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BSc THESIS

Implementing Quantum Fourier Transform via variational quantum circuits

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ΕΘΝΙΚΟ ΚΑΙ ΚΑΠΟΔΙΣΤΡΙΑΚΟ ΠΑΝΕΠΙΣΤΗΜΙΟ ΑΘΗΝΩΝ

**ΣΧΟΛΗ ΘΕΤΙΚΩΝ ΕΠΙΣΤΗΜΩΝ
ΤΜΗΜΑ ΠΛΗΡΟΦΟΡΙΚΗΣ ΚΑΙ ΤΗΛΕΠΙΚΟΙΝΩΝΙΩΝ**

ΠΤΥΧΙΑΚΗ ΕΡΓΑΣΙΑ

**Υλοποίηση Κβαντικού Μετασχηματισμού Fourier μέσω
μεταβαλλόμενων κβαντικών κυκλωμάτων**

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ABSTRACT

Variational quantum circuits are quantum circuits which contain gates with adjustable parameters. Such circuits are already physically realizable in small scale and there is a wide range of possible applications for these promising structures. In this thesis, I develop the idea of tuning a variational quantum circuit to simulate the important for quantum computing, operation of Quantum Fourier Transform. I use algebraic arguments, so called an ansatz, for reducing the depth of the variational quantum circuit and use different classical algorithms to optimize the parameters. The results of this thesis concerning 3-qubit circuits can be possibly extended to a higher number of qubits.

SUBJECT AREA: Quantum computing

KEYWORDS: quantum circuits, quantum Fourier transform, optimization algorithms

ΠΕΡΙΛΗΨΗ

Τα μεταβαλλόμενα κβαντικά κυκλώματα είναι κυκλώματα που περιέχουν κβαντικές πύλες, με παραμέτρους που μεταβάλλονται κατά τη διάρκεια της εκτέλεσης. Τα συγκεκριμένα κυκλώματα είναι ήδη φυσικά εφαρμόσιμα σε μικρή κλίμακα και υπάρχει μια ευρεία γκάμα δυνατών εφαρμογών για αυτά. Σε αυτήν την εργασία, αναπτύσω την ιδέα της υλοποίησης ενός μεταβαλλόμενου κβαντικού κυκλώματος για να προσομοιώσω μια σημαντική για τον κβαντικό υπολογισμό λειτουργία, τον Κβαντικό Μετασχηματισμό Fourier (Quantum Fourier Transform). Χρησιμοποιώ με αλγεβρικά δεδομένα, έναν επονομαζόμενο ανσάτζ (ansatz) ώστε να μειώσω το βάθος του ποιοτικού κβαντικού κυκλώματος και χρησιμοποιώ διάφορους κλασικούς αλγορίθμους για να βελτιστοποιήσω τις παραμέτρους. Τα αποτελέσματα αυτής της εργασίας σχετικά με τα κυκλώματα τριών qubit μπορούν να επεκταθούν πιθανώς σε περισσότερα qubits.

ΕΥΧΑΡΙΣΤΙΕΣ

Η ολοκλήρωση της πτυχιακής αυτής εργασίας θα ήταν αδύνατη χωρίς την πολύτιμη υποστήριξη της καθηγήτριάς μου, Κατερίνας Μανδηλαρά, χάρη στην οποία επέλεξα να στραφώ προς το συγκεκριμένο θέμα, αλλά και να καταφέρω να μάθω τόσα πολλά αυτό το ακαδημαϊκό έτος. Της εκφράζω ένα βαθύ ευχαριστώ για όλη τη βοήθεια, και το κίνητρο που μου προσέφερε, αλλά και τον πολύτιμο χρόνο που μου διέθεσε.

Θα ήθελα ακόμη να ευχαριστήσω τους φίλους μου Σταυρούλα και Νικηφόρο, που έκαναν αισθητά πιο ευχάριστη αυτή την διαδικασία, και με στήριξαν καθ'όλη τη διάρκεια.

Δεν θα μπορούσα να παραλείψω τον Αλέξανδρο, και τον Θέμη, καθώς στις εβδομαδιαίες μας συναντήσεις, αλλά και σε τυχαίες στιγμές, με βοηθούσαν με ό,τι τρόπο μπορούσαν σε οποιαδήποτε δυσκολία προέκυπτε.

Επίσης, θα ήθελα να ευχαριστήσω τον Κωνσταντίνο, γιατί από την πρώτη στιγμή πίστεψε σε μένα, αλλά και για την κατανόηση, και την συμπαράσταση που μου προσέφερε.

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1. INTRODUCTION

In the era where quantum technologies become a reality, it is useful to seek into the new opportunities offered to a computer scientist. While the well-known quantum algorithms offering exponential advantage over classical ones are out of the scope of realization because of the requiring quantum resources, a new sort of algorithms has emerged which is suitable for the current experimental status and which is still promising for exhibiting a quantum advantage. These algorithms, the so called Variational Quantum Algorithms (VQA) are hybrid requiring low-depth Variational Quantum Circuits (VQC) but also a classical optimization loop. VQAs are addressing different problems, using different ansatzs, but their underlined structure is the same, resembling the one of a classical neural network where the weights/parameters are trained via a gradient descent method.

In this work, inspired from the work "Quantum Assisted Quantum Compiling"[1] we aim to train of a VQC to simulate approximately the effect of a circuit implementing Quantum Fourier Transform (QFT). We first build an ansatz in order to avoid to work with VQC of random structure and to reduce the depth. Then we show that with this ansatz a relatively low-depth circuit is able to approximate the QFT on three qubits. This result if it is extensible to a higher number of qubits can be of importance, since QFT is the basic block for Shor's quantum algorithm which threatens security of RSA cryptographic scheme. We also study another important aspect for this method and of VQA algorithms in general, that is the appropriate choice of the classical optimization algorithm. We see that for the case under study a gradient descent method performs better than a stochastic one.

In what follows, we first provide the basic elements of a quantum circuit. Then we give some information on the operation of a QFT and provide the initial circuit representation of it for 3 qubits. In chapter 4 an overview about VQCs and VQAs is provided, while finally in chapter 6 we present our results.

2. THE BASIC ELEMENTS OF QUANTUM CIRCUITS

Quantum circuits are a fundamental concept in quantum computation, similar to classical circuits. They consist of a sequence of quantum gates, measurements, qubit initializations, and other actions that enable quantum computation. [2]. It's important to note that quantum circuits differ from classical circuits as they operate on qubits, which can exist in superpositions of states. This allows for the exploration of multiple possibilities simultaneously, which is a key advantage of quantum computation.

2.1 Qubit

The first thing we need to define, is the basic quantum computational unit, the qubit that is the short for quantum bit. While classical bits can only have two possible states (0 or 1), qubits can exist in a superposition of both states simultaneously. This means that a qubit can be in a linear combination of the 0 and 1 states. A qubit is a two-level quantum system, with the two basis qubit states usually represented as $|0\rangle$ and $|1\rangle$. A qubit can be in state $|0\rangle$, $|1\rangle$, or in a superposition of both states as $\cos \theta |0\rangle + \sin \theta e^{i\phi} |1\rangle$. The superposition property allows a quantum computer to be in multiple states at once, which leads to the exponential growth of possible states as the number of qubits increases [3]. It is also convenient for calculations to represent qubit as vectors

$$|0\rangle = \begin{bmatrix} 1 \\ 0 \end{bmatrix}$$

and

$$|1\rangle = \begin{bmatrix} 0 \\ 1 \end{bmatrix}$$

To understand the concept of a qubit, it's helpful to think about examples from the physical world. A simple analogy, is polarized light. Polarized light can be thought of as a qubit because it exist in two mutually exclusive/orthogonal states: vertically polarized or horizontally polarized. However, if in a general state a single measurement will give both two answers, each one with a partial weight. In contrast, a qubit can be asked many different questions, but each question can only have one of two possible outcomes [4].

In practice, qubits are realized using various physical systems, such as the spin of an electron or the polarization of a photon. The spin of an electron is a common example of a qubit. The two levels of the electron's spin can be taken as spin up and spin down, which correspond to the 0 and 1 states of a qubit [5]. Similarly, the polarization of a single photon can be used to represent the 0 and 1 states of a qubit.

It's important to note that qubits are not limited to two-level systems. Qudits and qutrits are terms used to describe quantum systems with more than two levels. A qudit is a unit of quantum information that can be realized in suitable d-level quantum systems, where d is an integer [6].

The behavior of qubits is governed by the principles of quantum mechanics, such as superposition and entanglement. Superposition allows qubits to exist in multiple states simultaneously, while entanglement enables the correlation of qubits even when they are physically separated. These properties are fundamental to quantum computing and enable the potential for exponential speedup in certain computational tasks compared to classical computers.

We can use the Bloch sphere to represent the state of a single qubit, see Figure 6.4. Any state in a quantum computation can be represented as a vector that begins at the origin and terminates on the surface of the unit Bloch sphere. By applying unitary operators to the state vectors, we can move the state around the surface of the sphere. We take as convention that the poles of the sphere are $|0\rangle$ on the top and $|1\rangle$ on the bottom [7].

We can also represent superposition states such as $\frac{|0\rangle + |1\rangle}{\sqrt{2}}$ that lies along the X axis on Figure 6.4.

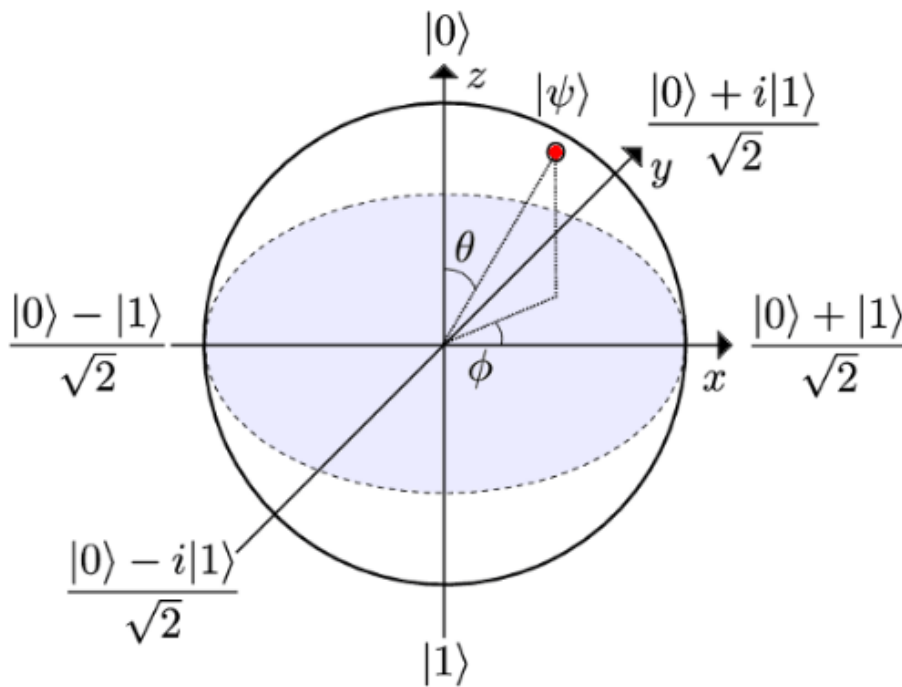


Figure 2.1: The Bloch Sphere

[8]

2.2 Quantum Gates

The building blocks of quantum circuits, are the quantum gates. They are operations performed on qubits, such as rotations, flips, and entangling gates. Quantum gates can manipulate the quantum state of the qubits, enabling various computations and transformations. Quantum logic gates can be derived from classical logic gates, but the Hilbert-space structure of qubits allows for an infinite number quantum gates that are not induced by classical gates.

Quantum gates are described as unitary operators, represented by unitary matrices relative to some basis. The action of a quantum gate on a qubit can be represented by a matrix multiplication of the gate's unitary matrix with the qubit's state vector. Hermitian gates, such as the Pauli gates, Hadamard gate, CNOT gate, SWAP gate, and Toffoli gate, are examples of gates that are both Hermitian and unitary.

The representation of Pauli gates and other important gates as matrices is the following, [2],:

- Pauli-X gate (NOT gate):

$$X = \sigma_x = \sigma_1 = \begin{bmatrix} 0 & 1 \\ 1 & 0 \end{bmatrix}$$

- Pauli-Y gate:

$$Y = \sigma_y = \sigma_2 = \begin{bmatrix} 0 & -i \\ i & 0 \end{bmatrix}$$

- Pauli-Z gate:

$$Z = \sigma_z = \sigma_3 = \begin{bmatrix} 1 & 0 \\ 0 & -1 \end{bmatrix}$$

- Pauli-I gate (identity gate):

$$I = \sigma_4 = \begin{bmatrix} 1 & 0 \\ 0 & 1 \end{bmatrix}$$

- Hadamard gate:

$$H = \frac{1}{\sqrt{2}} \begin{bmatrix} 1 & 1 \\ 1 & -1 \end{bmatrix}$$

We will be mentioning these gates a lot as we move on in this project. Lets study an example of the X-gate matrix applied to a qubit in state $|0\rangle$:

$$X|0\rangle = \begin{bmatrix} 0 & 1 \\ 1 & 0 \end{bmatrix} \begin{bmatrix} 1 \\ 0 \end{bmatrix} = \begin{bmatrix} 0 \\ 1 \end{bmatrix} = |1\rangle$$

Unlike classical logic gates, quantum gates are reversible. This means that the input state can be recovered from the output state by applying the same gate in reverse. Reversibility is a fundamental property of quantum gates and is a consequence of the unitary nature of quantum operations. Reversibility allows for the efficient simulation of classical computation using quantum circuits.

To prove it, we will apply the X-gate on the previous result, and we will notice that the initial input is the initial state:

$$X|1\rangle = \begin{bmatrix} 0 & 1 \\ 1 & 0 \end{bmatrix} \begin{bmatrix} 0 \\ 1 \end{bmatrix} = \begin{bmatrix} 1 \\ 0 \end{bmatrix} = |0\rangle$$

2.2.1 Single Qubit Gates

In the field of quantum computing, single qubit gates and 2-qubit gates play a crucial role in manipulating and transforming the quantum states of qubits. These gates are used to perform operations on individual qubits and multiple qubits, respectively.

Starting with single qubit gates, they act on individual qubits and can be used to change their state or perform specific operations. Some commonly used single qubit gates, are the ones we mentioned before, the Pauli-X gate (bit-flip), Pauli-Y gate (bit and phase flip), Pauli-Z gate (phase flip), Hadamard gate (superposition), and the identity gate. Each of these gates is represented by a unitary matrix and operates on the state vector of a single qubit. We showed their application above. Single qubit gates can be physically realized using techniques such as laser pulses or microwave pulses to manipulate the state of individual qubits [9].

2.2.2 2-Qubit and 3-qubit Gates

Moving on to 2-qubit gates, these act on two qubits simultaneously inducing interaction between them. The controlled-NOT (CNOT) gate is a commonly used 2-qubit gate. It performs a controlled operation where the second qubit (target qubit) is flipped if and only if the first qubit (control qubit) is in the state $|1\rangle$. In matrix representation in the computational basis, controlled-NOT (CNOT) gate writes as

$$CNOT = \begin{bmatrix} 1 & 0 & 0 & 0 \\ 0 & 1 & 0 & 0 \\ 0 & 0 & 0 & 1 \\ 0 & 0 & 1 & 0 \end{bmatrix}$$

$$CNOT(|0\rangle \otimes |0\rangle) = \begin{bmatrix} 1 & 0 & 0 & 0 \\ 0 & 1 & 0 & 0 \\ 0 & 0 & 0 & 1 \\ 0 & 0 & 1 & 0 \end{bmatrix} \begin{bmatrix} 1 \\ 0 \\ 0 \\ 0 \end{bmatrix} = \begin{bmatrix} 1 \\ 0 \\ 0 \\ 0 \end{bmatrix} = |00\rangle$$

Lets apply the CNOT gate on $|10\rangle$:

$$CNOT(|1\rangle \otimes |0\rangle) = \begin{bmatrix} 1 & 0 & 0 & 0 \\ 0 & 1 & 0 & 0 \\ 0 & 0 & 0 & 1 \\ 0 & 0 & 1 & 0 \end{bmatrix} \begin{bmatrix} 0 \\ 0 \\ 1 \\ 0 \end{bmatrix} = \begin{bmatrix} 0 \\ 0 \\ 0 \\ 1 \end{bmatrix} = |11\rangle$$

As we can see, now that the control qubit is $|1\rangle$ the target qubit becomes from $|0\rangle$, $|1\rangle$. Concerning 3-qubit gates, the extension of controlled-NOT (CNOT) gate is the Toffoli gate, which is a reversible gate that performs a NOT operation on the target qubit if both control qubits are in the state $|1\rangle$. Similar to 2-qubit gates, 3-qubit gates can be physically implemented using techniques controlling interactions between qubits. Providing a couple of examples:

- Toffoli gate:

$$\begin{bmatrix} 1 & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\ 0 & 1 & 0 & 0 & 0 & 0 & 0 & 0 \\ 0 & 0 & 1 & 0 & 0 & 0 & 0 & 0 \\ 0 & 0 & 0 & 1 & 0 & 0 & 0 & 0 \\ 0 & 0 & 0 & 0 & 1 & 0 & 0 & 0 \\ 0 & 0 & 0 & 0 & 0 & 1 & 0 & 0 \\ 0 & 0 & 0 & 0 & 0 & 0 & 1 & 0 \\ 0 & 0 & 0 & 0 & 0 & 0 & 0 & 1 \end{bmatrix}$$

- Fredkin gate(controlled-swap):

$$\begin{bmatrix} 1 & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\ 0 & 1 & 0 & 0 & 0 & 0 & 0 & 0 \\ 0 & 0 & 1 & 0 & 0 & 0 & 0 & 0 \\ 0 & 0 & 0 & 1 & 0 & 0 & 0 & 0 \\ 0 & 0 & 0 & 0 & 1 & 0 & 0 & 0 \\ 0 & 0 & 0 & 0 & 0 & 1 & 0 & 0 \\ 0 & 0 & 0 & 0 & 0 & 0 & 1 & 0 \\ 0 & 0 & 0 & 0 & 0 & 0 & 0 & 1 \end{bmatrix}$$

In the context of quantum computing, gates can be combined in series or in parallel to create more complex operations and circuits. Series combinations of gates involve applying one gate after another, while parallel combinations involve applying gates simultaneously to different qubits. The order of gates in a circuit diagram is reversed to the the multiplication procedure written in formulas.

