

Red-QAOA: Efficient Variational Optimization through Circuit Reduction

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Abstract

The Quantum Approximate Optimization Algorithm (QAOA) addresses combinatorial optimization challenges by converting inputs to graphs. However, the optimal parameter searching process of QAOA is greatly affected by noise. Larger problems yield bigger graphs, requiring more qubits and making their outcomes highly noise-sensitive. This paper introduces Red-QAOA, leveraging energy landscape concentration via a simulated annealing-based graph reduction.

Red-QAOA creates a smaller (*distilled*) graph with nearly identical parameters to the original graph. The distilled graph produces a smaller quantum circuit and thus reduces noise impact. At the end of the optimization, Red-QAOA employs the parameters from the distilled graph on the original graph and continues the parameter search on the original graph. Red-QAOA outperforms state-of-the-art Graph Neural Network (GNN)-based pooling techniques on 3200 real-world problems. Red-QAOA reduced node and edge counts by 28% and 37%, respectively, with a mean square error of only 2%.

CCS Concepts: • **Computer systems organization** [→] **Quantum computing**; • **Mathematics of computing** → *Simulated annealing*; **Combinatorial optimization**.

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1 Introduction

Quantum computing, particularly with Noisy Intermediate-Scale Quantum (NISQ) computers, offers a powerful solution for tackling complex algorithms [1–9]. The Quantum Approximate Optimization Algorithm (QAOA), a widely recognized Variational Quantum Algorithm (VQA), addresses intricate optimization problems in graph theory, supply chain optimization, and machine learning [10–18]. To this end, QAOA treats inputs as graphs and maps nodes of the graph into qubits. However, modern NISQ machines, being inherently noisy, struggle to provide meaningful outcomes for large graphs as they use larger amounts of qubits [19, 20]. This leads to deep circuits that require a substantial number of qubits. Additionally, the execution of larger graphs tends to prolong processing times, reducing the overall throughput of the NISQ machine. This paper seeks to enhance the execution of larger graphs while ensuring meaningful outcomes.

Our paper examines QAOA applied to the MaxCut problem in graphs. These problems are NP-hard and are practically important. QAOA implementations harness the synergy between classical and NISQ computers. The classical computer furnishes quantum circuit parameters, while the NISQ computer maps the quantum circuit's graph nodes to qubits, executes it with the provided parameters, and generates optimization outcomes. The classical computer then updates circuit parameters based on these outcomes. This iterative feedback loop continues until the solution converges on optimal parameters [21]. The optimal parameters are then used to compute the Max-Cut of the graph. Ideally, regardless of the size of the input graph, the QAOA implementation should seamlessly operate on NISQ computers. However, practical NISQ machines encounter two critical concerns.

1. Noise-Induced Degradation: The approximation ratio, determining the closeness of QAOA outcomes to the ground truth, experiences significant degradation due to noise in the system. This is particularly high for larger problem instances. Figure 1 illustrates the QAOA convergence rate for 6-node and 10-node graphs. When an ideal optimizer is employed, the approximation ratio steadily increases for both graphs, surpassing 90%, indicating excellent performance. However,

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Figure 1. Comparing QAOA MaxCut approximation ratios (indicating outcome proximity to ground truth) between two graphs (6-node and 10-node) with noisy and ideal optimization. We perform 100 iterations for this comparison. The comparison illustrates (a) divergence from the ideal scenario as iterations increase (left) and (b) stagnating approximation ratios when scaling from 6-node to 10-node graphs (right).

with noisy optimization (NISQ), the QAOA optimization's approximation ratio is markedly affected as the number of iterations rises. Thus, the cumulative nature of noise influences the *usability* of NISQ computers [22]. Additionally, as the graph size scales from 6 nodes to 10 nodes, the required number of qubits also increases. As the impact of noise becomes severe at increased qubits, the approximation ratio for the 10-node graph remains stagnant at around 60%. In contrast, the smaller 6-node graph maintains a relatively higher ratio of approximately 80%. This highlights the heightened impact of noise for larger QAOA problem instances.

2. Distorted Solution Space: Noise distorts the underlying energy landscape [23–25]. It actively misguides the optimization process and results in suboptimal outcomes. Figure 2 compares the *ideal QAOA energy* landscape (left) with the noisy landscape executed on the 27-qubit *ibmq_kolkata* system (right) for a 13-node graph. Figure 2 highlights the substantial differences from noise-induced distortions.

Figure 2. A comparison of the ideal QAOA energy landscape (left) and the noisy landscape executed on the 27-qubit *ibmq_kolkata* system (right) for a 13-node graph. We can observe that noise-induced distortions result in significant differences between the two energy landscapes.

State-of-art approaches: One may substitute a large QAOA problem with an equivalent smaller one sharing the same optimal parameters. This leverages QAOA's unique property of similar instances having comparable energy landscapes [10, 26, 27]. Prior work has explored the practicality of transferring optimal parameters between graphs [26, 28, 29]. However, existing frameworks encounter scalability hurdles due to stringent preconditions, limiting transferability to small subgraphs. For larger graphs, parameters must be transferable across multiple subgraphs within the larger graph (donor graphs), which acceptor graphs can then utilize. These approaches have two limitations: 1) Identifying and transferring parameters for numerous subgraph combinations is computationally hard, and 2) strict mutual transfer conditions for large graphs restrict their practicality.

Our Proposal: We present Red-QAOA, a novel simulated annealing-based approach designed to overcome these limitations. Red-QAOA generates a reduced (distilled) graph for optimal parameter search. Once the parameters are determined using this *single distilled graph*, they are transferred back to the original graph. Subsequently, the optimization process proceeds using the original graph to converge on its precise optimal parameters. Red-QAOA ensures a less error-prone and more efficient optimization process. This is because using a distilled graph enables executing a smaller quantum circuit for a significant portion of the optimization.

This paper makes four key contributions:

- We propose Red-QAOA, a novel approach to search for optimal parameters with a reduced circuit for QAOA. Red-QAOA addresses noise challenges and outperforms state-of-the-art graph reduction methods.
- We comprehensively examine the theoretical definition of similar instances in QAOA. Subsequently, we integrate a dynamic simulated annealing-based graph reduction algorithm within Red-QAOA to enable a more generalized approach.
- We showcase the effectiveness of Red-QAOA by achieving a reduction of 28% nodes and 37% edges, along with a low Mean Squared Error (MSE) of 2% between ideal energy landscapes. These results are demonstrated across both real-world datasets and random graphs.
- We assess Red-QAOA on real quantum devices, demonstrating substantial improvements over noisy baselines. Moreover, Red-QAOA seamlessly integrates with existing optimization methods, enhancing the performance of quantum optimization algorithms.

2 Background

2.1 Basics of Quantum Computing

Quantum computing is a computational paradigm that leverages the principles of quantum mechanics to perform calculations. The fundamental building blocks of quantum computers are qubits, quantum gates, and quantum circuits.

2.1.1 Qubits. Unlike classical bits, which can take the value of either 0 or 1, a qubit can exist as a linear combination of its basis states, $|0\rangle$ and $|1\rangle$, as shown:

$$
|\psi\rangle = \alpha|0\rangle + \beta|1\rangle, \tag{1}
$$

where α and β are complex coefficients, subject to the normalization condition $|\alpha|^2 + |\beta|^2 = 1$.

2.1.2 Quantum Operations: Gates and Circuits. Quantum gates are unitary transformations used to manipulate the states of qubits. Common examples include the Pauli-X, Y, Z, Hadamard, and CNOT gates. For instance, the Pauli-X gate is represented by the following matrix:

$$
X = \begin{pmatrix} 0 & 1 \\ 1 & 0 \end{pmatrix}.
$$
 (2)

When the Pauli-X gate acts on a qubit $|\psi\rangle$, it flips the qubit's state such that $X|0\rangle = |1\rangle$ and $X|1\rangle = |0\rangle$.

