Data re-uploading for a universal quantum classifier

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A single qubit provides sufficient computational capabilities to construct a universal quantum classifier. This fact may be surprising since a single qubit only offers a simple superposition of two states and single qubit gates only make a rotation in the Bloch sphere. The key ingredient to circumvent these limitations is to allow for multiple data re-uploading. A quantum circuit can then be organized as a series of data re-uploading and single-qubit processing units. Furthermore, both data re-uploading and measurements can accommodate multiple dimensions in the input and multiple categories in the output, so as to conform a universal quantum classifier. The extension of this idea to several qubits enhances the efficiency of the strategy as entanglement expands the superpositions carried along the classification. Extensive benchmarking on different examples of the single- and multi-qubit quantum classifier validates its ability to describe and classify complex data.

I. INTRODUCTION

Quantum circuits that make use of a small amount of quantum resources are of most importance to the field of quantum computation. Indeed, algorithms that need few qubits may prove relevant even if they do not attempt any quantum advantage, as they may be useful parts of larger circuits.

A reasonable question to ask is what is the lower limit of quantum resources needed to achieve a given computation. A naive estimation for the quantum cost of a new proposed quantum algorithm is often made based on analogies with classical algorithms. But this may be misleading, as classical computation can play with memory in a rather different way as quantum computers do. The question then turns to the more refined problem of establishing the absolute minimum of quantum resources for a problem to be solved.

We shall here explore the power and minimal needs of quantum circuits to carry out a general supervised classification task. Three elements in the computation need renewed attention. The obvious first concern is to find the way to upload data in a quantum computer. Then, it is necessary to find an optimal processing of information, followed by an optimal measurement strategy. We shall revisit these three issues in turn. The non-trivial step we take here is to combine the first two, that is data uploading and processing, in a non-trivial way.

There exist several proposals to construct a quantum classifier. These can be divided into two families: those that are inspired in classical classification algorithms – i.e. support vector machines and perceptrons [1,7] – and those that use parametrized quantum circuits [8,10]. The latter are promising candidates to be implemented in a Noisy Intermediate Scale Quantum computer. In these proposals, the quantum circuit is designed in a way that will be hardly simulated by a classical computer with the goal to achieve quantum advantage once applied to a large amount of qubits. The quantum classifiers introduced in those papers are designed to perform binary classification or, equivalently, to represent Boolean functions. The desired non-linearities of the functions are introduced by using the properties of tensor product. In contrast, we focus on the smaller possible circuit that is capable of performing a classification task consisting on two or more classes. The data non-linearities are achieved by re-uploading the data points along the circuit.

The main result of this work is, indeed, to show that there is a trade-off between the number of qubits needed to perform a classification and multiple data re-uploading. That is, we may use less qubits at the price of re-entering data several times along the quantum computation. Quantum computers can not copy data, but classical devices can. Therefore, it makes sense to re-upload classical data along a quantum computation to bypass the limitations established by the no-cloning theorem on the quantum circuit. By following this line of thought, we present an equivalence between data re-uploading and the Universal Approximation Theorem applied to artificial neural networks [11]. Just as a network composed of a single hidden layer with enough neurons can reproduce any continuous function, a single-qubit classifier can in principle achieve the same by re-uploading the data enough times.

The single-qubit classifier illustrates the computational power that a single qubit can handle. This proposal is to be added to other few-qubit benchmarks in machine learning [12]. The input redundancy has also been proposed to construct complex encoding in parametrized quantum circuits [13]. These and other proposals mentioned in the previous paragraphs are focused in representing classically intractable or very complex Kernel functions with few qubits. On the contrary, the focus of this work is to distill the minimal amount of quantum resources needed for a given classification task quantified in terms of the number of qubits and unitary operations.
We shall illustrate the power of a single- and multi-qubit classifiers with data re-uploading with a series of examples. First, we classify points in a plane which is divided in two areas. Then, we extend successfully the number of regions on a plane to be classified. Next, we consider the classification of multi-dimensional patterns and, finally, we benchmark this quantum classifier with non-convex figures. For every example, we train a parametrized quantum circuit that carries out the task and we analyze its performance in terms of the circuit architecture, i.e. for single- and multi-qubit classifiers with and without entanglement between qubits.

This paper is structured as follows. First, in section II we present the basic structure of a single-qubit quantum classifier. Data and processing parameters are uploaded and re-uploaded using one-qubit general rotations. For each data point, the final state of the circuit is compared with the target state assigned to its class and the free parameters of the circuit are updated accordingly using a classical minimization algorithm. Second, in section III we motivate the data re-uploading approach by using the Universal Approximation Theorem of artificial neural networks. In section IV we introduce the extension of this classifier to multiple qubits. Then, in section V we detail the minimization methods used to train the quantum classifiers. Finally, in section VI we benchmark single- and multi-qubit quantum classifiers defined previously with problems of different dimensions and complexity and present a comparison with respect to classical classification techniques. The conclusions of this proposal for a quantum classifier are exposed in section VII.

II. STRUCTURE OF A SINGLE-QUBIT QUANTUM CLASSIFIER

The global structure of any quantum circuit can be divided into three elements: uploading of information onto a quantum state, processing of the quantum state, and measurement of the final state. It is far from obvious how to implement each of these elements in an optimal way to perform a specific operation. We shall now address them one at a time for the task of classification.

A. Re-uploading classical information

To load classical information onto a quantum circuit is a highly non-trivial task [1]. A critical example is the processing of big data. While there is no in-principle obstruction to upload large amounts of data onto a state, it is not obvious how to do it.

The problem we address here is not related to a large amount of data. It is thus possible to consider a quantum circuit where all data are loaded in the coefficients of the initial wave function $|\psi\rangle$. In the simplest of cases, data are uploaded as rotations of qubits in the computational basis. A quantum circuit would then follow that should perform some classification.

This strategy would be insufficient to create a universal quantum classifier with a single qubit. A first limitation is that a single qubit only has two degrees of freedom, thus only allowing to represent data in a two-dimensional space. No quantum classifier in higher dimensions can be created if this architecture is to be used. A second limitation is that, once data is uploaded, the only quantum circuit available is a rotation in the Bloch sphere. It is easy to prove that a single rotation cannot capture any non-trivial separation of patterns in the original data.

We need to turn to a different strategy, which turns out to be inspired by neural networks. In the case of feed-forward neural networks, data are entered in a network in such a way that they are processed by subsequent layers of neurons. The key idea is to observe that the original data are processed several times, one for each neuron in the first hidden layer. Strictly speaking, data are re-uploaded onto the neural network. If neural networks were affected by some sort of no-cloning theorem, they could not work as they do. Coming back to the quantum circuit, we need to design a new architecture where data can be introduced several times into the circuit.

The central idea to build a universal quantum classifier with a single qubit is thus to re-upload classical data along with the computation. Following the comparison with an artificial neural network with a single hidden layer, we can represent this re-upload diagramatically as shows Fig. 1. Data points in a neural network are introduced in each processing unit, represented with squares, which are the neurons of the hidden layer. After the neurons process these data, a final neuron is necessary to construct the output to be analyzed. Similarly, in the single-qubit quantum classifier, data points are introduced in each processing unit which this time correspond with a unitary rotation. However, each processing unit is affected by the previous one and the input data re-introduced. The final output is a quantum state to be analyzed as it will be explained in the next subsections.