To sum it up, single qubit gates, 2 and 3 qubit gates are fundamental components in quantum computing. They are used to manipulate and transform the quantum states of qubits, enabling the implementation of various quantum algorithms and protocols. Single qubit gates act on individual qubits, while 2 qubit gates allow for interactions between pairs of qubits. These gates can be combined to create more complex operations and circuits, as we will see later.

2.2.3 Variational Gates

Another category of gates, is what we call variational gates. These are gates with trainable/free parameters within, which are used in variational circuits. In this project, I use both single, and 2-qubit variational gates. Since, my results concern 3-qubit circuits I will explain here, via the generators of the $su(8)$ algebra of 3 qubits, the variational gates that I will use later. One can create the 64 generators of the group by taking the Kronecker product of 4 single qubit operators (X-gate, Y-gate, Z-gate and Identity gate). This way we can have up to 64 different generators $\hat{g}_{i,j,k}$ with $i, j, k = 1, \dots, 4$. For example a generator can be $\hat{g}_{1,4,4} = \hat{X} \otimes 1 \otimes 1$ or $\hat{g}_{1,4,2} = \hat{X} \otimes 1 \otimes \hat{Y}$ etc. Note that 1 represents the Identity matrix. The variational gates of the circuit are constructed by exponentiation of this generator $\hat{g}_{j,j,k}$ multiplied by a parameter x , as

$$\hat{R}_{\hat{g}_{j,j,k}}(x) = e^{ix\hat{g}_{j,j,k}}$$

In geometric space such gates $\hat{R}_{\hat{g}_{j,j,k}}(x)$ perform a rotation around the axis that is defined by the generator $\hat{g}_{j,j,k}$ while the rotation angle is proportional to the parameter x . Further information on this matter, is provided in chapter 5. Let me provide some examples, the variational gate

$$\hat{R}_{\hat{X} \otimes 1 \otimes 1}(x) = e^{ix\hat{X} \otimes 1 \otimes 1} \quad (2.1)$$

is a single qubits variational gate while

$$\hat{R}_{\hat{X} \otimes 1 \otimes \hat{Y}}(x) = e^{ix\hat{X} \otimes 1 \otimes \hat{Y}} \quad (2.2)$$

is a 2-qubit gate acting on the first and third qubit.

Let me here clarify how exponentiation and Kronecker product works using the examples in Eqs.(2.1)-(2.2). Knowing that

$$X = \begin{bmatrix} 0 & 1 \\ 1 & 0 \end{bmatrix}, \quad I = \begin{bmatrix} 1 & 0 \\ 0 & 1 \end{bmatrix}$$

then Kronecker product is computed as follows:

$$X \otimes I = \begin{bmatrix} 0 & 1 \\ 1 & 0 \end{bmatrix} \otimes \begin{bmatrix} 1 & 0 \\ 0 & 1 \end{bmatrix} = \begin{bmatrix} 0 & 0 & 1 & 0 \\ 0 & 0 & 0 & 1 \\ 1 & 0 & 0 & 0 \\ 0 & 1 & 0 & 0 \end{bmatrix}$$

$$X \otimes I \otimes I = \begin{bmatrix} 0 & 0 & 1 & 0 \\ 0 & 0 & 0 & 1 \\ 1 & 0 & 0 & 0 \\ 0 & 1 & 0 & 0 \end{bmatrix} \otimes \begin{bmatrix} 1 & 0 \\ 0 & 1 \end{bmatrix} = \begin{bmatrix} 0 & 0 & 0 & 0 & 1 & 0 & 0 & 0 \\ 0 & 0 & 0 & 0 & 0 & 1 & 0 & 0 \\ 0 & 0 & 0 & 0 & 0 & 0 & 1 & 0 \\ 0 & 0 & 0 & 0 & 0 & 0 & 0 & 1 \\ 1 & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\ 0 & 1 & 0 & 0 & 0 & 0 & 0 & 0 \\ 0 & 0 & 1 & 0 & 0 & 0 & 0 & 0 \\ 0 & 0 & 0 & 1 & 0 & 0 & 0 & 0 \end{bmatrix}$$

The corresponding single-qubit variational gate can be computed by the formula

$$\hat{R}_{\hat{X} \otimes 1 \otimes 1}(x) = e^{ix\hat{X} \otimes 1 \otimes 1} = \exp \left(ix \begin{bmatrix} 0 & 0 & 0 & 0 & 1 & 0 & 0 & 0 \\ 0 & 0 & 0 & 0 & 0 & 1 & 0 & 0 \\ 0 & 0 & 0 & 0 & 0 & 0 & 1 & 0 \\ 0 & 0 & 0 & 0 & 0 & 0 & 0 & 1 \\ 1 & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\ 0 & 1 & 0 & 0 & 0 & 0 & 0 & 0 \\ 0 & 0 & 1 & 0 & 0 & 0 & 0 & 0 \\ 0 & 0 & 0 & 1 & 0 & 0 & 0 & 0 \end{bmatrix} \right)$$

using spectral theorem or Taylor expansion.

Similarly for the equation 2.2 the generator is the Kronecker product of a X-Pauli gate, with the Identity gate, and the Y-Pauli gate. Following the same process as before, we can write this 2-qubit variational gate as

$$\hat{R}_{\hat{X} \otimes 1 \otimes \hat{Y}}(x) = e^{ix\hat{X} \otimes 1 \otimes \hat{Y}} = \exp \left(ix \begin{bmatrix} 0 & 0 & 0 & 0 & 0 & -i & 0 & 0 \\ 0 & 0 & 0 & 0 & i & 0 & 0 & 0 \\ 0 & 0 & 0 & 0 & 0 & 0 & 0 & -i \\ 0 & 0 & 0 & 0 & 0 & 0 & i & 0 \\ 0 & -i & 0 & 0 & 0 & 0 & 0 & 0 \\ i & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\ 0 & 0 & 0 & -i & 0 & 0 & 0 & 0 \\ 0 & 0 & i & 0 & 0 & 0 & 0 & 0 \end{bmatrix} \right)$$

2.3 Measurements

Measurement is another important aspect of quantum circuits. Measurements are used to extract information from qubits. They collapse the quantum state of a qubit to one of its basis states, and the measurement result is a classical value that can be used in classical computations. It's important to note that measurement in quantum mechanics is probabilistic, and the outcome of a measurement cannot be predicted with certainty but the probabilities can be inferred if the state of qubits is known.

2.4 Quantum circuits

Now using a combination of the gates mentioned before, we will create a circuit. Specifically, we will provide the a circuit, that entangles 3 qubits. As we can see, the circuit is composed of a Hadamard gate(H), followed by two $CNOT$ gates, and lastly the measurements of the 3 qubits. [11]



Figure 2.2: GHZ states circuit

GHZ states are named after Greenberger, Horne, and Zeilinger, who were the first to study them in 1989. GHZ states are also known as “Schrödinger cat states” or just “cat states.” [12]

3. QUANTUM FOURIER TRANSFORM (QFT)

One of the most useful ways of solving a problem in mathematics or computer science is to transform it into some other problem for which a solution is known. A great discovery of quantum computation has been that some such transformations can be computed much faster on a quantum computer than on a classical computer, a discovery which has enabled the construction of fast algorithms for quantum computers.

The Quantum Fourier Transform is a reversible transformation that operates on qubits and is used to convert a quantum state from the computational basis to the Fourier basis. It is implemented using a series of unitary gates, such as the Hadamard gate and the controlled phase shift gate [2]. The QFT is particularly powerful when combined with other algorithms, as it can be used to measure the period of a function, which is essential for cracking the RSA algorithm.

The algorithm implements a Discrete Fourier transform on the values of the amplitudes. As a reminder, the classical version of the Fourier transform takes as input a vector of complex numbers x_0, x_1, \dots, x_{N-1} , where the length N of the vector is a fixed parameter. It outputs the transformed data, a vector of complex numbers y_0, y_1, \dots, y_{N-1} , defined by

$$y_k \equiv \frac{1}{\sqrt{N}} \sum_{j=0}^{N-1} x_j e^{\frac{2\pi i j k}{N}} \quad (3.1)$$

The Quantum Fourier transform is exactly the same transformation, although the conventional notation for the quantum Fourier transform is somewhat different. The quantum Fourier transform on an orthonormal basis $|0\rangle, \dots, |N-1\rangle$ is defined to be a linear operator with the following action on the basis states

$$|j\rangle \rightarrow \frac{1}{\sqrt{N}} \sum_{k=0}^{N-1} x_j e^{\frac{2\pi i j k}{N}} |k\rangle \quad (3.2)$$

Equivalently, the action on an arbitrary state may be written

$$\sum_{j=0}^{N-1} x_j |j\rangle \rightarrow \sum_{k=0}^{N-1} y_k |k\rangle \quad (3.3)$$

where the amplitudes y_k are the discrete Fourier transform of the amplitudes x_j [2]. In other words, the quantum Fourier transform implements a discrete Fourier transform on the values of the amplitudes of an input state. This transformation is a unitary transformation, and thus can be implemented via a circuit of a quantum computer [20].

3.1 QFT on 3 qubits

QFT operation has been proposed together with a circuit implementation where the number of gates increases approximately quadratically with the number of qubits. On the other

hand this circuit, as we are going to show, contains Controlled-Phase gates whose design and implementation are hard in general. The QFT with 3 qubits requiring 6 gates, see Figure 6.4, is the first non-trivial case, and for this reason we have chosen to work with.

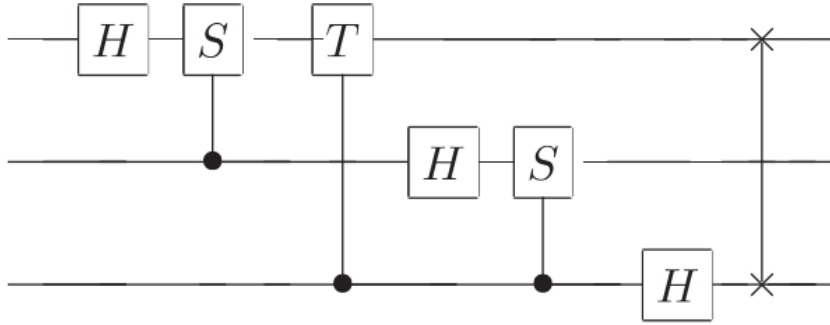


Figure 3.1: Quantum Fourier Transform Circuit

Recall that S and T are the phase and $\pi/8$ gates, and H is the Hadamard gate which we mentioned in the introductory sections [2].

- S gate(phase gate):

$$\begin{bmatrix} 1 & 0 \\ 0 & i \end{bmatrix}$$

- T gate($\pi/8$ gate):

$$\begin{bmatrix} 1 & 0 \\ 0 & e^{i\pi/4} \end{bmatrix}$$

It's important to note that controlled phase shift gates, which include Controlled- T and Controlled- S gates, are essential in the construction of a QFT circuit.

As with any single qubit gate one can build a controlled version of the phase shift gate. With respect to the computational basis, the 2-qubit controlled phase shift gate, shifts the phase with φ only if it acts on the state $|11\rangle$ [21].

$$|a, b\rangle \mapsto \begin{cases} e^{i\varphi}|a, b\rangle & \text{for } a = b = 1 \\ |a, b\rangle & \text{otherwise} \end{cases}$$

Controlled Phase gate:

$$CR(\phi) = \begin{bmatrix} 1 & 0 & 0 & 0 \\ 0 & 1 & 0 & 0 \\ 0 & 0 & 1 & 0 \\ 0 & 0 & 0 & e^{i\phi} \end{bmatrix}$$

The controlled- T gate is a specific case of the controlled phase gate(CR), where the phase angle is set to $\phi = \pi/4$.

$$CT = \begin{bmatrix} 1 & 0 & 0 & 0 \\ 0 & 1 & 0 & 0 \\ 0 & 0 & 1 & 0 \\ 0 & 0 & 0 & e^{i\frac{\pi}{4}} \end{bmatrix}$$

The controlled- S gate is the case where the phase angle is set to $\phi = \pi/2$ [2].

$$CS = \begin{bmatrix} 1 & 0 & 0 & 0 \\ 0 & 1 & 0 & 0 \\ 0 & 0 & 1 & 0 \\ 0 & 0 & 0 & i \end{bmatrix}$$

As a matrix the quantum Fourier transform in this instance may be written out explicitly, by introducing for simpler notation $\omega = e^{\frac{2\pi i}{8}} = \sqrt[4]{i}$ [2]

$$\hat{U}_{QFT} = \frac{1}{\sqrt{8}} \begin{bmatrix} 1 & 1 & 1 & 1 & 1 & 1 & 1 & 1 \\ 1 & \omega & \omega^2 & \omega^3 & \omega^4 & \omega^5 & \omega^6 & \omega^7 \\ 1 & \omega^2 & \omega^4 & \omega^6 & 1 & \omega^2 & \omega^4 & \omega^6 \\ 1 & \omega^3 & \omega^6 & \omega^1 & \omega^4 & \omega^7 & \omega^2 & \omega^5 \\ 1 & \omega^4 & 1 & \omega^4 & 1 & \omega^4 & 1 & \omega^4 \\ 1 & \omega^5 & \omega^2 & \omega^7 & \omega^4 & \omega^1 & \omega^6 & \omega^3 \\ 1 & \omega^6 & \omega^4 & \omega^2 & 1 & \omega^6 & \omega^4 & \omega^2 \\ 1 & \omega^7 & \omega^6 & \omega^5 & \omega^4 & \omega^3 & \omega^2 & \omega^1 \end{bmatrix}$$

=

$$\frac{1}{\sqrt{8}} \begin{bmatrix} 1 & 1 & 1 & 1 & 1 & 1 & 1 & 1 \\ 1 & e^{i\frac{\pi}{4}} & e^{i\frac{\pi}{2}} & e^{i\frac{3\pi}{4}} & e^{i\pi} & e^{i\frac{5\pi}{4}} & e^{i\frac{3\pi}{2}} & e^{i\frac{7\pi}{4}} \\ 1 & e^{i\frac{\pi}{2}} & e^{i\pi} & e^{i\frac{3\pi}{2}} & 1 & e^{i\frac{\pi}{2}} & e^{i\pi} & e^{i\frac{3\pi}{2}} \\ 1 & e^{i\frac{3\pi}{4}} & e^{i\frac{3\pi}{2}} & e^{i\frac{9\pi}{4}} & e^{i\pi} & e^{i\frac{5\pi}{4}} & e^{i\frac{7\pi}{2}} & e^{i\frac{15\pi}{4}} \\ 1 & e^{i\pi} & 1 & e^{i\pi} & 1 & e^{i\pi} & 1 & e^{i\pi} \\ 1 & e^{i\frac{5\pi}{4}} & e^{i\frac{\pi}{2}} & e^{i\frac{7\pi}{4}} & e^{i\pi} & e^{i\frac{\pi}{4}} & e^{i\frac{\pi}{2}} & e^{i\frac{3\pi}{4}} \\ 1 & e^{i\frac{3\pi}{2}} & e^{i\pi} & e^{i\frac{7\pi}{2}} & 1 & e^{i\frac{\pi}{2}} & e^{i\pi} & e^{i\frac{3\pi}{2}} \\ 1 & e^{i\frac{7\pi}{4}} & e^{i\frac{3\pi}{2}} & e^{i\frac{15\pi}{4}} & e^{i\pi} & e^{i\frac{3\pi}{4}} & e^{i\frac{7\pi}{2}} & e^{i\frac{15\pi}{4}} \end{bmatrix}$$

This representation of the Quantum Fourier Transform, is going to be very useful to us, when we get to the code implementation.