A quantum circuit is a sequence of gates applied to a set of qubits [30, 31]. The circuit represents a computation that transforms an input state into an output state.

2.2 Variational Quantum Algorithms

Variational Quantum Algorithms (VQAs) are a class of quantum algorithms that aim to solve optimization problems using quantum computers [16–18, 32]. They harness quantum parallelism and interference to search for optimal solutions efficiently. VQAs use a parameterized quantum circuit to find the optimal solution. This is done by minimizing a cost function using classical optimization techniques.

2.2.1 Quantum Approximate Optimization Algorithm.

Quantum Approximate Optimization Algorithm (QAOA) [10] is a popular VQA. QAOA is designed to solve combinatorial optimization problems [33]. These problems involve finding the best arrangement or order of objects or variables given certain constraints or criteria. QAOA aims to maximize the expected value of the cost Hamiltonian H_c and the mixer Hamiltonian H_m with respect to a trial state $|\psi(\gamma, \beta)\rangle$ obtained from a parameterized quantum circuit. The trial state is produced by applying *p* alternating layers of unitary operators (called QAOA layers) as shown in Equation 3:

$$
U(\gamma, \beta) = e^{-i\beta_p H_m} e^{-i\gamma_p H_c} \cdots e^{-i\beta_1 H_m} e^{-i\gamma_1 H_c} |s\rangle
$$
 (3)

where $|s\rangle$ is the uniform superposition over computational basis states shown in Equation 4,

$$
|s\rangle = \frac{1}{\sqrt{2^n}} \sum_{z} |z\rangle \tag{4}
$$

where n is the number of qubits. For the Maxcut problem, the cost Hamiltonian H_c and the mixer Hamiltonian H_m can be defined as follows:

$$
H_c = \sum_{\langle i,j\rangle \in E} \frac{1}{2} (I - \sigma_i^z \sigma_j^z),\tag{5}
$$

$$
H_m = \sum_{i=1}^n \sigma_i^x,\tag{6}
$$

where σ_i^z and σ_i^x represent the Pauli-Z and Pauli-X operators acting on the i -th qubit, respectively, and E signifies the set of edges in the input graph. The Maxcut solutions are encoded in the eigenstates of the problem Hamiltonian, where each qubit denotes a vertex of the graph, with its state indicating the partition (0 or 1) to which the vertex belongs.

2.2.2 Graph Reduction for QAOA. QAOA operates on graphs, so reducing the problem size entails shrinking the underlying graph representation. This reduction effectively transforms the circuit reduction problem into a graph reduction problem. Graph reduction is a well-explored field with techniques that simplify graphs while preserving their structural or functional properties [34].

Graph neural network (GNN)-based graph pooling methods offer promising techniques for graph reduction while maintaining graph structures. Attention-based Spectral Aggregative (ASA) pooling [35] and Self-Attention Graph (SAG) pooling [36, 37] utilize attention mechanisms to learn reduced representations of graphs. ASA pooling aggregates node features across scales using attention coefficients and multiscale Laplacian eigenvectors. In contrast, SAG pooling learns soft hierarchical node clustering by computing importance scores for each node. Another approach, Top-K pooling [36, 38, 39], is a pooling method that selects the top-K nodes based on learned importance scores, constructing a smaller graph retaining the most relevant information.

3 Motivation: Equivalent Graph Instances

VQA executions involve two primary steps to solve the underlying optimization problem: parameter optimization and solution-finding. The first step aims to determine the optimal parameters of a parameterized quantum circuit to minimize a cost function. On the other hand, the second step, called solution-finding, seeks to find the optimal solution using the parameters obtained from the previous step.

Prior works have focused on finding equivalent graph instances for the solution-finding step. However, we argue that one can more efficiently optimize QAOA by finding equivalent instances for parameter optimization.

3.1 Parameter Optimization

The parameter optimization step is significantly more prone to errors and requires more computational resources. It entails employing classical optimization algorithms, which involve iteratively evaluating circuits with varying parameter values on a NISQ machine. Errors within qubits may accumulate during each circuit evaluation due to noise and imperfections inherent in the quantum (NISQ) hardware. These accumulated errors can potentially impede the accuracy of the parameter optimization process.

Moreover, the optimization process might entail navigating through a vast parameter space, increasing the likelihood of encountering local minima—suboptimal solutions that are not the global minimum. This situation arises when the optimization algorithm becomes trapped within a specific region of the parameter space, neglecting exploration of other regions that could potentially harbour superior solutions. Furthermore, the combination of complex quantum circuits with the iterative classical optimization process makes the parameter optimization step more prone to errors.

3.2 Challenge: Finding Similar Graphs

The definition of equivalent instances differs between the parameter optimization and solution-finding steps in VQA due to their distinct objectives. In the solution-finding step, instances are considered equivalent if their outcomes are *exactly* the same [40]. This poses challenges, especially for large graphs, as it requires converging on a few reduced graphs that yield identical outcomes.

In contrast, instances are considered equivalent in the parameter optimization step if they share *similar* (or nearlyidentical) optimal parameters that minimize the cost function [10, 26, 28]. This broader criterion allows for operating on significantly more reduced graphs in this step. Thus, the equivalence requirements in parameter optimization are more flexible than the solution-finding step. This relaxed criterion enables the selection of instances that may not be exactly identical but possess similar properties, enabling more efficient optimization of larger problem instances.

3.3 Observation: Common Energy Landscape

When addressing the Maxcut problem for a specific graph using QAOA, the operators in Equation 3 can be partitioned into sub-terms. Each sub-term corresponds to an edge within the graph. By commuting the terms, the resulting subterm operators involve only qubits j and k and any other qubits with a graph distance from *j* or *k* no greater than p layers ¹.

The expected value can be expressed as a sum of subexpected values, each calculated on a subgraph. The subterm operators are also identical when comparing two graphs with identical subgraphs. Consequently, the overall QAOA for both graphs is indistinguishable, allowing for the direct transfer of optimal parameters from one graph to the other. Not only are the optimal parameters transferable in such cases, but the entire energy landscapes of the two graphs also coincide. In the context of QAOA, an energy landscape refers to the distribution of energy values (or objective function values) over possible solutions or configurations.

Figure 3 presents two energy landscapes acquired using 7 and 10-node cycle graphs. A cycle graph comprises a single closed loop of nodes and edges, with each node connecting

Figure 3. QAOA energy landscapes for 7-node and 10-node cycle graphs. β and γ are in the ranges of [0, π] and [0, 2π], respectively. Since cycle graphs share the same sub-graphs, they exhibit nearly identical energy landscapes.

to two other nodes. As a result, irrespective of the number of nodes, they always share the same subgraphs, leading to almost identical energy landscapes in the two cases.

3.4 Idea: Leverage Energy Landscape Similarity

When searching for optimal parameters for QAOA, we leverage the observation that we can substitute one energy landscape with another if the two landscapes are nearly identical. This holds even if the landscape is from a reduced graph. Thus, our key goal is to find the reduced graph efficiently.

We assess the similarity of the two energy landscapes' shapes by calculating the mean square error (MSE) between their normalized versions. Normalizing the energy landscapes ensures their energy ranges are comparable. Equation 12 presents the MSE calculation, discussed in more detail in Section 5.1. A small value of MSE implies that the two landscapes have a similar shape, while a large value of MSE suggests a significant difference in shape. In Figure 3, the MSE between the normalized energy landscapes is 1.6×10^{-5} , indicating that they are nearly identical.

