Quantum circuit architecture search for variational quantum algorithms

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Variational quantum algorithms (VQAs) are expected to be a path to quantum advantages on noisy intermediate-scale quantum devices. However, both empirical and theoretical results exhibit that the deployed ansatz heavily affects the performance of VQAs such that an ansatz with a larger number of quantum gates enables a stronger expressivity, while the accumulated noise may render a poor trainability. To maximally improve the robustness and trainability of VQAs, here we devise a resource and runtime efficient scheme termed quantum architecture search (QAS). In particular, given a learning task, QAS automatically seeks a near-optimal ansatz (i.e., circuit architecture) to balance benefits and side-effects brought by adding more noisy quantum gates to achieve a good performance. We implement QAS on both the numerical simulator and real quantum hardware, via the IBM cloud, to accomplish data classification and quantum chemistry tasks. In the problems studied, numerical and experimental results show that QAS cannot only alleviate the influence of quantum noise and barren plateaus but also outperforms VQAs with pre-selected ansatze.

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INTRODUCTION

The variational quantum learning algorithms (VQAs)1,2, including quantum neural network3 and variational quantum eigensolvers (VQEs)6–9, are a class of promising candidates to use noisy intermediate-scale quantum (NISQ) devices to solve practical tasks that are beyond the reach of classical computers10. Recently, the effectiveness of VQAs toward small-scale learning problems such as low-dimensional synthetic data classification, image generation, and energy estimation for small molecules has been validated by experimental studies11–14. Despite the promising achievements, the performance of VQAs will degrade significantly when the qubit number and circuit depth become large, caused by the tradeoff between the expressivity and trainability15. More precisely, under the NISQ setting, involving more quantum resources (e.g., quantum gates) to implement the ansatz results in both a positive and negative aftermath. On the one hand, the expressivity of the ansatz, which determines whether the target concept will be covered by the represented hypothesis space, will be strengthened by increasing the number of trainable gates16–19. On the other hand, a deep circuit depth implies that the gradient information received by the classical optimizer is full of noise and the valid information is exponentially vanished, which may lead to divergent optimization or barren plateaus20–24. With this regard, it is of great importance to design an efficient approach to dynamically control the expressivity and trainability of VQAs to attain good performance.

Initial studies have developed two leading strategies to address the above issue. The first one is quantum error mitigation techniques. Representative methods to suppress the noise effect on NISQ machines are quasi-probability25,26, extrapolation27, quantum subspace expansion28, and data-driven methods29,30. In parallel to quantum error mitigation, another way is constructing ansatz with a variable structure. Compared with traditional VQAs with the fixed ansatz, this approach cannot only maintain a shallow depth to suppress noise and trainability issues, but also keep sufficient expressibility to contain the solution. Current literature generally adopts brute-force strategies to design such a variable ansatz31–33. This implies that the required computational overhead is considerable, since the candidates of possible ansatze scale exponentially with respect to the qubits count and the circuit depth. How to efficiently seek a near-optimal ansatz remains largely unknown.

In this study, we devise a quantum architecture search scheme (QAS) to effectively generate variable structure ansatze, which considerably improves the learning performance of VQAs. The advantage of QAS is ensured by unifying the noise inhibition and the enhancement of trainability for VQAs as a learning problem. In doing so, QAS does not request any ancillary quantum resource and its runtime is almost the same as conventional VQA-based algorithms. Moreover, QAS is compatible with all quantum platforms, e.g., optical, trapped-ion, and superconducting quantum machines, since it can actively adapt to physical restrictions and weighted noise of varied quantum gates. In addition, QAS can seamlessly integrate with other quantum error mitigation methods35–37 and solutions for resolving barren plateaus34,36. Celebrated by the universality and efficacy, QAS contributes to a broad class of VQAs on various quantum machines.

RESULTS

The mechanism of VQAs

Before moving on to present QAS, we first recap the mechanism of VQAs. Given an input \( X \) and an objective function \( \mathcal{L} \), VQA employs a gradient-based classical optimizer that continuously updates parameters in an ansatz (i.e., a parameterized quantum
enhances the trainability of one typical heuristic ansatz throughout the whole study, we focus on exploring how QASware—efficient ansatz11,13. Such an ansatz is supposed to obey a reasonable runtime and memory usage. Intuitively, weight sharing strategy in QAS refers to correlating parameters among different ansatzes via weight sharing strategy. Notably, the performance of VQAs heavily relies on the circuit $U(\theta)$ to find the optimal $\theta^*$, i.e.,

$$\theta^* = \arg\min_{\theta \in \mathcal{C}} L(\theta, Z),$$

(1)