4. VARIATIONAL QUANTUM CIRCUITS AND ALGORITHMS

4.1 Variational Quantum Circuits (VQCs)

Variational quantum circuits, also known as “parametrized quantum circuits” or “adaptable quantum circuits”, are a type of quantum circuit that are used in various applications, such as supervised learning, reinforcement learning, and functional regression. They are designed to harness the potential advantages of quantum computing for solving complex problems. They involve the use of parameterized/variational gates and measurements to encode and manipulate quantum information.

One of the main motivations behind using variational quantum circuits is to take advantage of the expressive power of quantum computing to solve problems that are difficult for classical computers. These circuits are designed to be trainable, meaning that the parameters of the gates can be adjusted to optimize the circuit’s performance for a specific task. The optimization process typically involves finding the set of parameters that minimizes a cost or loss function, which is defined based on the specific problem being solved [13].

One common approach in variational quantum circuits is to use a hybrid model that combines classical and quantum components. This allows for the use of classical optimization algorithms to update the parameters of the circuit based on feedback from the classical part of the model. This hybrid approach helps to mitigate the noise and errors inherent in current quantum hardware, known as NISQ (Noisy Intermediate-Scale Quantum) devices [14].

The training process of variational quantum circuits involves updating the parameters of the gates using gradient descent optimization. The gradients of the loss function with respect to the circuit parameters are calculated using the backpropagation method, similar to the calculation in classical neural networks [15]. The gradients are then used to update the parameters in the direction that minimizes the loss function.

The performance of variational quantum circuits can be evaluated through numerical simulations and experiments on quantum hardware. Simulations can provide insights into the learning performance and efficiency of the circuits, while experiments on real quantum machines, such as IBMQ systems, can test the applicability and scalability of the circuits in real-world scenarios.

Variational quantum circuits are an active area of research, and there are ongoing efforts to explore their capabilities and limitations. Theoretical analysis, such as error performance analysis and optimization properties, can provide insights into the representation and generalization powers of variational quantum circuits. Experimental validation on different datasets and problem domains is crucial for understanding the practical usefulness of these circuits.

4.2 Variational Quantum Algorithms (VQAs)

Variational quantum algorithms (VQAs) are a type of hybrid quantum-classical optimization algorithm that harnesses the capabilities of both classical and quantum computers to solve complex problems [16]. VQAs have emerged as a promising approach in the Noisy Intermediate-Scale Quantum (NISQ) era, where quantum computers are characterized by noise and imperfections.

In VQAs, the objective function is typically encoded by a VQC, which serves as an ansatz or guess for the ground state of the system. The quantum computer evaluates the objective function by measuring the expectation value of an observable, often the Hamiltonian of the system. The classical optimizer then utilizes this evaluation to update the parameters of the circuit, aiming to minimize the associated cost function [16]. Classical optimization methods such as gradient descent can be employed to iteratively adjust the parameters and approach the optimal solution.

It is important to note, that the noise and errors inherent in current quantum hardware present challenges for implementing VQAs. NISQ devices are prone to errors, and the probability of failed operations is non-negligible. Consequently, VQAs often rely on shallow circuits or circuits with a significant fraction of failed operations to ensure reliability. However, this limitation can impact the performance of VQAs and restrict their effectiveness in the presence of noise [17].

One of the various proposed uses of VQCs in combination with an appropriate VQA is the task of quantum compiling [1]. The aim is to train a VQC to simulate the effect of a quantum operation by training the parameters in the variational gates. For this task one does not try to reach the ground state of a Hamiltonian but rather to minimize the overlap of the operation produced by the VQC with the target operation. The corresponding VQA consists of devising ansatzes on the VQA structure and designing an efficient construction of the overlap function that plays the role of the cost function. My work falls into this category of VQAs.

5. QFT VIA A VARIATIONAL QUANTUM CIRCUIT WITH 3 QUBITS

As we mentioned before, QFT is an essential operation in quantum computing. To be realized for n qubits this requires $\approx n^2$ gates that is an excellent record –taking into account that a general unitary operation is described by 4^n real parameters. On the other hand, this includes controlled-phase gates R_n each of which if to be decomposed into gates from a universal set, requires with the best known method up today ≈ 20 gates and an ancillary qubit.

In the NIST era the idea of employing fault-tolerant quantum computation has been set aside since deep-depth circuits are out of discussion. In this work we investigate the question of realizing the QFT operation with a variational circuit of relatively low depth. Instead of working with universal set of gates we employ Hamiltonians which can be in principle physically realizable. The initial procedure that we follow for a 3-qubit is a standard one:

- We build an ansatz on the structure of the circuit based on the algebraic properties of QFT.
- We define a cost function that measures the distance from the target operation i.e. QFT.
- We optimize the parameters in the circuit with a gradient descent so that the cost is minimized.

5.1 Building an Ansatz

In order to build an ansatz we first take the unitary matrix (in the computational basis), \hat{U}_{QFT} , and we decompose it onto the 64 generators of $SU(8)$ algebra, i.e. \hat{g}_{ijk} , as

$$O_{ijk} = \text{Tr} \left(\hat{U}_{QFT} \hat{g}_{ijk} \right). \quad (5.1)$$

We find out that only 20 generators give non-zero overlap O_{ijk} .

We commute these 20 generators and we find out that these together with first, second,.. order commutations form a closed subgroup of 32 elements. Then we search for a minimum set of single-qubit and 2-qubit operators able to generate the whole subgroup. We identify 11 of them (5 single qubit and 6 two-qubit Hamiltonians) but possibly this result could be improved.

We build a parametrized circuit that consists of N parametrized gates \hat{R}_{ijk} generated by these 11 operators, e.g. $\hat{R}_{ijk}(x) = (Ix\hat{g}_{ijk})$. Let us denote by $\hat{U}_c(\vec{x})$ the unitary operation describing the parametrized circuit (consisting by a product of N parametrized gates). By $\vec{x} = \{x_1, \dots, x_N\}$ we denote a vector built by the N parameters of the circuit. The aim is to find \vec{x} such that the overlap cost function is as low as possible:

$$C \left(\hat{U}_c(\vec{x}), \hat{U}_{QFT} \right) = 1 - \frac{1}{64} \left| \text{Tr} \left(\hat{U}_c(\vec{x}) \hat{U}_{QFT}^\dagger \right) \right|^2. \quad (5.2)$$

The cost function C is constructed such a way [1] so that this takes value in the interval $[0, 1]$ with $C = 0$ corresponding to the case where $\hat{U}_c(\vec{x}) = \hat{U}_{QFT}$ up to a global phase. By augmenting N naturally one expects better results since this way more ‘volume’ in the space of unitary matrices can be covered by the VQC. I also note here that the structure of circuit is not crucial provided that some simple logical rules are followed such that there is full connectivity of all qubits and that there is no successive single qubit gates on the same qubit.

5.2 Gradient Descent Optimizer

Before we move on to the code implementation, it is important to give some information about the optimization method I used for this project.

Gradient descent is an iterative optimization algorithm used to find the local minimum of a differentiable function. It is particularly useful in machine learning for minimizing the cost or loss function. The basic idea behind gradient descent is to take repeated steps in the opposite direction of the gradient of the function at the current point, because this is the direction of steepest descent. By iteratively updating the parameters in the direction of the negative gradient, the algorithm gradually converges towards a minimum point [18].

To understand gradient descent, let’s consider an analogy. Imagine a person who is stuck in the mountains and is trying to find the lowest point (i.e., the global minimum). Due to heavy fog, the person cannot see the path down the mountain. Instead, they must rely on local information to find the minimum. In this scenario, the person can use the method of gradient descent. They look at the steepness of the hill at their current position and proceed in the direction with the steepest descent (i.e., downhill). If they were trying to find the top of the mountain (i.e., the maximum), they would proceed in the direction of steepest ascent (i.e., uphill).

In the context of machine learning, gradient descent is used to update the parameters of a model in order to minimize the cost or loss function. The parameters refer to the coefficients in linear regression or the weights in neural networks [18]. The goal is to find the parameter values that minimize the cost function, which represents the discrepancy between the predicted values and the actual values in the training data [19].

There are different variations of gradient descent, including batch gradient descent, mini-batch gradient descent and stochastic gradient descent, which is also tested in this project.

In stochastic gradient descent, the algorithm calculates the gradient of the cost function with respect to a single training example in each iteration and updates the parameters based on this gradient. This method is computationally efficient but can have high variance in the parameter updates, leading to slower convergence. However, it can escape local minima and find better solutions [18].

The choice of which variation of gradient descent to use depends on the specific problem and the available computational resources. For example, batch gradient descent is typi-

cally used when the dataset can fit into memory and computational efficiency is not a major concern. On the other hand, stochastic gradient descent and mini-batch gradient descent are often preferred for large datasets or when computational efficiency is important [19].

6. RESULTS

In order to reach to conclusions, I run multiple tests, for different number of parameters, N , using a different combination of hyperparameters for the gradient descent optimizer. I start from a random initial point $\mathbf{X}_{\text{INITIAL}} = \vec{x}_{init}$ in the parameter space and the aim is to reach by gradient descent an optimum set of parameters $\mathbf{X}_{\text{FINAL}} = \vec{x}_{fin}$ so that the overlap cost function in Eq. (5.2) is minimized.

6.1 Hyperparameters

Another important thing to explain, is the purpose of the hyperparameters in this problem.

- **learning rate/gamma:** Controls how big the steps are taken in each iteration of the gradient descent algorithm. A smaller learning rate results in smaller steps, which might make the optimization process slower but more stable. On the other hand, a larger learning rate can make the optimization process faster but might also result in overshooting the optimal solution.
- **delta:** This is the perturbation value, which is used to calculate the numerical gradient approximation. It represents the small change in the x values used to estimate the gradient of the cost function.
- **epsilon:** It is small value used as a stopping criterion for the optimization process. The optimization stops when the change in cost between two consecutive iterations becomes smaller than epsilon.
- **threshold:** It is used to decide when to adjust the learning rate. If the change in cost between two consecutive iterations becomes smaller than threshold, the learning rate is adjusted using the learning rate step scheduler function.
- **step size:** This parameter determines the size of the adjustment made to the learning rate when it needs to be changed. It controls how much the learning rate is decreased during the optimization process, when the scheduler is called.

Overall, the purpose of these hyperparameters is to control the optimization process, find the optimal x values that minimize the cost function, and adjust the learning rate dynamically during the optimization process for better convergence. The appropriate values for these hyperparameters depend on the specific problem, and tuning them can significantly impact the performance and efficiency of the optimization algorithm.

6.2 18 parameters

A.

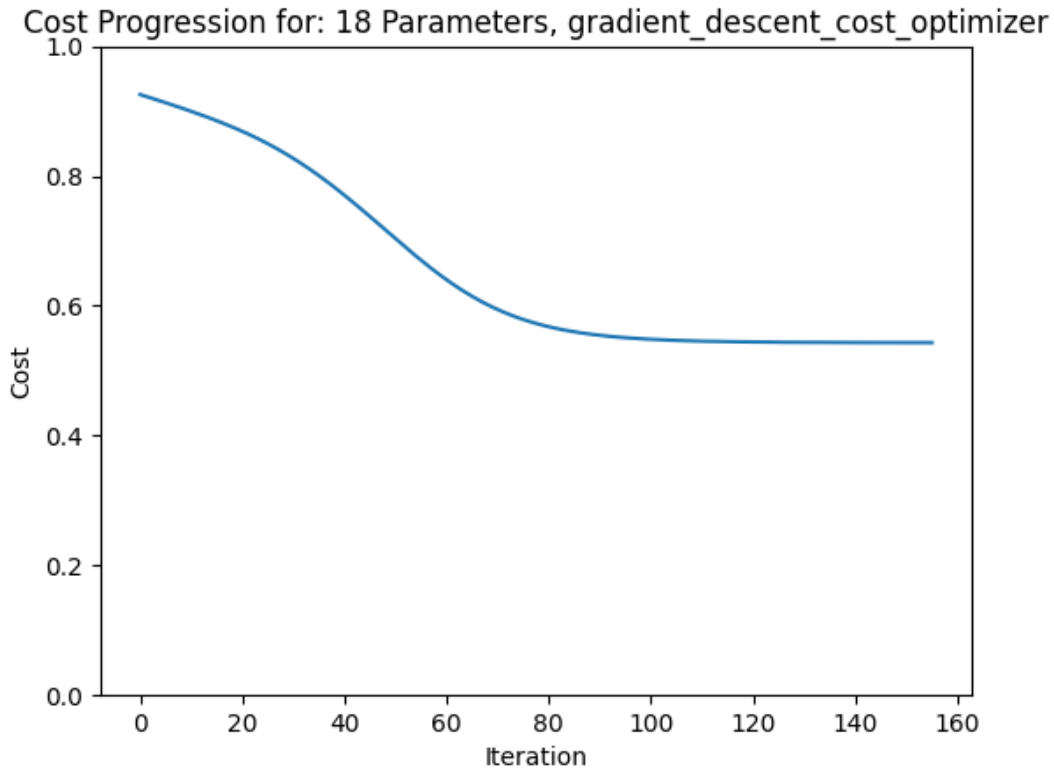


Figure 6.1: Cost progression plot, with 18 trainable parameters, using Gradient Descent optimizer(A)

It is obvious, that in this particular snapshot, the final cost is not the optimal result. Here are the initial and the final parameter values, as well as the initial and final costs.

X INITIAL is: tensor([1.7658, 3.4312, 4.1421, 5.3068, 0.1327, 5.7904, 4.1582, 1.2668, 5.6374, 6.0970, 2.9710, 1.6479, 0.6261, 2.5107, 3.8973, 3.1691, 0.1296, 4.8192])

initial cost: tensor(0.9279)

X FINAL is: tensor([1.4667, 4.2022, 4.2278, 5.2510, 0.1786, 6.2554, 3.9019, 1.3128, 5.7231, 6.0693, 3.4226, 1.6939, 0.3527, 2.5964, 4.2336, 2.5559, 0.2154, 4.8651]),

final cost: tensor(0.5432)

learning rate = 0.05

delta = 0.005

epsilon = 1e-08

threshold = 1e-05

step size = 0.1

More information about how the code is implemented, are in the appendix.

Some other implementations with 18 parameters gave similar results.

B.