4 Our Proposal: Red-QAOA

We propose Red-QAOA, a framework that addresses the challenges of noise and execution overhead in QAOA. Red-QAOA seeks reduced graphs that maintain the energy landscape of the original graph for efficient quantum computation.

Figure 4 presents a high-level overview of the Red-QAOA design. The main components, depicted in the dashed block, include the **graph reduction step** and **optimal parameter** searching step-using the reduced graph (G'). Red-QAOA minimizes errors and improves accuracy by executing the reduced graph on a smaller quantum processing unit (QPU). After determining the *final optimal parameters*, Red-QAOA executes the original graph (G) on a larger QPU. This allows the application of error mitigation techniques on G , as G executes only for the *final optimal and accurate parameters*.

Our design consists of two main components: (1) First, we identify the key metric for selecting smaller equivalent

¹For a more in-depth discussion, we refer the reader to the seminal QAOA paper by Farhi et al. [10].

Figure 4. High-level overview of the design of Red-QAOA. The dashed block indicates the main component of Red-QAOA, which includes a graph reduction step and optimal parameter searching with the reduced graph (G') . The reduced graph can be executed on a smaller quantum processing unit (QPU) and is less prone to errors. For the final step, we find the solution with the optimal parameter and the original graph (G) . This step is more prone to error as we execute this on a larger QPU. However, since we only need to execute this only for final optimal and more accurate parameters, we can apply error mitigation to improve the overall execution accuracy.

instances, and (2) Second, we develop a graph reduction method based on this identified metric.

4.1 Theoretical Foundation of Red-QAOA

The energy corresponding to QAOA parameter values can be expressed as a sum of local energies as shown in Equation 7:

$$
E(\gamma, \beta) = \sum_{\langle jk \rangle} E_{\langle jk \rangle}(\gamma, \beta) \tag{7}
$$

where γ , β are the QAOA parameters, <jk> is an edge in the input graph, and E_{jk} has the form:

$$
E_{< jk>} = U^{\dagger}(C, \gamma_1)...U^{\dagger}(B, \beta_p)C_{< jk>}U(B, \beta_p)...U(C, \gamma_1)
$$
 (8)

The optimization process aims to find γ', β' such that $E(y', \beta')$ is minimized. This is done with gradient descent and is guided by the gradient ∇E , as shown in Equation 9:

$$
\nabla E = \sum_{< jk>} \nabla E_{< jk>} \tag{9}
$$

In prior works [28, 29], they identified that, if two energy functions, E_1 and E_2 have their gradient functions meet the criteria shown in Equation 10:

$$
\nabla E_1 = n * \nabla E_2 \tag{10}
$$

then, these two functions share the same optima. Thus, the optimal parameters can be transferred between them. However, the likelihood of two random QAOA instances meeting the criteria in Equation 10 is extremely rare.

Red-QAOA: The key insight in the design of Red-QAOA is that we can potentially approximate ∇E by only a subset of terms $E_{< jk>}$, if their gradient behaves similarly. That is:

$$
\nabla E \approx \sum_{\langle jk \rangle \in S} \nabla E_{\langle jk \rangle} \tag{11}
$$

for some set $S \subset$ all edges. This would allow Red-QAOA to reduce the number of qubits and quantum gates required by eliminating local energy terms while adequately approximating the optimization landscape.

However, analytically determining which $E_{\leq jk>}$ terms can be eliminated is intractable due to their complex unitary structure, as shown in Equation 8. The entanglement between unitaries makes it challenging to rigorously model the impact of removing individual $E_{\leq i k>}$ terms. Therefore, Red-QAOA relies on an empirical node degree heuristic to select the dominant $E_{\leq jk>}$ terms.

4.2 Identifying Equivalent Instances

In Section 3.3, we discussed that the overall expectation value of QAOA is the sum of individual sub-terms. Each sub-term corresponds to a subgraph encompassing all nodes and edges within a distance of p from the central edge. This formulation leads to a generalized subgraph matching problem, where finding another graph with comparable subgraphs can be challenging due to the exponential search space.

The creation of each subgraph involves several steps. First, we select a main edge connected to the problem we are trying to solve, called the problem Hamiltonian. We then add nearby nodes and edges to form the subgraph. This expansion continues until a certain distance, denoted as p , is reached from the main edge. The number of nodes and edges added at each step depends on the degrees of the nodes already present in the subgraph. If two graphs exhibit similar average numbers of connections per node, called Average Node Degrees (AND), they likely possess identical subgraphs.

To demonstrate this relation, we select 15 graphs at random and extract *all of their* unique non-isomorphic subgraphs. We perform a 1-layer QAOA run for each subgraph using a grid search with a width of 30, resulting in 900 sets of parameters. We then normalize the expectation values

Figure 5. Scatter plot illustrating the correlation between the mean squared error (MSE) and the difference in Average Node Degrees (AND) of the subgraph with its original graph, the subgraphs are selected from a collection of all unique non-isomorphic subgraphs for 15 graphs. A 6th-degree polynomial was found to be the best-fit curse – essentially indicating a correlation between MSE and AND.

and compute the mean squared error (MSE) between the subgraph results and its corresponding original graph. Figure 5 shows the resulting plot, where the y-axis shows MSE and the x-axis shows the AND ratio, the proportion of the subgraph's AND to that of the original graph. The plot reveals a significant correlation between the variables, suggesting that smaller graphs with AND values comparable to the original graph can effectively identify optimal parameters.

4.3 Mean Square Error for Landscape Similarity

A lower Mean Squared Error (MSE) implies higher similarity among instances during QAOA parameter optimization. However, defining an ideal MSE target remains challenging.

To establish an acceptable MSE, we analyze the optimal points' positions on energy landscapes. Figure 6 displays energy landscapes for six randomly selected graphs. One graph serves as a reference, and we calculate the MSE relative to this baseline for the remaining graphs, with optimal points denoted by blue stars. The normalized MSE can be viewed as a percentage error – 0.01 MSE corresponds to a 1% error.

Our observations suggest that when the MSE exceeds 2% (0.02), optimal point placement significantly deviates from the reference landscape. In practice, we identify an equivalent subgraph with an energy landscape deviating less than 0.02 from the original graph. In Figure 6, this 0.02 threshold equates to a minimum acceptable AND ratio of 0.7, and this ratio is used as the default value in the experiments. Users can adjust this threshold to suit their specific needs and use cases for equivalent instance searching.

To demonstrate the effectiveness of the MSE metric in quantifying differences among energy landscapes, we conducted a case study using random 15-node graphs and their

Figure 6. Six QAOA energy landscapes compared, with optimal points marked by blue stars. Mean Squared Error (MSE) values indicate similarity or divergence from the baseline landscape. A low MSE value, such as that of the 2nd graph in the first row (MSE = 0.01), indicates a landscape closer to the reference landscape. In contrast, a high MSE value, such as that of the graph in the bottom left ($MSE = 0.11$), indicates a landscape farther from the reference landscape. Our paper aims to identify an equivalent subgraph with an energy landscape that deviates less than 0.02 from the original graph.

subgraphs. Energy landscapes were generated for the original graph and its subgraphs using a 2-layer QAOA with 2048 random parameter sets. We computed the MSE between each subgraph's normalized landscape and that of the original graph. Additionally, we determined the average distance between their optimal solutions. Figure 7 illustrates a strong correlation between MSE and the distance between optimal solutions. This correlation confirms that MSE accurately captures disparities in optimal solutions, making it a suitable metric for comparing energy landscapes in QAOA.