where $\mathcal{C} \subseteq \mathbb{R}^d$ is a constraint set, and $\theta$ are adjustable parameters of quantum gates$^{15,18}$. For instance, when VQA is specified as an eigen-solver$^6$, $Z$ refers to a Hamiltonian and the objection function could be chosen as $L = \text{Tr}(Z|\psi(\theta)\rangle\langle\psi(\theta)|)$, where $|\psi(\theta)\rangle$ is the quantum state generated by $U(\theta)$. For compatibility, throughout the whole study, we focus on exploring how QAS enhances the trainability of one typical heuristic ansatz—hard-ware-efficient ansatz$^{11,13}$. Such an ansatz is supposed to obey a multi-layer layout,

$$U(\theta) = \prod_{l=1}^{L} U_l(\theta) \in SU(2^N),$$

(2)

where $U_l(\theta)$ consists of a sequence of parameterized single-qubit and two-qubit quantum gates, and $L$ denotes the layer number. Note that the arrangement of quantum gates in $U_l(\theta)$ is flexible, enabling VQAs to adequately use available quantum resources and to accord with any physical restriction. Remarkably, the achieved results can be effectively extended to other representative ansate.

The scheme of quantum architecture search

Let us formalize the noise inhibition and trainability enhancement for VQAs as a learning task. Denote the set $S$ as the ansatze pool that contains all possible ansatze (i.e., circuit architectures) to build $U(\theta)$ in Eq. (2). The size of $S$ is determined by the qubits count $N$, the maximum circuit depth $L$, and the number of allowed types of quantum gates $Q$, i.e., $|S| = O(Q^NL)$. Throughout the whole study, when no confusion occurs, we denote $a$ as the $a$th ansatz $U(\theta, a)$ in $S$. Notably, the performance of VQAs heavily relies on the employed ansatz selected from $S$. Suppose the quantum system noise, induced by $a$, is modeled by the quantum channel $\mathcal{E}_a$.

Taking into account of the circuit architecture information and the related noise, the objective of VQAs can be rewritten as

$$\arg\min_{\theta \in \mathcal{C}, a \in S} L(\theta, a, Z, \mathcal{E}_a).$$

(3)

The learning problem formulated in Eq. (3) forces the optimizer to output the best quantum circuit architecture $a$ by assessing both the effect of noise and the trainability. Notably, Eq. (3) is intractable via the two-stage optimization strategy that is broadly used in previous literature$^{31-33}$, i.e., individually optimizing all possible ansatze from scratch and then ranking them to obtain $(\theta^*, a^*)$. This is because the classical optimizer needs to store and update $O(dQ^NL)$ parameters, which forbids its applicability toward large-scale problems in terms of $N$ and $L$.

The proposed QAS belongs to the one-stage optimization strategy. Different from the two-stage optimization strategy that suffers from the computational bottleneck, this strategy ensures the efficiency of QAS. In particular, for the same number of iterations $T$, the memory cost of QAS is at most $T$ times more than that of conventional VQAs. Meanwhile, their runtime complexity is identical. The protocol of QAS is shown in Fig. 1. Two key elements of QAS are supernet and weight sharing strategy. Both of them contribute to locate a good estimation of $(\theta^*, a^*)$ within a reasonable runtime and memory usage. Intuitively, weight sharing strategy in QAS refers to correlating parameters among different ansatze. In this way, the parameter space, which amounts to the total number of trainable parameters required to be optimized in Eq. (3), can be effectively reduced. As for supernet, it plays two significant roles in QAS: (1) supernet serves as the ansatz indicator, which defines the ansatze pool $S$ (e.g., determined by the maximum circuit depth and the choices of quantum gates) to be searched and (2) supernet parameterizes each ansatz in $S$ via the specified weight sharing strategy. All possible single-qubit gates are highlighted by the brown rectangle. In Step 2, QAS optimizes the trainable parameters for all candidate ansatze. Given the specified learning task $L$, QAS iteratively samples an ansatz $\theta_i \in S$ from $\mathcal{A}$ and optimizes its trainable parameters to minimize $L$. $\mathcal{A}$ correlates parameters among different ansatze via weight sharing strategy. After $T$ iterations, QAS moves to Step 3 and exploits the trained parameters $\theta_i$ and the predefined $L$ to compare the performance among $K$ ansatze. The ansatz with the best performance is selected as the output, indicated by a red smiley face. Last, in Step 4, QAS utilizes the searched ansatze and the parameters $\theta_i$ to retrain the quantum solver with few iterations.

