XY mixers: Analytical and numerical results for the quantum alternating operator ansatz

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The quantum alternating operator ansatz (QAOA) is a promising gate-model metaheuristic for combinatorial optimization. Applying the algorithm to problems with constraints presents an implementation challenge for near-term quantum resources. This paper explores strategies for enforcing hard constraints by using XY Hamiltonians as mixing operators (mixers). Despite the complexity of simulating the XY model, we demonstrate that, for an integer variable admitting κ discrete values represented through one-hot encoding, certain classes of the mixer Hamiltonian can be implemented without Trotter error in depth $O(\kappa)$. We also specify general strategies for implementing QAOA circuits on all-to-all connected hardware graphs and linearly connected hardware graphs inspired by fermionic simulation techniques. Performance is validated on graph-coloring problems that are known to be challenging for a given classical algorithm. The general strategy of using XY mixers is borne out numerically, demonstrating a significant improvement over the general X mixer, and moreover the generalized W state yields better performance than easier-to-generate classical initial states when XY mixers are used.

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I. INTRODUCTION

Prior to achieving full error correction, which likely requires large physical-to-logical qubit ratios and low error rates, the exploration of what near-term quantum resources can achieve is paramount. One of the main uses of nearterm quantum devices will be to evaluate quantum algorithms beyond the reach of classical simulation. One of the most exciting and anticipated potential uses of quantum computers is solving combinatorial optimization problems, with near-term quantum hardware providing unprecedented means for exploring and evaluating quantum algorithms for optimization. The quantum-approximation-optimization algorithm (QAOA) has risen to be the leading candidate to test the applicability of gate-model quantum resources at solving optimization problems on near-term quantum hardware prior to fault tolerance [1–3]. Studies using QAOA to obtain the $\Theta(\sqrt{2^n})$ query complexity on Grover's problem [4], to find approximate solutions to MAXCUT [1,5,6], MAXE3LIN2 [7], network detection [8], simple machine learning models [9,10], and sampling [11],

suggest that there is a path forward to obtaining high-quality solutions with QAOA under a noiseless environment. The hybrid nature of this algorithm implies that noise of physical qubits can be tolerated to some extent [3,6,12].

These initial findings led to the development of a general framework known as the quantum alternating operator ansatz (also QAOA) that extends the utility of the initial algorithm to a wide variety of optimization problems involving linear or nonlinear constraints and to a wider variety of mixing operators that can greatly increase the implementability of a QAOA approach to many combinatorial optimization problems [2]. Both frameworks are metaheuristics and so require further specification. Challenges include devising strategies for selecting angles with minimal computational overhead, efficient initial-state determination and preparation, and embedding high-dimensional graphs, e.g., nonplanar graphs, into physically realizable lattices of qubits. Prior work on components of the general QAOA algorithm for handling hard and soft optimization constraints includes lattice protein folding by changing the driver [13], classical and quantum embeddings for representing all-to-all connected graph problems [14], optimization strategies [6], and compilation strategies [15–19] and furthers our understanding of how to apply the QAOA heuristic.

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In this paper we explore the feasibility of expanding QAOA's scope of applicability to discrete optimization problems with integer variables, as conceptually proposed in Ref. [2]. Commonly, κ -ary variables facilitate simpler representations of combinatorial optimization problems and open the possibility of multiple encoding strategies. For example, integer variables can be directly encoded into a binary, redundantly encoded in a classical coding fashion, or encoded into a one-hot-encoded set if κ is small. In this paper we study the implementation and performance of one-hot encodings for graph-coloring problems with QAOA using mixers based on the XY Hamiltonian. Pairing the one-hot-encoding and XY mixers is a natural choice as XY mixers preserve the representation [2]. To validate the XY-mixing Hamiltonian consistently with the salient feature of QAOA—short-depth circuits—we provide short-depth circuit implementations for each term in QAOA. For any one-hot-encoded integer variable problem, the feasible subspace is spanned by all Hammingweight-1 bit strings. Given such a problem on an all-to-all connected hardware platform, we propose a scheme that can generate the exact evolution of the XY model on a complete graph in linear depth. Moreover, exploiting the fermionic transformation, we show that the XY model on a ring can be realized in logarithmic depth. Most notably, due to the commuting nature of the cost Hamiltonians [20] a SWAP network, akin to sorting networks, can be used to implement any two-local cost operator requiring all-to-all connectivity in linear depth on a linearly connected graph of qubits. Though the XY mixer is significantly more complicated than the standard X mixer, we demonstrate that under numerous scenarios this driver term can be implemented in linear depth by taking a fermionic perspective. If approximate evolution is found to be tolerable, for all-to-all connected architectures, the first-order Trotter implementation of the XY mixer drops to $O(\log(\kappa))$ circuit depth.

Through numerical simulations, we also compare performance of different XY mixers. In a noise-free scenario, the mixer based on the XY model on a complete graph K_{κ} for each node gives better performance than the mixer using the XY model on a ring for finite QAOA levels. This advantage needs to be considered as a tradeoff to the complexity of the circuit generating the mixing unitary; furthermore, the realistic performance will also depend on the effects of noise and gate infidelity. We also show that initial states play a crucial rule in QAOA with XY mixers. While an easy-to-generate classical state serves as a valid initial state in the feasible subspace, the generalized W state, i.e., the uniform superposition of all Hamming-weight-1 bit strings, yields significantly better performance.

The rest of the paper is structured as follows. Section II outlines the general QAOA framework, with the emphasis on an analysis showing that the approximation ratio for optimization problems in a discrete bounded domain can provide a lower bound on the typical case. Section III formulates the max- κ -colorable-subgraph problem in a binary representation and introduces the terminology required for comparing the XY and X mixers. Section IV describes methods for implementing various mixers in short-depth circuits. In Sec. V A, the XY mixer is demonstrated to outperform the X mixer, and in Secs. V B and V C we provide benchmark numerics on small

hard-to-color graphs and all κ -colorable graphs of given sizes. Circuit implementation strategies for W-state generation are relegated to the Appendices.

II. QAOA FRAMEWORK

The QAOA framework starts with the specification of a cost Hamiltonian (phase-separating Hamiltonian) H_{PS} such that its specification requires a polynomial number of k-local terms that all commute. Commonly, the z-computational basis states are used for problem encoding. That way, every basis state corresponds to an eigenstate of H_{PS} . The objective is to to find the lowest-energy eigenstate by a quantum evolution that effects transition between the eigenstates. The H_{PS} term serves to interfere various eigenstates and thus change the transition probabilities.

A QAOA circuit of level p consists of the following steps.

- (1) Prepare a suitable initial state $|\psi_0\rangle$.
- (2) Repeat the following steps p times: in the lth repetition, apply the phase-separating unitary $\exp[-i\gamma_l H_{PS}]$ and apply the mixing unitary $U_M(\beta_l)$.
 - (3) Measure in the computational basis.

The unitaries are parametrized by a set of real numbers $\{\gamma_j, \beta_j\}_{j=1}^p$, respectively. In a classical-quantum hybrid setup, Monte Carlo averaging for the expectation value of H_{PS} serves as the objective function for classical feedback on the angles $\{\gamma_j, \beta_j\}_{j=1}^p$. Efficient strategies for statistical estimation of $\langle \psi(\gamma, \beta) H_{PS} \psi(\gamma, \beta) \rangle$ and for noncommuting Hamiltonians have been discussed in Refs. [21,22].

In many QAOA case studies, analytical or numerical, the expectation value of H_{PS} instead of the probability of the lowest-energy solution has been used as a proxy for performance. Concern has been raised in using expectation values (or approximation ratio) as a sole figure of merit because a high expected value does not guarantee the quality of solutions upon measurement; the underlying distribution needs to be scrutinized. In a general setting, the variance of the distribution would be required to further indicate sufficient concentration of probability on the desired solution.

We point out that for problems with a domain of discrete real values, as most combinatorial optimization problems are, a high approximation ratio generally accompanies a high value for the typical case. To see this, we analyze how the tail probability is bounded by the mean when the domain is a set of bounded discrete integer values. Consider a random integer variable $X \in \{0, 1, \ldots, m\}$; if the mean value is μ then for any $l \leq \lfloor \mu \rfloor$, where $\lfloor \cdot \rfloor$ is the floor function, the probability of x taking a value larger than l is lower bounded as

$$\Pr(X > l) \geqslant \frac{\mu - l}{m - l}.\tag{1}$$

In Appendix A we provide a proof for Eq. (1) under more general assumptions. In Sec. VC we will see examples: for our QAOA results of the high approximation ratio, without examining the energy distribution, we can infer with high confidence that a typical solution will have high cost.

III. PROBLEM FORMULATION

In this section we formulate the max- κ -colorable-subgraph problem in binary form using a one-hot-encoding

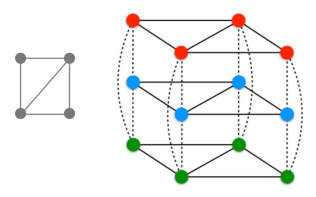


FIG. 1. Left: The original graph to be colored. Right: The qubit layout encoding the problem. Each vertex v is represented by κ qubits $x_{v,c}$ for $c=1,\ldots,\kappa$ representing its κ possible colors. The extended graph construction can be thought of taking the graph represented in its natural Euclidean space and then augmenting that space with another dimension and replicating the graph κ times for each of the colors. The phase-separation Hamiltonian is composed of two-qubit operations corresponding to edges on each surface, and the mixing operation is among the qubits in the augmented dimension.

representation for the possible colorings of each node. Using binary variable $x_{v,c}$ to indicate whether vertex v is assigned color c, the one-hot-encoding formulation requires solutions live in a subspace of the full Hilbert space that satisfies $\sum_c x_{v,c} = 1$, the *feasible subspace*. This results in two formulations of the coloring problem for QAOA—with and without a penalty term in the phase separator. We also recap nomenclature for various XY-Hamiltonian drivers introduced in Ref. [2] which becomes necessary when discussing their circuit implementations.

