

Quantum Approximate Optimization Algorithm: Performance, Mechanism, and Implementation on Near-Term Devices

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The Quantum Approximate Optimization Algorithm (QAOA) is a hybrid quantum-classical variational algorithm designed to tackle combinatorial optimization problems. Despite its promise for near-term quantum applications, not much is currently understood about QAOA’s performance beyond its lowest-depth variant. An essential but missing ingredient for understanding and deploying QAOA is a constructive approach to carry out the outer-loop classical optimization. We provide an in-depth study of the performance of QAOA on MaxCut problems by developing an efficient parameter-optimization procedure and revealing its ability to exploit non-adiabatic operations. Building on observed patterns in optimal parameters, we propose heuristic strategies for initializing optimizations to find quasi-optimal p -level QAOA parameters in $O(\text{poly}(p))$ time, whereas the standard strategy of random initialization requires $2^{O(p)}$ optimization runs to achieve similar performance. We then benchmark QAOA and compare it with quantum annealing, especially on difficult instances where adiabatic quantum annealing fails due to small spectral gaps. The comparison reveals that QAOA can learn via optimization to utilize non-adiabatic mechanisms to circumvent the challenges associated with vanishing spectral gaps. Finally, we provide a realistic resource analysis on the experimental implementation of QAOA. When quantum fluctuations in measurements are accounted for, we illustrate that optimization will be important only for problem sizes beyond numerical simulations, but accessible on near-term devices. We propose a feasible implementation of large MaxCut problems with a few hundred vertices in a system of 2D neutral atoms, reaching the regime to challenge the best classical algorithms.

I. INTRODUCTION

As quantum computing technology develops, there is a growing interest in finding useful applications of near-term quantum machines [1]. In the near future, however, the number of reliable quantum operations will be limited by noise and decoherence. As such, hybrid quantum-classical algorithms [2–4] have been proposed to make the best of available quantum resources and integrate them with classical routines. The Quantum Approximate Optimization Algorithm (QAOA) [2] and the Variational Quantum Eigensolver [3] are such algorithms put forward to address classical combinatorial optimization and quantum chemistry problems, respectively. Proof-of-principle experiments running these algorithms have already been demonstrated in the lab [5–8].

In these hybrid algorithms, a quantum processor prepares a quantum state according to a set of variational parameters. Using measurement outputs, the parameters are then optimized by a classical computer and fed back to the quantum machine in a closed loop. In QAOA, the state is prepared by a p -level circuit specified by $2p$ variational parameters. Even at the lowest circuit depth ($p = 1$), QAOA has non-trivial provable performance guarantees [2, 9] and is not efficiently simulatable

by classical computers [10]. It is thus an appealing algorithm to explore quantum speedups on near-term quantum machines.

However, very little is known about QAOA beyond the lowest depth. While QAOA is known to monotonically improve with depth and succeed in the $p \rightarrow \infty$ limit [2], its performance when $1 < p < \infty$ is largely unexplored. In fact, it has been argued that one needs to go beyond low-depth QAOA in order to compete with the best classical algorithm for some problems on bounded-degree graphs [11, 12]. It thus remains a critical problem to assess QAOA at intermediate depths where one may hope for a quantum computational advantage. One major hurdle lies in the difficulty to efficiently optimize in the non-convex, high-dimensional parameter landscape. Without constructive approaches to perform the parameter optimization, any potential advantages of the hybrid algorithms could be lost [13].

In this work, we contribute, in three major aspects, to the understanding and applicability of QAOA on near-term devices, with a focus on MaxCut problems. First, we develop heuristic strategies to efficiently optimize QAOA variational parameters. These strategies are found, via extensive benchmarking, to be quasi-optimal in the sense that they usually produce known global optima. The standard approach with random initialization generically require $2^{O(p)}$ optimization runs to surpass our heuristics. Secondly, we benchmark the performance of QAOA and compare it with quantum annealing. On difficult graph instances where the minimum