The explicit form of this single-qubit classifier is shown in Fig. 2. Classical data are re-introduced several times in a sequence interspaced with processing units. We shall consider the introduction of data as a rotation of the qubit. This means that data from a three-dimensional space, $\mathbf{x}$, can be re-uploaded using unitaries that rotate the qubit $U(\mathbf{x})$. Later processing units will also be rotations as discussed later on. The whole structure needs to be trained to the classification of patterns.

As we shall see, the performance of the single-qubit quantum classifier will depend on the number of re-uploads of classical data. This fact will be explored in the results section.
B. Processing along re-uploading

The single-qubit classifier belongs to the category of parametrized quantum circuits. The performance of the circuit is quantified by a figure of merit, some specific $\chi^2$ to be minimized and defined later. We need, though, to specify the processing gates present in the circuit in terms of a classical set of parameters.

Given the simple structure of a single-qubit circuit presented in Fig. 2, the data is introduced in a simple rotation of the qubit, which is easy to characterize. We just need to use arbitrary single-qubit rotations $U(\phi_1, \phi_2, \phi_3) \in SU(2)$. By an abuse of language, we will write $U(\hat{\phi})$ with $\hat{\phi} = (\phi_1, \phi_2, \phi_3)$. Then, the structure of the universal quantum classifier made with a single qubit is

$$U(\hat{\phi}, \vec{x}) \equiv U(\hat{\phi}_N)U(\vec{x}) \ldots U(\hat{\phi}_1)U(\vec{x}),$$

which acts as

$$|\psi\rangle = U(\hat{\phi}, \vec{x})|0\rangle.$$  

The final classification of patterns will come from the results of measurements on $|\psi\rangle$. We may introduce the concept of processing layer as the combination

$$L(i) \equiv U(\hat{\phi}_i)U(\vec{x}),$$

so that a more complex and more robust classifier corresponds to

$$U(\hat{\phi}, \vec{x}) = L(N) \ldots L(1),$$

where the depth of the circuit is $2N$. The more layers the more representation capabilities the circuit will have, and the more powerful the classifier will become. Again, this follows from the analogy to neural networks, where the size of the intermediate hidden layer of neurons is key to represent complex functions.

There is a way to compactify the quantum circuit into a shorter one. This can be done if we incorporate data and processing angles in a single step. Then, a layer would only need a single rotation to introduce data and tunable parameters, i.e. $L(i) = U(\hat{\phi}, \vec{x})$. In addition, each data point can be uploaded with some weight $w_i$. These weights will play a similar role as weights in artificial neural networks, as we will see in the next section. Altogether, each layer gate can be taken as

$$L(i) = U \left( \hat{\theta}_i + w_i \circ \vec{x} \right),$$

where $w_i \circ \vec{x} = (w_{i1} x^1, w_{i2} x^2, w_{i3} x^3)$ is the Hadamard product of two vectors. In case the data points have dimension lesser than three, the rest of $\vec{x}$ components are set to zero. Such an approach reduces the depth of the circuit by half. Further combinations of layers into a fewer rotations would be possible, but the nonlinearity inherent to subsequent rotations would be lost, and the circuit would not be performing well.

It is also possible to enlarge the dimensionality of the input space in the following way. Let us extend the definition of $i$-th layer to

$$L(i) = U \left( \hat{\theta}^{(k)}_i + w^{(k)}_i \circ \vec{x}^{(k)} \right) \ldots U \left( \hat{\theta}^{(1)}_i + w^{(1)}_i \circ \vec{x}^{(1)} \right),$$

where each data point is divided into $k$ vectors of dimension three. In general, each unitary $U$ could absorb as many variables as freedom in a $SU(2)$ unitary. Each set of variables act at a time, and all of them have been shown to the circuit after $k$ iterations. Then, the layer structure follows. The complexity of the circuit only increases linearly with the size of the input space.
C. Measurement

The quantum circuit characterized by a series of processing angles $\{\theta_i\}$ and weights $\{w_i\}$ delivers a final state $|\psi\rangle$, which needs to be measured. The results of each measurement are used to compute a $\chi^2$ that quantifies the error made in the classification. The minimization of this quantity in terms of the classical parameters of the circuit can be organized using any preferred supervised machine learning technique.

The critical point in the quantum measurement is to find an optimal way to associate outputs from the observations to target classes. The fundamental guiding principle to be used is given by the idea of maximal orthogonality of outputs [13]. This is easily established for a dichotomic classification, where one of two classes $A$ and $B$ have to be assigned to the final measurement of the single qubit. In such a case it is possible to measure the output probabilities $P(0)$ for $|0\rangle$ and $P(1)$ for $|1\rangle$. A given pattern could be classified into the A class if $P(0) > P(1)$ and into B otherwise. We may refine this criterion by introducing a bias. That is, the pattern is classified as A if $P(0) > \lambda$, and as B otherwise. The $\lambda$ is chosen so as to optimize the success of classification on a training set. Results are then checked on an independent validation set.

The assignment of classes to the output reading of a single qubit becomes an involved issue when many classes are present. For the sake of simplicity let us mention two examples for the case of classification to four distinct classes. One possible strategy consists on comparing the probability $P(0)$ to four sectors with three thresholds: $0 \leq \lambda_1 \leq \lambda_2 \leq \lambda_3 \leq 1$. Then, the value of $P(0)$ will fall into one of them and a classification is issued. A second, more robust assignment is obtained by computing the overlap of the final state to one of the states of a label states-set. This states-set is to be chosen with maximal orthogonality among all of them. This second method needs from the maximally orthogonal points in the Bloch sphere. Figure 3 shows the particular cases that can be applied to a classification task of four and six classes. In general, a good measurement strategy may need some prior computational effort and refined tomography of the final state. Since we are proposing a single-qubit classifier, the tomography protocol will only require three measurements.

It is possible to make an interpretation of the single-qubit classifier in terms of geometry. The classifier opens a 2-dimensional Hilbert space, i.e. the Bloch sphere. As we code data and classification within the parameters defining rotations, this Hilbert space is enough to achieve classification. Any operation $L(i)$ is a rotation on the surface of the Bloch sphere. With this point of view in mind, we can easily see that we can classify any point using only one unitary operation. We can transport any point to any other point on the Bloch sphere by nothing else than choosing the angles of rotation properly. However, this does not work for several data, as the optimal rotation for some data could be very inconvenient for some other.

1. A fidelity cost function

We propose a very simple cost function motivated by the geometrical interpretation introduced above. We want to force the quantum states $|\psi(\vec{\theta}, \vec{w}, \vec{x})\rangle$ to be as near as possible to one particular state on the Bloch sphere. The geometrical distance between the label state and the data state can be measured with the relative fidelity between the two states [15]. Thus, our aim is to maximize the average fidelity between the states at the end of the quantum circuit and the label states corresponding to their class. We define the following cost function that carries out this task,

$$
\chi^2_f(\vec{\theta}, \vec{w}) = \sum_{\mu=1}^{M} \left( 1 - |\langle \psi_{\mu} | \psi(\vec{\theta}, \vec{w}, x_{\mu}) \rangle|^2 \right),
$$

where $|\psi_{\mu}\rangle$ is the correct label state of the $\mu$ data point, which will correspond to one of the classes.

2. A weighted fidelity cost function

We shall next define a refined version of the previous fidelity cost function to be minimized. The set of maximally orthogonal states in the Bloch sphere, i.e. the label states, are written as $|\psi_c\rangle$, where $c$ is the class. Each of these label states represents one class for the classifier. Now, we will follow the lead usually taken in neural network classification.