4.4 Looking Beyond Pooling: Simulated Annealing

Simulated annealing (SA) is a stochastic optimization algorithm inspired by metallurgical annealing [41, 42]. SA begins

Figure 7. Scatter plot of MSE vs. average optimal point distance for random 15-node graphs and subgraphs, showing a strong correlation. This supports using MSE as a metric for energy landscape divergences.

with an initial solution and iteratively explores the solution space, accepting new solutions based on differences in objective function values and a decreasing 'temperature' parameter.

SA aligns well with our goal of identifying equivalent graphs. SA offers a more flexible and adaptive approach than graph pooling methods, which may impose rigid structures or rely on specific graph properties. This flexibility allows SA to discover subgraphs that better preserve the essential characteristics of the original graph, potentially making it a superior method. A key innovation of Red-QAOA is its utilization of SA to identify high-quality equivalent reduced graphs within the context of QAOA.

Red-QAOA utilizes an SA algorithm that supports 'constant' and 'adaptive' cooling schedules to dynamically construct reduced graphs and adjust the reduction ratio. The SA algorithm is executed multiple times. After each iteration, the average node degree (AND) of the resulting reduced graph is checked against the desired AND of the original graph. If the AND requirement is unmet, our algorithm adjusts the reduction ratio and reruns the SA algorithm until the desired AND is achieved. The resulting reduced graphs are then evaluated using the Mean Squared Error (MSE) metric to assess their similarity to the original graph.

Algorithm 1 shows the pseudocode of the proposed SA algorithm. The inputs to the algorithm include an input graph

 (G) , the desired subgraph size (k) , an initial temperature (T_0) , a cooling rate function or factor called α , a stopping temperature (T_f) , and a boolean flag called is *adaptive* that determines which cooling schedule to use. The algorithm starts by initializing a random subgraph as the initial solution and setting the initial temperature. Then, at each iteration, the algorithm explores a neighboring subgraph by replacing one of the nodes in the current subgraph with a node outside of it. The quality of the subgraph is measured using an objective function that calculates the difference between the ANDs of the subgraph and the original graph.

The algorithm uses a temperature-dependent probability function to accept or reject neighboring subgraphs. If a neighboring subgraph has a better objective function value, then, it is accepted as the new solution. Otherwise, if it has a worse or equal objective function value, it is accepted with a probability that decreases as the temperature decreases. This probability function allows the algorithm to escape local optima in the early stages of the search while converging towards the global optimum as the temperature drops. It then updates the current temperature either by a constant factor or adaptively based on the number of rejected subgraphs.

The adaptive cooling schedule is a crucial component of the algorithm, as it controls the *exploration*-*exploitation* tradeoff in the search process. By adjusting the cooling rate based on the current temperature, the algorithm can be fine-tuned to perform better on various graph pruning instances.

4.5 SA versus Graph Pooling Methods

We compare our SA-based methods to state-of-the-art graph neural network (GNN) pooling methods: Additive Self-Attention (ASA) [35], Set Attentional Aggregation (SAG) [36, 37], and Top-k Pooling (Top_k) [36, 38, 39]. These pooling methods use fixed reduction ratios and do not dynamically check if the reduced graph accurately approximates the original graph.

Figure 8. The mean square error (MSE) of subgraphs compared to the reduction ratio. We compare our Simulated Annealing (SA) methods to the state-of-the-art graph pooling techniques. The SA-based methods almost always provide lower MSE than prior techniques.

Figure 9. The performance of the proposed SA (simulated annealing) algorithm with varying node reduction ratios. The x-axis in each figure represents the mean square error, while the y-axis shows the percentage frequency. The algorithm, denoted as Red-QAOA, consistently identifies one of the most effective subgraphs across different reduction ratios. This suggests that Red-QAOA can consistently achieve desirable results for a given reduction ratio.

We test on the random graph dataset with $p = 3$, using a range of fixed reduction ratios from 0.1 to 0.7.

Figure 8 shows the experiment's results, including both 'constant' and 'adaptive' cooling versions. The results show that both these versions outperform all state-of-the-art GNNbased graph pooling methods, except for constant cooling when the reduction ratio is 0.7. However, this reduction ratio is too extreme and impractical in real-world applications. Overall, the adaptive cooling version of SA performs significantly better than the other methods. Given that adaptive cooling has a lower computational overhead, we equip Red-QAOA to employ the adaptive cooling method in all cases.

The superior performance of Red-QAOA compared to other GNN pooling methods highlights the importance of developing specialized graph reduction techniques tailored to the needs and constraints of QAOA and quantum computing.

4.6 Effectiveness of Simulated Annealing

To demonstrate the effectiveness of our proposed algorithm, we conducted an experiment using a random 15-node graph. We analyze its unique connected subgraphs for node reduction ratios of 0.67, 0.53, and 0.40, and for each subgraph, we perform a grid search with 900 data points and calculate the normalized MSE. Figure 9 shows the results as a histogram, with the x-axis representing MSE values and the y-axis indicating the frequency of subgraphs as a percentage. A dashed red line is marked to show the MSE obtained using the proposed SA algorithm. Across all reduction ratios, the SA algorithm consistently achieves the lowest MSEs.

5 Methodology

5.1 Figure of Merit

We utilize two key performance metrics - Mean Square Error (MSE) and Approximation Ratio - to evaluate Red-QAOA. The MSE is primarily used to measure the similarity between two QAOA instances, capturing discrepancies in the energy

landscapes. We apply MSE in two settings: ideal execution to compare the energy landscapes of the baseline graph and Red-QAOA graph, and noisy execution to compare both the noisy baseline and Red-QAOA landscapes against the ideal baseline. On the other hand, the Approximation Ratio is used to evaluate the quality of QAOA outcomes relative to the ground truth. This ratio assesses the performance of Red-QAOA against the baseline in ideal and noisy conditions.

5.1.1 Mean Square Error (MSE). Our study primarily uses the MSE to measure the similarity between two QAOA instances. MSE serves as a tool to quantify how closely two different QAOA-generated energy landscapes resemble each other. The MSE is defined by the equation:

$$
MSE = \frac{1}{N} \sum_{i=1}^{N} (E_i - \hat{E}_i)^2
$$
 (12)

where N is the total number of data points, the default is set to 1024 in our experiments. E_i represents the normalized energy at the *ith* data point in the first QAOA instance, and \hat{E}_i corresponds to the normalized energy at the same point in the second QAOA instance. This formula effectively captures the discrepancy between the two energy landscapes, providing a numerical measure of their similarity.

Our application of MSE occurs in two distinct settings. In the ideal execution setup, MSE is employed to compare the energy landscapes generated by the baseline graph and Red-QAOA's graph. The aim is to assess how closely the Red-QAOA graph's landscape mirrors the baseline under ideal conditions. In the noisy execution setup, on the other hand, two separate MSE values are computed. The first is between the noisy baseline landscape and the ideal baseline landscape. The second MSE is between the noisy Red-QAOA landscape and the ideal baseline landscape. In this context, the ideal baseline landscape serves as a benchmark, and our goal is to demonstrate that the Red-QAOA graph under noisy

conditions can produce a landscape more akin to this ideal baseline than what is achieved with the noisy baseline.

5.1.2 Approximation Ratio. In addition to MSE, we utilize the approximation ratio r to evaluate the quality of QAOA outcomes. This ratio is defined as the ratio of the optimal expectation value obtained by QAOA to the ground truth result, determined classically via brute force:

$$
r = \frac{\min_{\mathbf{y}, \boldsymbol{\beta}} \langle \psi(\mathbf{y}, \boldsymbol{\beta}) | C | \psi(\mathbf{y}, \boldsymbol{\beta}) \rangle}{C_{\text{ground truth}}} \tag{13}
$$

where $|\psi(\gamma, \beta)\rangle$ represents the trial state prepared by the QAOA operator given in Equation 3. Our experiments aim to compare the approximation ratios obtained from the baseline and Red-QAOA in both ideal and noisy execution setups. In the ideal execution setup, we seek approximation ratios for Red-QAOA that closely match the baseline. On the other hand, in the noisy execution setup, our objective is to assess the improvement brought by Red-QAOA in approximation ratio over the noisy baseline.