The max- κ -colorable-subgraph problem is formulated as follows.

Problem 1. Given a graph G = (V, E) with n vertices and m edges, and κ colors, maximize the size (number of edges) of a properly vertex-colored subgraph.

The max- κ -colorable-subgraph problem is encoded into qubits with a one-hot-encoding fashion to represent the colors. Each node of the graph G is expanded into κ qubits where each qubit occupation is used to represent a coloring of the node. For example, a three-coloring problem on a graph of four vertices requires 12 qubits depicted in Fig. 1.

In the feasible subspace where each vertex is assigned exactly one color, the cost or objective function

$$f_C = m - \sum_{i=1}^{\kappa} \sum_{v,v' \in F} x_{v,j} x_{v',j}$$
 (2)

counts the properly colored edges, and we aim at maximizing f_C . By the replacement $x \to (1 - \sigma^z)/2$ in Eq. (2), the corresponding quantum objective Hamiltonian is

$$H_C = \frac{1}{4}[(4 - \kappa)m\mathbf{1} + H_C'],$$
 (3)

where

$$H'_{C} = \sum_{v=1}^{n} d_{v} \sum_{j=1}^{\kappa} \sigma_{v,j}^{z} - \sum_{j=1}^{\kappa} \sum_{\{v,v'\} \in E} \sigma_{v,j}^{z} \sigma_{v',j}^{z}.$$
 (4)

Throughout this paper we use σ_x , σ_y , σ_z , and X, Y, Z interchangeably to refer to the Pauli operators. The approximation ratio we will adopt in the following is the ratio of the expectation value of the cost Hamiltonian, projected onto the feasible subspace, to the true maximal number of correctly colored edges:

$$r = \frac{\langle P_{\text{feas}} H_C P_{\text{feas}} \rangle}{C_{\text{max}}},\tag{5}$$

where P_{feas} is the projection operator onto the feasible subspace, and C_{max} is the number of edges in the true $\text{max-}\kappa$ -colorable subgraph. Note that by projecting to the feasible subspace the ratio of the feasible subspace to the full Hilbert space is also factored in. The numerator in Eq. (5) is equivalent to the average number of properly colored edges observed upon measurement, with the infeasible output valued zero.

A. Adding a penalty in the phase-separating Hamiltonian

The common practice for incorporate constraints is to add a penalty term to the cost function. For the one-hot-encoded problem we define a quadratic penalty to penalize the case that a node is assigned no color or multiple colors,

$$f_{\text{pen}} = \sum_{v} \left(1 - \sum_{j=1}^{\kappa} x_{v,j} \right)^2,$$
 (6)

which, up to a constant, corresponds to the penalty Hamiltonian.

$$H_{\text{pen}} = \frac{1}{2} \sum_{v} \left[(2 - \kappa) \sum_{j} \sigma_{v,j}^{z} + \sum_{j < j'} \sigma_{v,j}^{z} \sigma_{v,j'}^{z} \right], \quad (7)$$

that increases the energy of all states outside the subspace. The phase-separating Hamiltonian becomes a weighted sum of the cost and the penalty Hamiltonians:

$$H_{\rm PS} = H_C' - \alpha H_{\rm pen} \tag{8}$$

where the weight parameter $\alpha \in \mathcal{R}_+$. Note that in Eq. (8) the penalty Hamiltonian is subtracted because we aim to maximize the original cost function and minimize the penalty. In order for the penalized function to have the same optima as the original cost function, the penalty weight needs to be set above a critical value. In the current problem, assigning more than one color to a vertex is not energetically favorable, so it is the opposite, assigning no color to a vertex that may create fake maxima. Since for every no-color vertex there are at most $\lfloor d_v/\kappa \rfloor$ edges lifted from being improper, the penalty should satisfy $\alpha > \lfloor \max\{d_v\}/\kappa \rfloor$, and we can loosely take $\alpha \geqslant n/\kappa$. On the other hand, the range of possible values of f_C (and of spectral values of H_C) is κm . Therefore, any $\alpha > \kappa m$ will ensure an energy separation between all feasible states and all infeasible states.

It should be noted that, unlike the motivating situation in adiabatic computation, the energy gap plays no clear role in QAOA. Thus it should be expected that, while the introduction of a penalty into the cost Hamiltonian may alter the QAOA dynamics, perhaps manipulating the reachable set of unitary operators, the role of the penalty strength is unclear at best. This perspective is supported by the numerical results in

Sec. V A. Indeed, while for some problems, such as the one-hot-encoded problems under consideration, sophisticated mixers can be designed to satisfy the constraints [2], the design of general and systematic methods for incorporating constraints into QAOA remains an open problem.

In the penalty formulation the mixer can be either the standard X driver

$$H_X = \sum_{v=1}^{n} \sum_{c=1}^{K} \sigma_{v,c}^{X}$$
 (9)

or the XY Hamiltonian. If the XY Hamiltonian is selected the penalty parameter may help the variational optimizer maintain probability mass in the feasible subspace and is not strictly necessary. In QAOA, it is unclear how a penalty parameter helps maintain probability mass over the feasible subspace. The feasible space of a κ -coloring problem is the set of states that satisfy

$$Z_{\text{tot},v} \equiv \sum_{c=1}^{\kappa} \sigma_{v,c}^{z} = \kappa - 2, \tag{10}$$

i.e., a subspace spanned by states in the computational basis that correspond to bit strings of Hamming weight equal to 1.

Although formulating the penalty Hamiltonian facilitates the use of the standard X mixer in QAOA, which can be implemented in constant circuit depth, we emphasize that the relative size of the feasible space becomes exponentially small as the graph size grows and thus a penalty formulation is suboptimal. To see this, consider the size of the feasible subspace $\mathcal{H}_{\text{feas}}$; for each node, the feasible subspace can be spanned by states corresponding to κ Hamming-weight-1 bit strings, and hence is of dimension k, and the feasible subspace for the whole problem is of dimension k^n . The ratio of the feasible subspace sizes to the size of the full Hilbert space is

$$\frac{\dim(\mathcal{H}_{\text{feas}})}{\dim(\mathcal{H})} = \frac{\kappa^n}{2^{n\kappa}} = \left(\frac{\kappa}{2^{\kappa}}\right)^n,\tag{11}$$

which for any $\kappa \geqslant 1$ shrinks exponentially with the graph size n.

B. XY mixer: Enforcing evolution in the feasible subspace

The $Z_{\text{tot},v}$ constraint can be incorporated in a natural way by selecting a mixing term that preserves the feasible subspace. Here we use the XY Hamiltonian

$$H_{XY,v} = \frac{1}{2} \sum_{c,c' \in K}^{\kappa} H_{XY,v,c,c'},$$
 (12)

$$H_{XY,v,c,c'} = \sigma_{v,c}^{x} \sigma_{v,c'}^{x} + \sigma_{v,c}^{y} \sigma_{v,c'}^{y}, \tag{13}$$

which drives rotations in the $\{(0, 1), (1, 0)\}$ subspace of each color labeling. In the above equation the mixer applies to any color pair c, c' in a set K. It can be verified that, for any K, $[H_{XY,v}, Z_{\text{tot},v}] = 0$.

1. Complete vs ring mixing Hamiltonians

In Eq. (12), when the mixing set K includes all pairs, the mixer is termed a *complete-graph* mixer. An alternative is the *ring* mixer in which K takes a one-dimensional (1D) structure:

c'=c+1, and we apply the periodic boundary condition. In the same fashion, there are a variety of derivative mixers based on the XY Hamiltonian, depending on the underlying connectivity between colors. We focus on the complete-graph and ring mixers.

2. Simultaneous vs partitioned mixers

For a given mixing Hamiltonian, Eq. (12), for each node, a *simultaneous* mixer exactly applies the unitary $\exp[-i\beta H_{XY,v}]$ while a *partitioned* mixer applies the product of $\exp[-i\beta H_{XY,v,c,c'}]$ in some order of $\{(c,c')\}$. We define the *parity-partition* mixer such that a local XY Hamiltonian is applied on even pairs first and odd pairs next.

The parity-partitioned mixing unitary is a first-order approximation of the simultaneous mixing unitary. Employing the Zassenhaus formula through second order,

$$e^{it(H_{\text{even}} + H_{\text{odd}})} \approx e^{itH_{\text{even}}} e^{itH_{\text{odd}}} e^{\frac{t^2}{2}[H_{\text{even}}, H_{\text{odd}}]},$$
 (14)

allows us to characterize the leading error term $e^{-t^2/2[H_{\rm even},H_{\rm odd}]}$ as a function of κ . For simplicity, we consider even κ , so that $H_{\rm even}$ and $H_{\rm odd}$ contain n/2 commuting terms exactly. The parity-partitioned mixer can be represented as two separate Hamiltonians:

$$H_{\text{odd}} = H_{1,2}^{(XY)} + H_{3,4}^{(XY)} + \dots + H_{\kappa-1,\kappa}^{(XY)},$$

$$H_{\text{even}} = H_{2,3}^{(XY)} + H_{4,5}^{(XY)} + \dots + H_{\kappa,1}^{(XY)}$$
(15)

where $H_{j,j'}^{(XY)} = X_j X_{j+1} + Y_j Y_{j+1}$ and each term $H_{j,j+1}^{XY}$ in H_{even} commutes with all terms in H_{odd} except for $H_{j-1,j}^{(XY)}$ and $H_{j+1,j+2}^{(XY)}$. We can simply determine the term generated by the commutation

$$\left[H_{j-1,j}^{(XY)}, H_{j,j+1}^{(XY)}\right] = 2i(X_{j-1}Y_{j+1} - Y_{j-1}X_{j+1})Z_j$$
 (16)

to obtain the general form of the error term. Therefore, $[H_{\text{even}}, H_{\text{odd}}]$ is composed of $\kappa/2$ terms of the type of Eq. (16). Since $||X_{j-1}Y_{j+1}Z_j||$ is of order 1, we have $||[H_{\text{even}}, H_{\text{odd}}]|| \sim \kappa$ and therefore expect the difference between the simultaneous and the parity-partitioned mixing operators to be more prominent as κ grows.