Let us define the quantity

$$
F_c(\vec{\theta}, \vec{w}, \vec{x}) = |\langle \psi_c | \psi(\vec{\theta}, \vec{w}, \vec{x}) \rangle|^2,
$$

where $|\psi_c\rangle$ is the label state of the class $c$ and $|\psi(\vec{\theta}, \vec{w}, \vec{x})\rangle$ is the final state of the qubit at the end of the circuit.
This fidelity is to be compared with the expected fidelity of a successful classification, $Y_c(\vec{x})$. For example, given a four-class classification and using the vertices of a tetrahedron as label states (as shown in Fig. 3), one expects $Y_s(\vec{x}) = 1$, where $s$ is the correct class, and $Y_r(\vec{x}) = 1/3$ for the other $r$ classes. In general, $Y_c(\vec{x})$ can be written as a vector with one entry equal to 1, the one corresponding to the correct class, and the others containing the overlap between the correct class label state and the other label states.

With these definitions, we can construct a cost function which turns out to be inspired by the conventional cost function in artificial neural networks. By weighting the fidelities of the final state of the circuit with all label states, we define the weighted fidelity cost function as

$$\chi^2_{wf}(\vec{\alpha}, \vec{\theta}, \vec{w}) = \frac{1}{2} \sum_{\mu=1}^{M} \left( \sum_{c} \left( \alpha_c F_c(\vec{\theta}, \vec{w}, \vec{x}_\mu) - Y_c(\vec{x}_\mu) \right)^2 \right),$$

where $M$ is the total number of training points, $C$ is the total number of classes, $\vec{x}_\mu$ are the training points and $\vec{\alpha} = (\alpha_1, \ldots, \alpha_C)$ are introduced as class weights to be optimized together with $\vec{\theta}$ and $\vec{w}$. This weighted fidelity has more parameters than the fidelity cost function. These parameters are the weights for the fidelities.

The main difference between the weighted fidelity cost function of Eq. (9) and the fidelity cost function of Eq. (7) is how many overlaps do we need to compute. The $\chi^2_{wf}$ requires as many fidelities as classes every time we run the optimization subroutine, while the $\chi^2$ needs just one. This is not such a big difference for few classes and only one qubit. It is possible to measure any state with a full tomography process which, for one qubit, is achievable. However, for many different classes we expect that one measurement will be more efficient than many.

Besides the weighted fidelity cost function is costlier than the fidelity cost function, there is another qualitative difference between both. The fidelity cost function forces the parameters to reach the maximum in fidelities. As a way of speaking, this fidelity moves the qubit state where it should be. The weighted fidelity forces the parameters to be close to a specified configuration of fidelities. It moves the qubit state to where it should be and moves it away from it should not. Therefore, we expect that the weighted fidelity will work better than the fidelity cost function.

III. UNIVERSALITY OF THE SINGLE-QUBIT CLASSIFIER

After analyzing several classification problems, we obtain evidences that the single-qubit classifier introduced above is able to classify any function. In this section, we provide the motivation for this statement based on the Universal Approximation Theorem (UAT) of artificial neural networks [11].

A. Universal Approximation Theorem

Theorem—Let $I_m = [0, 1]^m$ be the $m$-dimensional unit cube and $C(I_m)$ the space of continuous functions in $I_m$. Let the function $\varphi : \mathbb{R} \rightarrow \mathbb{R}$ be a nonconstant, bounded and continuous function. Then, there exist an integer $N$ such that

$$h(\vec{x}) = \sum_{i=1}^{N} \alpha_i \varphi (\vec{w}_i \cdot \vec{x} + b_i),$$

for $\alpha_i, b_i \in \mathbb{R}$ and $\vec{w}_i \in \mathbb{R}^m$, is an approximate realization of the function $f : \mathbb{R}^m \rightarrow \mathbb{R}$, i.e.,

$$|h(\vec{x}) - f(\vec{x})| < \epsilon,$$

for all $\vec{x} \in I_m$ and $\epsilon > 0$.

In artificial neural networks, $\varphi$ is the activation function, $\vec{w}_i$ are the weights for each neuron, $b_i$ are the biases and $\alpha_i$ are the neuron weights that construct the output function. Thus, this theorem establishes that it is possible to reconstruct any continuous function with a single layer neural network of $N$ neurons. The proof of this theorem for sigmoidal activation function can be found in Ref. [16]. This theorem was generalized for any nonconstant, bounded and continuous activation function in Ref. [11]. Moreover, Ref. [11] presents the following corollary of this theorem: $\varphi$ could be a nonconstant finite linear combination of periodic functions, in particular $\varphi$ could be a nonconstant trigonometric polynomial.

B. Universal Quantum Circuit Approximation

The single-qubit classifier is divided into several layers which are general SU(2) rotational matrices. There exist many possible decompositions of a a SU(2) rotational matrix. In particular, we use

$$U(\vec{\phi}) = U(\phi_1, \phi_2, \phi_3) = e^{i\phi_2 \sigma_y} e^{i\phi_1 \sigma_z} e^{i\phi_3 \sigma_z}.$$

Using the SU(2) group composition law, we can rewrite the above parametrization in a single exponential,

$$U(\vec{\phi}) = e^{i\vec{\omega}(\vec{\phi}) \cdot \vec{\sigma}},$$

with $\vec{\omega}(\vec{\phi}) = (\omega_1(\vec{\phi}), \omega_2(\vec{\phi}), \omega_3(\vec{\phi}))$ and

$$\omega_1(\vec{\phi}) = d N \sin ((\phi_2 - \phi_3)/2) \sin (\phi_1/2),$$

$$\omega_2(\vec{\phi}) = d N \cos ((\phi_2 - \phi_3)/2) \sin (\phi_1/2),$$

$$\omega_3(\vec{\phi}) = d N \sin ((\phi_2 + \phi_3)/2) \cos (\phi_1/2),$$

where $d = \cos (((\phi_2 + \phi_3)/2) \cos (\phi_1/2)$ and $N = (\sqrt{1 - \cos^2 d})^{-1}$.

The single-qubit classifier codifies the data points into $\vec{\phi}$ parameters of the $U$ unitary gate. In particular, we
can re-upload data together with the tunable parameters as defined in Eq. (5), i.e.

$$\vec{\phi}(\vec{x}) = (\phi_1(\vec{x}), \phi_2(\vec{x}), \phi_3(\vec{x})) = \vec{\theta} + \vec{w} \circ \vec{x}. \quad (17)$$

Thus,

$$\mathcal{U}(\vec{x}) = U_N(\vec{x})U_{N-1}(\vec{x})\cdots U_1(\vec{x}) = \prod_{i=1}^{N} e^{i\vec{\omega}(\vec{\phi}_i(\vec{x})) \cdot \vec{\sigma}}, \quad (18)$$

Next, we apply the Baker-Campbell-Hausdorff (BCH) formula (17) to the above equation,

$$\mathcal{U}(\vec{x}) = \exp \left[ i \sum_{i=1}^{N} \vec{\omega}(\vec{\phi}_i(\vec{x})) \cdot \vec{\sigma} + \mathcal{O}_{\text{corr}} \right]. \quad (19)$$

Notice that the remaining BCH terms $\mathcal{O}_{\text{corr}}$ are also proportional to Pauli matrices due to $[\vec{\sigma}_i, \vec{\sigma}_j] = 2i\epsilon_{ijk}\vec{\sigma}_k$.