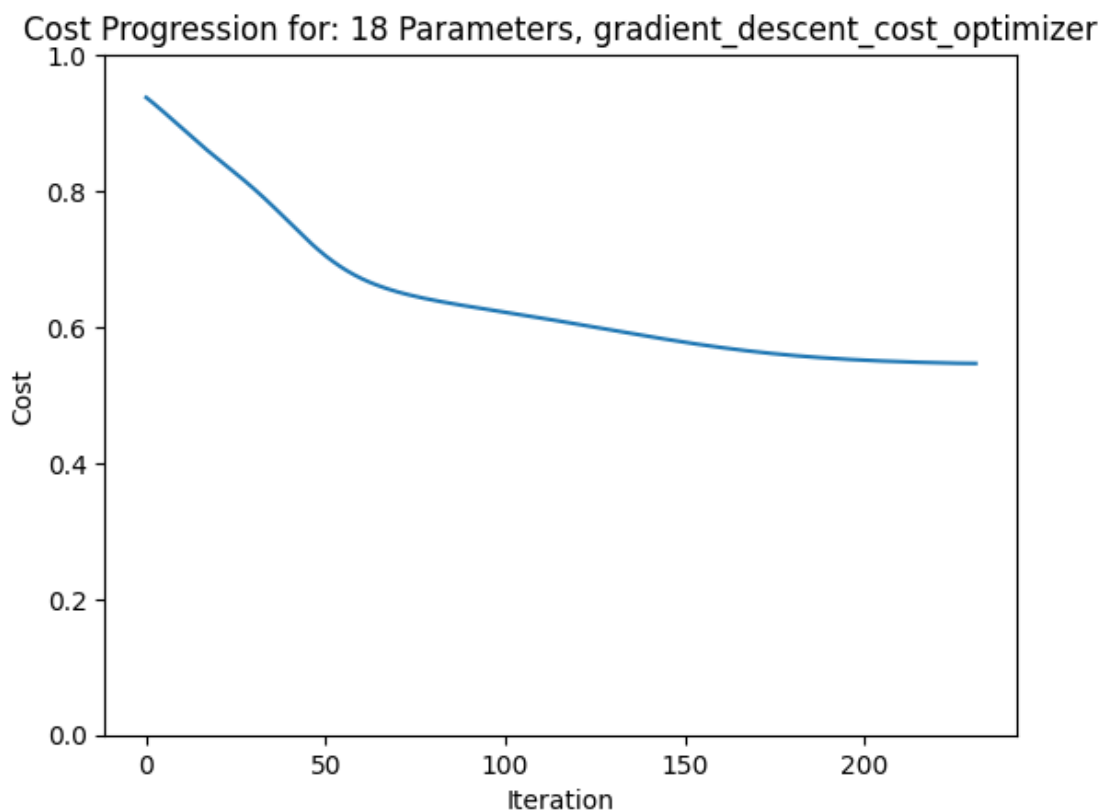


Figure 6.2: Cost progression plot, with 18 trainable parameters, using Gradient Descent optimizer(B)

X INITIAL is: tensor([4.8274, 1.5107, 1.5987, 3.6443, 5.9472, 2.8795, 0.4899, 4.3089, 1.4795, 4.2034, 3.8087, 0.3544, 0.4266, 2.9152, 0.2256, 1.8776, 4.5090, 0.9238])

initial cost: = tensor(0.9416)

X FINAL is: tensor([4.6312, 1.1150, 1.3784, 3.7199, 5.8616, 3.2902, 0.7585, 4.2233, 1.2592, 4.4295, 3.5401, 0.2688, 0.3703, 2.6949, 1.0287, 2.5523, 4.2887, 0.8382])

final cost: tensor(0.5464)

learning rate = 0.05

delta = 0.005

epsilon = 1e-08

threshold = 0.001

step size = 0.1

- C. In this last case, I decided to run the program with different parameters for the gradient descent. Instead of the previous "epsilon" value(1e-08) I choose to increase it to 1e-06.

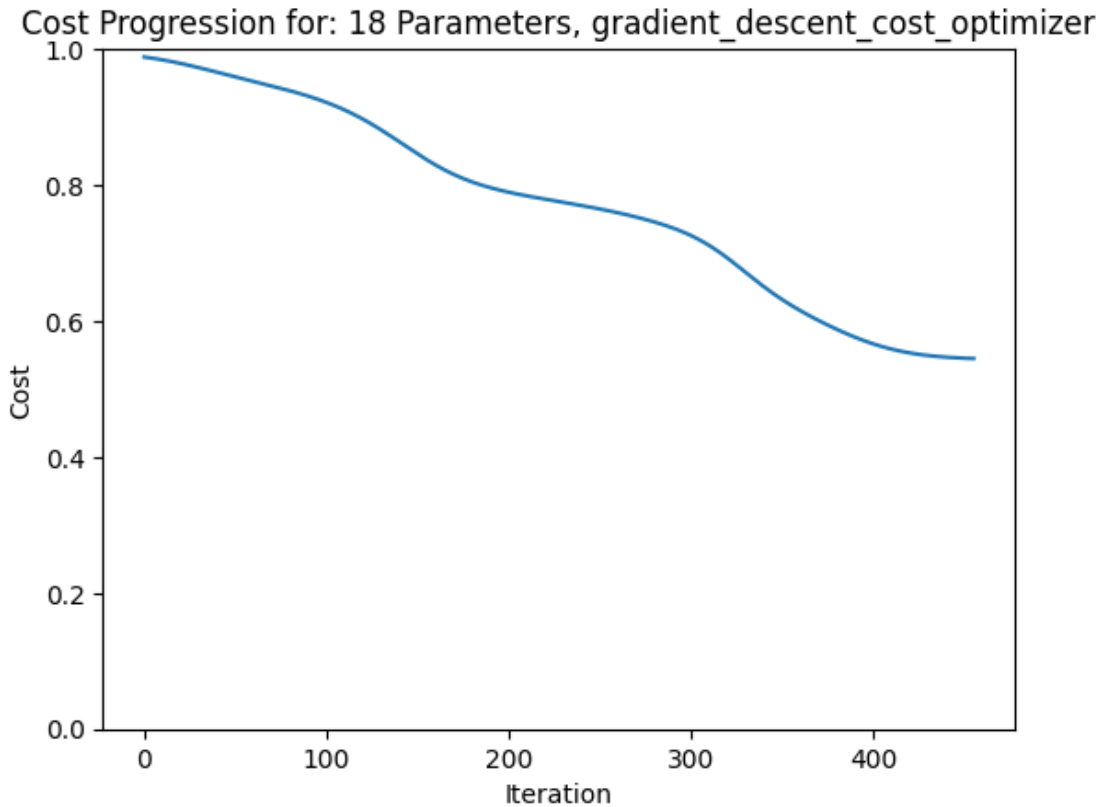


Figure 6.3: Cost progression plot, with 18 trainable parameters, using Gradient Descent optimizer(C)

X INITIAL is: tensor([1.4468, 6.1483, 5.0250, 2.7024, 1.6291, 0.2811, 4.5268, 0.7270, 2.9660, 4.9462, 4.7656, 0.4019, 0.4443, 6.2669, 3.6787, 3.4641, 5.6090, 2.2776])

initial cost: = tensor(0.9887)

X FINAL is: tensor([1.4848, 5.7350, 4.8197, 2.8080, 1.5974, 0.2750, 3.9014, 0.6953, 2.7608, 4.7029, 4.7299, 0.3702, -0.3118, 6.0617, 2.6975, 2.5539, 5.4037, 2.2459])

final cost: tensor(0.5449)

learning rate = 0.05

delta = 0.005

epsilon = 1e-06

threshold = 0.0001

step size = 0.1

After multiple executions with 18 parameters, I realized that the final cost would not get lower than 0.5. As a result, more parameters are needed to train this circuit and minimize the cost.

6.3 20 parameters

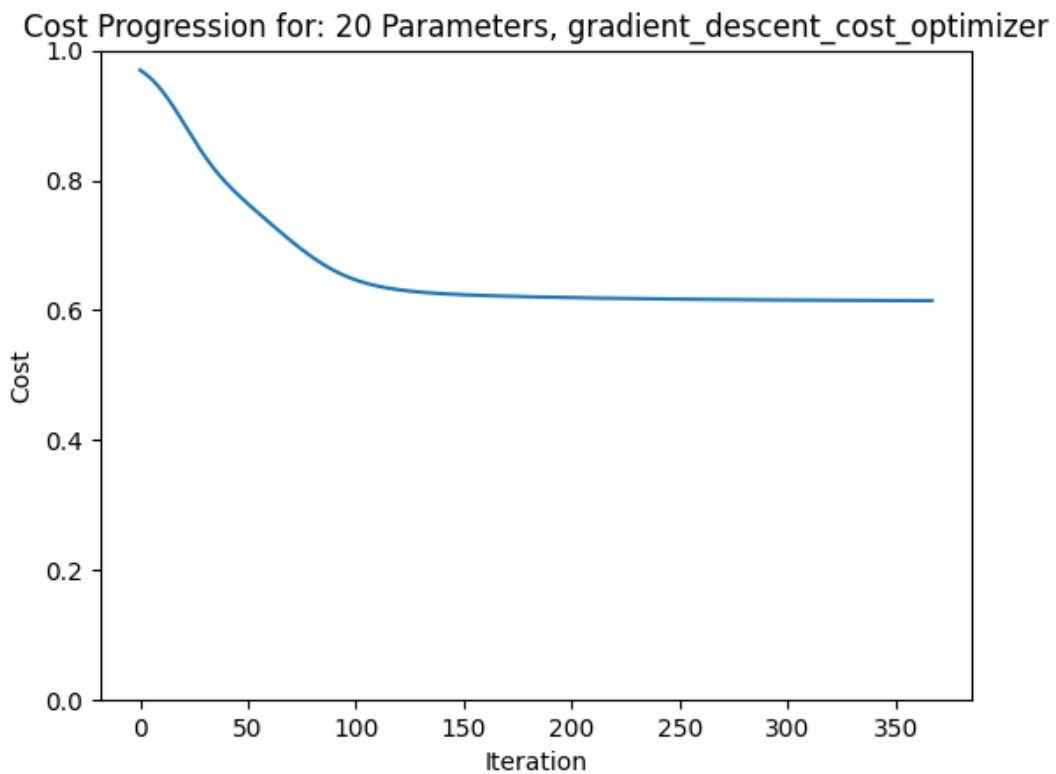


Figure 6.4: Cost progression plot, with 20 trainable parameters, using Gradient Descent optimizer

X INITIAL is: tensor([3.1948, 5.3759, 4.9216, 0.6639, 0.7274, 1.3457, 4.1867, 3.4602, 3.4542, 3.5626, 2.9046, 5.0324, 1.6932, 1.5336, 2.8378, 5.3266, 2.7702, 1.1606, 1.8269, 0.3608])

initial cost: tensor(0.9716)

X FINAL is: tensor([2.8576, 5.6991, 5.1387, 0.9144, 0.7338, 1.1805, 3.8612, 3.4666, 3.6713, 3.6662, 3.3033, 5.0389, 2.3572, 1.7507, 3.0914, 4.9087, 2.9874, 1.1671, 1.9527, 1.1274])

final cost: tensor(0.6147)

learning rate = 0.05

delta = 0.005
 epsilon = 1e-08
 threshold = 0.0001
 step size = 0.1

After multiple executions, I concluded that I need to increase the parameters a little more, since 20 parameters are not enough.

6.4 22 parameters

A.

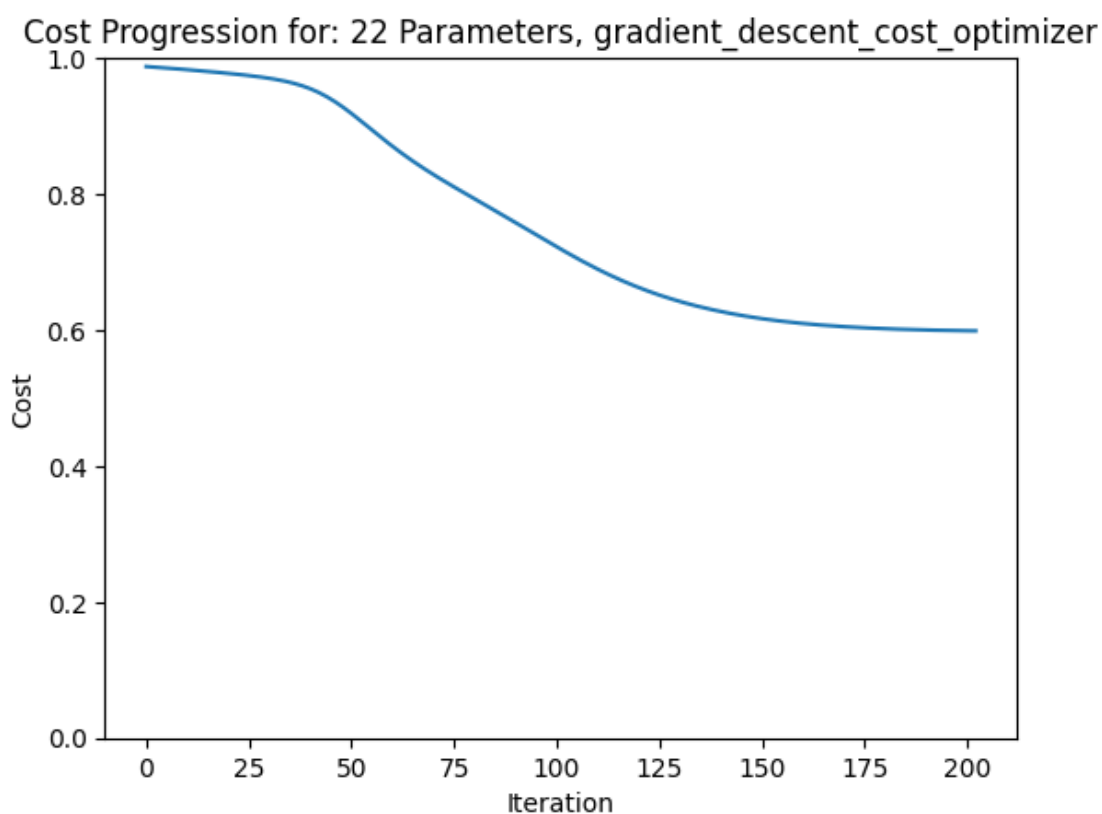


Figure 6.5: Cost progression plot, with 22 trainable parameters, using Gradient Descent optimizer(A)

X INITIAL is: tensor([5.5316, 4.8904, 5.3289, 0.0076, 3.5166, 1.9878, 4.1190, 3.4012, 2.7321, 3.2201, 5.3298, 1.9683, 1.6896, 3.9811, 0.0533, 3.2435, 1.0930, 5.2226, 2.2948, 0.1961, 1.6848, 4.0577])

initial cost: tensor(0.9880)

X FINAL is:

tensor([5.8946, 5.8026, 5.4321, 0.3726, 3.4180, 2.2436, 3.9729, 3.3026, 2.8353, 2.7521, 5.6147, 1.8697, 2.3123, 4.0844, -0.0451, 3.3367, 1.1963, 5.1240, 2.1425, 0.0551, 1.4876, 4.3096])

final cost: tensor(0.5989)

learning rate = 0.05

delta = 0.005

epsilon = 1e-08

threshold = 0.0001

step size = 0.1

B.

Instead of the previous "epsilon" value (1e-08), I chose to increase it to 1e-06.

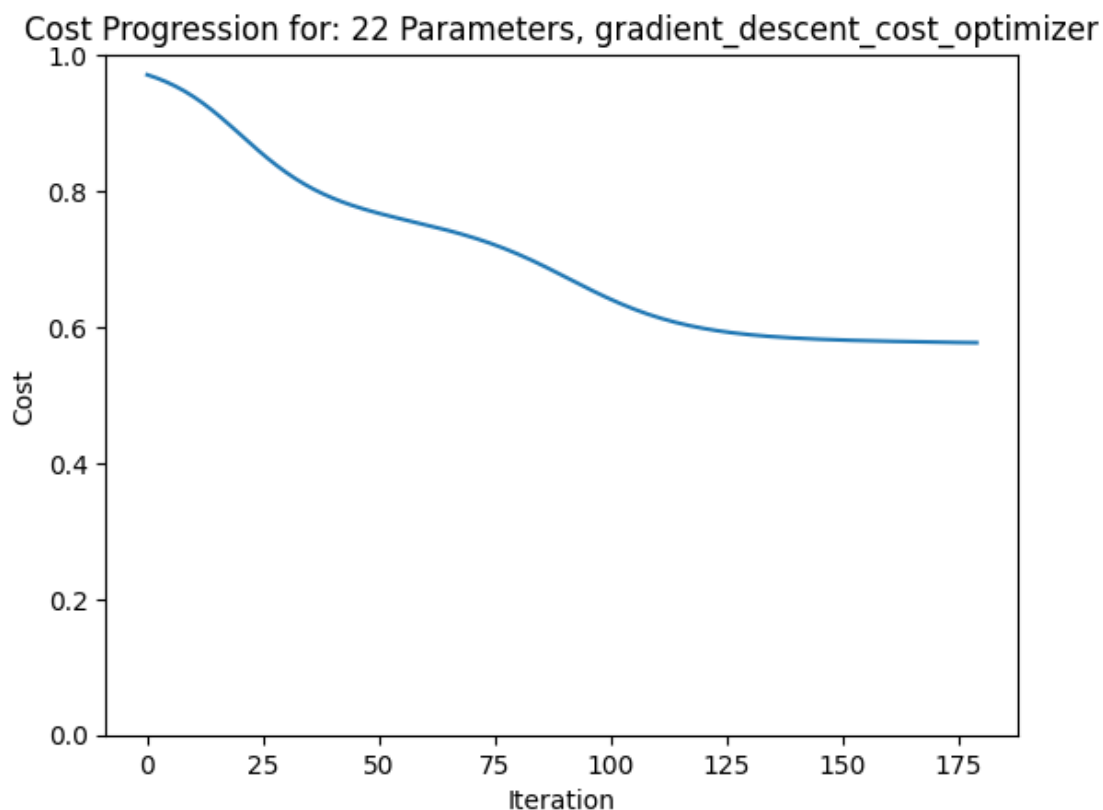


Figure 6.6: Cost progression plot, with 22 trainable parameters, using Gradient Descent optimizer(B)

X INITIAL is: tensor([3.4730, 2.6421, 5.1826, 1.7041, 5.8734, 3.2929, 1.7257, 2.0646, 2.5464, 6.2450, 0.6135, 3.7199, 2.8965, 3.6478, 2.8020, 1.7141, 0.3224, 5.4836, 1.1351, 5.5521, 1.8220, 3.1569])

initial cost: tensor(0.9731)

X FINAL is: tensor([3.0353, 2.8943, 5.0558, 1.5389, 5.8523, 3.0472, 2.2930, 2.0435, 2.4196, 6.0623, 0.5559, 3.6988, 3.1135, 3.5210, 3.0138, 0.9943, 0.1956, 5.4625, 1.4578, 5.3369, 1.7798, 3.2443])

final cost: tensor(0.5770)

learning rate = 0.05

delta = 0.005

epsilon = 1e-06

threshold = 0.0001

step size = 0.1

6.5 26 parameters

A.