In the above analysis, the two mixing operators in general do not commute. However, we only need to focus on their effects in the feasible subspace. Here we provide analysis on the commuting relations in the feasible subspace for general κ . Note that each $H_{j,j'}^{(XY)}$ operation in the feasible subspace corresponds to a 2×2 permutation matrix. Then H_{even} and H_{odd} can be identified with the two permutations π , $\sigma \in S_{\kappa}$ on κ letters:

$$\pi = (0\ 1)(2\ 3)\cdots((\kappa - 2)\ (\kappa - 1)),\tag{17a}$$

$$\sigma = (1\ 2)(3\ 4)\cdots((\kappa - 3)\ (\kappa - 2))((\kappa - 1)\ 0). \tag{17b}$$

Thinking of the letters arranged on a circle, these are the two possible permutations that consist of κ disjoint nearest-neighbor transpositions. It may be observed that

$$\sigma\pi = (0\ 2\ 4\ \dots\ (\kappa-2))(1\ 3\ 5\ \dots\ (\kappa-1))^{-1}$$
 (18a)

is a product of two disjoint cyclic permutations and therefore is of order $\kappa/2$, i.e., $(\sigma\pi)^{\kappa/2}$ is the identity permutation. Now

we see that π and σ satisfy exactly the relations necessary to generate D_{κ} , the dihedral group of order κ [23], i.e.,

$$D_{\kappa} = \langle \sigma, \pi | \sigma^2 = \pi^2 = (\sigma \pi)^{\kappa/2} = 1 \rangle.$$

In particular, we may note that, while D_4 is an Abelian group (isomorphic to the Klein four-group $\mathbb{Z}_2 \times \mathbb{Z}_2$), all D_{κ} for $\kappa > 4$ are non-Abelian. Therefore only for $\kappa = 4$, the simultaneous and the parity-partitioned ring mixers commute in the feasible subspace and hence are equivalent for QAOA.

3. Feasible initial states

The initial state in the standard QAOA framework with the X mixer is $|+\rangle^{\otimes n}$, the even superposition over all bit strings, which is a fair starting point given no prior knowledge about the optimal solution. This state is also the ground state of the X mixer, and can be generated by performing a single-qubit Hadamard transform on each qubit.

Under the new QAOA framework, which accepts a constraint, the full space spanned by all bit strings $\{0, 1\}^n$ is no longer a valid solution space. When the constraint, as in our case, dictates preserving the total magnetic quantum number,

$$Z_{\text{tot},v} = \sum_{c=1}^{\kappa} \sigma_c^z = C, \tag{19}$$

where $C \in [-\kappa, \kappa]$ is a constant integer, the feasible solution space is composed of Hamming-weight $(\kappa + C)/2$ bit strings, which correspond to states that satisfy Eq. (19). In analogy to the $|+\rangle^{\otimes n}$ state for the case where all bit strings are feasible solutions, a fair initial state should be the even superposition of all Hamming-weight $(\kappa + C)/2$ bit strings. Such a state is also an eigenstate of the XY mixer. In the graph-coloring problem, $C = \kappa - 2$, the generalized W state is the fair starting state. In Sec. C we survey circuit construction methods that can be used to create a W state.

IV. CIRCUIT REALIZATIONS

In this section we describe how to implement the various components of QAOA into short-depth circuits. We start by assuming the physical qubits are all-to-all connected, and show that the simultaneous complete-graph and ring mixers can be realized in depth linear and logarithmic in κ , respectively. In Sec. IV C we discuss the depth required due to limited connectivity between the physical qubits.

A. Logarithmic depth simultaneous ring mixer

The interacting spin-1/2 chain is one of the oldest problems of quantum mechanics. Stemming from the resemblance between spin-1/2 raising (resp. lowering) operators and fermionic creation (resp. annihilation) operators, in the detailed work of Ref. [24], the Jordan-Wigner transformation was introduced to convert spin-1/2 systems into problems of interacting spinless fermions. While in general spin-spin interactions map to nonlocal fermionic interactions, for the one-dimensional XY problem the transformation results in a particularly precise form involving only quadratic fermionic

couplings:

$$H_{XY} = \sum_{c=1}^{\kappa} \left(\sigma_c^{x} \sigma_{c+1}^{x} + \sigma_c^{y} \sigma_{c+1}^{y} \right)$$

$$\downarrow$$

$$H_{XY} = 2 \sum_{c=1}^{\kappa} (a_c^{\dagger} a_{c+1} + \text{H.c.}), \tag{20}$$

where \hat{a} and \hat{a}^{\dagger} are fermionic operators, and we assumed κ is even for simpler demonstration.

The quadratic Hamiltonian in Eq. (20) can be diagonalized by a basis rotation on the operators. For nearest-neighbor onebody coupling, the fermionic Fourier transform

$$\hat{a}_{c}^{\dagger} = \text{FFFT}^{\dagger} \hat{f}_{k}^{\dagger} \text{FFFT} \equiv \frac{1}{\sqrt{\kappa}} \sum_{c} e^{i2\pi ck} \hat{f}_{k}^{\dagger},$$

$$\hat{a}_{c} = \text{FFFT}^{\dagger} \hat{f}_{k} \text{FFFT} \equiv \frac{1}{\sqrt{\kappa}} \sum_{c} e^{-i2\pi ck} \hat{f}_{k}$$
(21)

is sufficient to diagonalize the Hamiltonian. We use the notation FFFT (fermionic fast Fourier transform) to denote the circuit for the operator Fourier transform and not the quantum Fourier transform [25]. The *XY* Hamiltonian on a ring is then exactly diagonalized as [26]

$$H_{XY} = \sum_{k=1}^{K} E_k f_k^{\dagger} f_k \tag{22}$$

where the eigenenergies $E_k = 2\cos(2k\pi/\kappa)$. Replacing the number operator $f_k^{\dagger} f_k$ with qubit operators $(1 - \sigma_k^z)/2$, the Hamiltonian can be expressed as

$$H_{XY}^{(k)} = \sum_{k=1}^{K} E_k (1 - \sigma_k^z) / 2$$
 (23)

where the upper index (k) is added as a reminder that we are in the momentum representation. In this representation evolving $e^{-i\beta H_{XY}^{(k)}}$ involves only single-qubit Z rotations.

The FFFT has emerged as a route to efficient simulation for fermions in tensor networks [27] and quantum circuits representing fermionic systems [25,28]. The circuit is constructed in a similar structure to the decimation-in time radix-2 classical Fourier transform and inherits the divide-and-conquer complexity. The FFFT circuit can be implemented with $O(\log(\kappa))$ [27] depth for a system with parallel arbitrary two-qubit interactions. For more realistic systems where only nearest-neighbor interactions are allowed fermionic swaps are required to swap the two modes together to perform the butterfly operation. This adds an additional overhead resulting in a $O(\kappa \log(\kappa))$ circuit depth and $O(\kappa^2 \log(\kappa))$ total gate count [25]. The gate depth required to implement the FFFT was further improved to $O(\kappa)$ in Ref. [29] by using a Givens rotation network and requires only linear connectivity.

We also point out that the Givens rotation network is a powerful tool for state preparation for general quadratic Hamiltonians. For pairing models, the linear depth network was used to prepare ground states [30]. This initial state can be used in the context where the hard constraint is of the form that qubits must appear paired up. We point this out as an example of how different flavor constraints can correspond to evolving a wide variety of constraint-preserving Hamiltonians.

B. Linear depth simultaneous complete-graph mixer

We consider the simultaneous mixer for a node, $e^{-i\beta H_{\text{complete}},v}$, with $H_{\text{complete},v} = \sum_{c < c' = 0}^{\kappa - 1} H_{XY,v,c,c'}$, which corresponds to a complete graph of variables corresponding to all colors for each vertex $v, \{x_{v,c}\}$. Beyond a one-dimensional layout, the analytical solution to the XY model is not known; therefore, exactly realizing the evolution of the XY model on a complete graph poses a challenge. In this section we show that within the subspace of total $Z_{\text{tot},v} = \pm (\kappa - 2)$ as in our case, when $\kappa = 2^m$, this unitary can be exactly implemented in a circuit of depth $\kappa - 1$, up to a constant factor accounting for breaking a generic two-qubit operator to any fixed universal set of single- and two-qubit operators.

We illustrate the process using $\kappa = 4$ and then show the general formula. For $\kappa = 4$, we consider three partitions of the full set of colors: $\{\{0, 1\}, \{2, 3\}\}, \{\{0, 2\}, \{1, 3\}\}\}$, and $\{\{0, 3\}, \{1, 2\}\}$; in the feasible subspace, we have

$$\exp\left[-i\beta \sum_{c,c'\in[0,3]} (XY)_{c,c'}\right]$$

$$= \exp\{-i\beta[(XY)_{0,1} + (XY)_{2,3}]\}$$

$$\exp\{-i\beta[(XY)_{0,2} + (XY)_{1,3}]\}$$

$$\exp\{-i\beta[(XY)_{0,3} + (XY)_{1,2}]\}.$$
 (24)

where for notational simplicity we use $(XY)_{j,j'}$ to refer to the XY Hamiltonian $H_{j,j'}^{(XY)}$ defined below Eq. (15). Note that this equivalence is approximate in general but exact if we consider only the action on the feasible subspace. The fact that these partitioned operators commute in the feasible subspace can be easily verified mathematically.