Each $\vec{\omega}$ term is trigonometric functions, unconstant, bounded and continuous. Then

$$\sum_{i=1}^{N} \vec{\omega}(\vec{\phi}_i(\vec{x})) = \sum_{i=1}^{N} (\omega_1(\vec{\phi}_1(\vec{x})), \omega_2(\vec{\phi}_2(\vec{x})), \omega_3(\vec{\phi}_3(\vec{x})))$$

$$= \sum_{i=1}^{N} (\omega_1(\vec{\theta}_i + \vec{w}_i \circ \vec{x}), \omega_2(\vec{\theta}_i + \vec{w}_i \circ \vec{x}), \omega_3(\vec{\theta}_i + \vec{w}_i \circ \vec{x}))$$

$$= (f_1(\vec{x}), f_2(\vec{x}), f_3(\vec{x})). \quad (20)$$

We still have to deal with the remaining terms $\mathcal{O}_{\text{corr}}$ of the BCH expansion. Instead of applying such expansion, we can use again the SU(2) group composition law to obtain the analytical formula of $\mathcal{U}(\vec{x}) = e^{i\vec{\xi}(\vec{x}) \cdot \vec{\sigma}}$, where $\vec{\xi}(\vec{x})$ will be an inextricably trigonometric function of $\vec{x}$. The $\mathcal{O}_{\text{corr}}$ terms are proportional to $\vec{\sigma}$ matrices, so $\mathcal{O}_{\text{corr}} = \vec{\xi}(\vec{x}) \cdot \vec{\sigma}$ for some function $\vec{\xi}(\vec{x})$. Then,

$$\mathcal{U}(\vec{x}) = e^{i\vec{\xi}(\vec{x}) \cdot \vec{\sigma}} = e^{i\vec{f}(\vec{x}) \cdot \vec{\sigma} + i\vec{g}(\vec{x}) \cdot \vec{\sigma}}. \quad (21)$$

Thus, $\mathcal{O}_{\text{corr}}$ terms can be absorbed in $\vec{f}(\vec{x})$.

For each data point $\vec{x}$, we obtain a final state that will contain these $\vec{\xi}(\vec{x})$ functions. With all training points, we construct a cost function that can include new parameters $\alpha_c$ for each class if we use the weighted fidelity cost function of Eq. (10).

The function obtained from the combination of $\vec{\xi}(\vec{x})$ and $\alpha_c$ is expected to be complex enough to probably represent almost any continuous function. However, more parameters are necessary in order to map this argument with the UAT expression.

If we compare the parameters of the UAT with the single-qubit circuit parameters, the $\vec{w}_i$ will correspond with the weights, the $\vec{\theta}_i$ with the biases $b_i$, the number of layers $N$ of the quantum classifier will correspond with the number of neurons in the hidden layer and $\vec{\omega}$ functions with the activation functions $\varphi$.

We have provided an explanation about why is necessary to re-upload the data at each layer and why a single qubit could be a universal classifier. As has been stated before, an artificial neural network introduces the data points in each hidden neuron, weights them and adds some bias. Here we cannot just copy each data point because the non-cloning theorem, so we have to re-upload it at each layer.

IV. FROM SINGLE- TO MULTI-QUBIT QUANTUM CLASSIFIER

The single-qubit classifier can not carry any quantum advantage in front of classical classification techniques such artificial neural networks. In the previous sections, we have defined a quantum mechanical version of a neural network with a single hidden layer. In general, it is necessary a huge amount of hidden neurons to approximate a target function with a single layer. To circumvent this inconvenience, more hidden layers are introduced, leading eventually to the concept of deep neural networks.

By using the single-qubit classifier formalism that we have introduced in the previous sections, we propose its generalization to more qubits. The introduction of multiple qubits to this quantum classifier may improve the performance of it in a similar way as more hidden layers improves the classification task of an artificial neural network. With the introduction of entanglement between these qubits, we reduce the number of layers of our classifier as well as propose a quantum classification method that can achieve quantum advantage.

A. Measurement strategy and cost function for a multi-qubit classifier

With a single-qubit classifier, the measurement strategy consisting on comparing the final state of the circuit with a pre-defined target state was achievable. Experimentally, one needs to perform a quantum state tomography protocol of only three measurements. However, if more qubits are to be considered, tomography protocols become exponentially expensive in terms of number of measurements.

We propose two measurement strategies for a multi-qubit classifier. The first one is the natural generalization of the single-qubit strategy, although it will become unrealizable for a large number of qubits. We compare the final state of the circuit with one of the states of the computational basis, one for each class. The second strategy consist on focusing in one qubit and depending on its state associate one or other class. This is similar to previous proposals of binary multi-qubit classifiers [10], although we add the possibility of multiclass classification by introducing several thresholds (see section II).

Another part that should be adapted is the definition of the cost function. In particular, we use different functions for each strategy explained above.
and similarly for a four-qubit classifier, we identify each work. For a two-qubit classifier without entanglement, circuit-design: how to introduce the entanglement. qubit quantum classifier there is an extra freedom in the edition, it is, in general, problem-dependent. For a multi-layers and neurons per layer to perform some task. In ad-

neural network. In artificial neural networks, it is far

reducing the number of parameters to be optimized.

Eventually, we can just measure one of these qubits, Q |ψ⟩ is the reduced density matrix of the qubit to be

considered to improve the performance of this method.

For the second strategy, we use the weighted fidelity cost function. As stated above, we just focus on one qubit, thus

F_{c,q}(\vec{\theta}, \vec{w}, \vec{x}) = \langle \tilde{\psi}_c | \rho_q(\vec{\theta}, \vec{w}, \vec{x}) | \tilde{\psi}_c \rangle, \quad (22)

where \( \rho_q \) is the reduced density matrix of the qubit to be measured. Then, the weighted fidelity cost function can be adapted as

\[
\chi^2_{wf}(\vec{\alpha}, \vec{\theta}, \vec{w}) = \frac{1}{2} \sum_{\mu=1}^{M} \sum_{c=1}^{C} \sum_{Q=1}^{Q} \left( \alpha_{c,q} F_{c,q}(\vec{\theta}, \vec{w}, \vec{x}_\mu) - Y_c(\vec{x}_\mu) \right)^2,
\] \quad (23)

where we average over all Q qubits that form the classifier. Eventually, we can just measure one of these qubits, reducing the number of parameters to be optimized.

For the first strategy, we use the fidelity cost func-

\[
F_{c,q}(\theta, \vec{w}, \vec{x}) = \langle \tilde{\psi}_c | \rho_q(\theta, \vec{w}, \vec{x}) | \tilde{\psi}_c \rangle, \quad (21)
\]

layer as parallel rotations on all qubits. We introduce the entanglement using CZ gates between each rotations that are absorbed in the definition of layer. For two-qubit classifier with entanglement, we apply a CZ gate after each rotation with the exception in the last layer. For a four-qubit classifier, two CZ gates are applied after each rotation with the exception in the last layer. For a multi-qubit classifier there is an extra freedom in the circuit-design: how to introduce the entanglement.