Fig. 1 Paradigm of the quantum architecture search scheme (QAS). In Step 1, QAS sets up supernet $\mathcal{A}$, which defines the ansatze pool $S$ to be searched and parameterizes each ansatz in $S$ via the specified weight sharing strategy. All possible single-qubit gates are highlighted by hexagons and two-qubit gates are highlighted by the brown rectangle. The unitary $U_l$ refers to the data encoding layer. In Step 2, QAS optimizes the trainable parameters for all candidate ansatze. Given the specified learning task $L$, QAS iteratively samples an ansatz $\theta_i \in S$ from $\mathcal{A}$ and optimizes its trainable parameters to minimize $L$. $\mathcal{A}$ correlates parameters among different ansatze via weight sharing strategy. After $T$ iterations, QAS moves to Step 3 and exploits the trained parameters $\theta_i$ and the predefined $L$ to compare the performance among $K$ ansatze. The ansatz with the best performance is selected as the output, indicated by a red smiley face. Last, in Step 4, QAS utilizes the searched ansatze and the parameters $\theta_i$ to retrain the quantum solver with few iterations.
1) Initialization: QAS employs a supernet $\mathcal{A}$ as an indicator for the ansatz pool $\mathcal{S}$. Concretely, the setup of the supernet $\mathcal{A}$ amounts to leveraging the indexing technique to track $\mathcal{S}$ using a linear memory cost. For instance, when $N = 4, L = 1$, and the choices of the quantum gates are $(R_y, R_y, R_y)$ with $Q = 3, \mathcal{A}$ indexes $R_y, R_y, R_y$ as $0^\circ, 1^\circ, 2^\circ$, respectively. With setting the range of $a, b, c, d$ as $(0, 1, 2)$, the index list $\{a^\circ, b^\circ, c^\circ, d^\circ\}$ tracks $\mathcal{S}$ e.g., $[0^\circ, 0^\circ, 0^\circ]$, $[0^\circ, 1^\circ, 0^\circ]$ describes the ansatz $\otimes_{i=1}^4 R_y(\theta_i)$ and $[2^\circ, 2^\circ, 2^\circ, 2^\circ]$ describes the ansatz $\mathbf{a_1} R_y(\beta_1)$. See Method for the construction of the ansatz pool $\mathcal{S}$ involving two-qubit gates. Meantime, as detailed below, $\mathcal{A}$ parameterizes all candidate ansatze via weight sharing strategy to reduce parameter space.

2) Optimization: QAS jointly optimizes $(\mathbf{a, \theta})$ in Eq. (3). Similar to conventional VQAs, QAS optimizes trainable parameters in an iterative manner. At the $t$th iteration, QAS uniformly samples an ansatz $\mathbf{a}^{(t)}$ from $\mathcal{S}$ (i.e., an index list indicated by $\mathcal{A}$). To minimize $\mathcal{L}$ in Eq. (3), the parameters attached to the ansatz $\mathbf{a}^{(t)}$ are updated to $\theta^{(t)} = \theta^{(t)} - \eta \partial \mathcal{L}(\mathbf{a}^{(t)}, \mathcal{S}, \mathcal{E}_\theta)/\partial \theta^{(t)}$, with $\eta$ being the learning rate. The total number of updating is set as $T$. Note that since the optimization of VQAs is NP-hard, empirical studies generally restrict $T$ to be less than $O(\text{poly}(\text{ON}))$ to obtain an estimation within a reasonable runtime cost.

To avoid the computational issue encountered by the two-stage optimization method, QAS leverages the weight sharing strategy developed in deep neural architecture search to parameterize ansatze in $\mathcal{S}$ via a specified correlation rule. Concretely, for any ansatz $\mathbf{a} \in \mathcal{S}$, if the layout of the single-qubit gates of the $\mathbf{a}$th layer between $\mathbf{a}$ and $\mathbf{a}^{(t)}$ is identical with $V \in [L]$, then $\mathcal{A}$ uses the training parameters $\theta^{(t)}$ assigned to $U(\mathbf{a}^{(t)}, \mathbf{a})$ to parameterize $U(\mathbf{a}, \mathbf{a}^{(t)})$, regardless of variations in the layout of other layers. We remark that the parameterization shown above is efficient, which can be accomplished by comparing the generated index list and the stored index lists. In addition, the above-correlated updating rule implies that the parameters of unsampled ansatze are never stored in classical memory. To this end, even though the size of the ansatz pool exponentially scales in terms of $N$ and $L$, QAS harnesses supernet and weight sharing strategy to guarantee its applicability toward large-scale problems.

3) Ranking: after $T$ iterations, QAS uniformly samples $K$ ansatze from $\mathcal{S}$ (i.e., $K$ index lists generated by $\mathcal{A}$), ranks their performance, and then assigns the ansatz with the best performance as the output to estimate $\mathbf{a}$. Mathematically, denoted $\mathcal{A}$ as the set collecting the sampled $K$ ansatze, the output ansatz is

$$\arg \min_{\mathbf{a} \in \mathcal{A}} \mathcal{L}(\mathbf{a}^{(T)}, \mathbf{a}, \mathcal{S}, \mathcal{E}_\theta).$$

(4)

In QAS, $K$ is a hyper-parameter to balance the tradeoff of efficiency and performance. To avoid the exponential runtime complexity of QAS, the setting of $K$ should polynomially scale with $N, L, Q$. Besides random sampling, other methods such as evolutionary algorithms can also be used to establish $K$ with better performance. See Supplementary D for details.