5.2 Benchmark Graph Datasets

To comprehensively evaluate Red-QAOA, we selected four diverse benchmark graph datasets: AIDS, Linux, IMDb, and a collection of Random graphs spanning different domains.

- AIDS: A set of 700 chemical compound graphs from the National Cancer Institute's repository. Each graph represents a chemical compound. Specifically, the nodes of the graph are the atoms and its edges are chemical bonds. The average graph size is eight nodes.
- **Linux:** Comprising 1,000 function call graphs extracted from the Linux kernel source code. Nodes represent functions, and edges denote function calls. The average graph size is ten nodes.
- **IMDb:** Consisting of 1,500 movie collaboration networks from the Internet Movie Database. Nodes represent actors, and edges indicate collaborations between them. The average graph size is six nodes, with most containing fewer than ten nodes.
- **Random graphs:** We generated ten random graphs using the NetworkX random graph generator, with node counts ranging from 7 to 20. These graphs provide a versatile testing platform for our experiments.

This diverse dataset selection allows us to evaluate our methods across various domains, offering insights into their scalability and adaptability to different graph structures. Table 1 summarizes the characteristics of benchmark graph datasets used in our experiments.

5.3 Circuit Simulation and Noise Modeling

We use the Qiskit [47] framework for circuit simulations. Ideal quantum circuit simulations are executed with the statevector backend, while noisy simulations are performed with the density matrix backend to account for potential noise

impact on quantum circuits. We employ the FakeToronto backend, which emulates the noise characteristics of the IBM Quantum Toronto device, incorporating gate errors, readout errors, and relaxation times. FakeToronto provides realistic quantum hardware conditions. Similar to prior work [48], we transpile circuits using the SABRE algorithm [49] to optimize depth and execution time. It selects the circuit with the shortest depth out of 100 repetitions. This process helps ensure optimal performance and resource usage.

5.4 Hardware Platform

Perlmutter's GPU nodes, each equipped with four Nvidia A100 GPUs (40GB VRAM per GPU), are used for efficient quantum circuit simulations, reducing the time required to obtain results. For executing quantum circuits on real hardware, we utilize the 27-qubit *ibmq_kolkata* backend from IBM and the larger 79-qubit *Aspen-M-3* from Rigetti. These backends allow us to validate our simulation results in realworld settings, providing insights into circuit performance under hardware constraints and error rates. By comparing the outcomes of our simulations with the results from the device backends, we can assess the accuracy and robustness of our methodology in the presence of hardware imperfections.

5.5 Graph Pooling Methods

We compared our proposed method, which uses dynamic checking to ensure accurate graph reduction, with three fixed-ratio graph pooling methods: Additive Self-Attention (ASA) [35], Set Attentional Aggregation (SAG) [36, 37], and Top-k Pooling (Top_k) [36, 38, 39]. These methods were chosen because they are widely used in the literature and have been shown to achieve state-of-the-art performance in various graph-related tasks [50].

All the graph pooling methods take the graph feature vector and a pooling ratio as inputs. In our case, the feature vector is generated from the input graph, which is a normalized vector that includes the node degrees, clustering coefficient, betweenness centrality, closeness centrality, and eigenvector centrality. These metrics provide insights into the node's connectivity, position within the network, and

Table 1. Description of Benchmark Graph Datasets

influence. The pooling ratio is determined dynamically based on the Red-QAOA reduced graph. We first generate a reduced graph using the Red-QAOA method. Then, we calculate the pooling ratio such that all pooled graphs are the same size as the Red-QAOA graph. By ensuring all reduced/pooled graphs are the same size, we enable a fair comparison between them.

5.6 Parameter Transfer

Previous studies on transferring optimal QAOA parameters focused on random regular graphs [29]. To evaluate parameter transferability on non-regular graphs, we start with random regular base graphs. We then randomly modify a small portion, 10% in our case, of the edges in these base graphs by removing some edges and adding new ones. This process makes the graphs slightly irregular while retaining similarities to the original regular base graphs. We generate two graphs to test parameter transfer: 1) a Red-QAOA graph, and 2) a smaller random regular graph with the same node degrees as the original unmodified base graph and the same number of nodes as the Red-QAOA graph. Comparing these two graphs allows us to evaluate how Red-QAOA performs on non-regular graphs compared to transferring parameters directly between similar non-regular graphs.

6 Results

6.1 Effectiveness of Recovering the Ideal Landscape

Our initial investigation assesses Red-QAOA's ability to restore the energy landscape distorted by noise. We compare the noisy MSE of the Red-QAOA landscape with the ideal QAOA landscape and contrast it with the noisy MSE between the baseline noisy QAOA landscape and the ideal QAOA landscape. Noisy MSE values evaluate energy landscape preservation by comparing noisy landscapes to ideal

Figure 10. Mean Squared Error (MSE) comparison of baseline and Red-QAOA noisy landscapes for graphs of 7-14 nodes. As the number of qubits increases, the noisy MSEs for both Red-QAOA and the baseline increase due to a higher device noise impact. Overall, with a more noise-tolerant circuit, Red-QAOA consistently performs better than the baseline.

Figure 11. Normalized energy landscapes of ideal, noisy baseline, and Red-QAOA for the best-case scenario of the 10 node graph. Blue stars indicate the globally optimal points on each landscape. Red-QAOA outperforms the noisy baseline by locating optimal points closer to the ideal scenario.

ones. Previously, we focused on ideal MSEs between different QAOA instances, capturing their equivalency. Therefore, the noisy and ideal MSEs should be interpreted separately.

We generate random graphs with 7 to 14 nodes and conduct noisy simulations. Figure 10 illustrates that Red-QAOA consistently outperforms the baseline noisy QAOA landscape in all scenarios. This improvement is primarily attributed to Red-QAOA' use of a reduced graph, reducing node counts by an average of 36% and edge counts by an average of 50%, significantly reducing the likelihood of noise interference.

Among the tested graphs, the 10-node graph exhibits the most substantial reduction in MSE from the baseline to Red-QAOA. Figure 11 displays the energy landscapes for the ideal, Red-QAOA, and noisy baseline, with blue stars marking the globally optimal points. The baseline noisy landscape is severely affected by noise, while the Red-QAOA energy landscape closely retains the ideal landscape's overall shape. Importantly, the location of optimal points is crucial, representing final solutions. The baseline's noise introduces false global points, whereas Red-QAOA maintains optimal points very close to the ideal landscape.

Figure 12. Normalized energy landscapes of ideal, noisy baseline, and Red-QAOA for the worst-case scenario of the 11-node graph. Blue stars indicate the globally optimal points on each landscape. The optimal points on Red-QAOA landscape begin to deviate from the ideal scenario; however, even with this deviation, Red-QAOA still outperforms the baseline, which exhibits a much greater deviation in terms of the location of optimal points from the ideal scenario.