For the second strategy, we use the weighted fidelity cost function. As stated above, we just focus on one qubit, thus

\[
F_{c,q}(\vec{\theta}, \vec{w}, \vec{x}) = \langle \tilde{\psi}_c | \rho_q(\vec{\theta}, \vec{w}, \vec{x}) | \tilde{\psi}_c \rangle, \quad (22)
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where \( \rho_q \) is the reduced density matrix of the qubit to be measured. Then, the weighted fidelity cost function can be adapted as

\[
\chi^2_{wf}(\vec{\alpha}, \vec{\theta}, \vec{w}) = \frac{1}{2} \sum_{\mu=1}^{M} \sum_{c=1}^{C} \sum_{Q=1}^{Q} \left( \alpha_{c,q} F_{c,q}(\vec{\theta}, \vec{w}, \vec{x}_\mu) - Y_c(\vec{x}_\mu) \right)^2,
\] \quad (23)

where we average over all Q qubits that form the classifier. Eventually, we can just measure one of these qubits, reducing the number of parameters to be optimized.

\[ F_{c,q}(\vec{\theta}, \vec{w}, \vec{x}) = \langle \tilde{\psi}_c | \rho_q(\vec{\theta}, \vec{w}, \vec{x}) | \tilde{\psi}_c \rangle, \quad (21) \]

layer as parallel rotations on all qubits. We introduce the entanglement using CZ gates between each rotations that are absorbed in the definition of layer. For two-qubit classifier with entanglement, we apply a CZ gate after each rotation with the exception in the last layer. For a four-qubit classifier, two CZ gates are applied after each rotation alternatively between (1)-(2) and (3)-(4) qubits and (2)-(3) and (1)-(4) qubits.

The number of parameters needed to perform the opti-

mization doubles the ones needed for a single-qubit classifier, and quadruples for the four-qubit classifier. For N layers, the circuit depth is N for the non-entangling classifiers and 2N for the entan-

gling classifiers.

V. MINIMIZATION METHODS

The practical training of a parametrized single-qubit or multi-qubit quantum classifier needs merging all the above structure with some machine learning algorithm to proceed with the minimization in the parameter space describing the circuit. This is often referred as a hybrid algorithm, where classical and quantum logic coexist and benefit from each other. To be precise, the set of \{\theta_i\} angles and \{w_i\} weights, together with \alpha_{q,t} parameters if applicable, forms a space to be explored in search of a minimum \( \chi^2 \). This may be non-trivial and the ap-
pearance of many false minima is likely. A bad classical minimizer can easily get trapped in those minima.

Our problem is reduced to minimize a function of many parameters. For a single-qubit classifier, the number of parameters is $(3 + d)N$ where $d$ is the dimension of the problem, i.e. the dimension of $\vec{x}$, and $N$ is the number of layers. Three of these parameters are the rotational angles and the other $d$ correspond with the $\vec{w}_i$ weight. If using the weighted fidelity cost function, we should add $C$ extra parameters, one for each class.

In principle, one does not know how is the landscape of the cost function to be minimized. If the cost function were, for example, a really convex function, a downhill strategy would be likely to work properly. The pure downhill strategy is known as gradient descent. In machine learning, the method commonly used is a Stochastic Gradient Descent (SGD) \[18\].

There is another special method of minimization known as L-BFGS-B \[19\]. This method has been used in classical machine learning with very good results \[20\].

The results we present from now are starred by the L-BFGS-B algorithm, as we found it is accurate and relatively fast. We used open source software \[21\] as the core of the minimization with own made functions to minimize. The minimizer is taken as a black box whose parameters are set by default. As this is the first attempt of constructing a single- or multi-qubit classifier, further improvements can be done on the hyperparameters of minimization.

Nevertheless we have also tested a SGD algorithm for the fidelity cost function. This whole algorithm has been developed by us following the steps from \[15\]. The details can be read in App. A. In general, we found that L-BFGS-B algorithm is better than SGD. This is something already observed in classical neural networks. When the training set is small, it is often more convenient to use a L-BFGS-B strategy than a SGD. We were forced to use small training sets due to computational capabilities for our simulations. Numerical evidences on this arise when solving the problems we face for this single- and multi-qubit classifiers with classical standard machine learning libraries \[20\]. This can be understood with a simple argument. Neural networks or our quantum classifier are supposed to have plenty of local minima. Neural networks have got huge products of non linear functions. The odds of having local minima are then large. In the quantum circuits side, there are nothing but trigonometric functions. In both cases, if there are a lot of training points it is more likely to find some of them capable of getting us out of local minima. If this is the case, SGD is more useful for being faster. On the contrary, when the training set is small, we have to pick an algorithm less sensitive to local minima, such as the L-BFGS-B.

VI. BENCHMARK OF A SINGLE- AND MULTI-QUBIT CLASSIFIER

We can now tackle some classification problems. We will prove that a single-qubit classifier can perform a multi-class classification for multi-dimensional data and that a multi-qubit classifier, in general, improves these results.

We construct several classifiers with different number of layers. We then train the circuits with a training set of data points to obtain the values of the free parameters $\{\theta_i\}$ and $\{w_{ij}\}$ for each layer and $\{\alpha_i\}$ when applicable. We perform these calculations for cost functions defined in Eq. \[9\] and Eq. \[7\].

We run a single-, two- and four-qubit classifiers, with and without entanglement, using the two cost functions described above. We benchmark several classifiers formed by $L = 1, 2, 3, 4, 5, 6, 8$ and 10 layers.

In the following subsections, we describe the particular problems addressed with these single- and multi-qubit classifiers with data re-uploading. We choose four problem types: a simple binary classification, a classification of a figure with multiple patterns, a multi-dimensional classification and a non-convex figure.

The code used to define and benchmark the single- and multi-qubit quantum classifier is open and can be found in Ref. \[22\].

A. Simple example: classification of a circle

Let us start with a simple example. We create a random set of data on a plane with coordinates $\vec{x} = (x_1, x_2)$ with $x_i \in [-1, 1]$. Our goal is to classify these points according to $x_1^2 + x_2^2 < r^2$, i.e. if they are inside or outside of a circle of radius $r$. The value of the radius is chosen in such a way that the areas inside and outside it are equal, that is, $r = \sqrt{\frac{2}{\pi}}$, so the probability of success if we label each data point randomly is 50%. We create a train dataset with 200 random entries. We then validate the single-qubit classifier against a test dataset with 4000 random points.

The results of this classification are written in Tab. I. With the weighted fidelity cost function, the single-qubit classifier achieves more than 90% of success with only two layers, that is, 12 parameters. The results are worse with the fidelity cost function. For a two-qubit and a four-qubit classifier, two layers are required to achieve 96% of success rate, that is, 22 parameters for the two-qubit and 42 for the four-qubit. The introduction of entanglement does not change the result in any case. The results show a saturation of the success rate. Considering more layers or more qubits does not change this success rate.

The characterization of a closed curved is a hard problem for an artificial neural network that work in a linear regime, although many different neurons, i.e. linear terms, can achieve a good approximation to any function.
Table I: Results of the single- and multi-qubit classifiers with data re-uploading for the circle problem. Numbers indicate the success rate, i.e. number of data points classified correctly over total number of points. Words "Ent." and "No Ent." refer to considering entanglement between qubits or not respectively. We have used the L-BFGS-B minimization method with the weighted fidelity and fidelity cost functions. For this problem, both cost functions lead to high success rates. The multi-qubit classifier increases this success rate but the introduction of entanglement does not affect it significantly.

On the contrary, the layers of a single-qubit classifier are rotational gates, which have an intrinsic non-linear behavior. In a sense, a circle becomes an easy function to classify as a linear function is for an artificial neural network. The circle classification is, in a sense, trivial for a quantum classifier. We need to run these classifiers with more complex figures or problems to really test their performance.