4) Fine-tuning: QAS employs the trained parameters $\theta^{(t)}$ to fine tune the output ansatz in Eq. (4).

We empirically observe fierce competition among different ansatze in $\mathcal{S}$ when optimizing QAS (see Supplementary B for details). Namely, suppose $\mathcal{S}$ can be decomposed into two subsets $\mathcal{S}_{\text{good}}$ and $\mathcal{S}_{\text{bad}}$, where the subset $\mathcal{S}_{\text{good}}$ (in $\mathcal{S}_{\text{bad}}$) collects ansatze in the sense that they all attain relatively good (bad) performance via independently training. For instance, in the classification task, the ansatz in $\mathcal{S}_{\text{good}}$ (in $\mathcal{S}_{\text{bad}}$) promises a classification accuracy above (below) 99%. However, when we apply QAS to accomplish the same classification task, some ansatze in $\mathcal{S}_{\text{bad}}$ may outperform certain ansatze in $\mathcal{S}_{\text{good}}$. This observation hints the hardness of optimizing correlated trainable parameters among all ansatze accurately, where the learning performance of a portion of ansatze in $\mathcal{S}_{\text{good}}$ is no better than training them independently.

To relieve fierce competition among ansatze in $\mathcal{S}$ and further boost performance of QAS, we slightly modify the initialization and optimization steps of QAS. Specifically, instead of exploiting a single supernet, QAS involves $W$ supernets to optimize the objective function in Eq. (3). The weight sharing strategy applied to $W$ supernets is independent of each other, where the parameters corresponding to $W$ supernets are separately initialized and updated. At the training and ranking stages, $W$ supernets separately utilize a weight sharing strategy to parameterize the sampled ansatz $\mathbf{a}^{(t)}$ to obtain $W$ values of $\mathcal{L}(\theta^{(m)}, \mathbf{a}^{(t)}, \mathcal{S}, \mathcal{E}_\theta)$, where $\theta^{(m)}$ refers to the parameters corresponding to the $m$th supernet. Then, the parameters applied to the ansatz $\mathbf{a}^{(t)}$ is categorized into the $w$th supernet when $w = \arg \min_{\theta^{(m)}} \mathcal{L}(\theta^{(m)}, \mathbf{a}^{(t)}, \mathcal{S}, \mathcal{E}_\theta)$.

We last emphasize how QAS enhances the learning performance of hardware-efficient ansatz $U(\theta)$ in Eq. (2). Recall that the central aim of QAS is to seek a good ansatz associated with optimized parameters to minimize $\mathcal{L}(\theta, \mathcal{A}, \mathcal{S}, \mathcal{E}_\theta)$ in Eq. (3). In other words, given $\mathcal{U} = \prod_{i=1}^L U_i(\theta)$, a good ansatz is located by dropping some unnecessary multi-qubit gates and substituting single-qubit gates in $U(\theta)$ for $\forall i \in [L]$. Following this routine, several studies have proved that removing multi-qubit gates to reduce the entanglement of the ansatz contributes to alleviate barren plateaus. In addition, a recent study unveiled that the choice of the quantum circuit architecture can significantly affect the expressive power of the ansatz and the learning performance.

Since the objective function of QAS implicitly evaluates the effect of different ansatze, our proposal can be employed as a powerful tool to enhance the learning performance of VQAs. Refer to Method for further explanation about the role of supernet, weight sharing, and analysis of the memory cost and runtime complexity of QAS.

Simulation and experimental results

The proposed QAS is universal and facilitates a wide range of VQA-based learning tasks, e.g., machine learning, quantum chemistry, and quantum information processing. In the following, we separately apply QAS to accomplish a classification task and a VQE task to confirm its capability toward the performance enhancement. All numerical simulations are implemented in Python in conjunction with the Pennylane and the Qiskit packages. Specifically, Pennylane is the backbone to implement QAS and Qiskit supports different types of noisy models. We defer the explanation of basic terminologies in machine learning and quantum chemistry in Appendices B and C.