The following perspective on the partitioning scheme allows us to derive a generalization for any κ . Consider an integer variable x the values of which are from 0 to 3; in the one-hot encoding, in the feasible space, the XY operation on a pair of qubits swaps the integer values the states represent. For example, $(XY)_{1,3}$ swaps between the variable taking value 1 and taking value 3. Now consider the two-bit binary encoding of x: $x = 2^1x_1 + 2^0x_0$ where x_0 and x_1 are bits. The swap between $\{0, 1\}$ and $\{2, 3\}$ corresponds to flipping the zeroth bit x_0 . The swap between $\{0, 2\}$ and $\{1, 3\}$ corresponds to flipping both bits x_0 and x_1 . Such operations can happen in any order without affecting the final value of x, hence the corresponding partitioned mixers commute.

For a general $\kappa=2^m$, the partition can be read out taking the inverse of this process: all pairs involved in each l-bit flipping form a partition, for $l=1,\ldots,m$. There are $\binom{m}{l}$ many l-bit flips, hence the total partitions $\sum_{l=1}^m \binom{m}{l} = 2^m - 1 = \kappa - 1$. Within each partition the pairwise XY operators commute and can be executed simultaneously. The simultaneous complete-mixer unitary can be accordingly executed in depth $\approx \kappa - 1$.

For example, partitions for $\kappa=8$ can be prescribed using the following equations. Here we use $\tilde{0}$, $\tilde{1}$, and $\tilde{2}$ to represent bits in the binary encoding. The left-hand side for each equa-

TABLE I. Demonstration of deriving the partition corresponding to operator IXX on the binary encoding, Eq. (25).

Decimal	One-hot	Apply IXX	Decimal
0	000	011	3
1	001	010	2
2	010	001	1
3	011	000	0
4	100	111	7
5	101	110	6
6	110	110	5
7	111	100	4

tion is a *l*-bit flip operation; all seven operations commute. The right-hand side is derived from reading off the effect of the operation on the numerical colors. The right-hand side gives the corresponding partition for the *XY* operators in the one-hot encoding, with the detailed procedure displayed in Table I:

$$I_{\bar{2}}I_{\bar{1}}X_{\bar{0}} = (XY)_{01} + (XY)_{23} + (XY)_{45} + (XY)_{67},$$

$$I_{\bar{2}}X_{\bar{1}}I_{\bar{0}} = (XY)_{02} + (XY)_{13} + (XY)_{46} + (XY)_{57},$$

$$X_{\bar{2}}I_{\bar{1}}I_{\bar{0}} = (XY)_{04} + (XY)_{15} + (XY)_{26} + (XY)_{37},$$

$$I_{\bar{2}}X_{\bar{1}}X_{\bar{0}} = (XY)_{03} + (XY)_{12} + (XY)_{47} + (XY)_{56},$$

$$X_{\bar{2}}I_{\bar{1}}X_{\bar{0}} = (XY)_{05} + (XY)_{14} + (XY)_{27} + (XY)_{36},$$

$$X_{\bar{2}}X_{\bar{1}}I_{\bar{0}} = (XY)_{06} + (XY)_{17} + (XY)_{24} + (XY)_{35},$$

$$(26)$$

$$X_{\bar{2}}X_{\bar{1}}X_{\bar{0}} = (XY)_{07} + (XY)_{16} + (XY)_{25} + (XY)_{34}.$$

C. On realistic layout of physical qubits

To achieve the above derived circuit depth for simultaneous ring and complete-graph mixers requires physical qubits to have a particular connectivity, for example, all-to-all connectivity for the simultaneous complete-graph mixer. For a physical-qubit-layout of lower connectivity, SWAP operations may be necessary to enable the pairwise operations [15,17].

We first note that in the feasible space an XY operation $\exp[-i\beta(XY)_{j,j'}]$ and a SWAP_{j,j'} executed consecutively are equivalent to a XY of different parameter:

$$e^{-i\beta(XY)_{j,j'}}$$
SWAP_{j,j'} = $ie^{-i(\beta+\frac{\pi}{2})(XY)_{j,j'}}$, (27)

or, in matrix form,

$$\begin{pmatrix} 1 & 0 & 0 & 0 \\ 0 & -i\sin\beta & \cos\beta & 0 \\ 0 & \cos\beta & -i\sin\beta & 0 \\ 0 & 0 & 0 & 1 \end{pmatrix}.$$
 (28)

This relation can be explored in circuit compilation to achieve a circuit on physical qubits of lower depth.

For example, in the $\kappa = 4$ example for the simultaneous complete-graph mixer in Sec. IV B, if the physical qubits form a ring 1-2-3-4-1, the three partitions can be executed in the following order:

$$\{\{0,1\},\{2,3\}\}, \{\{0,3\},\overline{\{1,2\}}\}, \{\{0,2\},\{1,3\}\}$$

where the pair with an overline indicates a SWAP is combined in the XY mixing in the fashion of Eq. (27). This SWAP

enables the operation of the last partition, and the whole circuit is of depth 3, the same as in the case of a complete graph.

For a general κ simultaneous mixer, this feature helps scheduling but cannot avoid SWAPs completely because the partitions need to be executed in a specific order.

On the other hand, executing the pairwise XY unitary in any given order would give a valid partitioned mixer, though not equivalent to the simultaneous one. Exploiting the feature Eq. (27) combined with a bubble sort scheme, we can completely avoid SWAPs and implement a partitioned complete-graph mixer in linear depth.

The same SWAP network circuit architecture, derived from a fermionic simulation perspective, can be used to implement the phase-separator Hamiltonian with no Trotter error [16,29]. Viewing the qubits as an array of κ sites the parallel bubble sort algorithm implements a swap network in κ depth where all elements (qubits) of the array that are swapped pass each other once. Since all terms in the phase separator commute, there is no incurred Trotter error. The swap interaction can be efficiently combined with the evolution of a $e^{-i\theta ZZ}$ nearestneighbor interaction by adding an $e^{-i\theta Z}$ in between the second and third controlled-NOT (CNOT) (denoted CX below) in the SWAP decomposition [17].

Explicitly,

$$SWAP_{0,1}e^{-i\theta ZZ_{0,1}} = CX_{0,1}CX_{1,0}CX_{0,1}e^{-i\theta ZZ_{0,1}}$$
 (29)

$$= CX_{0.1}CX_{1.0}e^{-i\theta Z_1}CX_{0.1}$$
 (30)

can be used as the swap interaction and simultaneously evolve a local ZZ-interaction term. For any encoding model that increases the dimensions of the graph, such as in the case where one-hot encoding is used, simulating the interaction term removes the necessity of encoding techniques such as minor embedding or classical logical encoding [31,32].

V. SIMULATION RESULTS

In this section, we present the results of numerical simulations of QAOA applied to the $\max \kappa$ -colorable-subgraph problem. We first compare the performance of the XY mixer to that of the X mixer with penalty. We then more deeply explore the behavior of XY mixers, looking at general features of their performance on small hard-to-color graphs, and comparing complete-graph XY mixers against ring XY mixers.

To acquire a good set of QAOA parameters, a stochastic optimizer is needed; in Appendix B we show a rugged landscape with local optima in the parameter space that would cause a problem for deterministic optimizing methods like gradient descent. We instead use basin hopping with Broyden–Fletcher–Goldfarb–Shanno (BFGS) algorithm to obtain (sub)optimal parameters.

A. Death of the X mixer

We use a simple example, two-coloring and three-coloring of a triangle, to demonstrate the performance comparison of XY and X mixers.

Note that the penalty weight α in general affects the performance of the algorithm. In Fig. 2 we show that for two-coloring the approximation ratio is optimized over the parameter set (γ, β) for each penalty weight α . The best

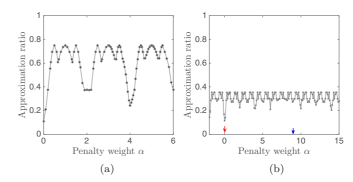


FIG. 2. (a) Two-coloring and (b) three-coloring of the triangle with level 1 QAOA. The highest approximation ratio across the parameter sets (γ_1, β_1) is plotted vs the penalty weight α . The red arrow at $\alpha=0$ indicates the minimum penalty that guarantees the optimal state being the optimal state in the feasible subspace, and the blue arrow at $\alpha=9$ indicates the penalty value that guarantees separation between energies of feasible and infeasible states.

approximation ratio, r, takes the value 0.75 while with the XY driver QAOA_{p=1} we have r=1.

In Fig. 2, while the penalty strength has an effect on the behavior of level 1 QAOA, there appears to be no clear intuition for choosing a good value. In particular, the minimum penalty that guarantees the optimal state being the optimal state in the feasible subspace, indicated by the red arrows on the plots, does not stand out, nor does the penalty value that guarantees separation between energies of feasible and infeasible states, indicated by the blue arrows. This supports our argument in Sec. III A that the role of the energy gap plays no clear role in QAOA.

For three-coloring, in Fig. 3 we plot how the approximation ratio varies in the two-dimensional (γ, β) space, for using the X mixer and for using the XY mixer. While with the X mixer the QAOA $_{p=1}$ gives approximation ratio ≈ 0.2 across the parameter value range, with the XY mixer parameter values that correspond to ≈ 0.8 can easily be found. This example thus shows significant performance advantage in using the XY as compared with the X mixer.

B. Small and hard-to-color graphs

For a fixed classical algorithm, a *slightly-hard-to-color* graph is a graph for which the algorithm will sometimes

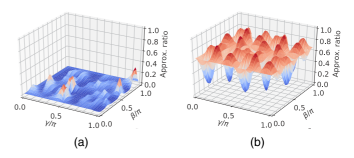
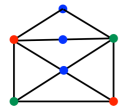


FIG. 3. Numerical results for level 1 QAOA on the problem of three-coloring of a triangle graph. (a) Using the X mixer along with the phase-separating Hamiltonian, Eq. (8), where the penalty weight is taken to be the numerically determined optimal value $\alpha^* = 1.7$. (b) Using the XY mixer with the W state being the initial state.