It is interesting to compare classifiers with different number of layers. Figure 6 shows the result of the classification for a single-qubit classifier of 1, 2, 3 and 10 layers. As with only one layer the best that the classifier can do is to divide the plane in half. With two layers, it catches the circular shape which is readjusted as we consider more layers.

B. Classification of multiple patterns

We want to show now that the single-qubit classifier is able to solve multi-class problems. We divide a 2D plane into several regions and assign a label to each one. We propose the following division: three regions corresponding to three circular sectors and the intermediate space between them. We call this problem the 3-circles problem. This is a hardly non-linear problem and, consequently, difficult to solve for a classical neural network in terms of computational power.

Table II shows the results for this four-class problem. For a single-qubit classifier, a maximum of 92% of success is achieved with 10 layers, i.e. 54 parameters. From these results, it seems that this problem also saturates around 91% of success. However, the introduction of more qubits and entanglement makes possible this result with less parameters. For two qubits with entanglement, 4 layers are necessary to achieve the same success as with a single-qubit, i.e. 34 parameters. For four qubits without entanglement also requires 4 layers. Notice also that, although the number of parameters increase significantly with the number of qubits, some of the effective operations are performed in parallel.

There is an effect that arises from this more complex classification problem: local minima. Notice that the success rate can decrease when we add more layers into our quantum classifier.

As with the previous problem, it is interesting to observe the explicit results of different classifiers with different number of layers. Figure 7 shows the results for a two-qubit classifier with no entanglement for 1, 2, 3 and 4 layers. Even with only one layer, the classifier identifies the four regions, being the more complicated to describe the central one. As we consider more layers, the classifier performs better and adjust these four regions.

C. Classification in multiple dimensions

As explained in section IV, there is no restriction in uploading multidimensional data. Actually, we can update up to three points per rotation since this is the degrees
of freedom of a SU(2) matrix. If the dimension of data is larger than that, we can just split the data vector into subsets and upload each one at a time, as described explicitly in Eq. (6). Therefore, there is no reason to limit the dimension of data to the numbers of degrees of freedom of a qubit. We can in principle upload any kind of data if we apply enough gates.

Following this idea we will now move to a more complicated classification using data with 4 coordinates. We use as a problem the four-dimensional hypersphere, i.e. classifying data points according to \(x^2 + x^2 + x^2 + x^2 < 2/\pi\). Similarly with the previous problems, \(x_i \in [-1, 1]\) and the radius has been chosen such that the volume of the hypersphere is half of the total volume. This time, we will take 1000 random points as the training set because the total volume increases.

Results are shown in Tab. III. With a single-qubit, it achieves the 97% of success with eight layers (82 parameters) using the weighted fidelity cost function. Results are better if we consider more qubits. For two qubits, the best result is 98% and it only requires three entangled layers (62 parameters). For four qubits, it achieves the 98% of success with two layers with entanglement, i.e. 82 parameters.

### D. Classification of non-convex figures

As a final benchmark, we propose the classification of a non-convex pattern. In particular, we classify the points of an annulus with radius \(r_1 = \sqrt{0.8 - 2/\pi}\) and \(r_2 = \sqrt{0.8}\). We fix three classes: points inside the small circle, points in the annulus and points outside the big circle.
E. Comparison with classical classifiers

It is important to check if our proposal is in some sense able to compete with actual technology of supervised machine learning. To do so we have used the standard machine learning library scikit [20] and solved the same problems as we have solved with the quantum classifier. We have included the four problems presented in the main paper plus five extra problems analyzed in App. B. The aim of this classical benchmarking is not to make an extended review of what classical machine learning is capable to perform. The aim is to compare our simple quantum classifier to simple models such as narrow neural networks and simple support vector machines.

The technical details of the classical classification are the following: the neural network has got one hidden layer with 100 neurons, a ReLu activation function and the solver lbfgs by scikit. The support vector machine is the default sklearn.svm.SVC. Some changes in the initialization parameters were tested with no significant differences.

Table V compares the best performance of a neural network, support vector classifier (SVC), single-qubit classifier with fidelity cost function and single-qubit classifier with weighted fidelity cost function. In all problems, the performance of the single-qubit classifier is, at least, comparable with the classical methods. Some problems like 3-circles problem and binary annulus problem the results of the single-qubit classifier are better than with these methods.

VII. CONCLUSIONS

We have proposed a single-qubit classifier that is able to represent multidimensional complex figures. The core of this quantum classifier is the data re-uploading. This formalism allows to circumvent the limitations of the no-cloning theorem in order to achieve a correct generalization of an artificial neural network with a single layer.

The structure of this classifier is the following. Data and processing parameters are uploaded multiple times with a single qubit rotation along the circuit. These processing parameters are different at each upload and
Figure 8: Results obtained with the single-qubit classifier for the annulus problem, using the weighted fidelity cost function during the training. The better results are obtained with a 10 layers classifier (93% of success rate). As we consider more qubits and entanglement, we can increase the success rate up to 96%, as shows Tab. IV.

should be optimized using a classical minimization algorithm. To do so, we have defined two cost functions: one inspired in the traditional neural networks cost functions (weighted fidelity cost function) and the other, simpler, consisting on the computation of the fidelity of the final state respect to a target state. These target states are defined to be maximally orthogonal between them. Then, the single-qubit classifier finds the optimal rotations to separate the data points into different regions of the Bloch sphere, each one corresponding with a class.

The single-qubit classifier can be generalized to a larger number of qubits. This allows the introduction of entanglement between these qubits by adding two-qubit gates between each layer of rotations.

We have benchmarked several quantum classifiers of this kind, made of different number of layers, qubits and with and without entanglement. The patterns chosen to test these classifiers are the points inside and outside of a circle (simple example) and similarly for a four-dimensional hypersphere (multidimensional example); a two dimensional region composed by three circles of different size (multiple classes example); and the points outside and inside of an annulus (non-convex example). In all cases, the single-qubit classifier achieves more than 90% of success rate. The introduction of more qubits and entanglement increases this success and reduces the number of layers required. The weighted fidelity cost function turns out to be more convenient to achieve better results than the fidelity cost function. In all problems, the probability to get stuck in a local minima increases with the number of layers, an expected result from an optimization problem involving several parameters.

In summary, we have proposed a quantum classifier model that seems to be universal by exploiting the non-linearities of the single-qubit rotational gates and by re-uploading data several times.

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[1] N. Wiebe, D. Braun, and S. Lloyd, Phys. Rev. Lett. 109, 050505 (2012).
[2] P. Rebentrost, M. Mohseni, and S. Lloyd, Phys. Rev. Lett. 113, 130503 (2014).
[3] M. Schuld, I. Sinayskiy, and F. Petruccione, Quantum Information Processing 13, 2567 (2014).
Appendix A: The Stochastic Gradient Descent method (SGD)

We propose an algorithm which is analogous to the stochastic gradient descent algorithm for neural networks. This kind of computation is not new, as similar proposals can be found in \cite{9, 10, 23}. We show that this algorithm can be interpreted as a back-propagation, as it takes intermediate steps of the circuit for exactly computing the gradient of the function $\chi_f^2$.