Here we first apply QAS to achieve a binary classification task under both the noiseless and noisy scenarios. Denote $\mathcal{D}$ as the synthetic dataset, where its construction rule follows the proposal of the quantum kernel classifier. The dataset $\mathcal{D}$ contains $n = 100$ samples. For each example $(x^{(i)}, y^{(i)})$, the feature dimension of the input $x^{(i)}$ is 3 and the corresponding label $y^{(i)} \in \{0, 1\}$ is binary. Examples of $\mathcal{D}$ are shown in Fig. 2. At the data preprocessing stage, we split the dataset $\mathcal{D}$ into the training set $\mathcal{D}_{\text{tr}}$, validation set $\mathcal{D}_{\text{val}}$, and test set $\mathcal{D}_{\text{te}}$ with size $m_{\text{tr}} = 100, m_{\text{val}} = 100$, and $m_{\text{te}} = 100$. The explicit form of the objective function is

$$\mathcal{L} = \frac{1}{m_{\text{tr}}} \sum_{i=1}^{m_{\text{tr}}} \left(y^{(i)}(\mathcal{A}, x^{(i)}, \theta) - y^{(i)}(\theta)\right)^2,$$

(5)

where $(x^{(i)}, y^{(i)}) \in \mathcal{D}_{\text{tr}}$ and $y^{(i)}(\mathcal{A}, x^{(i)}, \theta) \in [0, 1]$ is the output of the quantum classifier (i.e., a function taking the input $x^{(i)}$, the supernet $\mathcal{A}$, and the trainable parameters $\theta$). The training (validation and test) accuracy is measured by $\frac{1}{m_{\text{tr}}} \sum_{i=1}^{m_{\text{tr}}} g(y^{(i)})$ with $g(y^{(i)})$ being the predicted label for $x^{(i)}$. We also apply the quantum kernel classifier proposed by to learn $\mathcal{D}$ and compare its performance with QAS, where the implementation of such a quantum classifier is shown...
Parameterized gates are along all supernets is set as two types of quantum gates. Specifically, at each layer $U(\theta)$, the parameterized gates are fixed to be the rotational quantum gate along $Y$-axis $R_y$. For the two-qubit gates, denoted the index of three qubits as $(0, 1, 2)$, QAS explores whether applying CNOT gates to the qubits pair $(0, 1), (0, 2), (1, 2)$ or not. Hence, the size of $S$ equals to $|S| = 8^3$. The number of sampled ansatz for ranking is set as $K = 500$. The setting $K \approx |S|$ enables us to understand how the number of supernets $W$, the number of epochs $T$, and the system noise affect the learning performance of different ansatze in the ranking stage.

Under the noiseless scenario, the performance of QAS with three different settings is exhibited in Fig. 2d. In particular, QAS with $W=1$ and $T=10$ attains the worst performance, where the validation accuracy for most ansatze concentrates on 50–60%, highlighted by the green bar. With increasing the number of epochs to $T=400$ and fixing $W=1$, the performance is slightly improved, i.e., the number of ansatze that achieves validation accuracy above 90% is 30, highlighted by the yellow bar. When $W=5$ and $T=400$, the performance of QAS is dramatically enhanced, where the validation accuracy of 115 ansatze is above 90%. The comparison between the first two settings indicates the correctness of utilizing QAS to accomplish VQA-based learning tasks in which QAS learns useful feature information and achieves better performance with respect to the increased epoch number $T$. The varied performance of the last two settings reflects the fierce competition phenomenon among ansatze and validates the feasibility to adopt $W>1$ to boost the performance of QAS. We retain the output ansatz of QAS under the setting: $W=5$ and $T=400$, both the training and test accuracies converge to 100% within 15 epochs, which is identical to the original quantum kernel classifier.

The performance of the original quantum kernel classifier is evidently degraded when the depolarizing error for the single-qubit and two-qubit gates is set as 0.05 and 0.2, respectively. As shown in the lower plot of Fig. 2f, the training and test accuracies of the original quantum kernel classifier drop to 50% (almost conduct a random guess) under the noisy setting. The degraded performance is caused by the large amount of accumulated noise, where the classical optimizer fails to receive the valid optimization information. By contrast, QAS can achieve good performance under the same noise setting. As shown in Fig. 2e, with setting $W=5$ and $T=400$, the validation accuracy of 115 ansatze is above 90% under the noisy setting. The ansatz that attains the highest validation accuracy is shown in Fig. 2c. Notably, compared with the original quantum kernel classifier in Fig. 2b, the searched ansatz contains fewer CNOT gates. This implies that, under the noisy setting formulated above, QAS suppresses the noise effect and improves the training performance by adopting a few CNOT gates. When we retrain the obtained ansatz with 10 epochs, both the train and test accuracies achieve 100%, as shown in the upper plot of Fig. 2f. These results indicate the feasibility to apply QAS to achieve noise inhibition and trainability enhancement.

We defer the omitted simulation results and the exploration of fierce competition to Supplementary B. In particular, we assess the learning performance of the quantum classifier with the hardware-efficient ansatz and the ansatz searched by QAS under the noise model extracted from the real quantum device, i.e., “ibmq_lima”. The achieved simulation result indicates that the ansatz obtained by QAS outperforms the conventional quantum classifier.