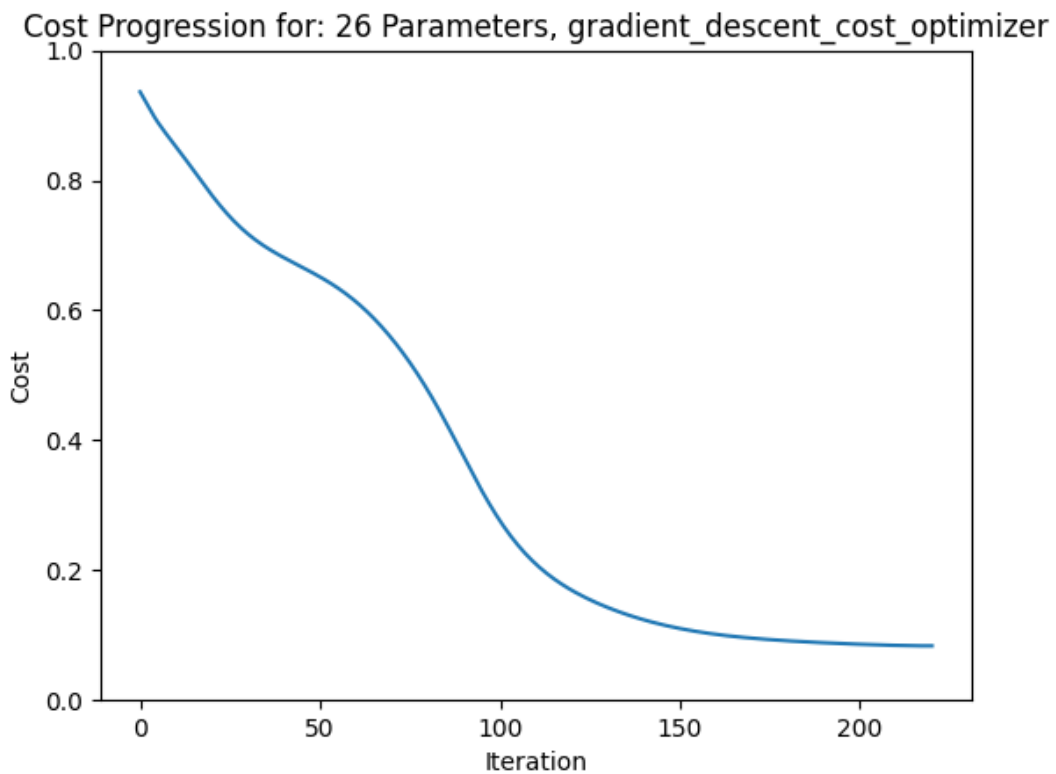


Figure 6.7: Cost progression plot, with 26 trainable parameters, using Gradient Descent optimizer(A)

X INITIAL is: tensor([3.2022, 5.6416, 4.8515, 3.3115, 3.6187, 5.0852, 2.9730, 5.0537, 1.5072, 4.7531, 0.5125, 5.5763, 5.8622, 1.1606, 0.6496, 2.2168, 3.9991, 5.3104,

4.2755, 3.8328, 1.0954, 1.9451, 4.8207, 4.8218, 4.6000, 1.4309])

initial cost: tensor(0.9451)

X FINAL is: tensor([2.3827, 5.2529, 4.6696, 3.5108, 3.6095, 5.0535, 3.9956, 5.2220,
1.1325, 4.1914, 0.7261, 5.7447, 5.8750, 0.7859, 0.9014, 2.5525, 3.6244, 5.4788,
3.8632, 4.2661, 0.9771, 2.2849, 5.1514, 4.4313, 4.2616, 1.4216])

final cost: tensor(0.0836)

learning rate = 0.05

delta = 0.005

epsilon = 1e-08

threshold = 0.0001

step size = 0.1

B.

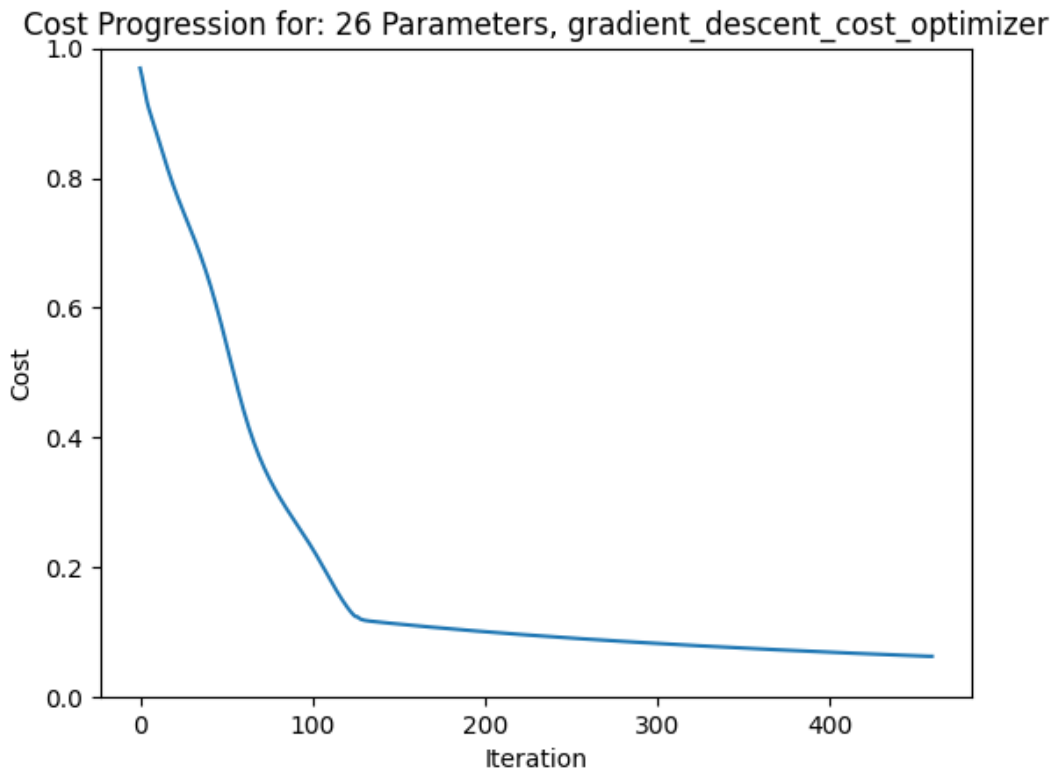


Figure 6.8: Cost progression plot, with 26 trainable parameters(b), using Gradient Descent optimizer(B)

X INITIAL is: tensor([2.8648, 5.4188, 0.7875, 4.8503, 4.0227, 1.7765, 1.3553, 0.6937,
5.7583, 0.8078, 4.9534, 1.9077, 6.2795, 0.9409, 3.3676, 0.6460, 3.6312, 4.4370,
5.8479, 5.9351, 5.4692, 2.7993, 3.0437, 3.2066, 5.3126, 2.4545])

initial cost: tensor(0.9781)

X FINAL is:

tensor([3.3034, 5.5917, 0.8085, 4.6238, 4.3980, 1.5872, 0.9654, 0.7109, 5.4160,
1.0719, 4.8915, 1.9250, 5.3130, 0.5986, 4.1386, 0.9808, 3.2889, 4.4543, 5.9470,
5.9129, 5.4806, 2.7109, 3.6705, 2.8900, 5.2372, 2.8299])

final cost: tensor(0.0631)

learning rate = 0.05

delta = 0.005

epsilon = 1e-08

threshold = 0.0001

step size = 0.1

C. Instead of the previous "epsilon" value(1e-08), I choose to increase it to 1e-06.

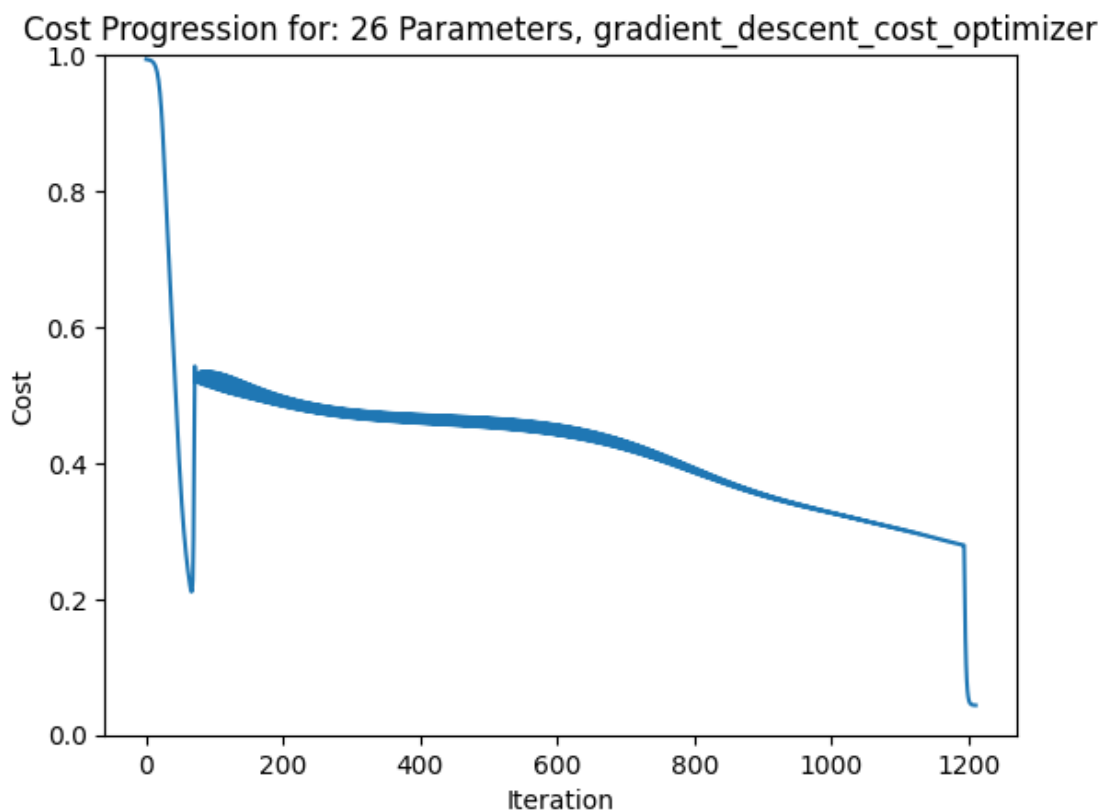


Figure 6.9: Cost progression plot, with 26 trainable parameters(c), using Gradient Descent optimizer(C)

X INITIAL is: tensor([2.8710, 4.7578, 3.3982, 1.5568, 5.3054, 0.7483, 4.1578, 3.5420,
2.4154, 4.7246, 5.3815, 0.3457, 2.7512, 2.6583, 5.3696, 4.0464, 1.9786, 3.5664,
5.2470, 1.9843, 6.2567, 0.6134, 1.5815, 5.8635, 5.9813, 2.5164])

initial cost: tensor(0.9940)

X FINAL is: tensor([1.3361, 5.7477, 3.3366, 2.1055, 5.4184, -0.1233, 4.2182, 3.1592, 2.1595, 5.3936, 5.6810, -0.0371, 1.3140, 2.4024, 5.8358, 4.1234, 1.7227, 3.1837, 5.0808, 2.3511, 6.2768, 0.7038, 1.5699, 6.0155, 5.3044, 2.6294])

final cost: tensor(0.0437)

learning rate = 0.05

delta = 0.005

epsilon = 1e-06

threshold = 0.0001

step size = 0.1

After testing the circuit for 18, 20, and 22 parameters, on 26 parameters, the result is optimal. The cost is minimized and is equal to 0.0437.

6.6 28 parameters

A.

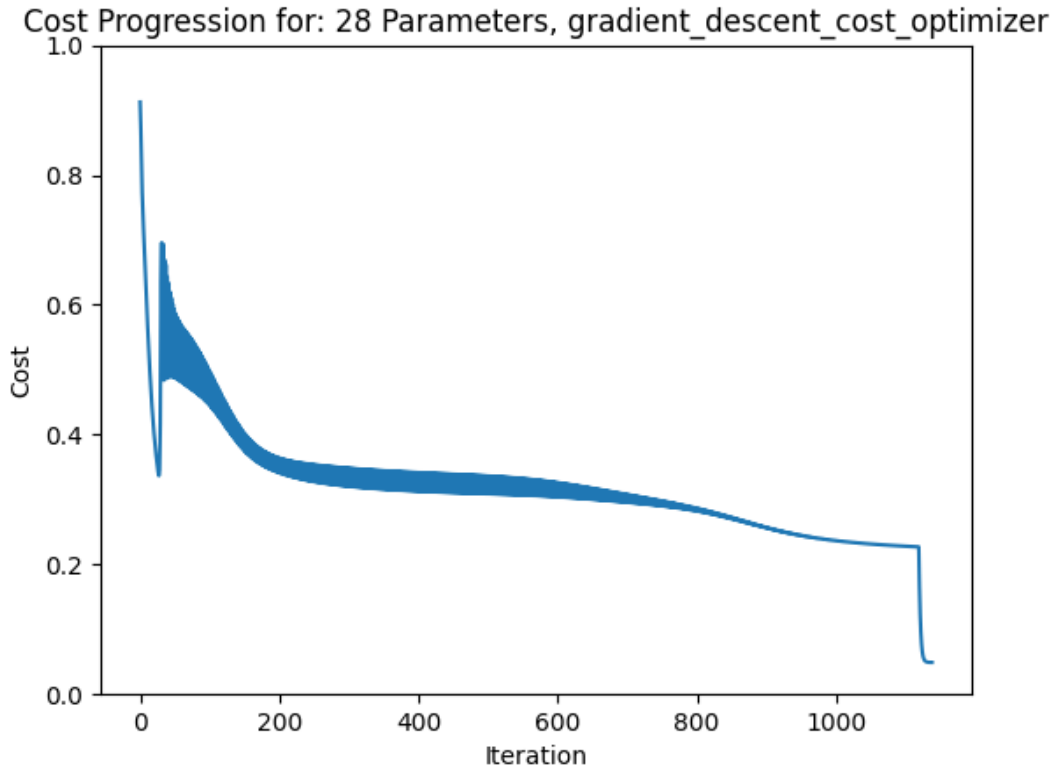


Figure 6.10: Cost progression plot, with 28 trainable parameters, using Gradient Descent optimizer(A)

X INITIAL is: tensor([3.6285, 0.5717, 1.3463, 0.2393, 3.4633, 1.5723, 6.1080, 5.0117, 4.5497, 1.1938, 5.7585, 1.8747, 2.4548, 3.4901, 4.9434, 3.3576, 0.2792, 4.1673, 3.8621, 2.6452, 4.2201, 5.2078, 4.3327, 5.5728, 5.9729, 0.1138, 2.8426, 4.2832])

initial cost: tensor(0.9368)

X FINAL is: tensor([2.7217, 0.4280, 0.1943, 0.3745, 3.3493, 1.4766, 6.7955, 5.2375, 4.6558, 0.7024, 5.9003, 2.1004, 2.4367, 3.5962, 3.9115, 3.3380, 0.3853, 4.3931, 3.5211, 2.5860, 3.9345, 5.3438, 3.9282, 5.4054, 5.5653, -0.0083, 3.1441, 4.0902])

final cost: tensor(0.0491)

learning rate = 0.05

delta = 0.005

epsilon = 1e-08

threshold = 0.0001

step size = 0.1

B.