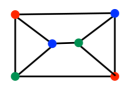


FIG. 4. The two small and hard-to-color graphs: envelope and prism. A valid three-coloring is shown on each graph.

yield the optimal solution. Similarly, a hard-to-color graph is one such that the chosen algorithm never yields the optimal solution. Two examples are the envelope and the prism graphs [33], sketched in Fig. 4. The prism graph is the smallest slightly-hard-to-color graph for the smallest-last sequential coloring method and the envelope graph is the smallest hardto-color graph for the largest-first sequential method. Note that these classical algorithms are aiming to compute the chromatic number, while in this paper we focus on finding the maximal colorable subgraph. Although finding the maxcolorable subgraph could serve as a subroutine for determining chromatic numbers, the chromatic number can also be directly attacked by QAOA using a much more complex mixer [2]. Nevertheless we are not aiming at doing side-by-side comparison of quantum and classical algorithms, and will use these small graphs only as a proof-of-principle demonstration of the QAOA with XY mixers.

1. Performance of QAOA with the simultaneous ring mixer

With the simultaneous ring mixer, Fig. 5 shows the results for QAOA levels 1 to 6. For each level, the *W* state is used as initial state, and a stochastic search (basin hopping with BFGS) is performed to optimize the expected value of the cost Hamiltonian over the angle sets. The approximation ratio corresponding to the optimal expectation value is plotted as filled circles. Even at level 1, the approximation ratio takes a high value 0.8, and this value quickly approaches 1 as the

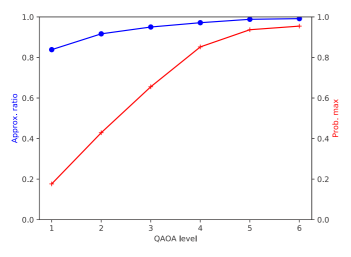


FIG. 5. The prism graph. Dots are approximation ratios and crosses are the expected probability of getting the optimal coloring. For each QAOA level, results are shown at the (sub)optimal angles resulted from a basin-hopping search.

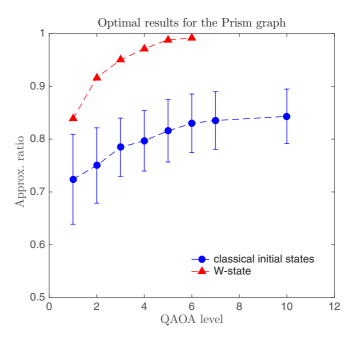


FIG. 6. The prism graph, showing the expected value of QAOA optimized over the angle sets. Triangles show the results with the *W* state as initial states. Circles show the results with a feasible classical initial state, averaged over the set of all feasible classical states; the error bar is the standard deviation. For each initial state, optimization over angles is derived from a basin-hopping search.

level increases. Furthermore, for each level, we computed the probability of getting the actual optimal solution (a valid three-coloring) upon measurement. At level 1, this probability is slightly lower than 0.2, and quickly goes above 0.6 at level 3, which implies that, repeating the experiment three times, one will find a valid coloring with probability >0.9.

2. Effect of initial states

The W state—as both an even superposition of all feasible classical states and the ground state of the simultaneous ring mixer—is a natural candidate for the initial state for QAOA. It involves multiple two-qubit gates to prepare. An easier-to-prepare state for each vertex can be defined via a randomly assigned coloring (feasible but not necessarily optimal), $|\psi_C\rangle$, i.e., a randomly drawn bit string of Hamming weight 1. Preparing such a state involves only n single-qubit gates.

We study both initial states for the prism graph with the simultaneous ring mixer. For level-1 QAOA, the best achievable optimization ratio [optimized over all angle sets (β, γ)] for the W state is higher than the classical Hamming-weight-1 state $|\psi\rangle_C$. Notice that for $|\psi\rangle_C$ the phase-separating unitary commutes with the density matrix of the state, and hence has no effect on the state evolution. As a result, the whole circuit for level-1 QAOA is equivalent to applying the mixing unitary followed by measurement. We further simulated higher levels, and in Fig. 6 we show the performance of QAOA with the W state vs a classical state as initial state. We found that with the classical initial state the performance of QAOA is significantly lower than using the W state as initial state. Even at level 10, $r_{\text{classical}}$ is still lower than r_W for level 1. Moreover, the approximation ratio with classical initial state shows a

TABLE II. Top: Benchmarking graph sets: each row indicates all χ -chromatic graphs of size n, and we solve the problem of κ coloring of such graphs choosing $\kappa = \chi$. Bottom: Benchmarking graph sets II for examining the simultaneous vs partitioned ring mixers on different ring sizes; each row indicates all connected graphs of size n, and we solve the problem of κ coloring of such graphs. Because the total number of qubits is $n\kappa$, which is the limiting factor to the simulation, we limit to small n to see κ varying up to 8.

χ	n	No. graphs
3	5	12
3	6	64
3	7	475
4	6	26
4	7	282
5	7	46
6	7	5
n	κ	No. graphs
4	4	6
4	6	6
4	8	6

tendency toward saturation around level 10—this could either be the nature of the algorithm or be due to increasing difficulty in finding the global optimum in the parameter subspace as the level increases, which poses another practical consideration for application. (Note that due to the optimization over parameter space for each initial state the average over the classical initial state is not equivalent to prepare the initial state in a mixed state for the ensemble.)

Because our simulation is noise free, due to ergodicity, in the limit of $p \to \infty$ the optimal performance should be independent of the initial state. But for practical implementation on a near-term hardware, where noise accumulates fast with circuit depth, such medium-level QAOA behavior is of high relevance. In Appendix C we survey methods to generate quantum circuits for preparing W states. It is shown that with certain methods it can be generated with $O(\kappa)$ CNOT gates. The overall performance of QAOA will be a tradeoff between the extra effort in preparing the W state and the damage that comes with circuit depth.

C. Benchmarking graph sets

To better understand the behavior of these QAOA graph-coloring algorithms, we make use of the sets of all κ -chromatic graphs of size n as the benchmarking sets for the XY mixers under consideration. See Table II for the number of instances in each benchmarking set.

1. Approximation ratio and probability of optimal solution

Using the *W* state as the initial state, for simultaneous ring and complete-graph mixers, the mean and median of the approximation ratio as well as the probability of optimal solution are evaluated across problem sets.

The following observations have been made on the typical performance for each problem set.

a. Consistent performance over instances. For all problem sets, the approximation ratio and the probability-of-optimal-

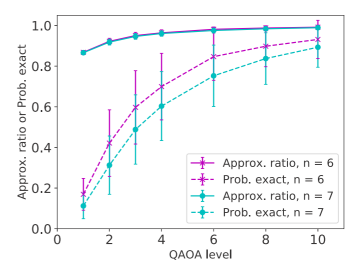


FIG. 7. Approximation ratio (solid lines) and probability to exact solution (broken lines) for QAOA with the ring simultaneous mixer, with n = 6 (crosses) vs n = 7 (filled circles).

solution curves as a function of the QAOA level are highly consistent across graphs, bearing the same shape for the prism and envelope graphs. For each problem set, the approximation ratio showed very little deviation from the mean (demonstrated by the small error bars in Fig. 7).

b. Larger graphs are harder to color. As expected, for the same κ , as n increases, the performance of QAOA with the same type of mixer decreases (see Fig. 7 for comparison of the simultaneous ring mixer for n=6 and 7).

c. The complete-graph mixer is better than the ring mixer. For the same problem size n, the simultaneous complete-graph mixer demonstrates better performance than the simultaneous ring mixer in QAOA levels from 1 to 10. See the scatter plot for QAOA level 2 and level 8 in Fig. 8. For small QAOA levels, this advantage is uniform across instances for smaller levels, as shown in Fig. 8(a) for level 2, where for all 282 instances the complete mixer generates a higher approximation ratio. The advantage is decreasing as QAOA level increases [cf. Figs. 8(a) and 8(b)]. This is possibly due to the approximation ratio getting close to 1. We also speculate

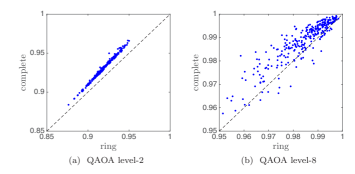


FIG. 8. QAOA with simultaneous mixers. Performance comparison between ring and complete-graph mixers applied to the same graph-coloring problems. The axes show the approximation ratio achieved using the labeled mixer type. The scatter plot shows the results for four-coloring of all connected chromatic-4 graphs of size n = 7. In (b), for better visibility, an outlier data point at (ring = 0.95, complete = 0.9) is not shown in the plot.

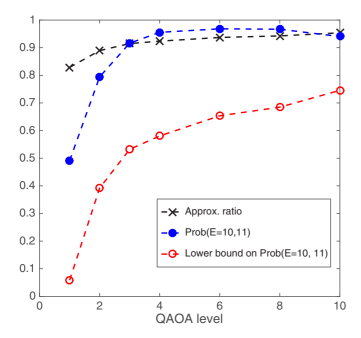


FIG. 9. Three-coloring of the envelope graph (11 edges). QAOA with the simultaneous ring mixer. For each QAOA level, the probability of getting the top two highest approximate results (cost 11 and 10) is shown in comparison to the bound given by Eq. (1) with the observed approximation ratio as parameter.

that the QAOA level where this closeup happens would vary with κ , the number of colors.

d. Similar performance between simultaneous and parity mixers for small κ . We also study κ coloring of all connected graphs (regardless of chromatic number) of size n=3 and 4, with varying κ to compare simultaneous vs partitioned ring mixers on different ring sizes. Since for $\kappa=4$ the simultaneous and the parity mixers are equivalent, we will need to go for higher κ to examine the difference; however, numerical power is limited by the number of qubits $n\kappa$, thus we examined two classically trivial cases: n=4, $\kappa=6$ and 8 (trivial coloring exists). Both approximation ratio and probability of exact solution are high due to the small problem size, and no noticeable difference is observed between the performance of partitioned and simultaneous mixers. Extensive studies on larger problem sizes are needed to further evaluate these two types of mixers.