We next apply QAS to find the ground state energy of the Hydrogen molecule$^{13,50}$ under both the noiseless and noisy scenarios. The molecular hydrogen Hamiltonian is formulated as

$$H_h = g + \sum_{i=0}^{3} g_i Z_i + \sum_{i=1, i<k}^{3} g_{i,j} Z_i Z_k + g_{y,x} X_1 Y_3 + g_{x,y} Y_1 X_3 + g_{x,x} X_1 Y_1 X_2 ,$$

where $(X, Y, Z)$ denote the Pauli matrices acting on the $i$th qubit and the real scalars $g$ with or without subscripts are efficiently computable functions of the hydrogen–hydrogen bond length (see Supplementary C for details about $H_h$ and $g$). The ground state energy calculation amounts to computing the lowest energy eigenvalues of $H_h$, where the accurate value is $E_{\text{min}} = -1.136$ Ha$^{19}$. To tackle this task, the conventional VQE$^{10}$ and its variants$^{19,50}$ optimize the trainable parameters in $U(\theta)$ to prepare the ground state $|\psi\rangle = U(\theta) |0\rangle$ of $H_h$, i.e., $E_m = \langle \psi | H_h | \psi \rangle$. The implementation of $H_h$ in QAS is achieved by applying QAS to the original quantum kernel classifier under the noisy setting.
of $U(\theta)$ is illustrated in Fig. 3a. Under the noiseless setting, the estimated energy of VQE fast converges to the target result $E_m$ within 40 iterations, as shown in Fig. 3c.

The hyper-parameters of QAS to compute the lowest energy eigenvalues of $H_h$ are as follows. The number of supernets has two settings, i.e., $W = 1$ and $W = 5$, respectively. The layer number for all ansatze is $L = 3$. The number of iterations and sampled ansatze for ranking is $T = 500$ and $K = 500$, respectively. The search space of QAS for the single-qubit gates is fixed to be the rotational quantum gates along $Y$ and $Z$ axis. For the two-qubit gates, denoted by the index of four qubits as $(0, 1, 2, 3)$, QAS explores whether applying CNOT gates to the qubits pair $(0, 1), (1, 2), (2, 3)$ or not. Therefore, the total number of ansatze equals to $|S| = 128^3$. The performance of QAS with $W = 5$ is shown in Fig. 3d. Through retraining the obtained ansatz of QAS with 50 iterations, the estimated energy converges to $E_m$, which is the same as the conventional VQE.

The performance between the conventional VQE and QAS is largely distinct when the noisy model described in the classification task is deployed. Due to a large amount of gate noise, the estimated ground energy of the conventional VQE converges to $-0.4$ Ha, as shown in Fig. 3c. In contrast, the estimated ground energy of QAE with $W = 1$ and $W = 5$ achieve $-0.93$ and $-1.05$ Ha, respectively. Both of them are closer to the target result $E_m$ compared with the conventional VQE. Moreover, as shown in Fig. 3e, a larger $W$ implies a better performance of QAS, since the estimated energy of most ansatze is below $-0.6$ Ha when $W = 5$, while the estimated energy of 350 ansatze is above $0$ Ha when $W = 1$. We illustrate the generated ansatz of QAS with $W = 5$ in Fig. 3b. In particular, to mitigate the effect of gate noise, this generated ansatz does not contain any CNOT gate, which is applied to a very large noise level. Recall that a central challenge in quantum computational chemistry is whether NISQ devices can outperform classical methods already available. The achieved results in QAS can provide good guidance to answer this issue. Concretely, the searched ansatz in Fig. 3, which only produces the separable states that can be efficiently simulated by classical devices, suggests that QAS method may not outperform classical methods when NISQ devices contain large gate noise.

Note that more simulation results are deferred to Supplementary. Specifically, in Supplementary C, we exhibit more results of the above task. Furthermore, we implement VQE with the hardware-efficient ansatz and the ansatz searched by QAS on the real superconducting quantum hardware, i.e., “ibmq_ourense”, to estimate the ground state energy of $H_h$. Due to the runtime issue, we complete the optimization and ranking using the classical backend and perform the final runs on the IBMQ cloud. The experimental result indicates that the ansatz obtained by QAS outperforms the conventional VQE, where the estimated energy of the former is $-0.96$ Ha while the latter is $-0.61$ Ha. Then, in Supplementary D, we exhibit that utilizing the evolutionary algorithms to establish $\mathcal{K}$ can dramatically improve the performance of QAS. Subsequently, in Supplementary E, we provide numerical evidence that QAS can alleviate the influence of barren plateaus. Last, we present a variant of QAS to tackle large-scale problems with the enhanced performance in Supplementary F.