1. To compute the gradient

Our goal is to find the gradient of $\chi_f^2$ for one data point of the training set. In terms of Eq. (2) and Eq. (4),

$$|\langle \psi_c | \psi \rangle|^2 = |\langle \psi_c | L(N) L(N-1) \cdots L(1) | 0 \rangle|^2.$$  \hspace{1cm} (A1)

Let us write explicitly an intermediate quantum state $|\psi_l\rangle$ of the single-qubit classifier,

$$|\psi_l\rangle = \prod_{k=0}^l L(k) |0\rangle = L(l) |\psi_{l-1}\rangle,$$  \hspace{1cm} (A2)

where $1 \leq l \leq N$ and $|\psi_0\rangle \equiv |0\rangle$. Next, notice that we can recursively define the counterpart of the label states $|\psi_c\rangle$ as

$$\langle \Delta_l | = \langle \tilde{\psi}_c | L(N) L(N-1) \cdots L(l+1) = \langle \Delta_{l+1} | L(l+1),$$  \hspace{1cm} (A3)

where $1 \leq l \leq N$ and $\langle \Delta_N | \equiv \langle \tilde{\psi}_c |$. A way to understand this notation is by checking that $|\psi_l\rangle$ counts gates from the beginning of the circuit, starting from first layer $L(1)$, and $\langle \Delta_l |$ counts backwards from the end of the circuit, starting from $N$th layer. Thus, Eq. (A1) can be rewritten as

$$|\langle \psi_c | \psi \rangle|^2 = |\langle \Delta_l | \psi_l \rangle|^2,$$  \hspace{1cm} (A4)

where $l \in [1, N]$ defines the circuit layer where the gradient will be computed.

Each layer $L(l)$ contains as much as six tunable parameters: three rotational angles represented with the vector $\vec{\theta}_l$ and three weights $\vec{w}_l$. For a problem of dimension $d > 3$, each layer is composed by sublayers with up to six tunable parameters, as shown in Eq.(6). Then, when we derive with respect to one parameter, only the corresponding layer gate will be affected by it. We can split the set of gates in two parts: those before the derivative gate and those after. Using the recursive definitions presented above, the derivative with respect to $\theta^i_l$ of $\chi_f^2$ can be written as

$$\frac{\partial \chi_f^2}{\partial \theta^i_l} = -2 Re \left\{ \left( \langle \Delta_l | \frac{\partial L(l)}{\partial \theta^j_l} | \psi_{l-1} \rangle \right) \langle \psi_l | \Delta_l \rangle \right\},$$  \hspace{1cm} (A5)

for $i = 1, 2, 3$ and similarly with $\partial \chi_f^2 / \partial \vec{w}_l$. This derivative is not unitary. Nevertheless, this fact has not importance in our algorithm since we will use its value to
update the values of the new parameters in the next iteration classically.

The computation of the partial derivatives of $L(l)$ is straightforward do to its matrix unitary structure. In general, $L(l)$ is a general SU(2) matrix which can be parametrized as

$$L(l) = U\left(\varphi_l\right) = \begin{pmatrix} \cos\left(\frac{\varphi_l}{2}\right)e^{i\left(\sigma_2^0 + \sigma_2^0\right)} & -\sin\left(\frac{\varphi_l}{2}\right)e^{-i\left(\sigma_2^0 + \sigma_2^0\right)} \\ \sin\left(\frac{\varphi_l}{2}\right)e^{i\left(\sigma_2^0 + \sigma_2^0\right)} & \cos\left(\frac{\varphi_l}{2}\right)e^{-i\left(\sigma_2^0 + \sigma_2^0\right)} \end{pmatrix},$$

where $\varphi_l = (\varphi_l^1, \varphi_l^2, \varphi_l^3) = \tilde{\varphi}_l + \tilde{w}_l \circ \tilde{x}$. Thus,

$$\frac{\partial L(l)}{\partial \theta_l^i} = \frac{\partial L(l)}{\partial \phi_l^i},$$

$$\frac{\partial L(l)}{\partial w_l^i} = \frac{\partial L(l)}{\partial \chi_l^i} x^i,$$

The key feature of taking the $L(l)$ parametrization of Eq. (A6) is that its derivatives are almost itself, which simplifies its computation significantly. Notice that

$$\frac{\partial L(l)}{\partial \theta_l^i} = \frac{\partial U(\tilde{\varphi}_l + \tilde{w}_l \circ \tilde{x})}{\partial \theta_l^i} = \frac{1}{2} U(\tilde{\varphi}_l + \tilde{w}_l \circ \tilde{x} + \pi \tilde{\varphi}_l),$$

where $\tilde{\varphi}_l$ is a vector of dimension three which components are all zeros except the $i$-th one. The calculations for the derivative become simpler in this case, as we just have to shift the proper parameter by $\pi$, and use it to measure the derivative.

Once we have computed the gradient of the cost function $\chi^2_l$, we can update the parameters by setting

$$\theta_l^i \leftarrow \theta_l^i - \eta \frac{\partial \chi^2_l}{\partial \theta_l^i},$$

$$w_l^i \leftarrow w_l^i - \eta \frac{\partial \chi^2_l}{\partial w_l^i},$$

where $\eta$ is an adjustable learning rate.

2. To update parameters

There are several ways of updating the parameters. One naive option is to update parameters every time we compute the gradient of the function. Then we start all over again and repeat until the parameters become stable. This is not a good practice. If we do so, we are moving point by point, optimizing and classifying every data from the training set of our classifier. This classifies all points slowly. At late steps of the minimization process, we will have to face plenty of points which are already classified and do not have important gradients. In summary, this is a very inefficient algorithm.

Another option is to compute the gradients for every point, and then take the average of all gradients and update parameters with it. However, this method is likely to achieve poor accuracies. Some gradients can be cancelled out by some others and the stability is achieved before it should be.

The method that is usually used is the so-called batch optimization. This method is a mix of the two previous ways of tackling the problem. First, we have to split the training set into smaller subsets or batches. We compute the averaged gradient of the first batch with respect to all points in the batch and then update. Then, proceed similarly with the second batch. We keep going until we finish all batches. The crucial step is to shuffle the training set and split it into new different mixed batches. This way, we optimize the classifier with respect to subsets of the original training set that are constantly changing which is more efficient than a point-by-point optimization, but statistically equivalent. In our SGD algorithms we took batches with 20 points out of a total amount of 200 points for the two-dimensional problems, 500 for three-dimensional problems and 1000 for four-dimensional problems.

Although we tested this SGD algorithm, we obtained worse results than with the L-BFGS-B method. In addition, the computational efforts for this SGD algorithm were larger as could not be fully optimized. Thus we discard for the moment this SGD minimization method.

Appendix B: Classification results in other problems

We tested our classifier in several different problems. Here, we show a summary of the results obtained in the other problems we tackled. As the ones defined in the main paper, we constrained the data space to $x \in [-1,1]$ and $y \in [0,1]$. The aim of these extra classification tasks is to cover different kinds of training data, and show that our quantum classifier is able of adapt itself to large varieties of problems.

The datasets have 200 random entries – with the exception of the sphere, which have 500 – and test the performance of the classifier against 4000 random points. All problems were designed without biases, i.e. a blind classifier will get successes of $\sim 50\%$ for a binary classifier, $\sim 33\%$ for a ternary classifier and $\sim 25\%$ for a quaternary classifier.

In general, we can observe that the classifier improves its performance as we increase the number of layers until it comes into a stationary regime. More qubits and entanglement usually allows entering in the stationary regime with a lesser number of layers.

In the following subsections, we define these extra problems and present the results of each one summarized in a table. The best results obtained in each problem are plotted in Fig. [9].
The non-convex problem is made for testing two mutually non-convex zones dividing a 2D area. The border of both zones lies on the function $x_2 = -2x_1 + 3/2 \sin(\pi x_1)$. This border is chosen in a way that there is no area small enough as to leave it unclassified, forcing the classifier to catch these smaller zones.

