phase factor containing the corresponding return. Consequently, the expected return is a high-dimensional polynomial in the parameters $x_{sa}$. By that definition, the policy is properly normalized and all $x_{sa} \in [0,1]$. Consequently, the expected return is a high-dimensional polynomial in the parameters $x_{sa}$. For taking the derivative of this objective, Jordan’s quantum gradient algorithm [Jor05] in its advanced form [GAW19] is employed. This leads to a finite-difference approximation of the gradients, written in a phase factor, which can be read out after applying phase estimation. Following Ref. [Gil+19], significant amount of work is devoted to transform the probability oracle for the policy and the transition matrix described above into a phase oracle. Once the superposition of state-action sequences is prepared, an oracle call multiplies each state in the superposition by the corresponding discounted reward. Consecutively, the gradient estimation algorithm is applied and the gradients can be read out. This step is followed by adapting the policy through gradient ascent. It is concluded in the thesis that this policy optimization algorithm does not necessarily lead to quantum speed-up. However, as the author argues, it is conceivable that modifications and improvement of the algorithm might lead to a quantum speed-up.

5 Outlook

We have given a rather detailed account of the various instances QRL that have appeared throughout the literature. We observed, that the dichotomy found at the hardware level, i.e., currently available
NISQ devices vs. fault-tolerant and error-corrected QPUs, manifests also at the algorithmic level. With NISQ devices in mind, VQCs have been suggested as function approximators. These replace their classical counterparts in RL algorithms with function approximation in policy space, value space, or both. Here, one typically replaces a classical learning heuristic by a learning heuristic with a quantum component. Any sort of potential quantum advantage, however, is not immediately apparent. We eventually can obtain theoretical insight into the properties of VQCs viewed as ML models and function approximators. However, a direct comparison to their classical cousins, such as neural networks, is anything but easy and might strongly depend on the chosen metric. How can we meaningfully deploy an agent trained with VQC-components? What are the requirements for quantum advantage in such a heuristic setting? What does non-simulability of quantum circuits imply for e.g. generalization bounds of VQCs as ML models? Can we scale VQCs while maintaining their desirable properties? What is the intrinsic inductive bias of VQCs viewed as ML models? What are the implications for RL and its application domains? All these questions are currently being investigated in the research community, and we are looking forward to new results.

While quantum algorithms for fault-tolerant and error-corrected QPUs have been put forward, we are still far from being able to deploy these algorithms for meaningful problem sizes. Given the necessary advancements of hardware platforms, it will be exciting to see whether these types of quantum algorithms will become competitive with classical learning approaches in practice.

We hope that our survey on the QRL literature and the various types of QRL algorithms will help guide newcomers to the field and will serve as a valuable reference for researchers.

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List of Abbreviations

CNN convolutional neural network. 19, 35
CPTP completely positive trace preserving. 48
CTDE centralized training with decentralized execution. 29, 30
CV continuous-variable. 31
DDQL double deep Q-learning. 16–19, 21
DistRL distributional reinforcement learning. 32, 33
DL deep learning. 3
DLP discrete logarithm problem. 21
DNN deep neural network. 11, 16, 22, 27
DQL deep Q-learning. 16, 17, 35, 37
DRL deep reinforcement learning. 3, 11, 15
FIM Fisher information matrix. 23, 24
MARL multi-agent reinforcement learning. 29
MDP Markov decision process. 3, 4, 33, 44, 46, 50
ML machine learning. 2, 7, 10, 11, 21, 23, 36, 39, 51
MPS matrix product state. 36, 37
MSE mean square error. 16
NISQ noisy intermediate-scale quantum. 2, 3, 10–12, 22, 35, 37, 43, 45, 47, 51
NN neural network. 3, 6, 10, 12, 16, 18–20, 23, 24, 28, 29, 32–34, 36–39, 41, 42
PDF probability density function. 4, 21
POMDP partially observable Markov decision process. 33
PPO proximal policy optimization. 37, 38
PS projective simulation. 11, 12, 40, 41, 43
QC quantum computing. 2, 3, 7, 12, 27, 31, 35, 43
QCNN quantum convolutional neural network. 19
QDDPG quantum deep deterministic policy gradient. 26, 27
QiRL quantum-inspired reinforcement learning. 2, 11, 13, 14, 35, 37, 38
QMARL quantum multi-agent reinforcement learning. 28–31
QML quantum machine learning. 7, 10, 11, 21–23
QNN  quantum neural network. 30–32, 36
QPG  quantum policy gradient. 21, 22
QPU  quantum processing unit. 3, 11, 16, 51
QR   quantile regression. 32
QRL  quantum reinforcement learning. 2, 3, 7, 10–12, 14, 15, 20, 28, 30, 33, 37, 40, 41, 43, 46, 47, 50, 51
RL   reinforcement learning. 2–5, 7, 10–14, 16, 18, 20–22, 24–28, 30–33, 35–37, 39, 41–43, 50, 51
SAC  soft actor-critic. 27, 28, 30, 31
TD   temporal difference. 13, 32
TN   tensor network. 35–37
TSP  traveling salesman problem. 39
VQA  variational quantum algorithm. 11
VQC  variational quantum circuit. 2, 6, 7, 10–12, 14, 16–31, 33–39, 51
VQE  variational quantum eigensolver. 26
VRP  vehicle routing problem. 38, 39
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