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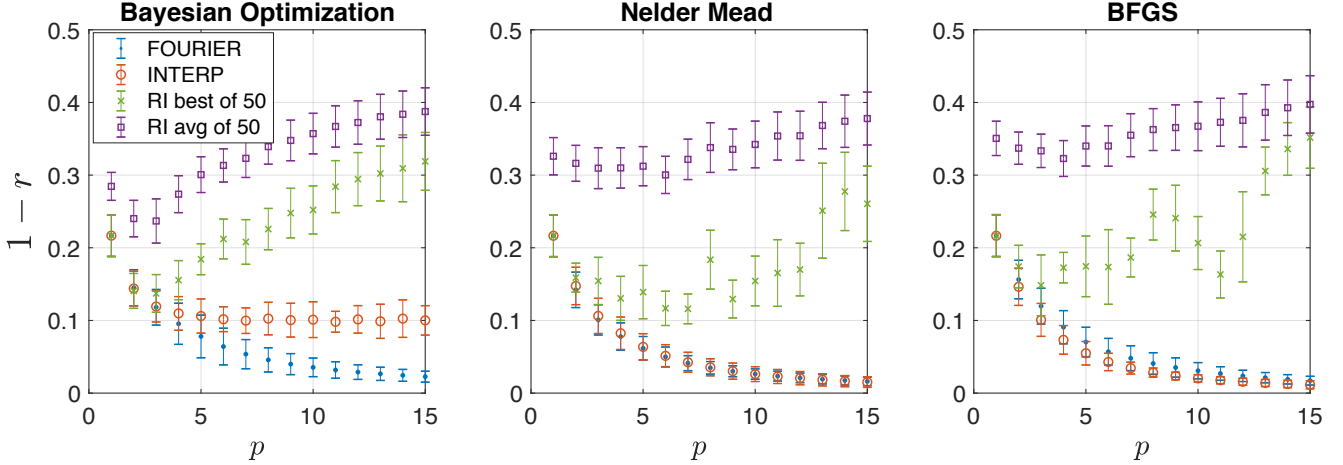


FIG. 14. Comparison of three different optimization routines applied to 10 instances of 14-vertex w3R graphs. The initial point of optimization is generated with either our heuristics (FOURIER or INTERP) or random initialization (RI). We plot the fractional error, $1 - r$, averaged over instances, for each optimization routine and each initialization strategy at each p . The error bars are sample standard deviations from the 10 instances. For our heuristic strategy, the optimization starts with a single initial point generated using our FOURIER[$\infty, 0$] or INTERP heuristics as described in Appendix B. For the RI strategy, we generate 50 random initial points uniformly in the parameter space, and optimize from each initial point; both the best and the average of 50 RI runs are plotted.

The first equality can be derived from Eq. (D1). This produces the standard adiabatic condition $T = O(1/\Delta_{\min}^2)$. As we discussed in section VB, the minimum gap for some graphs can be exceedingly small, so the adiabatic limit is not practical. However, it may be possible to choose an appropriate run time T , which breaks adiabaticity, but is long enough such that only few excited states are effectively involved in the dynamics. This is the regime where the diabatic bump operates and one can understand the dynamics by truncating Eq. (D4) to the first few basis states.

As an example, we plot in Fig. 13(a) the instantaneous eigenstate populations of the first few states. It is simulated with the full Hilbert space, but effectively the same dynamics will be generated if the simulation is restricted to the first few basis states in Eq. (D4). Fig. 13(b) shows the strength of the couplings between the instantaneous ground state and the low excited states. By comparing Fig. 13(a) and Fig. 13(b), one can see that $T = T^* = 40$ allows the time evolution to break the adiabatic condition before the anticrossing: population leaks to the first excited state, which becomes the ground state after the anticrossing. Thus, the time scale of T^* for the diabatic bump represents a delicate balance between allowing population to leak out of the ground state and suppressing excessive population leakage, which explains why it happens at a certain range of time scale.

Appendix E: Comparing different classical optimization routines

In this appendix, we compare three different classical routines that can be used to optimize QAOA parameters: Bayesian Optimization [37], Nelder-Mead [36], and BFGS [33]. This comparison is done by a numerical experiment where we apply these optimization routines to 10 instances of 14-vertex weighted 3-regular (w3R) graphs. To compare them on equal footing, we terminate each optimization run after a budget of $20p$ objective function evaluations is used. In the gradient-based routine, BFGS, we include the cost of gradient estimation via the finite-difference method into the budget of $20p$ objective function evaluations. For each routine, we start at $p = 1$ and gradually increment p , and perform optimization where the initial point is generated using either our heuristic strategies (FOURIER and INTERP) or the standard strategy of random initialization (RI). We use the versions of these optimization routines implemented in MATLAB R2017b as `bayesopt`, `fminsearch`, and `fminunc`, respectively. The objective function at each set of parameter is evaluated to floating point precision. The tolerance in both objective function value and step size in parameter space, as well as the finite-difference-gradient step size, are chosen to be 0.01.

The result of our numerical experiment is plotted in Fig. 14. Similar to Fig. 3(a), we see that the average quality of local optimum found from 50 RI runs is much worse than the best, indicating the difficulty of optimizing in the QAOA parameter landscape without a good initial point. We also see, regardless of the classical routines chosen, one run of optimization from an initial point