2. Typical solution upon measurements

Note that our optimization over the set of angles is designed to maximize the expected value, and the high approximation ratio discussed in Sec. V C 1 is also about the expected value. For approximate optimization, the expectation value of the approximation ratio is not the whole story. One also cares about the probability of getting the optimal or near-optimal states upon measurement. We apply the argument and analysis on the tail probability in Sec. II, Eq. (1), on the case of three-coloring of the envelope graph (11 edges), and show in Fig. 9 the theoretical lower bound in probability of getting a solution with costs 10 or 11, i.e., the valid coloring or only one improperly colored edge. The true probability from evaluating

the wave function is shown for comparison. For QAOA level 3 and up, the bound inferred from the approximation ratio gives us confidence that with greater than 50% probability we will get the optimal or the next best solutions.

Viewing the QAOA as an exact solver, as observed in the case of small hard-to-color graphs, for the benchmarking problem sets, we also see that as p increases, along with the increase in r^* , there is a more drastic increase in the probability of optimal solution. In Fig. 7 we also plot the mean probability of optimal solution as p changes, with error bars indicating the standard deviation over the graphs in the set.

VI. CONCLUSION

Exploring the range of applications of the QAOA provides insight into what can be achieved with near-term quantum resources. While the general search for applications of QAOA is important, the detailed specification of the algorithm can be the difference between success and failure when running the algorithm on a real device. These details can change if the gate-model computing substrate is switched, e.g., switching from superconducting qubits to ions. For example, constant factors in circuit depth gained by switching gate sets can drastically change circuit depth and thus the success probability of the overall algorithm.

In this paper we explored applying QAOA to optimization over κ -ary variable sets. Our representative example of this optimization was the max- κ -colorable-subgraph problem on small hard-to-color graph problems. We numerically demonstrate and theoretically motivate that the XY-mixer Hamiltonian is a natural choice for this particular set of problems. Part of the numerical analysis was providing circuit implementations for the phase-separator and the XY mixer under various qubit topologies. Though there is higher implementation cost of the XY mixer in comparison to the standard X mixer, the benefits of eliminating a penalty term and restricting dynamics to the feasible space can potentially outweigh the linear-depth implementation cost.

Along with the circuit analysis the bound on tail effects based on the mean value we provide in Eq. (1) suggests that a high mean value is sufficient to guarantee performance without having to analyze the variance of the distributions produced by QAOA.

This paper establishes the possibility of using more sophisticated drivers in a QAOA framework for naturally including constraints. We expect this analysis is helpful for nearterm experimental validations of the QAOA algorithm and, hopefully, inspires alternative constraint encodings that would lower the representational cost of real-world optimization problems.

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APPENDIX A: PROOF ON FINITE TAIL PROBABILITIES

In what follows, we will take $\{a_j\}_{j=0}^K \subset \mathbb{R}$ to be a strictly ordered finite set, i.e., $a_0 < a_1 < \cdots < a_K$. Suppose that p_j is the probability of a random variable X having outcome a_j , with $\sum_{j=0}^K p_j = 1$. We will assume below that we know the values $\{a_j\}$ and the mean μ of this probability distribution, but not the entire distribution itself.

Lemma 1. Given $\mu = \sum_{j=0}^{K} p_j a_j$, with $\mu \geqslant a_l$ for some $0 \leqslant l \leqslant K$.

$$\Pr(X > a_l) \geqslant \frac{\mu - a_l}{a_K - a_l} = 1 - \frac{a_K - \mu}{a_K - a_l}.$$
 (A1)

Proof. We will prove the bound using the method of Lagrange multipliers. To ensure the probabilities are nonnegative, we will represent them as squares: $p_j = q_j^2$. Then the Lagrangian is

$$\mathcal{L} = \sum_{j=l+1}^{K} q_j^2 + \lambda \left(\sum_{j=0}^{K} q_j^2 - 1 \right) + \gamma \left(\sum_{j=0}^{K} q_j^2 a_j - \mu \right).$$
 (A2)

Differentiating and setting the derivatives equal to zero, we find the conditions

$$2q_i(\delta_{>l}(j) + \lambda + a_i\gamma) = 0, \tag{A3}$$

i.e.,

for
$$j \leq l$$
: $q_j = 0$ or $\lambda + a_j \gamma = 0$; (A4a)

for
$$j > l$$
: $q_j = 0$ or $1 + \lambda + a_j \gamma = 0$. (A4b)

First, consider the case that $q_j=0$ for all j>l. Then $p_j=q_j^2=0$ for all j>l, so the only way $\mu\geqslant a_l$ can be satisfied is if $\mu=a_l$. Then $\Pr(X>a_l)=\sum_{j=l+1}^K p_l=0=\frac{\mu-a_l}{a_K-a_l}$, so (A1) is satisfied. Similarly, if $q_j=0$ for all $j\leqslant l$, then $\Pr(X>a_l)=1$, so this case represents the maximum, rather than the minimum of $\Pr(X>a_l)$, and in any case (A1) is satisfied.

Now, if $q_m \neq 0$ and $q_n \neq 0$ for $0 \leqslant m < n \leqslant l$, then it follows that $\lambda = \gamma = 0$ and therefore $q_j = 0$ for all j > l, so that, as just argued, (A1) is satisfied. If, on the other hand, $q_m \neq 0$ and $q_n \neq 0$ for $l < m < n \leqslant K$, then it follows that $\gamma = 0$ and $\lambda = -1$, so that $q_j = 0$ for all $0 \leqslant j \leqslant l$, and therefore (A1) is satisfied.

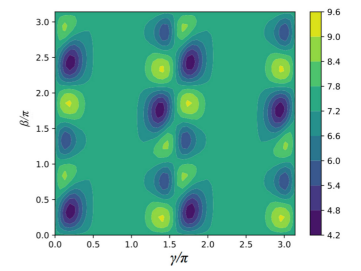
What remains is the case that exactly one $q_m \neq 0$ for $0 \leq m \leq l$ and exactly one $q_n \neq 0$ for $l < n \leq K$ and all other q_j are zero. Then $\Pr(X > a_l) = p_n$, $p_m = 1 - p_n$, and $\mu = p_m a_m + p_n a_n = (1 - p_n) a_m + p_n a_n$. Solving for p_n , we find

$$\Pr(X > a_l) = p_n = \frac{\mu - a_m}{a_n - a_m} = 1 - \frac{a_n - \mu}{a_n - a_m}.$$
 (A5)

It is easily seen that this expression for p_n decreases as a_n increases and decreases as a_m increases, so the minimum p_n obtained in this way is when m = l and n = K:

$$Pr(X > a_l) = p_n = \frac{\mu - a_l}{a_K - a_l} = 1 - \frac{a_K - \mu}{a_K - a_l}, \quad (A6)$$

from which it follows that (A1) is satisfied.



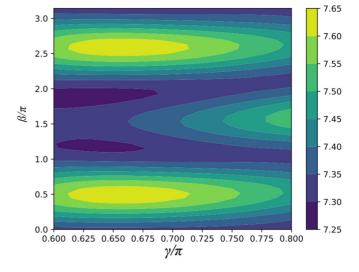


FIG. 10. Landscape for level-1 QAOA for the envelope graph. Top: Full landscape. Bottom: Zoom-in of the seemingly flat area that contains local maxima.

APPENDIX B: RUGGED LANDSCAPE

The landscape of the parameter space plays an important role in quantum control. In the case of MAXCUT for a ring graph (equivalent to binary encoding for a two-coloring of the ring), in Ref. [5] it has been observed that the landscape contains only the global maximum. In the current case of QAOA with the XY driver on graph coloring, we notice that even for level-1 QAOA the control landscape is rugged and contains the local maximum. In Fig. 10 the landscape for the envelope graph is plotted, and the inset reveals local optima. Stochastic optimization is therefore needed to perform a parameter search.

APPENDIX C: W-STATE GENERATION

The W state is a well-known multipartite-entangled state that plays an important role in quantum information theory. Here, we survey methods to produce a generalized W state using quantum gates.

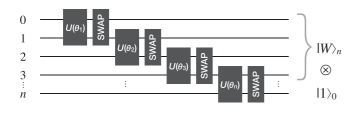




FIG. 11. Circuit for generating the $|W\rangle$ state for arbitrary n. SWAP gates are added for illustration on 1D architecture of physical qubits. Qubit n is discarded at the end.