**DISCUSSION**

In this study, we devise QAS to dynamically and automatically design ansatz for VQAs. Both simulation and experimental results validate the effectiveness of QAS. Besides good performance, QAS only requests similar computational resources to conventional VQAs with fixed ansatz and is compatible with all quantum systems. Through incorporating QAS with other advanced error mitigation and trainability enhancement techniques, it is possible to seek more applications that can be realized on NISQ machines with potential advantages.

There are many critical questions remaining in the study of QAS. Our future work includes the following several directions. First, we will explore better strategies to sample ansatz at each iteration. For example, the reinforcement learning techniques, which are used to construct optimal sequences of unitaries to accomplish quantum simulation tasks, may contribute to this goal. Next, we will design a more advanced strategy to shrink the parameter space while not degrading the learning performance. Subsequently, to further boost the performance of QAS, we will leverage some prior information on the learning problem such as the...
The classical analog of the learning problem in Eq. (3) is the neural network architecture search. Recall that the success of deep learning is largely attributed to novel neural architectures for specific learning tasks, e.g., the convolutional neural networks for image processing tasks. However, deep neural networks designed by human experts are generally time-consuming and error-prone. To tackle this issue, the neural architecture search approach, i.e., the process of automating architecture engineering, has been widely explored, and achieved state-of-the-art performances in many learning tasks. Despite having a similar aim, naively generalizing classical results to the quantum scenario to accomplish Eq. (3) is infeasible due to the distinct basic components: neurons versus quantum gates, classical correlation versus entanglement, the barren plateau phenomenon, the quantum noise effect, and physical hardware restrictions. These differences and extra limitations further intensify the difficulty of searching the optimal quantum circuit architecture, compared with the classical setting. In the following, we explain the omitted implementation details of QAS demonstrated in the lower panel visualize the trainable parameters of 64 ansatze. The label $\theta$ refers to parameters assigned to the different ansatze. The parameter space reduces to a reasonable runtime complexity.

We remark that through adjusting the correlation criteria applied to the weight sharing strategy, the parameter space can be further reduced. For instance, when all parameters in an ansatze are correlated, the size of the parameter space reduces to $O(1)$. With this regard, another feasible correlation rule for QAS is unifying the single-qubit gates for all ansatze as $U_3 = R_z(a)R_z(b)R_y(\gamma)$. In other words, QAS only adjusts the arrangement of two-qubit gates to enhance the learning performance. From the practical perspective, this setting is reasonable since the gate error introduced by the single-qubit gates is much less than that of two-qubit gates.

Super-net

We next elucidate super-net used in QAS. As explained in the main text, super-net has two important roles, which are constructing the ansatze pool $S$ and parameterizing each ansatze in $S$ via the specified weight sharing strategy. In other words, super-net defines the search space, which subsumes all candidate ansatze, and the candidate ansatze in $S$ are evaluated through inheriting weights from the super-net. Rather than training numerous separate ansatze from scratch, QAS trains super-net just once (Step 2 in Fig. 1), which significantly cuts down the search cost.

We next explain how QAS leverages the indexing technique to construct $S$ when the available quantum gates include both single-qubit and two-qubit gates. Following notation in the main text, suppose that $N = S, L = 1, Q = 2$. The total number of ansatze is $Q^{2N} = 64$. The upper right panel illustrates how to use the indexing technique to accomplish the weight sharing. The label $a$ refers to the ansatze $a$. Namely, for any two ansatze, if the indexes in the $i$th layer are identical (highlighted by the blue and brown regions), then their trainable parameters in the $i$th layer are the same. The two heatmaps demonstrated in the lower panel visualize the trainable parameters of 64 ansatze. The label $\theta$ refers to the parameter assigned to the $i$th rotational quantum gate. Note that when the weight sharing strategy is applied, the trainable parameters are reused for different ansatze, as indicated by the dashed circles.

**METHODS**

**The classical analog of QAS**

The classical analog of the learning problem is Eq. (3) is the neural network architecture search. Recall that the success of deep learning is largely attributed to novel neural architectures for specific learning tasks, e.g., the convolutional neural networks for image processing tasks. However, deep neural networks designed by human experts are generally time-consuming and error-prone. To tackle this issue, the neural architecture search approach, i.e., the process of automating architecture engineering, has been widely explored, and achieved state-of-the-art performances in many learning tasks. Despite having a similar aim, naively generalizing classical results to the quantum scenario to accomplish Eq. (3) is infeasible due to the distinct basic components: neurons versus quantum gates, classical correlation versus entanglement, the barren plateau phenomenon, the quantum noise effect, and physical hardware restrictions. These differences and extra limitations further intensify the difficulty of searching the optimal quantum circuit architecture, compared with the classical setting. In the following, we explain the omitted implementation details of QAS.