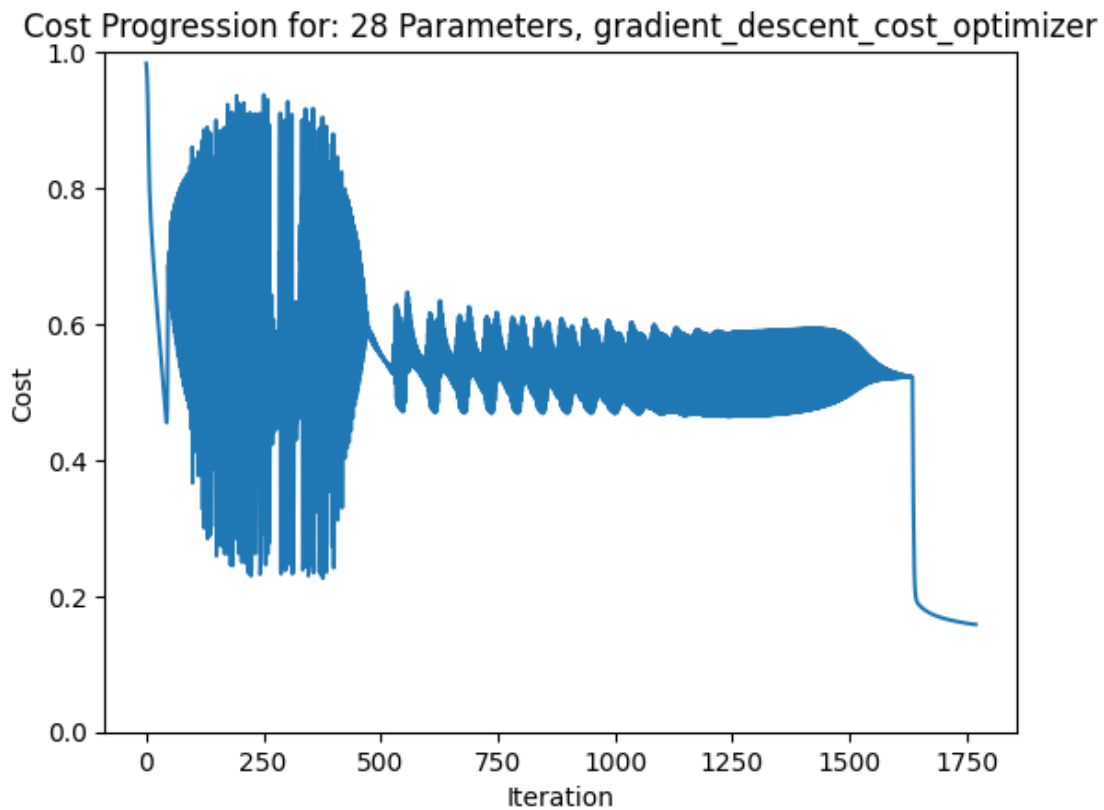


Figure 6.11: Cost progression plot, with 28 trainable parameters, using Gradient Descent optimizer(B)

X INITIAL is: tensor([4.5330, 1.2127, 5.8686, 3.9970, 6.0627, 3.8046, 3.1711, 3.8535, 0.4302, 4.0036, 2.4540, 5.2031, 2.1072, 1.0062, 4.4468, 0.7581, 1.1217, 2.6496, 4.4254, 2.6519, 1.4644, 3.2951, 4.1166, 5.0768, 0.5094, 2.3641, 4.8506, 3.9727])

initial cost: tensor(0.9884)

X FINAL is: tensor([4.9584, 1.3110, 6.6644, 3.9844, 6.1405, 4.0523, 3.8551, 2.9820, 0.6069, 4.5222, 1.5677, 4.3316, 2.9369, 1.1830, 4.8590, 0.1916, 1.2984, 1.7781, 4.4359, 2.7650, 1.8271, 3.5744, 4.3506, 5.0568, 0.6657, 1.9018, 4.3499, 3.3173])

final cost: tensor(0.1581)

learning rate = 0.05

delta = 0.005

epsilon = 1e-08

threshold = 0.0001

step size = 0.1

C. Lets test it with "epsilon" = 1e-06.

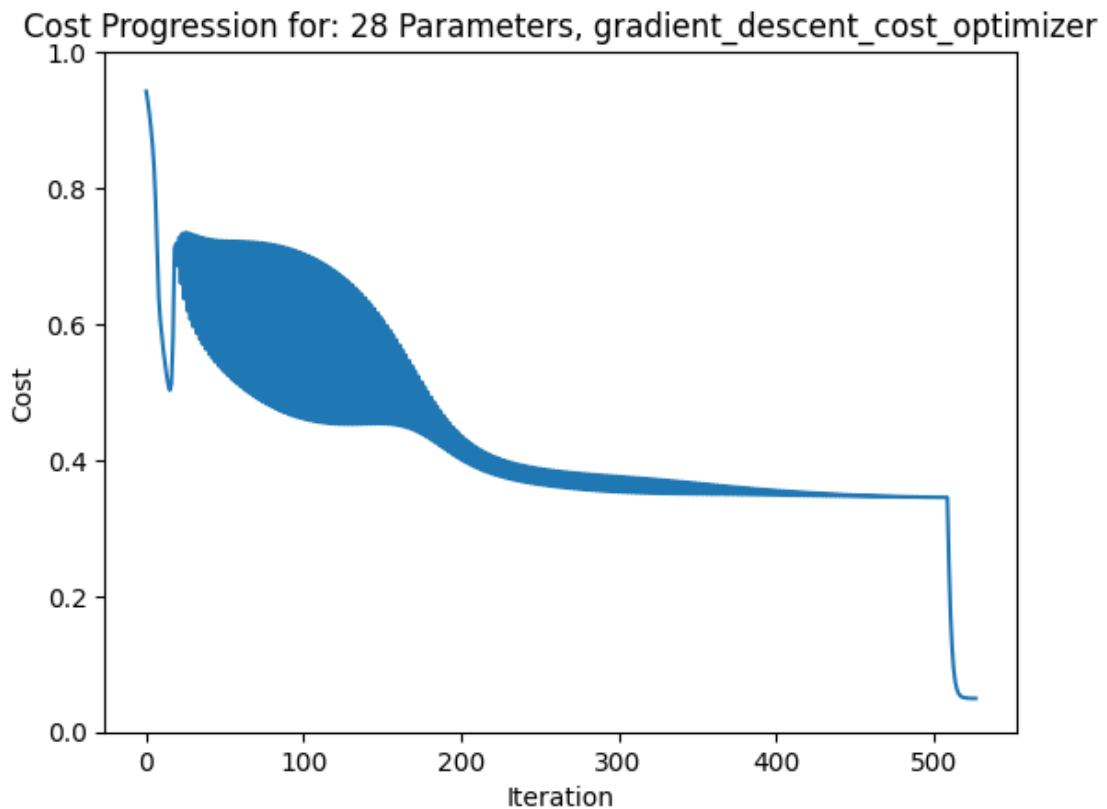


Figure 6.12: Cost progression plot, with 28 trainable parameters, using Gradient Descent optimizer(C)

X INITIAL is: tensor([2.0370, 1.5601, 2.9670, 5.9147, 1.7438, 4.8593, 0.8470, 0.8450, 5.3775, 1.0312, 4.9395, 2.6428, 1.3595, 4.7149, 1.6253, 2.5752, 3.9885, 3.9920, 5.5750, 2.4976, 4.4143, 0.8474, 6.2534, 5.4128, 3.5603, 3.2367, 2.3206, 1.7052])

initial cost: tensor(0.9539)

X FINAL is: tensor([2.4495, 1.5559, 3.4123, 5.9498, 2.6323, 3.9259, 1.4310, 0.4180, 5.1273, 0.5496, 5.4894, 2.2158, 1.0248, 4.4646, 1.7050, 2.5476, 3.7383, 3.5650, 5.6853, 2.5584, 4.7759, 0.8392, 6.1957, 5.5991, 4.2603, 2.7112, 2.9145, 1.8037])

final cost: tensor(0.0494)

learning rate = 0.05

delta = 0.005

epsilon = 1e-06

threshold = 0.0001

step size = 0.1

6.7 36 parameters

A.

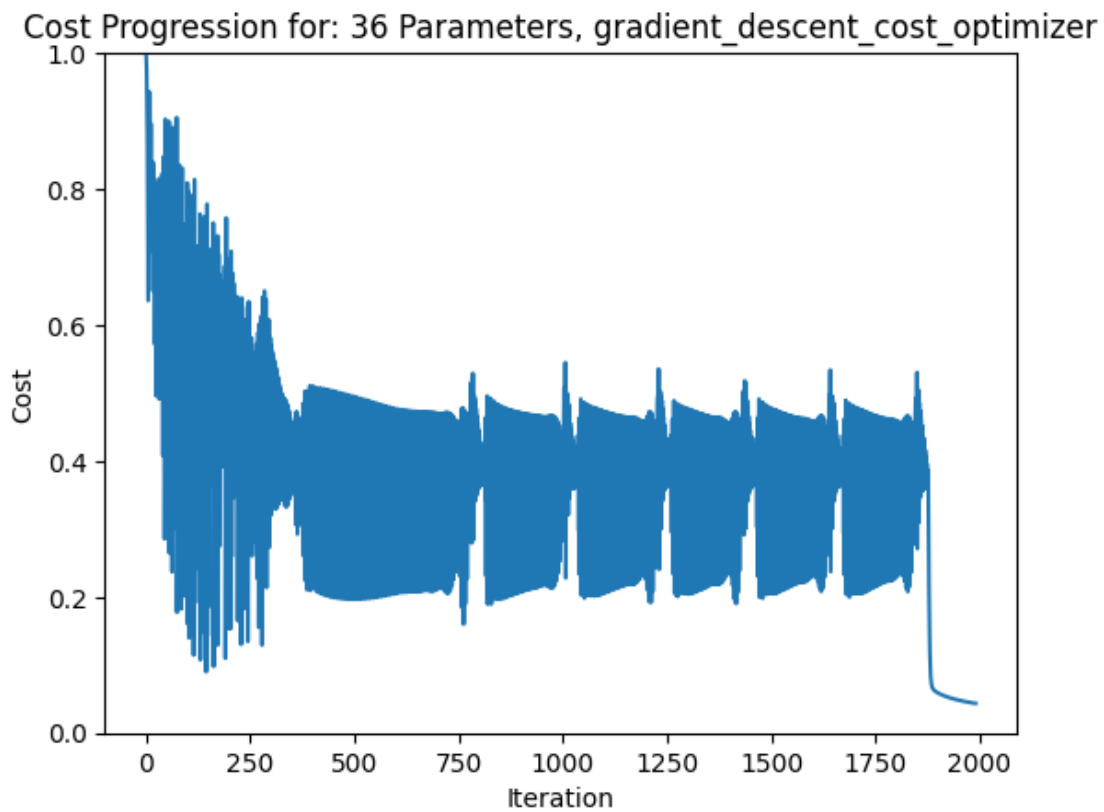


Figure 6.13: Cost progression plot, with 36 trainable parameters, using Gradient Descent optimizer (A)

X INITIAL is: tensor([4.8362, 3.6321, 3.6455, 5.0998, 3.9771, 3.4398, 2.6750, 1.2670, 4.5698, 0.5346, 4.9464, 5.9116, 5.6344, 2.2204, 1.3853, 2.7208, 4.4199, 2.4230,

0.8827, 5.6453, 5.6098, 6.0475, 5.0129, 5.9373, 3.6265, 1.8172, 0.2852, 3.4911, 4.7342, 2.6652, 0.2896, 3.0127, 0.6324, 3.9121, 5.5221, 2.0697])

initial cost: tensor(0.9984)

X FINAL is: tensor([5.0624, 3.4483, 3.9738, 4.6945, 4.5981, 2.7576, 2.7066, 1.1965, 4.7613, 0.6210, 4.8822, 5.8412, 5.8972, 2.4119, 1.2588, 2.7085, 4.6114, 2.3525, 1.0755, 5.5077, 5.8030, 5.8436, 4.6022, 5.8765, 3.9150, 1.7130, 0.5636, 3.0456, 4.8265, 2.0078, 0.1334, 2.5234, 0.5314, 3.8751, 4.9492, 2.6907])

final cost: tensor(0.0437)

learning rate = 0.05

delta = 0.005

epsilon = 1e-08

threshold = 0.0001

step size = 0.1

B. Again changing epsilon to 1e-06:

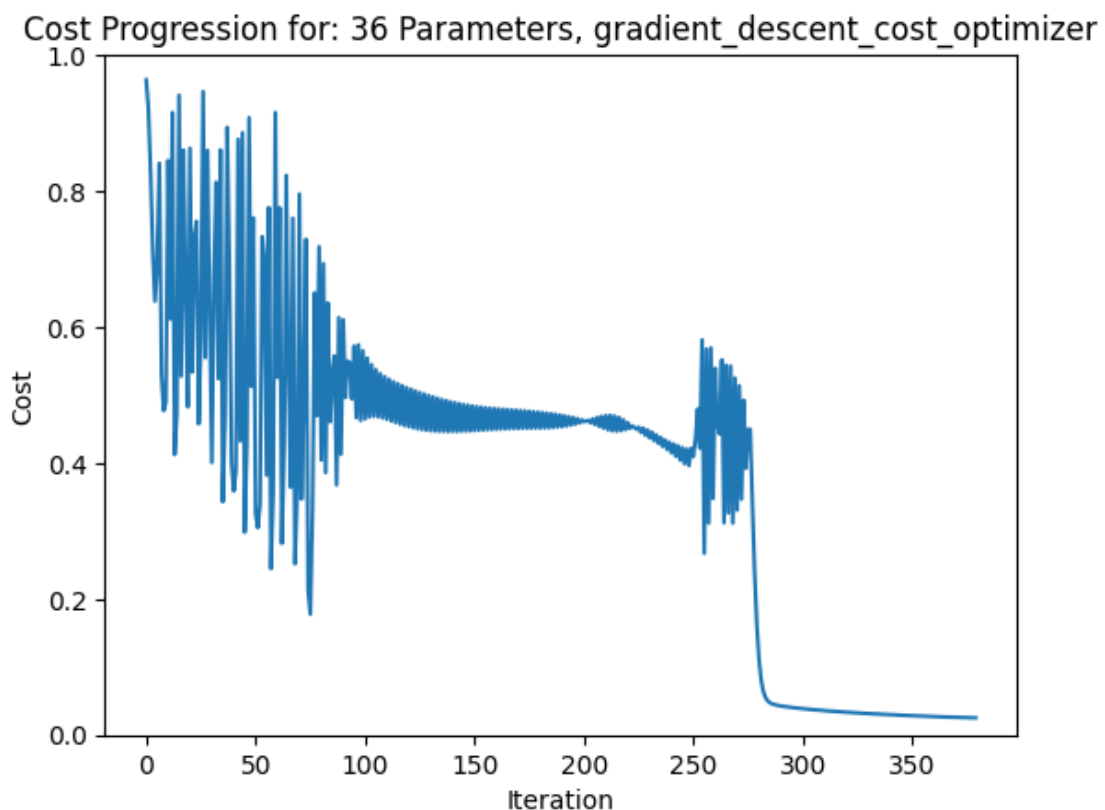


Figure 6.14: Cost progression plot, with 36 trainable parameters, using Gradient Descent optimizer (B)

X INITIAL is: tensor([5.9952, 5.5263, 0.6751, 6.0429, 2.1922, 0.3648, 5.7690, 6.0980, 3.3402, 1.8612, 2.8357, 2.1689, 5.5096, 4.7071, 5.0906, 4.2524, 2.3970, 0.8527,

3.8382, 4.1531, 3.7767, 1.5861, 5.9046, 4.4120, 0.5971, 2.0312, 1.8445, 5.6879, 4.9407, 4.7796, 5.3012, 4.8495, 5.8104, 0.8958, 3.2829, 6.1114])

initial cost: tensor(0.9840)

X FINAL is: tensor([5.9893, 5.3949, 0.6761, 5.9251, 2.5392, 0.2661, 5.6742, 6.2167, 3.2759, 1.5737, 2.7046, 2.2876, 5.5921, 4.6428, 4.4534, 4.2063, 2.3327, 0.9713, 3.6880, 4.0220, 3.6738, 1.6063, 5.5940, 4.3295, 0.9447, 1.7410, 2.2376, 5.3814, 4.9522, 4.8468, 5.0590, 4.5405, 6.0542, 0.7577, 3.0983, 6.4584])

final cost: tensor(0.0253)

learning rate = 0.05

delta = 0.005

epsilon = 1e-06

threshold = 0.0001

step size = 0.1

6.8 Execution time comparison

System specs

CPU: i7 9750h (6 cores - 12threads)

RAM: 16GB DDR4

For these results, different initialization values are used for each epsilon testing, for each parameter set.

Table 6.1: Optimization Results for different number, N of parameters and different Epsilon Values

# of parameters	Epsilon	Iterations	Final Cost	Execution time
18	1e-08	231	0.5464	12998.27 (3h 36m 38s)
	1e-06	455	0.5449	5266.01 (1h 27m 46s)
22	1e-08	202	0.5989	3767.74 (1h 2m 47s)
	1e-06	179	0.5770	3209.74 (53m 29s)
26	1e-08	220	0.0836	26892.91 (7h 28m 12s)
	1e-06	1211	0.0437	27805.79 (7h 43m 25s)
28	1e-08	1770	0.1581	45584.77 (12h 39m 45s)
	1e-06	527	0.0494	12425.25 (3h 27m 5s)
36	1e-08	1991	0.0437	84463.23 (23h 27m 43s)
	1e-06	379	0.0253	16445.67 (4h 34m 6s)

For the following results, each execution had the same initialization values.