1. Sequential generation of the W state

In Ref. [34], it was shown a W-type state, which is any state living in the subspace spanned by states corresponding to Hamming-weight-1 bit strings, can be generated using an auxiliary qubit by sequentially entangling it with each qubit. Here we detail the case for the W state using this method. Consider an auxiliary qubit q_0 and an n-qubit register (q_1 to q_n), initialized in the tensor product state $|0\rangle \otimes |0\rangle^{\otimes n}$. Entanglement between q_0 and q_j is introduced by unitary

$$U_{0,j}(\theta_j, \phi_j) = |01\rangle\langle 01| + |10\rangle\langle 10| + c_j|00\rangle\langle 00| + c_j|11\rangle\langle 11| + s_j|11\rangle\langle 00| - s_i^*|00\rangle\langle 11|,$$
(C1)

where $c_j \equiv \cos \theta_j$ and $s_j \equiv e^{i\phi_j} \sin \theta_j$. This unitary generates superposition between $|00\rangle$ and $|11\rangle$ states in the subspace of q_0 and q_1 , and acts as identity to the orthogonal subspace. Unitary $U_{0,1}$ acting on the initial state yields

$$U_{0,1}(\theta_1, \phi_1)|00\rangle = c_1|00\rangle + s_1|11\rangle.$$
 (C2)

This unitary can be realized by a circuit as $U(\theta)$ in Fig. 11. The state for the whole system is now

$$c_1|000\cdots0\rangle + s_1|110\cdots0\rangle.$$
 (C3)

Further applying $U_{0,2}(\theta_2, \phi_2)$ on q_0 and q_2 yields

$$c_1c_2|000\cdots 0\rangle + c_1s_2|101\cdots 0\rangle + s_1|110\cdots 0\rangle.$$
 (C4)

In this fashion we apply $U_{0,k}$ sequentially on the initial state for $k \in [1,n]$. Each application introduces amplitudes in $|0\rangle \otimes X_k |0\rangle^{\otimes n}$ where the register qubit state corresponds to the Hamming-1 bit string with the one on the kth qubit. In order for all Hamming-weight-1 register states to be of the same amplitude, and decoupled from the ancilla, the following conditions are imposed:

$$c_1 c_2 \cdots c_n = 0, \tag{C5}$$

$$c_i|s_{i+1}| = |s_i|,$$
 (C6)

which has a solution $\sin \theta_j = \frac{1}{\sqrt{n+1-j}}$. By using such angles, the $|0\rangle \otimes |0\rangle^{\otimes n}$ state is removed in the *n*th step because $\cos \theta_n = 0$, resulting in the final tensor-product state:

$$|1\rangle \otimes \frac{1}{\sqrt{n}} (e^{i\phi_1} |10 \cdots 0\rangle + e^{i\phi_2} |010 \cdots 0\rangle + \cdots + e^{i\phi_n} |0 \cdots 01\rangle).$$
 (C7)

Setting all $\phi_j = 0$ further leads to the exact W state on the register:

$$|1\rangle \otimes |W\rangle_n$$
. (C8)

Figure 11 shows the corresponding circuit.

2. Reverse engineering for W-state preparation

Wang, Ashhab, and Nori [35] outline a procedure to produce arbitrary states with fixed particle number. The approach is a recursive approach and scales as $O(2^m n^m/m!)$ where m is the number of spin-up or $|1\rangle$ states and n is the number of qubits. For the W-state case, m=1, the number of CNOT gates scales as O(2n).

A target state that we want to prepare is transformed to the $|0\rangle^{\otimes n}$ by using a series of generalized Hadamard gates and controlled generalized Hadamard gates. The generalized Hadamard has the form $\tilde{H} = C^{\dagger}XC$, where

$$C = \begin{pmatrix} u & -v^* \\ v & u^* \end{pmatrix} \tag{C9}$$

is a unitary matrix. For any single-qubit state $\cos \theta |0\rangle + \sin \theta e^{i\phi} |1\rangle$, it is easy to determine a \tilde{H} that takes it to $|0\rangle$:

$$\tilde{H} = \begin{pmatrix} u^*v + uv^* & (u^*)^2 - (v^*)^2 \\ u^2 - v^2 & -(u^*v + uv^*) \end{pmatrix} \begin{pmatrix} \cos \theta \\ \sin \theta e^{i\phi} \end{pmatrix} = \begin{pmatrix} 1 \\ 0 \end{pmatrix}.$$
(C10)

Consider the set of states in the computational basis that correspond to Hamming-weight-1 bit strings, $\{X_i|0\rangle^{\otimes n}\}_{i=1}^n$; the W state is a uniform superposition of these states:

$$|W_n\rangle = \frac{1}{\sqrt{n}} \sum_{i=1}^n X_i |0\rangle^{\otimes n}, \tag{C11}$$

where n indicates the number of qubits. This wave function can be expressed as the first qubit and the remaining n-1 qubits:

$$|W_n\rangle = c_0|0\rangle|W_{n-1}\rangle + c_1|1\rangle|0\rangle^{\otimes(n-1)} \tag{C12}$$

where $c_1 = 1/\sqrt{n}$ and $c_0 = \sqrt{(n-1)/n}$. Define an operator Q_{n-1} that takes $|W_{n-1}\rangle$ to $|0\rangle^{\otimes (n-1)}$:

$$Q_{n-1}|W_{n-1}\rangle = |0\rangle^{\otimes (n-1)}. (C13)$$

An X operation on the first qubit in Eq. (C12) followed by a controlled- Q_{n-1} operation (the first qubit being the controlling qubit) creates the state $(c_0|0\rangle + c_1|1\rangle)|0\rangle^{(n-1)}$, to which we can perform the generalized Hadamard to evolve to the zero state $|0\rangle^{\otimes n}$. Analogously, we can define Q_{n-1} using controlled- Q_{n-2} and so on until the base case for the recursion, $|W_2\rangle$, which is also the Bell state,

$$|\beta\rangle = a|10\rangle + b|01\rangle. \tag{C14}$$

which can be brought to $|00\rangle$ by an X gate followed by a generalized Hadamard. Reversing the whole circuit above gives the circuit to prepare the W state from the $|0\rangle^{\otimes n}$ state.

Reference [35] outlines a procedure to produce arbitrary states with fixed particle number through a recursive approach that scales as $O(2^m n^m/m!)$ where m is the number of spin-up

or $|1\rangle$ states and n is the number of qubits. When considering creation of single-excitation states, or W states, the circuits have O(n) CNOT single-qubit gates with only nearest-neighbor physical coupling.

As an example for constructing an even superposition of three excitations, a W state, we provide the QUIL [36] and CIRQ code below

(i) QUIL code for the W state

RY(acos(-1/3)) 2 PHASE(-pi/2) 2 RY(pi/4) 1 CNOT 2 1 RY(-pi/4) 1 RZ(pi/2) 1 CNOT 2 1 RZ(pi/2) 1 CNOT 1 0 CNOT 2 1 X 2

(ii) CIRQ code for the W state

qubits = cirq.LineQubit.range(3)
w_state = cirq.Circuit().from_ops([
cirq.Ry(numpy.arccos(-1/3)).
on(qubits[2]),
cirq.ZPowGate(exponent=-1/2).
on(qubits[2]),
cirq.Ry(pi/4).on(qubits[1]),
cirq.CNOT(qubits[2], qubits[1]),
cirq.Ry(-pi/4).on(qubits[1]),
cirq.Rz(pi/2).on(qubits[1]),
cirq.CNOT(qubits[2], qubits[1]),
cirq.CNOT(qubits[2], qubits[1]),
cirq.CNOT(qubits[1], qubits[1]),
cirq.CNOT(qubits[1], qubits[0]),
cirq.CNOT(qubits[2], qubits[1]),
cirq.X.on(qubits[2])])

3. Preparing the generalized W state via projective measurements

This section is about preparing generalized *W* states through projective measurements proposed in Ref. [37].

a. Procedure outline

We start with the *n*-qubit state $|0\rangle^{\otimes n}$ and apply the biased Hadamard gate:

$$H = \begin{pmatrix} \sqrt{1 - \frac{w}{n}} & \sqrt{\frac{w}{n}} \\ \sqrt{\frac{w}{n}} & -\sqrt{1 - \frac{w}{n}} \end{pmatrix}^{\otimes n}.$$
 (C15)

The biased Hadamard gate will drive the initial state to $|\psi\rangle=H|0\rangle^{\otimes n}=(\sqrt{1-p}|0\rangle+\sqrt{p}|1\rangle)^{\otimes n}$ with p=w/n. Consider $|\psi\rangle$ measured in the computational basis; since the probability of each qubit being in $|0\rangle$ is p, the probability of getting a state of Hamming weight w is $\Pr(w)=C_n^wp^w(1-p)^{n-w}$, which as a function of w has a minimum at w=n/2 with $\Pr(w=n/2)=\frac{n!}{2^n(n/2)!(n/2)!}\approx\sqrt{\frac{2}{n\pi}}$, where the \approx refers to the large-n limit. Therefore $\Pr(w)\gtrsim\sqrt{\frac{2}{n\pi}}$ for any w. Specifically,

for our interest of w = 1, we have p = 1/n and

$$\Pr(w=1) = \left(1 - \frac{1}{n}\right)^{n-1} = \left(1 + \frac{1}{n-1}\right)^{-(n-1)} \approx \frac{1}{e},$$
(C16)

which is a fairly high probability.

If we can conduct a projective measurement on the Hamming weight (instead of measuring σ^z on each qubit), then instead of collapsing to a state in the computational basis the system is projected to the subspace of Hamming weight w. Given Eq. (C16), with only a few repetitions one is expected to get w=1, accompanying a generalized W state.

Now we describe the circuit to perform the projective measurement on the Hamming weight illustrated in Ref. [38]. By definition the Hamming weight is the number of 1's of a state in the computational basis. Computing the Hamming weight can be done by introducing an auxiliary register W, and apply a "controlled-add-1" gate on it. The "add-1" is conditioned on the qubit being in state 1. Since the Hamming weight is at most n, $\log n$ qubits are sufficient to encode W. Upon measuring the auxiliary qubits, one gets a generalized W state when the measured W = 1. Now we consider how the controlled-add-one is implemented. First, we review a more general operation controlled-add-k for any constant k presented in Ref. [39]. The idea is to use a n-bit auxiliary register C to record whether a carry will happen for the next bit. Populate C bit by bit. In a second loop, do the real addition.

b. Program Conditional_Add_k

Notation

X: n-bit register

B: bit register (control)

C: n-bit auxiliary register (initialized and finalized to 0)

Pseudo-code

Conditional_Add_k :

for
$$i = 1$$
 up to $n - 1$ (C17)

$$C_i \leftarrow C_i \oplus \text{MAJ}(k_{i-1}, X_{i-1}, C_{i-1}) \tag{C18}$$

endfor

for
$$i = n - 1$$
 down to 1 (C19)

$$X_i \leftarrow X_i \oplus (k_i \wedge B)$$
 (C20)

$$X_i \leftarrow X_i \oplus (C_i \wedge B)$$
 (C21)

$$C_i \leftarrow C_i \oplus \text{MAJ}(k_{i-1}, X_{i-1}, C_{i-1})$$
 (C22)

endfor

$$X_0 \leftarrow X_0 \oplus k_0$$
 (C23)

where the "majority" gate MAJ takes value true when at least two out of the three bits are true:

$$MAJ(O, P, Q) = \begin{cases} P \wedge Q & \text{if } O = 0 \\ P \vee Q & \text{if } O = 1 \end{cases}$$
 (C24)

We now explain the above pseudo-code.