In contrast, Figure 12 illustrates a worst-case scenario for 11 nodes where we observe the slightest reduction in MSE compared to the baseline. In this case, Red-QAOA landscape's overall shape is noticeably off from the ideal case. However, its globally optimal points are still very close to the true optimal, indicating the superior performance of Red-QAOA. Interestingly, the ideal landscapes of the 10-node and 11 node graphs are quite similar since they have very close node and edge counts. This implies that the reduced graph found by Red-QAOA for the 10-node case can be used for the 11-node case to achieve a better result. However, due to the constraints on searching for subgraphs, the reduced graph was rejected by Red-QAOA. This indicates a potential opportunity for further improvements.

Overall, with a smaller circuit, Red-QAOA is more robust to noise than the baseline approach and can perform better in practical quantum computing implementations.

6.2 Reductions and Ideal MSEs on Small Graphs

To thoroughly assess Red-QAOA performance, we selected real-world graphs with up to 10 nodes from Aids, Linux, and IMDb datasets for graph reduction and MSE experiments. We analyze larger IMDb graphs in Section 6.3 to evaluate Red-QAOA scalability and effectiveness. Figure 13 shows the three graph datasets' node and edge reduction ratios. Each dataset is represented by two bars, one for the node reduction ratio and one for the edge reduction ratio. On average, 28% of nodes were reduced, and 37% of edges were eliminated, resulting in substantially smaller graphs for execution.

Figure 14 displays the Mean Squared Error (MSE) for the three graph datasets, plotting results for different QAOA circuit layer parameter values: $p = 1$, $p = 2$, and $p = 3$. For each p value, we randomly selected 1024 parameter sets and computed the MSE relative to the baseline. Each dataset is represented in the figure by three bars corresponding to the distinct p values. The data indicates that as p increases, the MSE experiences a slight increase, suggesting that adding

Figure 13. Graph reductions, measured in terms of node and edge reduction ratios, for graphs from Aids, Linux, and IMDb graph datasets with up to 10 nodes. On average, Red-QAOA achieves 28% node reductions and 37% edge reductions.

Figure 14. Mean Squared Error (MSE) for Aids, Linux, and IMDb Graph Datasets with Up to 10 Nodes for $p = 1$, $p = 2$, and $p = 3$. The MSE achieved with the Aids and Linux datasets is below 0.01, while for IMDb, it is around 0.05. Section 6.3 presents more detailed information about this.

more layers to the QAOA circuit introduces complexity into the graph reduction process. However, this growth rate remains relatively low and manageable.

6.3 Scaling Up to Larger Graphs

Figure 13 and Figure 14 reveal a significant observation: the IMDb dataset has the lowest reduction ratios among the three datasets and its MSE values are notably higher. This difference primarily stems from IMDb's considerably higher average node degrees than AIDS and Linux datasets. Since our analysis focuses on graphs with a maximum of 10 nodes, removing a single node from IMDb's graphs results in a larger loss of edges. This discrepancy is evident in Figure 13, where the gap between edge and node reduction ratios is about 5% for AIDS and Linux but exceeds 10% for IMDb.

Figure 15 and Figure 16 show the results for IMDb graphs in two categories: IMDb small (up to 10 nodes) and IMDb

Figure 15. Node and edge reduction ratios for IMDb, classified into small (up to 10 nodes) and medium (up to 20 nodes) graph categories. When scaling from small to medium graphs, the node reduction ratio increased from 15% to 25% and the edge reduction ratio rose from 28% to 35%.

Figure 16. Mean Squared Error (MSE) for IMDb datasets for $p = 1$, $p = 2$, and $p = 3$. Graphs are divided into small (up to 10 nodes) and medium (up to 20 nodes) graphs. We observed a notable reduction in overall MSEs, dropping from approximately 0.05 to below 0.02.

medium (10 to 20 nodes). As graph size increases, the reduction ratio improves, and MSEs decrease. In this case, the IMDb medium graph set exhibits a similar performance as compared to Aids and Linux datasets. This suggests that Red-QAOA has a relatively worse performance for small, dense graphs. However, for these graphs, the noise impact and execution overhead are considerably smaller than in larger graphs, making it beyond the scope of Red-QAOA.

6.4 Scalability of Red-QAOA

6.4.1 Effectiveness. To assess the scalability of Red-QAOA, we conducted tests on 100 randomly generated graphs, each containing 30 nodes. We utilized the COBYLA classical optimizer with 20, 50, and 150 restarts for 1, 2, and 3 QAOA layers, respectively. Two key performance metrics were evaluated: the best result among all restarts for a given layer and the average result across restarts. Figure 17 demonstrates that Red-QAOA consistently achieved near-optimal best results,

Figure 17. Red-QAOA achieves near-optimal solutions for all testing cases when considering the best results achieved. On average, Red-QAOA achieves over 97% performance relative to baseline QAOA on a set of 100 large 30-node graphs.

exceeding 99% across all cases compared to the baseline. Remarkably, even with an average reduction of 30.7% in nodes and 44.3% in edges, Red-QAOA maintained over 97% of the baseline's average performance.

6.4.2 Runtime Analysis. Red-QAOA imposes minimal preprocessing overhead, scaling as $n \log n$ due to its binary search approach over graph sizes. Figure 18 confirms this asymptotically efficient complexity on random graphs ranging from 10 to 1000 nodes. For a small 10-node graph, Red-QAOA requires just 0.004 seconds of preprocessing time. In contrast, executing a corresponding 1-layer QAOA circuit on ibm_sherbrooke processor takes 4.2 seconds. Therefore, Red-QAOA has negligible overhead, around 0.1% of the total QAOA runtime, even for modest problem sizes. Theoretical $n \log n$ scaling suggests Red-QAOA will maintain efficient preprocessing as problem dimensions and quantum system sizes increase. This negligible overhead makes Red-QAOA well-suited for time-critical applications.

Figure 18. The runtime overhead of Red-QAOA scales as $O(n \log n)$, showing asymptotic efficiency. Using limited benchmark data from [51], we extrapolated the per-circuit execution time up to 65 qubits and compared Red-QAOA against it. Overall, Red-QAOA introduces a *negligible* graph processing overhead as compared to circuit execution time.

6.5 End to End Evaluation

To evaluate graph pooling methods for generating surrogate problems to train QAOA, we tested four techniques: Red-QAOA, SAG, Top-K, and ASAPooling. We generated 10-node random graphs and reduced versions using each method.

We performed grid searches to find the optimal QAOA parameters on the original and surrogate graphs. The key metric is the relative improvement in the approximation ratio over the original graphs under the noisy execution setup.

Figure 19 shows that Red-QAOA provides consistent positive improvements, with a 4.2% median increase over the baseline. This outperformed the other techniques, with SAG

and Top-K pooling showing high variability. ASAPooling consistently decreased performance.

Figure 19. Box plots of relative improvement in approximation ratio over noisy baselines for Red-QAOA and three GNN-based graph pooling techniques. Red-QAOA showed consistently positive improvements across all testing cases, outperforming the highly variable SAG and Top-K pooling. The ASAPooling performs the worst overall.

To evaluate the convergence rate improvement of QAOA using Red-QAOA, a 10-node random graph was selected, and classical optimization with five random restarts was performed using the COBYLA optimizer [52]. Figure 20 illustrates the convergence behaviour of noisy QAOA simulations on this problem instance using both the original graph and the Red-QAOA graph. To enable a direct comparison of convergence rates, the QAOA parameters were recorded at each iteration and then used to re-calculate the expected energy with an ideal noiseless simulator. Overall, Red-QAOA shows substantially faster and better convergence to high-energy solutions than the standard QAOA optimization.

Figure 20. Comparison of convergence between standard QAOA and Red-QAOA on noisy quantum simulations. Five restarts are performed using the baseline and reduced graphs from Red-QAOA. Red-QAOA demonstrates substantially faster and better convergence to optimal-energy solutions compared to the standard QAOA optimization.