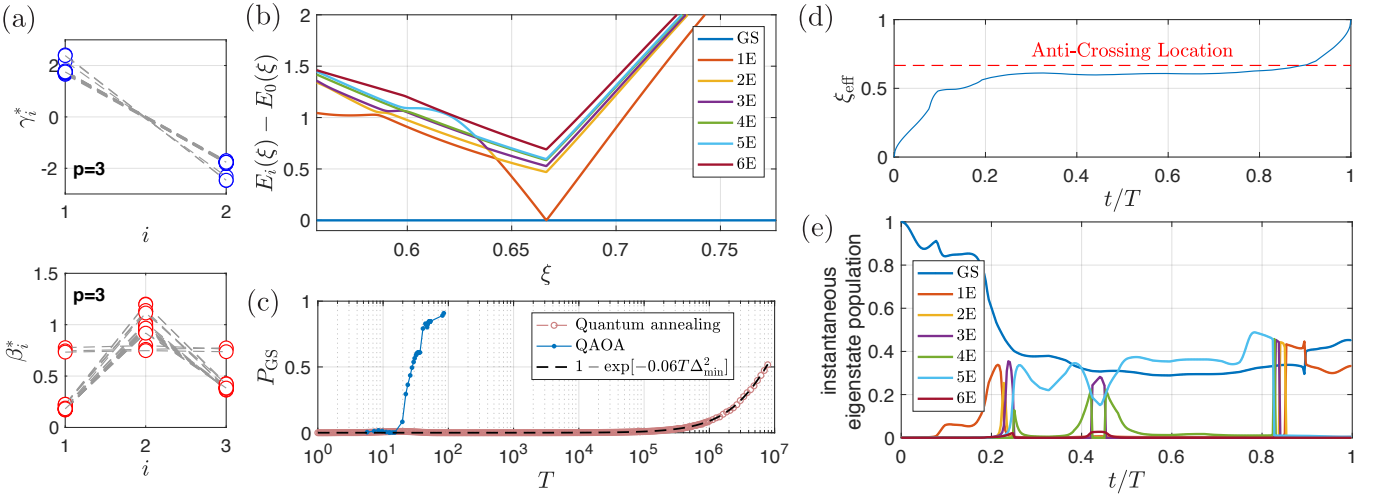


FIG. 15. (a) Pattern in the optimal QAOA parameters at level $p = 3$ for 20 random instances of Maximum Independent Set (MIS). Each dashed line connects parameters for one particular graph instance. (b) The energy difference relative to the ground state in the quantum annealing protocol of Eq. (H6) for an example 32-vertex MIS instance. The annealing protocol progresses as the parameter ξ increases from 0 to 1. The minimum spectral gap between the ground state (GS) and the first excited state (1E) is $\Delta_{\min} = 0.0012$ at $\xi = 0.6666$. (c) Comparing performance of quantum annealing and QAOA on the example instance, in terms the ground state population at the end of the quantum evolution. The equivalent evolution time for QAOA is calculated via $T_{\text{QAOA}} = \sum_{i=1}^{p-1} |\gamma_i| + \sum_{i=1}^p |\beta_i|$. (d) The effective annealing schedule converted from optimized 25-level QAOA parameters for the example MIS instance. (e) The population of the system in the instantaneous eigenstates, during the effective annealing schedule that approximates the dynamics under 25-level QAOA. Here, we observe that the algorithm attempts to transport the system to the fifth excited state, keeping it there before it undergoes a series of anti-crossings towards the ground state.

We further analyze the performance and mechanism of QAOA for MIS by focusing on example instances that are difficult for adiabatic quantum annealing due to small spectral gaps. In Fig. 15(b), we show the level-crossing structure for such an example instance, where the minimum spectral gap is $\Delta_{\min} = 0.0012$. The same instance is studied in Ref. [45]. To study the performance of QAOA in deeper-depth circuits, we use the interpolation-based heuristic strategy outlined in Appendix B 1 to optimize QAOA parameters for this example instance starting at level $p = 3$. The performance of QAOA and quantum annealing are then compared in Fig. 15(c), where we see that QAOA is able to obtain a much larger ground state population in much shorter time compared to the adiabatic time scale of $1/\Delta_{\min}^2 \approx 10^6$. We then study the mechanism of QAOA by converting its parameters at level $p = 25$ to a smooth annealing path $(f_P^{\text{QAOA}}, f_Q^{\text{QAOA}})$ in a similar

procedure as in Sec. V B. This annealing path is visualized in Fig. 15(d), where we plot the effective $\xi_{\text{eff}}(t)$ defined by $f_P^c(\xi_{\text{eff}}(t))/f_Q^c(\xi_{\text{eff}}(t)) = f_P^{\text{QAOA}}(t)/f_Q^{\text{QAOA}}(t)$ for the QAOA-like schedule. We then monitor populations in the instantaneous eigenstates during the evolution to gain insights into the mechanism of QAOA. As shown in Fig. 15(e), QAOA is able to learn to navigate a very complicated level-crossing structure by a combination of adiabatic and non-adiabatic operations: the system diabatically couples to the excited states, lingers to maximize population in the fifth excited state, and then exploits a series of anti-crossings to return to the ground state. Our results here demonstrate that the non-adiabatic mechanisms observed in QAOA for MaxCut can play a significant role in more general problems, such as difficult MIS instances.