Initial Values: [4.7603, 1.7550, 2.5326, 4.6162, 0.1840, 5.0257, 2.4953, 4.7399, 3.5783, 2.7569, 4.0129, 3.2966, 4.2890, 1.9173, 2.9125, 2.8588, 3.5969, 3.1290]
Initial Cost: 0.9753

Table 6.2: Optimization Results (Parameters: 18)

Epsilon	Iterations	Final Cost	Time Taken (s)	Time Taken (hh:mm:ss)
1e-08	203	0.5456	3164.09	0 h 52 min 44 s
1e-06	201	0.5456	2882.87	0 h 48 min 2 s

Initial Values: [5.8879, 4.1192, 1.9716, 1.2442, 2.6150, 1.7865, 2.1349, 3.2920, 5.0144, 4.8492, 0.0705, 5.0891, 4.0192, 6.1216, 5.2152, 0.2792, 0.1545, 1.6263, 5.9003, 2.6183, 4.4861, 1.6817] **Initial Cost:** 0.9807

Table 6.3: Optimization Results (Parameters: 22)

Epsilon	Iterations	Final Cost	Time Taken (s)	Time Taken (hh:mm:ss)
1e-08	158	0.5194	3269.40	0 h 54 min 29 s
1e-06	157	0.5194	3175.23	0 h 52 min 55 s

Initial Values: [6.2242, 1.8124, 5.4976, 3.1788, 1.4865, 4.7564, 1.4740, 4.0655, 2.2344, 2.7972, 0.1213, 1.6437, 4.8463, 2.3779, 6.2708, 5.6599, 2.9945, 1.0446, 5.0547, 4.1166, 1.1108, 5.1822, 5.0489, 5.9279, 1.3805, 2.6245]
Initial Cost: 0.9858

Table 6.4: Optimization Results (Parameters: 26)

Epsilon	Iterations	Final Cost	Time Taken (s)	Time Taken (hh:mm:ss)
1e-08	1377	0.1439	38592.14	10 h 43 min 12 s
1e-06	1374	0.1439	34051.83	9 h 27 min 31 s

Initial Values: [2.0370, 1.5601, 2.9670, 5.9147, 1.7438, 4.8593, 0.8470, 0.8450, 5.3775, 1.0312, 4.9395, 2.6428, 1.3595, 4.7149, 1.6253, 2.5752, 3.9885, 3.9920, 5.5750, 2.4976, 4.4143, 0.8474, 6.2534, 5.4128, 3.5603, 3.2367, 2.3206, 1.7052]
Initial Cost: 0.9539

Table 6.5: Optimization Results (Parameters: 28)

Epsilon	Iterations	Final Cost	Time Taken (s)	Time Taken (hh:mm:ss)
1e-08	526	0.0495	16444.75	4 h 34 min 4 s
1e-06	525	0.0495	16672.29	4 h 37 min 52 s

7. CONCLUSIONS AND FUTURE WORK

7.1 Conclusions

The results showed that increasing the number of trainable parameters in the quantum circuit generally improved the result of optimization process. Starting from 18 parameters, the cost decreased as more parameters were added, indicating that a larger parameter space allows for better optimization and lower cost values. Significant results, were achieved for the first time, on 26 parameters. Usually, more parameters produced better results. Another important thing to notice, is the significant impact of the initial parameter values, on the optimization process, affecting the convergence rate, the final cost achieved, and the overall performance of the optimization algorithm. An illustrative example is the results of the executions with 26 parameters. We observe costs as low as 0.0836 and 0.0437 (**Table 6.1**), to 0.1439 (**Table 6.4**), resulting from different initial values, which proves the impact they have on the optimization process. Different initialization values resulted in different convergence rates and final cost values, even though for more than 26 parameters, the cost was relatively successfully optimized. This is an interesting point since it implies that even though the number of local minima is increasing with the number of parameters, these minima are of equal depth/value. This effect has not been mentioned in other optimization tasks attacked with VQCs where usually barren plateaus appear.

The findings also reveal another interesting aspect of the optimization process: When dealing with a considerable number of parameters (in this case $N = 28$ and 36), and as the number of parameters increases, adjusting the epsilon value from $1e-08$ to $1e-06$ had a notable impact on both execution time and the number of iterations required for convergence, depending on the initialization values. In the examples of **Table 6.1**, the execution is about 4 times faster in the case of 28 parameters, and almost 6 times faster in the case of 36 parameters. On the other hand, when dealing with the example of **Table 6.5**, there is not a significant improvement on the overall performance of the algorithm. This indicates the point stated before. The importance of the initial values, cannot be overlooked. In conclusion, epsilon values less than $1e-06$, set a high "sensitivity" to the algorithm, which makes the convergence slower, and lower the performance, while not improving the final cost.

Adopting a greater epsilon value accelerated the optimization process significantly. By increasing epsilon, the optimizer could take more substantial steps during each iteration, expediting the convergence towards an optimal solution. As a result, fewer iterations were needed to reach a satisfactory cost value, leading to a reduction in the overall execution time of the optimization algorithm. To summarize, the epsilon value played a crucial role not only in influencing the final cost but also in significantly improving the optimization process's speed and resource efficiency. Fine-tuning this hyperparameter allowed for a more balanced trade-off between solution quality and computational requirements, making the overall optimization procedure more viable and scalable for real-world quantum computing

applications.

The cost progression plots provided a useful way to visualize the optimization process, showing how the cost evolved over iterations, and whether the algorithm was converging or getting stuck in local minima. For most configurations, the cost progression plots showed that the optimization process was converging towards a minimum cost value, indicating that the quantum circuit was being trained successfully.

The experiments suggest that to efficiently train this specific quantum circuit, 26 parameters were found to be sufficient. Adding more parameters beyond this threshold did not always lead to significant improvements in the final cost. Even though the best result was achieved with 36 trainable parameters, with the lowest cost value of 0.0253 using Gradient Descent optimizer (B) and an epsilon value of $1e-06$, the difference between the cost produced by 26 parameters (0.0437), or 28 parameters (0.094) (**Table 6.1**) is not significant, especially if optimal initial parameter values are chosen. A generic random unitary would require a VQC with at least **64** parameters to reach such a value of cost function and thus I can safely conclude that the algebraic ansatz which I have used has proven advantageous reducing the number of parameters and consequently the depth of quantum circuit in half. Notably, I performed an experiment with random generators outside from the set provided by the ansatz and the cost reached was larger by an order of magnitude (for $N = 26$ parameters).

In conclusion, the experiments showed that the optimization process of a quantum circuit heavily relies on the the number of parameters, their initial values and the hyperparameter tuning. It is essential to find the right balance between complexity and computational efficiency to achieve an optimal solution for the specific quantum circuit under consideration.

7.2 Future work

As a future work I would like to proceed with 4 and 5 qubit VQC and investigate how my observations keep up in these cases. Even though my VQC consists of a circuit of larger depth than the initial compilation of QFT (see Figure 6.4), one should take into account that the proposed VQC in this work can compile other operations as well with the same architecture and thus is more generic and powerful. In conclusion my results give some promise that this important operation in quantum computing could be realized by VQC obeying certain algebraic structure. In addition the training of such VQC is executable on a classical computer if the optimization methods and parameters withing are thoughtfully selected.

8. APPENDIX

8.1 Python implementation

This project is implemented in python, using mostly the **PyTorch** library.

Gradient descent, as well as stochastic gradient descent are implemented from scratch following the mathematical formula, since the optimizers included in **PyTorch** library, were not suitable for this type of model. The final circuit is a 8×8 matrix, that is imposible to be optimized by using simply the **torch.optim** package. The full code, implemented in python, is stated below in **Google Collab** notebook form. The stochastic gradient descent is implemented in the code bellow, but it is not used since it turned out to be ineffective for the needs of this project.

Quantum Fourier transform via variational quantum circuits

Steiropoulou Evangelia

1 Code implementation

```
[ ]: import torch
import numpy as np
import scipy.linalg
import sys
import matplotlib.pyplot as plt
import numpy as np
import time
```

1.1 Pauli Matrices:

1.1.1 Pauli-X matrix:

```
[ ]: ss1 = [[0,1],[1,0]]
ss1 = torch.tensor(ss1)
```

1.1.2 Pauli-Y matrix:

```
[ ]: ss2 = [[0,-1j],[1j,0]]
ss2 = torch.tensor(ss2)
```

1.1.3 Pauli-Z matrix:

```
[ ]: ss3 = [[1,0],[0,-1]]
ss3 = torch.tensor(ss3)
```

1.1.4 Identity matrix:

```
[ ]: ss4 = torch.eye(2)
```


1.2 QFT matrix, row by row:

```
[ ]: #fill a tensor 1*8 with 1/(2*sqrt(2))
QF3 = torch.full((1,8),1/(2*torch.sqrt(torch.tensor(2.0))))
QF3 = torch.tensor(QF3)

[ ]: #fill a tensor 1*8 with 1/(2*sqrt(2), exponential(1j*pi/4)/2*sqrt(2), (1j)/
    ↪ 2*sqrt(2), exponential(3j*pi/4)/2*sqrt(2), - 1/(2*sqrt(2), exponential(-3j*pi/
    ↪ 4)/2*sqrt(2), -(1j)/2*sqrt(2), exponential(-1j*pi/4)/2*sqrt(2)
QF4 = [[1/(2*torch.sqrt(torch.tensor(2.0))), (torch.exp(1j*torch.tensor(np.pi/
    ↪ 4)))/(2*torch.sqrt(torch.tensor(2.0))), 1j/(2*torch.sqrt(torch.tensor(2.0))), ↵
    ↪ torch.exp(3j*torch.tensor(np.pi/4))/(2*torch.sqrt(torch.tensor(2.0))), -1/
    ↪ (2*torch.sqrt(torch.tensor(2.0))), torch.exp(-3j*torch.tensor(np.pi/4))/
    ↪ (2*torch.sqrt(torch.tensor(2.0))), -1j/(2*torch.sqrt(torch.tensor(2.0))), ↵
    ↪ torch.exp(-1j*torch.tensor(np.pi/4))/(2*torch.sqrt(torch.tensor(2.0)))]
QF4 = torch.tensor(QF4)

[ ]: #fill a tensor 1*8, with 1/(2*sqrt(2), (1j)/2*sqrt(2), - 1/(2*sqrt(2), -(1j)/
    ↪ 2*sqrt(2), 1/(2*sqrt(2), (1j)/2*sqrt(2), - 1/(2*sqrt(2), -(1j)/2*sqrt(2)
QF5 = [[1/(2*torch.sqrt(torch.tensor(2.0))), 1j/(2*torch.sqrt(torch.tensor(2.
    ↪ 0))), -1/(2*torch.sqrt(torch.tensor(2.0))), -1j/(2*torch.sqrt(torch.tensor(2.
    ↪ 0))), 1/(2*torch.sqrt(torch.tensor(2.0))), 1j/(2*torch.sqrt(torch.tensor(2.
    ↪ 0))), -1/(2*torch.sqrt(torch.tensor(2.0))), -1j/(2*torch.sqrt(torch.tensor(2.
    ↪ 0)))]
QF5 = torch.tensor(QF5)

[ ]: #fill a tensor 1*8, with 1/(2*sqrt(2), exponential(3j*pi/4)/2*sqrt(2), -(1j)/
    ↪ 2*sqrt(2), exponential(1j*pi/4)/2*sqrt(2), - 1/(2*sqrt(2), exponential(-1j*pi/
    ↪ 4)/2*sqrt(2), (1j)/2*sqrt(2), exponential(-3j*pi/4)/2*sqrt(2)
QF6 = [[1/(2*torch.sqrt(torch.tensor(2.0))), torch.exp(3j*torch.tensor(np.pi/4))/
    ↪ (2*torch.sqrt(torch.tensor(2.0))), -1j/(2*torch.sqrt(torch.tensor(2.0))), ↵
    ↪ torch.exp(1j*torch.tensor(np.pi/4))/(2*torch.sqrt(torch.tensor(2.0))), -1/
    ↪ (2*torch.sqrt(torch.tensor(2.0))), torch.exp(-1j*torch.tensor(np.pi/4))/
    ↪ (2*torch.sqrt(torch.tensor(2.0))), 1j/(2*torch.sqrt(torch.tensor(2.0))), ↵
    ↪ torch.exp(-3j*torch.tensor(np.pi/4))/(2*torch.sqrt(torch.tensor(2.0)))]
QF6 = torch.tensor(QF6)

[ ]: #fill a tensor 1*8 with 1/(2*sqr(2) and -1/(2*sqr(2)
QF7 = [[1/(2*torch.sqrt(torch.tensor(2.0))), -1/(2*torch.sqrt(torch.tensor(2.
    ↪ 0))), 1/(2*torch.sqrt(torch.tensor(2.0))), -1/(2*torch.sqrt(torch.tensor(2.
    ↪ 0))), 1/(2*torch.sqrt(torch.tensor(2.0))), -1/(2*torch.sqrt(torch.tensor(2.
    ↪ 0))), 1/(2*torch.sqrt(torch.tensor(2.0))), -1/(2*torch.sqrt(torch.tensor(2.
    ↪ 0)))]
QF7 = torch.tensor(QF7)

[ ]:
```

```

QF8 = [[1/(2*torch.sqrt(torch.tensor(2.0))), torch.exp(-3j*torch.tensor(np.pi/
→4))/(2*torch.sqrt(torch.tensor(2.0))), 1j/(2*torch.sqrt(torch.tensor(2.0))),
→torch.exp(-1j*torch.tensor(np.pi/4))/(2*torch.sqrt(torch.tensor(2.0))), -1/
→(2*torch.sqrt(torch.tensor(2.0))), torch.exp(1j*torch.tensor(np.pi/4))/
→(2*torch.sqrt(torch.tensor(2.0))), -1j/(2*torch.sqrt(torch.tensor(2.0))),
→torch.exp(3j*torch.tensor(np.pi/4))/(2*torch.sqrt(torch.tensor(2.0)))]
QF8 = torch.tensor(QF8)

```

```

[ ]: #fill a tensor 1*8, with 1/(2*sqrt(2), (-1j)/2*sqrt(2), - 1/(2*sqrt(2), (1j)/
→2*sqrt(2), 1/(2*sqrt(2), (-1j)/2*sqrt(2), - 1/(2*sqrt(2), (1j)/2*sqrt(2)
QF9 = [[1/(2*torch.sqrt(torch.tensor(2.0))), -1j/(2*torch.sqrt(torch.tensor(2.
→0))), -1/(2*torch.sqrt(torch.tensor(2.0))), 1j/(2*torch.sqrt(torch.tensor(2.
→0))), 1/(2*torch.sqrt(torch.tensor(2.0))), -1j/(2*torch.sqrt(torch.tensor(2.
→0))), -1/(2*torch.sqrt(torch.tensor(2.0))), 1j/(2*torch.sqrt(torch.tensor(2.
→0)))]
QF9 = torch.tensor(QF9)

```

1.2.1 QFT matrix:

```

[ ]: #fill a tensor 1*8 with 1/(2*sqrt(2), exponential(-1j*pi/4)/2*sqrt(2), (-1j)/
→2*sqrt(2), exponential(-3j*pi/4)/2*sqrt(2), - 1/(2*sqrt(2), exponential(3j*pi/
→4)/2*sqrt(2), (1j)/2*sqrt(2), exponential(1j*pi/4)/2*sqrt(2)
QF10 = [[1/(2*torch.sqrt(torch.tensor(2.0))), torch.exp(-1j*torch.tensor(np.pi/
→4))/(2*torch.sqrt(torch.tensor(2.0))), -1j/(2*torch.sqrt(torch.tensor(2.0))),
→torch.exp(-3j*torch.tensor(np.pi/4))/(2*torch.sqrt(torch.tensor(2.0))), -1/
→(2*torch.sqrt(torch.tensor(2.0))), torch.exp(3j*torch.tensor(np.pi/4))/
→(2*torch.sqrt(torch.tensor(2.0))), 1j/(2*torch.sqrt(torch.tensor(2.0))),
→torch.exp(1j*torch.tensor(np.pi/4))/(2*torch.sqrt(torch.tensor(2.0)))]
QF10 = torch.tensor(QF10)

```

$$\frac{1}{\sqrt{8}} \begin{bmatrix} 1 & 1 & 1 & 1 & 1 & 1 & 1 & 1 \\ 1 & e^{i\frac{\pi}{4}} & e^{i\frac{\pi}{2}} & e^{i\frac{3\pi}{4}} & e^{i\pi} & e^{i\frac{5\pi}{4}} & e^{i\frac{3\pi}{2}} & e^{i\frac{7\pi}{4}} \\ 1 & e^{i\frac{\pi}{2}} & e^{i\pi} & e^{i\frac{3\pi}{2}} & 1 & e^{i\frac{\pi}{2}} & e^{i\pi} & e^{i\frac{3\pi}{2}} \\ 1 & e^{i\frac{3\pi}{4}} & e^{i\frac{3\pi}{2}} & e^{i\frac{9\pi}{4}} & e^{i\pi} & e^{i\frac{5\pi}{4}} & e^{i\frac{7\pi}{2}} & e^{i\frac{15\pi}{4}} \\ 1 & e^{i\pi} & 1 & e^{i\pi} & 1 & e^{i\pi} & 1 & e^{i\pi} \\ 1 & e^{i\frac{5\pi}{4}} & e^{i\frac{\pi}{2}} & e^{i\frac{7\pi}{4}} & e^{i\pi} & e^{i\frac{\pi}{4}} & e^{i\frac{\pi}{2}} & e^{i\frac{3\pi}{4}} \\ 1 & e^{i\frac{3\pi}{2}} & e^{i\pi} & e^{i\frac{7\pi}{2}} & 1 & e^{i\frac{\pi}{2}} & e^{i\pi} & e^{i\frac{3\pi}{2}} \\ 1 & e^{i\frac{7\pi}{4}} & e^{i\frac{3\pi}{2}} & e^{i\frac{15\pi}{4}} & e^{i\pi} & e^{i\frac{3\pi}{4}} & e^{i\frac{7\pi}{2}} & e^{i\frac{15\pi}{4}} \end{bmatrix}$$

```

[ ]: #make tensor with all the above tensors in it
QF = torch.cat((QF3, QF4, QF5, QF6, QF7, QF8, QF9, QF10), 0)

```

1.3 Generators:

Here we create the generators, the quantum gates that are going to be used in the circuit. The generators, are combinations of Kronecker products of the gates we mentioned above.