6.6 Comparison: Parameter Transfer

Previous research [28] demonstrated QAOA parameter transferability on random *regular* graphs. Our experiments evaluated this transferability on various graphs up to 60 nodes,

Figure 21. MSE between ideal and transferred QAOA landscapes using Red-QAOA and parameter transfer, evaluated on real-world and non-regular graphs. Red-QAOA reliably outperforms transfer across graph types.

including real-world and non-regular graphs from AIDS, Linux, and IMDb datasets. We also tested modified regular and non-regular star/4-array graphs. We transferred optimal parameters between graphs with even/odd degree nodes for parameter transfer. To ensure fairness, we initially reduced the graph using Red-QAOA and then created a random regular graph with a similar node count and average degree. Figure 21 displays the MSE between ideal and transferred landscapes for both methods. Parameter transfer works well for regular or near-regular graphs but struggles with increased randomness. In contrast, Red-QAOA consistently maintains a low MSE across all graph types, demonstrating robust performance regardless of regularity.

6.7 Execution on Real Quantum Devices

IBM Device: We demonstrate Red-QAOA's effectiveness in addressing noise and hardware limitations on a real quantum device. We employ QAOA on the 27-qubit ibmq_kolkata backend with a 13-node random graph. Scaling to larger graphs is currently challenging due to the significant device

Figure 22. Normalized energy landscapes of an ideal scenario, a noisy baseline, and Red-QAOA for a random 13-node graph executed on the 27-qubit ibmq_kolkata backend are compared. The optimal points on the Red-QAOA energy landscape are much closer to the ideal case, whereas the baseline shows a significant deviation.

error rate. Though we have not extensively optimized the circuit beyond transpile the circuit using SABRE [49] multiple times and selecting the minimum-depth one. Further optimizations [53, 54] and error mitigation strategies [55] can potentially reduce noise impact for larger circuits. However, our primary aim is to showcase the improvement achieved with Red-QAOA over noisy baseline under the same execution conditions. This enhancement is expected to persist for larger instances and with more rigorous optimizations.

Figure 22 presents normalized energy landscapes for the ideal scenario, a noisy baseline, and Red-QAOA. This set of results showcases Red-QAOA's accuracy and reliability in identifying equivalent instances and reducing noise impact in QAOA parameter optimization. Red-QAOA outperforms the noisy baseline in identifying optimal points in the energy landscape, substantiating its effectiveness in addressing noise and hardware constraints during QAOA parameter optimization. This noise reduction leads to more accurate and reliable energy landscapes, resulting in more efficient solutions for large-scale optimization problems.

Figure 23. Red-QAOA outperforms noisy baseline QAOA on the 79-qubit Rigetti *Aspen-M-3 device* by achieving a lower MSE across all evaluated cases.

Rigetti Device: To further evaluate the performance of Red-QAOA on near-term quantum hardware, we employed it on the Rigetti Aspen-M-3 system with 79 qubits. Due to this device's higher error rates and limited access time, we benchmarked smaller graphs, 5 to 10 nodes, with a 1-layer QAOA. We compared the MSE between the ideal and noisy energy landscapes for Red-QAOA versus the baseline QAOA. As shown in Figure 23, Red-QAOA consistently achieved lower MSE across all evaluated cases on the Rigetti machine. By using a reduced QAOA circuit and obtaining enhanced performance, Red-QAOA showcases higher noise resilience even on today's noisy quantum devices.

6.8 Varying Noise Models

To demonstrate the noise tolerance of Red-QAOA, we conducted an experiment using a random 10-node test graph and 1-layer QAOA with 1024 parameter sets. We calculated the mean squared error (MSE) between the noise-free energy

Figure 24. ours demonstrates enhanced noise tolerance versus baseline QAOA across devices with a wide range of error rates. MSE between the noise-free and noisy energy landscapes is consistently lower for Red-QAOA.

landscape and landscapes generated under different noise models. The noise models were sampled from real IBM quantum device backends covering a wide range of error rates. As shown in Figure 24, Red-QAOA consistently achieves a lower MSE than the baseline across all noise levels. This included noise models from the Kolkata backend, which has one of the lowest error rates among the existing IBM devices, and the retired Toronto device with substantially higher errors. By using a smaller QAOA circuit, Red-QAOA is inherently more tolerant of all types of noise.

6.9 Increased Execution Throughput

Figure 25. Expected throughput improvement of Red-QAOA compared to the baseline on 27, 33, 65, and 127-qubit devices using aids, Linux, and IMDb benchmarks. We observed significant improvements: around 1.85x for aids, 2.1x for Linux, and 1.4x for IMDb in terms of system throughput.

Figure 25 shows the substantial throughput enhancement achieved with Red-QAOA, resulting in reduced execution time. We evaluated this improvement on large-scale quantum devices running multiple quantum circuits concurrently to optimize their utilization. The analyzed devices include Falcon 27-qubit, Eagle 33-qubit, Hummingbird 65-qubit, and Eagle 127-qubit. Notably, we observed throughput improvements of 1.85× for AIDS, 2.1× for Linux, and 1.4× for IMDb.

7 Related Work

7.1 Parameter Transfer

Previous research [26, 28, 29] has observed similar findings regarding input instances. However, their typical approach involves using a set of small, computationally feasible input graphs and optimized parameters as a reference. When encountering a new input, they search the pre-optimized dataset, find the most similar graph, and apply its parameters directly. This approach for transferring optimal parameters between graphs has two significant limitations. Firstly, identifying transferability for a vast set of subgraphs becomes impractical due to the exponential growth of possible subgraph combinations with increasing graph size. Secondly, achieving mutual transfer conditions for large graphs is challenging, as it necessitates meeting the condition for every subgraph pair, limiting the concept's applicability. Therefore, these limitations raise doubts about the effectiveness and feasibility of existing theoretical frameworks for optimal QAOA parameter transferability.

Previous studies have shown transferability between regular graphs with uniform degree distribution. The precondition is automatically satisfied for two regular graphs with the same degree of parity. However, real-world data typically consists of irregular graphs. In particular, only **1.14%** and **0%** of the graphs from the AIDS and LINUX datasets that are used in this study are regular graphs respectively, while about **54%** of the graphs in the IMDb dataset are regular.

7.2 Enhancing QAOA Performance

Various warm start techniques are proposed to enhance quantum optimization algorithms by initializing them with educated guesses, leading to faster convergence and better solutions. Jain et al. [56] proposed a graph neural networkbased warm start for QAOA that is effective on various combinatorial optimization problems. CAFQA[57] is a hybrid classical-quantum algorithm that improves solutions and convergence by finding parameters for a variational quantum algorithm. Egger et al. [58] introduced warm-starting quantum optimization using classical relaxations of optimization problems and showed effectiveness in portfolio optimization and MAXCUT problems. Other works [54, 59, 60] have also utilized domain-specific knowledge of QAOA to enhance execution fidelity, while some works [61, 62] proposed efficient Hamiltonian transitions and expectation calculations.

Our work focuses on a complementary approach to these warm start and domain-specific techniques. By combining our approach with these methods, we can improve the performance of quantum optimization algorithms even further.

7.3 Classical Optimization

Using reduced or surrogate models to expedite optimization and learning is a well-established concept in classical domains. It has found applications in hyperparameter tuning

for machine learning [63], approximating expensive blackbox functions [64], discovering suitable initializations in topology optimization [65], serving as substitutes for costly real-world experiments [66], and facilitating policy transfer from simpler to more complex environments in reinforcement learning [67]. Additionally, the challenge posed by saddle points in high-dimensional non-convex optimization has led to the development of saddle-free Newton approaches to optimize neural networks more efficiently [68]. While these reduced model methods expedite optimization and transfer in complex domains, Red-QAOA is specifically tailored to address the challenges of quantum tasks.