Line (C18), assuming B = 1, i.e., the addition will happen, determines whether a carry will happen for the next bit, and record in C_i . A Toffli (controlled-controlled-NOT) gate is

applied to C_i , with the control condition being that at least two bits in $\{k_{i-1}, X_{i-1}, C_{i-1}\}$ take value 1. Because C is initialized to be all zeros, the control condition being true will set $C_i = 1$.

Line (C20) and line (C21) implement the real addition of k_i and C_i to X_i , controlled by B.

Line (C22) is exactly the same as line (C18), hence C resets to zero.

Line (C23) adds k_0 to X_0 .

Now we consider the special case k=1. We only need a single-bit register for k: $k_0=1$. Line (C18) reduces to $C_1=C_1\oplus (X_0\vee C_0)$ and $C_i=C_i\oplus (X_{i-1}\wedge C_{i-1})$ for i>1. Therefore the pseudo-code is

$Conditional_Add_1:$

for
$$i = 1$$
 up to $n - 1$ (C25)

endfor

for
$$i = n - 1$$
 down to 1 (C27)

$$X_i \leftarrow X_i \oplus (C_i \wedge B)$$
 (C28)

$$SET_C$$
 (C29)

endfor

$$X_0 \leftarrow \bar{X}_0 \tag{C30}$$

with subroutine

$$if i == 1 (C32)$$

$$C_i \leftarrow C_i \oplus (X_{i-1} \vee C_{i-1})$$
 (C33)

else

$$C_i \leftarrow C_i \oplus (X_{i-1} \wedge C_{i-1})$$
 (C34)

c. Number of gates

The circuit for computing the Hamming weight would involve n Controlled_add_1 gates on the Hamming weight register W, each controlled by one data qubit. Since W is composed of $\log n$ bits, each Controlled_add_1 requires $5\log n$ Toffoli (or Toffoli-like) gates. The total circuit requires $5n\log n$ Toffoli gates. With the probability 1/e of getting W=1 in the measurement, it will take on average three repetitions, hence the expected number of Toffoli gates is $\approx 15n\log n$, which translates into $\approx 90n\log n$ CNOT gates.

Note that when measurement is expensive it would be more efficient to hold the measurement on W after performing the computation of W, apply the following target algorithm, and measure W in the end. This will triple the run time of the whole algorithm.

4. Applying the XY Hamiltonian

The XY Hamiltonian on a 1D chain or ring can be exactly implemented in the gate model. Applying the XY Hamiltonian on a state $|0\rangle$ can generate a certain superposition of states in the subspace S expanded by states corresponding to Hamming-weight-1 bit strings, but the exact W state may not

be generated this way. The Hamiltonian for a 1D XY model with nearest-neighbor coupling reads

$$H_{XY} = \sum_{c=1}^{m} \sigma_c^x \sigma_{c+1}^x + \sigma_c^y \sigma_{c+1}^y$$
 (C36)

$$= \frac{1}{2} \sum_{c=1}^{m} (\sigma_c^+ \sigma_{c+1}^- + \sigma_c^- \sigma_{c+1}^+)$$
 (C37)

where periodic boundary condition (PBC) $\sigma_{m+1} = \sigma_1$ is implied. We now examine how well the ring transfers the classical Hamming-weight-1 state.

a. State transfer using a XY chain with open boundary condition

If we remove the constraint of PBC, i.e., work on an open-end chain instead of a ring, it is known that perfect state transfer can be achieved along a XY chain only for n = 2 and 3. The fidelity can be computed by diagonalizing the Hamiltonian; in this case the eigenvector reads

$$|k\rangle = \sqrt{\frac{2}{m+1}} \sum_{n=1}^{m} \sin\left(\frac{nk\pi}{N+1}\right) |n\rangle$$
 (C38)

and the eigenvalues are $E_k = 2\cos(\frac{k\pi}{m+1})$. Furthermore, if inhomogeneous coupling between nearest-neighbor qubits along the chain is allowed, perfect transfer can be realized for any chain length [40].

b. State transfer using a XY chain with PBC

We now study how the state $|0\rangle$ is transferred along a ring. We apply Jordan-Wigner transformation [26,41] to $H_{B,j}$:

$$a_c = S_c^- e^{i\phi_c},\tag{C39}$$

$$a_c^{\dagger} = S_c^+ e^{-i\phi_c} \tag{C40}$$

where $S_c^+ = (\sigma_c^x + i\sigma_c^y)/2$, $S_c^- = (\sigma_c^x - i\sigma_c^y)/2$, and the phase factor $\phi_c = \pi \sum_{c' < j} (\sigma_{c'}^z + 1)/2$ is long ranged involving all operators for c' < c. The new operators a_c and a_c^\dagger can be verified to obey the fermion anticommutation relations, $\{a_c, a_{c'}^+\} = a_c a_{c'}^\dagger + a_{c'}^\dagger a_c = \delta_{c,c'}$ and $\{a_c, a_{c'}\} = \{a_c^\dagger, a_{c'}^\dagger\} = 0$. The inverse transformation reads

$$S_c^+ = a_c^{\dagger} e^{i\phi_c}, \tag{C41}$$

$$S_c^- = a_c e^{-i\phi_c},\tag{C42}$$

$$\sigma_c^z = 2a_c^{\dagger} a_c - 1, \tag{C43}$$

and the phase factor in the fermionic representation is $\phi_c = \pi \sum_{c' < c} a_{c'}^{\dagger} a_{c'}$. The Jordan-Wigner transformation is a convenient tool for one-dimensional spin systems, particularly for nearest-neighbor couplings because in the product of the neighboring spin operators like $S_c^+ S_{c+1}^-$ the phase factors drop out, leaving a concise expression with short-ranged coupling.

Applying the transformation to our problem, for simplicity, we omit the index j in the Pauli operators for this section, and without loss of generality we use $\mathcal{P}_j = \{1, 2, ..., m\}$. We get

$$H_{XY} = 2\sum_{c=1}^{m-1} a_c^{\dagger} a_{c+1} - a_m^{\dagger} a_1 G + \text{H.c.},$$
 (C44)

and the initial state $|1,0,\cdots,0\rangle$ in the fermionic representation becomes

$$|\Phi\rangle_0 = a_1^{\dagger} |\mathbf{0}\rangle \tag{C45}$$

where $|\mathbf{0}\rangle$ denotes the vacuum state (the zero eigenstate of the total particle number operator $\hat{N}_{\text{tot}} = \sum_{c} a_{l}^{\dagger} a_{c}$). Here we introduced gauge operator $G = \exp[i\pi \sum_{l=1}^{m} a_{l}^{\dagger} a_{l}] = \prod_{l=1}^{m} (-)\sigma_{l}^{z}$. The initial state has only one spin up, and hence corresponds to G = -1. Because G commutes with both H_{B} and H_{C} , it is a constant of motion and its value stays -1 throughout the evolution. We can therefore replace G with -1 in Eq. (C44):

$$H_{XY} = 2\sum_{c=1}^{m} a_c^{\dagger} a_{c+1} + \text{H.c.}$$
 (C46)

This Hamiltonian can be diagonalized by introducing the Fourier transformation $f_k = \frac{1}{\sqrt{m}} \sum_{c=1}^m \exp[-ick\omega] a_c$, with $\omega = \frac{2\pi}{N}$. The diagonalized Hamiltonian is

$$H_{XY} = \sum_{k=1}^{m} E_k f_k^{\dagger} f_k \tag{C47}$$

where the eigenenergies $E_k = 2\cos(2k\pi/m)$ and the eigenstates are

$$|\psi_k\rangle = f_k^{\dagger}|0\rangle = \frac{1}{\sqrt{m}} \sum_{c=1}^m e^{ick\omega} a_c^{\dagger}|0\rangle.$$
 (C48)

We now measure the expectation value of the occupation operator $\hat{n}_c = a_c^{\dagger} a_c$ for each site c. The occupation operator \hat{n}_c taking value 1 indicates the the spinless fermion particle is at site c, and correspondingly in the spin problem the spin on site c is up.

At time t the fidelity of state transfer at site c can be analytically derived to be

$$\mathcal{F} \equiv \text{Tr}[\hat{n}(c)e^{-itH_{XY}}|\Phi_0\rangle\langle\Phi_0|e^{itH_{XY}}]$$
 (C49)

$$= \left| \frac{1}{m} \sum_{k} e^{i(c-1)k\omega} e^{-itE_k} \right|^2. \tag{C50}$$

Figure 12 shows the problability evolution of the transfer fidelity at each site. For W-state generation, we are looking for a time when population on all sites has equal probability,

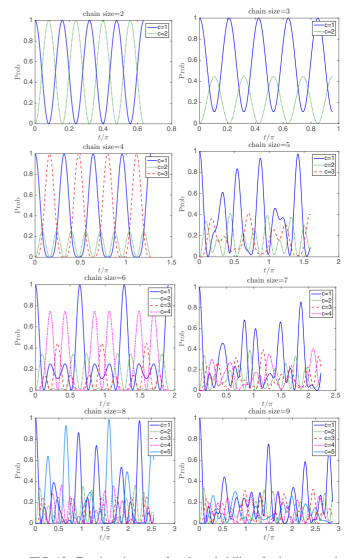


FIG. 12. For ring size m = 2 to 9, probability of spin-up at each site as a function of time t, under the evolution of the XY Hamiltonian on the ring.

i.e., when curves of different colors interact at 1/n. We see for n up to 4 that an exact W state can be generated by choosing the right time t.

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