1.3.1 Single qubit gates:

(1,4,4)

```
[ ]: c1 = torch.kron(ss1, ss4)
     c1 = torch.kron(c1, ss4)
```

(4,2,4)

```
[ ]: c2 = torch.kron(ss4, ss2)
     c2 = torch.kron(c2, ss4)
```

(4,3,4)

```
[ ]: c3 = torch.kron(ss4, ss3)
     c3 = torch.kron(c3, ss4)
```

(4,1,4)

```
[ ]: c4 = torch.kron(ss4, ss1)
     c4 = torch.kron(c4, ss4)
```

(4,4,3)

```
[ ]: c5 = torch.kron(ss4, ss4)
     c5 = torch.kron(c5, ss3)
```

1.3.2 Two - qubit gates:

(4,3,3)

```
[ ]: c6 = torch.kron(ss4, ss3)
     c6 = torch.kron(c6, ss3)
```

(4,1,3)

```
[ ]: c7 = torch.kron(ss4, ss1)
     c7 = torch.kron(c7, ss3)
```

(1,1,4)

```
[ ]: c8 = torch.kron(ss1, ss1)
     c8 = torch.kron(c8, ss4)
```

(1,2,4)

```
[ ]: c9 = torch.kron(ss1, ss2)
     c9 = torch.kron(c9, ss4)
```

(3,4,2)

```
[ ]: c10 = torch.kron(ss3, ss4)
     c10 = torch.kron(c10, ss2)
```

(1,4,3)

```
[ ]: c11 = torch.kron(ss1, ss4)
      c11 = torch.kron(c11, ss3)
```

In vv3 we will add all the generators, for later use.

```
[ ]: #c1 - c5 are single qubit gates, c6 - c11 are two qubit gates
      vv3 = torch.stack((c1, c2, c3, c4, c5, c6, c7, c8, c9, c10, c11))
```

1.4 Variational Gates:

```
[ ]: #Gi_, j_, k_, x_ := MatrixExp[I x KroneckerProduct[Assi, ssj, ssk]]
      G = torch.zeros(11, 8, 8, dtype=torch.complex64)
      for i in range(G.size(dim = 0)):
          G[i] = torch.linalg.matrix_exp(1j*vv3[i])
```

```
[ ]: def Gx(x):
      Gx = torch.zeros(11, 8, 8, dtype=torch.complex64)
      for i in range(Gx.size(dim = 0)):
          Gx[i] = torch.tensor(scipy.linalg.fractional_matrix_power(G[i], x)) #Gi
      ↪ to the power of x
      return Gx
```

```
[ ]: #find conjugate transpose of QF
      B = torch.conj(torch.transpose(QF, 0,1))
      B.requires_grad = True
```

1.5 Circuit generator:

The circuit created below, was firstly designed by hand, in order to combine both single and 2-qubit variational gates.

```
[ ]: def create_circuit(x_var, parameters_num):
      Gm = []
      # loop over the x values to generate the corresponding G matrices
      for i in range(x_var.size(dim=0)):
          Gx_i = torch.zeros(11, 8, 8, dtype=torch.complex64)
          Gx_i = Gx(x_var[i].item())
          Gm.append(Gx_i)

      # multiply the 18 G matrices to get the final G matrix/circuit
      i = 0

      G1 = Gm[i][5] #get the first 2-qubit gate(of the first x-modified
      ↪ Gx_i(i==0)), 4-3-3
      G2 = Gm[i+1][1] #get the second single qubit gate, 4-2-4
      G3 = Gm[i+2][4] #get the last single qubit gate 4-4-3
      G4 = Gm[i+3][7] #1-1-4
      G5 = Gm[i+4][0] #1-4-4
```

```

G6 = Gm[i+5][2] #4-3-4
G7 = Gm[i+6][9] #3-4-2
G8 = Gm[i+7][4] #4-4-3
G9 = Gm[i+8][0] #1-4-4
G10 = Gm[i+9][6] #4-1-3
G11 = Gm[i+10][2] #4-3-4
G12 = Gm[i+11][4] #4-4-3
G13 = Gm[i+12][8] #1-2-4
G14 = Gm[i+13][0] #1-4-4
G15 = Gm[i+14][3] #4-1-4
G16 = Gm[i+15][10] #1-4-3
G17 = Gm[i+16][0] #1-4-4
G18 = Gm[i+17][4] #4-4-3

G_final = G1@G2@G3@G4@G5@G6@G7@G8@G9@G10@G11@G12@G13@G14@G15@G16@G17@G18

#In order to expand the initial 18-parameter circuit to a n-parameter
↪ circuit, we need to add more gates at the end of the circuit.
#As additional gates we will use those in the initial 18-parameter circuit.
↪ For example if the desired circuit has 20 parameters,
#we need to add 2 additional gates to the initial 18-parameter circuit. In
↪ order to do that, we will add the first 2 gates of the initial circuit
#at the end of the circuit. If we want a 28-parameter circuit, we will add
↪ the first 10 gates of the initial circuit, at the end etc.

#Initial gate indices
gate_indices = [5, 1, 4, 7, 0, 2, 9, 4, 0, 6, 2, 4, 8, 0, 3, 10, 0, 4] #
↪ Example gate indices
# Additional gates
G_additional = torch.eye(8, dtype=torch.complex64) # identity matrix

# Multiply additional gates based on the number of parameters
for i in range(parameters_num - 18):
    gate_idx = gate_indices[i % 18] # Cycle through the gate_indices list
    G_additional = G_additional @ Gm[i+18][gate_idx]
    G_final = G_final @ G_additional # Multiply G_final with G_additional

return G_final

```

1.6 Cost function:

```

[ ]: def cost_function(x_var):
    G_final = create_circuit(x_var, len(x_var))
    cost = 1 - 1/64 * ((torch.abs(torch.trace(G_final @ B)))**2)
    return cost

```

1.7 Optimization methods:

```
[ ]: def learning_rate_step_scheduler(learning_rate, step_size):  
    return learning_rate * step_size
```

1.7.1 Gradient Descent optimizer:

```
[ ]: #function performs gradient descent of cost to find the optimal x values  
#x_var is the initial x values, gamma is the learning rate, delta is the  
↳perturbation value  
def optimize_parameters(x_var, gamma, delta):  
    #print("x initial is: \n\n", x_var)  
    #print("cost initial = ", cost_function(x_var))  
    x_new = x_var.clone()  
  
    for i in range(len(x_var)):  
        x_var_sum = x_var.clone() #create a copy of the x_var tensor  
        x_var_sum[i] = x_var[i] + delta  
        cost_sum = cost_function(x_var_sum)  
  
        x_var_diff = x_var.clone()  
        x_var_diff[i] = x_var[i] - delta  
        cost_diff = cost_function(x_var_diff)  
        x_new[i] = x_var[i] - gamma * ((cost_sum - cost_diff) / (2* delta))  
  
    return x_new, cost_function(x_new)  
  
[ ]: #function that calls the optimize_parameters function until the cost stops  
↳changing more than a certain value(epsilon)  
def gradient_descent_cost_optimizer(x_var, learning_rate, delta, epsilon,  
    ↳threshold, step_size):  
    iterations = 0  
    x_init, cost_init = optimize_parameters(x_var, learning_rate, delta) #get  
↳the initial cost after the first optimization  
    x_old = x_init.clone()  
    cost_old = cost_init.clone()  
    cost_history = [cost_init] # List to store the cost at each iteration  
  
    while True:  
        #print("ITERATION = \n", iterations)  
        x_new, cost_new = optimize_parameters(x_old, learning_rate, delta)  
        #print("new cost = ", cost_new)  
        if torch.abs(cost_new - cost_old) < epsilon:  
            break  
        else:  
            if(torch.abs(cost_new - cost_old) < threshold and iterations != 0):
```

```

        learning_rate = learning_rate_step_scheduler(learning_rate,
↪step_size)
        #print("ITERATION = ", iterations, "    LEARNING RATE = ",
↪learning_rate, "\n")
        x_old = x_new.clone()
        cost_old = cost_new.clone()
        iterations += 1
        cost_history.append(cost_new) # Add the current cost to the history

    return x_new, cost_new, iterations, cost_history

```

1.7.2 Stochastic gradient descent:

```

[ ]: # Perform optimization on a single data point -> this will be used in the
↪stochastic gradient descent
def optimize_stochastic_parameters(x_var, learning_rate, delta, data_point): #
↪Perform optimization on a single data point
    x_var_sum = x_var.clone()
    x_var_sum[data_point] = x_var[data_point] + delta
    cost_sum = cost_function(x_var_sum)

    x_var_diff = x_var.clone()
    x_var_diff[data_point] = x_var[data_point] - delta
    cost_diff = cost_function(x_var_diff)

    x_new = x_var.clone()
    x_new[data_point] = x_var[data_point] - learning_rate* ((cost_sum -
↪cost_diff) / (2 * delta))

    return x_new, cost_function(x_new)

```

```

[ ]: def stochastic_gradient_descent(x_var, learning_rate, delta, epsilon, threshold,
↪step_size, scheduler, num_epochs):
    iterations = 0
    num_data_points = len(x_var)
    x_init, cost_init = optimize_stochastic_parameters(x_var, learning_rate,
↪delta, np.random.randint(num_data_points))
    x_old = x_init.clone()
    cost_old = cost_init.clone()
    cost_difference = torch.abs(cost_old - cost_init)
    cost_history = [cost_init] # List to store the cost at each iteration

    while True:
        #print("ITERATION =", iterations)
        data_point = np.random.randint(num_data_points)

```

```

        x_new, cost_new = optimize_stochastic_parameters(x_old, learning_rate,
↪delta, data_point)
        #print("x new =", x_new)
        #print("new cost =", cost_new)

        if torch.abs(cost_new - cost_old) != cost_difference:
            cost_difference = torch.abs(cost_new - cost_old)
            #print("cost difference =", cost_difference)
        if iterations > num_epochs and cost_difference < epsilon:
            break
        else:
            if cost_difference < threshold and iterations != 0:
                #print("Scheduler called", scheduler)
                learning_rate = scheduler(learning_rate, step_size)
                #print("ITERATION =", iterations, " LEARNING RATE =",
↪learning_rate, "\n")

            x_old = x_new.clone()
            cost_old = cost_new.clone()
            iterations += 1
            cost_history.append(cost_new) # Add the current cost to the history

    return x_new, cost_new, iterations, cost_history

```

1.8 Execution of the program, and cost optimization

```

[297]: num_parameters = [18, 22, 26, 28]
iterations = [100, 150, 200, 300] # Number of iterations for each num of
↪parameters. Used only in stochastic gradient descent.
#In this implementation, we perform only the gradient descent algorithm

counter = 1 # Initialize the counter

# Create an empty list to store the results
results = []
for algorithm in [gradient_descent_cost_optimizer]:
    for i, j in zip(num_parameters, iterations):

        # Create an empty list to store the results
        results_algorithm = []

        # Count time for each iteration
        start = time.time()
        x_var = torch.rand(i, dtype=torch.float32) * 2 * np.pi
        print("Algorithm is: ", algorithm.__name__, "\n")
        print("Number of parameters is: ", i, "\n")
        print("x initial is: \n", x_var, "\n")

```



```

if algorithm == stochastic_gradient_descent:
    learning_rate = 0.05
    delta = 0.0005
    epsilon = 1e-08
    threshold = 0.00001
    step_size = 0.1
    x, cost, iters, cost_history = stochastic_gradient_descent(x_var,
↪learning_rate, delta, epsilon, threshold, step_size,
↪learning_rate_step_scheduler, j)
else:
    learning_rate = 0.05
    delta = 0.005
    epsilon = 1e-08
    threshold = 0.0001
    step_size = 0.1
    x, cost, iters, cost_history =
↪gradient_descent_cost_optimizer(x_var, learning_rate, delta, epsilon,
↪threshold, step_size)

results_algorithm.append((x_var, cost_function(x_var), x, iters, cost))
end = time.time()

print("Parameters are: \n" , i, " X INITIAL is:\n", x_var)
print("initial cost: ", cost_function(x_var))
print("X FINAL is:\n\n", x)
print("iterations =", iters, "final cost: ", cost)
print("time taken =", end - start, "\n\n")
print("learning_rate = ", learning_rate, "\n")
print("delta = ", delta, "\n")
print("epsilon = ", epsilon, "\n")
print("threshold = ", threshold, "\n")
print("step_size = ", step_size, "\n")

cost_history_np = np.array([cost.detach().numpy() for cost in
↪cost_history])

# Plot the cost progression
plt.figure()
plt.plot(cost_history_np)
plt.xlabel('Iteration')
plt.ylabel('Cost')
plt.title(f'Cost Progression for: {i} Parameters, {algorithm.__name__}')

plt.ylim(bottom=0.0, top=1)

# Show the plot without blocking program execution

```

```

plt.show(block=False)

# Save the figure as a PNG file
filename = f'cost_progression_{i}_{algorithm.__name__}_{counter}.png'
plt.savefig(filename)

# Increment the counter
counter += 1

# Append the results to the list
for result in results_algorithm:
    results.append({
        'Algorithm': algorithm.__name__,
        'Number of Parameters': i,
        'Initial Values': result[0].detach().numpy(),
        'Final Values': result[2].detach().numpy(),
        'Iterations': result[3],
        'Initial Cost': result[1].item(),
        'Final Cost': result[4].item(),
        'Execution Time': end - start
    })

# Open the file in write mode
output_file = f'{i}_{algorithm.__name__}_{counter}.txt'
with open(output_file, 'w') as file:
    # Iterate over the 'results' list and write each element to the
    ↪file
    for result in results:
        file.write("Algorithm: {}\n".format(result['Algorithm']))
        file.write("Number of Parameters: {}\n".
        ↪format(result['Number of Parameters']))
        file.write("Initial Values: {}\n".format(result['Initial_
        ↪Values']))
        file.write("Final Values: {}\n".format(result['Final_
        ↪Values']))
        file.write("Initial Cost: {}\n".format(result['Initial_
        ↪Cost']))
        file.write("Final Cost: {}\n".format(result['Final Cost']))
        file.write("Iterations: {}\n".format(result['Iterations']))
        file.write("Execution Time: {}\n".format(result['Execution_
        ↪Time']))

```

ABBREVIATIONS - ACRONYMS

CNOT controlled-NOT. 25–27, 31

GHZ Greenberger-Horne-Zeilinger. 31

QFT Quantum Fourier Transform. 21, 33, 34, 39

qubit quantum bit. 9, 11, 21, 23–29, 31, 33, 34, 39

RSA Rivest–Shamir–Adleman. 21, 33

VQA Variational Quantum Algorithms. 21, 38

VQC Variational Quantum Circuits. 21, 61

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