8 Conclusion

In the NISQ era, noise-resilient techniques are crucial, especially for variational algorithms like Quantum Approximate Optimization Algorithm (QAOA). This paper introduces Red-QAOA, a framework designed to mitigate noise effects on QAOA in NISQ devices. Red-QAOA utilizes a reduced graph for parameter optimization, enhancing resilience to errors and yielding superior results. Experimental findings demonstrate a substantial improvement in the noise-affected energy landscape, resulting in improved outcomes.

This overall improvement can be attributed to 28% reduction in node counts and 37% reduction in edge counts. We tested Red-QAOA on real quantum devices and showcased promising results. Our findings highlight the efficacy of our methodology in optimizing quantum circuits, offering valuable insights for future research in quantum computing applications. By integrating our method with complementary techniques, we can further boost the efficiency and accuracy of quantum optimization algorithms, paving the way for new advancements in quantum computing.

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A Artifact Appendix

A.1 Abstract

Red-QAOA introduces an innovative approach to optimizing the Quantum Approximate Optimization Algorithm (QAOA) by substituting the standard QAOA circuit with a smaller, less error-prone version. This reduced circuit maintains similar optimal circuit parameters while offering enhanced efficiency and reduced computational errors, making it more suitable for execution on quantum devices and classical simulations. Red-QAOA is implemented in Python, utilizing the Qiskit framework for quantum circuit operations and NetworkX for graph-related computations.

A.2 Description

- **How to access:** https://github.com/meng-ubc/Red-QAOA
- **Hardware Dependencies:** The experiments can be accelerated using NVIDIA GPUs, though this is not a requirement for basic execution.
- **Software Dependencies:** The following Python packages are essential for running the experiments:
	- **Qiskit:** Used for circuit simulation.
	- **Networkx:** Provides graph-related utilities.
	- **Scipy:** Employed for classical optimization tasks.
	- **– (Optional) torch-geometric:** This package offers Graph Neural Network (GNN) based graph pooling methods that are compared to Red-QAOA.
- **Data Sets:** The following graph data sets are included within the repository and are utilized in our experiments:
	- **– Linux:** A dataset representing Linux kernel interaction networks.
	- **– AIDS:** A dataset involving molecular structures related to AIDS research.
	- **– IMDb:** Contains data from the IMDb movie database.
- **Installation:** To install the necessary software for running the experiments, follow these steps:
	- 1. **Python Version:** The scripts and tools have been tested with Python 3.11.
- 2. **Required Packages:** A requirements.txt file is provided for installing required python packages.
- 3. **Optional GPU Support:** For systems with CUDA capabilities, the optional package 'qiskit-aer-gpu' can be installed separately to enable GPU acceleration. Note that the default 'qiskit-aer' package has to be uninstalled before installing 'qiskit-aer-gpu'.

A.3 Experiment workflow

Running Experiments: Each experiment is provided with a Python script. The script requires specific command-line arguments and can be executed as:

python experiment_script.py [--arguments]

Output and Analysis: The script outputs the numerical result that can be directly compared to the numbers reported in the paper.

A.4 Evaluation and expected results

Our study employs two key metrics: Mean Square Error (MSE) and Approximation Ratio.

- **Mean Square Error (MSE)**: Utilized to measure the differences between two QAOA landscapes, this metric is applied in:
	- 1. *Ideal Execution with MSE*: Assessing the MSE between ideal QAOA landscapes.
- 2. *Noisy Execution with MSE*: Comparing the MSE in noisy QAOA landscapes to evaluate the impact of noise.
- **Approximation Ratio**: This metric assesses the endto-end performance of Red-QAOA, with distinct objectives in ideal and noisy scenarios:
	- 1. *Ideal Execution with Approximation Ratio*: In ideal conditions, Red-QAOA aims to achieve results as close to the baseline QAOA performance.

In evaluating Red-QAOA, we have prepared a suite of experiments. Among these, three key experiments are crucial for understanding the efficacy and robustness of Red-QAOA, and they are:

A.4.1 MSE Analysis of Red-QAOA under Noisy Execution. This experiment reproduces Section 6.1 results in the paper.

Script and Arguments:

- **Script:** mse_noisy.py
- **Required Arguments:**
	- **–** -n: Specifies the number of nodes, ranging from 7 to 14, as used in the paper.
	- **–** –width: Sets the width of the landscape, defaulting to 32 (totalling 1024 executions).
	- **–** –shots: Defines the number of circuit executions, with a default of 8192.
	- **–** –use_gpu: An optional flag to utilize GPU computing, with CPU as the default.

Result Analysis: The script produces two MSE values:

- 1. The MSE between the noisy and ideal baseline landscapes.
- 2. The MSE between the noisy Red-QAOA and ideal baseline landscapes.

This analysis examines the relative difference between these MSE values. Due to the randomness in graph generation, absolute MSE values may vary; however, the focus should be on comparing the relative performance of Red-QAOA against the baselines under noisy conditions.

A.4.2 MSE Analysis of Red-QAOA in Ideal Conditions. This experiment reproduces results presented in Sections 6.2 and 6.3 in the paper.

Script and Arguments:

- **Script:** mse_ideal.py
- **Required Arguments:**
	- **–** –graph_set: Specifies the graph dataset; options include aids, Linux, and IMDb.
	- **–** –num_graphs: Defines the number of graphs for testing (it is recommended to test with at least ten graphs). **–** –p: Sets the number of QAOA layers.
- **Optional Arguments:**
	- **–** –num_points: The number of points sampled for the landscape, defaulting to 1024.
	- **–** –shots: The number of shots for circuit execution, with a default of 8192.
	- **–** –use_gpu: Indicates whether to use a GPU backend; the default is CPU.
	- **–** –min_nodes and –max_nodes: Specifies the range of nodes, defaulting to 0 to 10 for Section 6.2 and 10 to 20 for Section 6.3.

Interpreting Results: The output should include MSE values along with node and edge reductions, which can be directly compared with the figures and data presented in the paper.

A.4.3 End-to-End Performance Evaluation of Red-QAOA in Ideal Conditions. This experiment evaluates the end-to-end performance of the Red-QAOA under ideal, noise-free conditions, which is discussed in Section 6.4.1 in the paper.

Script and Arguments:

- **Script:** end_to_end.py
- **Required Argument: –** –p: Specifies the number of QAOA layers.
- **Optional Arguments:**
	- **–** –num_graphs: Sets the number of graphs for testing, with a default of 100.
	- **–** –num_nodes: Determines the number of nodes in each graph; the default is 30 to align with the paper's setting. A smaller value, like 10, is suggested for reduced computational overhead.
	- **–** –shots: Number of shots for circuit execution, defaulting to 8192.
	- **–** –use_gpu: Allows the option to use a GPU backend, with CPU as the default.

Interpreting Results: The script conducts multiple optimization restarts for each test case. It reports the average optimization result for Red-QAOA across all restarts, comparing these to the baseline and the optimal results. These values can be directly compared with those reported in the paper to validate Red-QAOA's performance in ideal scenarios.

A.4.4 Reproducing the Figures. The figures in the paper are generated using matplotlib package. The repository contains documentation (README_plot.md) to reproduce the corresponding figures in the paper.

A.5 Experiment Customization

The experiments in our paper, including variations like different numbers of QAOA layers, are designed with flexibility in mind. Key experiment parameters are set as required arguments in our scripts, ensuring consistency with the study's main findings. Additionally, a range of optional arguments is available, allowing for fine-tuning and more detailed, sensitive testing scenarios.

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