**Weight sharing strategy**

The role of the weight sharing strategy is to reduce the parameter space to enhance the learning performance of QAS. The classical analog of the learning problem is Eq. (3) is the neural network architecture search. Recall that the success of deep learning is largely attributed to novel neural architectures for specific learning tasks, e.g., the convolutional neural networks for image processing tasks. However, deep neural networks designed by human experts are generally time-consuming and error-prone. To tackle this issue, the neural architecture search approach, i.e., the process of automating architecture engineering, has been widely explored, and achieved state-of-the-art performances in many learning tasks. Despite having a similar aim, naively generalizing classical results to the quantum scenario to accomplish Eq. (3) is infeasible due to the distinct basic components: neurons versus quantum gates, classical correlation versus entanglement, the barren plateau phenomenon, the quantum noise effect, and physical hardware restrictions. These differences and extra limitations further intensify the difficulty of searching the optimal quantum circuit architecture, compared with the classical setting. In the following, we explain the omitted implementation details of QAS.

**Optimizing the weight sharing strategy**

The upper left panel depicts the potential ansatze when $N = 3, L = 2$, and the choices of quantum gates are $\{R_x, R_y\}$ with $Q = 2$. The total number of ansatze is $Q^{2N} = 64$. The upper right panel illustrates how to use the indexing technique to accomplish the weight sharing. The label $\theta$ refers to the ansatze $a$. Namely, for any two ansatze, if the indexes in the $i$th layer are identical (highlighted by the blue and brown regions), then their trainable parameters in the $i$th layer are the same. The two heatmaps demonstrated in the lower panel visualize the trainable parameters of 64 ansatze. The label $\theta$ refers to the parameter assigned to the $i$th rotational quantum gate. Note that when the weight sharing strategy is applied, the trainable parameters are reused for different ansatze, as indicated by the dashed circles.

**Fig. 4** A visualization of weight sharing strategy. The upper left panel depicts the potential ansatze when $N = 3, L = 2$, and the choices of quantum gates are $\{R_x, R_y\}$ with $Q = 2$. The total number of ansatze is $Q^{2N} = 64$. The upper right panel illustrates how to use the indexing technique to accomplish the weight sharing. The label $\theta$ refers to the ansatze $a$. Namely, for any two ansatze, if the indexes in the $i$th layer are identical (highlighted by the blue and brown regions), then their trainable parameters in the $i$th layer are the same. The two heatmaps demonstrated in the lower panel visualize the trainable parameters of 64 ansatze. The label $\theta$ refers to the parameter assigned to the $i$th rotational quantum gate. Note that when the weight sharing strategy is applied, the trainable parameters are reused for different ansatze, as indicated by the dashed circles.

**Symmetric property and some post-processing strategies that remove redundant gates of the searched ansatze.** In addition, we will delve into theoretically understanding the fierce competition. In the end, it is intriguing to explore applications of QAS beyond VQAs such as optimal quantum control and the approximation of the target unitary using the limited quantum gates.
describes the ansatz \( Q \). In the last step, QAS samples with a required runtime is identical to conventional VQAs, which satisfies the memory cost in this step. To better understand how the computational complexity scales with the index space, which requests at most \( K_d \), the index list that describe the circuit architecture of \( Q \) is identical to the conventional VQAs with a fixed ansatz, which satisfies \( O(d) \). The total runtime complexity of QAS is hence \( O(T + T_{NL} + K_d) \).

To better understand how the computational complexity scales with \( N, L, \) and \( Q \), in the following, we set the total number of iterations in Step 2 and the number of sampled ansatz in Step 3 as \( T = O(QNL) \) and \( K = O(QNL) \), respectively. Note that since the size of \( S \) becomes indefinite, it is reasonable to set \( K \) to \( O(QNL) \) instead of a constant used in the numerical simulations. Under the above settings, we conclude that the runtime complexity and the memory cost of QAS are \( O(dQNL) \) and \( O(dQNL + QM^{L+L}) \), respectively.

We remark that when \( W \) supernets are involved, the required memory cost and runtime complexity of QAS linearly scales with respect to \( W \). Moreover, employing heterogeneous bandit learning techniques\(^{28}\) can exactly remove this overhead (see Supplementary A for details).

**DATA AVAILABILITY**

The datasets generated and/or analyzed during the current study are available from Y.D. on reasonable request.

**CODE AVAILABILITY**

The source code of QAS to reproduce all numerical experiments is available on the GitHub repository (https://github.com/yuxuan-du/Quantum_architecture_search/).

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Y. Du et al.
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**AUTHOR CONTRIBUTIONS**

Y.D. and D.T. conceived this work. Y.D., S.Y., and M.-H.H. accomplished the theoretical analysis. Y.D. and T.H. conducted numerical simulations. All authors reviewed and discussed the analysis and results, and contributed to writing the manuscript.

**COMPETING INTERESTS**

The authors declare no competing interests.

**ADDITIONAL INFORMATION**

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