Complete results can be read in Tab. VI. The weighted fidelity cost function gets the best performance, 98%, with 6 layers and 1 qubit, (32 parameters). Fidelity cost gets a result of 96% for the same conditions, while the best, 97%, is achieved for 2 entangled qubits with 8 layers (80 parameters). The final figure obtained after the classification can be seen in Fig. 9(a).

We can compare this result with the one obtained in Tab. V. Neural networks work really well as they can approximate our non convex function properly. It achieves the 0.98 success with two qubits, entanglement and four layers.

b. Binary annulus

This a binary version of the annulus problem. The two classes are defined as being inside or outside the annulus with radii $r_1 = \sqrt{0.8 - 2/\pi}$ and $r_2 = \sqrt{0.8}$. This geometry, as happens with the three-class annulus problem, is interesting because the classifier finds a way to connect areas of the problem which are disconnected. This involves a to and fro path for the parameters.

The fidelity cost function reaches the best result at 94%, with 2 qubits with entanglement and 6 layers, that is 60 parameters. For the weighted fidelity we reach up to 97% with 2 qubits with no entanglement and 4 layers, 40 parameters. Results are written in Tab. VII and the best result is plotted in Fig. 9(b).

If we compare these results to the classical ones in Tab. V we see again that the SVC method is way worse than the NN, which can be interpreted as the sign of non convexity this problem has.

c. Sphere

This quantum classifier is able to classify multidimensional data, as we have shown with the four-dimensional hypersphere. We also tested a three-dimensional figure, a sphere with radius $r = \sqrt{33}/\pi$.

For this problem, fidelity cost function reaches its maximum, 93%, with a single-qubit classifier of 10 layers (60 parameters). The same success is obtained with a two-qubit entangled classifier and 6 layers (72 parameters). With the weighted fidelity, this success rate grows up to 96% for two- and four- qubit classifier of 2 layers (24 and 48 parameters respectively) with and without entanglement. All results are written in Tab. VIII.

d. Squares

This problem divides a 2D area in four quadrants with straight lines. This is one of the easiest problems for a neural network. By construction, neural networks can
The best performance is plotted in Fig. 9(c).

Table VIII: Results of the single- and multi-qubit classifiers with data re-uploading for the three-dimensional sphere problem.

| Qubits | Layers | $\chi^2_f$ | No Ent. | Ent. | No Ent. | Ent. | No Ent. | Ent. |
|--------|--------|-----------|--------|------|--------|------|--------|------|
| 1      | 1      | 0.53      | 0.70   | 0.53 | 0.70   | 0.70 | 0.70   | 0.70 |
| 2      | 0.77   | 0.73      | 0.53   | 0.78 | 0.94   | 0.96 | 0.96   | 0.96 |
| 3      | 0.76   | 0.74      | 0.77   | 0.78 | 0.92   | 0.96 | 0.94   | 0.96 |
| 4      | 0.84   | 0.83      | 0.78   | 0.89 | 0.92   | 0.95 | 0.95   | 0.93 |
| 5      | 0.89   | 0.85      | 0.82   | 0.90 | 0.94   | 0.93 | 0.95   | 0.92 |
| 6      | 0.90   | 0.89      | 0.93   | 0.92 | 0.87   | 0.94 | 0.94   | 0.92 |
| 8      | 0.89   | 0.87      | 0.84   | 0.93 | 0.92   | 0.93 | 0.94   | 0.92 |
| 10     | 0.93   | 0.91      | 0.90   | 0.93 | 0.94   | 0.95 | 0.92   | 0.92 |

Table IX: Results of the single- and multi-qubit classifiers with re-uploading data for the four-classes squares problem.

| Qubits | Layers | $\chi^2_f$ | No Ent. | Ent. | No Ent. | Ent. | No Ent. | Ent. |
|--------|--------|-----------|--------|------|--------|------|--------|------|
| 1      | 1      | 0.58      | 0.48   | 0.70 | 0.92   | 0.90 | 0.90   | 0.90 |
| 2      | 0.76   | 0.96      | 0.96   | 0.74 | 0.91   | 0.94 | 0.95   | 0.95 |
| 3      | 0.90   | 0.96      | 0.97   | 0.90 | 0.94   | 0.95 | 0.95   | 0.94 |
| 4      | 0.89   | 0.98      | 0.96   | 0.88 | 0.94   | 0.96 | 0.95   | 0.94 |
| 5      | 0.91   | 0.97      | 0.95   | 0.89 | 0.94   | 0.94 | 0.95   | 0.94 |
| 6      | 0.92   | 0.99      | 0.97   | 0.93 | 0.94   | 0.94 | 0.95   | 0.94 |
| 8      | 0.93   | 0.98      | 0.96   | 0.93 | 0.94   | 0.94 | 0.95   | 0.94 |
| 10     | 0.94   | 0.97      | 0.95   | 0.94 | 0.94   | 0.94 | 0.94   | 0.95 |

Table X: Results of the single- and multi-qubit classifiers with re-uploading data for the four-classes wavy lines problem.

| Qubits | Layers | $\chi^2_f$ | No Ent. | Ent. | No Ent. | Ent. | No Ent. | Ent. |
|--------|--------|-----------|--------|------|--------|------|--------|------|
| 1      | 2      | 0.70      | 0.52   | 0.76 | 0.75   | 0.88 | 0.88   | 0.91 |
| 2      | 0.86   | 0.75      | 0.80   | 0.84 | 0.89   | 0.88 | 0.91   | 0.92 |
| 3      | 0.74   | 0.82      | 0.88   | 0.84 | 0.92   | 0.92 | 0.92   | 0.92 |
| 4      | 0.80   | 0.85      | 0.91   | 0.87 | 0.89   | 0.91 | 0.92   | 0.94 |
| 5      | 0.85   | 0.90      | 0.91   | 0.87 | 0.92   | 0.92 | 0.93   | 0.93 |
| 6      | 0.92   | 0.92      | 0.91   | 0.88 | 0.93   | 0.93 | 0.93   | 0.93 |
| 8      | 0.90   | 0.91      | 0.92   | 0.92 | 0.92   | 0.93 | 0.93   | 0.94 |
| 10     | 0.92   | 0.91      | 0.94   | 0.90 | 0.93   | 0.92 | 0.93   | 0.93 |

Table IX: Results of the single- and multi-qubit classifiers with re-uploading data for the four-classes wavy lines problem.

This problem is the four-class version of the non convex problem. Now the area is divided in four regions by two different functions. The borders’ equations are $x_2 = \sin(\pi x_1) \pm x_1$. The important feature of this problem is that there are some areas in the problem too small to be caught by the classifier.

As can be seen in Fig. 9(d) most of failed points are in these small non convex areas. The classifier would rather adjust the rest of the points instead tuning those zones and losing everything else. The results for this problem are not as good as for other problems, but we still get 94%
Figure 9: Best results for the 2D problems analyzed in this appendix. The caption of each figure includes the architecture of the classifier, i.e. number of qubits, layers and the use of entanglement. In case of equal results with different classifiers, the simplest architecture has been chosen. Colors in left part of each figure represent the different classes obtained from the classifier outputs. Right part of each figure prints the points correctly classified (in green) and bad classified (in red). Black solid lines define the problem boundaries. Notice that all missed points are located near the borders of the problem. This means that the classifier is understanding properly which are the problem most important features, but it lacks of more training points. This can be easily corrected if we increase the number of training points.